

## Mystran Analysis Interface Module (AIM)

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## 0.1 Introduction

### 0.1.1 MYSTRAN AIM Overview

A module in the Computational Aircraft Prototype Syntheses (CAPS) has been developed to interact (primarily through input files) with the finite element structural solver MYSTRAN [1]. MYSTRAN is an open source, general purpose, linear finite element analysis computer program written by Dr. Bill Case. Available at, <http://www.mystran.com/>, MYSTRAN currently supports Linux and Windows operating systems.

An outline of the AIM's inputs, outputs and attributes are provided in [AIM Inputs](#) and [AIM Outputs](#) and [MYSTRAN AIM attributes](#), respectively.

The MYSTRAN AIM can automatically execute MYSTRAN, with details provided in [AIM Execution](#).

Details of the AIM's automated data transfer capabilities are outlined in [MYSTRAN Data Transfer](#)

### 0.1.2 Examples

An example problem using the MYSTRAN AIM may be found at [Mystran AIM Basic Example](#).

### 0.1.3 Clearance Statement

This software has been cleared for public release on 05 Nov 2020, case number 88ABW-2020-3462.

## 0.2 MYSTRAN AIM attributes

The following list of attributes are required for the MYSTRAN AIM inside the geometry input.

- **capsAIM** This attribute is a CAPS requirement to indicate the analysis the geometry representation supports.
- **capsGroup** This is a name assigned to any geometric body. This body could be a solid, surface, face, wire, edge or node. Recall that a string in ESP starts with a \$. For example, attribute `capsGroup $Wing`.
- **capsLoad** This is a name assigned to any geometric body where a load is applied. This attribute was separated from the `capsGroup` attribute to allow the user to define a local area to apply a load on without adding multiple `capsGroup` attributes. Recall that a string in ESP starts with a \$. For example, attribute `capsLoad $force`.
- **capsConstraint** This is a name assigned to any geometric body where a constraint/boundary condition is applied. This attribute was separated from the `capsGroup` attribute to allow the user to define a local area to apply a boundary condition without adding multiple `capsGroup` attributes. Recall that a string in ESP starts with a \$. For example, attribute `capsConstraint $fixed`.
- **capsIgnore** It is possible that there is a geometric body (or entity) that you do not want the MYSTRAN AIM to pay attention to when creating a finite element model. The `capsIgnore` attribute allows a body (or entity) to be in the geometry and ignored by the AIM. For example, because of limitations in OpenCASCADE a situation where two edges are overlapping may occur; `capsIgnore` allows the user to only pay attention to one of the overlapping edges.
- **capsBound** This is used to mark surfaces on the structural grid in which data transfer with an external solver will take place. See [MYSTRAN Data Transfer](#) for additional details.

## 0.3 AIM Inputs

The following list outlines the MYSTRAN inputs along with their default value available through the AIM interface. Unless noted these values will be not be linked to any parent AIMs with variables of the same name.

- **Proj\_Name = "mystran\_CAPS"**  
This corresponds to the project name used for file naming.
- **Tess\_Params = [0.025, 0.001, 15.0]**  
Body tessellation parameters used when creating a boundary element model. Tess\_Params[0] and Tess\_Params[1] get scaled by the bounding box of the body. (From the EGADS manual) A set of 3 parameters that drive the EDGE discretization and the FACE triangulation. The first is the maximum length of an EDGE segment or triangle side (in physical space). A zero is flag that allows for any length. The second is a curvature-based value that looks locally at the deviation between the centroid of the discrete object and the underlying geometry. Any deviation larger than the input value will cause the tessellation to be enhanced in those regions. The third is the maximum interior dihedral angle (in degrees) between triangle facets (or Edge segment tangents for a WIREBODY tessellation), note that a zero ignores this phase
- **Edge\_Point\_Min = 2**  
Minimum number of points on an edge including end points to use when creating a surface mesh (min 2).
- **Edge\_Point\_Max = 50**  
Maximum number of points on an edge including end points to use when creating a surface mesh (min 2).
- **Quad\_Mesh = False**  
Create a quadratic mesh on four edge faces when creating the boundary element model.
- **Property = NULL**  
Property tuple used to input property information for the model, see [FEA Property](#) for additional details.
- **Material = NULL**  
Material tuple used to input material information for the model, see [FEA Material](#) for additional details.
- **Constraint = NULL**  
Constraint tuple used to input constraint information for the model, see [FEA Constraint](#) for additional details.
- **Load = NULL**  
Load tuple used to input load information for the model, see [FEA Load](#) for additional details.
- **Analysis = NULL**  
Analysis tuple used to input analysis/case information for the model, see [FEA Analysis](#) for additional details.
- **Analysis\_Type = "Modal"**  
Type of analysis to generate files for, options include "Modal", "Static", and "Craig-Bampton".
- **Support = NULL**  
Support tuple used to input support information for the model, see [FEA Support](#) for additional details.
- **Mesh = NULL**  
A Mesh link.

## 0.4 AIM Execution

If auto execution is enabled when creating an MYSTRAN AIM, the AIM will execute MYSTRAN just-in-time with the command line:

```
mystran $Proj_Name.dat > Info.out
```

where preAnalysis generated the file Proj\_Name + ".dat" which contains the input information.

The analysis can be also be explicitly executed with `caps_execute` in the C-API or via `Analysis.runAnalysis` in the pyCAPS API.

Calling `preAnalysis` and `postAnalysis` is NOT allowed when auto execution is enabled.

Auto execution can also be disabled when creating an MYSTRAN AIM object. In this mode, `caps_execute` and `Analysis.runAnalysis` can be used to run the analysis, or MYSTRAN can be executed by calling `preAnalysis`, `system call`, and `posAnalysis` as demonstrated below with a pyCAPS example:

```
print ("\n\preAnalysis.....")
mystran.preAnalysis()
print ("\n\nRunning.....")
currentDirectory = os.getcwd() # Get our current working directory
os.chdir(mystran.analysisDir) # Move into test directory
os.system("mystran " + mystran.input.Proj_Name + ".dat > Info.out"); # Run via system call
os.chdir(currentDirectory) # Move back to top directory
print ("\n\postAnalysis.....")
mystran.postAnalysis()
```

## 0.5 AIM Outputs

The following list outlines the MYSTRAN outputs available through the AIM interface.

- **EigenValue** = List of Eigen-Values (  $\lambda$  ) after a modal solve.
- **EigenRadian** = List of Eigen-Values in terms of radians (  $\omega = \sqrt{\lambda}$  ) after a modal solve.
- **EigenFrequency** = List of Eigen-Values in terms of frequencies (  $f = \frac{\omega}{2\pi}$  ) after a modal solve.
- **EigenGeneralMass** = List of generalized masses for the Eigen-Values.

## 0.6 MYSTRAN Data Transfer

The MYSTRAN AIM has the ability to transfer displacements and eigenvectors from the AIM and pressure distributions to the AIM using the conservative and interpolative data transfer schemes in CAPS.

### 0.6.1 Data transfer from MYSTRAN (FieldOut)

- **"Displacement"**  
Retrieves nodal displacements from the \*.F06 file.
- **"EigenVector\_#"**  
Retrieves modal eigen-vectors from the \*.F06 file, where "#" should be replaced by the corresponding mode number for the eigen-vector (eg. `EigenVector_3` would correspond to the third mode, while `EigenVector_6` would be the sixth mode).

### 0.6.2 Data transfer to MYSTRAN (FieldIn)

- **"Pressure"**  
Writes appropriate load cards using the provided pressure distribution.

## 0.7 FEA Material

Structure for the material tuple = ("Material Name", "Value"). "Material Name" defines the reference name for the material being specified. The "Value" can either be a JSON String dictionary (see Section [JSON String Dictionary](#)) or a single string keyword (see Section [Single Value String](#)).

### 0.7.1 JSON String Dictionary

If "Value" is JSON string dictionary (e.g. "Value" = {"density": 7850, "youngModulus": 120000.0, "poissonRatio": 0.5, "materialType": "isotropic"}) the following keywords ( = default values) may be used:

- **materialType = "Isotropic"**  
Material property type. Options: Isotropic, Anisothotropic, Orthotropic, or Anisotropic.
- **youngModulus = 0.0**  
Also known as the elastic modulus, defines the relationship between stress and strain. Default if 'shearModulus' and 'poissonRatio' != 0,  $\text{youngModulus} = 2 * (1 + \text{poissonRatio}) * \text{shearModulus}$
- **shearModulus = 0.0**  
Also known as the modulus of rigidity, is defined as the ratio of shear stress to the shear strain. Default if 'youngModulus' and 'poissonRatio' != 0,  $\text{shearModulus} = \text{youngModulus} / (2 * (1 + \text{poissonRatio}))$
- **poissonRatio = 0.0**  
The fraction of expansion divided by the fraction of compression. Default if 'youngModulus' and 'shearModulus' != 0,  $\text{poissonRatio} = (2 * \text{youngModulus} / \text{shearModulus}) - 1$
- **density = 0.0**  
Density of the material.
- **thermalExpCoeff = 0.0**  
Thermal expansion coefficient of the material.
- **thermalExpCoeffLateral = 0.0**  
Thermal expansion coefficient of the material.
- **temperatureRef = 0.0**  
Reference temperature for material properties.
- **dampingCoeff = 0.0**  
Damping coefficient for the material.
- **youngModulusLateral = 0.0**  
Elastic modulus in lateral direction for an orthotropic material
- **shearModulusTrans1Z = 0.0**  
Transverse shear modulus in the 1-Z plane for an orthotropic material
- **shearModulusTrans2Z = 0.0**  
Transverse shear modulus in the 2-Z plane for an orthotropic material



### 0.7.2 Single Value String

If "Value" is a string, the string value may correspond to an entry in a predefined material lookup table. NOT YET IMPLEMENTED!!!!

## 0.8 FEA Property

Structure for the property tuple = ("Property Name", "Value"). "Property Name" defines the reference `capsGroup` for the property being specified. The "Value" can either be a JSON String dictionary (see Section [JSON String Dictionary](#)) or a single string keyword (see Section [Single Value String](#)).

### 0.8.1 JSON String Dictionary

If "Value" is JSON string dictionary (e.g. "Value" = {"shearMembraneRatio": 0.83, "bendingInertiaRatio": 1.0, "membraneThickness": 0.2, "propertyType": "Shell"}) the following keywords ( = default values) may be used:

- **propertyType = No Default value**  
Type of property to apply to a given `capsGroup` Name. Options: ConcentratedMass, Rod, Bar, Shear, Shell, Composite, and Solid
- **material = "Material Name" ([FEA Material](#))**  
"Material Name" from [FEA Material](#) to use for property. If no material is set the first material created will be used
- **crossSecArea = 0.0**  
Cross sectional area.
- **torsionalConst = 0.0**  
Torsional constant.
- **torsionalStressReCoeff = 0.0**  
Torsional stress recovery coefficient.
- **massPerArea = 0.0**  
Non-structural mass per unit length.
- **zAxisInertia = 0.0**  
Section moment of inertia about the element z-axis.
- **yAxisInertia = 0.0**  
Section moment of inertia about the element y-axis.
- **yCoords[4] = [0.0, 0.0, 0.0, 0.0]**  
Element y-coordinates, in the bar cross-section, of four points at which to recover stresses

- **zCoords[4] = [0.0, 0.0, 0.0, 0.0]**  
Element z-coordinates, in the bar cross-section, of four points at which to recover stresses
- **areaShearFactors[2] = [0.0, 0.0]**  
Area factors for shear.
- **crossProductInertia = 0.0**  
Section cross-product of inertia.
- **crossSecType = NULL**  
Cross-section type. Must be one of following character variables: BAR, BOX, BOX1, CHAN, CHAN1, CHAN2, CROSS, H, HAT, HEXA, I, I1, ROD, T, T1, T2, TUBE, or Z.
- **crossSecDimension = [0,0,0,...]**  
Cross-sectional dimensions (length of array is dependent on the "crossSecType"). Max supported length array is 10!
- **membraneThickness = 0.0**  
Membrane thickness.
- **bendingInertiaRatio = 1.0**  
Ratio of actual bending moment inertia to the bending inertia of a solid plate of thickness "membraneThickness"
- **shearMembraneRatio = 5.0/6.0**  
Ratio shear thickness to membrane thickness.
- **materialBending = "Material Name" (FEA Material)**  
"Material Name" from [FEA Material](#) to use for property bending. If no material is given and "bendingInertiaRatio" is greater than 0, the material name provided in "material" is used.
- **materialShear = "Material Name" (FEA Material)**  
"Material Name" from [FEA Material](#) to use for property shear. If no material is given and "shearMembraneRatio" is greater than 0, the material name provided in "material" is used.
- **massPerArea = 0.0**  
Non-structural mass per unit area.
- **zOffsetRel = 0.0**  
Relative offset from the surface of grid points to the element reference plane as a percentage of the thickness.  
 $zOffset = thickness * zOffsetRel / 100$
- **compositeMaterial = "no default"**  
List of "Material Name"s, ["Material Name -1", "Material Name -2", ...], from [FEA Material](#) to use for composites.
- **shearBondAllowable = 0.0**  
Allowable interlaminar shear stress.

- **symmetricLaminate = False**  
Symmetric lamination option. True- SYM only half the plies are specified, for odd number plies 1/2 thickness of center ply is specified with the first ply being the bottom ply in the stack, default (False) all plies specified.
- **compositeFailureTheory = "(no default)"**  
Composite failure theory. Options: "HILL", "HOFF", "TSAI", and "STRN"
- **compositeThickness = (no default)**  
List of composite thickness for each layer (e.g. [1.2, 4.0, 3.0]). If the length of this list doesn't match the length of the "compositeMaterial" list, the list is either truncated [ >length("compositeMaterial")] or expanded [ <length("compositeMaterial")] in which case the last thickness provided is repeated.
- **compositeOrientation = (no default)**  
List of composite orientations (angle relative element material axis) for each layer (eg. [5.0, 10.0, 30.0]). If the length of this list doesn't match the length of the "compositeMaterial" list, the list is either truncated [ >length("compositeMaterial")] or expanded [ <length("compositeMaterial")] in which case the last orientation provided is repeated.
- **mass = 0.0**  
Mass value.
- **massOffset = [0.0, 0.0, 0.0]**  
Offset distance from the grid point to the center of gravity for a concentrated mass.
- **massInertia = [0.0, 0.0, 0.0, 0.0, 0.0, 0.0]**  
Mass moment of inertia measured at the mass center of gravity.

## 0.8.2 Single Value String

If "Value" is a string, the string value may correspond to an entry in a predefined property lookup table. NOT YET IMPLEMENTED!!!!

## 0.9 FEA Constraint

Structure for the constraint tuple = ("Constraint Name", "Value"). "Constraint Name" defines the reference name for the constraint being specified. The "Value" can either be a JSON String dictionary (see Section [JSON String Dictionary](#)) or a single string keyword (see Section [Single Value String](#)).

### 0.9.1 JSON String Dictionary

If "Value" is JSON string dictionary (eg. "Value" = {"groupName": "plateEdge", "dofConstraint": 123456}) the following keywords ( = default values) may be used:

- **constraintType = "ZeroDisplacement"**  
Type of constraint. Options: "Displacement", "ZeroDisplacement".

|| NASTRAN || ASTROS || HSM || ABAQUS)

- **groupName = "(no default)"**  
Single or list of `capsConstraint` names on which to apply the constraint (e.g. "Name1" or ["Name1", "↔ Name2", "...]). If not provided, the constraint tuple name will be used.
- **dofConstraint = 0**  
Component numbers / degrees of freedom that will be constrained (123 - zero translation in all three directions).
- **gridDisplacement = 0.0**  
Value of displacement for components defined in "dofConstraint".

## 0.9.2 Single Value String

If "Value" is a string, the string value may correspond to an entry in a predefined constraint lookup table. NOT YET IMPLEMENTED!!!!

## 0.10 FEA Support

Structure for the support tuple = ("Support Name", "Value"). "Support Name" defines the reference name for the support being specified. The "Value" can either be a JSON String dictionary (see Section [JSON String Dictionary](#)) or a single string keyword (see Section [Single Value String](#)).

### 0.10.1 JSON String Dictionary

If "Value" is JSON string dictionary (eg. "Value" = {"groupName": "plateEdge", "dofSupport": 123456}) the following keywords ( = default values) may be used:

- **groupName = "(no default)"**  
Single or list of `capsConstraint` names on which to apply the support (e.g. "Name1" or ["Name1", "↔ Name2", ...]. If not provided, the constraint tuple name will be used.
- **dofSupport = 0**  
Component numbers / degrees of freedom that will be supported (123 - zero translation in all three directions).

### 0.10.2 Single Value String

If "Value" is a string, the string value may correspond to an entry in a predefined support lookup table. NOT YET IMPLEMENTED!!!!

## 0.11 FEA Connection

Structure for the connection tuple = ("Connection Name", "Value"). "Connection Name" defines the reference name to the `capsConnect` being specified and denotes the "source" node for the connection. The "Value" can either be a JSON String dictionary (see Section [JSON String Dictionary](#)) or a single string keyword (see Section [Single Value String](#)).

### 0.11.1 JSON String Dictionary

If "Value" is JSON string dictionary (e.g. "Value" = {"dofDependent": 1, "propertyType": "RigidBody"}) the following keywords ( = default values) may be used:

- **connectionType = RigidBody**  
Type of connection to apply to a given capsConnect pair defined by "Connection Name" and the "groupName".  
Options: Mass (scalar), Spring (scalar), Damper (scalar), RigidBody.
- **dofDependent = 0**  
Component numbers / degrees of freedom of the dependent end of rigid body connections (ex. 123 - translation in all three directions).
- **componentNumberStart = 0**  
Component numbers / degrees of freedom of the starting point of the connection for mass, spring, and damper elements (scalar) ( 0 <= Integer <= 6).
- **componentNumberEnd= 0**  
Component numbers / degrees of freedom of the ending point of the connection for mass, spring, and damper elements (scalar), rigid body interpolative connection ( 0 <= Integer <= 6).
- **stiffnessConst = 0.0**  
Stiffness constant of a spring element (scalar).
- **dampingConst = 0.0**  
Damping coefficient/constant of a spring or damping element (scalar).
- **stressCoeff = 0.0**  
Stress coefficient of a spring element (scalar).
- **mass = 0.0**  
Mass of a mass element (scalar).
- **glue = False**  
Turn on gluing for the connection.
- **glueNumMaster = 5**  
Maximum number of the masters for a glue connections.
- **glueSearchRadius = 0**  
Search radius when looking for masters for a glue connections.
- **weighting = 1**  
Weighting factor for a rigid body interpolative connections.
- **groupName = "(no default)"**  
Single or list of capsConnect names on which to connect the nodes found with the tuple name ("↔ Connection Name") to. (e.g. "Name1" or ["Name1","Name2",...]).

### 0.11.2 Single Value String

If "Value" is a string, the string value may correspond to an entry in a predefined connection lookup table. NOT YET IMPLEMENTED!!!!

## 0.12 FEA Load

Structure for the load tuple = ("Load Name", "Value"). "Load Name" defines the reference name for the load being specified. The "Value" can either be a JSON String dictionary (see Section [JSON String Dictionary](#)) or a single string keyword (see Section [Single Value String](#)).

### 0.12.1 JSON String Dictionary

If "Value" is JSON string dictionary (e.g. "Value" = {"groupName": "plate", "loadType": "Pressure", "pressureForce": 2000000.0}) the following keywords (= default values) may be used:

- **loadType = "(no default)"**  
Type of load. Options: "GridForce", "GridMoment", "Rotational", "Thermal", "Pressure", "PressureDistribute", "PressureExternal", "Gravity".
- **groupName = "(no default)"**  
Single or list of `capsLoad` names on which to apply the load (e.g. "Name1" or ["Name1", "Name2", ...]). If not provided, the load tuple name will be used.
- **loadScaleFactor = 1.0**  
Scale factor to use when combining loads.
- **forceScaleFactor = 0.0**  
Overall scale factor for the force for a "GridForce" load.
- **directionVector = [0.0, 0.0, 0.0]**  
X-, y-, and z- components of the force vector for a "GridForce", "GridMoment", or "Gravity" load.
- **momentScaleFactor = 0.0**  
Overall scale factor for the moment for a "GridMoment" load.
- **gravityAcceleration = 0.0**  
Acceleration value for a "Gravity" load.
- **pressureForce = 0.0**  
Uniform pressure force for a "Pressure" load (only applicable to 2D elements).
- **pressureDistributeForce = [0.0, 0.0, 0.0, 0.0]**  
Distributed pressure force for a "PressureDistribute" load (only applicable to 2D elements). The four values correspond to the 4 (quadrilateral elements) or 3 (triangle elements) node locations.

- **angularVelScaleFactor = 0.0**  
An overall scale factor for the angular velocity in revolutions per unit time for a "Rotational" load.
- **angularAccScaleFactor = 0.0**  
An overall scale factor for the angular acceleration in revolutions per unit time squared for a "Rotational" load.
- **coordinateSystem = "(no default)"**  
Name of coordinate system in which defined force components are in reference to. If no value is provided the global system is assumed.
- **temperature = 0.0**  
Temperature at a given node for a "Temperature" load.
- **temperatureDefault = 0.0**  
Default temperature at a node not explicitly being used for a "Temperature" load.

### 0.12.2 Single Value String

If "Value" is a string, the string value may correspond to an entry in a predefined load lookup table. NOT YET IMPLEMENTED!!!!

## 0.13 FEA Analysis

Structure for the analysis tuple = ('Analysis Name', 'Value'). 'Analysis Name' defines the reference name for the analysis being specified. The "Value" can either be a JSON String dictionary (see Section [JSON String Dictionary](#)) or a single string keyword (see Section [Single Value String](#)).

### 0.13.1 JSON String Dictionary

If "Value" is JSON string dictionary (e.g. "Value" = {"numDesiredEigenvalue": 10, "eigenNormaliztion": "MASS", "numEstEigenvalue": 1, "extractionMethod": "GIV", "frequencyRange": [0, 10000]}) the following keywords ( = default values) may be used:

- **analysisType = "Modal"**  
Type of load. Options: "Modal", "Static".
- **analysisLoad = "(no default)"**  
Single or list of "Load Name"s defined in [FEA Load](#) in which to use for the analysis (e.g. "Name1" or ["↔ Name1", "Name2", ...]).
- **analysisConstraint = "(no default)"**  
Single or list of "Constraint Name"s defined in [FEA Constraint](#) in which to use for the analysis (e.g. "Name1" or ["Name1", "Name2", ...]).

- **analysisSupport = "(no default)"**  
Single or list of "Support Name"s defined in [FEA Support](#) in which to use for the analysis (e.g. "Name1" or ["Name1","Name2",...]).
- **extractionMethod = "(no default)"**  
Extraction method for modal analysis.
- **frequencyRange = [0.0, 0.0]**  
Frequency range of interest for modal analysis.
- **numEstEigenvalue = 0**  
Number of estimated eigenvalues for modal analysis.
- **numDesiredEigenvalue = 0**  
Number of desired eigenvalues for modal analysis.
- **eigenNormalization = "(no default)"**  
Method of eigenvector renormalization. Options: "POINT", "MAX", "MASS"
- **gridNormalization = 0**  
Grid point to be used in normalizing eigenvector to 1.0 when using eigenNormalization = "POINT"
- **componentNormalization = 0**  
Degree of freedom about "gridNormalization" to be used in normalizing eigenvector to 1.0 when using eigenNormalization = "POINT"
- **lanczosMode = 2**  
Mode refers to the Lanczos mode type to be used in the solution. In mode 3 the mass matrix, Maa, must be nonsingular whereas in mode 2 the matrix  $K_{aa} - \sigma * M_{aa}$  must be nonsingular
- **lanczosType = "(no default)"**  
Lanczos matrix type. Options: DPB, DGB.
- **aeroSymmetryXY = "(no default)"**  
Aerodynamic symmetry about the XY Plane. Options: SYM, ANTISYM, ASYM. SYMMETRIC Indicates that a half span aerodynamic model is moving in a symmetric manner with respect to the XY plane. ANTISYMMETRIC Indicates that a half span aerodynamic model is moving in an antisymmetric manner with respect to the XY plane. ASYMMETRIC Indicates that a full aerodynamic model is provided.
- **aeroSymmetryXZ = "(no default)"**  
Aerodynamic symmetry about the XZ Plane. Options: SYM, ANTISYM, ASYM. SYMMETRIC Indicates that a half span aerodynamic model is moving in a symmetric manner with respect to the XZ plane. ANTISYMMETRIC Indicates that a half span aerodynamic model is moving in an antisymmetric manner with respect to the XZ plane. ASYMMETRIC Indicates that a full aerodynamic model is provided.

### 0.13.2 Single Value String

If "Value" is a string, the string value may correspond to an entry in a predefined analysis lookup table. NOT YET IMPLEMENTED!!!!



## 0.14 FEA DesignVariable

Structure for the design variable tuple = ("DesignVariable Name", "Value"). "DesignVariable Name" defines the reference name for the design variable being specified. This string will be used in the FEA input directly. The "Value" must be a JSON String dictionary (see Section [JSON String Dictionary](#)).

### 0.14.1 JSON String Dictionary

If "Value" is JSON string dictionary the following keywords ( = default values) may be used:

## 0.15 FEA DesignVariableRelation

Structure for the design variable tuple = ("DesignVariableRelation Name", "Value"). "DesignVariableRelation Name" defines the reference name for the design variable being specified. This string will be used in the FEA input directly. The "Value" must be a JSON String dictionary (see Section [JSON String Dictionary](#)).

### 0.15.1 JSON String Dictionary

If "Value" is JSON string dictionary the following keywords ( = default values) may be used:

## 0.16 FEA DesignConstraint

Structure for the design constraint tuple = ("DesignConstraint Name", "Value"). "DesignConstraint Name" defines the reference name for the design constraint being specified. The "Value" must be a JSON String dictionary (see Section [JSON String Dictionary](#)).

### 0.16.1 JSON String Dictionary

If "Value" is JSON string dictionary the following keywords ( = default values) may be used:

## 0.17 FEA DesignEquation

Structure for the design equation tuple = ("DesignEquation Name", ["Value1", ... , "ValueN"]). "DesignEquation Name" defines the reference name for the design equation being specified. This string will be used in the FEA input directly. The values "Value1", ... , "ValueN" are a list of strings containing the equation definitions. (see Section [List of equation strings](#)).

### 0.17.1 List of equation strings

Each design equation tuple value is a list of strings containing the equation definitions

## 0.18 FEA TableConstant

Structure for the table constant tuple = ("TableConstant Name", "Value"). "TableConstant Name" defines the reference name for the table constant being specified. This string will be used in the FEA input directly. The "Value" is the value of the table constant.

## 0.19 FEA DesignResponse

Structure for the design response tuple = ("DesignResponse Name", "Value"). "DesignResponse Name" defines the reference name for the design response being specified. This string will be used in the FEA input directly. The "Value" must be a JSON String dictionary (see Section [JSON String Dictionary](#)).

### 0.19.1 JSON String Dictionary

If "Value" is JSON string dictionary the following keywords ( = default values) may be used:

## 0.20 FEA DesignEquationResponse

Structure for the design equation response tuple = ("DesignEquationResponse Name", "Value"). "DesignEquationResponse Name" defines the reference name for the design equation response being specified. This string will be used in the FEA input directly. The "Value" must be a JSON String dictionary (see Section [JSON String Dictionary](#)).

### 0.20.1 JSON String Dictionary

If "Value" is JSON string dictionary the following keywords ( = default values) may be used:

## 0.21 FEA DesignOptParam

Structure for the design optimization parameter tuple = ("DesignOptParam Name", "Value"). "DesignOptParam Name" defines the reference name for the design optimization parameter being specified. This string will be used in the FEA input directly. The "Value" is the value of the design optimization parameter.

## 0.22 Mystran AIM Basic Example

This is a walkthrough for using MYSTRAN AIM to analyze a three-dimensional wing with internal ribs and spars.

### 0.22.1 Prerequisites

It is presumed that ESP and CAPS have been already installed, as well as MYSTRAN.

### 0.22.1.1 Script files

Two scripts are used for this illustration:

1. feaWingBEM.csm: Creates geometry, as described in the next section ([Creating Geometry using ESP](#)).
2. mystran\_PyTest.py: pyCAPS script for performing analysis, as described in [Performing analysis using pyCAPS](#).

## 0.22.2 Creating Geometry using ESP

The CSM script generates Bodies which are designed to be used by specific AIMs. The AIMs that the Body is designed for is communicated to the CAPS framework via the "capsAIM" string attribute. This is a semicolon-separated string with the list of AIM names. Thus, the CSM author can give a clear indication to which AIMs should use the Body. In this example, the list contains the structural finite element analysis tools that can analyze the body:

```
attribute capsAIM $nastranAIM;astrosAIM;mystranAIM;masstranAIM;egadsTessAIM
```

A typical geometry model can be created and interactively modified using design parameters. These design parameters are either design- or geometry- based. In this example, a wing configuration is created using following design parameters.

```
# Design Parameters for OML
despmtr thick 0.12 frac of local chord
despmtr camber 0.04 frac of local chord
despmtr area 10.0
despmtr aspect 6.00
despmtr taper 0.60
despmtr sweep 20.0 deg (of c/4)
despmtr washout 5.00 deg (down at tip)
despmtr dihedral 4.00 deg
# Design Parameters for BEM
cfgpmtr nrrib 11 number of ribs
despmtr spar1 0.20 frac of local chord
despmtr spar2 0.75 frac of local chord
```

After our design parameters are defined they are used to setup other local variables (analytically) for the outer model line (OML).

```
# OML
set span sqrt(aspect*area)
set croot 2*area/span/(1+taper)
set ctip croot*taper
set dxtip (croot-ctip)/4+span/2*tand(sweep)
set dytip span/2*tand(dihedral)
```

In a similar manner, local variables are defined for the ribs and spars.

```
# wing ribs
set Nrib nint(nrrib)
# wing spars
set eps 0.01*span
```

Once all design and local variables are defined, a full span, solid model is created by "ruling" together NACA series airfoils (following a series of scales, rotations, and translations).

```
mark
# Right tip
udprim naca Thickness thick Camber camber
scale ctip
rotatez washout ctip/4 0
translate dxtip dytip -span/2
# root
udprim naca Thickness thick Camber camber
scale croot
# left tip
udprim naca Thickness thick Camber camber
scale ctip
rotatez washout ctip/4 0
translate dxtip dytip +span/2
rule
attribute OML 1
```

Once complete, the wing is stored for later use under the name OML.

```
store      OML
```

Next, the inner layout of the ribs and spars are created using the waffle udprim.

```
udprim waffle Depth +6*thick*croot Filename «
  patbeg i Nrib
    point A at (span/2)*(2*i-Nrib-1)/Nrib -0.01*croot
    point B at (span/2)*(2*i-Nrib-1)/Nrib max(croot,dxtip+ctip)
    line AB A B tagComponent=rib tagIndex=!val2str(i,0)
  patend
  point A at -span/2-eps spar1*ctip+dxtip
  point B at 0 spar1*croot
  line AB A B tagComponent=spar tagIndex=1 tagPosition=left
  point A at span/2+eps spar1*ctip+dxtip
  point B at 0 spar1*croot
  line AB A B tagComponent=spar tagIndex=1 tagPosition=right
  point A at -span/2-eps spar2*ctip+dxtip
  point B at 0 spar2*croot
  line AB A B tagComponent=spar tagIndex=2 tagPosition=left
  point A at span/2+eps spar2*ctip+dxtip
  point B at 0 spar2*croot
  line AB A B tagComponent=spar tagIndex=2 tagPosition=right
»
```

An attribute is then placed on ribs and spars so that the geometry components may be reference by the MYSTRAN AIM.

```
attribute capsGroup $Ribs_and_Spars
```

Following a series of rotations and translations the ribs and spars are stored for later use.

```
translate 0 0 -3*thick*croot
rotatey 90 0 0
rotatez -90 0 0
store layoutRibSpar
```

Next, the layout of the ribs and spars are intersected the outer mold line of wing, which results in only keeping the part of layout that is inside the OML.

```
restore layoutRibSpar
restore OML
intersect
```

Finally, select faces (airfoil sections at the root) are tagged, so that a constraint may be applied later.

```
udprim editAttr filename «
  edge adj2face tagComponent=spar tagPosition=right
  and adj2face tagComponent=spar tagPosition=left
  set capsConstraint=Rib_Constraint
  node adj2face tagComponent=spar tagPosition=right
  and adj2face tagComponent=spar tagPosition=left
  set capsConstraint=Rib_Constraint
»
ifthen nint(mod(Nrib,2)) ne 0
  set midRib Nrib/2
  select face $tagComponent $rib $tagIndex val2str(midRib,0)
  attribute tagPosition $root

  udprim editAttr filename «
    face has tagComponent=rib tagPosition=root
    set capsConstraint=Rib_Constraint

    edge adj2face tagComponent=rib tagPosition=root
    set capsConstraint=Rib_Constraint

    node adj2face tagComponent=rib tagPosition=root
    set capsConstraint=Rib_Constraint
  »
endif
```

The above \*.csm file results in the follow geometry model:

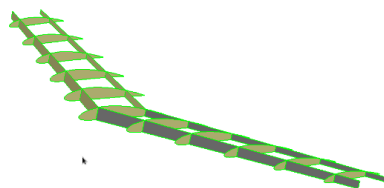


Figure 1 Wing built up element model

### 0.22.3 Performing analysis using pyCAPS

The first step in the pyCAPS script is to import the required modules. For this example the following modules are used,

```
# Import pyCAPS module
import pyCAPS
# Import os module
import os
import argparse
```

Similarly, local variables used throughout the script may be defined.

```
projectName = "MystranModalWingBEM"
workDir = os.path.join(str(args.workDir[0]), projectName)
```

Once the required modules have been loaded, a pyCAPS.Problem can be instantiated with the desired geometry file.

```
geometryScript = os.path.join(".", "csmData", "feaWingBEM.csm")
myProblem = pyCAPS.Problem(problemName=workDir,
                           capsFile=geometryScript,
                           outLevel=args.outLevel)
```

After the geometry is loaded, the MYSTRAN AIM needs to be instantiated.

```
mystranAIM = myProblem.analysis.create(aim = "mystranAIM",
                                       name = "mystran" )
```

Once loaded analysis parameters specific to MYSTRAN need to be set (see [AIM Inputs](#)). These parameters are automatically converted into MYSTRAN specific format and transferred into the MYSTRAN configuration file. One will note in the following snippet the instance of the AIM is referenced in two different manners: 1. Using the returned object from load call and 2. Using the "name" key in the Problem.analysis Sequence. While syntactically different, these two forms are essentially identical.

```
# Link Surface_Mesh
mystranAIM.input["Mesh"].link(myProblem.analysis["tess"].output["Surface_Mesh"])
# Set project name so a mesh file is generated
mystranAIM.input.Proj_Name = projectName
```

Along the same lines of setting the input values above the "Analysis" (see [FEA Analysis](#)), "Material" (see [FEA Material](#)), "Property" (see [FEA Property](#)), and "Constraint" (see [FEA Constraint](#)) tuples are used to set more complex information. The user is encouraged to read the additional documentation on these inputs for further explanations. Once provided this information is converted into MYSTRAN specific syntax and set in the MYSTRAN configuration file.

```
# Set analysis
eigen = { "extractionMethod" : "Lanczos",
          "frequencyRange"   : [0, 50],
          "numEstEigenvalue" : 1,
          "eigenNormaliztion" : "MASS"}
mystranAIM.input.Analysis = {"EigenAnalysis": eigen}
# Set materials
unobtainium = { "youngModulus" : 2.2E11 ,
                "poissonRatio" : .33,
                "density"      : 7850}
madeupium   = { "materialType" : "isotropic",
                "youngModulus" : 1.2E9 ,
                "poissonRatio" : .5,
                "density"      : 7850}
mystranAIM.input.Material = {"Unobtainium": unobtainium,
                             "Madeupium" : madeupium}
# Set property
shell = { "propertyType" : "Shell",
          "membraneThickness" : 0.2,
          "bendingInertiaRatio" : 1.0, # Default
          "shearMembraneRatio" : 5.0/6.0} # Default }
mystranAIM.input.Property = {"Ribs_and_Spars": shell}
```

Once all the inputs have been set, outputs can be directly requested. The MYSTRAN analysis will be automatically executed just-in-time ([AIM Execution](#)).

Finally, available AIM outputs (see [AIM Outputs](#)) may be retrieved, for example:

```
print ("\nGetting results for natural frequencies.....")
natrualFreq = mystranAIM.output.EigenFrequency
for mode, i in enumerate(natrualFreq):
    print ("Natural freq ( Mode", mode, ") = ", i, "(Hz)")
```

results in,

```
Natural freq (Mode 1) = 1.89166 (Hz)
Natural freq (Mode 2) = 6.33335 (Hz)
Natural freq (Mode 3) = 6.51397 (Hz)
Natural freq (Mode 4) = 23.88463 (Hz)
Natural freq (Mode 5) = 24.98205 (Hz)
Natural freq (Mode 6) = 28.23676 (Hz)
Natural freq (Mode 7) = 32.53667 (Hz)
Natural freq (Mode 8) = 33.92054 (Hz)
Natural freq (Mode 9) = 43.49964 (Hz)
```

#### **0.22.4 Executing pyCAPS script**

Issuing the following command executes the script:

```
python mystran_PyTest.py
```

# Bibliography

- [1] William Case. *MYSTRAN General Purpose Finite Element Structural Analysis Computer Program (Linux Version 6.35) [Software Manual]*, Nov. 2011. Available from <http://www.MYSTRAN.com>. 1

