YLVIS v4.2 User Manual

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This document is a user manual for the pYthon-based consoLe finite-Volume meshIng Software, for generation of simple 2D and 3D quad-based and hexa-based domains and 3D spherical domains for computational fluid dynamics simulations.

This program is an intellectual property of KU Leuven. Please, cite it appropriately if used in published work and inform us of its application. For questions and concerns, contact the author of the repository.

This program has two main modes of operation:

- Generation of simple blocked 2D and 3D quad based domains
- Generation of a spherical grid based on a surface mesh

These modes were developed in parallel due to the applications on which the code was used (fundamental physics modelling and solar corona modelling, respectively).

Separate python scripts are used for the two modes. With each of them, there is a respective fortran output generating executable (write.exe) and also the source code in case you want to modify / create your own output format. The procedure is two-fold:

- 1. First, create a native mesh file format using the python script
- 2. Second, transform this native format to GAMBIT neutral format with the fortran script

The reason why the second script is in fortran is due to the fact that the GAMBIT neutral format writing instructions are formulated in fortran, thus this is the most reliable way to produce a proper output file. The write.exe executable can be used if the is no need to modify the format, so in that case, the user does not have to work with fortran programming at all.

More details about how to run the program depending on the mode of operation are given in the following two sections.

1. Simple blocked 2D and 3D quad based domains

The main purpose of the code is to create simple quad-based (or hexagonal-based in 3D) meshes in which the user is in complete control of mesh stretching and other properties. For this, the code main.py is used.

The name of the native file is saved in the variable name. Each mesh type (whether it is the entire mesh or a block - blocking will be discussed later) has several params that must be defined:

block_type: the dimension and type of mesh, e.g. 2D_rect_quad or 3D_box_quad
 (currently the only supported, later we plan to extend to triangular and prism types)

- refinement: which consists of the number of nodes in each dimension, the type of stretching in each dimension (the stretching function) and the parameters for each stretching function if required
- xmin, xmax, ymin, ymax (, zmin, zmax): minimum and maximum coordinates in each dimension, giving the mesh size
- start_n, start_e: these denote from which index should the indexing of the nodes and elements of the block start, can be easily obtained by taking the length of the nodes and elements array prior to adding of the block (or set to 0 in case of a single block mesh)

The stretching functions currently implemented are the following (s is spacing, N is number of elements, idx is the current cell index and a the parameter, $\zeta = idx/(N-1)$ unless stated otherwise):

- uniform stretching

$$s = \zeta / I$$

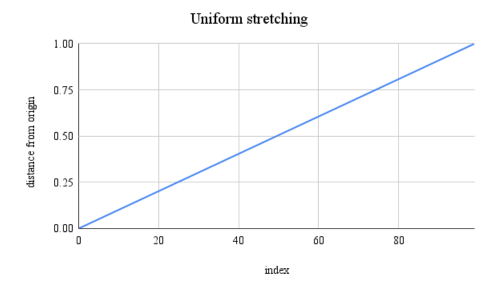


Figure 1: Uniform stretching

- negatsin stretching, type I (if a = 1):

$$\zeta = 198.0 \text{idx}/(N-1)$$

$$s = -0.0213 + 0.0149\zeta - 0.000151\zeta^2 + 0.000000515\zeta^3$$

$$\zeta_{\text{max}} = 198.$$

$$\min = -0.0213$$

$$\max = -0.0213 + 0.0149\zeta_{\text{max}} - 0.000151\zeta_{\text{max}}^2 + 0.000000515\zeta_{\text{max}}^3$$

range =
$$(max - min)$$

 $s = s - min$
 $s = s/range$

- negatsin stretching, type II (if a = 2):

$$s = -0.0199 + 2.41\zeta - 4.18\zeta^{2} + 2.81\zeta^{3}$$

 $min = -0.0199$
 $max = 1.0201$
 $range = (max - min)$
 $s = s - min$
 $s = s/range$

- negatsin stretching, type III (if $a \neq 1 \& \& a \neq 2$):

$$s = 0.00712 + 1.5\zeta + 2.53\zeta^2 - 16.0\zeta^3 + 22.7\zeta^4 - 10.7\zeta^5 + 0.966\zeta^6$$

$$min = 0.00712$$

$$max = 1.00312$$

$$range = (max - min)$$

$$s = s - min$$

$$s = s/range$$

Negative sinus stretching types

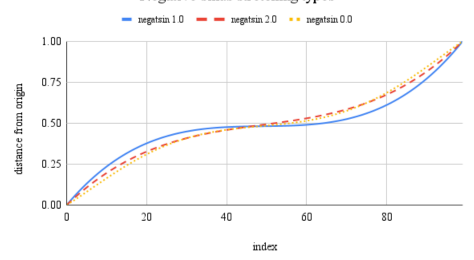


Figure 2: Negative sine stretching types

Double sided hyperbolic tan stretching

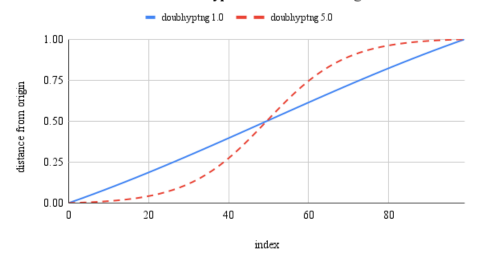


Figure 3: Double sided hyperbolic tan stretching

- doubhyptng stretching

$$s = \frac{1}{2} \frac{1.0 + \tanh(a(\zeta/I - 1.0/2.0))}{\tanh(a/2.0)}$$

hyptng stretching

$$s = \frac{1.0 + \tanh(a(\zeta/I - 1.0))}{\tanh(a)}$$

Hyperbolic tan stretching

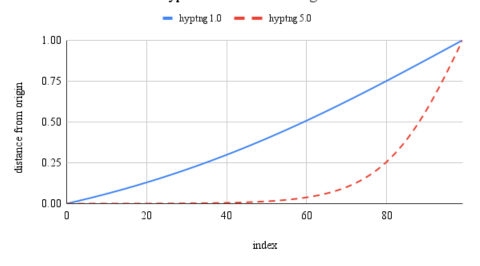


Figure 4: Single sided hyperbolic tan stretching

- douhbypsin stretching

$$s = \frac{1}{2} \frac{1.0 + \sinh(a(\zeta/I - 1.0/2.0))}{\sinh(a/2.0))}$$

Double sided hyperbolic sin stretching

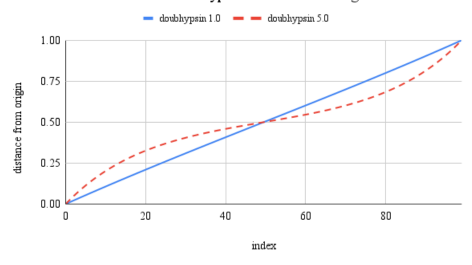


Figure 5: Double sided hyperbolic sin stretching

- hypsin

$$s = \frac{1.0 + \sinh(a(\zeta/I - 1.0))}{\sinh(a)}$$

Hyperbolic tan stretching

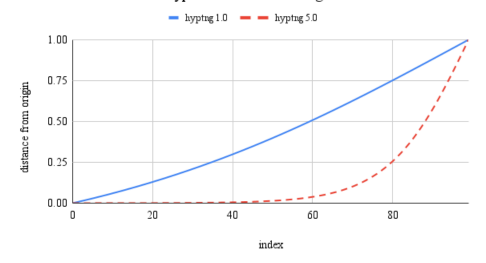


Figure 6: Single sided hyperbolic sin stretching

The negative sine distribution has three types, where the type can be selected by the stretching factor (setting it to 1, 2 or something else). These were determined by fitting negative sine distributions raised to different powers and tailored for a specific application (magnetic reconnection simulations) and are there as a demonstration how the user can also add their own fitted spacing functions. The other functions follow the conventional definitions. The functions shown above, for two different parameters (1.0 and 5.0) are shown in Figures 1, 2, 3, 4, 5 and 6 for reference.

The user can easily add their own stretching functions if they follow the principle in which the other functions are handled, especially the negative sine distributions which are based on fitting.

With the mesh being properly defined, the code is run by simply invoking it, without any arguments:

```
python spherical_main.py
```

and afterwards, the output file in the native format must be transformed to the GAMBIT file:

```
./write.exe 2d [input_file] [output_file]
for 2d domains, and:
./write.exe 3d [input_file] [output_file]
```

for 3d domains.

Keep in mind that the 3d domains (especially with blocking) were not yet properly tested.

Without blocking, the original example given in the code with the comments can be simply followed and modified according to user's wishes. The output file format will have boundaries marked as x0, x1, y0, y1 (, z0, z1). This all can be modified in the fortran source code.

Blocking is also possible, where several blocks of different mesh types can be added together to create one mesh. Adding multiple blocks is shown in the commented-out portion of the code (follow the comments in the code). Here, however, the user must be cautious with each block, re-indexing of the nodes must be done and connectivity adjusted. This is currently done through a double for-loop searching for the nodes residing at the same position, which can be very computationally heavy for finer meshes. In case for finer meshes, for this reason, avoid blocking if possible.

For blocking, the following instructions must be followed after adding each block:

- Define the new block params (refinement, boundaries, block_type)
- 2. Add the block by:

```
nodes_add, elements_add = add_block(block_type, params)
nodes = nodes + nodes_add
```

```
elements = elements + elements_add
```

3. Delete repeated nodes (on the boundaries of the touching blocks) by:

```
nodes, elements = delete_repeated_nodes(nodes, elements, tol)
```

note here that the tolerance can be adjusted depending on the scale of the mesh you are building. This tolerance is used to determine whether two points overlap or not, so it should be much smaller than the scale of the smallest mesh cell.

4. Recompute the starting node and element index if you wish to not overwrite the previous block:

```
start_n = len(nodes)
start_e = len(elements)
```

Please also note that there is no automatic gap detection. If the mesh blocks do not overlap according to their coordinates, the resulting mesh will be degenerate. Contact us in case of troubleshooting or concerns.

2. Spherical grid based on a surface mesh

The second mode can create a spherical 3D mesh by taking a surface spherical mesh and extending it radially outwards. By default, surface triangles are used, but also quad-based meshes are supported. For this functionality, use the spherical_main.py script in the spherical folder (for the quad-based surface meshes, use the scripts in quad_examples.

This mode requires an input spherical surface mesh. This mesh must be in the .ply (Stanford) format to be legible to the script. Such meshes can be easily generated by, for example, Blender. Some examples for icospheric surface meshes are included in the folder example_surface_meshes, including their blender setup files .blend and the output .ply formats read by the mesher. The input files can include double points or disconnected points (which is frequently the case for Blender-generated meshes) - all of this is corrected for in the code.

The principle is simple. The surface connectivity (in the Stanford format) is preserved, but radially extended outwards. From the subsequent layers, the 3D elements are formed (so for example, a triangular surface mesh creates prism elements while a quadrilateral surface mesh creates hexagonal elements). Thus, at least one additional layer is needed, in which case we create a mesh of one layer of 3D elements.

The radial spacing between the layers is prescribed in ReturnSpacing_idx(i). Here, depending on which layer is added (layer + 1 = i), the radial distance can be defined. In this function, it is defined as an absolute radius, but also spacing can be used, in which case one should return Rs[i] instead of the spacing variable. The number of layers that will be added can be independent of the Rs array and is defined in n_layers variable.

Keep in mind that the first layer is added separately, since the procedure is a bit different (the elements and connectivity have to be formed from scratch). Thus, for the first layer, define the spacing in the respective delta_r variable as commanded by the comments (read the comments).

Finally, set the overall output path (without the name) in the variable path. To start the program, follow the syntax:

```
python spherical_main.py [surface_ply_file] [output_file]
```

The arguments will be printed onscreen by the code as a way of verification. In addition, also the spacing of the layers will be printed (apart from the very first layer) so that the user can check whether the correct mesh is being built.

This procedure produces a native-formatted file. To convert this file to one that has the information formatted according to the GAMBIT neutral format, follow this syntax:

```
./write.exe pr [input_file] [output_file]
```

where the output file should have the .neu extension. The pr argument tells the code that prisms will be created. The boundaries will be marked inlet (the inner sphere) and outlet (the outer sphere), but this can be easily modified by the user. Once can either do this directly in the output file, or in the write.f90 and recompiling.

Here, we only discussed generation of a prism mesh (that is why the pr argument) where the surface mesh is made out of triangles, but in the quad_example folder, one can find this procedure also for a quad-based surface mesh (thus, giving hexagonal elements). In that case, still do use the pr when invoking the fortran function, but use the correct executable (in the respective file) so that the fortran writer knows to write hexagonal elements.

Again, in case of any further questions or concerns, contact us directly.

This code is open source and free to use for anyone. It has been so far used for:

- US Air Force award NO. FA9550-18-1-0093
- C1 project TRACESpace
- HPM rapid hypersonic modelling software

Please cite the code accordingly in case you use it, inform us know about the application and let us know of your suggestions for further development.