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function [DEFL,REACT,ELE_FOR,AFLAG] = ud_3d1el(...

nnodes,coord,concen,fixity,nele,ends,A,Izz,Iyy,J,Cw,IsSym,Ysc,Zsc,Betay,Betaz,Betaw
,Zzz,Zyy,Ayy,Azz,...
    E,v,Fy,YldSurf,Wt,webdir,beta_ang,w,thermal,truss,anatype);

% Code developed by Mrunmayi Mungekar and Devasmit Dutta
%
% UD_3D1EL performs a user defined three-dimensional
% first-order elastic analysis of a structural system.
%
%
%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%
%
% Functions Called
%     < to be defined by the student >
%
% Dictionary of Variables
%     Input Information:
%         nnodes      == total number of nodes
%         coord(i,1:3) == node i's coordinates
%                        coord(i,1) = X coordinate
%                        coord(i,2) = Y coordinate
%                        coord(i,3) = Z coordinate
%         concen(i,1:6) == concentrated loads for node i's 6 d.o.f.
%                        concen(i,1) = force in global X direction
%                        concen(i,2) = force in global Y direction
%                        concen(i,3) = force in global Z direction
%                        concen(i,4) = moment about global X axis
%                        concen(i,5) = moment about global Y axis
%                        concen(i,6) = moment about global Z axis
%         fixity(i,1:6) == prescribed displacements for node i's 6 d.o.f.
%                        Note: A free d.o.f. will have a value of NaN
%                        and hence, you will find the Matlab function
%                        isnan very useful.
%                        Examples: If fixity(15,3) is set to NaN, then node 15's
%                                Z-disp component is free;
%                                If fixity(2,6) is set to 0.0, then node 2's
%                                Z-rotation component is supported;
%                                If fixity(5,2) is set to -2.1, then node 5's
%                                Y-disp component is supported and defined
%                                with a settlement of -2.1 units.
%         fixity(i,1) = prescribed disp. in global X direction
%         fixity(i,2) = prescribed disp. in global Y direction
%         fixity(i,3) = prescribed disp. in global Z direction
%         fixity(i,4) = prescribed rotation about global X axis
%         fixity(i,5) = prescribed rotation about global Y axis
%         fixity(i,6) = prescribed rotation about global Z axis
%         nele      == total number of elements
%         ends(i,1:14) == element i's nodal information
%                        ends(i,1) = start node #
%                        ends(i,2) = finish node #
%                        ends(i,3) = flag to indicate whether or not flexural
%                                moments are released at start node. ends(i,3)=0 both
not
%                                released (rigid connection); ends(i,3)=1 both flexural
%                                moments are released (pinned connection); ends(i,3)=2
%                                at least one of the flexural moments are partially or

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[illegible]

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%           YldSurf(i,3) = maximum My/Mpy value
%       Wt(i)           == element i's material weight density
%                       (Assume that gravity is directed in the negative global
Y dir)
%       webdir(i,1:3) == element i's unit web vector. This is a unit vector
%                       that defines the element's local y-y axis with respect
%                       to the global coordinate system. It is based on the
%                       structure's undeformed geometry.
%                       webdir(i,1) = x component of element's unit web
vector
%                       webdir(i,2) = y component of element's unit web
vector
%                       webdir(i,3) = z component of element's unit web
vector
%
%       NOTE: An element's 3x3 rotation matrix, [g], is
constructed
%       as follows: First, calculate a unit vector, x_vect, that
%       describes the element's local x-axis. Second, take the
%       cross product of x_vect and webdir(i,:) to obtain
z_vect,
%       i.e. z_vect = cross(x_vect,webdir(i,:)). Third, set
z_vect
%       to a unit vector, i.e. z_vect = z_vect/norm(z_vect).
%       Finally, the first row of [g] is x_vect, its second row
is
%       webdir(i,:), and its third row is z_vect.
%       beta_ang(i) == element i's web rotation angle. These values are
%       provided for those students who are required to
calculate
%       their own unit web vectors (see above). It is based
%       on the structure's undeformed geometry.
%       Note: MASTAN2 uses the following convention for
%       defining a member's default web orientation:
%       A vector defining the element's local y-axis
%       with respect to the global coordinate system
%       will have a positive component in the global
%       Y direction. If the element's local x-axis,
%       its length axis, is aligned with the global Y
%       axis, then element's local y-axis is aligned
%       with global negative X axis. After this initial
%       orientation, element i may be rotated about
%       its local x-axis by the amount defined by
%       its web rotation angle, beta_ang(i). The
%       angle is in radians and assumes a right-hand
%       convention about the local x-axis which runs from
%       the element's start node to its finish node.
%       w(i,1:3) == element i's uniform load which references its
%       local coordinate system
%       w(i,1) = x component of uniform load
%       w(i,2) = y component of uniform load
%       w(i,3) = z component of uniform load
%       thermal(i,1:4) == element i's thermal strain effects which reference its
%       local coordinate system
%       thermal(i,1) = coefficient of thermal expansion
%       thermal(i,2) = change in temperature at centroid
%       thermal(i,3) = linear temperature gradient in local
y-dir
%
%                       = (T_up_y - T_btm_y) / depth_y
%       thermal(i,4) = linear temperature gradient in local

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%                                AFLAG = 1      Successful
%                                AFLAG = 0      Unstable Structure
%                                AFLAG = inf    No analysis code available
%
%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%
%
%
% Start by defining all output arrays to be empty
%
DEFL=[]; REACT=[]; ELE_FOR=[];

memb_id = MD_member_id(nnodes, nele, ends);

disp('member id');
disp(memb_id);

load_dof = MD_load_dof(concen, nnodes);

disp('load_dof');
disp(load_dof);

AFLAG = inf;
%
% STUDENT NOTE:
%   In order for this routine to become fully active AFLAG
%   must be changed.
%
% Student's code starts here...
%
%
% Good luck CE Student!!!
%

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