

HYDRAN-XR

Command Reference

Hydrodynamic Response Analysis
with
Integrated Structural Finite Element Analysis

Version 25.1

NumSoft Technologies

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1. Index of Commands

The following listing of commands is available on-line via the commands `index hydrodynamics`, `index general`, `index database`, `index matrix`, `index functions`, `index fe_commands`, `index_fe_elements`, and `index misc`. All categories are printed by the single command `index`.

1.1. Hydrodynamic Commands

| | |
|--|---|
| <code>hyd_analysis</code> | added mass, hydrodynamic damping, exciting forces |
| <code>hyd_analysis_response</code> | wave-induced hydroelastic response |
| <code>hyd_analysis_response_drag</code> | wave-induced hydroelastic response |
| <code>hyd_analysis_response_P</code> | external force hydroelastic response |
| <code>hyd_assign_mooring</code> | assign mooring stiffness to a body |
| <code>hyd_body_check</code> | check body data |
| <code>hyd_close_files</code> | close HYDRAN-XR output files |
| <code>hyd_convert_fea_mesh</code> | convert FEA mesh to hydrodynamic panel mesh |
| <code>hyd_coordaxs</code> | specify coordinate axes |
| <code>hyd_coord_trans</code> | transform input coordinates to inertial coordinates |
| <code>hyd_export_graphics</code> | export hydro. panel mesh to graphics program |
| <code>hyd_export_graphics_th</code> | export time history motion to graphics program |
| <code>hyd_flex_modes</code> | input flexible modes |
| <code>hyd_genmodes</code> | transform to generalized coordinates |
| <code>hyd_irregular</code> | short-term extreme response |
| <code>hyd_modal_pressure</code> | print exciting and modal pressures |
| <code>hyd_mooring_line</code> | input mooring line data |
| <code>phyd_mooring_line</code> | print mooring line data |
| <code>hyd_mooring_stiffness</code> | input mooring stiffness |
| <code>phyd_mooring_stiffness</code> | print mooring stiffness |
| <code>hyd_nodes</code> | input nodal coordinates |
| <code>phyd_nodes</code> | print nodal coordinates |
| <code>hyd_node_gen</code> | node generation |
| <code>hyd_node_tolerance</code> | check nodes for still water and symmetry planes |
| <code>hyd_panel</code> | define 4-node (or 3-node) panel elements |
| <code>phyd_panel</code> | print panel elements |
| <code>hyd_panel_rmap</code> | create reverse mapping of panel numbers |
| <code>hyd_parameters</code> | input global control parameters |
| <code>hyd_postresponse</code> | obtain some post-processed responses |
| <code>hyd_postresponse_P</code> | obtain some post-processed responses |
| <code>hyd_rigid_modes</code> | generate rigid body modes |
| <code>hyd_rmass</code> | input mass matrix for user modes |
| <code>hyd_surf_elevation</code> | determine the "free" surface elevation |
| <code>hyd_surf_nodes</code> | input surface nodal coordinates |
| <code>phyd_surf_nodes</code> | print surface nodal coordinates |
| <code>hyd_surf_node_gen</code> | surface node generation |
| <code>hyd_surf_node_tolerance</code> | check surface nodes |
| <code>hyd_surf_panel</code> | define 4-node (or 3-node) surface elements |
| <code>phyd_surf_panel</code> | print surface elements |
| <code>hyd_tf</code> | calculate transfer functions |
| <code>hyd_velocity</code> | calculate fluid velocity at user-specified points |
| <code>hyd_velocity_nodes</code> | define velocity point coordinates |
| <code>phyd_velocity_nodes</code> | print coordinates |
| <code>hyd_velocity_node_gen</code> | generate velocity point coordinates |
| <code>hyd_velocity_node_tolerance</code> | check velocity nodes |
| <code>hyd_wave</code> | input wave frequencies and incidence angles |

| | |
|---------------------|------------------------------------|
| hyd_wave_dispersion | solve for wave length, wave number |
| hyd_wave_spectra | input wave spectra |
| phyd_wave_spectra | print wave spectra |
| hyd_wet | estimate "wet" natural frequencies |

1.2. General Commands

| | |
|----------------|---|
| break_loop | break do/while loop |
| date | prints the current date and time |
| do | do loop |
| filein | read commands from file |
| flush | flush the buffer for a file |
| help | on-line help |
| if | conditional if |
| index | index of commands by category |
| logfile | turn log file on/off |
| login | read commands from a log file |
| name? | echo current project name |
| new_project | start a new project (also newproj or newprob) |
| palias | print command aliases |
| quit | quit MANOA |
| read | read an array from file |
| rename_file | rename a file |
| return | return from batch mode |
| rm_file | remove a file |
| savequit | save database and quit |
| system_command | execute a system command |
| time | returns the system seconds |
| while | conditional while |
| write | write an array to file |

1.3. Database Commands

| | |
|--------------|-------------------------------------|
| clear | initialize the database |
| ls or dir | short listing of arrays in database |
| ll | long listing of arrays in database |
| memory | report memory used by database |
| mv or rename | rename an array |
| readdb | read a database from file |
| rm or del | remove an array from the database |
| rm* or del* | remove members from the database |
| save | save the database to file |

1.4. Matrix Commands

| | |
|------------------------|----------------------------------|
| add | matrix add |
| arpack | eigenvalue solver |
| array3d_slice | slice a 3-D array |
| <u>array3d_unslice</u> | insert a matrix into a 3-D array |
| cp | copy |
| cpdg | copy diagonal elements |
| diag_mult | multiply by a diagonal matrix |
| dim_reduce | reduce dimensions of a matrix |
| eigval | eigenvalue solver |
| extract | extract rows of a matrix |

| | |
|---------------------|---|
| fft | compute Fourier or inverse Fourier transform |
| fft_helper | multiply transfer functions with Fourier coefficients |
| ftopro | full to profile storage |
| gauss | Gauss elimination |
| get_dim | get dimensions of a matrix |
| ident | create real identity matrix |
| identc | create complex identity matrix |
| identi | create integer identity matrix |
| input | input a real matrix |
| inputc | input a complex matrix |
| inputch | input a character matrix |
| inputi | input an integer array |
| input3d | input a real 3-D array |
| input3dc | input a complex 3-D array |
| inputi | input an integer 3-D array |
| interpolate | interpolate discrete data |
| invert | invert a square matrix |
| jacobi | jacobi eigenvalue solver |
| joinh | join two matrices horizontally |
| joinv | join two matrices vertically |
| max | find the maximum in a matrix column |
| min | find the minimum in a matrix column |
| mult | matrix multiply |
| mult_col | multiply columns of two matrices |
| mult_elem | multiply elements of two matrices |
| norm | matrix norm |
| pmult | matrix multiply for profile storage |
| print | print a matrix |
| printf | print a matrix in F format |
| printi | print a matrix in integer format |
| psolve | profile equation solver |
| psolve16 | profile equation solver for real*16 |
| ptoful | convert profile to full storage |
| ptosparse | convert profile to sparse storage |
| put | put matrix inside another |
| putdg | put vector on diagonal of a matrix |
| scale | scale an array |
| series | create a 1d series of real data |
| series2d | create a 2d series of real data |
| seti | set a character string |
| seti | set an integer scalar |
| seti | set a real scalar |
| sort | sort a matrix |
| sparse_matrix_clean | remove sparse matrix storage |
| sparse_mult | matrix multiply for sparse storage |
| split | split a matrix based on columns |
| sub | matrix subtraction |
| subcol | subtract columns of two matrices |
| sumcol | sum columns of a matrix |
| tmult | multiply by transpose |
| to_complex | convert to complex |
| to_complex2 | convert to complex with two arguments |
| to_int | convert to integer |
| to_real | convert to real |
| to_reall6 | convert to real*16 |
| to_vector | convert to vector |
| trans | matrix transpose |
| unsplit | unsplit a matrix |

| | |
|--------|-----------------------------|
| unwrap | unwrap the rows of a matrix |
| wrap | wrap the rows of a matrix |
| xprint | extended print |
| zero | zero a real matrix |
| zeroc | zero a complex matrix |
| zeroi | zero an integer matrix |

1.5. Mathematical Functions

| | |
|-----------------------|--|
| abs | absolute value of matrix elements |
| acos | arccosine of matrix elements |
| asin | arcsine of matrix elements |
| atan | arctangent of matrix elements |
| bessel_j or bessel_y | Bessel function of 1st/2nd kind of matrix elements |
| conjugate | complex conjugate of matrix elements |
| cosine | cosine of matrix elements |
| cosh | hyperbolic cosine of matrix elements |
| epsilon | returns a small value relative to 1 |
| erf | error function |
| erfc (see erf) | complementary error function |
| erfc_scaled (see erf) | scaled complementary error function |
| exp | exponential of matrix elements |
| gamma | gamma function of matrix elements |
| log | natural log of matrix elements |
| log10 | common log of matrix elements |
| pi | creates a scalar with the value of pi |
| power | power of matrix elements |
| sine | sine of matrix elements |
| sinh | hyperbolic sine of matrix elements |
| sqrt | square root of matrix elements |
| tan | tangent of matrix elements |
| tanh | hyperbolic tangent of matrix elements |

1.6. Finite Element Commands

| | |
|--------------------|---|
| bcid | displacement boundary conditions |
| body_frc2d | define 2D body forces |
| check_diag | check diagonals of K for zero |
| consolidation | carry out consolidation analysis |
| dampers | nodal dampers |
| direct_th | determines dynamic time history response |
| disp_cntl | displacement control |
| el_iso_matl | elastic, isotropic material matrix |
| elem_alias | print element aliases |
| elem_grp | add/delete element group |
| elemXX | input for element type XX |
| eq_direction | generate earthquake direction vector |
| export_graphics | export to graphics program |
| fem_error | estimate finite element error |
| form_c | form global damping |
| form_k | form global stiffness |
| form_m | form global mass |
| imposed_displ | imposed nodal displacements |
| initial_conditions | specify initial conditions for time history |
| load_summary | print load summary |
| lsolve | linear equation solver |

| | |
|------------------|---|
| mass | nodal mass definition |
| mass_summary | summary of total structural mass |
| merge_nodes | merge coincident nodes |
| modal_th | determines dynamic modal response |
| nodal_constraint | impose nodal constraints |
| nodal_disp | arrange displacements on node basis |
| nodal_pressure | nodal pressure definition |
| nodef | nodal load definition |
| nodes | node definition |
| node_gen | generate nodes |
| node_order | orders nodes for equation numbering |
| node_str | average nodal stresses |
| nsolve | nonlinear equation solver |
| num_eqs | number equations |
| pbcid | print nodal restraints |
| pbody_frc2d | print 2D body forces |
| pcurrentvelocity | print current velocity |
| pdampers | print nodal dampers |
| pdisp | print displacements |
| pelemXX | print data for element type XX |
| peqns | print equation numbers |
| pimposed_displ | print imposed nodal displacements |
| pmass | print input nodal masses |
| pndisp | print nodal displacements |
| pndisp_th | print time history of nodal displacements |
| pnodef | print input nodal loads |
| pnodes | print nodes |
| presponse | print element response |
| pstate | print element state |
| response | determine element response |
| rigid_modes | generate rigid body modes |
| state | determine element state |
| water_waves | input water waves |

1.7. Finite Elements

| | |
|----------------|--|
| beam3d | linear, 3-D beam element |
| biot1d234 | 1-D element for linear, elastic consolidation |
| biot2d3to9 | 2-D element for linear, elastic consolidation |
| cable | elastic catenary cable |
| contact_spring | nonlinear, contact spring |
| d11234 | 1-D, linear, elastic element |
| d11234v2 | 1-D, linear, elastic element, v.2 |
| d213to9 | 2-D, linear elastic element |
| d21tri | 2-D, linear elastic triangular element |
| iFEM2D | 2-D, 3 to 9 node element for nonlinear iFEM |
| interface | quadrilateral (and triangular) interface element |
| isomin6 | linear, triangular Mindlin plate |
| min3s | Mindlin 3-D triangular, linear shell |
| min5s | Mindlin 3-D quadrilateral, linear shell |
| min4t | quadrilateral, linear, Mindlin shell |
| min6 | linear, triangular Mindlin plate |
| nbeam2d | large displacement, elastic 2D beam |
| ntruss | large displacement truss |
| spring | nonlinear, elastic spring |
| stiff2n | linear elastic 2-node stiffness element |
| truss | linear truss |

1.8. Miscellaneous Commands

| | |
|---------------------------|---|
| <code>fortran_kind</code> | prints the number of bytes for standard types |
| <code>gauss_int</code> | Gauss integration |
| <code>gauss_pts</code> | Gauss integration points |
| <code>poly</code> | evaluate a 1-D or 2-D polynomial |
| <code>tri_intpts</code> | integration points for triangle |
| <code>userf</code> | user-defined functions |

2. Command Reference

2.1. Hydrodynamic Commands

hyd_analysis

Command to determine the added mass, damping, and wave excitation forces

Command Syntax

```
hyd_analysis [#added_freqs=? -periods] [-no_pot] [-source]
```

#added_freqs is the number of wave frequencies to add to a previous analysis. The frequencies, in radians/sec, are to be listed on the input line immediately following this command. If -period is specified, the input values are interpreted to be wave periods. The frequencies will be inserted in numerical sequence into the frequencies previously specified by the hyd_wave command.

If -no_pot is specified, the velocity potentials are not saved.
WARNING: Although this option saves memory in the database, specifying -no_pot means that the potentials and/or pressures cannot be calculated in the hyd_analysis_response command.

If -source is specified, the radiation source strengths are written to file *.rad by the Fortran statement

```
write(f_rad)omega, strength
```

and the diffraction source strengths are written to file *.dif by the Fortran statement

```
write(f_dif)omega, angle, strength
```

For the latter, the outer loop is on the wave frequency. The * in the above file names represents the project name. Contact NumSoft Technologies for more complete details regarding the information written to these files. Note that this option is not compatible at this time with the #added_freqs option.

Some basic commands to carry out the hydrodynamic analysis can include:

```
hyd_parameters  
hyd_coordaxs  
hyd_wave  
hyd_nodes  
hyd_coord_trans  
hyd_panel  
hyd_body_check  
hyd_rigid_modes  
hyd_rmass  
hyd_flex_modes  
hyd_genmodes  
hyd_analysis  
hyd_analysis_response  
hyd_postreponse
```

Other commands can be used, as needed. In addition, the sequence of the commands as given above may be altered somewhat, but this command must

precede the `hyd_analysis_response` command. The `hyd_nodes` command (and the `hyd_coord_trans` command, if it is required) must precede the `hyd_panel` command. Also, `hyd_rigid_modes`, if used, must be given before `hyd_rmass` and `hyd_flex_modes`. The `hyd_flex_modes` command is not needed to carry out the hydrodynamic analysis of single or multiple rigid bodies.

The hydrodynamic analysis is based on linear potential theory. The Green function method with constant source strengths over each fluid panel is used to solve for the hydrodynamic variables. All calculations are carried out in double precision, except the Green functions are evaluated to single precision accuracy and the solutions of the equations to obtain the radiation and diffraction potentials are carried out in single precision arithmetic. The program is applicable for both infinite and finite water depths. Results from this command are the added mass, hydrodynamic damping, and generalized wave exciting forces.

Single and double symmetry of the structure are exploited by using the composite source distribution method. For single symmetry, the panels should be generated corresponding to the $y > 0$ region. For double symmetry, the panels should be generated on the $x > 0, y > 0$ region of the body.

The results are written to files, whose names are the project name with an extension. In addition to the basic *.out file, the following files may be created:

```
*.pan --> panel data
*.adm --> added mass coefficients
*.adp --> hydrodynamic damping coefficients
*.exf --> generalized wave excitation forces
*.ot2 --> additional output
*.pot --> velocity potentials at panel centers
```

The following main arrays are created in the database:

```
.hyd_modesym(nmode)      -> symmetry code for modes
.hyd_twnk(nfreq,2)      -> wave period, wave number
.hyd_un(nmode,npanel)   -> generalized normals
.hyd_xr(npanel,4)       -> x coordinates of panels
.hyd_yr(npanel,4)       -> y coordinates of panels
.hyd_zr(npanel,4)       -> z coordinates of panels
.hyd_xyzc(npanel,3)     -> x,y,z coordinates of panel centers
.hyd_panela(npanel)     -> area of fluid panels
.hyd_paneln(3,npanel)   -> x,y,z components of panel normals
.hyd_addm(nmode,nmode,nfreq) -> added mass coefficients
.hyd_damp(nmode,nmode,nfreq) -> hydrodynamic damping coefficients
.hyd_uz(nmode,npanel)   -> vertical, z, displacements
.hyd_fd(nmode,nbeta,nfreq) -> generalized diffraction forces
.hyd_fi(nmode,nbeta,nfreq) -> generalized Froude-Krylov forces
.hyd_rhs(nmode)         -> RHS of generalized equations
.hyd_dynk(nmode,nmode)  -> dynamic stiffness (LHS)
.hyd_gencor(nmode,nfreq,nbeta) -> generalized coordinates
.hyd_potincm(npanel,nfreq,nbeta) -> incoming potentials
.hyd_potdiff(npanel,nfreq,nbeta) -> diffraction potentials
.hyd_potrad(npanel,nfreq,nmode) -> radiation potentials
```

where `nmode` is the number of modes, `npanel` is the number of active panels, `nfreq` is the number of wave frequencies, and `nbeta` is the number of wave angles. These quantities are specified by the `hyd_parameters` and `hyd_panel`

commands. Additional, temporary, arrays are also created and deleted. They all begin with either ".hyd_" or "\$". To avoid a collision in array names, the user should not create any other array with these prefixes.

See Also

hyd_analysis_response hyd_panel hyd_parameters

hyd_analysis_response

Command to determine the wave-induced response

Command Syntax

```
hyd_analysis_response [-residual] [symmetrize=on|off]
```

This command solves the equations of motion to determine the generalized response based on the added mass, damping, and exciting forces calculated by the hyd_analysis command, which must precede this command.

If -residual is specified, the residual error in the solution of the equations of motion is calculated. The maximum norm of the error is reported.

If symmetrize=on, the added mass and hydrodynamic damping matrices are symmetrized by averaging the corresponding off-diagonal terms. If symmetrize=off, no averaging is done (default). Theoretically, these matrices should be symmetric. However, because of discretization errors, some corresponding terms may not be symmetric. The unsymmetry reduces as the mesh is refined.

This command can be executed any number of times, for example, to evaluate changing the moments of inertia of a body. The hyd_analysis command does not need to be repeated as long as the added mass, hydrodynamic damping, and exciting forces are not affected. Before this command, the structural mass matrix must be defined as a database member and named hyd_mstr. For single or multiple rigid bodies, this matrix is created easily with the hyd_rmass command. If a viscous structural damping matrix (hyd_cstr) and a structural stiffness matrix (hyd_kstr) are not defined, they will be created and zeroed. (Constant structural hysteretic damping can be easily specified via the hyd_parameters command.) The dimensions of the structural matrices should be (nmode,nmode), where nmode is the number of modes. The hyd_flex_modes command is not needed to carry out the hydrodynamic analysis of single or multiple rigid bodies.

The results are written to files, whose names are the project name with an extension. In addition to the basic *.out file, the following file will be created:

```
*.cor --> generalized coordinates
```

The following main arrays are created in the database:

```
.hyd_rhs(nmode)           -> RHS of generalized equations  
.hyd_dynk(nmode,nmode)    -> dynamic stiffness (LHS)  
.hyd_gencor(nmode,nfreq,nbeta) -> generalized coordinates
```

where nmode is the number of modes, nfreq is the number of wave frequencies, and nbeta is the number of wave angles. Additional, temporary, arrays are also created and deleted. They all begin with either ".hyd_" or "\$". To avoid a collision in array names, the user should not create any other array with these prefixes.

See Also

```
hyd_analysis hyd_postresponse
```

hyd_analysis_response_drag

Command to determine the wave-induced response including linearized quadratic drag

Command Syntax

```
hyd_analysis_response_drag arg1 arg2 beta=? conv=? [maxit=?]  
                          [wave=?] [symmetrize=on|off]
```

This command solves the equations of motion to determine the generalized response based on the added mass, damping, and exciting forces calculated by the hyd_analysis command, which must precede this command. The command linearizes the Morison-type drag term as described below. Only one wave angle is considered.

arg1 is the vector of diagonal terms in the "viscous" damping matrix that is used for the v^2 drag term. Note that this matrix is required to be diagonal, and hence only the diagonal values must be specified. The vector can be either real or complex.

arg2 is the modal transformation matrix that is used to transform structural mass, damping and stiffness matrices from physical displacements to modal coordinates.

beta is the integer wave angle number (not the wave angle) to be used (default=1)

conv is the convergence criterion

maxit is the maximum number of iterations (default=10)

wave is the wave amplitude (default = 1)

If symmetrize=on, the added mass and hydrodynamic damping matrices are symmetrized by averaging the corresponding off-diagonal terms. If symmetrize=off, no averaging is done (default). Theoretically, these matrices should be symmetric. However, because of discretization errors, some corresponding terms may not be symmetric. The unsymmetry reduces as the mesh is refined.

The equations of motion are of the form

$$[-w^2 M + i w (C + C_d) + K]q = P + P_m$$

M and K are the total mass and stiffness matrices (structure + fluid), C is the structure plus radiation damping, C_d is the linearized drag term, P is the linearized exciting forces, and P_m are the exciting forces from the Morison-type drag term. q are the modal coordinates. This equation is identical to the equations solved by hyd_analysis_response, except for the terms C_d and P_m.

C_d is obtained as follows. The diagonal damping matrix represented by arg1 is multiplied with $(v-u)$ to obtain effective damping terms. v is the velocity at each structural degree-of-freedom based on the incoming wave, and u is the corresponding structural velocity from the previous iteration. The modified diagonal damping matrix is then transformed to modal coordinates using arg2, to obtain C_d.

The modified damping matrix is multiplied by v to obtain the force in

physical coordinates. The transformation matrix `arg2` is used to transform this physical force vector to modal coordinates, giving `P_m`.

The iteration continues until the maximum difference between displacements `u` from one iteration to the next is less than `conv`. If convergence is not reached in `maxit` iterations, a warning is issued but the unconverged results will be treated as converged and the solution will proceed.

This command can be executed any number of times, for example, to evaluate multiple wave angles.

The results are written to files, whose names are the project name with an extension. In addition to the basic `*.out` file, the following file will be created:

```
*.cor --> generalized coordinates
```

If they do not exist, the following main arrays are created in the database:

```
.hyd_rhs(nmode)           -> RHS of generalized equations  
.hyd_dynk(nmode,nmode)    -> dynamic stiffness (LHS)  
.hyd_gencor(nmode,nfreq,nbeta) -> generalized coordinates
```

where `nmode` is the number of modes, `nfreq` is the number of wave frequencies, and `nbeta` is the number of wave angles. Additional, temporary, arrays are also created and deleted. They all begin with either `".hyd_"` or `"$"`. To avoid a collision in array names, the user should not create any other array with these prefixes.

The results from this command will be inserted in `.hyd_gencor(nmode,nfreq,beta)`. Hence, all wave angles can be dealt with one at a time.

See Also

```
hyd_analysis  hyd_postresponse
```

hyd_analysis_response_P

Command to determine response to harmonic loading

Command Syntax

```
hyd_analysis_response_P [-eq] [-residual] [symmetrize=on|off]
```

This command is very similar to `hyd_analysis_response`, which is used for wave-induced motion, whereas this command determines the response to predefined load pattern "P", contained in `hyd_Peiwt(nmode)`. This command uses the added mass and hydrodynamic damping determined by the `hyd_analysis` command, which must precede this command. Only the radiation potentials (i.e., added masses and hydrodynamic damping) are considered. The excitation frequencies are specified by the `hyd_wave` command. Specify one wave angle (0 degrees, for example). The wave exciting forces are ignored. Although it is a bit inefficient to calculate wave exciting forces and then ignore them, the computational effort to obtain the wave exciting forces for one wave angle is relatively small.

For earthquake ground motion, specify option `-eq`. In a usual earthquake analysis, the effective load vector `Peiwt` would be defined as the negative of the transpose of the generalized modes times the combined mass (structure mass M_s + added mass M_f) times an influence vector r (i.e., $-\psi^T * (M_s + M_f) * r$). r specifies which displacement degrees of freedom move with the ground motion. However, the approach used here is to form the generalized added mass ($M_f^* = \psi^T * M_f * \psi$) directly, and hence M_f is not available. It can be shown that the effective load vector can be approximated as

$$-(I + M_f^*) * \psi^T * M_s * r$$

if the generalized structural mass is the identity matrix; i.e., the modes are orthonormal with respect to the structure mass matrix. Therefore, when `-eq` is specified, `Peiwt` should be specified as $\psi^T * M_s * r$, which will be multiplied internally by $-(I + M_f^*)$ for each frequency.

For earthquake ground motion, there is an additional term to the effective load vector because the hydrodynamic damping depends on the total velocity (as compared to viscous structure damping, which is assumed to depend on the relative velocity). Again assuming that the generalized structure mass matrix is the identity matrix, the additional term for the effective load vector is

$$+ i/w C_f^* * \psi^T * M_s * r$$

in which C_f^* is the generalized hydrodynamic damping matrix.

The above expressions for the effective load vector are based on the assumption that the influence vector r can be represented by the generalized modes. There is typically an error in this representation, which results from using a truncated subspace of generalized modes. The error is

$$e = r - \psi * p_r = r - \psi * (M_s^*)^{-1} \psi^T * M_s * r$$

in which $\psi * p_r$ is the approximation of r by the generalized modes. The more modes one uses, the better this approximation should be.

If `-residual` is specified, the residual error in the solution of the equations of motion is calculated. The maximum norm of the error is reported.

If `symmetrize=on`, the added mass and hydrodynamic damping matrices are symmetrized by averaging the corresponding off-diagonal terms. If `symmetrize=off`, no averaging is done (default). Theoretically, these matrices should be symmetric. However, because of discretization errors, some corresponding terms may not be symmetric. The unsymmetry reduces as the mesh is refined.

Some basic commands to carry out the analysis can include:

```
hyd_parameters
hyd_coordaxs
hyd_wave
hyd_nodes
hyd_coord_trans
hyd_panel
hyd_body_check
hyd_rigid_modes
hyd_rmass
hyd_flex_modes
hyd_genmodes
hyd_analysis
hyd_analysis_response_P
hyd_postresponse_P
```

Note that this list does not include the creation of the modal forces in `hyd_Peiwt(nmode)`.

Other commands can be used, as needed. In addition, the sequence of the commands as given above may be altered somewhat. However, the `hyd_analysis` command must precede this command. Before this command, the structural mass matrix must be defined as a database member and named `hyd_mstr`. For single or multiple rigid bodies, this matrix is created easily with the `hyd_rmass` command. If a viscous structural damping matrix (`hyd_cstr`) and a structural stiffness matrix (`hyd_kstr`) are not defined, they will be created and zeroed. (Constant structural hysteretic damping can be easily specified via the `hyd_parameters` command.) The dimensions of the structural matrices should be `(nmode,nmode)`, where `nmode` is the number of modes. The `hyd_flex_modes` command is not needed to carry out the hydrodynamic analysis of single or multiple rigid bodies.

The results are written to files, whose names are the project name with an extension. In addition to the basic `*.out` file, the following file may be created:

```
*.cor --> generalized coordinates
```

See the commands `hyd_analysis` and `hyd_analysis_response` for more details. This command creates the following array in the database:

```
.hyd_gencor(nmode,nfreq) -> generalized coordinates
```

where `nmode` is the number of modes and `nfreq` is the number of frequencies. Additional, temporary, arrays are also created and deleted. They all begin

with either ".hyd_" or "\$". To avoid a collision in array names, the user should not create any other array with these prefixes.

See Also

hyd_analysis hyd_analysis_response hyd_postresponse_P

hyd_assign_mooring

Assign mooring stiffnesses to bodies

Command Syntax

```
hyd_assign_mooring  
body=? mooring=? attach=?,?,?,? [theta=?] [T=?,?,?,?,?,,?,?,,?,?]
```

body is the body number

mooring is the stiffness number in .hyd_mooring_K

attach is the x,y,z body coordinates of the attachment point on the body

theta is the rotation in degrees about the z-axis (see below)

T is a 3x3 transformation matrix in the order

T(1,1),T(2,1),T(3,1),T(1,2),etc.

Assigns ("attaches") a mooring stiffness to a body. The stiffness is assembled into the structural stiffness matrix `hyd_kstr`. If `hyd_kstr` does not exist or is not the proper size (nmode x nmode), it is created.

Prior to adding the mooring stiffness to the structural stiffness, it is transformed to body coordinates. If the z-axis of the mooring stiffness coordinate system is parallel to the z-axis of the body coordinates, then the transformation is conveniently specified by `theta`, which is the z-rotation from the mooring x-axis to the body x-axis. Otherwise, the general 3x3 orthogonal coordinate transformation matrix, `T`, that transforms a vector from the body-fixed coordinate system to the mooring coordinate system may be input in column order.

The mooring stiffness is used to determine the contribution of the mooring stiffness to the stiffness of the rigid body modes, as defined by the command `hyd_rigid_modes`. A mooring stiffness assigned, for example, to body 2 will have stiffness contributions to rigid body modes 7 - 12, the surge, sway, heave, roll, pitch, and yaw of body 2. The mooring stiffnesses must be assigned prior to any transformation of the modes to assumed modes, for example by the command `hyd_genmodes`. Also, any contribution to flexible modes is not included by this command. The user must define that contribution "manually" when specifying `hyd_kstr`.

Every mooring line must be assigned explicitly. In particular, even for a symmetric mooring arrangement, each mooring line must be specified.

See Also

`hyd_mooring_stiffness` `phyd_mooring_stiffness`

hyd_body_check

Perform some checks on the panel mesh

Command Syntax

```
hyd_body_check [body=?] [nodes=?,?] [panels=?,?] [body_sym=?]
```

This command performs some checking of the body panel mesh. The arguments are only required in the case of multiple bodies, as defined by the parameter `nbodies` in the `hyd_parameters` command. In this case, the checking will be for the body number specified by the parameter `body`. Hence, one `hyd_body_check` command should be issued for each body for which a mesh is specified explicitly. The parameters `nodes` and `panels` specify the ranges of nodes and panels corresponding to body. For example, `body=1 nodes=1,100 panels=1,81` mean that body 1 is represented by nodes 1 to 100 and panels 1 to 81. If symmetry in the command `hyd_parameters` is 1 or 2, then the parameter `body_sym` must be specified here. Its value must be: 0, if the mesh for the particular body is for the entire body; 1, if the mesh for the particular body is for 1/2 the body; and 2 if the mesh for the particular body is for 1/4 the body. The default for `body_sym` is the system symmetry specified by `hyd_parameters`.

This command will check that the nodal coordinates are consistent with the symmetry specification. E.g., in the case of double symmetry, only the body in the (+x,+y) quadrant should be meshed. A warning will be issued for all nodes outside this quadrant.

The volume of the mesh is reported, as calculated by the x, y, and z projections.

The inertial coordinates of the center of buoyancy are calculated, based on the average of the three volume calculations.

hyd_close_files
Close output files

hyd_convert_fea_mesh

Convert FEA mesh to a hydrodynamic panel mesh

Command Syntax

```
hyd_convert_fea_mesh
```

Import the fea mesh defined in HYDRAN-XR to a hydrodynamic panel mesh. All FEA nodes are converted to "hydrodynamic" nodes.

A hydrodynamic panel is created for each "wet" interface, min5s and min3s element. If the "positive" side of the finite element is wet, then the node order is reversed when defining the panel to make it consistent with the clockwise specification of nodes for the panels.

The interface elements are converted first, followed by the min5s elements and then the min3s elements.

This command means that a mesh does not have to be essentially redefined by the hyd_nodes and hyd_panel commands as long as a compatible finite element mesh has been defined.

NOTE: the fea nodal coordinates must be specified in the inertial coordinate system to use this command.

See Also

```
hyd_nodes hyd_panel interface min3s min5s
```

hyd_coordaxs

Define the coordinate systems for the hydrodynamic analysis.

Command Syntax

```
hyd_coordaxs [origin=?,?,?] [angles=?,?,?] [xb=?] [yb=?] [zb=?] &  
             [theta=?] [zcg=?] [body=?]
```

origin = (x,y,z) inertial coordinates of the origin of the input coordinate system
angles = Euler (Bryant) angles that define the orientation of the input axes (degrees)
xb = inertial x-coordinate of the origin of the body fixed coordinate system
yb = inertial y-coordinate of the origin of the body fixed coordinate system
zb = inertial z-coordinate of the origin of the body fixed coordinate system
theta = angle in degrees between the inertial x-axis and the body-fixed x-axis (see below)
zcg = body-fixed z-coordinate of the center of gravity
body = the number of the body (see below)

Although the default for all values is 0, this command is required.

There is one inertial (global) coordinate system, and for each body there is an input coordinate system and a body-fixed coordinate system. The number of bodies is specified by the nbodies parameter in the hyd_parameters command. There is always at least one body. For a given body, the input coordinate system is used to specify the nodal coordinates for that body. The body-fixed coordinate system is used principally to define the rigid body modes for the body.

The inertial coordinate system is located on the still-water plane, with the z-axis positive upward. The hydrodynamic calculations are carried out in the inertial coordinate system.

The input coordinate system for a body is for input convenience. If the input and inertial coordinates systems are not the same, then after the nodes have been defined the user must transform the coordinates to the inertial system by the hyd_coord_trans command. The input coordinate system is defined relative to the inertial system by a shift (specified by the origin parameter) and a rotation (specified by the angles parameter). The orientation of the input coordinate system is obtained by sequentially imposing the rotations about first the input x-axis, then the rotated y-axis after 1 rotation, and finally the z-axis after 2 rotations.

The origin of the body-fixed coordinate system is specified by the parameters xb, yb, and zb. The origin of the body axes need not be at the center of gravity of the body. However, it must be on the same vertical line as the CG of the body. The z-axis of the body fixed coordinate system is positive upward. The angle theta is the angle between the inertial x-axis and the body x-axis, measured from the inertial x-axis with counterclockwise positive. The inertial z-axis and the body z-axis are parallel.

In the case of multiple bodies, as defined by nbodies in the command hyd_parameters, one hyd_coordaxs command must be input for each body, and the body number ranges from 1 to nbodies. In this case, the data specified

here define the location and orientation of each body's input coordinates and body-fixed coordinates.

In the case of symmetry, it is possible that a body is a complete reflection of another body that has been input. There are no nodes or panels that are input explicitly for such a body. This command is still required, however, to specify zcg. Although the values are not used for calculations, it is recommended that the origin of the body-fixed coordinate system be given as well so that the correct values will be printed in the project summary.

The data are stored as:

```
.hyd_coordaxs(11,nbodies) -> origin, angles, xb, yb, zb, theta, zcg
```

See Also

```
hyd_coord_trans hyd_nodes hyd_rmass
```

hyd_coord_trans

Transform the nodal coordinates to the inertial coordinate system.

Command Syntax

```
hyd_coord_trans [body=?] [nodes=?,?]
```

This command is required if the input coordinate system and the inertial coordinate system are not the same, as defined by the command `hyd_coordaxs`. For multiple bodies, as specified by the `nbodies` parameter in the `hyd_parameters` command, the parameters `body` and `nodes` are required. `body` is the body number and `nodes` define the range of nodes that correspond to that body. E.g., `body=1 nodes=1,100` would mean that the coordinates for nodes 1 to 100 would be transformed using the coordinate transformation data for body 1 that was specified in the `hyd_coordaxs` command. For multiple bodies, the command must be issued as many times as necessary to transform the nodal coordinates to the inertial coordinate system.

If this command is required, it must be issued after all the nodal coordinates have been defined (`hyd_nodes`) and before the panels are defined (`hyd_panel`).

See Also

```
hyd_coordaxs hyd_nodes hyd_panel
```

hyd_export_graphics

Export hydrodynamic panel mesh to graphics program input file

Command Syntax

```
hyd_export_graphics -target [-modes T=?] &  
                    [-displ freq=? angle=? steps=?] &  
                    [-freesurface] [-wetonly] [file=filename]
```

Export the hydrodynamic panel mesh to a graphic program's input text file.

The graphics program is specified by the argument `-target`. Only the program Gmsh (<http://www.geuz.org/gmsh/>) is supported at this time. That is, the available option is `-Gmsh`.

The default is to plot the undeformed mesh.

If `-modes` is specified, the modes in `.hyd_umx`, `.hyd_umy`, and `.hyd_umz` (see `hyd_rigid_modes` and `hyd_flex_modes` command) are exported as displacements. `T=` is the name of a real vector of frequencies or periods, which is used for identification in the output file. The size of the vector is the number of modes. If a vector is not specified, one is created with the values 1,2, ...

If `-displ` is specified, then a time sequence of displacements will be determined for a given wave frequency and wave angle. `freq` is the integer number of the wave frequency/period, `angle` is the integer number of the wave angle, and `steps` is the number of steps in the wave period for which the real displacements will be determined.

If `-freesurface` is specified, then the free surface panels are exported as well. If `modes` is selected, then the free surface is flat. If `deformed` is selected, then the free surface elevation is also plotted. The displacement of the free surface is determined by the `hyd_surf_elevation` command.

If `-wetonly` is specified, only "wet" panels are exported (panels with a W or V code); see command `hyd_panel` for information.

If `filename` is specified, the results will be written to the file `filename`; otherwise they will be written to the file `project_name.msh` (Gmsh).

hyd_export_graphics_th

Export time history of motion response to graphics program input file

Command Syntax

```
hyd_export_graphics_th [-target] time xdisp ydisp zdisp      &
                        [surfzdisp] [t1=?] [t2=?] [step=?]    &
                        [file=filename]
```

The graphics program is specified by the argument `-target`. Only Gmsh (<http://www.geuz.org/gmsh/>) is supported at this time. That is, the available option is `-Gmsh`, and it is the default.

The time history response must have been previously calculated, e.g., via a Fourier transform approach (see the command `fft`).

`time` is a vector of N time steps

`xdisp` is the $(N, \#nodes)$ matrix of x -displ. for the structure panels

`ydisp` is the $(N, \#nodes)$ matrix of y -displ. for the structure panels

`zdisp` is the $(N, \#nodes)$ matrix of z -displ. for the structure panels

`surfzdisp` is the $(N, \#surfnodes)$ matrix of z -displ. for the surface panels

If `surfzdisp` is not given, then the surface displacements are not plotted.

If `t1` is given, export begins at time(`t1`) (default = 1).

If `t2` is given, export will stop at time(`t2`) (default = N).

If `step` is given, every `step` time steps will be exported (default = 1).

If `filename` is specified, the results will be written to the file `filename`; otherwise they will be written to the file `project_name.msh` (Gmsh).

hyd_flex_modes

Input the flexible structural modes for the hydroelastic analysis. There are two Command Syntax options.

----- OPTION 1 -----

Command Syntax

```
hyd_flex_modes arg
```

----- OPTION 2 -----

```
hyd_flex_modes [-noread]
```

```
mode_j  x_z_sym  y_z_sym
```

```
i  ux(i,j)  uy(i,j)  uz(i,j)  thx(i,j)  thy(i,j)  thz(i,j)
```

Option 1

arg is the array in the database containing the structural mode shapes from the eigenvalue analysis; e.g., .phi (see command eigval). These correspond to the structural degrees-of-freedom, and the dimension is $n_{eq} \times \#modes$. For this option to work, the node numbers of the structural mesh and the node numbers of the panel mesh must be compatible. This can be ensured via the command hyd_convert_fea_mesh. Each column of arg is a mode shape. The first nmoder columns are skipped, and the next nmoder columns are processed. Hence, #modes must be greater than or equal to nmoder + nmoder. If nmoder > 0, then it is assumed that the arrays .hyd_umx, .hyd_uy, etc. (see hyd_rigid_modes for all the arrays expected) have been created and the first nmoder columns have been defined already (e.g., via the hyd_rigid_modes command or "manually"). This command will define the values for the nmoder modes. If symmetry is not 0, the values for the nmoder modes must be defined manually.

Option 2

mode_j = mode number

x_z_sym = port-starboard symmetry code for mode_j

0 -> port-starboard symmetric

1 -> port-starboard anti-symmetric

y_z_sym = fore-aft symmetry code for mode_j

0 -> fore-aft symmetric

1 -> fore-aft anti-symmetric

ux,uy,uz(i,j) = translation of node i in the j-th mode

thx,thy,thz(i,j)= rotation of node i in the j-th mode

Ranges of the indices are: i = 1, nnode; j = nmoder+1, nmode.

End input for each mode with a blank line. That is, the modes are separated by a blank line.

The use of -noread is expected to be uncommon. If the flag -noread is present, the data are not read and only the one command line should be specified. In this case it is assumed that the seven arrays

.hyd_modesym, .hyd_umx, ..., .hyd_thz (see below) have been defined elsewhere and the data are in them for further processing. .hyd_kf, .hyd_un, and .hyd_uz will be created.

The data are stored as:

.hyd_umx(nnode,nmode) -> translational x displacements

.hyd_uy(nnode,nmode) -> translational y displacements

.hyd_umz(nnode,nmode) -> translational z displacements

```
.hyd_thx(nnode,nmode) -> rotational x displacements  
.hyd_thy(nnode,nmode) -> rotational y displacements  
.hyd_thz(nnode,nmode) -> rotational z displacements
```

The symmetry code for mode *j* is stored in `.hyd_modesym(j)`. These codes are only used if the structural symmetry parameter on the `hyd_parameters` command is 1 or 2, indicating single (x-z) or double (x-z and y-z) structural symmetry, respectively. For single symmetry, the modal symmetry code is:

```
1 -> symmetric  
2 -> antisymmetric
```

For double symmetry, the modal symmetry code is:

```
1 -> symmetric/antisymmetric  
2 -> antisymmetric/symmetric  
3 -> symmetric/symmetric  
4 -> antisymmetric/antisymmetric
```

where, e.g., symmetric/antisymmetric means symmetric with respect to the x-z plane and antisymmetric with respect to the y-z plane.

Note: The modal displacements are given in the inertial coordinate system.

This command defines `.hyd_un` and `.hyd_uz` for the flexible modes and it forms an estimate - for the flexible modes - of the hydrostatic stiffness `.hyd_kf` based on the fluid term only. If a better hydrostatic stiffness matrix is available, it should replace the one created by this command (after the command finishes).

See Also

```
hyd_coordaxs  hyd_coord_trans  hyd_nodes  hyd_rigid_modes
```

hyd_genmodes

Transform matrices to generalized coordinates.

Command Syntax

```
hyd_genmodes [1=?/? 2=?/?]
```

This command transforms the input modes to assumed modes, as explained below. There must be the same number of assumed modes as input modes (nmoder + nmodef from hyd_parameters command).

This command is meant primarily for a system of multiple rigid bodies, in which the initial modes are specified to be the traditional surge, sway, heave, etc. of each body. This command allows the transformation to symmetric and antisymmetric modes. The array hyd_psi(nmode,nmode) must be defined prior to this command (e.g., by the input command). Array hyd_psi is defined such that

$$\{d\} = [\text{psi}] \{u\}$$

in which [psi] is hyd_psi, {d} is the displacement vector, and {u} is the vector of generalized displacements. For the cases of single and double symmetry of the system (as defined by the symmetry parameter on the hyd_parameters command), the symmetry/antisymmetry of the modes are specified by the parameters i=?/?, where i is the number of the mode (1 to nmode) and ?/? is of the form S or A for single structural symmetry and S/S, S/A, A/S, and A/A for double structural symmetry. S and A refer to symmetric and antisymmetric, respectively. This input is used to define the vector .hyd_modesym. See the help on commands hyd_rigid_modes and hyd_flex_modes for more discussion on .hyd_modesym.

The transformation is carried out by replacing the system matrices as indicated below:

```
hyd_mstr <- [psi]^T * [hyd_mstr] * [psi]
hyd_kstr <- [psi]^T * [hyd_kstr] * [psi]
hyd_cstr <- [psi]^T * [hyd_cstr] * [psi]
.hyd_kf <- [psi]^T * [.hyd_kf] * [psi]

.hyd_umx <- [.hyd_umx] * [psi]
.hyd_omy <- [.hyd_omy] * [psi]
.hyd_umz <- [.hyd_umz] * [psi]
.hyd_thx <- [.hyd_thx] * [psi]
.hyd_thy <- [.hyd_thy] * [psi]
.hyd_thz <- [.hyd_thz] * [psi]
.hyd_un <- [psi]^T * [.hyd_un]
.hyd_uz <- [psi]^T * [.hyd_uz]
```

Note that the response given in the file *.cor will be the generalized coordinates. Usually, one will also want to define a "modal" matrix such that the response corresponding directly to the original modes will be determined by the command hyd_tf. If no other response components are desired, the modal matrix will be the same as hyd_psi. In this case, it is conveniently defined by the cp or mv commands.

See Also

```
hyd_flex_modes hyd_parameters hyd_rigid_modes hyd_tf
```

hyd_irregular

Calculate short-term extreme response in irregular seas

Command Syntax

```
hyd_irregular arg1 arg2 [ext=extension] [file=filename]
```

This command determines the short-term extreme response based on the transfer functions that have been determined with the `hyd_tf` command and the wave spectra specified with the `hyd_wave_spectra` command. `arg1` is the name of the array with the transfer functions. The extreme responses are put in the array whose name is specified by `arg2`. The command creates this array with dimensions $(n_{\text{spectra}}, n_{\text{beta}}, n_{\text{comp}})$, where n_{spectra} and n_{beta} are the number of wave spectra and wave angles, respectively.

Note: The extreme values are defined as $3.72 \times \text{square root of the variance of the response}$, i.e., $3.72 \times \text{square root of the area of the } \text{RAO}^2 \times \text{wave spectrum}$.

The extreme responses are written to a file. If extension is specified, the file name is `project_name.extension`. Otherwise, if filename is specified, the file will have the name specified by filename.

See Also

```
hyd_tf hyd_wave_spectra
```

hyd_modal_pressure

Print modal pressures

Command Syntax

```
hyd_modal_pressure [digits=?]
```

For each panel, prints the incoming + diffraction pressure and the pressure in each mode. The results are in file `project_name.prs2`. For this command to function, the potentials must have been saved in the `hyd_analysis` command.

`digits` is the number of significant digits to print (default is 5).

Note: The pressures determined by the `hyd_analysis_response` command are the total hydrodynamic pressures. This command prints the pressures in each mode, for a unit displacement of that mode. Therefore, for large problems, the file created may be quite large.

See Also

```
hyd_analysis hyd_analysis_postresponse
```

hyd_mooring_line

Mooring line stiffness calculated from elastic catenary cable

Command Syntax

```
hyd_mooring_line m=? n=? [maxiter=?] [tol=?]
m=seg_prop e=emodulus a=area w=wx,wy,wz (m records)
n=nel anchor=x1,z1 end=x2,z2 [#segs=#segs] [tension=tenX,tenY,tenZ]
seg=seg mat=seg_prop L=length (#segs records)
```

m is the number of different mooring line properties

n is the number of mooring stiffnesses

maxiter is the max. # of iterations on the tension (default=30)

tol is the relative tolerance on the end point position (default=1.e-5)

For each set of mooring line segment properties:

seg_prop is the segment property number

emodulus is the modulus of elasticity

area is the effective cross sectional area

wx,wy,wz are the weight/unit length components in global coordinates

For each element:

nel is the stiffness number

x1,z1 are the x and z coordinates of the anchor point

x2,z2 are the x and z coordinates of the top point

#segs is the number of different segments (default=1)

tension is the initial estimate of the tension

seg is the segment number

seg_prop is the segment property number

length is the unstretched segment length

This command will calculate the 3x3 stiffness at the end (attachment) point for a mooring line. The stiffnesses are put in the array .hyd_mooring_K(3,3,n). The geometry of the mooring lines are defined in the plane of the line, and therefore only two coordinates are used to specify the anchor point and end point. The lines actual orientation in the global inertial coordinate system and their attachment to a particular body are specified by the command hyd_assign_mooring. The same stiffness can be assigned multiple times and to multiple bodies. Therefore, it is only necessary to define unique mooring lines once.

The element is based on small strain elastic catenary theory. A shooting method is used to solve the two-point boundary value problem. Specifically, iteration on the tension at the anchor point is carried out until the distance between the calculated position of the end and the specified position of the end, divided by the element length, is less than or equal to the tolerance (tol). For information on the formulation, see H.R. Riggs and T. Leraand, "Efficient Static Analysis and Design of Flexible Risers," J. Off. Mech. Arctic Engrg., Vol. 113, pp. 235-240, 1991, and H.R. Riggs and T. Leraand, "A Robust Element for Static Analysis of Marine Cables," Proc. Third International Offshore and Polar Engineering Conference, Singapore, Vol. 2, pp. 357-363, 1993. Note: the element described in those papers includes fluid drag; this element does not.

See Also

hyd_assign_mooring hyd_mooring_stiffness phyd_mooring_line

phyd_mooring_line

command Syntax

phyd_mooring_line

Print mooring line data as specified by hyd_mooring_line

See Also

hyd_mooring_line

hyd_mooring_stiffness

Specify mooring stiffnesses

Command Syntax

```
hyd_mooring_stiffness #=nstiff
```

```
n=?
```

(Input 3x3 stiffness matrix - 1 row/input record)

Reads nstiff 3x3 mooring stiffness matrices. n is the mooring stiffness number, which must be in the range 1 to nstiff. Multiple stiffness matrices may not be separated by blank or comment lines.

Each 3x3 matrix relates the displacements (u1,u2,u3) and forces of the mooring line at the point where it will be connected to a body. The forces and displacements are defined in a "mooring line coordinate system." A typical situation is: u1 is the horizontal displacement in the plane of the mooring line, u2 is the horizontal displacement normal to u1, and u3 is the vertical displacement. Only unique mooring stiffnesses must be defined.

End input with a blank line.

The command stores the stiffnesses in the array `.hyd_mooring_K(3,3,nstiff)`.

See Also

`hyd_assign_mooring` `hyd_mooring_line` `phyd_mooring_stiffness`

phyd_mooring_stiffness

Print mooring stiffnesses

Command Syntax

phyd_mooring_stiffnesses

Print the mooring stiffnesses.

See Also

hyd_assign_mooring hyd_mooring_line phyd_mooring_stiffness

hyd_nodes

Command Syntax

```
hyd_nodes  #=?  
n=node_no  x=x-coor  y=y-coor  z=z-coor  [lgen=lgen]
```

Reads and generates nodal coordinates. The value specified by # is used to define storage requirements, and it must be greater than or equal to the maximum node number. If this value is missing or 0, it is assumed that existing nodes are being changed or added to, and the previous value applies. lgen is the node number increment for linear generation. Nodes are generated equally spaced along a straight line if two adjacent records do not have sequential node numbers and if lgen on the second line is not zero or blank. Nodes need not be input in sequence.

End input with a blank line.

This command must precede the hyd_panel command.

The coordinates are stored in array .hyd_xyz(3,#), and the maximum possible node number (specified by #) is stored in .hyd_#nodes_tot.

Active nodes are those that are defined explicitly either by this command or another command that creates nodes. The node number of the maximum defined node is stored in .hyd_#nodes. The character vector .hyd_node_active has an "A" for active nodes. Only active nodes can be used.

The nodes command need not be executed as long as the coordinates, which could be generated by another program, are put in the array .hyd_xyz, .hyd_#nodes and .hyd_#nodes_tot are set, and .hyd_node_active is created.

See Also

```
hyd_coordaxs  hyd_coord_trans  hyd_node_tolerance  phyd_nodes
```

phyd_nodes

Command Syntax

```
phyd_nodes [nodes=?,?] [-screen]
```

Print nodal coordinates. A range of node numbers can be specified by nodes=. The first value is the first node number in the range, and the second value is the last number in the range. The default is to print the coordinates for all nodes defined. The default is to print to output file only; if -screen is present, then output is to the screen as well.

See Also

hyd_nodes

hyd_node_tolerance

Command Syntax

```
hyd_node_tolerance [z_tol=?] [x_tol=?] [y_tol=?]
```

Checks for and corrects slight errors in nodal coordinates. This command checks for nodes that should be on the still water plane or planes of symmetry, but numerical precision-related errors cause them to be slightly off these planes. This can happen when coordinates are generated by a mesh generation or CAD program, or when input coordinates are transformed. If a nodal z-coordinate is within `z_tol` (plus or minus) of the still water plane, it is set to 0. For single or double symmetry, if a nodal y-coordinate is within `y_tol` of the x-z plane, it is set to 0. For double symmetry, if a nodal x-coordinate is within `x_tol` of the y-z plane, it is set to 0. The default for each tolerance is 0.001. Tolerances should be positive; if a negative value is specified, the default of .001 is used. (Note: the symmetry type must have been specified by the `hyd_parameters` command for `x_tol` and `y_tol` to be used.

See Also

`hyd_nodes`

hyd_panel

Constant pressure flat fluid panel

Command Syntax

```
hyd_panel n=?  
n=nel nodes=node1,node2,node3,node4 [gen=gen inc=inc1,inc2] &  
  [gen_2d=gen_2d inc_2d=inc1_2d,inc2_2d inc_el=inc_el]
```

n is the maximum panel (element) number

nel is the element number

node1 thru node4 are node numbers

inc1, inc2 are node increments in a "linear sequence"

gen is the number of elements to generate in a sequence

inc1_2d, inc2_2d are node increments between sequences

gen_2d is the number of linear sequences to generate

inc_el is the element increment between sequences

Nodes 1 to 4 are the corner nodes for quadrilateral elements. For triangular elements, node 4 should either be 0 or equal to node 3. The nodes are specified clockwise, looking from the fluid (see sketch below).

A "linear sequence" of elements can be generated by specifying inc1, inc2 and gen. In a linear sequence, nodes 1 and 2 are incremented by inc1; and nodes 3 and 4 are incremented by inc2. gen is the number of elements to generate, so a sequence will have gen+1 elements. To generate a 2D patch of elements, multiple sequences can be specified; inc1_2d and inc2_2d are used to increment the node numbers from one sequence to the next. gen_2d is the number of additional sequences. The element numbers in two successive sequences differ by inc_el (default = numgen+1).

End input with a blank line.

On input, created arrays are:

```
.hyd_panel_el(4,n)    -> node1 - node4  
.hyd_#panels(1)      -> total # of defined panels  
.hyd_#wetpanels(1)   -> total # of wet panels  
.hyd_panelsa(nwet)   -> panel areas  
.hyd_panelsn(3,nwet) -> components of panel normals  
.hyd_panels_type(nwet) -> = 0 for quad; = 1 for triangle  
.hyd_panel_code(n)   -> wet/dry panel codes  
.hyd_panel_#map(n)   -> map panel numbers to internal numbers  
.hyd_xr(nwet,4)      -> x coord. of the panel corners  
.hyd_yr(nwet,4)      -> y coord. of the panel corners  
.hyd_zr(nwet,4)      -> z coord. of the panel corners  
.hyd_xyzc(nwet,3)    -> x,y,z coords. of the panel centers
```

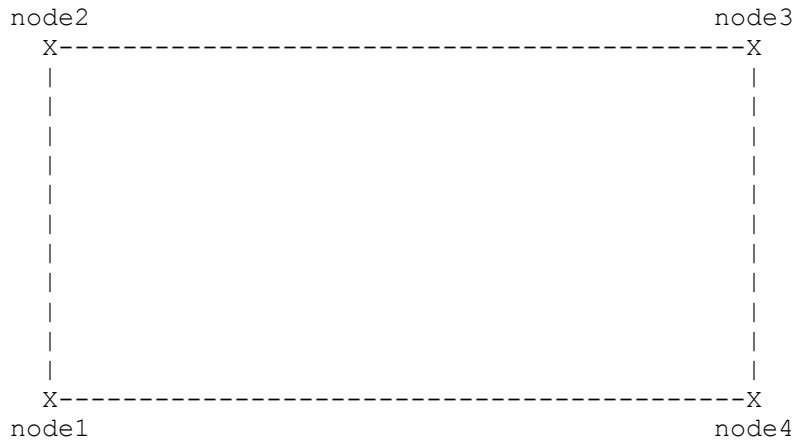
where nwet are the number of wet panels (see below).

Panels that are out of the fluid are ignored for hydrodynamic calculations. The vector .hyd_panels_code contains a character code: blank -> panel not defined; W -> wet panel; D -> dry panel; V -> wet panel that is partially dry; and E -> dry panel that is partially wet. A panel with all nodes below the still water plane is W. A panel with all nodes above the still water plane is D. A panel with some nodes above the still water plane and some below is either V or E. It is V if the panel center is below the still water line. It is E if the center is above the

still water line. Only wet panels (W and V) are included in the hydrodynamic calculations. These panels are numbered internally. The mapping from the "external" numbering to the internal numbering is in the vector `.hyd_panels_#map`.

If the 4 nodes of a quadrilateral panel are not coplanar, it is replaced by an "equivalent" flat panel; it is the data for this flat panel that are stored in the above arrays.

The `hyd_nodes` command must precede this command. Also, because the panel normals and the coordinates of the panel corners and the centroids are in the inertial coordinate system, the `hyd_coords_trans` command, if it is required, must precede this command.



See Also
`phyd_nodes` `phyd_panel`

phyd_panel

Command Syntax

```
phyd_panel [-wet_only]
```

Print hyd_panel element data.

If `-wet_only` is specified, only "wet" panels will be printed.

See Also

hyd_panel

hyd_panel_rmap

Create reverse mapping of panel numbers

Command Syntax

```
hyd_panel_rmap
```

Create the integer vector `.hyd_panel_rmap(#wetpanels)` that contains the input panel number for each wet panel. The wet panel number is assigned internally, and ranges from 1 to `#wetpanels`.

This command is called automatically by `hyd_analysis` if the potentials or pressures are requested. The mapping is needed to print the potentials and pressures.

See Also

```
hyd_analysis  hyd_panel
```

hyd_parameters

Input control parameters for the hydrodynamic analysis.

Command Syntax

```
hyd_parameters h=? [symmetry=?] [nmoder=?] [nmodef=?] [nbodies=?] &  
                [gauge=?] [grav=?] [rho=?] [kh=?] [hdamp=?]
```

h is the water depth

symmetry specifies the symmetry of the panel mesh (see note below)

= 0 -> no symmetry (default)

= 1 -> port-stbd. (x-z) symmetry

= 2 -> port-stbd. and fore-aft (x-z and y-z) symmetry

nmoder is the number of "user-defined" modes (see below)

nmodef is the number of flexible modes

nbodies is the number of multiple bodies

gauge is used for the panel characteristic length (default = 2.5)

grav is the gravitational acceleration (default = 9.80665)

rho is the mass density of the fluid (default = 1025)

kh is the limiting depth beyond which "deep" water is assumed (k is the wave number; default kh = 10)

hdamp is the structural hysteretic damping (.02 means 2% damping)

Dimensional quantities, such as rho and grav, should be specified using consistent units throughout.

The number of user-defined modes are defined by nmoder. The default value for nmoder is 6. Typically, these modes are the traditional rigid body modes of surge, sway, heave, etc, which are defined with the hyd_rigid_modes command. However, any user-defined modes can be specified, including "flexible" modes. To define other than the traditional rigid body modes, the user must supply the matrices required to define these modes (e.g., mass matrix, stiffness matrix, etc.). See the help on the hyd_rigid_modes command for a description of the matrices that must be supplied.

nmodef is the number of flexible modes that are to be input with the command hyd_flex_modes.

The parameter nbodies may be specified if the analysis of multiple bodies (connected or unconnected) is to be carried out. In this case, nbodies is the total number of bodies. Such an analysis can be carried out without specifying nbodies, but defining the rigid body modes is simplified if this option is used. If nbodies is given, then nmoder is set to 6*nbodies.

Important note regarding symmetry:

The parameter symmetry specifies the structural symmetry with respect to the inertial coordinate system. Symmetry is used both to reduce user input and to reduce the computations of the added mass, hydrodynamic damping, and exciting forces. However, these quantities are determined for the entire structure, and the equations of motion are for the entire structure. Hence, the user must specify the structural matrices, such as mass and damping, for the entire structure. For a system of multiple bodies, the system is considered to be the "structure," and it is the structure that must be symmetric to exploit single or double symmetry. For additional discussion of symmetry as used in HYDRAN-XR, see the Getting Started section of the manual.

The characteristic length of a panel is defined as the product of gauge and the square root of the panel area, nondimensionalized with respect to the deep water wave number. Different integration techniques are used to compute the surface integrals involving the Green function and its derivative, depending on whether the distance between the field point and the source point is less than or greater than the characteristic length of the panel. In general, $2 < \text{GAUGE} < 4$. Computational experiments by C.J. Garrison have indicated that gauge = 2.5 is an appropriate choice for the approximations to the surface integrations to be valid (see reference 10).

The data are stored as:

```
.hyd_depth      -> h
.hyd_symmetry   -> symmetry
.hyd_nmoder     -> nmoder
.hyd_nmoder     -> nmoder
.hyd_nmoder     -> nmoder
.hyd_nbodies    -> nbodies
.hyd_deep_water -> kh
.hyd_gauge      -> gauge
.hyd_grav       -> grav
.hyd_hysdamp    -> hdamp
.hyd_fluid_rho  -> rho
```

See Also

```
hyd_analysis  hyd_flex_modes  hyd_rigid_modes  hyd_rmass
```

hyd_postresponse

Command to determine responses

Command Syntax

```
hyd_postresponse [-disp] [-potential] [-pressure]
```

This command determines response quantities based on the solution of the equations of motion as determined by the `hyd_analysis_response` command, which must precede this command. The existing generalized coordinates from that command will be used.

If `-disp` is specified, the nodal displacements are determined based on the generalized displacements.

If `-pressure` is specified, the pressures are calculated based on the generalized displacements. The potentials must have been saved from the `hyd_analysis` command for this option to be possible.

If `-potential` is specified, the potentials for the entire structure are written. This means that for single and double symmetry, they are written for the "actual" panels and also for the "reflected panels." The file indicates how the reflected panels are identified. The potentials must have been saved from the `hyd_analysis` command for this option to be possible.

Results are written to files with names of the form `*.ext`, where `*` represents the project name. Output is:

```
-potential  -> velocity potentials at panel centers (*.pot)
-pressure    -> pressures at panel centers (*.prs and *.hpr)
-disp        -> nodal displacements (*.dis and *.res - a binary file)
```

If `-pressure` is specified, the hydrodynamic pressures are written to file `*.prs` and the change in hydrostatic pressures are written to file `*.hpr`. The change in hydrostatic pressure is defined as $-\rho * \text{grav} * u_z$, where u_z is the displacement in the inertial z direction.

The pressures for the entire structure are written. This means that for single and double symmetry, they are written for the "actual" panels and also for the "reflected panels." The files indicate how the reflected panels are identified.

If `-disp` is specified, the nodal displacements are written to the formatted file `*.dis` and to the unformatted file `*.res`. The file `*.res` is written by the Fortran statement

```
write(f_res)omega, angle, disp
```

Omega and angle are the wave frequency (rad/sec) and wave angle (degrees). The displacements are in the vector `disp(ndof)`, with `ndof = 6*#nodes`. The outer loop is on the wave frequency.

See Also

```
hyd_analysis  hyd_analysis_response
```

hyd_postresponse_P

Command to determine responses

Command Syntax

```
hyd_postresponse_P [-disp] [-potential] [-pressure]
```

This command determines response quantities based on the solution of the equations of motion as determined by the `hyd_analysis_response_P` command, which must precede this command. The existing generalized coordinates from that command will be used.

If `-disp` is specified, the nodal displacements are determined based on the generalized displacements.

If `-pressure` is specified, the pressures are calculated based on the generalized displacements. The potentials must have been saved from the `hyd_analysis` command for this option to be possible.

If `-potential` is specified, the potentials for the entire structure are written. This means that for single and double symmetry, they are written for the "actual" panels and also for the "reflected panels." The file indicates how the reflected panels are identified. The potentials must have been saved from the `hyd_analysis` command for this option to be possible.

Results are written to files with names of the form `*.ext`, where `*` represents the project name. Output is:

```
-potential  -> velocity potentials at panel centers (*.pot)
-pressure   -> pressures at panel centers (*.prs and *.hpr)
-disp       -> nodal displacements (*.dis and *.res - a binary file)
```

If `-pressure` is specified, the hydrodynamic pressures are written to file `*.prs` and the change in hydrostatic pressures are written to file `*.hpr`. The change in hydrostatic pressure is defined as $-\rho * \text{grav} * u_z$, where u_z is the displacement in the inertial z direction.

The pressures for the entire structure are written. This means that for single and double symmetry, they are written for the "actual" panels and also for the "reflected panels." The files indicate how the reflected panels are identified.

If `-disp` is specified, the nodal displacements are written to the formatted file `*.dis` and to the unformatted file `*.res`. The file `*.res` is written by the Fortran statement

```
write(f_res)omega, angle, disp
```

Omega and angle are the wave frequency (rad/sec) and wave angle (degrees). The displacements are in the vector `disp(ndof)`, with `ndof = 6*#nodes`. The outer loop is on the wave frequency.

See Also

```
hyd_analysis  hyd_analysis_response
```

hyd_rigid_modes

Generate the traditional rigid body modes

Command Syntax

```
hyd_rigid_modes [body=?] [nodes=?,?] [panels=?,?] [body_sym=?] &  
               [reflected=?,?,?,?]
```

This command generates surge, sway, heave, roll, pitch, and yaw rigid body modes relative to the body-fixed coordinate system. For a single body, no parameters are required. If this command is used, it must be given before the `hyd_flex_modes` command.

For multiple bodies, as defined by the parameter `nbodies` in the `hyd_parameters` command, it will generate the rigid body modes for the body number specified by the parameter `body`. Hence, one `hyd_rigid_modes` command must be issued for each body for which a mesh is specified explicitly. The first command must be for body 1. The parameters `nodes` and `panels` specify the ranges of nodes and panels corresponding to body. For example, `body=1 nodes=1,100 panels=1,81` mean that body 1 is represented by nodes 1 to 100 and panels 1 to 81. If symmetry in the command `hyd_parameters` is 1 or 2, then the parameter `body_sym` must be specified here. It will be: 0, if the mesh for the particular body is for the entire body; 1, if the mesh for the particular body is for 1/2 the body; and 2 if the mesh for the particular body is for 1/4 the body. This specification is required so that the hydrostatic stiffness will be calculated correctly. Furthermore, if another body is represented by a reflection of this body, then that body number is specified by the parameter `reflected`. For single symmetry, at most one other body can be given. For double symmetry, either 0, 1, or 3 bodies can be specified.

Modes 1 - 6 correspond to body 1. For multiple bodies, modes 7 - 12 correspond to body 2, etc.

The following arrays are created, where `nmode = nmoder + nmoder`. This command populates the arrays with the appropriate data for the first `nmoder` modes.

```
.hyd_modesym(nmode)    -> symmetry codes for each mode  
.hyd_kf(nmode,nmode)   -> hydrostatic stiffness matrix  
.hyd_umx(nnode,nmode)  -> nodal x-displs.  
.hyd_umy(nnode,nmode)  -> nodal y-displs.  
.hyd_umz(nnode,nmode)  -> nodal z-displs.  
.hyd_thx(nnode,nmode)  -> nodal x-rotations  
.hyd_thy(nnode,nmode)  -> nodal y-rotations  
.hyd_thz(nnode,nmode)  -> nodal z-rotations  
.hyd_un(nmode,npanel)  -> panel normal displacements  
.hyd_uz(nmode,npanel)  -> panel z-displacements
```

The symmetry code for mode `j` is stored in `.hyd_modesym(j)`. These codes are only used if the structural symmetry parameter on the `hyd_parameters` command is 1 or 2, indicating single (x-z) or double (x-z and y-z) structural symmetry, respectively. For single symmetry, the modal symmetry code is:

```
1 -> symmetric  
2 -> antisymmetric
```

For double symmetry, the modal symmetry code is:

```
1 -> symmetric/antisymmetric
2 -> antisymmetric/symmetric
3 -> symmetric/symmetric
4 -> antisymmetric/antisymmetric
```

where, e.g., symmetric/antisymmetric means symmetric with respect to the x-z plane and antisymmetric with respect to the y-z plane.

If user-defined modes are used but this command is not, then these arrays should be created with the appropriate data.

See Also

```
hyd_flex_modes hyd_rmass
```


hyd_rmass

Input the structural mass matrix.

Command Syntax

```
hyd_rmass [body=?]
```

The mass matrix is specified "row-wise" immediately following the command, with 1 record per row. Note that only the mass associated with the user-defined modes is input with this command. This command is usually used to input a rigid body mass matrix. The mass matrix should be specified with respect to the same coordinates used to define the modes. If `hyd_rigid_modes` is used to define the rigid modes, then the mass matrix in the body-fixed coordinate system should be given here.

For a single body, the parameter `body` is not required. The command expects the (nmoder,nmoder) matrix to be input.

For multiple bodies, as specified by the parameter `nbodies` in the command `hyd_parameters`, the 6x6 mass matrix for body is expected. Hence, one `hyd_rmass` command must be given for each body. The first command must be for body 1.

The mass matrix is stored in the array `hyd_mstr(nmode,nmode)`. If this matrix does not exist, it will be created. This command is not required. If it is not used, the structural mass matrix (`hyd_mstr`) must be defined in some other way.

See Also

```
hyd_coordaxs hyd_rigid_modes
```

hyd_surf_elevation

Command to determine the "free" surface elevation (i.e., the potentials) after a hydrodynamic analysis.

Command Syntax

```
hyd_surf_elevation [incoming=?] [diffraction=?] [radiation=?] &
                  [dif=difname] [rad=radname]
```

```
incoming    = 1 -> include the incoming potential (default)
              = 0 -> do not include the incoming potential
diffraction = 1 -> include the diffraction potential (default)
              = 0 -> do not include the diffraction potential
radiation   = 1 -> include the radiation potential (default)
              = 0 -> do not include the radiation potential
```

If dif= is specified, the diffraction source strengths are read from file difname; otherwise they are read from file project_name.dif.

If rad= is specified, the radiation source strengths are read from file radname; otherwise they are read from file project_name.rad.

The hyd_analysis and hyd_surf_nodes commands must precede this command. As of now, this command has only been implemented for the case of no symmetry (hyd_parameters command). Also, the -source option must have been specified on the hyd_analysis command to create the files with the source strengths.

The surface elevation is stored in the array .hyd_surf_disp(nnode,nfreq,nbeta), where nnode is the number of surface nodes (hyd_surf_nodes command).

The purpose of this command is to generate the surface elevations for export to graphics programs for visualization; see the command hyd_export_graphics.

NOTE: this command is only implemented for the case of no symmetry.

See Also

```
hyd_analysis  hyd_export_graphics  hyd_parameters  hyd_surf_nodes
hyd_surf_panel
```

hyd_surf_nodes

Command Syntax

```
hyd_surf_nodes  #=?  
n=node_no  x=x-coor  y=y-coor  z=z-coor  [lgen=lgen]
```

Reads and generates surface nodal coordinates in inertial coordinates. The surface nodal coordinates are used to plot surface elevation only. The value specified by # is used to define storage requirements, and it must be greater than or equal to the maximum node number. If this value is missing or 0, it is assumed that existing nodes are being changed or added to, and the previous value applies. lgen is the node number increment for linear generation. Nodes are generated equally spaced along a straight line if two adjacent records do not have sequential node numbers and if lgen on the second line is not zero or blank. Nodes need not be input in sequence.

End input with a blank line.

This command must precede the hyd_surf_panel command.

The coordinates are stored in array .hyd_surf_xyz(3,#), and the maximum possible node number (specified by #) is stored in .hyd_#surf_nodes_tot.

Active nodes are those that are defined explicitly either by this command or another command that creates surface nodes. The node number of the maximum defined surface node is stored in .hyd_#surf_nodes. The character vector .hyd_surf_node_active has an "A" for active nodes. Only active nodes can be used.

The hyd_surf_nodes command need not be executed as long as the coordinates, which could be generated by another program, are put in the array .hyd_surf_xyz, and .hyd_#surf_nodes and .hyd_#surf_nodes_tot are set, and .hyd_surf_node_active is created.

NOTE: Surface nodes cannot be on body panels (as specified by the hyd_nodes and hyd_panel commands. Locate these surface nodes (and panels) a small distance away from the actual body. If a surface node does lie on the body, the surface elevation for that node will likely be reported as NaN.

See Also

```
hyd_surf_node_gen  hyd_surf_node_tolerance  phyd_surf_nodes
```

phyd_surf_nodes

Command Syntax

```
phyd_surf_nodes [nodes=?,?] [-screen]
```

Print surface nodal coordinates. A range of node numbers can be specified by nodes=. The first value is the first node number in the range, and the second value is the last number in the range. The default is to print the coordinates for all nodes defined. The default is to print to output file only; if -screen is present, then output is to the screen as well.

See Also

hyd_surf_nodes

hyd_surf_node_tolerance

Command Syntax

```
hyd_surf_node_tolerance [z_tol=?] [x_tol=?] [y_tol=?]
```

Checks for and corrects slight errors in nodal coordinates. This command checks for nodes that should be on the still water plane or planes of symmetry, but numerical precision-related errors cause them to be slightly off these planes. This can happen when coordinates are generated by a mesh generation or CAD program, or when input coordinates are transformed. Surface nodes must have z-coordinate of 0, so this command enforces that requirement. A warning is printed if the input z-coordinate is further away than ztol. For single or double symmetry, if a nodal y-coordinate is within y_tol of the x-z plane, it is set to 0. For double symmetry, if a nodal x-coordinate is within x_tol of the y-z plane, it is set to 0. The default for each tolerance is 0.001. Tolerances should be positive; if a negative value is specified, the default of .001 is used. (Note: the symmetry type must have been specified by the hyd_parameters command for x_tol and y_tol to be used.

See Also

hyd_surf_nodes

hyd_surf_panel

Surface panel for surface elevation. There are two Command Syntax options.

Command Syntax (option 1)

```
hyd_surf_panel n=?  
n=nel nodes=node1,node2,node3,node4 [gen=gen inc=inc1,inc2] &  
[gen_2d=gen_2d inc_2d=inc1_2d,inc2_2d inc_el=inc_el]
```

n is the maximum panel (element) number

nel is the element number

node1 thru node4 are node numbers

inc1, inc2 are node increments in a "linear sequence"

gen is the number of elements to generate in a sequence

inc1_2d, inc2_2d are node increments between sequences

gen_2d is the number of linear sequences to generate

inc_el is the element increment between sequences

Nodes 1 to 4 are the corner nodes for quadrilateral elements. For triangular elements, node 4 should either be 0 or equal to node 3. The nodes are specified clockwise, looking at the surface from above (see sketch below).

Command Syntax (option 2)

```
hyd_surf_panel -subdivide range=?,? nxm=?,?
```

Option 2 subdivides previously defined panels

range specifies a range of panel ID numbers; all panels in the range are divided

nxm specifies how many panels to subdivide each panel into.

For example, nxm=2,3 would subdivide each panel into 6 elements; 2 in the 1-2 direction and 3 in the 1-4 direction.

A "linear sequence" of elements can be generated by specifying inc1, inc2 and gen. In a linear sequence, nodes 1 and 2 are incremented by inc1; and nodes 3 and 4 are incremented by inc2. gen is the number of elements to generate, so a sequence will have gen+1 elements. To generate a 2D patch of elements, multiple sequences can be specified; inc1_2d and inc2_2d are used to increment the node numbers from one sequence to the next. gen_2d is the number of additional sequences. The element numbers in two successive sequences differ by inc_el (default = numgen+1).

End input with a blank line.

On input, created arrays are:

```
.hyd_surf_panel_el(4,n) -> node1 - node4  
.hyd_#surf_panels(1) -> total # of defined panels  
.hyd_surf_panel_code(n) -> see below  
.hyd_surf_panelpype(nwet) -> = 0 for quad; = 1 for triangle  
.hyd_surf_panel_#map(n) -> map panel numbers to internal numbers
```

where nwet are the number of wet panels (see below).

These panels are only used when exporting the surface elevation to graphics programs for visualization. They are not used for any calculations. The surface elevations are calculated at the surface nodes. The vector .hyd_panel_code contains a character code: blank -> panel not

defined; W -> wet panel (defined); These panels are numbered internally.
The mapping from the "external" numbering to the internal numbering is in
the vector .hyd_surf_panel_#map.

The hyd_surf_nodes command must precede this command.



See Also

phyd_surf_nodes phyd_surf_panel

phyd_surf_panel

Command Syntax

phyd_surf_panel

Print hyd_surf_panel element data.

Only defined panels will be printed.

See Also

hyd_surf_panel

hyd_tf

Calculate transfer functions

Command Syntax

```
hyd_tf [arg1 arg2] [ext=extension file=filename] [-radians] &  
      [-period] [-RAO]
```

After the response analysis has been carried out with the `hyd_analysis_response` command, transfer functions for any response quantity can be obtained with this command. `arg1` is the name of a user-defined 'modal matrix' of dimension (ncomp,nmode), where ncomp is the number of response components and nmode is the number of assumed modes used in the analysis. Column `i` of `arg1` contains the response of each component in mode `i`. The default name for `arg1` is `hyd_modemat`. The transfer functions are obtained by multiplying `arg1` with the generalized coordinates calculated previously by the `hyd_analysis` command. The transfer functions are put in the complex array whose name is specified by `arg2` (default name is `hyd_tf`). The command creates this array with dimensions (nfreq,nbeta,ncomp), where nfreq and nbeta are the number of frequencies and wave angles, respectively.

The transfer functions (magnitude and phase angle) are written to a file. If extension is specified, the file name is `project_name.extension`. If filename is specified, the file will have the name specified by filename. The default file is `project_name.tf`. If `-radians` is specified, the phase angle will be written in radians; otherwise, it will be written in degrees. If `-period` is specified, the transfer functions will be written as a function of wave period rather than wave frequency. If `-RAO` is specified, only the RAO (magnitude) will be printed. The default is to print both RAO and phase angle.

Note: transfer functions for the generalized displacements can be generated by specifying the matrix `arg1` as an nmode x nmode identity matrix. If `arg1` is not specified, and the default matrix (`hyd_modemat`) does not exist, then an identity matrix named `hyd_modemat` will be created and the transfer functions for the generalized displacements will be generated. If this is done for the 6 rigid body modes, for example, the transfer functions for rotations will be in radians. To obtain transfer functions in degrees, `arg1` (`hyd_modemat`) should have the appropriate conversion factors. For example, the following commands would create such a matrix:

```
ident hyd_modemat r=6 c=6  
pi PI  
setr tmp v=180  
scale tmp PI -inv  
put tmp hyd_modemat r=4 c=4  
put tmp hyd_modemat r=5 c=5  
put tmp hyd_modemat r=6 c=6  
rm tmp PI
```

Because of the defaults, the command

```
hyd_tf
```

is equivalent to

```
hyd_tf hyd_modemat hyd_tf ext=tf
```

As many `hyd_tf` commands as desired can be used.

See Also

`hyd_analysis_response` `hyd_irregular`

hyd_velocity

Determine the fluid velocity after a hydrodynamic analysis.

Command Syntax

```
hyd_velocity del=delx,dely,delz [incoming=?] [diffraction=?] &  
            [radiation=?] [dif=difname] [rad=radname] [-keep_pot]
```

delx,dely,delz are the differences in the coordinates to obtain the gradient of the potentials. If only delx is specified, dely and delz are set to delx.

```
incoming      = 1 -> include the incoming potential (default)  
              = 0 -> do not include the incoming potential  
diffraction   = 1 -> include the diffraction potential (default)  
              = 0 -> do not include the diffraction potential  
radiation     = 1 -> include the radiation potential (default)  
              = 0 -> do not include the radiation potential  
-keep_pot     if present, then the velocity potentials are kept
```

If dif= is specified, the diffraction source strengths are read from file difname; otherwise they are read from file project_name.dif.

If rad= is specified, the radiation source strengths are read from file radname; otherwise they are read from file project_name.rad.

The hyd_analysis and hyd_velocity_nodes commands must precede this command. As of now, this command has only been implemented for the case of no symmetry (hyd_parameters command). Also, the -source option must have been specified on the hyd_analysis command to create the files with the source strengths.

The del values are stored in .hyd_vel_del(3).

The velocities are calculated for the nodes in .hyd_vel_xyz(3,nnodes) as follows. First, six points around the nodes, at +-delx, +-dely, and +-delz (in that order) are defined and stored in .hyd_vel6_xyz(3,6*nnodes). If a node is within delz of the still water plane or seafloor, the node is used as the point instead. The potentials for these 6 points are then calculated. The velocities are obtained as, for example, $velocity_x = (\phi(x+delx,y,z) - \phi(x-delx,y,z))/(2 * delx)$. (The expression is adjusted if the points are not 2*delx apart.) The x,y,z velocity components are stored in .hyd_vel_nodes(3,nnodes,nfreq,nbeta), which is complex. nfreq and nbeta are the number of wave frequencies and angles, respectively. The velocity potentials are stored in .hyd_vel6pot(6*nnodes,nfreq,nbeta). This array is usually destroyed, but if -keep_pot is used, it will be kept in the database.

NOTE: this command is only implemented for the case of no symmetry.

See Also

hyd_analysis hyd_parameters hyd_velocity_nodes

phyd_velocity

Command Syntax

`phyd_velocity [-period]`

Print fluid velocities. If `-period` is specified, the velocities will be written as a function of wave period rather than wave frequency.

See Also

`hyd_velocity`

hyd_velocity_nodes

Command Syntax

```
hyd_velocity_nodes  #=?  
n=node_no  x=x-coor  y=y-coor  z=z-coor  [lgen=lgen]
```

Reads and generates inertial coordinates of "nodes" at which to calculate fluid velocities. The value specified by # is used to define storage requirements, and it must be greater than or equal to the maximum node number. If this value is missing or 0, it is assumed that existing nodes are being changed or added to, and the previous value applies.

lgen is the node number increment for linear generation. Nodes are generated equally spaced along a straight line if two adjacent records do not have sequential node numbers and if lgen on the second line is not zero or blank. Nodes need not be input in sequence.

End input with a blank line.

This command must precede the hyd_velocities command.

The coordinates are stored in array .hyd_vel_xyz(3,#), and the total number of defined nodes is put in the scalar .hyd_#vel_nodes. This may be less than # specified on the command line, which is stored in .hyd_#vel_nodes_tot. Currently, the defined nodes are assumed to be sequential from 1 to .hyd_#vel_nodes.

The hyd_velocity_nodes command need not be executed as long as the arrays created by this command are defined "manually".

NOTE: Velocity nodes cannot be on body panels (as specified by the hyd_nodes and hyd_panel commands). Locate the velocity nodes over del away from the actual body (see help on hyd_velocity). In addition, they cannot be above the still water line or below the sea floor. It is recommended to run the command hyd_velocity_node_tolerance before hyd_velocity.

See Also

```
hyd_velocity_node_gen  hyd_velocity_node_tolerance  phyd_velocity_nodes
```

phyd_velocity_nodes

Command Syntax

```
phyd_velocity_nodes [nodes=?,?] [-screen]
```

Print velocity nodal coordinates. A range of node numbers can be specified by nodes=. The first value is the first node number in the range, and the second value is the last number in the range. The default is to print the coordinates for all nodes defined. The default is to print to output file only; if -screen is present, then output is to the screen as well.

See Also

hyd_velocity_nodes

hyd_velocity_node_tolerance

Command Syntax

```
hyd_velocity_node_tolerance [z_tol=?] [x_tol=?] [y_tol=?]
```

Checks for and corrects slight errors in nodal coordinates. This command checks for nodes that are above the still water plane, below the seafloor, or should be on the planes of symmetry, but numerical precision-related errors cause them to be slightly off these planes. This can happen when coordinates are generated by a mesh generation or CAD program, or when input coordinates are transformed. Velocity nodes must have z-coordinates in the range $-\text{water depth} \leq z \leq 0$. This command enforces that requirement. A warning is printed if the input z-coordinate is adjusted. For single or double symmetry, if a nodal y-coordinate is within y_tol of the x-z plane, it is set to 0. For double symmetry, if a nodal x-coordinate is within x_tol of the y-z plane, it is set to 0. The default for each tolerance is 0.001. Tolerances should be positive; if a negative value is specified, the default of .001 is used. (Note: the symmetry type must have been specified by the `hyd_parameters` command for x_tol and y_tol to be used.

See Also

`hyd_velocity_nodes`

hyd_wave

Specify wave frequencies (or periods) and angles for the hydrodynamic analysis.

Command Syntax

```
hyd_wave nbeta=? nfreq=? [-freq] [-period] [-descending] [-delete]
```

nbeta = the number of wave angles (default = 1)

nfreq = the number of incoming wave frequencies (default = 1)

If -period is specified, input wave periods in seconds (default)

If -freq is specified, input wave frequencies in rad/sec

If -descending is specified, the input frequencies or periods are sorted in descending order; otherwise they are put in ascending order.

Following the command, input the wave angles (in degrees) in 1 record and the frequencies/periods in a second record.

If -delete is specified, then nfreq wave frequencies or periods entered on the next line are deleted from the list of frequencies/periods previously input (wave angles are not input and cannot be deleted). At least one frequency must remain after deletion. Corresponding data such as added mass, potentials, and generalized coordinates are also deleted. However, transfer functions generated by the hyd_tf command are not modified. hyd_tf should be re-run after frequencies are deleted.

The data are stored as:

```
.hyd_wave(2)      -> nbeta, nfreq  
.hyd_beta(nbeta)  -> wave angles in radians  
.hyd_freq(nfreq)  -> wave frequencies in radians/sec
```

See Also

hyd_analysis

hyd_wave_dispersion

Command to determine the wave number and wave length

Command Syntax

hyd_wave_dispersion

Given the water depth from the hyd_parameters command, and the wave frequencies from the hyd_wave command, this command solves the dispersion equation to report the wave number and the wave length for each frequency.

See Also

hyd_parameters hyd_wave

hyd_wave_spectra

Specify wave spectra

Command Syntax

```
hyd_wave_spectra #=nspectra
```

```
n=? name=? (Additional data depends on wave spectrum; see below)
```

Reads the data to specify nspectra wave spectra. n is the spectrum number (1 to nspectra) and name is the spectrum name. Allowable spectra and their input are:

```
name=Bretschneider Hs=significant_wave_height To=modal_period
```

```
name=ISSC Hs=significant_wave_height Tv=visual_period
```

```
name=JONSWAP To=modal_period gamma=gamma Xo=Xo  
(default gamma=3.3; default Xo=0)
```

```
name=ITTC Hs=significant_wave_height k=k (default=1)
```

```
name=P-M-Wind U=wind_speed (Pierson-Moskowitz)
```

```
name=P-M-Hs Hs=significant_wave_height (Pierson-Moskowitz)
```

End input with a blank line.

For those spectra that require the gravitational constant, the value specified via the hyd_parameters command is used.

Note: the modal period is the inverse of the (cyclic) frequency at which the frequency spectrum is a maximum.

This command stores the data in the array .hyd_wave_spectrum(6,nspectra).

See Also

```
hyd_irregular hyd_parameters phyd_wave_spectra
```

phyd_wave_spectra

Print wave spectra

Command Syntax

```
phyd_wave_spectra [-spectrum]
```

Print the wave spectra information specified by the `hyd_wave_spectra` command. If the wave frequencies have been defined, e.g., by the `hyd_wave` command, then some statistics, such as calculated significant wave height, for the wave spectra are also reported. The statistics are approximate based on the wave frequencies and the trapezoidal rule of integration. In addition, if `-spectrum` is specified, the wave spectrum is also printed to the output file.

See Also

```
hyd_wave hyd_wave_spectra
```

hyd_wet

Command to estimate the "wet" natural frequencies and mode shapes

Command Syntax

```
hyd_wet [shift=?] [error=?] [form=format]
```

The "wet" natural frequencies and mode shapes are estimated by solving the eigenvalue problem, for each wave frequency, using the added mass that was saved in the database from a previous hyd_analysis command.

Shift is an eigenvalue shift that is used to handle zero eigenvalues. The default shift is 1. Error is the maximum error allowed for a wet frequency, as explained below. Form is the Fortran format in which to write the mode shapes (no spaces are allowed in the character string).

An estimate of the error in the calculated wet frequency ω_n is

$$|\omega - \omega_n|/\omega * 100\%$$

where ω is the wave frequency. Only those frequencies ω_n for which the estimated error is less than or equal to the value of error on the command line are reported. The default value for error is very large, so that all ω_n are reported.

Before the hyd_wet command is given, the structural mass and stiffness matrices must be defined as database members and named hyd_mstr and hyd_kstr, respectively. In addition, the hydrostatic stiffness .hyd_kf must be defined (do not forget the dot!). The dimensions of these matrices should be (nmode,nmode), where nmode is the number of modes. In the typical case, these matrices will have been defined already as part of the wave response analysis.

The "wet" natural frequencies are written to the output file as a function of wave frequency. Both the natural frequencies and mode shapes are also written to file probname.wet. In general, the mode shapes are in terms of the generalized coordinates. However, if the matrix hyd_modemat(ncomp,nmode) exists, the modes will be expressed in terms of the ncomp components. That is, the mode shapes will be multiplied by hyd_modemat, and the result will be written to *.wet. If hyd_modemat exists, the default format for writing is (i15,3x,1pe14.4); otherwise, the default format is (i17,1x,1pe14.4). The integer refers to the component/mode number. This format can be overwritten by specifying a format in the command line. The format must be of the same form as the default and without any blanks. This option is meant to allow the output of more digits for postprocessing. An example is: form=(i15,3x,1pe20.10). Note that the parentheses are required.

The following main arrays are created in the database:

```
.hyd_wet_freq(nmode)           -> wet natural frequencies
.hyd_wet_modes(nmode,nmode)    -> wet mode shapes
.hyd_wet_freq_cumulate(*)      -> converged wet natural frequencies
.hyd_wet_modes_cumulate(nmode,*) -> converged wet modes
```

Note that the first two matrices will contain the data only for the last wave frequency considered, while the second two accumulate the converged natural frequencies and mode shapes over all wave frequencies, i.e., those results that meet the convergence criterion.

Of course, the number of converged frequencies is unknown prior to the analysis.

See Also
hyd_analysis

2.2. General Commands

break_loop

Command Syntax

```
break_loop
```

Break out of a do/while loop. Note: in the case of nested loops, this command breaks out of the entire loop.

See Also

```
do while
```

date

Command Syntax

```
date [arg] [-noprint]
```

Prints the current date and time right justified on an 80 column page. If arg is specified, the date and time are also stored in a character array. If -noprint is specified, the date is not printed.

See Also

```
time
```

do

Command Syntax

```
do arg1 f=? l=? [s=?]  
    block of commands  
end_do
```

arg1 is the name of the loop variable. f is the first value, l is the last value, and s is the increment (default = 1). If arg1 is a real or integer matrix in the database, the first value is used, and the comparison is based on the type of arg1. If arg1 does not exist, an integer scalar with that name is created; it will not be deleted at completion.

While and do commands may have a combined nesting of 15 levels. The following restrictions apply:

Do/while loops in separate files may not be active at one time. Specifically, if a do/while loop is in an input file, the filein command must not be issued from inside a do/while loop.

A break_loop command causes the exit of an entire do/while loop, not just the "subloop" in the case of nested loops.

A return command must not occur inside the body of a do/while loop. A return command may occur in an input file read inside a loop, however, to cause the input file to be exited.

See Also

```
break_loop if while
```

filein

Command Syntax

```
filein [filename] [-noecho]
```

Opens the file "filename" and reads commands for "batch" execution.

If filename is not given, the program tries to open a file with the same name as the project, but with the extension .txt; i.e., .txt is appended to the project name. If that fails, it tries to open the file without the extension.

If filename is specified, the extension .txt, if it exists, can be omitted.

NOTE: The input file must be a plain text file with Windows/Macintosh line endings on Windows/Macintosh platforms, respectively.

If the flag -noecho is present, the input is not echoed to the screen.

Execution returns to the previous input file or to interactive mode when "return" or end-of-file is read.

Although the actual number depends on the number of files the operating system/compiler allows to be open at one time, the program supports a nesting of filein commands up to 28 levels; that is, a filein command can occur in another input file.

The login command is ignored in input files.

See Also
login return

flush

Command Syntax

```
flush [unit=]
```

Flush the buffer for unit. If unit is not specified, standard out buffer is flushed (i.e., the buffer for file project_name.out).

This command is useful, for example, to force writing to the output file without closing the program.

help

Command Syntax

```
help (h) command [-f] [l=?]
```

On-line help for command. For an index of available commands use the command index.

If -f is given, then output is written to the output file; otherwise output is to the screen only. If l is specified, then l lines will be scrolled at a time.

Optional parameters are indicated as such by being enclosed in []. The [] are not part of the command.

if

Command Syntax

```
if arg1 [operator arg2]  
    block of commands  
[else]  
    [block of commands]  
endif
```

arg1 must be a real or integer matrix in the database. The first value in the matrix is used in the comparison.

The simplest form is when only arg1 is specified. In this case, a value of 0 is considered false, and any other value is true.

The more general case is when operator and arg2 are specified. Operator is one of the following FORTRAN conditional operators:

.eq. .ne. .lt. .le. .gt. .ge.

If arg2 is a real or integer matrix in the database, then the comparison is with the first element of arg2. If arg1 and arg2 are of different types, the conditional is evaluated as a real. If arg2 is not found in the database, it is interpreted as a number of the same type as arg1. In this case, only one space may separate the operator and arg2.

If commands may be nested.

See Also
do while

index

Command Syntax

index [topic] [-f] [l=?]

Provides a one-line summary of available commands. If topic is given, topics are general, database, matrix, functions, fe_commands, fem_elements, and misc.

If -f is given, then output is written to the output file; otherwise output is to screen only. If l is specified, then l lines will be scrolled at a time.

logfile

Command Syntax

logfile [-on] [-off]

Turns writing to the log file on or off. Writing is always turned on at the start of a project, and the default option is to turn it on.

See Also
login

login

Command Syntax

login [-noecho]

The file "project_name.log" contains a log of all commands entered in interactive mode. Since the log file is always appended, it contains a complete history of a previous run or sequence of runs (unless logging of commands has been turned off).

The log file can be processed via the login command, which will open the file "project_name.log" as an input file. In this case, a quit in an

input file causes a return to the log file, while it is ignored if it is in the log file itself. A login command is only accepted in interactive mode.

If the flag `-noecho` is given, the input is not echoed to the screen.

See Also
filein logfile

name?

Command Syntax
name? [-f]

Prints project name. If `-f` is given, then output is written to the output file; otherwise output is to screen only.

new_project

Command Syntax
new_project (or newproj or newprob)
Initiates a new project.

See Also
quit save savequit

palias

Command Syntax
palias
Print command aliases.

quit

Command Syntax
quit (q)
Stops execution. This command is ignored if it is read from the log file via the login command.

See Also
login save savequit

read

Command Syntax
read arg filename [-ext]
Read array arg from filename. If `-ext` is present, filename is taken to be an extension, and the file `project_name.filename` is read. It is assumed that the file was created by the write command (unformatted option) and is in the proper form.

See Also
write

rename_file

Command Syntax
rename_file filename1 filename2
Rename filename1 to filename2.

WARNING: Do not rename an open file. In general, commands close files once they finish. Files `project_name.out` and `project_name.log` remain open until the program terminates or a new project is opened.

See Also
[rm_file](#)

return

Command Syntax
`return`

A "batch" command that closes the input file opened with the `filein` command. Causes a return to wherever the `filein` command was issued (the previous input file or interactive mode).

See Also
`filein` `login`

rm_file

Command Syntax
`rm_file filename [-ext]`

Remove file `filename`. If `-ext` is present, `filename` is taken to be an extension, and the file `project_name.filename` is deleted.

See Also
`read` `rename_file` `write`

savequit

Command Syntax
`savequit (sq)`

Saves the data base in the file "`project_name.db`" and stops execution, if not read from the log file via the `login` command.

See Also
`save` `quit`

system_command

Command Syntax
`system_command (term or comm) character_string`

Issues a command to the operating system. The aliases `term` (for terminal) and `comm` (for command prompt) are provided because typing `system_command` is a bit unwieldy. The format of `character_string`, which contains the command, is different between Macintosh and Windows.

Macintosh

An example of the command is

```
term 'cp name.out "name copy.out"'
```

Note that single quotes are required if there are spaces in `character_string`, but they are not allowed within `character_string`.

Windows

An example of the command is

```
term 'cmd.exe /c cp name.out "name copy.out"'
```

Note that single quotes are required at the beginning and end of `character_string`, and they are not allowed within `character_string`. The `/c` is required to terminate the command. The command should also work without the `".exe"`.

time

Command Syntax

```
time arg1 [arg2 arg3]
```

Puts the number of seconds since the start of execution in `arg1`. If `arg2` and `arg3` are given, `arg2` is assumed to be the result of an earlier call to `time`, and the difference between `arg1` and `arg2` will be put in `arg3`. The arguments are real scalars.

Note: This command uses the Fortran function `cpu_time`.

See Also

`date`

while

Command Syntax

```
while arg1 [operator arg2]
```

```
    block of commands
```

```
end_while
```

`arg1` must be a real or integer matrix in the database. The first value in the matrix is used in the comparison.

The simplest form is when only `arg1` is specified. In this case, a value of 0 is considered false, and any other value is true.

The more general case is when `operator` and `arg2` are specified. `Operator` is one of the following FORTRAN conditional operators:

```
.eq. .ne. .lt. .le. .gt. .ge.
```

If `arg2` is a real or integer matrix in the database, then the comparison is with the first element of `arg2`. If `arg1` and `arg2` are of different types, the conditional is evaluated as a real. If `arg2` is not found in the database, it is interpreted as a number of the same type as `arg1`. In this case, only one space may separate the operator and `arg2`.

While and do commands may have a combined nesting of 15 levels. The following restrictions apply:

Do/while loops in separate files may not be active at one time. Specifically, if a do/while loop is in an input file, the `filein` command must not be issued from inside a do/while loop.

A `break_loop` command causes the exit of an entire do/while loop, not just the "subloop" in the case of nested loops.

A `return` command must not occur inside the body of a do/while loop. A `return` command may occur in an input file read inside a loop, however, to cause the input file to be exited.

See Also

break_loop do if

write

Command Syntax

```
write arg filename [-ext] [-f] [-tab] [-a]
```

Write array *arg* to *filename*. If *-ext* is present, *filename* is assumed to be an extension to the project name, and the file *project_name.filename* will be written. The default file type is binary in which the information is written: array type (4 byte integer), dimension (4 byte integer), dimension integers specifying # rows, # cols, etc (each 4 bytes), and then the data. If *-f* is specified, then a formatted write of the data only is done. If *-tab* is also specified, then the columns will be tab separated. If *-a* is specified, then the file will be appended.

See Also

read rm_file

2.3. Database Commands

clear

Command Syntax

```
clear
  Clear (initialize) the data base.
```

See Also

rm save

dir

Command Syntax

```
dir (or ll or ls) [arg] [-f] [c=?]
  dir (or ls) or ll produce a short or long listing of arrays in memory,
  including the dimensions.
  If arg is given, information on that array only is given.
  If -f is given, writes to output file. Otherwise, output is only to
  screen.
  The output for dir (ls) will be in c columns (default = 4)
```

See Also

memory

ll

Command Syntax

```
dir (or ll or ls) [arg] [-f] [c=?]
  dir (or ls) or ll produce a short or long listing of arrays in memory,
  including the dimensions.
  If arg is given, information on that array only is given.
  If -f is given, writes to output file. Otherwise, output is only to
  screen.
  The output for dir (ls) will be in c columns (default = 4)
```

See Also

memory

memory

Command Syntax

```
memory
  Prints data memory used in database
```

mv

Command Syntax

```
mv (rename) arg1 arg2
  Move (rename) arg1 to arg2. If arg2 exists, it is first removed.
```

See Also

rm

readdb

Command Syntax

```
readdb [arg]
  Reads the database in the file "arg.db" and adds to the existing
  database. If arg is not given, the file "project_name.db" is read.
```

See Also

save

rm

Command Syntax

```
rm (del) arg1
```

Remove (delete) arg1 from database. Multiple arguments can be specified.

See Also

```
clear mv save
```

rm*

Command Syntax

```
rm* (del*) arg*
```

Remove (delete) all members from database that begin with "arg". For example,

```
rm* mat*
```

will delete all members that start with "mat". Note that the last character of the argument must be *.

The command syntax requiring * in both the command and the argument is a bit awkward. For example, it would be easier not to include the * in the argument. However, this command could be a bit dangerous in that it might make it easy to delete members accidentally. The syntax is meant to ensure that the intention is to delete everything that matches.

See Also

```
clear mv rm save
```

save

Command Syntax

```
save [arg1]
```

Saves the data base to the file "arg1.db." If arg1 is not give, it is saved to the file "project_name.db."

See Also

```
quit readdb savequit
```


2.4. Matrix Commands

add

Command Syntax

```
add arg1 arg2 [arg3] [r=?] [n=?] [c=?] [m=?]
```

Adds arg2 to arg1.

If arg3 is input, result is put in new matrix arg3.

If r is input, start adding at row r.

If n is input, only add n rows.

If c is input, start adding at column c.

If m is input, only add m columns.

Note: For 3-D arrays, only straight addition is supported. If any parameters are specified, they are ignored.

See Also

sub subcol sumcol

arpack

Command Syntax

```
arpack [#=?] [which=?] [maxit=?] [digits=?] [shift=?] [minv=?] [-freq]
```

By using ARPACK routines, determine the eigenvalues and eigenvectors of the generalized eigenvalue problem:

$$[K] \{\phi\} = \omega^2 [M] \{\phi\}$$

= # of eigenvalues to determine

which = Specify which of the Ritz values to compute

LA - compute # smallest (algebraic) eigenvalues

SA - compute # largest (algebraic) eigenvalues

LM - compute # smallest (in magnitude) eigenvalues (default)

SM - compute # largest (in magnitude) eigenvalues

BE - compute # eigenvalues, half from each end of the spectrum. When # is odd, compute one more from the high end than from the low end.

maxit = max. # of iterations (default = 30)

digits = convergence tolerance = $10^{-\text{digits}}$ (default = 8)

minv = minimum number of subspace vectors (default = $\min(2\#, \#+8)$)

shift = frequency shift (default = 0)

-freq -> the frequencies (ω) are determined

K and M are assumed to be symmetric matrices in sparse storage. The diagonals of K and M are in `.sparse_diag` and `.sparse_diag2`; the off-diagonals in `.sparse_off` and `.sparse_off2`, and the index arrays are `.sparse_ptr` and `.sparse_indx`. A diagonal M may be used, wherein `.mstr` is the vector of diagonal masses (do not provide `.sparse_diag2` and `.sparse_off2` in this case).

The eigenvalues (frequencies) are put in `.omega2` (`.omega`), depending on the parameter `-freq`. The eigenvectors are put in `.phi`.

If K is singular then a shift must be applied prior to calling this command. For this command, the shifted eigenvalues are equal to the actual eigenvalues + shift. (That is, if a shift is needed for frequency analysis, use a positive shift.)

The ARPACK routines were obtained from Rice University; for more

information see <http://www.caam.rice.edu/software/ARPACK/index.html>

See Also

eigval jacobi ptosparse

array3d_slice

Command Syntax

```
array3d_slice arg1 arg2 [r=?] [c=?] [t=?]
```

Extracts a "plane" of a 3-D array.

arg1(n,m,p) is a 3-D array and arg2 is a matrix. arg2 is created as follows:

If r is specified, $\text{arg2}(1:m,1:p) = \text{arg1}(r,1:m,1:p)$

If c is specified, $\text{arg2}(1:n,1:p) = \text{arg1}(1:n,c,1:p)$

If t is specified, $\text{arg2}(1:n,1:m) = \text{arg1}(1:n,1:m,t)$

Hence, if t is specified, arg2 is "table" t from arg1. This can also be obtained from the cp command.

See Also

array3d_unslice cp

array3d_unslice

Command Syntax

```
array3d_unslice arg1 arg2 [r=?] [c=?] [t=?]
```

Inserts 2-D matrix arg1 into a "plane" of 3-D array arg2. This operation is the opposite of array3d_slice. The two dimensions of arg1, n and m, must agree with two of the three dimensions of arg2 as follows.

If r is specified, $\text{arg2}(r,1:n,1:m) = \text{arg1}(1:n,1:m)$

If c is specified, $\text{arg2}(1:n,c,1:m) = \text{arg1}(1:n,1:m)$

If t is specified, $\text{arg2}(1:n,1:m,t) = \text{arg1}(1:n,1:m)$

Note: Only one of r, c, and t should be given. If two or more are specified, only one is used, in the priority order r, c, t.

See Also

array3d_slice

cp

Command Syntax

```
cp (copy) arg1 arg2 [r=?] [n=?] [c=?] [m=?] [table=?]
```

Copies arg1 to arg2. For a straight copy of any dimension array, do not include any parameters. A partial copy of an array can be made by specifying parameters.

If r is input, start copying at row r.

If n is input, only copy n rows.

If c is input, start copying at column c.

If m is input, only copy m columns.

If table is input, operate on one table of a 3-D array; i.e., extract the table.

For 3D arrays, this command can only duplicate the array (no parameters specified) or extract one table, or part thereof.

See Also

cpdg put putdg

cpdg

Command Syntax

cpdg arg1 arg2 [r=?] [n=?]

Copies the diagonal elements of arg1 to arg2.

If r is input, start copying at row r.

If n is input, only copy n rows.

See Also

cp put putdg

diag_mult

Command Syntax

diag_mult arg1 arg2 arg3

Multiply arg1 times arg2. arg1 is a diagonal matrix stored as a vector.

The result is stored in arg3. The dimension of arg1 must equal the number of rows of arg2.

See Also

mult tmult mult_elem

dim_reduce

Command Syntax

dim_reduce arg1 arg2 index=?

Copies all but the last dimension of a multi-dimensional array. If arg1 is an n-dimensional array, arg2 will be an (n-1)-dimensional array that is obtained by setting the last index of arg1 to the value specified by index. The default for index is 1.

eigval

Command Syntax

eigval [#=?] [maxit=?] [digits=?] [shift=?] [ss_size=?] &
[rigid=?,?,?,?,? cg=?,?,?,? node_range=?,?] [-vec_init] &
[-freq] [-disk] [-sparse] [-random] [seed=?]

Eigenvalue solution by either subspace (SS) iteration or Jacobi method of the generalized eigenvalue problem:

$$[K] \{\text{phi}\} = \text{omega}^2 [M] \{\text{phi}\}$$

= 0 -> use Jacobi and solve for all eigenvalues
> 0 -> use SS to solve for # of lowest eigenvalues
maxit = max. # of iterations (default = 30)
digits = convergence tolerance = 10^-digits (default = 8)
ss_size = number of subspace vectors (SS only; default=max(2#,#+8))
shift = frequency shift (default = 0)
rigid = six values, corresponding to 6 rigid body modes in global coordinates (SS iteration only)

```

    0 -> do not form corresponding rigid body mode
    1 -> form corresponding rigid body mode
cg      = point about which rigid modes are calculated
          (default = 0,0,0)
node_range = range of nodes to calculate rigid modes
          (default is to include all nodes)
-vec_init -> use #vecs initial vectors in matrix
          eigval_init(neq,#vecs)- SS only
-freq     -> the frequencies (omega) are determined
-disk     -> the factored and original stiffnesses will be swapped
          to/from disk for SS iteration (profile only)
-sparse   -> a sparse stiffness matrix is used

-random   -> use a random number generator to create
          the starting vectors (SS iteration only)
seed     .ge. 0 -> seed for generator (default = 1)
        .lt. 0 -> use next random number

```

K and M are assumed to be symmetric matrices. The default option is to use an upper profile storage scheme. In this case, K and M are assumed to be in .kstr and .mstr, respectively. The locations of the diagonal elements are assumed to be in .kdiag_loc. If a sparse storage option is chosen, then the diagonals of K and M are in .sparse_diag and .sparse_diag2; the off-diagonals in .sparse_off and .sparse_off2, and the index arrays are .sparse_ptr and .sparse_indx. In either case a diagonal M may be used, wherein .mstr is the vector of diagonal masses.

The eigenvalues (frequencies) are put in .omega2 (.omega), depending on the parameter -freq. The eigenvectors are put in .phi.

For subspace iteration, K and M are unchanged if there is no shift; if there is a shift, .kstr will be the shifted matrix.

For both SS and Jacobi, if K is singular then a shift must be applied. For this command, the shifted eigenvalues are equal to the actual eigenvalues + shift. (In other words, for frequency analysis typically use a positive shift.)

For the jacobi solution, M must be positive definite.

See Also

arpack jacobi ptosparse

extract

Command Syntax

```
extract arg1 arg2 c=? v=? [tol=?] [-eq -ne -gr -gt -le -lt]
```

Extracts rows in arg1 for which the value (v=) in column (c=) matches the criterion:

```

-eq -> equal to value +- tol (default)
-ne -> not equal to value
-ge -> greater than or equal to value
-gt -> greater than value
-le -> less than or equal to value
-lt -> greater than or equal to value

```

The result is put in arg2. A row vector is treated as a column vector.

For complex numbers, value is complex but the comparison is done on absolute values, except for equality and inequality.

fft

Command Syntax

```
fft arg1 [-ferziger] [-ooura] [-forward] [-inverse]
```

The default is to determine the forward (direct) Fourier transform of a function `arg1`. `arg1` is replaced with the frequency coefficients. If `-inverse` is specified, the inverse transform is determined, in which case the input `arg1` should be the frequency coefficients and the output `arg1` is the time history.

`arg1` must be an $N \times m$ dimensional array of equally spaced function values. N is the number of sample points and m is the number of sequences to be transformed.

Default Algorithm

The default is to use `fftpack` available at netlib.org that was originally written by Paul N. Swarztrauber, as modified by P. Bjorstad (version 3, June 1979). The command creates the work array `$fft_wsave`, which must not be changed between calls to `fft`. That is, the inverse transform will use the previously determined `$fft_wsave` vector created by the forward transform. This array can be deleted if no further inverse transformations with the same value of N are to be done. This command assumes `arg1` is a real matrix for a real function.

Ooura Algorithm

If option `-ooura` is specified, the Ooura FFT routines available at <http://faculty.prairiestate.edu/skifowit/fft/> are used (July 2010). `arg1` can be either double precision real or double precision complex. N must be a power of two. The work arrays `$fft_ip` and `$fft_w` are used, which can be deleted when they are no longer needed.

Ferziger Algorithm

If option `-ferziger` is specified, the FFT routine by Wouk based on the algorithm by Ferziger in Numerical Methods for Engineering Applications is used, but converted to double precision. The original code is available at <http://www.netlib.no/netlib/misc/fft.f> (July 2010). `arg1` is double precision complex. N must be a power of two. No work array is used.

You should clearly understand the scaling, the convention, and the storage scheme that these algorithms use. A first step is to refer to the simple test file in the examples folder.

See Also

```
fft_helper
```

fft_helper

Command Syntax

```
fft_helper H P HP
```

This is a special purpose function to multiply transfer functions H with frequency components P , with the result HP , which is stored in a form consistent with P .

The use of this routine is designed for dynamic structural analysis, where the work flow is as follows: 1) use the command `fft` to obtain the frequency coefficients `P` from the time history of the loading; 2) multiply the transfer functions `H` with the corresponding `P`, forming `HP`; 3) use `fft` to obtain carry out an inverse Fourier transform to obtain the time history of the response.

In theory both `H` and `P` are complex and the same size, in which case a simple element-by-element multiply is sufficient. But this is not usually the case. It is more likely that `H` is complex of dimension $N/2+1$, in which N is the number of points used for the discrete Fourier transform (this command requires that N is even). If the command `fft` with the default algorithm is used to obtain `P`, then the complex `P` is stored in a real matrix of dimension N . The storage scheme for `P` is revealed by the example `fft` in the folder `distr_examples`. This command will do the multiplication directly. The complex product is stored in the real matrix `HP`, using the same storage scheme as used for `P`. This is the storage scheme required by the default algorithm in the command `fft` to obtain the inverse Fourier transform.

Multiple transfer functions can be processed at the same time. In summary, the input and output are as follows:

```
H(N/2+1,#func) - complex, #func is the number of transfer functions
P(N)           - real
HP(N,#func)   - real
```

N must be even.

A typical analysis would then involve three commands:

```
fft P
fft_helper H P HP
fft HP -inverse
```

In the first command, input `P` is the loading sampled at N equidistant time points and output `P` is the frequency components. The second command is as explained above. In the third command, output `HP` is the time history of the response at the N equidistant time points.

See Also
`fft`

ftopro

Command Syntax

```
ftopro arg1 arg2 arg3
```

Store "full" symmetric matrix `arg1` in row vector `arg2` using upper profile storage. `arg3` is the vector of diagonal locations in `arg2`. Length of `arg3` is `arg1_rows + 1`. Length of `arg2` is `arg3(arg1_rows+1) - 1`.

See Also
`ptoful` `pmult`

gauss

Command Syntax

```
gauss arg1 arg2 [-pp] [ #digits=?]
```

Uses elimination (LU factorization) to solve a system of linear equations (real or complex). arg1 is the coefficient (square) matrix and arg2 is the "RHS." The solution is put in arg2.

For real matrices, a warning is printed if the number of "digits" lost in a diagonal is greater than or equal to #digits (default = 7).

If -pp is specified, partial pivoting with implicit scaling is carried out; otherwise, elimination without pivoting is done.

For large, symmetric matrices, use psolve.

See Also

```
ftopro invert psolve
```

get_dim

Command Syntax

```
get_dim arg1 arg2 [-rows] [-cols]
```

Gets the dimensions of arg1 and stores them in arg2. If -rows is present, only the number of rows is stored. If -cols is present, only the number of columns is stored.

ident

Command Syntax

```
ident (identi or identc) arg [r=?] [c=?]
```

Transform matrix arg such that all terms are zero except arg(i,i), the "diagonals", which are one. If arg is square, it becomes an identity matrix.

If r and c are input, integer (identi), real (ident), or complex (identc) matrix arg(r,c) is created.

See Also

```
input zero
```

input

Command Syntax

```
input (inputi or inputch or inputc) arg [r=?] [c=?] [-null]
```

Input a real (input), integer (inputi), character (inputch) or complex (inputc) matrix or vector.

r is the number of rows (default = 1).

c is the number of columns (default = 0).

For character matrices and vectors, the rows are null-terminated if the parameter -null is specified.

See Also

```
ident input3d zero
```

input3d

Command Syntax

`input3d` (`input3di` or `input3dc`) `arg` [`r=?`] [`c=?`] [`t=?`]
Input a real (`input3d`), integer (`input3di`), or complex (`input3dc`) 3-D array with dimensions (`r,c,t`).
`r` is the number of rows (default = 1).
`c` is the number of columns (default = 1).
`t` is the number of "tables" (default = 1).

A 3-D array is a collection of 2-D matrices, each of which is considered a "table." That is, `arg(:,:,1)` is table 1, `arg(:,:,2)` is table 2, and so forth. Each table is input as a 2-D matrix (see the command `input`), and the tables are input sequentially.

See Also
`Input`

interpolate

Command Syntax

`interpolate tin fin tout fout` [`prepad=?`] [`postpad=?`]
Interpolate using the data set (`tin,fin`) to create (`tout,fout`).

The input data set is defined by `tin(n)` and `fin(n,nfunc)`, where `tin` is a vector of real "time" values, `fin` is a matrix of real or complex function values. The output values are defined at the discrete "times" specified in `tout(N)`. The real (or complex) vector `fout(N,nfunc)` is created based on linear interpolation of the data in [`tin,fin`].

If the range of `tout` is larger than the range of `tin`, the output values are pre-padded and/or post-padded. The default is to prepad with the 1st value in `fin` and to postpad with the last value in `fin`. If a value is given for the argument `prepad`, prepadding will use that value. If a value is given for the argument `postpad`, postpadding will use that value. (Use these options to prepad and postpad with 0, for example.)

It is assumed that `tin` and `tout` are in ascending order.

See Also
`series`

invert

Command Syntax

`invert arg1 arg2`
The inverse of `arg1` is calculated and stored in `arg2`. Straight Gauss elimination is used. The matrix `arg1` is modified. For large, symmetric matrices, use `psolve`.

See Also
`ftopro` `psolve`

jacobi

Command Syntax

`jacobi k m phi lambda` [`d=?`] [`maxit=?`] [`digits=?`] [`-freq`]
Eigenvalue solution by Jacobi method for the generalized eigenvalue problem:

$$[K] \{\phi\} = \lambda [M] \{\phi\}$$

where K and M are symmetric, square, positive definite matrices, which will be modified, and

phi = matrix of all eigenvectors (to be created)
lambda = vector of all eigenvalues (to be created)
d = 0 -> mass matrix is input as 2-D matrix(default)
= 1 -> diagonal mass "vector" (not implemented)
maxit = maximum # of iterations (default = 30)
digits = # digits accuracy in eigenvalues (default = 8)
-freq = if present, then lambda is the square root of the eigenvalues.

See Also
Eigval

join

Command Syntax

joinh (or joinv) arg1 arg2 arg3
arg3 is formed by joining (concatenating) arg1 and arg2
joinh (horizontal joining) puts the matrices "side-by-side."
joinv (vertical joining) puts arg1 "on top of" arg2.

max

Command Syntax

max arg1 arg2 [c=?] [-abs]
Finds the maximum value in column c=? of arg1; the corresponding row is put in vector arg2. The default is c=1. If the flag -abs is present, the element with the maximum absolute value is found. If arg1 is a row vector, it is treated as a column vector.

See Also
min norm

min

Command Syntax

min arg1 arg2 [c=?]
Finds the minimum value in column c=? of arg1; the corresponding row is put in vector arg2. If arg1 is a row vector, it is treated as a column vector.

See Also
max

mult

Command Syntax

mult (or tmult) arg1 arg2 arg3
For mult, arg3 = arg1 * arg2
For tmult, arg3 = Transpose(arg1) * arg2

See Also
mult_col mult_elem scale

mult_col

Command Syntax

```
mult_col arg1 arg2 arg3
```

Multiply the columns of arg1 and the columns of arg2, and store the result in the row vector arg3. Hence, arg3(i) is the dot product of columns i of arg1 and arg2. The dimensions of arg1 and arg2 must be identical.

See Also

```
mult mult_elem
```

mult_elem

Command Syntax

```
mult_elem arg1 arg2 [arg3] [-inv]
```

Multiply each element of arg1 with the corresponding element of arg2, and store the result in arg3. If arg3 is not given, the result is stored in arg1. If -inv is specified, then the elements of arg1 are divided by arg2. The dimensions of arg1 and arg2 must be identical.

See Also

```
mult mult_col
```

norm

Command Syntax

```
norm arg1 arg2 [-max] [-l2]
```

Finds the vector "norm" of the columns of arg1; results are put in vector arg2. The default is to find the maximum value in each column. If the flag -max is present, the maximum absolute value of the elements is found. Otherwise, if the flag -l2 is present, the square root of the sum of the squares is found. If arg1 is a row vector, it is treated as a column vector.

See Also

```
sumcol
```

pmult

Command Syntax

```
pmult arg1 arg2 arg3 arg4
```

Multiply $\text{arg1} * \text{arg2} = \text{arg3}$, where arg1 is a symmetric matrix stored in upper profile form. arg4 is the vector of diagonal locations in arg1. Length of arg4 is arg2_rows + 1

See Also

```
ptoful ftopro
```

print

Command Syntax

```
print or printf (p or pf) arg1 [r=?] [c=?] [cols=?] [l=?] &  
[-nohead] [row#=?] [-f] [form=format] [-phase] [table=?]
```

Prints arg1. Printing begins at row r and column c (default = 1). If l is given, then l lines will be scrolled at a time. If -nohead is given, then the usual row/column headings will not be printed. The default is to print every fifth row number (1, 5, 10, 15, etc.) in the heading;

this can be overridden by specifying a different value with row#=. If -f is specified, then printing is to the output file only (and not to the screen). In this case, scrolling is not done. If -phase is specified and arg1 is complex, then the magnitude and phase angle (degrees) will be printed rather than the real and imaginary components.

The default output formats are ni6 and lpne12.4, where n=12, n=6, or n=3 for integer, real, and complex matrices, respectively. If cols is specified, n=cols. (pf prints using nf12.5 format.) Alternative formats can be specified by the parameter form=. In this case, format must be a valid FORTRAN format, enclosed in (), without any embedded blanks and with a maximum length of 160 characters. If a format is specified, the parameters cols=, l=, and -nohead are ignored (row/column headings are not printed).

If arg1 is a 3 dimensional array, then it is printed as a sequence of 2 dimensional arrays, called "tables." For example, table 1 is arg1(*,*,1). If table is specified, only that table is printed. For dimensions greater than 4, only the first table is printed.

See Also
xprint

psolve

Command Syntax

```
psolve arg1 arg2 arg3 [digits=?]
```

Solves $\text{arg1} * \text{arg2} = \text{arg3}$, where arg1 is a symmetric matrix stored in upper profile form. arg2 is the "load" vector, with neq rows, which is replaced by the solution. arg3 is the vector of diagonal locations in arg1. Length of arg3 is neq + 1. The size of arg1 = arg3(neq+1) - 1. A warning is printed if the loss of precision in a diagonal element is greater than or equal to digits (default = 7).

See Also
ftopro gauss invert pmult ptoful

psolve16

This command is not implemented, although it would be simple to do so.

ptoful

Command Syntax

```
ptoful arg1 arg2 arg3
```

Put symmetric matrix arg1 stored in profile form into the square matrix arg2. arg3 is the vector of diagonal locations in arg1. The length of arg3 is assumed to be the # diagonals of arg1 + 1. The dimensions of arg2 are arg2(#diag,#diag).

See Also
ftopro gauss invert pmult

ptosparse

Command Syntax

ptosparse arg1 arg2 [arg3]

Put symmetric matrix arg1 stored in profile form, with diagonal locations specified in arg2, into sparse matrix. The diagonals are stored in .sparse_diag and the nonzero off-diagonals are stored in .sparse_off. The number of nonzeros in each row are stored in .sparse_ptr, and the column number for each nonzero is stored in .sparse_indx. The number of nonzero off-diagonals is stored in .sparse_size. If arg3 is specified, then it is interpreted as a matrix with the same profile as arg1. Its values are stored in .sparse_diag2 and .sparse_off2.

See Also
sparse_mult

put

Command Syntax

put arg1 arg2 [r=?] [c=?]

Puts arg1 into arg2.

If r is input, starts putting at row r.

If c is input, starts putting at col c.

See Also
putdg

putdg

Command Syntax

putdg arg1 arg2 [r=?]

Puts arg1 into diagonals of arg2.

If r is input, starts putting at row r.

See Also
put

scale

Command Syntax

scale arg1 [arg2 e=?,?] [v=?] [-inv]

Multiplies arg1 by the scalar arg2(e1,e2) or by the value v. If -inv is present, multiplies by the inverse of the scalar.

See Also
mult

series

Command Syntax

series arg x=x1,x2 [inc=?] [w=?] [max=?]

Creates a real data series starting at x1 and ending at x2. The series is put in the row vector arg. The initial spacing is inc (default=1); the spacing between points 2 and 3 is inc * w, and the spacing between points n and n+1 is inc * w^(n-1). However, the spacing between the last two points may differ, as the last value will not be larger than x2. If w < 1, care must be taken to ensure that the series will reach x2. However, at most max (default=100000) number of points are generated. By specifying a large enough x2, one can generate a series with exactly max points.

See Also
series2d

series2d

Command Syntax

```
series2d arg p1=x1,y1,z1 p2=x2,y2,z1 p3=x3,y3,z1 p4=x4,y4,z1 &  
           [w=?,?] [n=?,?] [dim=?]
```

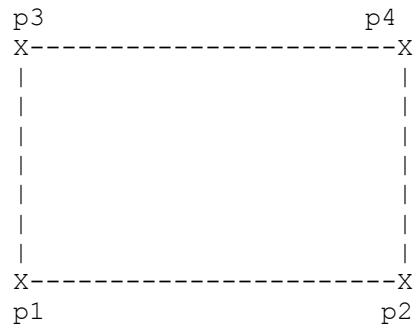
Creates a real data series on a 2-d grid, which is put in the matrix `arg(n1*n2,dim)`. The series data can be viewed as points in an X-Y-Z coordinate system, which are generated within the "quadrilateral" defined by the corner points p_j with coordinates (x_j, y_j, z_j) , $j=1,4$; see sketch below. A grid of $n_1 \times n_2$ points is generated, as follows. Linear generation is used to generate n_1 points between p_1 and p_2 , and between p_3 and p_4 , based on w_1 . Linear generation also is used to generate n_2 points between p_1 and p_3 , and between p_2 and p_4 , based on w_2 . The points generated between p_1 and p_3 are connected through linear generation, based on n_1 and w_1 , with the points generated between p_2 and p_4 . A gridlike ordering of points is carried out such that p_1 is point 1, p_2 is point n_1 , p_3 is point $n_1*(n_2-1)+1$, and p_4 is point n_1*n_2 . If n_2 is zero, then generation is along the line p_1 to p_2 only.

`dim` can be 1, 2 or 3 (default=2). Only the first `dim` coordinates are kept in `arg`.

If `arg` exists, and if the # rows = n_1*n_2 and the # columns is at least `dim`, then the data are put in the first `dim` columns of the existing matrix.

Linear generation is defined such that the initial spacing between points is equidistant unless the positive weight w (default=1) is specified, in which case the spacing between points k and $k+1$ is equal to w times the spacing between $k-1$ and k .

An alternative type of generation can be carried out using the commands `nodes` and `node_gen`.



See Also
series

seti

Command Syntax

```
seti (setr) arg v=? or
```

setch arg s=? [-null]

Set scalar arg to value. Seti and setr set integer and real scalars, respectively, equal to the value v. Setch sets a character string (a row matrix with a maximum length of 80, no blanks allowed) specified by s. If -null is present, the character string in arg will be null-terminated.

This command is more convenient than input and inputi for scalars and inputch for character constants.

sort

Command Syntax

sort arg1 [arg2] [key=?,?,?] [-reverse]

Sorts arg1 in ascending order. A maximum of three sort keys (columns on which sorting is done) may be specified by the identifier key; the first value is the first key, etc. If no keys are specified, the first key is taken to be column 1. The result is put in arg2, if it is given; otherwise arg1 is changed. If -reverse is present, sort is in descending order. If arg1 is a row vector, it is sorted as a column vector.

sparse_matrix_clean

Command Syntax

sparse_matrix_clean

Removes arrays created by the command ptospars as well as some arrays that are created internally as part of the sparse solver. Arrays that will be removed if they exist are: .sparse_mmik, .sparse_mmim, .sparse_mmf, .sparse_addr, .sparse_ptr, .sparse_indx, .sparse_diag, .sparse_diag2, .sparse_off, .sparse_off2, .sparse_ordr, and .sparse_size.

See Also

lsolve ptospars

sparse_mult

Command Syntax

sparse_mult arg1 arg2

Premultiplies the matrix arg1 by a (symmetric) sparse matrix and puts the result in arg2. The sparse matrix is assumed to be in a form compatible with the command ptospars, and the arrays .sparse_diag, .sparse_off, .sparse_ptr, .sparse_indx, and .sparse_size are assumed to exist. The array .sparse_ordr defines the reordering of the matrix; if it does not exist, it is assumed that the matrix has not been reordered.

See Also

ptospars

split

Command Syntax

split arg1 [arg2] [c=?]

Splits arg1 based on columns and joins the matrices vertically into arg2, with c columns (default=1). For example, let arg1 have 10 rows

and 8 columns and let $c=4$. Then arg2 will have 20 rows and 4 columns, such that the first 10 rows will be the same as the first 4 columns of arg1 , and the last 10 rows will be the same as the last 4 columns of arg1 . If arg2 is not given, then arg1 is replaced with the new matrix. If c does not divide evenly into the # of columns of arg1 , then the additional columns in the last rows of arg2 will be zero. If c is greater than or equal to the # of columns of arg1 , a simple copy of arg1 is done. This command could be used to form a column vector from a row vector by specifying $c=1$. (The transpose command - `trans` - is preferred in this case.)

See Also
unsplit

sub

Command Syntax

```
sub arg1 arg2 [arg3] [n=?] [r=?] [c=?] [m=?]
```

Subtracts arg2 from arg1 .

If arg3 is input, result is put in new matrix arg3 .
If r is input, start subtracting at row r .
If n is input, only subtract n rows.
If c is input, start subtracting at column c .
If m is input, only subtract m columns.

See Also
subcol add sumcol

subcol

Command Syntax

```
subcol arg1 arg2 arg3 [c=?]
```

Subtracts arg2 from arg1 .

Result is put in new matrix arg3 . If c is input, start subtracting at column c . This command is used to subtract all but the first $c-1$ columns of two equal size matrices.

See Also
sub add put putdg

sumcol

Command Syntax

```
sumcol arg1 arg2 [-abs]
```

Sums the columns of arg1 ; results are put in vector arg2 . If the flag `-abs` is present, the absolute values of the elements are summed. If arg1 is a row vector, it is treated as a column vector.

See Also
add sub subcol

to_complex

Command Syntax

```
to_complex arg1 [arg2] or  
to_complex2 arg1 arg2 arg3
```

The command `to_complex` converts arg1 into a complex matrix. arg1

becomes the real part of arg2, if it is given; otherwise arg1 is replaced.

The command `to_complex2` creates the complex array agr3 using arg1 as the real part and arg2 as the imaginary part.

See Also
`to_int` `to_real`

to_complex2

Command Syntax

```
to_complex arg1 [arg2] or  
to_complex2 arg1 arg2 arg3
```

The command `to_complex` converts arg1 into a complex matrix. arg1 becomes the real part of arg2, if it is given; otherwise arg1 is replaced.

The command `to_complex2` creates the complex array agr3 using arg1 as the real part and arg2 as the imaginary part.

See Also
`to_int` `to_real`

to_int

Command Syntax

```
to_int arg1 [arg2] [-trunc]
```

Converts arg1 into an integer matrix. The values in arg1 are converted to the nearest 4 byte integers and placed in arg2, if it is given; otherwise arg1 is replaced. If `-trunc` is given, the values are truncated instead (for complex arg1, the values are always truncated).

See Also
`to_complex` `to_real`

to_real

Command Syntax

```
to_real arg1 [arg2]
```

Converts agr1 into a real matrix. The values in arg1 are converted to 8 byte reals and placed in arg2, if it is given; otherwise arg1 is replaced. If arg1 is complex, the real part is returned.

See Also
`to_complex` `to_int`

to_real16

Command Syntax

```
to_real16 arg1 [arg2]
```

Converts agr1 into a real*16 matrix. The values in arg1 are converted to 16 byte reals and placed in arg2, if it is given; otherwise arg1 is replaced.

Note: Only limited support is provided for real*16 data types.

See Also

psolve16 to_complex to_int to_real

to_vector

Command Syntax
to_vector arg1

Converts a matrix with only one dimension greater than 1 to an N-dimensional vector. For example, A(N,1) and A(1,N) will both be converted to A(N).

tmult

Command Syntax
mult (or tmult) arg1 arg2 arg3
For mult, $\text{arg3} = \text{arg1} * \text{arg2}$
For tmult, $\text{arg3} = \text{Transpose}(\text{arg1}) * \text{arg2}$

See Also
mult_col mult_elem scale

trans

Command Syntax
trans arg1 [arg2]
Transposes arg1 to arg2, if it is given. Otherwise, replaces arg1 with its transpose.

unsplit

Command Syntax
unsplit arg1 [arg2] [r=?]

Unsplints arg1 based on rows and joins the matrices horizontally into arg2, with r rows (default=1). For example, let arg1 have 20 rows and 4 columns and let r=10. Then arg2 will have 10 rows and 8 columns, such that the first 4 columns will be the same as the first 10 rows of arg1, and the last 4 columns will be the same as the last 10 rows of arg1. If arg2 is not given, then arg1 is replaced with the new matrix. If r does not divide evenly into the # of rows of arg1, then the additional rows in the last columns of arg2 will be zero. If r is greater than or equal to the # of rows of arg1, a simple copy of arg1 is done. This command could be used to form a row vector from a column vector by specifying r=1. (The transpose command - trans - is preferred in this case.)

See Also
split

unwrap

Command Syntax
unwrap arg1 [arg2] [r=?]

"Unwraps" the rows of arg1 puts the result into arg2, which will have r rows (default=1). For example, let arg1 have 20 rows and 4 columns and let r=10. Then arg2 will have 10 rows and 8 columns, such that the first row will be the same as the first 2 rows of arg1, row 2 will be the same as rows 3 and 4 of arg1, etc. If arg2 is not given, then arg1

is replaced with the new matrix. If *r* is greater than or equal to the # of rows of *arg1*, a simple copy of *arg1* is done. Otherwise, *r* must divide evenly into the number of rows in *arg1*.

See Also

`split` `unsplit` `wrap`

wrap

Command Syntax

`wrap arg1 [arg2] [c=?]`

"Wraps" the rows of *arg1* and puts the result into *arg2*, which will have *c* columns (default=1). For example, let *arg1* have 10 rows and 8 columns and let *c*=4. Then *arg2* will have 20 rows and 4 columns, such that the first 2 rows will be the same as the first row of *arg1*, rows 3 and 4 will be the same as the second row of *arg1*, etc. If *arg2* is not given, then *arg1* is replaced with the new matrix. If *c* does not divide evenly into the # of columns of *arg1*, then the additional columns in *arg2* will be zero. If *c* is greater than or equal to the # of columns of *arg1*, a simple copy of *arg1* is done.

See Also

`split` `unsplit` `unwrap`

xprint

Command Syntax

`xprint (xp) arg1 [cols=?] [row#=?]`

Expanded print of *arg1* with virtually all significant digits. If *cols* is given, then a maximum of *cols* columns will be printed at a time. *row#* sets the increment used to print row headings (default = 5).

If *t* is given, *arg* is a 3-D array rather than a matrix, but the rules above apply.

See Also

`print`

zero

Command Syntax

`zero (zeroi or zeroc) arg [v=value] [r=? c=? t=?]`

Zeros *arg*.

If *value* is given, the matrix is filled with *value* rather than zero. If *r* or *c* is input, matrix *arg*(*r*,*c*) is created of type real (*zero*), integer (*zeroi*), or complex (*zeroc*).

See Also

`input` `ident`

2.5. Mathematical Functions

abs

Command Syntax

`abs arg1 [arg2] [x=?] [c=?]`

Takes absolute value of elements in *arg1*. The result is put in *arg2*, if

it is given; otherwise the values in arg1 are replaced.

The default is to operate on every element in arg1. However, to operate on a single column only, use the parameter x to specify the column number. If x is specified and arg2 is not specified, the result is put in column y of arg1 (default=x). If c is specified and x is not, the first c-1 columns are the same as arg1.

bessel_j or bessel_y

Command Syntax

bessel_j (or bessel_y) [n=] arg1 [arg2] [x=?] [c=?]

Takes Bessel function of 1st kind (bessel_j) or Bessel function of the 2nd kind (bessel_y) of order n of elements in arg1. The result is put in arg2, if it is given; otherwise the values in arg1 are replaced.

n=0 is the default

The default is to operate on every element in arg1. However, to operate on a single column only, use the parameter x to specify the column number. If x is specified and arg2 is not specified, the result is put in column y of arg1 (default=x). If c is specified and x is not, the first c-1 columns are the same as arg1.

conjugate

Command Syntax

conjugate arg1 [arg2] [x=?] [c=?]

Calculates the complex conjugates of elements in arg1. The result is put in arg2, if it is given; otherwise the values in arg1 are replaced.

The default is to operate on every element in arg1. However, to operate on a single column only, use the parameter x to specify the column number. If x is specified and arg2 is not specified, the result is put in column y of arg1 (default=x). If c is specified and x is not, the first c-1 columns are the same as arg1.

epsilon

Command Syntax

epsilon arg1 arg2

Returns a positive small value relative to 1 of the same real kind as arg1. arg2 will be a scalar.

erf

Command Syntax

erf (or erfc or erfc_scaled) arg1 [arg2] [x=?] [c=?]

Takes error function (erf), complementary error function (erfc) or the scaled complementary error function (erfc_scaled) of elements in arg1. The result is put in arg2, if it is given; otherwise the values in arg1 are replaced.

The default is to operate on every element in arg1. However, to operate on a single column only, use the parameter x to specify the column number. If x is specified and arg2 is not specified, the result is put in column y of arg1 (default=x). If c is specified and x is not, the first c-1 columns are the same as arg1.

exp

Command Syntax

```
exp arg1 [arg2] [x=?] [c=?]
```

Calculates the exponential of elements in arg1. The result is put in arg2, if it is given; otherwise the values in arg1 are replaced.

The default is to operate on every element in arg1. However, to operate on a single column only, use the parameter x to specify the column number. If x is specified and arg2 is not specified, the result is put in column y of arg1 (default=x). If c is specified and x is not, the first c-1 columns are the same as arg1.

gamma

Command Syntax

```
gamma arg1 [arg2] [x=?] [c=?]
```

Takes the gamma function of elements in arg1. The result is put in arg2, if it is given; otherwise the values in arg1 are replaced.

The default is to operate on every element in arg1. However, to operate on a single column only, use the parameter x to specify the column number. If x is specified and arg2 is not specified, the result is put in column y of arg1 (default=x). If c is specified and x is not, the first c-1 columns are the same as arg1.

log

Command Syntax

```
log arg1 [arg2] [x=?] [c=?]
```

Calculates the natural log of elements in arg1. The result is put in arg2, if it is given; otherwise the values in arg1 are replaced.

The default is to operate on every element in arg1. However, to operate on a single column only, use the parameter x to specify the column number. If x is specified and arg2 is not specified, the result is put in column y of arg1 (default=x). If c is specified and x is not, the first c-1 columns are the same as arg1.

log10

Command Syntax

```
log10 arg1 [arg2] [x=?] [c=?]
```

Calculates the common log of elements in arg1. The result is put in arg2, if it is given; otherwise the values in arg1 are replaced.

The default is to operate on every element in arg1. However, to operate on a single column only, use the parameter x to specify the column number. If x is specified and arg2 is not specified, the result is put in column y of arg1 (default=x). If c is specified and x is not, the first c-1 columns are the same as arg1.

pi

Command Syntax

```
pi arg1
```

Creates the scalar arg1 with the value of pi.

power

Command Syntax

```
power arg1 [arg2] [p=?] [-r] [x=?] [c=?]
```

Raises each element in `arg1` to the power `p` (default = 2). If `-r` is present, the power `p` is real; otherwise, it is an integer (default). The result is put in `arg2`, if it is given; otherwise the values in `arg1` are replaced.

The default is to operate on every element in `arg1`. However, to operate on a single column only, use the parameter `x` to specify the column number. If `x` is specified and `arg2` is not specified, the result is put in column `y` of `arg1` (default=`x`). If `c` is specified and `x` is not, the first `c-1` columns are the same as `arg1`.

sqrt

Command Syntax

```
sqrt arg1 [arg2] [x=?] [c=?]
```

Takes square root of elements in `arg1`. The result is put in `arg2`, if it is given; otherwise the values in `arg1` are replaced.

The default is to operate on every element in `arg1`. However, to operate on a single column only, use the parameter `x` to specify the column number. If `x` is specified and `arg2` is not specified, the result is put in column `y` of `arg1` (default=`x`). If `c` is specified and `x` is not, the first `c-1` columns are the same as `arg1`.

trig

Command Syntax

```
"trig_function" arg1 [arg2] [x=?] [c=?]
```

Evaluates the function "trig_function" for each element in `arg1`. The result is put in `arg2`, if it is given; otherwise the values in `arg1` are replaced. Valid functions are

```
cos sin tan asin acos atan sinh cosh tanh
```

The default is to operate on every element in `arg1`. However, to operate on a single column only, use the parameter `x` to specify the column number. If `x` is specified and `arg2` is not specified, the result is put in column `y` of `arg1` (default=`x`). If `c` is specified and `x` is not, the first `c-1` columns are the same as `arg1`.

2.6. Finite Element Commands

bcid

Command Syntax

```
bcid [#=?] [r=?,?,?...] [print=?]  
n=node_no r=?,?,?... [print=?] [inc=inc]
```

Specifies restraints on nodal degrees-of-freedom (DOFs).

```
# = number of DOFs per node (default = 6)  
r = comma-separated string of 0's and 1's, # values long  
  0 -> DOF is free (value to be calculated)  
  1 -> DOF is "fixed", i.e., value is set to 0  
print = print code  
  0 -> displacements will not be printed by pdisp  
  1 -> displacements will be printed by pdisp (default)
```

Values specified on the `bcid` line are nodal defaults. The most common situation is when there are 6 DOFs per node. `r=0,0,0,0,0,0` would then correspond such that 1 to 3 are for the translations along the X, Y, and Z axes, and 4 to 6 are the corresponding rotations. For example, `r=0,0,1,1,1,1` would imply only X and Y displacements are non-zero. Some elements require more than 6 DOFs per node; this can be specified by `#=`. Note that if a number less than 6 is specified, 6 will be used. That is, there are at least 6 DOFs per node.

Subsequent lines can be used to specify different values for specific nodes. However, all nodes have the same maximum number of DOFs, which is stored in `.#dofs_per_node`. If a restraint value < 0 is specified instead of a 0 or 1, then the corresponding displacement for the node is the same as for node number `|value|`. Generation occurs if two records do not have sequential nodes and `inc`, the node number increment, is not 0. The generated nodes have the same restraints as the last node entered.

Nodes need not be input in numerical order.

End input with a blank line.

The displacement restraints are stored in array `.bcid(,#,#nodes)`, which is required by the `num_eqs` or `form_k` command to number the equations, and the `pdisp` command to print displacements. The print codes are stored in `.bcid_print(#nodes)`.

The term "displacement" is used for the nodal variables, but actually it can be any quantity (e.g., temperature, etc.).

See also

```
form_k nodes num_eqs pbcid pdisp
```

body_frc2d

Command Syntax

```
body_frc2d n=?  
n=distribution# c=coefficients
```

Input polynomial coefficients used to define `n` "distributions" of 2d body forces for elements in a plane. A maximum of a 5th degree

polynomial in two variables (21 coefficients) is allowed. The coefficients are stored in the array `.body_f2d(21,n)`. These can then be used by elements to define body forces.

End input with a blank line.

See Also

`d213to9` `pbody_frc2d`

check_diag

Command Syntax

```
check_diag arg1 arg2 [zero=?] [v=?] [-noprnt]
```

Check the diagonal elements of matrix `arg1`, stored in profile form, for zeroes. `arg2` is the vector of diagonal locations in `arg1`. The length of `arg2` is assumed to be the # diagonals of `arg1` + 1. If `zero` (default = 0.0) is given, then any diagonal with an absolute value less than `zero` is replaced by `v` (default = 1.0). The equation numbers corresponding to the changed diagonals are reported unless the flag `-noprnt` is given. This command is useful to prevent a stiffness matrix from being singular as a result of "unconnected" nodes.

See Also

`form_k`

consolidation

Command Syntax

```
consolidation steps=? [dt=?] [theta=?] [dprin=?] [fprin=?] [k=?] [time=?]  
[history=?]
```

Consolidation time integration command

```
steps = number of time steps  
dt     = time step (default is to use existing time step)  
theta  = 2-step integration parameter (default=1.0 -> forward Euler)  
dprin  = print flag for nodal displacements  
        = 0 -> not printed from command (default)  
        = N -> print every N time steps  
fprin  = print flag for element state  
        = 0 -> not printed from command (default)  
        = N -> print every N time steps  
k      = 0 -> form stiffness by calling routine form_k (default)  
        = 1 -> use existing stiffness  
time   = initial time (default = 0)  
history = time increment at which results are saved (see below)
```

A `form_k` command must precede the first consolidation command. It is also important that the `nodef` command follow the `form_k` command.

The increment in external loads is determined from the `.load_pat` array created by the command `nodef`. For this command, the load "patterns" refer to the time step. Imposed displacements and pressures are specified by the `imposed_displ` command. Again, for this case the pattern number is the time step. This command interprets the values specified in the `nodef` and the `imposed_displ` command as incremental values.

At each time step, the calculated displacements and pressures are placed in `.disp`. If `history` is not zero, then the displacements at every `history` time

units (e.g., seconds) is saved in `.disp_history(neq+1,*)`. The last value in each column is the time of the results. For each node, DOF 4 is the pressure.

Note: Using an existing stiffness (`k=1`) should only be done if the time step and theta parameter are unchanged and the degrees-of-freedom with imposed displacements have not changed.

See Also
imposed_displ nodef

dampers

Command Syntax
dampers [-noecho]
n=node_no c=cx,cy,cz,cxx,cyy,czz [inc=]

Reads nodal dampers.

If `-noecho` is specified, the data will not be echoed to the output file.

`cj` and `cjj` are the nodal dampers with respect to the `j` axis. Generation occurs if two records do not have sequential nodes and `inc`, node number increment, is greater than 0. The generated nodal data will have the same value as the record with `inc`.

Dampers for repeated nodes are accumulated.

End input with a blank line.

The data are stored in the array `.dampers_inp(7,*)`, in the order node #, 6 damping components.

The nodal dampers are assembled into the structure damping by the `form_c` command.

See Also
form_c pdamper

pdampers

Command Syntax
pdampers

Print input nodal dampers.

See also
dampers

direct_th

Command Syntax
direct_th arg1 dt=? endtime=? [-linear] [damping=?,?] [begintime=]
[save_disp=?] [save_restart=?] [beta=] [gamma=] [conv=?,?]
[maxit=?] [k_iter=?] [-morison] [-strict_conv]
[conv_option=1]

Determine the direct time history response.


```

arg1      = the time function (nptsx2 matrix)
dt        = time step
endtime   = simulation time
-linear   = response is linear; no iteration (see below)
damping   = mass and stiffness coefficients for Rayleigh damping
begintime = time to begin the simulation (default=0)
save_disp = integer to specify that the displ. are saved every
           disp_save time steps (default=1)
save_restart= integer to specify that the displ., velocities and
           accelerations are saved every save_restart time steps
           (see below)
beta      = coefficient in Newmark method (default=1/4)
gamma     = coefficient in Newmark method (default=1/2)
conv      = convergence tolerances on displacement and forces
           (see below)
maxit     = maximum # of iterations within a time step (default=10)
k_iter    = # of iterations with constant stiffness (default=2)
-morison  = determine incoming wave velocities and accelerations
-strict_conv= requires force and displacement convergence
conv_option = if specified, then "small" displacements are ignored
           for convergence

```

This command integrates the equations of motion directly, using Newmark's method. The default beta, gamma coefficients correspond to the unconditionally stable constant average acceleration method. The command assumes the stiffness matrix is in .kstr, in profile storage, and the mass matrix is in .mstr and is either diagonal or in profile storage. The damping matrix is in .cstr and can be either diagonal or in profile storage. If it does not exist, the values specified by damping are used to form a Rayleigh damping matrix. The first value for damping is the mass coefficient, and the second value is the stiffness coefficient. Provide non-zero initial displacements in u_init(neq), nonzero initial velocities in v_init(neq), and nonzero initial accelerations in a_init(neq). Otherwise, they are assumed to be zero.

The default is to begin the simulation at time = 0. In this case, any existing time history results created by previous calls of this command are deleted. A nonzero time can be specified with begintime. In this case, if there are existing displacements from previous calls of this command, then the new nodal displacements are appended to those results.

The displacements are saved every save_disp time steps. The displacements, velocities, and accelerations will be saved every save_restart time steps and can be used for a restart analysis. These are always saved at endtime.

The number of time steps is the nearest integer to (endtime-begintime)/dt. Hence, the simulation may be slightly longer or shorter than endtime.

The load patterns are defined by the nodef command. If the array load_comb exists, it is interpreted as load combination factors to be applied with the "load patterns" in .load_pat. Otherwise, the first load pattern is used. Element loads are included if they correspond to the load pattern used.

The time variation of the loads is defined by arg1, which is a 2D matrix defining f(t). The first column is time, and the second column is the function value. If the time function is not defined out to endtime, the

function is continued at the last defined value out to endtime.

The displacements, that is the time history nodal response, is stored in `.nodal_th_disp(neq,nsaved)`, where `neq` is the number of equations. The value for `nsaved = (endtime-begintime)/(dt*save_disp)+1`. The times are saved in `.nodal_times(nsaved)`.

The displacements, velocities and accelerations are saved in `.th_restart_disp(neq,*)`, `.th_restart_vel(neq,*)`, and `.th_restart_acc(neq,*)`, where the number of columns is the number of times at which the results are saved. For linear analysis, the results are only saved at endtime. For nonlinear analysis, the number of columns depends on the parameter `save_restart`. The default is to save these only at endtime. In any event, the last columns are the results at endtime. The times are saved in `.th_restart_times(*)`. Because nodal displacements, velocities, and accelerations are saved, multiple `direct_th` commands can be used in a "restart" fashion.

If `-linear` is specified, no iteration is carried out and no check on equilibrium is made. It assumes strictly linear behavior. In this case, `maxit`, `save_restart`, `conv`, and `k` are ignored. Displacements, velocities, and accelerations are saved only at endtime. However, `save_disp` is used to save the displacements.

For a nonlinear analysis, the time stepping is done as follows. Within a time step, at the end of the first iteration the unbalance in the equation of motion is calculated. If the maximum absolute value in the unbalance is less than `conv(2)`, then no iteration is carried out; the analysis proceeds to the next time step and any unbalance is applied in that time step. Otherwise, iteration is carried out based on the unbalance. Iteration continues until the displacement increment for each DOF in the current iteration is not greater than `conv(1) * the previously calculated displacement increment in the current time step`, up to `maxit` iterations. If `-strict_conv` is specified, then both displacement and force convergence must be satisfied. Sometimes, the solver can get stuck trying to satisfy the convergence criterion for "small" displacements, for example, those that should be nearly zero. Specifying `conv_option` will ignore the DOFs for which the displacement increment in the time step is less than 10^{-4} times the maximum displacement increment in the time step. The stiffness is kept constant for `k_iter` iterations.

WARNING: If nonzero initial conditions are provided, including nonzero forces at start time, they should reflect dynamic equilibrium. The nonlinear solver may be able to adapt errors in the initial conditions, but the linear solver cannot. This warning is mostly applicable to user-defined initial conditions. Initial conditions based on a previous analysis should already satisfy dynamic equilibrium.

See Also

`form_c` `form_k` `form_m` `initial_conditions` `modal_th` `nodef` `nsolve`

disp_cntl

Command Syntax

`disp_cntl`

Read displacement control information. The data is input immediately following the command in the format:

Node DOF Factor

The command creates the vector `.disp_cntl_vec`, which is used to calculate the generalized displacement in a displacement control solution.

The command is terminated by a blank line.

See Also
nsolve

el_iso_mat1

Command Syntax

```
el_iso_mat1 arg1 e=? nu=? [-stress] [-strain] [-axisym]
arg1 = 2D stress-strain constitutive matrix
e = modulus of elasticity
-stress -> plane stress
-strain -> plane strain
-axisym -> axisymmetric case
```

Creates the stress-strain constitutive matrix for a linear elastic, isotropic material.

elem_alias

Command Syntax

```
elem_alias
Print aliases for elements.
```

elem_grp

Command Syntax

```
elem_grp e=? code=?
Adds or deletes element groups to be processed
```

```
e = element group number
code = integer code used by the element during processing
```

The group number is the same as the element "number" (1 for `elem01`, 2 for `elem02`, etc.). The code for group "i" is stored in location "i" of the vector `.elem_grp`. Groups with a nonzero code will be processed by element processing commands, such as `form_k`, `state`, and `response`. The array and codes are set up during element definition, but this command allows specific groups to be removed and/or added from processing.

See Also
`elem_alias` `form_k` `form_m` `response` `state`

eq_direction

Command Syntax

```
eq_direction arg1 [-x] [-y] [-z]
```

Define the seismic direction vector `arg1` (often denoted "r" in structural dynamics). `arg1` is a vector of 1's and 0's. If `-x` is specified, x degrees-of-freedom have a 1; other degrees-of-freedom have a 0. `-y` and `-z` are similar. Only one of the coordinate directions can

be specified.

The equations must have been numbered before this command can be used. It expects the number of equations to be in `.#eqs` and the equation numbers to be in `.node_eqs`.

See Also

`form_k` `num_eqs`

export_graphics

Export mesh to graphics program input file

Command Syntax

```
export_graphics [-target] [-deformed case=? T=? -autoscale scale=? &  
steps=?] [file=?]
```

Export the mesh to a graphic program's input text file.

The graphics program is specified by the argument `-target`. Gmsh (<http://www.geuz.org/gmsh/>) is the only option supported at this time. That is, the available option is `-Gmsh`, and it is automatically chosen.

The default is to plot the undeformed mesh.

If `-deformed` is specified, then the displaced shapes, one for each displacement set (column) in `.disp` will be created. If `case` is specified, only the column number specified by the integer `case` is processed. In that case, the displacements can be scaled if `steps` is omitted. If `-autoscale` is specified, then a scale factor will be determined automatically for the displacement set. The scale factor will be determined such that the displacement is `scale*length`, where `length` is the maximum length of a bounding box enclosing the model. The default is 2% (i.e., `scale=2`). If `-autoscale` is not specified, then `scale` specifies the scaling factor. The default is no scaling.

If `-displ` is specified, then the `x,y,z` displacements (`u,v,w`) in `.disp` are the plot variables.

`T=` is the name of a real vector of "times". The size of the vector is the same as the number of columns in `.disp`. This option can be used as follows. If mode shapes are being plotted, then `T` could contain the natural periods. If `.disp` contains a time sequence of displacements, then `T` could contain the time for each shape. If `.disp` contains static displacements, then `T` could contain the load case number. If a vector is not specified, one is created with the values `1,2, ...`

The normal mode shapes from an eigenvalue analysis can also be displayed. Just copy the modes to `.disp`; i.e., run the following command prior to this command:

```
cp .phi .disp
```

The HYDRAN-XR GUI can display the mesh and static displaced shapes. Only one displaced shape per file is allowed, however (i.e., use `case=` option). It cannot display animations.

The displacements in `.disp` (including mode shapes) can be animated to

more easily visualize the shape. Use case to specify the column number and provide a value for steps, the number of frames per cycle that will be generated. These can be viewed in an external program such as Gmsh.

If filename is specified, the results will be written to the file filename; otherwise they will be written to the file project_name.msh (Gmsh).

fem_error

Command Syntax

```
fem_error [-SE] [-user]
```

Estimate the error in a finite element solution.

The total error (initialized to -1) is put in .fem_error. In general, the elements also calculate their relative error (element error/global error) in response to this command.

The parameter -SE specifies that the estimate of the error in strain energy is used (default).

The parameter -user specifies that the error is based on a user-defined error function (see the appropriate element for details).

Each element is responsible for consistency with the specification of error type. See the appropriate element help page.

form_c

Command Syntax

```
form_c [-diag]
```

Form global damping and put in vector .cstr. The default is to form the consistent damping matrix and store it in upper profile form. The command expects the location of the diagonals to be in the vector .kdiag_loc, which is formed when the equations are numbered by the command form_k or num_eqs. The element groups to be included in the damping assembly are specified by nonzero codes in the array .elem_grp, which is defined during element definition.

If the flag -diag is specified, a diagonal damping matrix is formed.

The scalar .cstr_type is created with a value of 0 for consistent damping and 1 for diagonal damping matrix.

If a diagonal damping matrix is requested and an element returns a consistent matrix, the element damping is "lumped" by scaling the diagonals of C_e , the element consistent damping, by the factor $(u^T C_e u) / (\text{sum of diagonals})$, where u is a vector of ones. Hence, the lumped damping and the consistent damping will have the same value for $(u^T C u)$. This lumping procedure is not appropriate for all types of elements.

See Also

```
elem_grp form_m num_eqs
```

form_k

Command Syntax

```
form_k [-#eqs] [-stiff] [-loads]
```

Form global stiffness and loads. The element groups to be included in the stiffness assembly are specified by nonzero codes in the array `.elem_grp`, which is defined during element definition. The default operation is to number the equations, if necessary, form the stiffness, and form the loads. If any flag is specified, then the operation is determined by the flags specified, as follows:

```
-#eqs    -> number the equations
-stiff   -> form stiffness
-loads   -> form nodal loads
```

If `-#eqs` is not specified, the nodal equation numbers in `.node_eqs` are used. If `.node_eqs` does not exist, or if `-#eqs` is specified, the equations are numbered based on the restraint codes in `.bcid`, which is established by the `bcid` command. The nodal equation numbers are put in `.node_eqs(#dofs_per_node,#nodes)`, and the number of equations is put in the scalar `.#eqs`. The locations of the diagonal elements of the stiffness are stored in `.kdiag_loc` and the size of the stiffness matrix is stored in `.kstr_size`; hence, the elements must have been defined.

If the stiffness is formed, the upper profile is stored in the vector `.kstr`. A scalar `.k_status` is also created and zeroed. This value is modified when the stiffness is factored.

If the loads are formed, the load array `.load_pat(#eqs,#patterns)`, where `#patterns` is the number of load patterns defined in `.load_inp` (see command `nodef`). If this array does not exist, then the zero load pattern array `.load_pat(#eqs,1)` is created. Equivalent nodal loads from element loads are assembled into `.load_pat`.

See Also

```
bcid  elem_grp  load_summary  nodef  num_eqs  peqs
```

form_m

Command Syntax

```
form_m [-diag]
```

Form global mass and put in vector `.mstr`. The default is to form the consistent mass matrix and store it in upper profile form. The command expects the location of the diagonals to be in the vector `.kdiag_loc`, which is formed when the equations are numbered by the command `form_k` or `num_eqs`. The element groups to be included in the mass assembly are specified by nonzero codes in the array `.elem_grp`, which is defined during element definition.

If the flag `-diag` is specified, a diagonal mass matrix is formed.

The scalar `.mstr_type` is created with a value of 0 for consistent mass and 1 for diagonal mass.

Nodal masses input with the mass command are assembled into the global mass matrix, also.

If a diagonal mass matrix is requested and an element returns a consistent matrix, the element mass is "lumped" by scaling the diagonals of M_e , the element consistent mass, by the factor $(u^T M_e u) / (\text{sum of diagonals})$, where u is a vector of ones. Hence, the

lumped mass and the consistent mass will have the same value for ($u^T M u$). This lumping procedure is not appropriate for all types of elements.

See Also

elem_grp form_k mass mass_summary num_eqs

imposed_displ

Command Syntax

```
imposed_displ  
n=node_no dof=dof disp=p1,p2,...
```

Reads imposed nodal displacements.

n is the node number
dof indicates the nodal degree of freedom, that is, 1 for x translational displacements, 2 for y, ..., and 6 for z rotational displacements
pi is the displacement corresponding to load pattern "i"

The last values input for a node and dof are used. Degrees of freedom for which displacements are not specified are assumed to be "free", that is, displacements are not imposed. However, although different displacements can be imposed for different load patterns, a degree of freedom with specified displacements in one or more patterns are assumed to be constrained in all patterns. If the displacement value is not specified for a load pattern, it is assumed to be zero.

If the optional identifiers (those in [], e.g., n=) are used in an input record, all data in that record must have the correct identifier.

End input with a blank line.

The data are stored in the array `.disp_inp(2+#patterns,*)`. Each column contains the node #, DOF # (1-6), displacements in each load pattern. As many columns as necessary are used.

The `nodef` command, which defines the number of load patterns, must precede this command.

Caution: Displacements may not be imposed with all solution commands, and some commands may interpret these values differently. Check the help for the particular solution command.

See also

nodef pimposed_displ

initial_conditions

Command Syntax

```
initial_conditions arg [-noecho]  
n=node_no u=x,y,z,xx,yy,zz [inc=]
```

Reads initial values for nodal displacements, velocities or accelerations.

If `-noecho` is specified, the data will not be echoed to the output

file.

arg is the name of the neq vector to save the initial conditions. u are the six values for node_no. Generation occurs if two records do not have sequential nodes and inc, node number increment, is greater than 0.

The last values specified for a node is used.

End input with a blank line.

The data are stored in the vector arg(neq). The equations must have been numbered prior to this command; see the form_k command.

This command is meant to specify nonzero initial displacements and/or velocities and/or accelerations for a time history analysis. To specify multiple quantities, repeat the command with different names for arg.

Note: initial conditions should be consistent with the equation of motion.

See Also

direct_th form_k modal_th

load_summary

Command Syntax

load_summary

The sum of the loads in .load_pat in each coordinate direction is reported. The sums of the nodal moments are also printed. The summary only considers specified nodal loads and equivalent nodal loads from the elements. The effect of imposed displacements is not included.

See Also

form_k

lsolve

Command Syntax

lsolve [k=?] [-sparse]

Linear solution

k = 0 -> form stiffness by calling routine form_k

= 1 -> use existing, unfactored stiffness

= 2 -> use existing, factored stiffness

For an existing stiffness, the profile stiffness is expected in array .kstr and the locations of the diagonals are assumed to be in array .kdiag_loc. Loads are assumed to be in .load_pat. If the array load_comb exists, it is interpreted as load combination factors to be applied with the "load patterns" in .load_pat. The solution is placed in .disp.

If -sparse is specified, a sparse equation solver is used. If k > 0, the arrays as defined in the help for command ptospase are expected. The data for the factored sparse stiffness are in arrays .sparse_mmik, .sparse_mmim, .sparse_mmf, and .sparse_addr. These arrays can be deleted when the factored stiffness is no longer needed.

Note 1: The sparse option is recommended for all large problems because

it is much more efficient.

Note 2: To include geometric stiffness in a linear analysis, precede this command with command `form_k` and then use `k=1` or `2` here.

The profile solver modifies `.kstr`. Although the sparse solver puts the factored matrix in a copy, it does reorder the original arrays.

See Also

`form_k` `nodef` `ptosparse`

mass

Command Syntax

```
mass [-noecho]
n=node_no m=mx,my,mz,Ixx,Iyy,Izz [inc=]
```

Reads nodal masses.

If `-noecho` is specified, the data will not be echoed to the output file.

`mj` and `Ijj` are the nodal mass/inertia with respect to the `j` axis. Generation occurs if two records do not have sequential nodes and `inc`, node number increment, is greater than 0. The generated nodal data will have the same value as the record with `inc`.

Masses for repeated nodes are accumulated.

End input with a blank line.

The data are stored in the array `.mass_inp(7,*)`, in the order node #, 6 mass components.

The nodal masses are assembled into the structure `mass` by the `form_m` command.

See Also

`form_m` `pmass`

mass_summary

Command Syntax

```
mass_summary [shift=?,?,?] [-save]
```

A 6x6 "rigid body" mass matrix M is determined as follows. Six "rigid body modes" are formed in the matrix $u(\#dof,6)$, and then the 6x6 mass matrix is formed as $u^T * .mstr * u$. The default is to form the modes with respect to the origin, but if $shift=x,y,z$ is specified, they are formed with respect to the point (x,y,z) .

For an unconstrained structure, the modes are true rigid body modes. If degrees-of-freedom have been restrained to be zero, any mass associated with those degrees of freedom are not included.

If `-save` is specified, the 6x6 mass matrix is stored in `.mass_summary6x6` and the estimates of the center of mass (see below) are stored in the 3x3 matrix `.mass_summaryCG`.

Note: The `form_m` command must precede this command. Additionally, the command assumes the mass is in either diagonal or profile form.

The masses in the different directions, i.e. x-mass, y-mass, and z-mass - $M(1,1)$, $M(2,2)$ and $M(3,3)$ - may not be the same, e.g., because of different nodal boundary conditions. Three different estimates of the center of mass (\bar{x} , \bar{y} , \bar{z}) relative to the shift point are calculated based on the 3 different masses:

x-mass: sum of (nodal x-masses times x, y, z coordinates) divided by x-mass

y-mass: sum of (nodal y-masses times x, y, z coordinates) divided by y-mass

z-mass: sum of (nodal z-masses times x, y, z coordinates) divided by z-mass

Because calculations involving the center of mass can be sensitive to the number of significant figures, these values are reported to high precision to aid the user. Additionally, the option to save the results in the database for further use is provided. Arrays can be printed to higher precision than the default using the options on the `print` command or the command `xprint`.

Note: This command operates on the mass matrix, `.mstr`. Calculation of the center of gravity will be affected by any masses associated with constrained nodes. These masses will have already been transformed to the master node, and will be associated with the coordinates for that node.

See Also

```
form_m rigid_modes
```

merge_nodes

Command Syntax

```
merge_nodes [tol=?] [xtol=?] [ytol=?] [ztol=?] [-noprnt]
```

Merges the equation numbers of coincident nodes. Specifically, if the

x, y, and z coordinates of node j are within tolerance tol (default = 10^{-8}) of the coordinates of node i, $i < j$, then the equations for node j will be the same as for node i. This is carried out via the array .bcid. If node numbering arrays exist, they will be removed. Different tolerances for the coordinates may be specified via xtol, ytol, and ztol. If -noprnt is specified, the report of merged nodes will be suppressed.

See also
bcid

modal_th

Command Syntax

```
modal_th arg1 dt=? endtime=? [xi=] [begintime=] [save_velocities=?]
```

Determine the modal time history response.

arg1 is the time function (nptsx2 array)

dt = time step

endtime = simulation time

xi = constant modal damping (xi=.02 means 2%)

begintime = time to begin the simulation (default=0)

save_velocities is an integer to specify the time step increment to save the modal velocities

This command integrates the modal equations exactly, assuming that the time variation of the force varies linearly between the points in arg1. The command assumes the natural frequencies are in .omega(nmodes) and the normal modes are in .phi(neq,nmodes). It also assumes the modes have been mass-normalized. If xi is missing or zero, and an array xi(nmodes) exists, those values are used for the damping for each mode. For non-zero initial conditions, the initial displacements should be given in u_init(nmodes) and the initial velocities should be given in v_init(nmodes). If either or both of these arrays don't exist, they are assumed to be zero.

The default is to begin the simulation at time = 0. In this case, any existing time history results created by previous calls of this command are deleted. A nonzero time can be specified with begintime. In this case, if there are existing modal displacements from previous calls of this command, then the new modal displacements are appended to those results.

The modal velocities will be saved every save_velocities time steps. The modal velocities at endtime are always saved.

The load patterns are defined by the nodef command. If the array load_comb exists, it is interpreted as load combination factors to be applied with the "load patterns" in .load_pat. Otherwise, the first load pattern is used. Element loads are included if they correspond to the load pattern used.

The modal "coordinates", that is the time history modal response, are stored in .modal_disp(nmodes,nsteps), where nmodes is the number of frequencies and modes in .omega and .phi. The minimum value for nsteps = (endtime-begintime)/dt+1. However, the solution is also determined at each time for which the time function is defined (up to endtime). If the time function is not defined out to endtime, the function is continued at the last defined value out to endtime. The times are saved in .modal_times(nsteps). Note that regardless of dt, the solution is calculated at each time for which the time function has been defined. At each of these points, the next time is obtained by adding dt.

The modal velocities are saved in .modal_vel(nmodes,*), where the number of columns depends on the parameter save_velocities. The last column is the velocities at endtime. The times corresponding to saved velocities are saved in .modal_vel_times(*). Because both the modal displacements and velocities are saved, multiple modal_th commands can be used in a "restart" fashion.

See Also

```
initial_conditions lsolve nodef
```

nodal_constraint

Command Syntax

```
nodal_constraint maxnodes=?  
type=general c=n cdof=cdof r=n1,n2,... rdof=r1,r2,... factor=f1,f2,...  
type=body c=n r=n1  
type=body_trans c=n r=n1
```

Input linear kinematic constraints on the nodal degrees-of-freedom. There are three possible constraint types: general, body, and body_trans.

maxnodes is the maximum number of nodes that appears in any single constraint equation (default=4). For example, an equation for a body constraint involves 4 nodes, in which case maxnodes should be no less than 4.

type is the type of the constraint

Type general:

c is the node number that has a constrained DOF
cdof is the node-local, constrained degree-of-freedom (1-6)
r is the node numbers, nj, with independent DOFs
rdof is the degree of freedom, rj, for node nj
factor is the numerical factor, fj, in the constraint equation

Type body (rigid body constraint),

c is the node number that is constrained
r is the node number with the independent DOFs ("master" node)

Type body_trans (rigid body but only translational DOFs are constrained):

c is the node number that is constrained
r is the node number with the independent DOFs ("master" node)

End input with a blank line.

The constraint information is stored columnwise for each constraint in

```
.const_nodes(maxnodes,#constraints) = n,n1,n2,...  
.const_dofs(maxnodes,#constraints) = cdof,r1,r2,...  
.const_factor(maxnodes,#constraints) = 1,f1,f2,...
```

A constrained degree-of-freedom depends on the independent degrees-of-freedom via a constraint equation. For the general constraint, the equation is

$$\text{constrained dof} = f1*n1(r1) + f2*n2(r2) + \dots$$

in which ni(rj) represents the rj_th displacement of node ni. These displacements must be independent; i.e., they cannot be constrained by another constraint equation. For body constraints, the constraint equations are similar, but they express the rigid body constraint. The factors are calculated automatically based on the nodal coordinates.

For type=general, these values are specified directly. For type=body, the factors for each degree-of-freedom are calculated based on a rigid body constraint. Hence, each body constraint actually corresponds to 6 constraint equations. For type=body_trans, each constraint corresponds to 3 constraint equations.

The constraint information is stored columnwise for each constraint equation in

```
.const_nodes(maxnodes,#constraint_eqs) = n,n1,n2,...  
.const_dofs(maxnodes,#constraint_eqs) = cdof,r1,r2,...  
.const_factor(maxnodes,#constraint_eqs) = 1,f1,f2,...
```

This is an optional command. The command `bcid` must be processed prior to this command. The command modifies the array `.bcid(7,#nodes)`, created by the `bcid` command, such that the value 2 is inserted in `.bcid` corresponding to constrained DOFs. If a DOF has been previously constrained by the `bcid` command, to be either 0 or identical to another DOF, the constraint specified by this command is ignored. This command, if used, must be processed prior to `num_eqs` and `form_k`.

See also

```
bcid form_k num_eqs pdisp
```

nodal_disp

Command Syntax

```
nodal_disp arg [nodes=?,?] [-node#]
```

Arrange the nodal displacements in the first column of `.disp` into a rectangular array `arg(#active_nodes,6)`, where the displacements for each active node are in a row. A range of node numbers can be specified by `nodes=`. The first value is the first node number in the range, and the second value is the last number in the range. If the flag `-node#` is specified, then `arg` will have 7 columns, the first of which is the node number.

Note: Only active nodes are processed.

See Also

```
pdisp pndisp pndisp_th
```

nodal_pressure

Command Syntax

```
nodal_pressure  
n=first,last,incr pat=pat [p=pressure] [-hydro density=density] &  
[z=Z0] [-no_negative]
```

Reads nodal pressures.

`first` is the first node number in the sequence
`last` is the last node number in the sequence (blank for a single node)
`incr` is the node number increment in the sequence (default = 1)
`pat` is the load pattern to which the pressure is assigned
`pressure` is the pressure for the node (constant for the node sequence)

If `-hydro` is specified, hydrostatic pressure is generated as follows. The pressure is calculated as $-\text{density} \cdot (z - Z_0)$, where z is the z -coordinate of the node. Z_0 is the z -coordinate at which the pressure is zero. If `-no_negative` is specified, then calculated negative pressures are set to zero. (This option is active only if hydrostatic pressures are calculated.)

If `n=all` is specified instead of a node range, then it is applied to

all nodes.

End input with a blank line.

The data are stored in the array `.nodal_pressure(nodes,*)`, in which `nodes` is the largest active node number and the number of columns is the number of load patterns with pressures assigned. The integer vector `.nodal_pressure_pat(*)` is created as well; the *i*th element holds the pattern number for the *i*th column of `.nodal_pressure`. Note: the largest allowable load pattern is defined by the `nodef` command. If a larger pattern is specified by this command, the program will generate an error when the loads are formed.

See Also
nodef

nodef

Command Syntax

```
nodef [#=?]  
n=node_no p=pat f=?,?,?...
```

Reads nodal load patterns.

= number DOFs per node (default = 6)
pat is the pattern to which the load is assigned
f specifies the loads in a comma-separated list of # values

Loads for repeated nodes are accumulated.

End input with a blank line.

The data are stored in the array `.load_inp(#dofs_per_node+2,*)`, in the order node #, pattern, load components.

The maximum load pattern number specified in this command becomes the maximum load pattern number. If no nodal loads exist, but all loads come from element loads, the maximum load pattern number is still defined by this command, in which case a zero load should be specified. (If the number of load patterns equals 1, this is unnecessary.)

See Also
pnodef

nodes

Command Syntax

```
nodes #=?  
n=node_no x=x-coor y=y-coor z=z-coor [lgen=lgen]
```

Reads and generates nodal coordinates. The value specified by # is used to define storage requirements, and it must be greater than or equal to the maximum node number. If this value is missing or 0, it is assumed that existing nodes are being changed or added to, and the previous value applies. `lgen` is the node number increment for linear generation. Nodes are generated equally spaced along a straight line if two adjacent records do not have sequential node numbers and if `lgen` on the second line is not zero or blank. For other generation options, see the

command `node_gen`. Nodes need not be input in sequence.

End input with a blank line.

The coordinates are stored in array `.xyz(3,#)`, and the maximum possible node number (specified by `#`) is stored in `.#nodes_tot`.

Arrays associated with the equation numbers are deleted if `#` is specified; i.e., `.bcid`, `.node_eqs`, `.#eqs`, `.node_eq_order`, `.node_active` are deleted.

Active nodes are those that are defined explicitly either by this command or another command that creates nodes. The node number of the maximum defined node is stored in `.#nodes`. The character vector `.node_active` has an "A" for active nodes. Only active nodes can be used.

The nodes command need not be executed as long as the coordinates, which could be generated by another program, are put in the array `.xyz`, `.#nodes` and `.#nodes_tot` are set, `.node_active` is created, and any other associated arrays are deleted.

See also

`node_gen` `pnodes` `bcid`

node_gen

Command Syntax

`node_gen`

`linear=node1,node2 [inc=?] [w=?]`

`quad=node1,node2,node3,node4 [inc=inc1,inc2] [w=w1,w2]`

Generates nodal coordinates. The nodes command must precede this command.

Linear generation is specified by the identifier `linear`. Nodes are generated from `node1` to `node2` with a node increment of `inc` (default=1). The spacing of nodes is equidistant unless the positive weight `w` (default=1) is specified, in which case the spacing between node `n` and `n+1` is equal to `w` times the spacing between `n-1` and `n`.

The identifier `quad` specifies node generation within the "quadrilateral" defined by the four nodes (see sketch below). Linear generation is done between `node1` and `node2`, and between `node3` and `node4`, based on `inc1` and `w1`. Linear generation is also done between `node1` and `node3`, and between `node2` and `node4`, based on `inc2` and `w2`. Then, linear generation is done between the nodes generated from 1-3 and 2-4, based on `inc1` and `w1`. The number of interior lines generated is the same as the number of nodes generated from 1-3. If the number of nodes generated from 1-3 is larger than the number generated from 2-4, the "extra" lines will not be generated, as this would result in a redefinition of nodal coordinates along 3-4. If `node3` and `node4` are identical, the generated nodes are within a triangular domain. The nodes need not be coplanar.

End input with a blank line.

```
node3                                node4
X-----O-----O-----O-----X
```


tol is the tolerance on individual coordinates to determine to which node a particular point corresponds (default=1.e-8).

node is the node number

inc is the node increment used for generation

gen is the number of nodes to generate.

End input with a blank line.

This command determines the arithmetic average of stresses at the nodes specified. If the number of nodes on the command line is left blank, all nodes are used. If it is given, as many generation commands as necessary may be used to define the node numbers.

Nodal coordinates are expected in array .xyz. Coordinates and stresses are in array arg1 in the form coord1, ..., stresses. Which coordinates appear in this array are specified by the flags -x, -y, and -z. For example, if -x and -z are given, then the first column in arg1 contains x-coordinates, the second column contains z-coordinates, and the remaining columns are the stresses at the corresponding (x,z) coordinates. Coordinates that are not specified (y in this example) are assumed to be zero.

The node numbers for which nodal stress averages are calculated are kept in array arg3(n).

The average stresses are kept in arg2(n,*), where the number of columns is the same as in arg1. The rows of arg2 may be less than n, because nodes for which no stress values are found are eliminated from arg2 and arg3.

nsolve

Command Syntax

```
nsolve steps=? tol=?,? maxit=? [k=?] [control=?] [dlim=?,?] [gdisp=?]
      [dprin=?] [fprin=?] [state=?]
```

Nonlinear solution command

```
steps  = number of load steps
tol    = load and displacement tolerances
maxit  = maximum # of iterations within a load step
k      = # of iterations with constant stiffness (K)
        < 0 -> form tangent stiffness and use for abs(k) iter.
        = 0 -> form and use initial stiffness iter.
        > 0 -> use existing stiffness for k iter., then reform
control = 0 -> load control
        = 1 -> displacement control
dlim   = max. transl. and rotational displ. increment in iteration
        for load control
gdisp  = magnitude of the generalized displacement for displ.
        control; see command disp_cntl
state  = 1 -> do state determination in first step
-morison= determine incoming current velocities for drag

dprin  = print flag for nodal displacements
        = 0 -> not printed from nsolve (default)
        = 1 -> print at each load step
        = 2 -> print at each iteration
```

```
fprin    = print flag for element forces
          = 0 -> not printed from nsolve (default)
          = 1 -> print at each load step
          = 2 -> print at each iteration
```

A `form_k` command must precede this command.

The increment in external loads is determined from the `.load_pat` array created by the command `nodef`. If the array `load_comb` exists, it is assumed that it is a vector of load combination factors, and the load patterns are combined to form the load increment. If `load_comb` does not exist, the first load pattern is taken as the load increment. The displacements are placed in `.disp`, and a history of displacements are kept in `.disp_history`.

For additional information on the displacement control strategy, see Powell, G.H. and Simons, J., "Improved Iteration Strategy for Nonlinear Structures," *IJNME*, 17:1455-1467 (1981).

num_eqs

Command Syntax

```
num_eqs [-#eqs]
```

Number nodal equations and determine stiffness memory requirements. The element groups to be included are specified by nonzero codes in the array `.elem_grp`, which is defined during element definition. The default operation is to number the equations if the array `.node_eqs` does not exist. If the flag `-#eqs` is specified, the equations are numbered regardless.

The equations are numbered based on the restraint codes in `.bcid`, which is established by the `bcid` command and modified by the `nodal_constraint` command. The nodal equation numbers are put in `.node_eqs(#dof_per_node,#nodes)`, and the number of equations is put in the scalar `.#eqs`. The locations of the stiffness diagonals are determined based on the element connectivity and stored in `.kdiag_loc`; hence, the elements must have been defined. The size of the stiffness matrix is stored in the scalar `.kstr_size`.

This command is not normally used. The command `form_k` will number the equations by default, and that is the preferred approach. The command is provided in case a `form_k` command is not used, such as when only the mass matrix is formed. If used, it must follow the definition of all nodes and nodal boundary conditions, elements, and nodal constraints.

See Also

```
bcid elem_grp form_k nodal_constraints peqs
```

pbcid

Command Syntax

```
pbcid
Print nodal displacement restraints.
```

See also

```
bcid
```

pbody_frc2d

Command Syntax
pbody_frc2d

Print the body force coefficients defined in .body_f2d.

See Also
body_frc2d

pdisp

Command Syntax

pdisp [nodes=?,?] [-screen] [form=format] [-file] [-append]

Print nodal displacements in array .disp for all active nodes with a print code of 1. A range of node numbers can be specified by nodes=. The first value is the first node number in the range, and the second value is the last number in the range. The default is to print the displacements for all active nodes. The default is to print to the output file only; if -screen is present, the displacements will also be printed to the screen. The default format is (i5,2x,1p6e12.3). Alternative formats can be specified by the parameter form=. In this case, format must be a valid FORTRAN format, enclosed in () with a maximum length of 160 characters and without any blank spaces.

If -file is specified, the displacements will also be written to the unformatted file project_name.dis. If -append is present, this file will be appended.

See Also
bcid pndisp pndisp_th

peqns

Command Syntax

peqns
Print equation numbers.

See also
bcid form_k num_eqs

pimposed_displ

Command Syntax
pimposed_displ

Print imposed displacements.

See also
imposed_displ

pmass

Command Syntax
pmass

Print input nodal masses.

See also
mass

pndisp

Command Syntax

pndisp n=?

Print the "history," in terms of load increments and load steps, of displacements for node n. The displacements are stored in `.disp_history`, which is created by command `nsolve`.

See Also

nsolve pdisp pndisp_th

pndisp_th

Command Syntax

pndisp_th n=? arg1 arg2

Print the time history of displacements, velocities, or accelerations for node n as determined by the `direct_th` command. `arg1` is the array of displacements (or velocities, or accelerations) and `arg2` is the vector of corresponding times. For example, to print the time history of displacements for node 10 at the saved time steps the command would be

```
pndisp_th n=10 .nodal_th_disp .nodal_times
```

and to print the velocities at the restart times it would be

```
pndisp_th n=10 .th_restart_vel .th_restart_times
```

See Also

direct_th pdisp pndisp

pnodef

Command Syntax

pnodef

Print input nodal loads.

See also

nodef

pnodes

Command Syntax

pnodes [nodes=?,?] [-screen]

Print nodal coordinates of active nodes. A range of node numbers can be specified by `nodes=`. The first value is the first node number in the range, and the second value is the last number in the range. The default is to print the coordinates for all active nodes. The default is to print to output file only; if `-screen` is present, then output is to the screen as well.

See also

nodes

presponse

Command Syntax

```
presponse (or presp) [-file] [-append]
  Print element response to output file.
```

If `-file` is specified, the response will also be written to an unformatted file. If `-append` is present, this file will be appended. Not all elements support this option. Check the help for individual elements.

See Also

```
elem_grp response state
```

pstate

Command Syntax

```
pstate
  Print element state determined from state command.
```

See Also

```
elem_grp state response presponse
```

response

Command Syntax

```
response
  Calculate element response.
  The element groups for which the response is determined are specified by nonzero codes in the array .elem_grp, which is defined during element definition, and can be modified by the command elem_grp.
```

See Also

```
elem_grp state presponse
```

rigid_modes

Command Syntax

```
rigid_modes arg [cg=?,?,?,?] [rigid=?,?,?,?,?]? [node_range=first,last]
```

Generate rigid body modes relative to the "center of gravity" specified by `cg=x-coor,y-coor,z-coord`. The default is the origin. The modes are defined relative to axes that are parallel to the global coordinate axes. The default is to define six modes (surge, sway, heave, roll, pitch, yaw, in naval architecture parlance), but this can be controlled by the optional parameter `rigid`:

```
rigid    = six values, corresponding to 6 rigid body modes
          0 -> do not form corresponding rigid body mode
          1 -> form corresponding rigid body mode
```

A subset of nodes can be defined by the `node_range` option. Nodes not in this range will have zero displacement.

The modes are placed in the array specified by `arg`. The dimensions of this array are `#dofs` by `#modes` (default = 6).

This command expects the nodal coordinates in `.xyz` and the equation numbers in `.node_eqs`.

WARNING: This command has unreliable results when constrained nodes are involved and its use in that situation is not recommended. The problem is that the two constraints may not be compatible.

See Also
nodes num_eqs

state

Command Syntax
state

Calculate element state.

The element groups for which state determination is carried out are specified by nonzero codes in the array `.elem_grp`, which is defined during element definition, and can be modified by the command `elem_grp`.

See Also
elem_grp response pstate

water_waves

Command Syntax

```
water_waves  #=#waves  [h=?]  grav=?  [-current #pts=?  current_beta=?]  
n=?  ampl=?  period=?  phase=?  beta=?
```

End wave data with a blank line. If current exists, then enter current data, #pts records:

```
n=?  z=?  v=?
```

Reads wave and current data.

#waves is the total number of wave components.
h is the water depth. h=0 means deep water (default).
grav is the acceleration of gravity.

For each wave component:

n ranges from 1 to #waves
ampl is the component amplitude
period is the component period (s)
phase is the component phase angle (rad)
beta is the component wave angle (degrees)

If current, for #pts records:

n is the record number (between 1 and #pts)
z is the z-coordinate
v is the current velocity

End input with a blank line.

It is assumed that the origin of the global coordinate axes is on the free surface and that the global Z axis is directed upwards.

If there is a current, use `-current`. #pts must be at least 2. The first value must be for z=0 and the last value (negative) must be below the last node. `current_beta` is the current angle in degrees.

If there is a current but no wave, enter a wave with zero amplitude

Note: Only the first wave input is used at the present time (i.e., only a single regular wave is used).

The following arrays are created:

```
.water_wavesg(2)      -> h, grav  
.water_waves(#waves,5) -> ampl, period, phase, beta (rad), wave number  
.water_current_beta  -> current angle (rad)  
.water_current(#pts,2) -> z, velocity v
```

See Also
direct_th

2.7. Finite Element Library

beam3d

Linear, 3-D beam element

There are two Command Syntax options.

```
----- OPTION 1 -----  
Command Syntax (option 1)  
beam3d m=? n=? [-kg]  
m=matl e=emodulus g=gmodulus a=area j=jsec iy=iyy iz=izz           &  
[asy=asy asz=asz] [mbar=density mxyz=mx,my,mz]                   &  
[mI=mIxx,mIyy,mIzz] [cbar=cdense cxyz=cx,cy,cz] (1 record/matl)  &  
n=nel mat=mat nodes=node1,node2 node3=node3 [print=print]       &  
[gen=gen inc=inc inc2=inc2]                                       &  
[tension=tension tension_last=tension_last]
```

m is the number of different materials
n is the number of elements
-kg is a flag to include geometric stiffness

matl is the material number
emodulus is the modulus of elasticity
gmodulus is the shear modulus
density or mx,my,mz is the mass/unit length
area is the cross sectional area
jsec is the torsional constant
iyy,izz are area moments of inertia in local coordinates
asy, asz are the shear areas in y and z, respectively
(0 -> the corresponding shear deformation is ignored)
mIxx, mIyy, mIzz are mass moments of inertia (per unit length)
in local coordinates
cdense or cx,cy,cz is the damping/unit length

nel is the element number (ID#)
node1 and node2 are the node numbers
mat is the material number for the element
print .ne. 0, element results not printed
inc is the node 1 increment used for generation
inc2 is the node 2 increment used for generation (default=inc)
gen is the number of elements to generate
node3 lies in the local x-z plane
tension is the initial tension (for geometric stiffness only)
tension_last is the initial tension for the last element in a series

For generated elements, the initial tensions for the geometric stiffness are interpolated linearly, from tension in the first element to tension_last in the last element. If tension_last is not specified, it is set equal to tension.

There are two options to specify mass density. The usual option is to specify density with the mbar= identifier. In some special cases, it may be useful to specify a different mass for different directions of the local axes. In that case, use the identifier mxyz= to specify the three values separately. If both mbar= and mxyz= are specified, mxyz= will be ignored if density is a nonnegative value. The element creates a lumped, diagonal mass matrix in local coordinates. However, if the mxyz are not equal, and/or the mI are not equal, then the mass

matrix when transformed to global coordinates may not be diagonal.

There are two options to specify damping, which is analogous to specifying mass and the formulation is similar. The usual option is to specify `cdense` with the `cbar=` identifier. It may be useful to specify a different damping for different directions of the local axes. In that case, use the identifier `cxyz=` to specify the three values separately. If both `cbar=` and `cxyz=` are specified, `cxyz=` will be ignored if `cdense` is a nonnegative value. The element creates a lumped, diagonal damping matrix in local coordinates. However, if the `cxyz` are not equal, then the damping matrix when transformed to global coordinates may not be diagonal.

End input with a blank line.

----- OPTION 2 -----

Command Syntax (option 2)

```
beam3d -cylinder mat=mat node3=node3 [print=print] &
       p1=x1,y1,z1 p2=x2,y2,z2 R=R1,R2 CxL=Cseg,Lseg [face=face] &
       tension=tension_first tension_last=tension_last]
```

Option 2 generates a cylindrical mesh interface elements around a 'spine' of beam elements.

`p1` are the center coordinates of the cylinder start
`p2` are the center coordinates of the cylinder end
`R1,R2` are the radii at the start and end, respectively
`Cseg` are the number of interface elements around the circumference
`Lseg` are the number of interface elements along the length

All stiffness and mass properties are modeled by the 'spine' of beam elements along the center of the cylinder. The nodes to the interface elements are constrained via rigid body constraints to the nodes of the spine. For this option, the material must have been previously defined by an option 1 command, even if no elements were specified.

Interface nodes and elements are numbered around the circumference and then down the length. The interface elements are defined such that the local x-axis is down the length of the cylinder and the positive z-face is on the inside of the cylinder. Nodes are numbered clockwise looking from the outside, i.e., looking at the -1 face. See the interface element for details.

There is one cylinder per command line. To generate multiple cylinders, use multiple commands.

----- ALL OPTIONS -----

The local (principal) axes of the beam are defined as follows:

The local x-axis is directed from node1 to node2
The local y-axis = (x-axis) X (vector from node1 to node3)
The local z-axis = (x-axis) X (y-axis)

If node3 is -1, -2, or -3, then the "vector to node3" is a unit vector in the direction of the negative X, Y, or Z global axes, respectively.

On input, created arrays are:

```
.beam3d_mp(m,19) -> emodulus, gmodulus, unused, area, jsect,
                  iyy, izz, asy, asz, mIxx, mIyy, mIzz, mx, my, mz
```

```

                                cdense, cx, cy, cz
.beam3d_el(6,n)  -> node1, node2, material, print code, node3, ID#
.beam3d_len(n)  -> element length
.beam3d_st(12,n) -> Axial Force, Vy, Vz, Torque, My, Mz at node1
                  Axial Force, Vy, Vz, Torque, My, Mz at node2
.beam3d_kg      -> 0 or 1; w/o or w/ geometric stiffness

```

This element calculates a lumped mass matrix in local coordinates.

For state calculation, element forces in local coordinates are put in .beam3d_st. Positive forces follow the right hand rule, not "beam" sign convention.

For response calculation, element does nothing.

For state output, results in .beam3d_st are printed, but using beam sign convention for shear and moment; torque at nodej is positive in the local x-axis.

For response output, no results are printed.

The element can be used with nonlinear elements, but the element response will be linear.

See Also

interface pbeam3d pstate presponse

pbeam3d

Command Syntax

```

pbeam3d
  Print beam3d element data

```

See Also

beam3d

biot1d234

1-D element for consolidation of a linear, elastic medium

Implementation assumes element is directed along the positive X-axis.

Command Syntax

```

biot1d234 m=? n=? [disp=?] [pressure=?]
m=mat1 e=emodulus a=area kx=kx gammaw=gw gammas=gs (1/mat1)
n=nel nodes=node1,node2 mat=mat [print=print] [inc=inc gen=gen]

```

m is the number of different materials

n is the number of elements

mat1 is the material number

disp is linear (default), quad, or cubic (variation of displacement)

pressure is linear (default), quad, or cubic (variation of pressure)

emodulus is the uniaxial strain modulus of elasticity

area is the cross sectional area

kx is the soil permeability

gw is the weight density of water

gs is the effective weight density of the soil

nel is the element number

node1 and node2 are end node numbers
mat is the material number for the element
print .ne. 0, element results not printed
inc is the node increment used for generation
gen is the number of elements to generate

End input with a blank line.

Whether the elements have linear, quadratic, or cubic displacement and/or pressure variation, only the two end nodes are specified. The variation for displacements is specified by disp, and the variation for the pressure is specified by pressure. For quadratic and cubic elements, the interior nodes are generated automatically. E.g., disp=cubic pressure=quad would mean an element would have 4 displacement nodes and 3 pressure nodes. For a quadratic variation, node 3 is placed in the center of the element. For a cubic variation, nodes 3 and 4 are placed at the third points. If a physical node already exists at this location, then that node is used. If a node does not exist, then a new node is created. Therefore, be sure to specify a sufficient number of nodes in the nodes command to include generated nodes.

Prior to forming the stiffness of this element, the time step must be defined in the variable .biot_dt and the integration factor theta (2-step family from forward Euler (theta = 0) to backward Euler (theta = 1) must be defined in .biot_theta. Note: if the boundary conditions involve specified nonzero flow, then use theta = 1; otherwise, errors will result.

Regardless of the number of nodes, 3-pt Gauss integration is used to calculate the stiffness. This scheme is exact for all combinations of displacement and pressure variation.

On input, created arrays are:

```
.biot1d234_mp(m,5) -> modulus, area, kx, gw, gs  
.biot1d234_el(10,n) -> nodes, material #, print code, #disp_nodes,  
                    # pressure_nodes  
.biot1d234_len(n)  -> element length  
.biot1d234_st(n,15)-> coordinate, stress, excess pore pressure, flow,  
and liquifaction ratio for each gauss point
```

For stiffness calculation, 3-pt Gauss integration is used.

For state calculation, global coordinate, effective stress, excess pore pressure, flow and liquifaction ratio at each integration point are put in .biot1d234_st.

For response calculation, global coordinate, displ., effective stress, excess pore pressure, and flow are calculated for local coordinates in biot1d234_lc. Results are put in .biot1d234_resp(*,5).

For state output, results in .biot1d234_st are printed.

For response output, results in .biot1d234_resp are printed.

See Also

pbiot1d234 pstate presponse

pbiot1d234

Command Syntax

pbiot1d234

Print biot1d234 element data

See Also

biot1d234

biot2d3to9

2-D element for consolidation of a linear, elastic medium

Implementation assumes element is in the X-Z plane, with gravity acting in the negative Z direction and the medium's surface is at Z=0.

Command Syntax

```
biot2d3to9 m=? n=? [type=?] [gauss=?]
m=mat# e=emodulus nu=nu kx=kx kz=kz &
      gammaw=gw gammas=gs (1 record/mat1)
n=nel nodes=node1,node2,...,node9 mat=mat [print=print] &
      [gauss=gauss] [inc=inc1,inc2,inc3 gen=gen] &
      [inc_2d=inc1_2d,inc2_2d,inc3_2d gen_2d=gen_2d inc_el=inc_el] &
      [nodesp=nodep1,nodep2,...,nodep9]
```

m is the number of different materials

n is the number of elements

type is the element type

2 -> plane strain (default)

3 -> axisymmetric

gauss is the order of Gauss integration for the stiffness matrices

1 -> 1 x 1

2 -> 2 x 2

3 -> 3 x 3 (default)

...

10 -> 10 x 10

mat# is the material number

emodulus is the modulus of elasticity

nu is the Poisson ratio

kx and kz are the coefficients of permeability

gw is the weight density of water

gs is the effective weight density of the soil

nel is the element number

node1 thru node9 are node numbers (3 to 9 nodes)

mat is the material number for the element

print .ne. 0 -> element results not printed

gauss overrides the previously specified integration order

inc1, inc2, inc3 are node increments in a "linear sequence"

gen is the number of elements to generate in a sequence

inc1_2d, inc2_2d, inc3_2d are node increments between sequences

gen_2d is the number of linear sequences to generate

inc_el is the element increment between sequences

nodep1 thru nodep9 are the nodes with pressure DOFS; if these are the same as the displacement nodes, they are not given. That is, the default is nodep1 = node1, etc.

Nodes 1 to 4 are the corner nodes for quad elements and are specified counterclockwise. Nodes 5 to 8 are the midnodes on the edges (see

sketch below), while node 9 is the center node. For triangular elements, only the first three nodes are to be specified. The same holds true for the pressure nodes.

A "linear sequence" of elements can be generated by specifying `incl1`, `inc2`, `inc3`, and `gen`. In a linear sequence, nodes 1, 2, and 5 are incremented by `incl1`; nodes 6, 8, and 9 are incremented by `inc2`; and nodes 3, 4, and 7 are incremented by `inc3`. `gen` is the number of elements to generate, so a sequence will have `gen+1` elements. To generate a 2D patch of elements, multiple sequences can be specified; `incl1_2d`, `inc2_2d`, and `inc3_2d` are used to increment the node numbers from one sequence to the next. `gen_2d` is the number of additional sequences. The element numbers in two successive sequences differ by `inc_el` (default = `numgen+1`).

End input with a blank line.

On input, created arrays are:

```
.biot2d3to9_et(1)  -> 2 or 3 for plane strain or axisymmetric
.biot2d3to9_mp(m,7) -> emodulus, Poisson ratio, thickness,
                    kx, kz, gw, gs
.biot2d3to9_el(21,n) -> node1 - node9, nodep1 - nodep9, material #,
                    print code, # gauss pts
```

This element uses a value of 1 for the thickness.

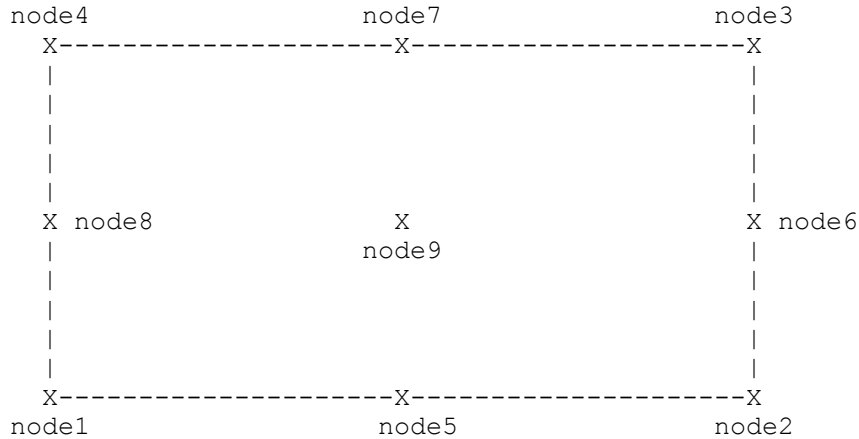
Prior to forming the stiffness of this element, the time step must be defined in the variable `.biot_dt` and the integration factor `theta` (2-step family from forward Euler (`theta = 0`) to backward Euler (`theta = 1`)) must be defined in `.biot_theta`. Note: if the boundary conditions involve specified nonzero flow, then use `theta = 1`; otherwise, errors will result.

For state calculation, X-Z coordinates, effective stress, excess pore pressure, flow, and liquefaction ratio at each Gauss point are put in `.biot2d3to9_st(n,10*gauss^2)`. The stresses are in the global coordinate system and are stored in order `Sxx`, `Syy`, `Sxy`, `Szz` for each point. Similarly, the flow is a vector with X and Z components.

For response calculation, the same quantities as for state are calculated, but this time for local coordinates in `biot2d3to9_lc(#pts,2)`. Results are put in `.biot2d3to9_resp(#pts*n,10)`. If instead global coordinates are given in `biot2d3to9_gc(#pts,2)`, then results for those points are put in `.biot2d3to9_resp(#pts,10)`. The vector `.biot2d3to9_index(#pts)` maps the data points to the element in which it falls.

For state output, results in `.biot2d3to9_st` are printed.

For response output, results in `.biot2d3to9_resp` are printed.



See Also
 pbiot2d3to9 presponse pstate

pbiot2d3to9

Command Syntax
 pbiot2d3to9
 Print biot2d3to9 element data

See Also
 biot2d3to9

cable

Elastic catenary cable element

Command Syntax
 cable m=? n=? [maxiter=?] [tol=?] [-restart]
 m=seg_prop e=emodulus a=area w=wx,wy,wz (m records)
 n=nel nodes=node1,node2 [#segs=#segs] [tension=tenX,tenY,tenZ]
 seg=seg mat=seg_prop L=length (#segs records)

m is the number of different cable properties
 n is the number of elements
 maxiter is the max. # of iterations on the tension (default=30)
 tol is the relative tolerance on the end point position (default=1.e-5)

If -restart is specified, then the data from the database is used to initialize the element. This option must be used if a database file containing cable data is read.

For each set of cable properties:
 seg_prop is the segment property number
 emodulus is the modulus of elasticity
 area is the cross sectional area
 wx,wy,wz are the weight/unit length components in global coordinates

For each element:
 nel is the element number
 nodes are the two nodes of the element
 #segs is the number of different segments (default=1)

tension is the initial estimate of the tension

seg is the segment number

seg_prop is the segment property number

length is the unstretched segment length

The element is based on small strain elastic catenary theory. A shooting method is used to solve the two-point boundary value problem. Specifically, iteration on the tension at end 1 is carried out until the distance between end 2 and node 2, divided by the element length, is less than or equal to the tolerance (tol). For information on the formulation, see H.R. Riggs and T. Leraand, "Efficient Static Analysis and Design of Flexible Risers," J. Off. Mech. Arctic Engrg., Vol. 113, pp. 235-240, 1991, and H.R. Riggs and T. Leraand, "A Robust Element for Static Analysis of Marine Cables," Proc. Third International Offshore and Polar Engineering Conference, Singapore, Vol. 2, pp. 357-363, 1993. The element described in those papers includes fluid drag; this element does not.

See Also

pcable

pcable

command Syntax

pcable

Print cable element data

See Also

cable

contact_spring

Nonlinear, contact spring element

Command Syntax

contact_spring m=? n=?

m=mat k=Kx,Ky,Kz

(1 record/material)

n=nel node=node mat=mat [print=print] [dist=distance] [dir=lx,ly,lz]

m is the number of different materials

n is the number of elements

mat is the material number

Kx is the spring stiffness in the local x-direction

Ky is the spring stiffness in the local y-direction

Kz is the spring stiffness in the local z-direction

nel is the element number

node is the node number to which the contact spring is attached

distance is the distance from the node to 'ground'

mat is the material number

print .ne. 0 -> element printout suppressed

lx, ly, lz element orientation vector (see below)

End input with a blank line.

The contact spring element can be assigned to a node that may contact rigid 'ground'. The distance in the local x-direction from the original

position of the node to ground is defined by dist. The local x-direction is defined by the orientation vector, which is directed from ground to the node. The default values for this direction vector are dir=0,0,1. In this case, the node is dist "above" ground in the global Z direction. (At present, only dir=0,0,1 is supported.) Once a node contacts ground, the spring stiffness specified by Kx, Ky, and Kz is introduced. The stiffness Kx tries to keep the node from going below ground. Ky and Kz prevent slipping along the ground (a plane normal to the direction vector). Clearly, Kx, Ky, and Kz function as penalty parameters, and therefore they should be relatively large. The element deformations are defined as the 'distance' below ground of the node, and the amount of slip along the normal plane.

On input, created arrays are:

```
.contactspring_mp(m,3)  -> Kx, Ky, Kz
.contactspring_el(n,3)  -> node, material, print
.contactspring_dr(3,n)  -> lx, ly, lz
.contactspring_dist(n)  -> distance
.contactspring_st(5,n)  -> element deformations
```

For state calculation, element deformations are put in .contactspring_st.

For response calculation, element does nothing.

For state output, results in .spring_st are printed.

For response output, results in .spring_st are printed for those springs that are in contact. If the unformatted write options on the presponse command are specified, the results for all elements are written to the file project_name.cspr. The data are written: element #, node, indentation, slip-1, slip-2, displ-1 at slip, and displ-2 at slip.

See Also

pcontact_spring pstate presponse

pcontact_spring

Command Syntax

```
pcontact_spring
  Print contact spring element data
```

See Also

contact_spring

d11234

1-D, linear, 2,3, or 4 node, isoparametric "rod" element
Implementation assumes element is parallel to X-axis.

Command Syntax

```
d11234 m=? n=?
m=mat1 e=emodulus a=area [mbar=density] [k=kdsp] (1 record/mat1)
n=nel nodes=node1,node2,node3,node4 mat=mat [print=print] &
[inc=inc gen=gen]
```

m is the number of different materials

n is the number of elements

mat1 is the material number
emodulus is the modulus of elasticity
area is the cross sectional area
density is density/unit length (unused)
kdsp is a distributed spring stiffness along length of member

nel is the element number
node1 thru node4 are node numbers (2,3 or 4 nodes)
mat is the material number for the element
print .ne. 0, element results not printed
inc is the node increment used for generation
gen is the number of elements to generate

End input with a blank line.

On input, created arrays are:

.d11234_mp(m,4) -> modulus, area, dens, ksp
.d11234_el(n,7) -> node1 - node4, material, print code, #nodes
.d11234_len(n) -> element length
.d11234_st(n,8) -> coordinate and force for each gauss point

For stiffness calculation, exact integration is used.

For state calculation, coordinates and stresses are put in .d11234_st.

For response calculation, global coordinate, displ., and force are calculated for local coordinates in d11234_lc. Results are put in .d11234_resp(*,3).

For state output, results in .d11234_st are printed.

For response output, results in .d11234_resp are printed.

See Also

pd11234 pstate presponse

pd11234

Command Syntax

pd11234
Print d11234 element data

See Also

d11234

d11234v2

1-D, linear, 2,3, or 4 node, isoparametric "rod" element, version 2
Implementation assumes element is parallel to X-axis.

Command Syntax

d11234v2 m=? n=? [-linear] [-quad] [-cubic]
m=mat1 e=emodulus a=area [mbar=density] [k=kdsp] (1 record/mat1)
n=nel nodes=node1,node2 mat=mat [print=print] [inc=inc gen=gen]

m is the number of different materials
n is the number of elements

-linear indicates a 2-node, linear displ element
-quad indicates a 3-node, quadratic displ element
-cubic indicates a 4-node, cubic displ element
mat1 is the material number
emodulus is the modulus of elasticity
area is the cross sectional area
density is density/unit length (unused)
kdsp is a distributed spring stiffness along length of member

nel is the element number
node1 and node2 are the two end node numbers
mat is the material number for the element
print .ne. 0, element results not printed
inc is the node increment used for generation
gen is the number of elements to generate

End input with a blank line.

Whether the elements have linear, quadratic, or cubic displacement variation, only the two end nodes are specified. For quadratic and cubic elements, the interior nodes are generated automatically. For quadratic elements, node 3 is placed in the center of the element. For cubic elements, nodes 3 and 4 are placed at the third points. If a physical node already exists at this location, then that node is used. If a node does not exist, then a new node is created. Therefore, be sure to specify a sufficient number of nodes in the nodes command to include generated nodes. The automatic generation of interior nodes is the main difference between this element and d11234 (in addition to extensive code changes).

On input, created arrays are:

.d11234_mp(m,4) -> modulus, area, dens, ksp
.d11234_el(7,n) -> node1 - node4, material, print code, #nodes
.d11234_len(n) -> element length
.d11234_st(n,8) -> coordinate and force for each gauss point

For stiffness calculation, exact integration is used.

For state calculation, coordinates and stresses are put in .d11234_st.

For response calculation, global coordinate, displ., and force are calculated for local coordinates in d11234_lc. Results are put in .d11234_resp(*,3).

For state output, results in .d11234_st are printed.

For response output, results in .d11234_resp are printed.

See Also

pd11234v2 nodes pstate presponse

pd11234v2

Command Syntax

pd11234v2

Print d11234v2 element data

See Also

d11234v2

d213to9

2-D, linear, 3 to 9 node, isoparametric element for 2-D elasticity.
Implementation assumes element is in the X-Y plane.

Command Syntax

```
d213to9 m=? n=? [type=?] [gauss=?]
m=mat# e=emodulus [nu=nu t=thickness [mass=mass] &
pat=pat bx=distr_x by=distr_y [gaussf=gaussf] (1 record/mat1)
n=nel nodes=node1,node2,...,node 9 mat=mat [print=print] &
[gauss=gauss] [inc=inc1,inc2,inc3 gen=gen] &
[inc_2d=inc1_2d,inc2_2d,inc3_2d gen_2d=gen_2d inc_el=inc_el]
```

m is the number of different materials

n is the number of elements

type is the element type

1 -> plane stress (default)

2 -> plane strain

3 -> axisymmetric

gauss is the order of Gauss integration for stiffness and mass

1 -> 1 x 1

2 -> 2 x 2

3 -> 3 x 3 (default)

...

10 -> 10 x 10

mat# is the material number

emodulus is the modulus of elasticity

nu is the Poisson ratio

thickness is the element thickness

mass is the mass density (per unit volume)

pat is the load pattern number for the body forces

distr_x and distr_y specify distributions for body forces (per unit volume) in the X-Y coordinates. The distributions are defined by the body_frc2d command.

gaussf is the integration order for the body forces (default = 3).

nel is the element number

node1 thru node9 are node numbers (3 to 9 nodes)

mat is the material number for the element

print .ne. 0 -> element results not printed

gauss overrides the previously specified integration order

inc1, inc2, inc3 are node increments in a "linear sequence"

gen is the number of elements to generate in a sequence

inc1_2d, inc2_2d, inc3_2d are node increments between sequences

gen_2d is the number of linear sequences to generate

inc_el is the element increment between sequences

Nodes 1 to 4 are the corner nodes for quad elements and are specified counterclockwise. Nodes 5 to 8 are the midnodes on the edges (see sketch below), while node 9 is the center node. For triangular elements, only the first three nodes are to be specified.

A "linear sequence" of elements can be generated by specifying inc1, inc2, inc3, and gen. In a linear sequence, nodes 1, 2, and 5 are incremented by inc1; nodes 6, 8, and 9 are incremented by inc2; and nodes 3, 4, and 7 are incremented by inc3. gen is the number of

elements to generate, so a sequence will have gen+1 elements. To generate a 2D patch of elements, multiple sequences can be specified; inc1_2d, inc2_2d, and inc3_2d are used to increment the node numbers from one sequence to the next. gen_2d is the number of additional sequences. The element numbers in two successive sequences differ by inc_el (default = numgen+1).

End input with a blank line.

On input, created arrays are:

```
.d2l3to9_et(1)    -> 1,2,3 for plane stress, strain, or axisym.  
.d2l3to9_mp(m,8) -> emodulus, Poisson ratio, thickness, mass,  
                  load pattern, bx, by, gaussf  
.d2l3to9_el(12,n) -> node1 - node9, material #, print code, # gauss pts
```

This element forms a consistent mass matrix.

For state calculation, X-Y coordinates and stress at each Gauss point are put in .d2l3to9_st(n,6*gauss^2). The stresses are in the global coordinate system and are stored in order Sxx, Syy, Sxy, Szz for each point.

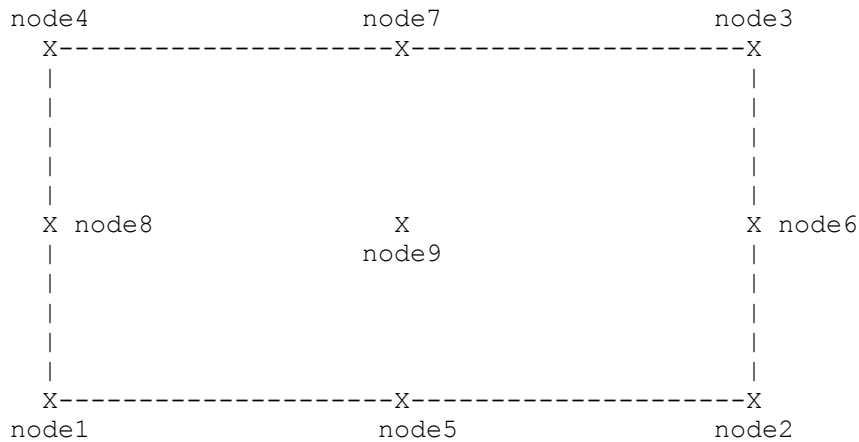
For response calculation, global coordinates and stresses are calculated for local coordinates in d2l3to9_lc(#pts,2). Results are put in .d2l3to9_resp(#pts*n,6). See the explanation for state calculation for the order of stresses. If instead global coordinates are given in d2l3to9_gc(#pts,2), then results for those points are put in .d2l3to9_resp(#pts,6). The vector .d2l3to9_index(#pts) maps the data points to the element in which it falls.

For state output, results in .d2l3to9_st are printed.

For response output, results in .d2l3to9_resp are printed.

For error estimation, the strain-energy option and the "user" option are supported. With strain-energy based error estimation, the "exact" stresses are expected in .d2l3to9_xst(n,6*#pts) in the same format as .d2l3to9_st; these values may be the result of some smoothing procedure. The finite element stresses are expected in .d2l3to9_st(n,6*#pts). The number of integration points used for the element error is based on the number of columns of these matrices. If the same integration points used for the stiffness calculation are used, .d2l3to9_st can be generated by the state command; otherwise, it can be generated by the response command and rearranged by the unwrap command. The X,Y coordinates and error of each integration point are put in .d2l3to9_err(n,3*#pts). The integrated element error and relative error (element error/global error) are put in .d2l3to9_elerr(n,2).

The "user" error estimation option functions similarly, except that the error function is expected in .d2l3to9_ruser(n,3*#pts), where #pts is the number of integration points and the first two columns contain the global coordinates. This option creates the same arrays as the strain-energy option. Because .d2l3to9_ruser can contain the values of any function, this option can be used for the numerical evaluation of an integral over a two-dimensional domain. For example, if .d2l3to9_ruser contains all ones, then the element and total volumes will be calculated.



See Also
gauss_pts pd213to9 presponse pstate unwrap

pd213to9

Command Syntax
pd213to9
Print d213to9 element data

See Also
d213to9

d21tri

2-D, linear, 3 to 6 node, isoparametric triangular element for 2-D elasticity.

Implementation assumes element is in the X-Y plane.

Command Syntax

```
d21tri m=? n=? [type=?] [intpt=?] [tol=?]
m=mat# e=emodulus nu=nu t=thickness [mass=mass] &
pat=pat bx=distr_x by=distr_y [intf=intf] (1 record/mat1)
n=nel nodes=node1,node2,...,node 6 mat=mat [print=print] &
[intpt=intpt] [inc=inc1,inc2,inc3 gen=gen] &
[inc_2d=inc1_2d,inc2_2d,inc3_2d gen_2d=gen_2d inc_el=inc_el]
```

m is the number of different materials

n is the number of elements

type is the element type

1 -> plane stress (default)

2 -> plane strain

3 -> axisymmetric

intpt is the order of integration

1 -> 1 pt. integration

3 -> 3 pt. integration (default)

4 -> 4 pt. integration

7 -> 7 pt. integration

9 -> 9 pt. integration

tol is a tolerance on nodal coordinates (see below)

mat# is the material number

emodulus is the modulus of elasticity
 nu is the Poisson ratio
 thickness is the element thickness
 mass is the mass density (per unit volume)
 pat is the load pattern number for the body forces
 distr_x and distr_y specify distributions for body forces (per unit volume) in the X-Y coordinates. The distributions are defined by the body_frc2d command
 intf is the integration order for the body forces and mass(default = 4)

nel is the element number
 node1 thru node6 are node numbers (3 or 6 nodes)
 mat is the material number for the element
 print .ne. 0 -> element results not printed
 intpt overrides the previously specified integration order
 incl1, inc2, inc3 are node increments in a "linear sequence"
 gen is the number of elements to generate in a sequence
 incl_2d, inc2_2d, inc3_2d are node increments between sequences
 gen_2d is the number of linear sequences to generate
 inc_el is the element increment between sequences

Nodes 1 to 3 are the corner nodes for the elements and are specified counterclockwise. Nodes 4 to 6 are the midnodes on the edges (see sketch below). If a negative value is input for a midside node, then the coordinates of the node are calculated midway between the corresponding vertex nodes. If a node does not exist at that location (within tolerance of tol), a node will be generated with a node number one greater than the previous maximum defined node number. In this case, the restraint codes for the node of absolute value of the number specified will be used for the new node (e.g., if -10 is specified, then restraint codes for node 10 will be used). Normally, it will be convenient to use the negative of one of the vertex nodes.

When specifying intf to calculate the mass matrix, one should be aware that with the 6-node element and 3-pt integration, the integration points are at the midside nodes. This leads to a diagonal mass matrix with zeroes for the vertex nodes. An intf of 1, 4, or higher will avoid this.

A "linear sequence" of elements can be generated by specifying incl1, inc2, inc3, and gen. In a linear sequence, nodes 1, 2, and 4 are incremented by incl1; nodes 5 and 6 are incremented by inc2; and node 3 is incremented by inc3. gen is the number of elements to generate, so a sequence will have gen+1 elements. Multiple sequences can be specified; incl_2d, inc2_2d, and inc3_2d are used to increment the node numbers from one sequence to the next. gen_2d is the number of additional sequences. The element numbers in two successive sequences differ by inc_el (default = numgen+1). Note: Node increments are added to positive node numbers and subtracted from negative node numbers.

End input with a blank line.

On input, created arrays are:

```

.d2ltri_et(1) -> 1, 2, 3 for plane stress, strain, or axisym.
.d2ltri_mp(m,8) -> emodulus, Poisson ratio, thickness, mass,      &
                    load pattern, bx, by, intf
.d2ltri_el(n,9) -> node1 - node6, material #, print code, #integ. pts
  
```

This element forms a consistent mass matrix.

For state calculation, X,Y coordinates and stress (Sx, Sy, Sxy, Sz) at each integration point are put in .d2ltri_st(n,6*#pts).

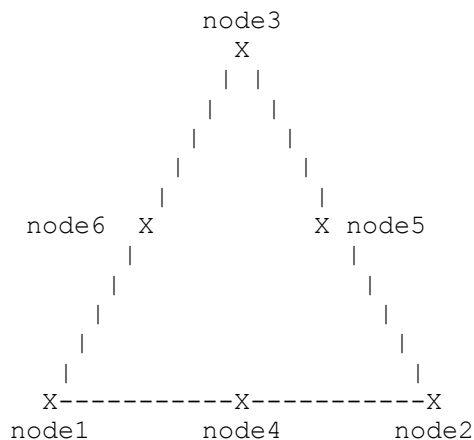
For response calculation, X,Y coordinates and stresses are calculated for area coordinates (3 values/point) in d2ltri_lc(#pts,3). Results are put in .d2ltri_resp(n*#pts,6).

For state output, results in .d2ltri_st are printed.

For response output, results in .d2ltri_resp are printed.

For error estimation, the strain-energy option and the "user" option are supported. With strain-energy based error estimation, the "exact" stresses are expected in .d2ltri_xst(n,6*#pts), i.e., in the same form as .d2ltri_st; these values may be the result of some smoothing procedure. The finite element stresses are expected in .d2ltri_st(n,6*#pts). The number of integration points used for the element error is based on the number of columns of these matrices. If the same integration points used for the stiffness calculation are used, .d2ltri_st can be generated by the state command; otherwise, it can be generated by the response command and rearranged by the unwrap command. The X,Y coordinates and error of each integration point are put in .d2ltri_err(n,3*#pts). The integrated element error and relative error (element_error/global error) are put in .d2ltri_elerr(n,2).

The "user" error estimation option functions similarly, except that the error function is expected in .d2ltri_ruser(n,3*#pts), where #pts is the number of integration points and the first two columns contain the global coordinates. This option creates the same arrays as the strain-energy option. Because .d2ltri_ruser can contain the values of any function, this option can be used for the numerical evaluation of an integral over a two-dimensional domain. For example, if .d2ltri_ruser contains all ones, then the element and total volumes will be calculated.



See Also

pd2ltri presponse pstate tri_intpts unwrap

pd2ltri

Command Syntax

```
pd2ltri
  Print d2ltri element data
```

SeeAlso

d2ltri

iFEM2D

2-D, 3 to 9 node, isoparametric element for 2-D nonlinear iFEM.

Implementation assumes element is in the X-Y plane.

Command Syntax

```
iFEM2D n=? [gauss=?,?]
n=nel nodes=node1,node2,...,node 9 print=print inc=inc1,inc2,inc3 &
gen=gen inc_2d=inc1_2d,inc2_2d,inc3_2d gen_2d=gen_2d &
inc_el=inc_el
```

n is the number of elements

gauss is the order of Gauss integration for stiffness and "loads"

n,m -> n x m in the xi and eta directions (default = 4x4)

nel is the element number

node1 thru node9 are node numbers (3 to 9 nodes)

print .ne. 0 -> element results not printed

inc1, inc2, inc3 are node increments in a "linear sequence"

gen is the number of elements to generate in a sequence

inc1_2d, inc2_2d, inc3_2d are node increments between sequences

gen_2d is the number of linear sequences to generate

inc_el is the element increment between sequences

Nodes 1 to 4 are the corner nodes for quad elements and are specified counterclockwise. Nodes 5 to 8 are the midnodes on the edges (see sketch below), while node 9 is the center node. For triangular elements, only the first three nodes are to be specified. Note: this element is meant to be used with 6 nodes, with quadratic interpolation in the longitudinal direction and linear interpolation in the transverse direction.

A "linear sequence" of elements can be generated by specifying inc1, inc2, inc3, and gen. In a linear sequence, nodes 1, 2, and 5 are incremented by inc1; nodes 6, 8, and 9 are incremented by inc2; and nodes 3, 4, and 7 are incremented by inc3. gen is the number of elements to generate, so a sequence will have gen+1 elements. To generate a 2D patch of elements, multiple sequences can be specified; inc1_2d, inc2_2d, and inc3_2d are used to increment the node numbers from one sequence to the next. gen_2d is the number of additional sequences. The element numbers in two successive sequences differ by inc_el (default = numgen+1).

End input with a blank line.

On input, created arrays are:

```
.ifem2d_int(2) -> # gauss pts in xi and eta directions
.ifem2d_el(10,n) -> node1 - node9, print code
```

For state calculation, X-Y coordinates, X-Y displacements, and strain at

each Gauss point are put in `.ifem2d_st(n,7*gauss_xi*gauss_eta)`. The strains are in the global coordinate system and are stored in order `Exx`, `Eyy`, `Exy` for each point.

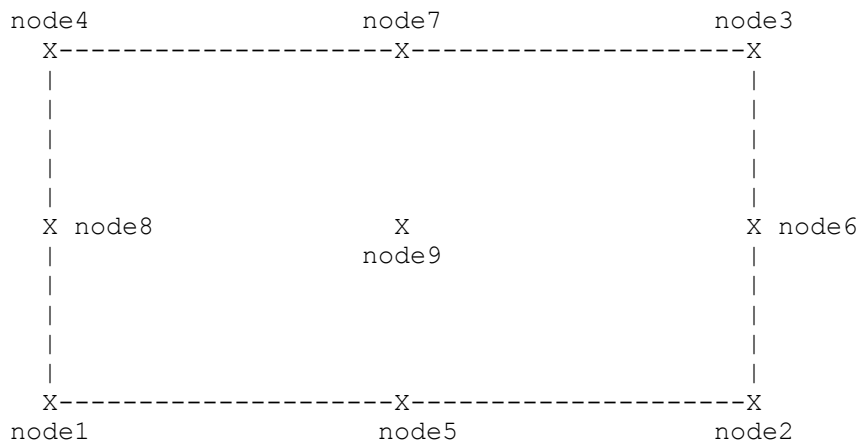
For response calculation, global coordinates and strains are calculated for local coordinates in `ifem2d_lc(#pts,2)`. Results are put in `.ifem2d_resp(#pts*n,7)`. See the explanation for state calculation for the data and order of strains. If instead global coordinates are given in `ifem2d_gc(#pts,2)`, then results for those points are put in `.ifem2d_resp(#pts,7)`. The vector `.ifem2d_index(#pts)` maps the data points to the element in which it falls.

For state output, results in `.ifem2d_st` are printed.

For response output, results in `.ifem2d_resp` are printed.

No error estimation feature is supported.

See Paczkowski, K., Riggs, H.R., 2007, "An inverse finite element strategy to recover full-field, large displacements from strain measurements," Proc., 26th International Offshore Mechanics and Arctic Engineering Conference, paper OMAE2007-29730 for details on this element.



See Also
`gauss_pts` `piFEM2D` `presponse` `pstate`

piFEM2D

Command Syntax
`piFEM2D`
Print iFEM2D element data

See Also
`iFEM2D`

interface

Quadrilateral (and triangular) interface element

Command Syntax (option 1)

```

interface  n=?
n=nel  nodes=node1,node2,node3,node4  [print=print]  [face=face]      &
      [gen=gen inc=inc1,inc2]                                           &
      [gen_2d=gen_2d  inc_2d=inc1_2d,inc2_2d  inc_el=inc_el]

```

n is the maximum element number specified

nel is the element number

node1 thru node4 are node numbers

mat is the material number for the element

print .ne. 0 -> element results not printed

inc1, inc2 are node increments in a "linear sequence"

gen is the number of elements to generate in a sequence

inc1_2d, inc2_2d are node increments between sequences

gen_2d is the number of linear sequences to generate

inc_el is the element increment between sequences

face specifies the face on which the pressure acts

= 0 -> no pressure acts ("dry" element) - default

= -1 -> pressure acts on negative face (-z)

= 1 -> pressure acts on positive face (+z)

Command Syntax (option 2)

```
interface -subdivide  range=?,?  nxm=?,?
```

Option 2 subdivides previously defined elements

range specifies a range of element ID numbers; all elements
in the range are divided

nxm specifies how many elements to subdivide each element into.

For example, nxm=2,3 would subdivide each element into 6
elements; 2 in the 1-2 direction and 3 in the 1-4 direction.

Nodes 1 to 4 are the corner nodes for quadrilateral elements, specified
in a counterclockwise order when looking at the positive face. For
triangular elements, if node 4 is not specified it will be set equal to
node 3.

A "linear sequence" of elements can be generated by specifying inc1,
inc2, and gen. In a linear sequence, nodes 1 and 2 are incremented by
inc1; nodes 3 and 4 are incremented by inc2. gen is the number of
elements to generate, so a sequence will have gen+1 elements. Multiple
sequences can be specified; inc1_2d and inc2_2d are used to increment
the node numbers from one sequence to the next. gen_2d is the number of
additional sequences. The element numbers in two successive sequences
differ by inc_el (default = numgen+1).

The local x-axis (axis-1) is directed from node 1 to node 2. The local
y-axis (axis-2) lies in the plane defined by nodes 1-2-4, is normal to
the x-axis, and is directed "toward" node 4. The local z-axis (axis-3)
follows from the right-hand-rule.

The normal pressures act in the local z-direction.

End input with a blank line.

On input, created arrays are:

```

.interface_el(8,n) -> node1 - node4, print code, face, ID#
.interface_pressure -> 1 if -pressure specified, otherwise 0

```

The columns of `.interface_el` are based on the number of defined elements, not the maximum ID# (i.e., ID numbers don't have to be sequential).

This element does very little. It has no stiffness or mass. The following is an example of one possible use. For visualization purposes, it can be used to represent the surface of a rigid body with the nodes of this element kinematically constrained to the CG of the rigid body so that the displacements are kinematically constrained. The face parameter is included to allow one side to be distinguished from the other (for example, as 'wet').

pinterface

Command Syntax

```
pinterface
  Print interface element data
```

See Also

interface

isomin6

Linear, 6-node, 18 DOF triangular, isoparametric Mindlin plate element. Implementation assumes element is in the X-Y plane.

Command Syntax

```
isomin6 m=? n=? [q=q_vec]
m=mat# e=E1,E2,E3 g=G12,G23,G13 nu=nu12,nu23,nu13          &
  t=thickness mass=mass iso=iso (1 record/mat1)
n=nel nodes=node1,node2,node3,node4,node5,node6 mat=mat print=print &
  inc=inc1,inc2,inc3 gen=gen inc_2d=inc1_2d,inc2_2d,inc3_2d &
  gen_2d=gen_2d inc_el=inc_el pat=pat [q=q1,q2,q3,q4,q5,q6]
```

m is the number of different materials

n is the number of elements

q_vec is the name of a vector in the database in which the ith element is the normal pressure at node i

mat# is the material number

E1, E2, E3 are the moduli of elasticity

G12, G23, G13 are the shear moduli

nu12, nu23, nu13 are the Poisson ratios

thickness is the element thickness

mass is the mass per unit volume

iso = 0 -> isotropic material (default)

 = 1 -> orthotropic material

nel is the element number

node1 thru node6 are node numbers

mat is the material number for the element

print .ne. 0 -> element results not printed

inc1 and inc2 are node increments in a "linear sequence"

gen is the number of elements to generate in a sequence

inc1, inc2 inc3 are node increments in a "linear sequence"

gen is the number of elements to generate in a sequence

inc1_2d, inc2_2d, inc3_2d are node increments between sequences

gen_2d is the number of linear sequences to generate

inc_el is the element increment between sequences

pat is the load pattern number for the normal pressures
q1,q2,q3,q4,q5,q6 are normal pressures for nodes (these values override those defined by q_vec, if any.)

Nodes 1 to 3 are the corner nodes and are specified counterclockwise.
Nodes 4 to 6 are the midnodes.

A "linear sequence" of elements can be generated by specifying inc1, inc2, inc3, and gen. In a linear sequence, nodes 1, 2, and 4 are incremented by inc1; nodes 5, 6 are incremented by inc2; and nodes 3 is incremented by inc3. gen is the number of elements to generate, so a sequence will have gen+1 elements. To generate a 2D patch of elements, multiple sequences can be specified; inc1_2d, inc2_2d, and inc3_2d are used to increment the node numbers from one sequence to the next. gen_2d is the number of additional sequences. The element numbers in two successive sequences differ by inc_el (default = numgen+1).

End input with a blank line.

The element is restricted to lie in an x-y plane (z=constant). For an orthotropic material, the material parameters are specified in the global coordinate system.

The normal pressures act in the z-direction.

On input, created arrays are:

```
.isomin6_mp(m,14) -> Ei, Gij, nuij, thickness, mass, unused, unused,  
iso  
.isomin6_el(8,n) -> nodel - node6, material #, print code  
.isomin6_q(7,n) -> pat, q1, q2, q3, q4, q5, q6
```

If a nonzero mass density is specified, uniform (gravitational) body forces are applied if the vector gravity(4) has been defined. The 4 components of gravity are: load pattern number, gx, gy, and gz, where gi is the gravitational acceleration in the global i direction.

For state calculation, global coordinates and bending stress resultants are stored in .isomin6_stb(n, 4*8) in the order x, y, z, Mx, My, Mxy, Qx, and Qy. These values are calculated at the 4 Gauss points of the triangle.

The element does not calculate the equivalent nodal forces in equilibrium with its stress state, and therefore cannot be used in a nonlinear analysis.

The response option has not been implemented.

The error estimation option has not been implemented.

For state output, the stress resultants in .isomin6_stb are printed.

See Also

pisomin6 min3s min5s min6 pstate

pisomin6

Command Syntax
pmin6

Print isomin6 element data

See Also
isomin6

min3s

Mindlin 3-D triangular, linear shell element.
Membrane and shear relaxations are included in the implementation.

Command Syntax

```
min3s m=? n=? [-pressure] [-kg] [-kf]
m=mat# e=E1,E2,E3 g=G12,G23,G13 nu=nu12,nu23,nu13 t=thickness &
      [tb=tbending] [mass=mass] [global=global] &
      [C_s=relxs] [C_m=relxm] [drill=drill_stiff] &
      [fdensity=fdensity] (1 record/mat1)
n=nel mat=mat nodes=node1,node2,...,node 6 [print=print] &
      [face=face] [gen=gen inc=inc1,inc2,inc3] &
      [gen_2d=gen_2d inc_2d=inc1_2d,inc2_2d,inc3_2d inc_el=inc_el
```

m is the number of different materials

n is the number of elements

-pressure is a flag to apply nodal pressures in the array
.nodal_pressure

-kg is a flag to include geometric stiffness

-kf is a flag to include hydrostatic stiffness (includes geometric
stiffness)

mat# is the material number

E1, E2, E3 are the moduli of elasticity

G12, G23, G13 are the shear moduli

nu12, nu23, nu13 are the Poisson ratios

thickness is the element thickness (see below)

tbending is the element bending thickness (default=thickness)

mass is the mass per unit volume

global = 0 -> calc. stress resultants in local coordinates (default)

= 1 -> calc. stress resultants in global coordinates

relxs is the shear relaxation factor (default = 0.5)

relxm is the membrane relaxation factor (default = 1.0)

drill_stiff is an artificial drilling stiffness (default = 1.e-5)

fdensity is the fluid weight density to calculate the hydrostatic
stiffness

Note: Defaults are obtained for the above factors not by specifying
0.0, but by omitting the input for the specific factors or by
specifying a negative value.

nel is the element number

node1 thru node6 are node numbers

mat is the material number for the element

print .ne. 0 -> element results not printed

inc1, inc2, inc3 are node increments in a "linear sequence"

gen is the number of elements to generate in a sequence

inc1_2d, inc2_2d, inc3_2d are node increments between sequences

gen_2d is the number of linear sequences to generate

inc_el is the element increment between sequences

face specifies the face on which the pressure acts

= 0 -> no pressure acts ("dry" element) - default

= -1 -> pressure acts on negative face (-z)

= 1 -> pressure acts on positive face (+z)

Nodes 1 to 3 are the vertex nodes and are specified counterclockwise. Nodes 4 to 6 are the midnodes on the edges, which are used to define the initial curvature of the element, and do not have any degree of freedom. If the element is "flat", then nodes 4 to 6 may be specified as zero.

A "linear sequence" of elements can be generated by specifying `inc1`, `inc2`, `inc3`, and `gen`. In a linear sequence, nodes 1, 2, and 4 are incremented by `inc1`; nodes 5 and 6 are incremented by `inc2`; and node 3 is incremented by `inc3`. `gen` is the number of elements to generate, so a sequence will have `gen+1` elements. Multiple sequences can be specified; `inc1_2d`, `inc2_2d`, and `inc3_2d` are used to increment the node numbers from one sequence to the next. `gen_2d` is the number of additional sequences. The element numbers in two successive sequences differ by `inc_el` (default = `numgen+1`).

The local x-axis (1) is directed from node 1 to node 2. The local y-axis (2) lies in the plane defined by the vertex nodes, is normal to the x-axis, and is directed "toward" node 3. The local z-axis (3) follows from the right-hand-rule.

The material parameters are specified in the local coordinate system. `thickness` is used for the membrane and shear stiffnesses, while `tbending` is used for the bending stiffness. If `tbending` is not input, it will be set equal to `thickness`. `thickness` is also used to determine the mass (from the mass density). The element has no drilling dof stiffness, and so an artificial stiffness is added to these dofs that is equal to `drill_stiff` times the minimum of the element diagonal stiffnesses.

The normal pressures act in the local z-direction.

End input with a blank line.

On input, created arrays are:

```
.min3s_mp(m,17) -> Ei, Gij, nuij, thickness, mass, global, C_s, C_m,
                  tbending, drill_stiff, fdensity
.min3s_el(9,n)  -> nodel - node6, material #, print code, face
.min3s_pressure -> 1 if -pressure specified, otherwise 0
```

For stiffness calculation, the shear and membrane relaxation factors are stored in `.min3s_rlx(2,n)` for later use in calculating the element state.

If a nonzero mass density is specified, uniform (gravitational) body forces are applied if the vector `gravity(4)` has been defined. The 4 components of `gravity` are: load pattern number, `gx`, `gy`, and `gz`, where `gi` is the gravitational acceleration in the global `i` direction.

If hydrostatic stiffness is to be calculated, gravity is assumed to act in the negative global `z` direction. The hydrostatic stiffness is only calculated for "wet" elements, i.e., with `face = +-1`. The flag `-pressure` must be specified as well. The hydrostatic pressure is assumed to be in the first column of `.nodal_pressure`, as specified by the command `nodal_pressure`. In this case, the load pattern number should be 1 for this pressure. If multiple load patterns are used, they should all have the same pressures specified.

For state calculation, global coordinates and the stress resultants in the local coordinate system are put in `.min3s_st(n,11)` in the order `x, y, z, Nx, Ny, Nxy, Mx, My, Mxy, Qx, and Qy`. These values are calculated at the element centroid. If the value of `global` on the material card is specified to be 1, then the resultants in global coordinates are calculated instead. (This option gives correct results only if the element is in the global X-Y plane.) The element does not calculate the equivalent nodal forces in equilibrium with its stress state, and therefore cannot be used in a nonlinear analysis.

For response calculation, global coordinates and local stress resultants are calculated for local coordinates in `min3s_lc(#pts,3)`. Results are put in `.min3s_resp(#pts*n,11)`. See the explanation for state calculation for the order and for the option for resultants in global coordinates.

For state output, stress resultants in `.min3s_st` are printed.

For response output, coordinates and stress resultants in `.min3s_resp` are printed.

For error estimation, the strain-energy option and the "user" option are supported. With strain-energy based error estimation, the "exact" stress resultants are expected in `.min3s_xst(n,11*#pts)`, i.e., in the same form as `.min3s_st`; these values may be the result of some smoothing procedure. The finite element resultants are expected in `.min3s_st(n,11*#pts)`. The number of integration points used for the element error is based on the number of columns of these matrices. If one-point integration is used, `.min3s_st` can be generated by the state command. The X,Y,Z coordinates and error of each integration point are put in `.min3s_err(n,4*#pts)`. The integrated element error and relative error (element error/global error) are put in `.min3s_elerr(n,2)`.

The "user" error estimation option functions similarly, except that the error function is expected in `.min3s_ruser(n,4*#pts)`, where `#pts` is the number of integration points and the first three columns contain the global coordinates. This option creates the same arrays as the strain-energy option. Because `.min3s_ruser` can contain the values of any function, this option can be used for the numerical evaluation of an integral over a two-dimensional domain. For example, if `.min3s_ruser` contains all ones, then the element and total volumes will be calculated.

For the theory of this element, see Tessler, A., "A C0 anisoparametric three-node shallow shell element," *Computer Methods in Applied Mechanics and Engineering*, v. 78, 1990, pp. 89-103.

The basic element has been provided courtesy of Dr. Alex Tessler, Computational Mechanics Branch, NASA Langley Research Center, Hampton, VA. For the hydrostatic stiffness formulation, see Huang, L.L. and Riggs, H.R., "The hydrostatic stiffness of flexible floating structures for linear hydroelasticity," *Marine Structures*, v. 13, 2000, pp. 91-106.

See Also

`nodal_pressure` `pmin3s` `pstate`

pmin3s

Command Syntax

pmin3s
Print min3s element data

See Also
min3s

min5s

Mindlin 3-D quadrilateral, linear shell element consisting of 4 min3s (triangular) elements. Membrane and shear relaxations are included in the implementation. There are three Command Syntax options.

```
----- OPTION 1 -----
min5s m=? n=? [-pressure] [-kg] [-kf]
m=mat# e=E1,E2,E3 g=G12,G23,G13 nu=nu12,nu23,nu13 &
      t=thickness [tb=tbending] [mass=mass] [local=local] &
      [C_s=relxs] [C_m=relxm] [drill=drill_stiff] &
      [iso=iso] [fdensity=fdensity] (1 record/mat1)
n=nel nodes=node1,node2,...,node5 mat=mat [print=print] [face=face] &
      [gen=gen inc=inc1,inc2,inc3] &
      [gen_2d=gen_2d inc_2d=inc1_2d,inc2_2d,inc3_2d inc_el=inc_el]
```

m is the number of different materials
n is the maximum element number specified
-pressure is a flag to apply nodal pressures in the array
.nodal_pressure
-kg is a flag to include geometric stiffness
-kf is a flag to include hydrostatic stiffness (includes geometric stiffness)

mat# is the material number
E1, E2, E3 are the moduli of elasticity
G12, G23, G13 are the shear moduli
nu12, nu23, nu13 are the Poisson ratios
thickness is the element thickness (see below)
tbending is the element bending thickness (default=thickness)
mass is the mass per unit volume
local = 0 -> calculate stress resultants in quad coordinates (default)
 = 1 -> calculate stress resultants in triangle coordinates
 Note: local must be 1 for -kg and -kf
relxs is the shear relaxation factor (default = 0.5)
relxm is the membrane relaxation factor (default = 1.0)
iso = 0 -> isotropic material (default)
 = 1 -> orthotropic material
drill_stiff is the drilling dof stiffness factor (default=1.e-5)
fdensity is the fluid weight density to calculate the hydrostatic stiffness

Note 1: For an isotropic material, the values E1, G12, and nu12 are used.

Note 2: Defaults are obtained for the above factors not by specifying 0.0, but by omitting the input for the specific factors or by specifying a negative value.

nel is the element number
node1 thru node5 are node numbers
mat is the material number for the element
print .ne. 0 -> element results not printed

inc1, inc2, inc3 are node increments in a "linear sequence"
gen is the number of elements to generate in a sequence
inc1_2d, inc2_2d, inc3_2d are node increments between sequences
gen_2d is the number of linear sequences to generate
inc_el is the element increment between sequences
face specifies the face on which the pressure acts
= 0 -> no pressure acts ("dry" element) - default
= -1 -> pressure acts on negative face (-z)
= 1 -> pressure acts on positive face (+z)

----- OPTION 2 -----

Command Syntax (option 2)

```
min5s -subdivide range=?,? nxm=?,?
```

Option 2 subdivides previously defined elements
range specifies a range of element ID numbers; all elements
in the range are divided
nxm specifies how many elements to subdivide each element into.
For example, nxm=2,3 would subdivide each element into 6
elements; 2 in the 1-2 direction and 3 in the 1-4 direction.

----- OPTION 3 -----

Command Syntax (option 3)

```
min5s -cylinder n=nel mat=mat [print=print] [face=face]      &
      p1=x1,y1,z1 p2=x2,y2,z2 R=R1,R2 CxL=Cseg,Lseg         &
      [-ring_stiffeners LRseg=LRseg ring_mat=ring_mat       &
      ring_node3=ring_node3]                                 &
      [-spine spine_mat=spine_mat spine_node3=spine_node3   &
      tension_first=tension_first tension_last=tension_last]
```

Option 3 generates a cylindrical mesh (min5s must have been initialized
previously with option 1, even if 0 elements were defined)
nel is the first element number (of a sequential sequence)
mat, print and face have the same meaning as in option 1
p1 are the center coordinates of the cylinder start
p2 are the center coordinates of the cylinder end
R1,R2 are the radii at the start and end, respectively
Cseg are the number of elements around the circumference
Lseg are the number of elements along the length

If -ring_stiffeners is specified, generate circumferential beam
stiffeners:

LRseg is the number of segments along the length separated by
stiffeners
ring_mat is the beam material number (define by the beam3d command)
ring_node3 (see beam3d for definition of node3)

If ring stiffeners exist, Lseg must be an integer multiple of LRseg.
If ring_node3 is blank, the node3 direction is along the length of
the cylinder on the surface

If -spine is specified, the cylinder is modeled by beam elements along
the center of the cylinder. The nodes to the shell elements are
constrained via rigid body constraints to the nodes of the spine. The
mass and stiffness of the combined structure can be specified either by
the material parameters on the shell elements or the beam elements. Be
careful not to include the properties twice. Specifically, if the mass
and/or stiffness properties are specified by the beam elements, then

the shell elements should have zero mass and very small (but not zero) values for modulus and thickness (and vice versa). For this option,

spine_mat is the beam material number (defined by the beam3d command)
spine_node3 corresponds to node3 for the beam (for a vertical cylinder -1 is usually convenient)
tension_first and tension_last, see beam3d

Nodes and elements are numbered around the circumference and then down the length. The shell elements are defined such that the local x-axis is down the length of the cylinder, in the direction from p1 to p2, that is, the line from node1 to node2 is parallel to the axis and in the direction of p2. Nodes are numbered clockwise looking from the outside.

----- ALL OPTIONS -----

The 5-node min5s element is formed by four 3-node triangular (min3s) elements. Nodes 1 to 4 define the quadrilateral and are specified counterclockwise. Node 5 is the "interior" node, which is common to the four triangles. The connectivity of the triangles in terms of the quadrilateral nodes is 1-2-5, 2-3-5, 3-4-5, and 4-1-5. Normally, node 5 is not specified, in which case it is located at the intersection of the diagonals (straight lines connecting nodes 1 and 3 and nodes 2 and 4). If a node does not exist at this location, one is created. The restraint conditions are the same as for node "1" of the element, if the restraints have already been defined. Note that this "cross-diagonal" pattern is the preferred meshing strategy. Although nodes 1 to 4 are not forced to be coplanar, the element is meant to be used as a flat shell element.

A "linear sequence" of elements can be generated by specifying incl, inc2, inc3, and gen. In a linear sequence, nodes 1 and 2 are incremented by incl; nodes 3 and 4 are incremented by inc2; and node 5, if specified, is incremented by inc3. gen is the number of elements to generate, so a sequence will have gen+1 elements. Multiple sequences can be specified; incl_2d, inc2_2d, and inc3_2d are used to increment the node numbers from one sequence to the next. gen_2d is the number of additional sequences. The element numbers in two successive sequences differ by inc_el (default = numgen+1).

Each triangle in the quadrilateral has its own "triangle" coordinate system. For each triangle, defined by nodes 1-2-3, the x-axis (axis-1) is directed from node 1 to node 2. The local y-axis (axis-2) lies in the plane defined by the vertex nodes, is normal to the x-axis, and is directed "toward" node 3. The local z-axis (axis-3) follows from the right-hand-rule. The quad coordinate system is the same as the triangle coordinates for triangle 1.

The material parameters are specified in the quad coordinate system. thickness is used for the membrane and shear stiffnesses, while tbending is used for the bending stiffness. If tbending is not input, it will be set equal to thickness. thickness is also used to determine the mass (from the mass density).

The normal pressures act in the local z-direction.

End input with a blank line.

On input, created arrays are:

```
.min5s_mp(m,18) -> Ei, Gij, nuij, thickness, mass, local, C_s, C_m,  
                  iso, tbending, drill_stiff, fdensity  
.min5s_el(9,n)  -> node1 - node5, material #, print code, face, ID#  
.min5s_pressure -> 1 if -pressure specified, otherwise 0
```

The columns of `.min5s_el` are based on the number of defined elements, not the maximum ID# (i.e., ID numbers don't have to be sequential).

For stiffness calculation, the shear and membrane relaxation factors are stored in `.min5s_rlx(8,n)` for later use in calculating the element state.

If a nonzero mass density is specified, uniform (gravitational) body forces are applied if the vector `gravity(4)` has been defined. The 4 components of gravity are: load pattern number, `gx`, `gy`, and `gz`, where `gi` is the gravitational acceleration in the global `i` direction.

The hydrostatic stiffness is only calculated for "wet" elements, i.e., with `face = 1`. The flag `-pressure` must be specified as well. The hydrostatic pressure is assumed to be in the first column of `.nodal_pressure`, as specified by the command `nodal_pressure`. In this case, the load pattern number should be 1 for this pressure. If multiple load patterns are used, they should all have the same pressures specified.

For state calculation, global coordinates and the stress resultants in the quad coordinate system are put in `.min5s_st(4*n,11)` in the order `x`, `y`, `z`, `Nx`, `Ny`, `Nxy`, `Mx`, `My`, `Mxy`, `Qx`, and `Qy`. These values are calculated at the element centroid. If the value of `local` on the material card is specified to be 1, then the resultants in triangle coordinates are calculated instead. The element does not calculate the equivalent nodal forces in equilibrium with its stress state, and therefore cannot be used in a nonlinear analysis.

The optimal stress resultants are stored in `.min5s_op(n,11)` in the same order as for `.min5s_st`. For a parallelogram, which consists of 4 constant stress elements in a cross diagonal pattern, the optimal stress at the intersection of the diagonals is the simple average of the stresses in the 4 triangles. For general quadrilaterals, this is an approximation. It is also an approximation for the shears, `Qx` and `Qy`, because they vary linearly within each triangle.

For response calculation, global coordinates and stress resultants are calculated for local coordinates in `min5s_lc(#pts,3)`. Results are put in `.min5s_resp(#pts*4*n,11)`. See the explanation for state calculation for the order and for the option for resultants in triangle coordinates.

For state output, stress resultants in `.min5s_st` are printed.

For response output, coordinates and stress resultants in `.min5s_resp` are printed.

For error estimation, the strain-energy option and the "user" option are supported. With strain-energy based error estimation, the "exact" stress resultants are expected in `.min5s_xst(4*n,11*#pts)`, i.e., in the same form as `.min5s_st`; these values may be the result of some smoothing procedure. The finite element resultants are expected in `.min5s_st(4*n,11*#pts)`. The number of integration points used for the

element error is based on the number of columns of these matrices. If one-point integration is used, `.min5s_st` can be generated by the state command. The X,Y,Z coordinates and error of each integration point are put in `.min5s_err(4*n,4*#pts)`. The integrated element error, relative error (element error/global error), and error density (element error/element area) are put in `.min5s_elerr(n,3)`.

The "user" error estimation option functions similarly, except that the error function is expected in `.min5s_ruser(4*n,4*#pts)`, where `#pts` is the number of integration points and the first three columns contain the global coordinates. This option creates the same arrays as the strain-energy option. Because `.min5s_ruser` can contain the values of any function, this option can be used for the numerical evaluation of an integral over a two-dimensional domain. For example, if `.min5s_ruser` contains all ones, then the element and total volumes will be calculated.

Note: If nodes 3 and 4 are equal, then the element degenerates to the `min3s` element. In general it is better to use `min3s` elements for triangles, but this option is included for convenience. However, the optimal stresses are not calculated correctly for triangular elements, and the error estimation is not implemented for them.

For the theory of this element, see Tessler, A., "A C0 Anisoparametric Three-Node Shallow Shell Element," *Computer Methods in Applied Mechanics and Engineering*, v. 78, 1990, pp. 89-103. For the hydrostatic stiffness formulation, see Huang, L.L. and Riggs, H.R., "The hydrostatic stiffness of flexible floating structures for linear hydroelasticity," *Marine Structures*, v. 13, 2000, pp. 91-106.

See Also

`min3s` `nodal_pressure` `pmin5s` `prresponse` `pstate`

pmin5s

Command Syntax

```
pmin5s
  Print min5s element data
```

See Also

`min5s`

min4t

Quadrilateral, linear, Mindlin shell element. Consists of 4 `min3s` (triangular) elements with the interior node kinematically constrained to the 4 vertex nodes. Shear relaxation is included. (Element `min5s` is recommended over this element.)

Command Syntax

```
min4t m=? n=? [q=q_vec] [version=version#] [constraint=constraint#]
m=mat# e=E1,E2,E3 g=G12,G23,G13 nu=nu12,nu23,nu13
  t=thickness [mass=mass] [local=local] [gamma=] [C_s=C_s] &
  [iso=iso] (1 record/mat1)
n=nel nodes=node1,node2,node3,node4 mat=mat [print=print] &
  [pat=pat q=q1,q2,q3,q4] [inc=inc1,inc2 gen=gen]
  [inc_2d=inc1_2d,inc2_2d gen_2d=gen_2d inc_el=inc_el]
```

`m` is the number of different materials

n is the number of elements
q_vec is the name of a vector in the database for which the ith element is the normal pressure at node i

version# controls application of shear relaxation & constraints
version# = 1 -> relax each triangle, then apply constraints (default)
 2 -> apply the constraints on the 4 triangles
 and relax the quad (not implemented)
 3 -> apply the constraints on the 4 triangles and
 relax each triangle (in quad DOFs) separately
constraint# specifies the constraints for the 2 theta DOFs:
 = 1 -> use least squares (default)
 = 2 -> use 2 constraints only
 = 3 -> use interior constraints only

mat# is the material number
E1, E2, E3 are the moduli of elasticity
G12, G23, G13 are the shear moduli
nu12, nu23, nu13 are the Poisson ratios
thickness is the element thickness
mass is the mass per unit volume
local = 0 -> stress resultants in quad coordinates (default)
 = 1 -> stress resultants in triangle coordinates
gamma is the penalty parameter for the membrane drilling DOFs
 default value is $G_{12} * 10^{-4}$
C_s is the shear relaxation factor (default = 0.6)
iso = 0 -> isotropic material
 = 1 -> orthotropic material.

nel is the element number
node1 thru node4 are node numbers
mat is the material number for the element
print .ne. 0 -> element results not printed
incl1 and inc2 are node increments in a "linear sequence"
gen is the number of elements to generate in a sequence
incl_2d and inc2_2d are node increments between sequences
gen_2d is the number of linear sequences to generate
inc_el is the element increment between sequences
pat is the load pattern number for the normal pressures
q1,q2,q3,q4 are normal pressures for nodes (these values override those defined by q_vec, if any.)

The 4 node min4t element is formed by four 3 node triangular elements. Nodes 1 to 4 define the quadrilateral. "Node 5" is an internal "virtual" node and is common to the four triangles. The connectivity of the triangles in terms of the quadrilateral nodes is 1-2-5, 2-3-5, 3-4-5, and 4-1-5. Node 5 is located at the intersection of the diagonals (straight lines connecting nodes 1 and 3 and nodes 2 and 4). Although nodes 1 to 4 are not forced to be coplanar, the element is meant to be used as a flat shell element.

A "linear sequence" of elements can be generated by specifying incl1, inc2, and gen. In a linear sequence, nodes 1 and 2 are incremented by incl1; and nodes 3 and 4 are incremented by inc2. gen is the number of elements to generate; hence, a sequence will have gen+1 elements. Multiple sequences can be specified; incl_2d and inc2_2d are used to increment the node numbers from one sequence to the next. gen_2d is the number of additional sequences. The element numbers in two successive

sequences differ by `inc_el` (default = `numgen+1`).

For each triangle, the local x-axis (1) is directed from node 1 to node 2. The local y-axis (2) lies in the plane defined by the vertex nodes, is normal to the x-axis, and is directed "toward" node 3. The local z-axis (3) follows from the right-hand-rule. The "quad" coordinate system is the same as local coordinate system for triangle 1.

For an orthotropic material, the parameters are specified in the quad coordinate system.

The normal pressures act in the local z-direction.

End input with a blank line.

On input, created arrays are:

```
.min4t_mp(m,14) -> Ei, Gij, nuij, thickness, mass, local, C_s, iso
.min4t_el(6,n) -> node1 - node4, material #, print code
.min4t_node5(3,n) -> x, y, z coordinates of "node 5"
.min4t_q(5,n) -> pat, q1, q2, q3, q4
.min4t_ver(1) -> version#
.min4t_const(1) -> constraint#
```

During stiffness calculation, the shear relaxation factors are stored in `.min4t_rlx(4*n)` for later use in calculating the element state.

If a nonzero mass density is specified, uniform (gravitational) body forces are applied if the vector gravity(4) has been defined. The 4 components of gravity are: load pattern number, `gx`, `gy`, and `gz`, where `gi` is the gravitational acceleration in the global `i` direction.

For state calculation, global coordinates and bending stress resultants are stored in `.min4t_stb(5*n,8)` in the order `x`, `y`, `z`, `Mx`, `My`, `Mxy`, `Qx`, and `Qy`. These values are calculated at the triangle centroids. Each 5th row is reserved for "optimal" stresses, which are the average of the 4 triangle centroidal values and are located at the cross diagonal. If the value of local for the material is 1, then the bending resultants are calculated in the local triangle coordinate systems. The global coordinates and the membrane stress resultants at the 2x2 Gauss points are stored in `.min4t_stm(n,24)` in the order `x`, `y`, `z`, `Nx`, `Ny`, `Nxy`. The membrane resultants are always in quad coordinates.

The element does not calculate the equivalent nodal forces in equilibrium with its stress state, and therefore cannot be used in a nonlinear analysis.

The response option has not been implemented.

The error estimation option has not been implemented.

For state output, the stress resultants in `.min4t_stb` and `.min4t_stm` are printed.

For the theory of this element, see Liu, J., Riggs, H.R., and Tessler, A. (2000). "A Four-Node, Shear-Deformable Shell Element Developed via Explicit Kirchhoff Constraints." *International Journal for Numerical Methods in Engineering*, 49(8), 1065-1086.

See Also

min3s min5s pmin4t pstate

pmin4t

Command Syntax

```
pmin4t
  Print min4t element data
```

See Also

min4t

min6

6-node, 18-DOF, linear, triangular Mindlin plate bending element.
Implementation assumes element is in the X-Y plane.

Command Syntax

```
min6 m=? n=? [q=q_vec]
m=mat# e=E1,E2,E3 g=G12,G23,G13 nu=nu12,nu23,nu13      &
      t=thickness [mass=mass] [local=local] [C_s=C_s]
      [iso=iso] (1 record/mat1)
n=nel mat=mat nodes=node1,node2,node3,node4,node5,node6  &
      [print=print] [inc=inc1,inc2,inc3 gen=gen]        &
      [inc_2d=inc1_2d,inc2_2d,inc3_2d gen_2d=gen_2d    &
      inc_el=inc_el] [pat=pat q=q1,q2,q3,q4,q5,q6]
```

m is the number of different materials

n is the number of elements

q_vec is the name of a vector in the database for which the ith element
is the normal pressure at node i

mat# is the material number

E1, E2, E3 are the moduli of elasticity

G12, G23, G13 are the shear moduli

nu12, nu23, nu13 are the Poisson ratios

thickness is the element thickness

mass is the mass per unit volume

local = 0 -> stress resultants in quad coordinates (default)

C_s is the shear relaxation factor (default = 0.0)

iso = 0 -> isotropic material (default)

= 1 -> orthotropic material

nel is the element number

node1 thru node6 are node numbers

mat is the material number for the element

print .ne. 0 -> element results not printed

inc1, inc2 inc3 are node increments in a "linear sequence"

gen is the number of elements to generate in a sequence

inc1_2d, inc2_2d, inc3_2d are node increments between sequences

gen_2d is the number of linear sequences to generate

inc_el is the element increment between sequences

pat is the load pattern number for the normal pressures

q1,q2,q3,q4,q5, q6 are normal pressures for nodes (these values
override those defined by q_vec, if any.)

Nodes 1 to 3 are the corner nodes and are specified 3 node triangular

elements. Nodes 1 to 3 define the counterclockwise. Nodes 4 to 6 are the midnodes.

A "linear sequence" of elements can be generated by specifying `incl1`, `inc2`, `inc3`, and `gen`. In a linear sequence, nodes 1, 2, and 4 are incremented by `incl1`; nodes 5, 6 are incremented by `inc2`; and nodes 3 is incremented by `inc3`. `gen` is the number of elements to generate, so a sequence will have `gen+1` elements. To generate a 2D patch of elements, multiple sequences can be specified; `incl1_2d`, `inc2_2d`, and `inc3_2d` are used to increment the node numbers from one sequence to the next. `gen_2d` is the number of additional sequences. The element numbers in two successive sequences differ by `inc_el` (default = `numgen+1`).

End input with a blank line.

The global coordinate system is the same as local coordinate system for triangle.

For an orthotropic material, the parameters are specified in the global coordinate system.

The normal pressures act in the local z-direction.

On input, created arrays are:

```
.min6_mp(m,14) -> Ei, Gij, nuij, thickness, mass, local, Cs, iso  
.min6_el(6,n) -> node1 - node6, material #, print code  
.min6_q(7,n) -> pat, q1, q2, q3, q4, q5, q6
```

During stiffness calculation, the shear relaxation factors are stored in `.min6_rlx(4*n)` for later use in calculating the element state.

If a nonzero mass density is specified, uniform (gravitational) body forces are applied if the vector `gravity(4)` has been defined. The 4 components of gravity are: load pattern number, `gx`, `gy`, and `gz`, where `gi` is the gravitational acceleration in the global `i` direction.

For state calculation, global coordinates and bending stress resultants are stored in `.min6_stb(n, 4*8)` in the order `x`, `y`, `z`, `Mx`, `My`, `Mxy`, `Qx`, and `Qy`. These values are calculated at the 4 Gauss points of the triangle.

The element does not calculate the equivalent nodal forces in equilibrium with its stress state, and therefore cannot be used in a nonlinear analysis.

For response calculation, global coordinates and bending stress resultants at the nodes are stored in `.min6_respb(n,6*8)` in the order `x`, `y`, `z`, `Mx`, `My`, `Mxy`, `Qx`, and `Qy`.

The error estimation option has not been implemented.

For state output, the stress resultants in `.min6_stb` are printed.

For response output, the stress resultants in `.min6_respb` are printed.

MIN6 is an anisoparametric Mindlin plate bending element with a cubic variation of transverse displacement and quadratic variation for rotational displacements.

See Also

min3s min5s min4t pmin6 presponse pstate

pmin6

Command Syntax

pmin6
Print min6 element data

See Also

min6

nbeam2d

Large displacement, elastic 2D beam element

```
nbeam2d m=? n=? [nl=?] [-xy] [-xz] [-yz]
m=matl e=emodulus g=gmodulus [mbar=density] [mxy=mx,my] [mI=mI] &
[wbar=wbar] a=area i=moi [as=as] &
[Cm=cmx,cmy] [Cd=cdx,cdy] (1 record/material)
n=nel mat=mat nodes=node1,node2 [print=print] [gen=gen inc=inc] &
[tension=tension] [-xy] [-xz] [-yz]
```

m is the number of different materials
n is the number of elements
nl = 1 -> linear stiffness
= 2 -> nonlinear and geometric stiffness (default)
-xy or -xz or -yz specifies the plane in global coordinates

matl is the material number
emodulus is the modulus of elasticity
gmodulus is the shear modulus
density or mx,my is the mass/unit length
mI is mass moment of inertia (per unit length) in local coordinates
wbar is the weight density (per unit length)
area is the cross sectional area
moi is the moment of inertia
as is the shear area (0 -> shear deformation is ignored)
Cm are the effective added mass coefficients (see below)
Cd are the effective drag coefficients (see below)

nel is the element number
node1 and node2 are the node numbers
mat is the material number for the element
print .ne. 0, element results not printed
inc is the node increment used for generation
gen is the number of elements to generate
tension is the initial tension (for stiffness calculation only)

End input with a blank line.

The 2-D large displacement, elastic beam element assumes small strains. Hence, the forces are calculated as for a linear beam element, except that first the rigid body rotation is removed from the displacements.

The element must lie in a plane parallel to the global X-Y (default), X-Z, or Y-Z planes. The default for all elements is on the nbeam2d

record; this can be overwritten on a member basis on the member input record. The element local z-axis is in the global Z, -X, and Z directions, respectively.

On input, created arrays are:

```
.nbeam2d_mp(m,13) -> emodulus, gmodulus, wbar, area, moi, as, mx,
                    my, mI, cmx, cmy, cd, cy
.nbeam2d_el(n,4)  -> nodel, node2, material, print code
.nbeam2d_len(n)   -> element length
.nbeam2d_st(n,4)  -> Axial Force, V, M at nodel, M at node2
.nbeam2d_dir(n)   -> 1 -> X-Y; 2 -> X-Z; 3 -> Y-Z
```

mx and my are the mass densities per unit length in local coordinates. If density is specified, then $mx=my=density$. This element computes a lumped mass matrix in local coordinates. However, when transformed to global coordinates, it will no longer be diagonal unless $mx = my$ or the element is parallel to one of the global axes. If neither of these conditions is met, then a global diagonal mass matrix should not be used. Note that for large displacements, the mass matrix will also need to be reformulated if mx and my are not the same, and even if the element were initially parallel to a global axis, it will not in general remain parallel.

If the weight density is specified, it always acts in the -Z (global) direction. Therefore, if it is used the element should be in the X-Z or the Y-Z planes. The weight is always applied if it is specified.

cmx and cmy are the effective mass coefficients for a "Morison" treatment of fluid loading. These are effectively densities per unit length in local coordinates and will typically be equal to $1/2 \cdot \rho \cdot C_m \cdot D$, where ρ is the fluid density, C_m is the actual mass coefficient, and D is the "diameter". The program will multiply these by $L/2$ for a lumped formulation, where L is the original length. Similarly, cdx and cdy are effective damping coefficients. Note: both the added mass and drag terms are on the right-hand-side only. The mass term is multiplied by the fluid acceleration to obtain a load, and the drag term is multiplied by $abs(v-u)(v-u)$ to obtain a quadratic drag loading. The component of the added mass that is multiplied by the structure acceleration is included by specifying structure mass densities (mx,my) that include the added mass; the user is responsible for providing these modified mass densities.

For state calculation, element forces are put in .nbeam2d_st.

For response calculation, element does nothing.

For state output, results in .nbeam2d_st are printed.

For response output, no results are printed.

See Also

pnbeam2d pstate presponse

pnbeam2d

Command Syntax

pnbeam2d

Print nbeam2d element data

See Also
nbeam2d

ntruss

Two node large displacement truss element

Command Syntax

```
ntruss m=? n=? [nl=?]  
m=mat1 e=emodulus a=area [mbar=density] [-no_compression] (1 rec/mat1)  
n=nel nodes=node1,node2 mat=mat [print=print] &  
[inc=inc] [gen=gen] [tension=tension] [Lo=length] &  
[ecc1=delta_x,delta_y,delta_z] [ecc2=delta_x,delta_y,delta_z]
```

m is the number of different materials

n is the number of elements

nl = 1 -> linear stiffness

= 2 -> nonlinear and geometric stiffness (default)

mat1 is the material number

emodulus is the modulus of elasticity

area is the cross sectional area

density is the mass/unit length

-no_compression indicates tension-only material

nel is the element number

node1 and node2 are the node numbers

mat is the material number for the element

print .ne. 0, element results not printed

inc is the node increment used for generation

gen is the number of elements to generate

tension is the initial tension

length is the unstretched length

ecc1 are the offsets, in global coordinates, of the element start from node1

ecc2 are the offsets, in global coordinates, of the element end from node2

End input with a blank line.

The tension specified by tension= is only used to provide an initial stiffness to stabilize a slack initial configuration. The value is not used subsequently. The length specified by Lo= is the unstretched length (corresponding to zero force). The default unstretched length is the initial distance between the two ends. If Lo is specified, then any value specified for tension is not used; rather the initial tension is calculated.

If -no_compression is specified for a material, then the corresponding elements can only resist tension.

On input, created arrays are:

```
.ntruss_mp(m,4) -> modulus, area, density, compression code  
.ntruss_el(4,n) -> node1, node2, material, print code  
.ntruss_len(n) -> unstretched length  
.ntruss_st(n) -> element force  
.ntruss_ecc(n) -> eccentricity code  
.ntruss_ecc2(6,n)-> eccentricities (delta_x,delta_y,delta_z)
```

The element calculates a diagonal mass matrix only.
For state calculation, element forces are put in .ntruss_st.
For response calculation, element does nothing.
For state output, the axial forces in .ntruss_st are printed.

For response output, the element state is printed. If the unformatted write options on the presponse command are specified, the results are written to the file project_name.ntr. The data is written: element #, axial force.

constraints associated with an offset of the element end with the corresponding node are based on linear kinematics. Specifically, for nonzero offsets, the element end is 'slaved' to the node, and the displacements at the element end are determined based on rigid body, linear kinematics.

See Also
pntruss pstate presponse

pntruss

Command Syntax
pntruss
Print ntruss element data

See Also
ntruss

spring

Nonlinear, elastic spring element

----- OPTION 1 -----

Command Syntax
spring m=? n=?
m=mat e=Ko,Fyt,Fyc,Kpt,Kpc (1 record/material)
n=nel type=type nodes=node1,node2 mat=mat [print=print] l=lx,ly,lz

m is the number of different materials
n is the number of elements

mat is the material number
Ko is the initial spring stiffness
Fyt is the tensile yield strength
Fyc is the compressive yield strength (typically negative)
Kpt is the post-yield stiffness in tension
Kpc is the post-yield stiffness in compression

End input with a blank line.

----- OPTION 2 -----

Command Syntax
spring m=? n=?
m=mat fv=d1,f1,d2,f2,etc (1 record/material)
n=nel type=type nodes=node1,node2 mat=mat [print=print] l=lx,ly,lz

m is the number of different materials

n is the number of elements

mat is the material number

fv is a sequence of deformation,force (d,f) defining the deformation-force curve for the spring. The d1, d2, etc must be in ascending order. A minimum of 2 points must be specified, and a maximum of 10 points may be specified.

End input with a blank line.

----- BOTH OPTIONS -----

nel is the element number

type is the specifier for spring type

= 1 -> translational spring

= 2 -> rotational spring

node1 and node2 are the node numbers

mat is the material number

print .ne. 0 -> element printout suppressed

lx, ly, lz are direction cosines that specify the element orientation

Option 1 has a bilinear force-deformation relation (bilinear for each of the positive and negative deformation regimes. Option 2 is multilinear, with 10 datapoints for the whole spectrum (positive and negative spectrum). Option 1 is kept for backward compatibility. For option 2, if the calculated deformation is beyond the range specified, extrapolation is used.

If the spring is connected to "ground", leave node2 blank. If node2 is specified, the spring can be used to resist relative translation or rotation of nodes, whether they are separated or co-located.

For a translational spring, node2 is ignored (it is taken as "ground"). For a rotational spring, both node1 and node2 are used, and hence can be used to resist relative rotation of two otherwise identical nodes.

The direction cosines lx, ly, lz are with respect to the global coordinate system, and define the element orientation in the following sense. A grounded translational spring in the positive x-direction has a direction vector of 1, 0, 0. A grounded rotational spring with direction vector 1, 0, 0 means it resists rotation about the x-axis, with the right-hand-rule used to determine positive rotation. If the direction vector is not one of the global axes, the deformation of the element is calculated by taking the dot product of the displacement vector and the direction vector. If the spring connects two nodes, the direction is defined by the order of the nodes and the direction vector, not by the vector between the two nodes (which may be zero). Specifically, the relevant displacements are defined by the direction vector, and then the deformation is obtained by subtracting the displacement of node2 from node1. Note that this may be opposite from what might normally be expected. In both cases it can be imagined that node1 is at the head of the direction vector and node2, or ground, is at the tail.

On input, created arrays are:

.spring_mp(m,5) -> Ko, Fyt, Fyc, Kpt, Kpc

or

.spring_mp(m,21) -> d1,f1...d10,f10, number of actual points

.spring_el(n,5) -> spring type, node1, node2, material, print

.spring_dr(n,3) -> lx, ly, lz

`.spring_st(n,2)` -> element force and deformation

For state calculation, element forces are put in `.spring_st`.

For response calculation, element does nothing.

For state output, results in `.spring_st` are printed.

For response output, no results are printed.

See Also

`pspring` `pstate` `prresponse`

pspring

Command Syntax

`pspring`
Print spring element data

See Also

`spring`

stiff2n

Linear elastic 2-node stiffness element

Command Syntax

`stiff2n n=?`
`n=nel nodes=node1,node2 k=stiff__name [print=?]`

`n` is the number of elements

`nel` is the element number

`node1` and `node2` are the node numbers

`stiff_name` is the name of the pre-defined 6x6 stiffness matrix

`print.ne. 0` -> element printout suppressed

End input with a blank line.

This element allows a 6x6 stiffness matrix to be used to connect two nodes. It may find most usefulness in connecting two nodes that are at identical locations. The "deformation" of the element is defined as the 6 displacements of node2 minus the 6 displacements of node1. The element forces are then obtained by multiplying the deformation by the stiffness matrix. One use of the element is to specify a diagonal matrix, each diagonal with the spring stiffness connecting the two nodes.

On input, created arrays are:

`.stiff2n_el(3,n)` -> node1, node2, print code
`.stiff2n_k(32,n)` -> names of the matrices (each up to 31 characters)
`.stiff2n_st(6,n)` -> element force

For state calculation, element forces are put in `.stiff2n_st`.

For response calculation, element does nothing.

For state output, results in `.stiff2n_st` are printed.

For response output, no results are printed.

See Also

pstiff2n pstate presponse

pstiff2n

Command Syntax

```
pstiff2n
  Print 2-node stiffness element data
```

See Also

stiff2n

truss

Two node linear truss element

Command Syntax

```
truss m=? n=? [-kg]
m=mat1 e=emodulus a=area [mbar=density] (1 rec/mat1)
n=nel nodes=node1,node2 mat=mat [print=print] &
  [inc=inc] [gen=gen] [tension=tension] [Lo=length] &
  [ecc1=delta_x,delta_y,delta_z] [ecc2=delta_x,delta_y,delta_z]
```

m is the number of different materials

n is the number of elements

-kg is a flag to include geometric stiffness

mat1 is the material number

emodulus is the modulus of elasticity

area is the cross sectional area

density is the mass/unit length

nel is the element number

node1 and node2 are the node numbers

mat is the material number for the element

print .ne. 0, element results not printed

inc is the node increment used for generation

gen is the number of elements to generate

tension is the initial tension

length is the unstretched length

ecc1 are the offsets, in global coordinates, of the element start from

node1

ecc2 are the offsets, in global coordinates, of the element end from

node2

End input with a blank line.

This element is identical to the ntruss element with option nl=1 (linear stiffness), except this element also uses linear kinematics to determine the axial force. The tension specified by tension= is only used for a geometric stiffness matrix. The value is not used subsequently. The length specified by Lo= is the unstretched length (corresponding to zero force). The default unstretched length is the initial distance between the two ends. If Lo is specified, then any value specified for tension is not used; rather the initial tension is calculated.

On input, created arrays are:

```
.truss_mp(m,4)  -> modulus, area, density, unused  
.truss_el(4,n)  -> nodel, node2, material, print code  
.truss_len(n)   -> unstretched length  
.truss_st(n)    -> element force  
.truss_ecc(n)   -> eccentricity code  
.truss_ecc2(6,n)-> eccentricities (delta_x,delta_y,delta_z)
```

The element calculates a diagonal mass matrix only.

For state calculation, element forces are put in .truss_st.

For response calculation, element does nothing.

For state output, the axial forces in .truss_st are printed.

For response output, the element state is printed. If the unformatted write options on the prresponse command are specified, the results are written to the file project_name.tr. The data is written: element #, axial force.

See Also

ntruss ptruss pstate prresponse

ptruss

Command Syntax

ptruss

Print truss element data

See Also

truss

2.8. Miscellaneous Commands

fortran_kind

Command Syntax

```
fortran_kind
```

Prints the number of bytes for standard data types (Absoft compiler).

gauss_int

Command Syntax

```
gauss_int arg1 arg2 arg3 n=?
```

arg1 = function values at gauss points - (n*nel,1) matrix

arg2 = element lengths - (nel,1) matrix

arg3 = integration result/element - (nel,1) matrix

n = number of integration points (1 to 10)

This function numerically evaluates the integral of an arbitrary function over a one-dimensional domain. For two-dimensional domains, see the finite element `d2l3to9`, which can be used as well to integrate an arbitrary function as a special case of "error estimation".

See Also

`d2l3to9` `fem_error`

gauss_pts

Command Syntax

```
gauss_pts arg1 [arg2] [n=?,?,?]
```

The array `arg1` is created with the natural coordinates corresponding to $n_1 \times n_2 \times n_3$ Gauss quadrature. The dimension of the space, and the number of columns of `arg1`, is specified by the number of parameters specified by `n`. That is, if only n_1 is given, then `arg1(n1)` will be created; if n_1 and n_2 are specified, then `arg1(n1*n2,2)` will be created; etc. The maximum value of the n 's is 10. If `arg2` is given, the weights will be stored in it.

See Also

`gauss_int`

poly

Command Syntax

```
poly arg d=? [n=?] [option=?] c=?,?,?,...,...
```

Evaluate a polynomial defined by the coefficients specified by `c`. `arg` is an $(npts, q)$ array, where `npts` is the number of points at which to evaluate the function. If `option=0` (default), then the function only will be evaluated; if `option=1`, then the function and the first derivatives will be evaluated.

One-Dimensional Polynomial (d=1):

An arbitrary degree 1-D polynomial can be defined. The number of coefficients is specified by `n` (degree = $n-1$). The coefficients are specified in the order 1, x , x^2 , etc. The first column of `arg` contains the X -coordinates at which to evaluate the function. The function values will be placed in the second column. If requested, the first derivatives will be put in the third column.

Two-Dimensional Polynomial (d=2):

A maximum fifth degree polynomial (21 coefficients) can be specified. The coefficients are given in the order of Pascal's triangle, i.e., 1, x, y, x², xy, y², etc. The X,Y coordinates are expected in the first two columns of arg. The function values will be put in the third column. If requested, the first partial derivatives wrt X and Y will be put in the 4th and 5th columns of arg, respectively.

In either case, if the coefficients have been defined previously (e.g., by the input command) use the form `c=~array_name`, where `array_name` is the array with the coefficients.

tri_intpts

Command Syntax

```
tri_intpts arg1 [arg2] n=?
```

The array `arg1(n,3)` is created with the area coordinates corresponding to the `n` integration points for a triangular domain. If `arg2` is given, the weights are put in it. The number of points may be 1, 3, 4, 7, 9, or 12.

See Also

`gauss_pts`

userf

Command Syntax

```
userf f=?
```

User-defined functions. `f` specifies which function. The arguments and parameters depend on the function. Help is only provided if the developer of the function provides a help routine. This option should only be used for prototyping and routines of short term use. Often, help is not available for such routines.