



This document describes *manifem*, a software for solving partial differential equations through the finite element method. The name comes from “finite elements on manifolds”. *manifem* has been designed with the goal of coping with very general meshes, in particular meshes on Riemannian manifolds, even manifolds which cannot be embedded in  $\mathbb{R}^3$ , like the torus  $\mathbb{R}^2/\mathbb{Z}^2$ . Also, *manifem* has been written with the goal of being conceptually clear and easy to read.

*manifem* is written in C++ and uses the Standard Template Library (STL). It uses Eigen for storing matrices and solving systems of linear equations, see [http://eigen.tuxfamily.org/index.php?title=Main\\_Page](http://eigen.tuxfamily.org/index.php?title=Main_Page).

Some of the drawings in this manual have been obtained by using *gmsh*, see ....

To use *manifem*, go to <https://github.com/cristian-barbarosie/manifem>, copy all .cpp and .h files to some directory in your computer, together with the Makefile. Then try `make run-1.1` for the example in paragraph 1.1, `make run-1.3` for the example in paragraph 1.3, and so on.

This manual is divided in six sections. The first two describe *manifem* from the viewpoint of the user who wants to use it as a black box, without bothering with too much detail. In particular, they should be accessible to readers who have some knowledge of C++ but are not necessarily experts in C++ programming. The third section gives more details about how cells and meshes are implemented in *manifem*. It is useful for users who want finer control on the mesh, e.g. for implementing their own remeshing algorithms. The fourth section dwells on fields and functions, while the fifth describes how finite elements work in *manifem*. The sixth section gives technical details, mainly for those interested in developing and extending *manifem*.

Do not read paragraphs marked as “outdated”.

TO DO :

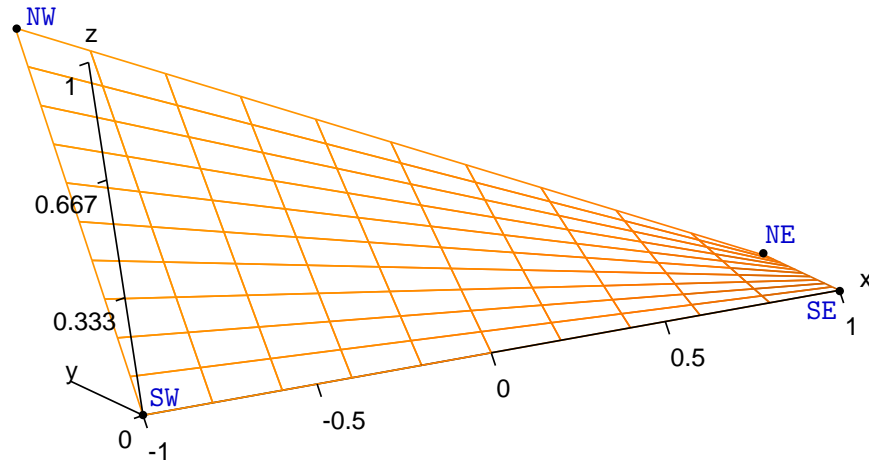
- mesh a triangle, starting from its three sides
- mesh a tetrahedron, starting from its four faces
- build a mesh of cubes (done through cartesian product, but we would like to build just one cube based on its six faces and be able to join cubes afterwards)
- implement Dirichlet boundary conditions
- solve the system of linear equations
- visualize the solution of the PDE
- define a Gauss quadrature on segments
- define a Lagrange P1 finite element on triangles
- eliminate memory leaks

# 1. General description

The first two sections of this manual describe *manil3d* from the viewpoint of the user who is not interested in too much technical detail about its implementation. In this section, we show how to build meshes on (rather simple) domains in  $\mathbb{R}^2$  and  $\mathbb{R}^3$  and how to solve (rather simple) partial differential equations.

## 1.1. An elementary example

In this paragraph, we show how to build a rectangular mesh on a surface in  $\mathbb{R}^3$  and then compute the integral of a given function. See paragraph 1.3 for a purely two-dimensional example. See paragraph 1.9 for a more complex example of solving a partial differential equation.



```
#include <iostream>
#include "Mesh.h"
using namespace std;

int main () {

    // we choose our (geometric) space dimension :
    auto RR3 = Manifold::euclid (3);
    // xyz is a map defined on our future mesh with values in RR3 :
    auto & xyz = RR3.coordinate_system ("Lagrange degree one");

    // We can extract components of xyz using the [] operator
    auto & x = xyz[0], & y = xyz[1], & z = xyz[2];

    // Let's build a rectangular mesh. First, the four corners :
    auto & SW = Cell::point ("SW"); x == -1; y == 0; z == 0;
    auto & SE = Cell::point ("SE"); x == 1; y == 0; z == 0;
    auto & NE = Cell::point ("NE"); x == 1; y == 1; z == 0;
    auto & NW = Cell::point ("NW"); x == -1; y == 1; z == 1;

    // we access the coordinates of a point using the () operator :
    cout << "coords of NW : " << x(NW) << " " << y(NW) << " " << z(NW) << endl;
```

```

// Now build the four sides of the rectangle :
auto & south = Mesh::segment ( SW, SE, 10 );
auto & east  = Mesh::segment ( SE, NE, 10 );
auto & north = Mesh::segment ( NE, NW, 10 );
auto & west  = Mesh::segment ( NW, SW, 10 );

// And now the rectangle :
auto & rect_mesh = Mesh::rectangle ( south, east, north, west );

// We may want to visualize the resulting mesh.
// Here is one way to export the mesh in the "msh" format :
rect_mesh.export_msh ("rectangle.msh");

// Let's define a symbolic function to integrate
auto & f = x*x+1/(5+y);

// and compute its integral on the rectangle,
// using Gauss quadrature with 9 points :
auto & integ = Integrator::gauss ("Q9");
cout << "integral = " << f.integrate ( rect_mesh, integ ) << endl;

}

```

In the present section, we often use the keyword `auto` in order to avoid declaring explicitly the type. This makes things easier for the final user. However, in the other examples presented in this section, we also include the type of the object as a comment, for those users who want to understand more of the underlying details. See paragraph 3.1 for an explanation about the ampersand following `auto`.

The string argument of `Cell::point` is an optional name.

See paragraph 1.3 for an explanation about why we build the rectangle based on its four sides rather than on its four vertices.

Note that in this example we do not have exact control on the surface being meshed. It is defined rather vaguely by the position of the four vertices. See paragraphs 2.6 and 2.7 for ways to precisely define a submanifold in  $\mathbb{R}^3$  and mesh it.

## 1.2. Cells and meshes

In *manifold*, all basic constituents of meshes are called “cells”. Points are zero-dimensional cells, segments are one-dimensional cells, triangles are two-dimensional cells, and so on.

A mesh is roughly a collection of cells of the same dimension. Internally, *manifold* keeps lists of cells of lower dimension, too. For instance, the mesh built in paragraph 1.1 is roughly a list of two-dimensional cells (quadrilaterals), but lists of segments and points are also kept.

A cell of dimension higher than zero is defined by its boundary, which in turn is a mesh of lower dimension. The boundary of a segment is a (zero-dimensional) mesh made of two points. The boundary of a triangle is a one-dimensional mesh made of three segments. Thus, a segment is essentially a pair of points, a triangle is essentially a triplet of segments, and so on.

Cells and meshes are oriented. An orientation of a mesh is just an orientation for each of its component cells (of course these orientations must be mutually compatible).

Although this is not how the orientation is implemented internally (see paragraph 6.4), an oriented point can be conceived simply as a point with a sign attached (1 or -1). The orientation of a cell of dimension higher than zero is given by an orientation of its boundary, which is a lower-dimensional mesh.

Thus, an oriented segment is essentially a pair of points, one of which has a -1 attached, the other having a 1. We call the former “base” and the latter “tip”. These signs are related to integration of functions along that segment. The integral of a function of one variable is equal to the value of the primitive function at one end of the segment minus the value of the primitive at the other end.

An oriented triangle is essentially a triplet of segments, each one with its own orientation. The orientations must be compatible to each other in the sense that each vertex must be seen as positive by one of the segments and as negative by another one. An oriented tetrahedron can be identified with four triangles, each one with its own orientation. In such a tetrahedron, each segment must be seen as positive by one of the triangles and as negative by another one.

Paragraph 3.6 gives more details about orientation of cells.

See paragraph 6.1 for some implementation details.

Cells are topological entities; they carry no geometric information. In particular, points do not have coordinates. Coordinates are stored externally, see paragraph 1.7.

### 1.3. Joining meshes

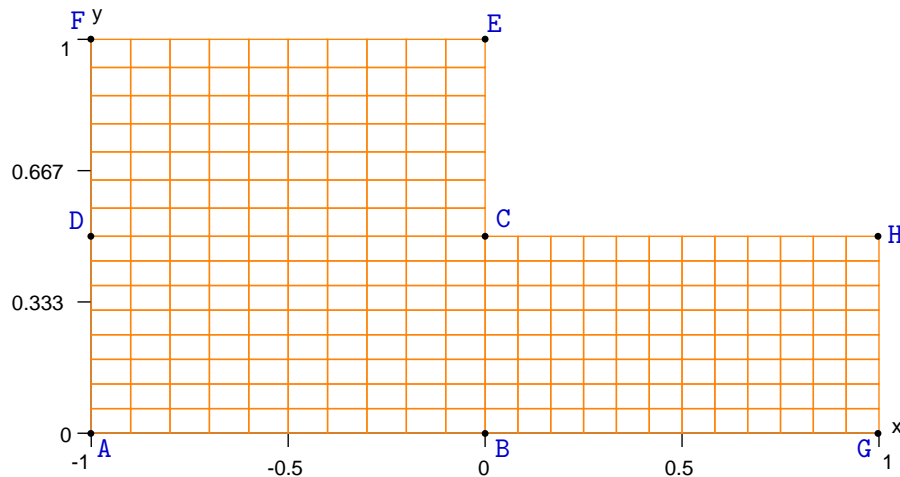
The example in paragraph 1.1 could have been shortened had we used the overloaded version of `Mesh::rectangle` which accepts the four corners as arguments. This overloaded version exists in *manifold*, but we prefer to build meshes in a structured way, first corners, then sides and then the plane region. This has the advantage that one can build more complex meshes from simple components. For instance, one can build an L-shaped mesh by joining three rectangular meshes :

```
auto RR2 = Manifold::euclid (2); // Manifold RR2
auto & xy = RR2.coordinate_system ("Lagrange degree one");
// FunctionOnMesh::Function & xy
auto & x = xy[0], & y = xy[1]; // FunctionOnMesh::Function & x, & y
auto & A = Cell::point ("A"); x == -1; y == 0; // Cell & A
auto & B = Cell::point ("B"); x == 0; y == 0;
auto & C = Cell::point ("C"); x == 0; y == 0.5;
auto & D = Cell::point ("D"); x == -1; y == 0.5;
auto & E = Cell::point ("E"); x == 0; y == 1;
auto & F = Cell::point ("F"); x == -1; y == 1;
auto & G = Cell::point ("G"); x == 1; y == 0;
auto & H = Cell::point ("H"); x == 1; y == 0.5;
auto & AB = Mesh::segment ( A, B, 10 ); // Mesh & AB
auto & BC = Mesh::segment ( B, C, 8 );
auto & CD = Mesh::segment ( C, D, 10 );
auto & DA = Mesh::segment ( D, A, 8 );
auto & CE = Mesh::segment ( C, E, 7 );
auto & EF = Mesh::segment ( E, F, 10 );
auto & FD = Mesh::segment ( F, D, 7 );
auto & BG = Mesh::segment ( B, G, 12 );
```

```

auto & GH = Mesh::segment ( G, H, 8 );
auto & HC = Mesh::segment ( H, C, 12 );
auto & ABCD = Mesh::rectangle ( AB, BC, CD, DA ); // Mesh & ABCD
auto & CEFD = Mesh::rectangle ( CE, EF, FD, CD.reverse() );
auto & BGHC = Mesh::rectangle ( BG, GH, HC, BC.reverse() );
auto & L_shaped = Mesh::join ( ABCD, CEFD, BGHC ); // Mesh & L_shaped

```



Note that, in *manim*, cells and meshes are oriented. To build CEFD one must use not CD but its reverse; to build BGHC one must use not BC but its reverse.

Note also that if we define the rectangles based on their vertices instead of their sides, the `Mesh::join` method does not work properly. For instance, the two rectangles defined by

```

auto & ABCD = Mesh::rectangle ( A, B, C, D, 10, 8 );
auto & CEFD = Mesh::rectangle ( C, E, F, D, 7, 10 );

```

cannot be joined\* because the side CD of ABCD has nothing to do with the side DC of CEFD. These two sides are one-dimensional meshes both made of 10 segments but with different interior points (only C and D are shared) and different segments. In contrast, CD and `CD.reverse()` share the same 11 points and the same 10 segments (reversed).

See paragraph 3.2 for a more complex use of `Mesh::join`.

Incidentally, note that the `Mesh::rectangle` method accepts any position for the vertices. Thus, you can use it to build any quadrilateral; the inner vertices' coordinates are simply interpolated from the coordinates of vertices on the boundary, as shown paragraphs 1.5 and 2.1. This can be done even in more than two (geometric) dimensions, like in paragraphs 1.1 and 2.7.

See also paragraph 6.5.

## 1.4. Triangular meshes

We can also build meshes on triangular domains and join them as we wish :

```

auto & A = Cell::point ("A"); x == -1. ; y == 0. ; // Cell & A

```

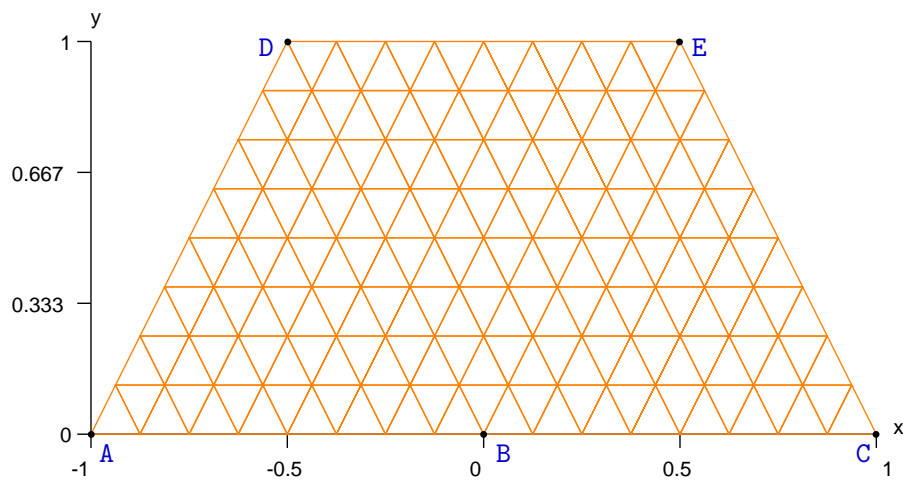
---

\* Actually, they can be joined but the resulting mesh will have a crack along CD – probably not what the user wants.

```

auto & B = Cell::point ("B"); x == 0. ; y == 0. ;
auto & C = Cell::point ("C"); x == 1. ; y == 0. ;
auto & D = Cell::point ("D"); x == -0.5; y == 1. ;
auto & E = Cell::point ("E"); x == 0.5; y == 1. ;
auto & AB = Mesh::segment ( A, B, 8 ); // Mesh & AB
auto & BC = Mesh::segment ( B, C, 8 );
auto & AD = Mesh::segment ( A, D, 8 );
auto & BD = Mesh::segment ( B, D, 8 );
auto & BE = Mesh::segment ( B, E, 8 );
auto & CE = Mesh::segment ( C, E, 8 );
auto & ED = Mesh::segment ( E, D, 8 );
auto & ABD = Mesh::triangle ( AB, BD, AD.reverse() ); // Mesh & ABD
auto & BCE = Mesh::triangle ( BC, CE, BE.reverse() );
auto & BED = Mesh::triangle ( BE, ED, BD.reverse() );
auto & three_tri = Mesh::join ( ABD, BCE, BED ); // Mesh & three_tri

```



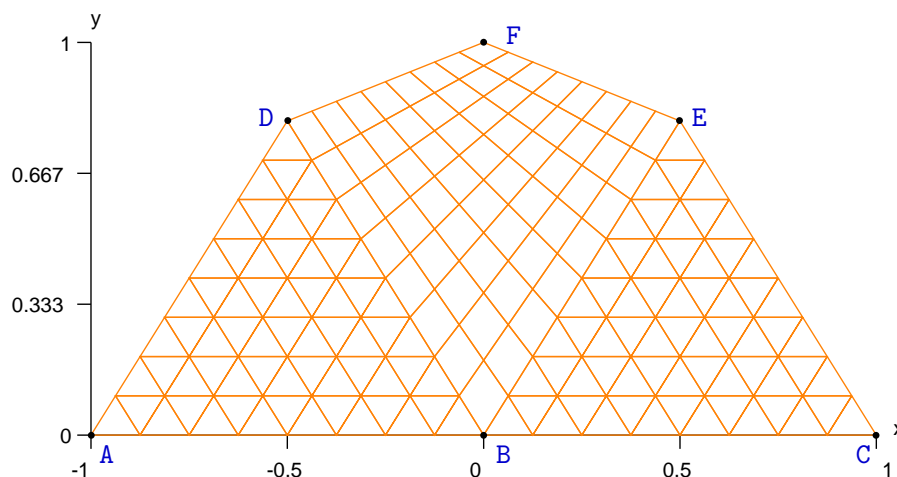
## 1.5. Mixing triangles and rectangles

It is possible to have triangles and quadrilaterals mixed in the same mesh :

```

auto & A = Cell::point ("A"); x == -1. ; y == 0. ; // Cell & A
auto & B = Cell::point ("B"); x == 0. ; y == 0. ;
auto & C = Cell::point ("C"); x == 1. ; y == 0. ;
auto & D = Cell::point ("D"); x == -0.5; y == 0.8;
auto & E = Cell::point ("E"); x == 0.5; y == 0.8;
auto & F = Cell::point ("F"); x == 0. ; y == 1. ;
auto & AB = Mesh::segment ( A, B, 8 ); // Mesh & AB
auto & BC = Mesh::segment ( B, C, 8 );
auto & AD = Mesh::segment ( A, D, 8 );
auto & BD = Mesh::segment ( B, D, 8 );
auto & BE = Mesh::segment ( B, E, 8 );
auto & CE = Mesh::segment ( C, E, 8 );
auto & EF = Mesh::segment ( E, F, 8 );
auto & FD = Mesh::segment ( F, D, 8 );
auto & ABD = Mesh::triangle ( AB, BD, AD.reverse() ); // Mesh & ABD
auto & BCE = Mesh::triangle ( BC, CE, BE.reverse() );
auto & BEFD = Mesh::rectangle ( BE, EF, FD, BD.reverse() );
auto & two_tri_one_rect = Mesh::join ( ABD, BEFD, BCE ); // Mesh & two_tri_one_rect

```



Paragraph 2.2 shows another example of mixed mesh.

## 1.7. Fields (outdated)

*ManiE* uses `NumericField` objects to store values associated to points (or to other cells). For instance, coordinates of points can be stored in a `NumericField` (see the example in paragraph 1.1). Other values, like the solution of a PDE, can also be stored in a `NumericField`.

`NumericField` objects can live not only on points. For instance, if we work with curved finite elements we may want to store, for each segment, information about the curvature of the segment. We may keep the coordinates of the middle point (a pair of values in 2D, a triplet of values in 3D). Such a `NumericField` should be declared (before the creation of any segment) like this :

```
int d = 2; // or 3
auto middle_coord = NumericField::multi_dim("size", d, "lives on", "segments");
```

After this declaration, every segment will be created with a heap space for `d` real numbers. These values can then be accessed as `middle_coord(seg)`.

Another example is when we have a coefficient which depends on the space variable but is piecewise constant. We may model such a function by declaring a `NumericField` which lives on two-dimensional cells (if the topological dimension is 2) :

```
auto coef = NumericField::one_dim("lives on", "2d cells");
```

or on three-dimensional cells (if the topological dimension is 3) :

```
auto coef = NumericField::one_dim("lives on", "3d cells");
```

In contrast, a function of the space variable can be modelled by a `FunctionOnMesh` object, see paragraph 1.8.

We should be careful to declare all `NumericField` objects before building the meshes. This is because, when creating a cell, *maniE* reserves a heap space for keeping values, accordingly to the existing `NumericField` objects. If we create a point, say, `P`, and afterwards declare a `NumericField` object, say, `x`, which lives on points, then any reference like `x(P)` will produce an error.

As mentioned above, the `operator()` is used to access the value associated to a particular cell. In contrast, the `operator[]` is used to access a particular coordinate of a multi-dimensional field. The opposite operation, concatenation of two or more fields, can be achieved through the `&&` operator. For instance, the two pieces of code below are equivalent :

```
auto xy = NumericField::multi_dim ("size", 2, "lives on", "points");
auto x = xy[0], y = xy[1];
```

and

```
auto x = NumericField::one_dimension ("lives on", "points");
auto y = NumericField::one_dimension ("lives on", "points");
auto xy = x && y;
```

## 1.8. Functions, derivation and integration (outdated)

Unlike `NumericFields`, `FunctionOnMesh` objects are used to model functions which vary in a continuum (a cell or a whole mesh). They might be just variables like `x` and `y`, or arithmetic expressions of such variables.

They can be differentiated; for instance, the expression `(xx*yy-1/xx).deriv(yy)` will produce the result `xx`.

They can also be integrated with the use of an `Integrator` object, like in the example in paragraph 1.1.

## 1.9. Partial differential equations (outdated)

Let's consider again the rectangular mesh from paragraph 1.1, this time with only two geometric dimensions.

```
Mesh::intended_dimension = 2; // topological dimension
Mesh::initialize();
auto xy = NumericField::multi_dim ("size", 2, "lives on", "points");
auto x = xy[0], y = xy[1];
auto u = NumericField::one_dim ("lives on", "points");
auto xxyy = FunctionOnMesh::from_field ( xy, "Lagrange degree one" );
auto xx = xxyy[0], yy = xxyy[1];
auto SW = Cell::point ("SW"); x(SW) = -1.1; y(SW) = 0.3;
auto SE = Cell::point ("SE"); x(SE) = 1; y(SE) = 0;
auto NE = Cell::point ("NE"); x(NE) = 1; y(NE) = 1;
auto NW = Cell::point ("NW"); x(NW) = -1; y(NW) = 1;
auto south = Mesh::segment ( SW, SE, 4, xy );
auto east = Mesh::segment ( SE, NE, 2, xy );
auto north = Mesh::segment ( NE, NW, 4, xy );
auto west = Mesh::segment ( NW, SW, 2, xy );
auto rect_mesh = Mesh::rectangle ( south, east, north, west, xy );
```

There is a new `NumericField`, `u`, which is meant to hold the values (at vertices) of the final solution of the partial differential equation.

In order to introduce the PDE, by means of a variational formulation, we declare an unknown function `uu`, related to the field `u`, and a test function `w` :



```

auto uu = FunctionOnMesh::unknown ( u, "Lagrange degree one");
auto w = FunctionOnMesh::test ( uu );

```

We are now in a position to define the variational formulation of our problem, by means of methods `.deriv` and `.integrate`, using a syntax mimicking the mathematical notation of variational calculus :

```

auto var_pb =
  ( uu.deriv(xx)*w.deriv(xx) + uu.deriv(yy)*w.deriv(yy) ) .integrate(rect_mesh)
  == w.integrate(rect_mesh) + w.integrate(south);

```

Dirichlet-like boundary conditions are declared as

```

var_pb.prescribe_on (north); uu == 0.;      w == 0.;
var_pb.prescribe_on (east); uu == xx*(1.-yy); w == 0.;
var_pb.prescribe_on (west); uu == 0.;      w == 0.;

```

We then declare a finite element (Lagrange Q1), a method of integrating functions (Gauss quadrature with 3 nodes on segments and 9 nodes on rectangles)

```

auto fe = FiniteElement::Lagrange ("Q1");
fe.set_integrator ("on cells of dimension", 1, "gauss", 3, "nodes");
fe.set_integrator ("on cells of dimension", 2, "gauss", 9, "nodes");

```

The following line of code triggers a complex process of discretizing the variational formulation `var_pb` and solving the associated linear system :

```

fe.discretize ( var_pb );

```

## 2. Meshes and manifolds

The first two sections of this manual describe *manil* from the viewpoint of the user who is not interested in too much technical detail about its implementation. This section describes several examples of (more complex) meshes, some of them built on specific manifolds.

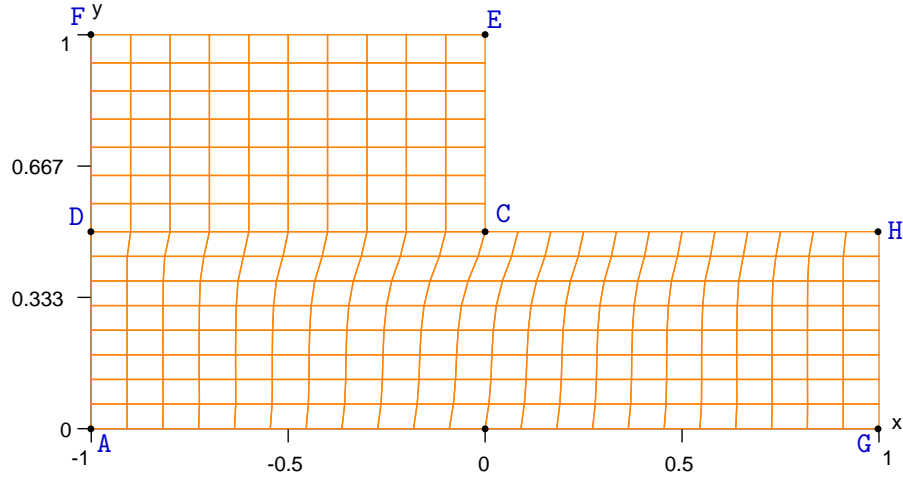
Paragraphs 2.1, 2.2, 2.9 and 2.10 show examples of domains in  $\mathbb{R}^2$ . Paragraphs 2.4 and 2.5 deal with one-dimensional meshes in  $\mathbb{R}^2$  (curves). Paragraphs 2.6 – 2.8 focus on two-dimensional meshes in  $\mathbb{R}^3$  (surfaces).

Paragraphs 2.1 – 2.10 are about building meshes of regular shape and then joining them together like patches, while paragraphs 2.11 – ... show how to directly mesh a domain starting from its boundary only.

### 2.1. Joining segments and rectangles

Here is another way of building the same L-shaped mesh as in paragraph 1.3 :

```
auto & RR3 = Manifold::euclid (2); // Manifold & RR3
auto & xy = RR3.coordinate_system ("Lagrange degree one");
// FunctionOnMesh::Function & xy
auto & x = xy[0], & y = xy[1]; // FunctionOnMesh::Function & x, & y
auto & A = Cell::point ("A"); x == -1; y == 0; // Cell & A
auto & C = Cell::point ("C"); x == 0; y == 0.5;
auto & D = Cell::point ("D"); x == -1; y == 0.5;
auto & E = Cell::point ("E"); x == 0; y == 1;
auto & F = Cell::point ("F"); x == -1; y == 1;
auto & G = Cell::point ("G"); x == 1; y == 0;
auto & H = Cell::point ("H"); x == 1; y == 0.5;
```



```
auto & AG = Mesh::segment ( A, G, 22 ); // Mesh & AG
auto & GH = Mesh::segment ( G, H, 8 );
auto & HC = Mesh::segment ( H, C, 12 );
auto & CD = Mesh::segment ( C, D, 10 );
auto & HD = Mesh::join ( HC, CD ); // Mesh & HD
```

```

auto & DA = Mesh::segment ( D, A, 8 );
auto & CE = Mesh::segment ( C, E, 7 );
auto & EF = Mesh::segment ( E, F, 10 );
auto & FD = Mesh::segment ( F, D, 7 );
auto & AGHD = Mesh::rectangle ( AG, GH, HD, DA ); // Mesh & ABCD
auto & CEFD = Mesh::rectangle ( CE, EF, FD, CD.reverse() );
auto & L_shaped = Mesh::join ( AGHD, CEFD ); // Mesh & L_shaped

```

The only difference between the two meshes is a slight distortion in the lower half of the domain, due to the non-uniform distribution of the vertices along HD.

See also paragraph 6.5.

## 2.2. Triangular meshes on rectangles

On a rectangular domain, we can build a mesh of triangles by adding an option to the `Mesh::rectangle` method. For instance, in the example in paragraph 1.3, if we rewrite the definition of BGHC as

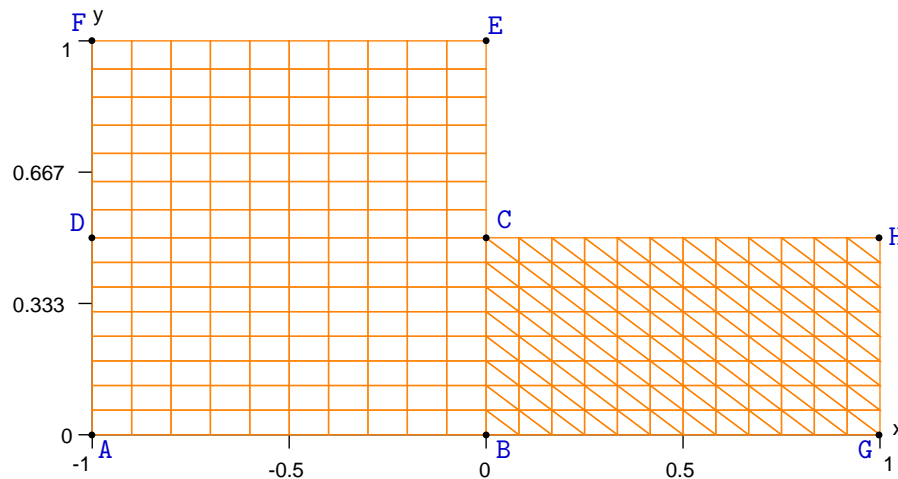
```

auto & BGHC = Mesh::rectangle ( BG, GH, HC, BC.reverse(), tag::with_triangles );

```

we get the mesh shown below.

See paragraph 6.8 for more details about tags.



If we give the sides of the rectangle in a different order, like in

```

auto & BGHC = Mesh::rectangle ( GH, HC, BC.reverse(), BG, tag::with_triangles );

```

the rectangles will be cut along the other diagonal (check it yourself).

The mesh in paragraph 1.4 could have been built like this :

```

auto & ABD = Mesh::triangle ( AB, BD, AD.reverse() ); // Mesh & ABD
auto & BCED = Mesh::rectangle ( CE, ED, BD.reverse(), BC, tag::with_triangles );
auto & one_tri_one_rect = Mesh::join ( ABD, BEFD, BCE ); // Mesh & one_tri_one_rect

```

## 2.3. Cartesian product of meshes

[...]

## 2.4. A manifold defined as a level set in $\mathbb{R}^2$

*ManiCER* allows one to define manifolds and submanifolds, and this feature may be used to build domains of the desired shape.

Until now, we have only met the trivial Euclidian manifold, defined as `Manifold::euclid (n)` or as a cartesian product of Euclidian manifolds. One can define a submanifold in terms of an implicit equation, that is, as a level set, using the method `implicit` of the Euclidian manifold. The code below introduces a one-dimensional submanifold of  $\mathbb{R}^2$  (a hiperbola).

```
auto RR2 = Manifold::euclid (2); // Manifold RR2
auto & xy = RR2.coordinate_system ("Lagrange degree one");
// FunctionOnMesh::Function & xy
auto & x = xy[0], & y = xy[1]; // FunctionOnMesh::Function & x, &y
auto & hiperbola = RR2.implicit ( x*y - 1. ); // Manifold & hiperbola
auto & A = Cell::point(); x == 0.5; y == 2.; // Cell & A
auto & B = Cell::point(); x == 3; y == 0.3333333333333333;
auto & arc_of_hiperbola = Mesh::segment ( A, B, 15 );
// arc_of_hiperbola.draw_ps ("hiperbola.eps");
arc_of_hiperbola.export_msh ("hiperbola.msh");
```

Note that the vertices are not perfectly uniformly distributed along the curve because they are obtained as projections of points uniformly distributed along AB onto the hiperbola manifold.

Note also that, when defining individual points A and B we must be careful to set coordinates x and y within the hiperbola manifold. As an alternative, we might explicitly project them onto the hiperbola like this :

```
auto & P = Cell::point(); x == 0.6; y == 2.1;
hiperbola.project(P);
```

In contrast, the `Mesh::segment` factory function builds points in the RR2 space and then projects them onto the hiperbola without the user's assistance.

The projection is done by applying a few steps of Newton's method for undetermined (systems of) equations.

## 2.5. A circle defined by four curved segments

We can define several arcs of curve and join them, thus obtaining a closed curve :

```
auto RR2 = Manifold::euclid (2); // Manifold RR2
auto & xy = RR2.coordinate_system ("Lagrange degree one");
// FunctionOnMesh::Function & xy
auto & x = xy[0], & y = xy[1]; // FunctionOnMesh::Function & x, &y
auto & circle_manifold = RR2.implicit ( x*x + y*y - 1. ); // Manifold & circle_manifold
auto & N = Cell::point(); x == 0.; y == 1.; // Cell & A
auto & W = Cell::point(); x == -1.; y == 0.;
auto & S = Cell::point(); x == 0.; y == -1.;
auto & E = Cell::point(); x == 1.; y == 0.;
auto & NW = Mesh::segment ( N, W, 10 ); // Mesh & NW
auto & WS = Mesh::segment ( W, S, 10 );
auto & SE = Mesh::segment ( S, E, 10 );
```

```

auto & EN = Mesh::segment ( E, N, 10 );
auto & circle = Mesh::join ( NW, WS, SE, EN ); // Mesh & circle;
// circle.draw_ps ("circle.eps");
circle.export_msh ("circle.msh");

```

Again, the vertices are not perfectly uniformly distributed along the circle because they are obtained as projections (on the circle) of points along segments NW, WS and so forth.

Note that applying `Mesh::join` to four segments is very different from applying `Mesh::rectangle` to the same segments, see paragraph 2.9.

## 2.6. A hemisphere defined by three curved triangles

Let's look at a surface in  $\mathbb{R}^3$  :

```

auto RR3 = Manifold::euclid (3);
// xyz is a map defined on our future mesh with values in RR3 :
auto & xyz = RR3.coordinate_system ("Lagrange degree one");
// we can extract components of xyz using the [] operator :
auto & x = xyz[0], & y = xyz[1], & z = xyz[2];

auto & sphere = RR3.implicit ( x*x + y*y + z*z - 1. );
// Manifold & sphere

// let's mesh half of a sphere
auto & E = Cell::point(); x == 1.; y == 0.; z == 0.;
auto & N = Cell::point(); x == 0.; y == 1.; z == 0.;
auto & W = Cell::point(); x == -1.; y == 0.; z == 0.;
auto & S = Cell::point(); x == 0.; y == -1.; z == 0.;
auto & up = Cell::point(); x == 0.; y == 0.; z == 1.;
int n = 15;
auto & EN = Mesh::segment ( E, N, n );
auto & NW = Mesh::segment ( N, W, n );
auto & WS = Mesh::segment ( W, S, n );
auto & SE = Mesh::segment ( S, E, n );
auto & upE = Mesh::segment ( up, E, n );
auto & upN = Mesh::segment ( up, N, n );
auto & upW = Mesh::segment ( up, W, n );
auto & upS = Mesh::segment ( up, S, n );

// now four triangles
auto & ENup = Mesh::triangle ( EN, upN.reverse(), upE );
auto & NWup = Mesh::triangle ( NW, upW.reverse(), upN );
auto & WSup = Mesh::triangle ( WS, upS.reverse(), upW );
auto & SEup = Mesh::triangle ( SE, upE.reverse(), upS );

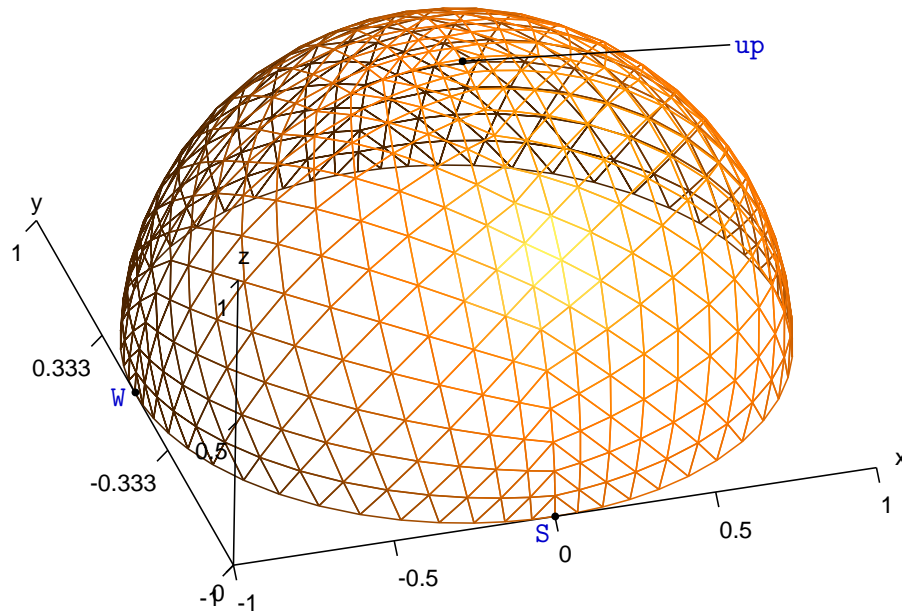
// and finally join the triangles :
auto & hemisphere = Mesh::join ( ENup, NWup, WSup, SEup );

hemisphere.export_msh ("hemisphere.msh");

```

Again, when we define individual points E, N, W and S we must be careful to provide coordinates within the sphere (or project them explicitly as in paragraph 2.7). In contrast, the `Mesh::segment`, `Mesh::rectangle` and `Mesh::triangle` factory functions build

points in the surrounding space ( $\mathbb{R}^2$  or  $\mathbb{R}^3$ ) and then project them onto the current manifold without the user's assistance.



Again, we notice that, within each triangle (ENup, NWup and so forth), the distribution of the vertices is not perfectly uniform, for the same reason as in paragraphs 2.4 and 2.5.

Note that, when we build the segments WS, upS and so on, we know that those segments will be (polygonal approximatin of) arcs of circle on the sphere. This is so due to the particular geometry of our manifold (we know that the projection of a straight line segment on the sphere is an arc of circle of radius equal to the radius of the sphere); the shape of these segments is less clear in other examples (see e.g. paragraph 2.7).

See paragraphs 2.11 and ...for another way of meshing a domain, having its boundary as starting point.

## 2.7. A more complex surface

If the surface is more “bumpy”, we must use smaller patches in order to get a mesh of good quality. Below we use 12 rectangles to get a deformed hemisphere.

```
auto & deformed_sphere = RR3.implicit ( x*x + y*y + z*z + 1.5*x*y*z - 1. );

// let's mesh a hemisphere (much deformed)
auto & E = Cell::point(); x == 1.; y == 0.; z == 0.;
auto & N = Cell::point(); x == 0.; y == 1.; z == 0.;
auto & W = Cell::point(); x == -1.; y == 0.; z == 0.;
auto & S = Cell::point(); x == 0.; y == -1.; z == 0.;
auto & up = Cell::point(); x == 0.; y == 0.; z == 1.;
auto & mWS = Cell::point(); x == -1.; y == -1.; z == 0.;
deformed_sphere.project ( mWS ); // midway between W and S
auto & mSup = Cell::point(); x == 0.; y == -1.; z == 1.;
deformed_sphere.project ( mSup ); // midway between S and up
auto & mWSup = Cell::point(); x == 1.; y == 1.; z == 1.;
deformed_sphere.project ( mWSup ); // somewhere between W, S and up
// ... and so forth ...
```

```

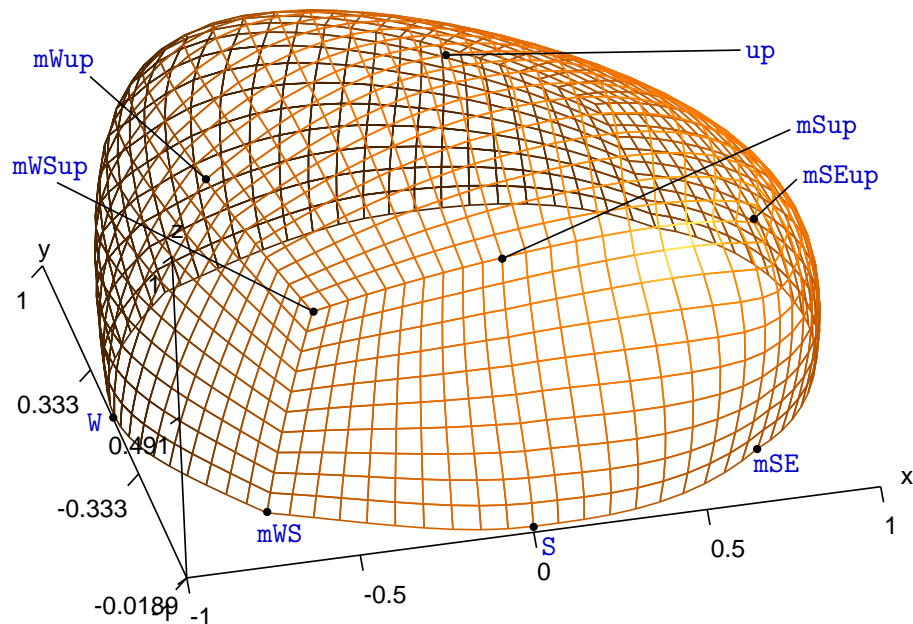
// now build segments :
int n = 10;
auto & W_mWS = Mesh::segment ( W, mWS, n );
auto & W_mWup = Mesh::segment ( W, mWup, n );
// ... and so forth ...

// now three rectangles in each octant :
auto & rect_W_to_S = Mesh::rectangle
( mWS_mWSup, mWup_mWSup.reverse(), W_mWup.reverse(), W_mWS );
auto & rect_S_to_W = Mesh::rectangle
( mSup_mWSup, mWS_mWSup.reverse(), S_mWS.reverse(), S_mSup );
auto & rect_up_to_WS = Mesh::rectangle
( mWup_mWSup, mSup_mWSup.reverse(), up_mSup.reverse(), up_mWup );
// ... and so forth ...

// and finally join the rectangles :
auto & hemisphere = Mesh::join ( list<Mesh*>
{ &rect_E_to_N, &rect_N_to_E, &rect_up_to_EN, &rect_N_to_W,
  &rect_W_to_N, &rect_up_to_NW, &rect_W_to_S, &rect_S_to_W,
  &rect_up_to_WS, &rect_S_to_E, &rect_E_to_S, &rect_up_to_SE } );

hemisphere.export_msh ("hemisphere.msh");

```

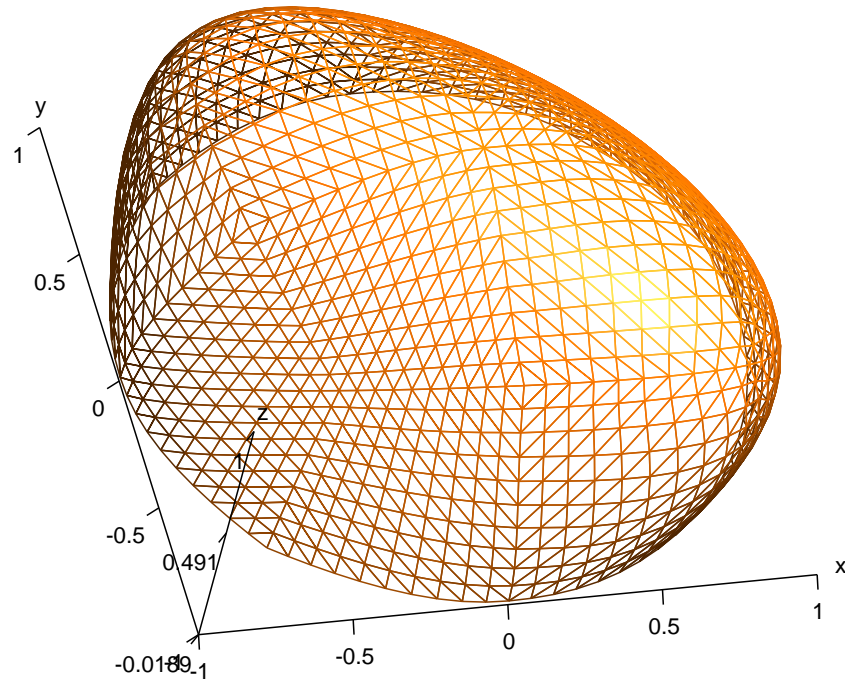


Unlike in paragraph 2.6, here we do not control the exact shape of the segments  $S\_mWS$ ,  $S\_mSup$  and so on. They are projections of straight line segments onto our surface but since the equation of the surface is rather complicated we do not know the exact shape of these projections. Due to the projection on the manifold, there is no guarantee that  $mWS$  lies in the plane  $z = 0$  (it probably doesn't). We notice an angle between  $W\_mWS$  and  $S\_mWS$  at  $mWS$ .

See paragraphs 2.11 and ...for another way of meshing a domain, having its boundary as starting point.

## 2.8. Exercise

Slightly change the code in paragraph 2.7 in order to obtain the mesh below. (Hint: see paragraph 2.2.)



## 2.9. Alternating between manifolds

Let's go back to the example in paragraph 2.5. Suppose we want to mesh the whole disk, not just its boundary. We can build the boundary of the disk just like in paragraph 2.5, by placing ourselves in the manifold `circle_manifold`. But if we want to mesh the interior of the disk, we must switch back to the original `RR2` manifold. The global variable `Mesh::environment` allows us to do that.

```
auto RR2 = Manifold::euclid (2); // Manifold RR2
auto & xy = RR2.coordinate_system ("Lagrange degree one");
// FunctionOnMesh::Function & xy
auto & x = xy[0], & y = xy[1]; // FunctionOnMesh::Function & x, & y
auto & circle_manifold = RR2.implicit ( x*x + y*y - 1. ); // Manifold & circle_manifold
auto & N = Cell::point(); x == 0.; y == 1.; // Cell & A
auto & W = Cell::point(); x == -1.; y == 0.;
auto & S = Cell::point(); x == 0.; y == -1.;
auto & E = Cell::point(); x == 1.; y == 0.;
auto & NW = Mesh::segment ( N, W, 10 ); // Mesh & NW
auto & WS = Mesh::segment ( W, S, 10 );
auto & SE = Mesh::segment ( S, E, 10 );
auto & EN = Mesh::segment ( E, N, 10 );
```



```
Mesh::environment = & RR2;  
auto & disk = Mesh::rectangle ( NW, WS, SE, EN );
```

See paragraphs 2.11 and ...for another way of meshing a domain, having its boundary as starting point.

## 2.10. Alternating between manifolds, again

Here is an example similar to the one in paragraph 2.9, this time with four pieces of hiperbolas.

See paragraphs 2.11 and ...for another way of meshing a domain, having its boundary as starting point.

## 2.11. Meshing a domain from its boundary

Consider again the example from paragraphs 2.5 and 2.9. Instead of meshing the disk as a (much deformed) rectangle, we can use ...in order to build a mesh gradually, having its boundary as a starting point.

### 3. A closer look at cells and meshes

This section gives more details about how cells and meshes are implemented in *manik3D*.

#### 3.1. Building cells and meshes

As with any C++ object, cells can be created by means of their constructor. For instance, the instruction `Cell P(0)` declares a cell of dimension 0 (a vertex), while `Cell * Q = new Cell(0)` declares a pointer to a vertex. We can even de-reference the pointer and declare `Cell & R = * ( new Cell(0) )`. In the subsequent code, P and R can be used in the same manner, the only meaningful difference being that P will be deleted at the end of the block while the cells created by `new` will continue to exist. However, both the cell pointed to by Q and the one referenced by R will become inaccessible outside the current syntactic block, unless we keep track of them in some other way (for instance, integrate them in a mesh and keep access to the mesh).

In *manik3D*, cells and meshes are implemented as persistent objects, built using `new` and thus having no syntactic scope. They are never destroyed; rather, if one is no longer used (e.g. during a remeshing process), it can be explicitly released by the user and is placed in a pool. Such unused cells and meshes will be later re-used. This is achieved through specific methods which build cells, meshes and other objects. We call these methods “factory functions”.

For cells, there are factory functions like `Cell::point`, `Cell::segment`, `Cell::triangle` and others. Except for the first one, the ordinary user will not need to use these factory functions, relying instead on factory functions for meshes (see below). A point may be declared as `Cell & R = Cell::point()`. We can even use the `auto` keyword which leaves to the C++ compiler the task of guessing the type : `auto & R = Cell::point()`. However, we should not forget the ampersand; if we declare `auto R = Mesh::point()`, a compilation error will emerge; see paragraph ? for technical details. The `auto` keyword is extensively used in sections 1 and 2.

For meshes, there are factory functions like `Mesh::segment`, `Mesh::loop`, `Mesh::rectangle` and others. Internally, these build new cells by using factory functions for cells. Note that `Mesh::segment` is quite different from `Cell::segment`. The latter builds a one-dimensional cell, the former produces a mesh which is a chain of one-dimensional cells.

The intention of the authors is that the final user employ (reference to) objects of types `Cell` and `Mesh` rather than pointers. However, in specific cases the use of pointers cannot be avoided, see e.g. paragraphs 3.2 and 3.8.

Recall that, in *manik3D*, cells and meshes are oriented. When a cell or mesh is declared, it is built as positive and has no reverse. Its reverse is a negative cell or mesh and will be built only if necessary. Cells and meshes have a method `reverse` which does the following. It checks if the reverse object has already been built; if yes, it returns that object; otherwise, it builds the reverse cell or mesh on-the-fly and returns it. Paragraph 3.6 gives a more detailed explanation about orientation of cells and meshes. See also paragraph 6.4.

The ordinary user will not need the `reverse` method for cells. However, we may see it occasionally for meshes. In the example in paragraph 1.3, only meshes CD and BC have reverse (because only these are needed).

Factory functions are intensively used in *manil*. Besides cells and meshes, they are used e.g. for building iterators (see paragraph 3.5), functions, variational formulations, finite elements and integrators.

As a general rule, we use lower-case letters for factory functions and capital initial(s) for class names; see also paragraph 6.8.

### 3.2. A ring-shaped mesh

If we want to create many meshes within a cycle, we must use pointers to `Meshes`. Since the factory functions return references to `Mesh` objects (see paragraph 3.1, we must keep the address of these objects in a pointer.

Note how we use a version of `Mesh::join` taking as argument a list of pointers to `Meshes`.

```

Manifold & RR3 = Manifold::euclid (2);
FunctionOnMesh::Function & xy =
    RR3.coordinate_system ("Lagrange degree one");
FunctionOnMesh::Function & x = xy[0], & y = xy[1];

short int n_sectors = 15;
double step_theta = 8*atan(1.)/n_sectors;
short int radial_divisions = 10;
short int rot_divisions = 5;

// start the process by building a segment
Cell & A = Cell::point (); x == 1.; y == 0.;
Cell & B = Cell::point (); x == 2.; y == 0.;
Mesh & ini_seg = Mesh::segment ( A, B, radial_divisions );
// we will need to vary ini_seg, A and B, so we use pointers :
Mesh * prev_seg_p = & ini_seg;
Cell * A_p = &A, * B_p = &B;
list <Mesh*> sectors;

for ( short int i = 1; i < n_sectors; i++ )
{ double theta = i * step_theta;
  // we build two new points
  Cell & C = Cell::point (); x == cos(theta); y == sin(theta);
  Cell & D = Cell::point (); x == 2.*cos(theta); y == 2.*sin(theta);
  // and three new segments
  Mesh & BD = Mesh::segment ( *B_p, D, rot_divisions );
  Mesh & DC = Mesh::segment ( D, C, radial_divisions );
  Mesh & CA = Mesh::segment ( C, *A_p, rot_divisions );
  Mesh & rect = Mesh::rectangle ( *prev_seg_p, BD, DC, CA );
  l.push_back ( &rect );
  prev_seg_p = & ( DC.reverse() );
  A_p = &C; B_p = &D;
}

// we now build the last sector, and close the ring
// prev_seg_p, A_p and B_p have rotated during the construction process
// but ini_seg, A and B are the same, initial, ones

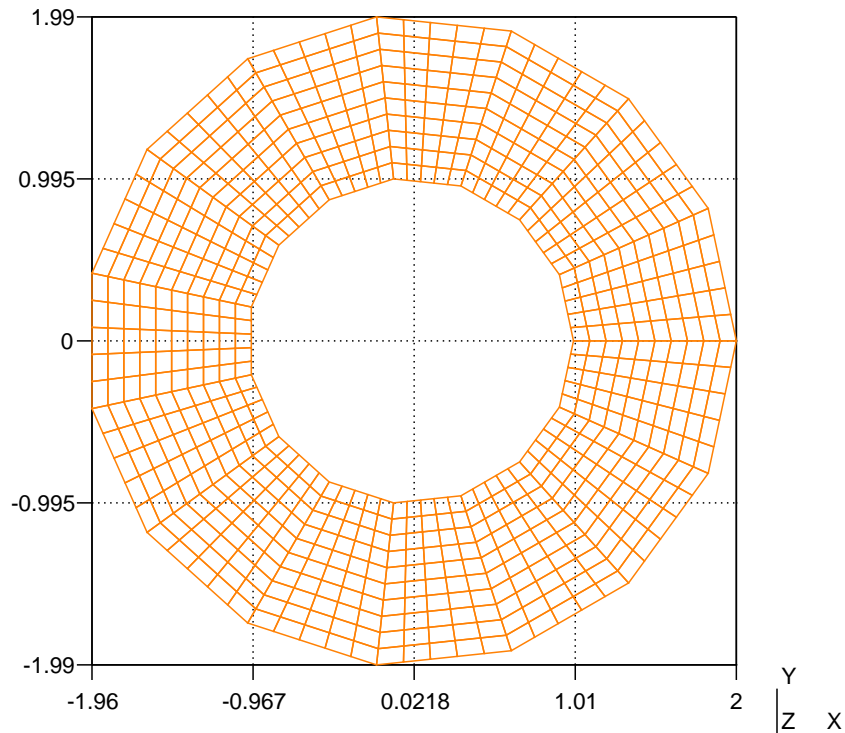
```

```

Mesh & outer = Mesh::segment ( *B_p, B, rot_divisions );
Mesh & inner = Mesh::segment ( A, *A_p, rot_divisions );
Mesh & rect = Mesh::rectangle ( outer, ini_seg.reverse(), inner, *prev_seg_p );
sectors.push_back ( &rect );

Mesh & ring = Mesh::join ( sectors );
ring.export_msh ("ring.msh");

```



See also paragraph 6.5.

### 3.3. Lists of cells inside a mesh

As explained in paragraph 1.2, a `Mesh` is roughly a list of cells. Internally, *manimash* keeps lists of cells of each dimension, up to the maximum dimension which is the dimension of the mesh. Thus, if `msh` is a `Mesh` object, modelling a mesh of triangles, then `msh.cells[0]` points to a list of pointers to cells holding all vertices of that mesh, `msh.cells[1]` points to a list of pointers to cells holding all segments and `msh.cells[2]` points to a list of pointers to cells holding all triangles.

Thus, if we want to iterate over all segments of the mesh, we can use a loop like below.

```

list<Cell*>::iterator it;
for ( it = msh.cells[1]->begin(); it != msh.cells[1]->end(); it++ )
{ Cell * seg = *it; do_something_to (*seg); }

```

Note that the above only works for a positive mesh. See paragraph 3.5 for nicer ways to iterate over cells of a mesh.

On the other hand, a cell is roughly defined by its boundary which in turn is a mesh of lower dimension. Thus, if `hex` is a `Cell` object modelling a hexagon, then `hex.boundary()` is a `Mesh` object modelling a one-dimensional mesh (a closed chain of six segments). So, if we want to iterate, say, over all vertices of that hexagon, we can use a loop like below.

```
list<Cell*> * li = hex.boundary().cells[0];
list<Cell*>::iterator it;
for ( it = li->begin(); it != li->end(); it++ )
{ Cell * P = *it; do_something_to (*P); }
```

Note that in the above we have no guarantee about the order in which the vertices will show up in the loop. See paragraph 3.5 for other ways of iterating over cells, some of which allow us to control the order.

### 3.4. Cells and segments

As explained in paragraph 1.2, the class `Cell` is used to hold points, segments, triangles and other types of cells.

However, in the particular case of segments, it would be nice if we had specific methods to get the base and the tip of that segment.

Recall that in *manifold* a segment is roughly a pair of points, one with a positive sign attached and the other one with a negative sign attached. We call “tip” the positive one and “base” the negative one (this convention is related to integration of a function along that segment).

If we wanted to get the tip of a segment `seg` without a dedicated method, we would have to search through the list `seg.boundary().cells[0]` and pick the positive one. This would be cumbersome and time-consuming, so *manifold* implements segments as a subclass `Segment` derived from `Cell`. The class `Segment` holds internally pointers to the base and the tip of the segment and provides two methods, `base` and `tip`, which return the respective vertex. Note that `base` returns always a negative zero-dimensional cell while `tip` returns a positive one.

Actually, the `base` and `tip` methods exist for any `Cell` object but, when in debug mode, they produce a run-time error if invoked for a cell of dimension other than 1.

### 3.5. Iterators over cells

As explained in paragraph 1.2, a mesh is essentially a list of cells. In many situations, we may want to iterate over all cells of a mesh.

Suppose we have a mesh `msh` of rectangles. If we want to do something to each rectangle, that is, to each two-dimensional cell, we should use a code like

```
list<Cell*>::iterator it;
for ( it = msh.cells[2]->begin(); it != msh.cells[2]->end(); it++ )
{ Cell * cll = *it; do_something_to (*cll); }
```

This style is slightly cumbersome and it doesn't work for negative meshes, so we provide specific methods and iterators. The code above is equivalent to

```
CellIterator it = msh.iter_over ( tag::cells, tag::of_dim, 2, tag::not_oriented );
for ( it.reset(); it.in_range(); it++ )
{ Cell & cll = *it; do_something_to (cll); }
```

See paragraph 6.8 for more details about tags.

If we want to iterate over all vertices of the mesh, we can use

```
CellIterator it = msh.iter_over ( tag::cells, tag::of_dim, 0, tag::not_oriented );
for ( it.reset(); it.in_range(); it++ )
{ Cell & P = *it; do_something_to (P); }
```

If we want to iterate over all segments of the mesh, we can use

```
CellIterator it = msh.iter_over ( tag::cells, tag::of_dim, 1, tag::not_oriented );
for ( it.reset(); it.in_range(); it++ )
{ Cell & seg = *it; do_something_to (seg); }
```

There is also an iterator which goes through oriented cells :

```
CellIterator it = msh.iter_over ( tag::cells, tag::of_max_dim, tag::oriented );
```

See paragraph 3.6 for a discussion about the orientation of cells inside a mesh.

If we only want to know how many cells there are in a certain mesh, instead of using `msh.cells[d]->size()` (`d` being the desired dimension of the cells) we should use the method `number_of` :

```
short int d = 2;
size_t n = msh.number_of ( tag::cells, tag::of_dim, d );
```

or, for cells of dimension equal to the dimension of the mesh :

```
size_t n = msh.number_of ( tag::cells, tag::of_max_dim );
```

Note that it is not safe to modify a mesh while iterating over its cells.

One-dimensional meshes have more specific iterators. Unlike the iterators above, which sweep the mesh in a rather unpredictable order, the iterators below follow the natural order of the cells (either vertices or segments) given by the topology of the mesh.

A one dimensional mesh can be either an open chain of segments or a closed one (a loop). We can iterate over all vertices along an open chain, beginning at one end and finishing at the other end :

```
CellIterator it1 = msh.iter_over ( tag::vertices, tag::along );
for ( it1.reset(); it1.in_range(); it1++ )
{ Cell & P = *it1; do_something_to (P); }
```

Or, we can iterate over all segments :

```
CellIterator it2 = msh.iter_over ( tag::segments, tag::along );
```

Note that `it1` produces positive points, while `it2` produces oriented segments (positive or negative).

There are also reversed versions of these iterators (they go backwards) :

```
CellIterator it = msh.iter_over ( tag::vertices, tag::along, tag::reverse );
CellIterator it = msh.iter_over ( tag::segments, tag::along, tag::reverse );
```

Similar iterators exist for a closed chain of segments (a loop) :

```
CellIterator it = msh.iter_over ( tag::vertices, tag::around );
CellIterator it = msh.iter_over ( tag::segments, tag::around );
CellIterator it = msh.iter_over ( tag::vertices, tag::around, tag::reverse );
CellIterator it = msh.iter_over ( tag::segments, tag::around, tag::reverse );
```

Iterators over closed chains (loops) will start the iteration process at some vertex or segment (the first one in the internal list). If we want to start at a specific location, we can make a `reset` call with one argument. For iterators over vertices, this argument should be a vertex, while for iterators over segments, this argument should be a segment. If such an argument is given, then the iteration process will begin at that particular vertex or segment. This special kind of `reset` can also be used for an open chain, but beware : the vertices or segments previous to the provided argument will not show up in the iteration process.

Due to their specific internal implementation, iterators over one-dimensional meshes (chains) have some limitations. For instance, on disconnected meshes they will fail to sweep the entire mesh (they get trapped in the current connected component).

A safe way to get the first vertex of a (connected) open chain is to use an iterator just once :

```
CellIterator it = msh.iter_over ( tag::vertices, tag::along );
it.reset();
Cell & first_vertex = *it;
```

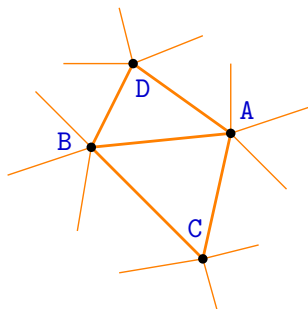
Similarly, we get the last vertex by adding `tag::reverse`. See paragraph 6.7 for some technical details.

Note that it is not safe to modify a mesh while iterating over its cells.

### 3.6. Orientation of cells inside a mesh

In *manifold*, all cells and meshes are oriented.

This can be confusing sometimes, so let's have a closer look at a particular example.



Consider a mesh `tri_mesh` made of triangles. Unless requested otherwise, `tri_mesh` will be a positive mesh and all triangles composing it will also be positive. So, `tri_mesh` will have no reverse mesh and the triangles composing it will have no reverse cell (there is no need for such).

However, the segments must have reverse. Consider triangle `ABC` for instance. Its boundary is made of three segments; let's look at `AB` for example, a segment having `A` as base and `B` as tip. Now, a triangle `BAD` (no offense intended) also exists as part of `tri_mesh`. The boundary of `BAD` is made of three segments, one of them being `BA`, which has `B` as base and `A` as tip. `AB` and `BA` are different `Cell` objects; each is the reverse of the other. One of them is considered positive and the other is considered negative. Which is which depends on which one was built first. So, all inner segments must have a reverse; segments on the boundary of `tri_mesh` will probably have no reverse.

For points (vertices), the situation is even more complex. Segment `AB` sees `A` as negative because `A` is its base, but other segments like `CA` see `A` as positive.

Let's look again at iterators described in paragraph 3.5. We now understand that there is no point to have an iterator over oriented segments, or over oriented vertices, of `tri_mesh`. That's why *manic* only provides an iterator over oriented cells of maximum dimension.

We also understand that there is no difference between these two iterators :

```
CellIterator it1 =
    tri_msh.iter_over ( tag::cells, tag::of_max_dim, tag::oriented );
CellIterator it2 =
    tri_msh.iter_over ( tag::cells, tag::of_dim, 2, tag::not_oriented );
```

because all triangles composing `tri_mesh` should be positive (better use `it1` because it is slightly faster). However, if some of the triangles are negative (which may happen only if `tri_mesh` is built in some fancy, unusual manner) `it1` will behave differently from `it2`. If you are aware that your mesh may contain negative cells but you want to run over their positive counterparts, use `it2`.

On the other hand, the two iterators below will probably have different behaviours, depending on which segments happen to be positive :

```
CellIterator it3 =
    ABC.boundary().iter_over ( tag::cells, tag::of_max_dim, tag::oriented );
CellIterator it4 =
    ABC.boundary().iter_over ( tag::cells, tag::of_dim, 1, tag::not_oriented );
```

Also, there is no difference between the two iterators below (both produce positive points).

```
CellIterator it5 =
    ABC.boundary().iter_over ( tag::vertices, tag::around, tag::reverse );
CellIterator it6 =
    ABC.boundary().reverse().iter_over ( tag::vertices, tag::around );
```

However, the two iterators below are quite different.

```
CellIterator it7 =
    ABC.boundary().iter_over ( tag::segments, tag::around, tag::reverse );
CellIterator it8 =
    ABC.boundary().reverse().iter_over ( tag::segments, tag::around );
```

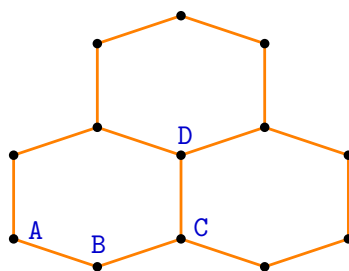
Iterator `it7` will produce segments `AB`, `CA`, `BC` (not necessarily beginning at `AB`), while `it8` will produce their reverses `BA`, `AC`, `CB` (not necessarily beginning at `BA`).



Incidentally, note that the mesh `ABC.boundary()` has no reverse initially; however, a reverse mesh is built on-the-fly when we call the `reverse` method in the declaration of `it6`.

### 3.7. Navigating inside a mesh

Objects in class `Mesh` have two methods, `cell_behind` and `cell_in_front_of`, which provide access to the neighbours of a given cell within the given mesh. Together with methods `base` and `tip` (see paragraph 3.4), they allow us to navigate inside a mesh.



Consider the mesh `hex_msh` shown above, made of three hexagons. Pick one of them, at random :

```
CellIterator it1 = hex_msh.iter_over ( tag::cells, tag::of_max_dim, tag::oriented
);
it1.reset();
Cell & hex_1 = *it1;
```

Now choose a random segment on the boundary of `hex_1` :

```
CellIterator it2 = hex_1.boundary().iter_over ( tag::segments, tag::around );
it2.reset();
Cell & AB = *it2;
```

Take its tip :

```
Cell & B = AB.tip();
```

Suppose now we want the next segment, within the boundary of `hex_1` :

```
Cell & BC = hex_1.boundary().cell_in_front_of(B);
```

And now we may continue by taking the tip of `BC` and then the segment following it :

```
Cell & C = BC.tip();
Cell & CD = hex_1.boundary().cell_in_front_of(C);
```

an so forth (this is how iterators over vertices and segments of one-dimensional meshes, described in paragraph 3.5, are implemented internally).

We can navigate towards a neighbour hexagon :

```
Cell & hex_2 = hex_msh.cell_in_front_of (CD);
```

Of course, since we picked `hex_1` at random within `hex_msh`, as well as `AB` within the boundary of `hex_1`, there is no guarantee that we actually are in the configuration shown in the picture above. That is, `CD` may be on the boundary of `hex_msh`; there may be no

neighbour hexagon `hex_2`. If that is the case, the code above will produce an execution error. See paragraph 3.8 for a way to check if there is actually a neighbour cell and thus avoid errors at execution time.

Note that faces point outwards. For instance, `CD` belongs to the boundary of `hex_1` and points outwards, towards `hex_2`. Thus, `hex_msh.cell_in_front_of(CD)` produces `hex_2`. On the other hand, `hex_msh.cell_behind(CD)` is `hex_1`.

Note also that `CD` does not belong to the boundary of `hex_2`. If we take `hex_2.boundary().cell_in_front_of(D)` we will obtain not `CD` but its reverse, a distinct cell which we may call `DC`. We have that `hex_msh.cell_in_front_of(DC)` is `hex_1` and `hex_msh.cell_behind(DC)` is `hex_2`.

### 3.8. Navigating at the boundary of a mesh

Consider the example in paragraph 3.7. Suppose you try to get a neighbour hexagon which does not exist :

```
Cell & no_such_hex = hex_msh.cell_in_front_of (AB);
```

You will get an execution error of the type

```
ManiFEM::Cell& ManiFEM::Mesh::cell_in_front_of(ManiFEM::Cell&,
const ManiFEM::tag::SurelyExists&): Assertion 'c11 != NULL' failed.
```

You may check previously the existence of the neighbour cell this by using a pointer to a `Cell` and a tag `may_not_exist` :

```
Cell * possible_hex = hex_msh.cell_in_front_of ( AB, tag::may_not_exist );
if ( possible_hex == NULL ) cout << "no neighbour !" << endl;
else do_something_to ( *possible_hex );
```

thus avoiding errors at execution time.

## 4. Fields, functions and variational formulations

### 4.1. Fields and functions (outdated)

Consider the mesh built by the code below.

```
Mesh::intended_dimension = 2; // topological dimension
Mesh::initialize();
auto & xy = NumericField::multi_dim ("size", 2, "lives on", "points");
auto & x = xy[0], & y = xy[1];
auto & u = NumericField::one_dim ("lives on", "points");
auto SW = Cell::point ("SW"); x(SW) = -1.1; y(SW) = 0.3;
auto SE = Cell::point ("SE"); x(SE) = 1; y(SE) = 0;
auto NE = Cell::point ("NE"); x(NE) = 1; y(NE) = 1;
auto NW = Cell::point ("NW"); x(NW) = -1; y(NW) = 1;
auto south = Mesh::segment ( SW, SE, 4, xy );
auto east  = Mesh::segment ( SE, NE, 2, xy );
auto north = Mesh::segment ( NE, NW, 4, xy );
auto west  = Mesh::segment ( NW, SW, 2, xy );
auto rect_mesh = Mesh::rectangle ( south, east, north, west, xy );
```

Note that, besides the `NumericField` objects `x` and `y` which we have already encountered in paragraph 1.1, we introduce another `NumericField`, `u`, which is meant to hold the values of the solution of some PDE.

We now declare two functions defined on the mesh, linked to the two fields `x` and `y`. We can declare each function individually, as we did in paragraph 1.1, or we can declare the pair and then extract each component :

```
auto & xxyy = FunctionOnMesh::from_field ( xy, "Lagrange degree one" );
auto & xx = xxyy[0], & yy = xxyy[1];
```

These functions are described as "Lagrange degree one", which means that they vary linearly along segments and also inside triangles. On quadrilaterals, they are polynomials of degree one, meaning they have a linear part plus a bi-linear one.

We can also declare an unknown function and a test function, to be used in a future variational problem :

```
auto & uu = FunctionOnMesh::unknown ( u, "Lagrange degree one");
auto & w = FunctionOnMesh::test ( uu );
```

The unknown `uu` is related to the `NumericField` previously declared, `u`, which provides space to hold, at each vertex of the mesh, a real value. Hopefully, at the end of the day these will be the values of the solution of our PDE. The test function does not need to hold any values. The test function is an abstract object whose only use is to express the variational formulation.

`FunctionOnMesh` objects obey to usual arithmetic rules. For instance,  $(xx+yy*w)/uu$  is a valid expression in *manil33*. They can also be differentiated, like in `(xx*yy).deriv(xx)`. Note that this expression will be evaluated right away, producing the result `yy`, while expressions involving an unknown function or a test function will produce a delayed

derivative object, to be evaluated later. Thus, `w.deriv(xx)` will be evaluated only after replacing the test function `w` by some function in a base of a discretized Hilbert space.

`FunctionOnMesh` objects can also be integrated through their method `integrate`, which produces a delayed integral expression. The integral is not evaluated right away, but only later, with the use of an `Integrator` object (which could be a Gauss quadrature).

Integral expressions are stored as objects belonging to the class `FunctionOnMesh::combinIntegrals`. These objects can be equated by using their `operator==`, which returns a `VariationalProblem` object. Thus, the operator `FunctionOnMesh::combinIntegrals::operator==` acts as a factory function for `VariationalProblem` objects. For instance, `x.integrate(rect_mesh) == y.integrate(south)` is a syntactically valid expression which would produce a `VariationalProblem` but actually gives a run-time error because the `operator==` checks that the right hand side of the variational problem actually contains a function declared as unknown and one declared as test, and that the left hand side contains a test function but no unknown. Here is a more meaningful example :

```
auto & var_pb =
  ( uu.deriv(xx)*w.deriv(xx) + uu.deriv(yy)*w.deriv(yy) ) .integrate(rect_mesh)
  == w.integrate(rect_mesh) + w.integrate(south);
```

Note how we can mix integrals on different domains; recall that `south` is a side of `rect_mesh`.

Dirichlet-type boundary conditions are implemented through the directive `prescribe_on` followed by one or more equalities.

```
var_pb.prescribe_on (north); uu == 0.;      w == 0.;
var_pb.prescribe_on (east); uu == xx*(1.-yy); w == 0.;
var_pb.prescribe_on (west); uu == 0.;      w == 0.;
```

## 5. Finite elements and integrators

Let's take over the example in paragraphs 1.9 and 4.1.

The last few lines of code look rather simple :

```
auto & fe = FiniteElement::Lagrange ("Q1");
fe.set_integrator ("on cells of dimension", 1, "gauss", 3, "nodes");
fe.set_integrator ("on cells of dimension", 2, "gauss", 9, "nodes");
fe.discretize ( var_pb );
```

This simplicity hides from the final user a complicated process : the discretization of the variational formulation by a finite element.

### 5.2. Finite elements

The notion of a finite element is quite complex. The purpose of a `FiniteElement` is to build a list of functions, say,  $\psi$ , defined on our mesh. The linear span of these functions will be a discretized Hilbert space. The `FiniteElement` should replace, in the variational formulation, the unknown function by one  $\psi$ , the test function by another  $\psi$  and, by evaluating the integrals, obtain the coefficients of a system of linear equations. Some external solver will then solve the system, and it is the job of the finite element to transform back the vector produced by the solver into a function defined on our mesh.

Computing each integral is a somewhat separate process; it's the job of an `Integrator` which could be a Gauss quadrature or some other procedure like symbolic integration. When a Gauss quadrature is used, the separation between a `FiniteElement`'s job and the `Integrator`'s job is not very sharp because often the Gauss quadrature is performed not on the physical cell but rather on a master element which is built and handled by the `FiniteElement`. The authors of *manil* have tried to separate these two concepts as much as possible, especially because some users may want to use a `FiniteElement` with no master element, or an `Integrator` acting directly on the physical cell.

Thus, there is a base class `FiniteElement` and a derived class `FiniteElement::withMaster` which keeps, as an extra attribute, the map transforming the master element to the current physical cell. This map depends of course on the geometry of the cell and thus it must be computed from scratch each time we begin integrating on a new cell. We say that the `FiniteElement` is docked on a new `Cell`; the method `dock_on` performs this operation. This method is element-specific, each type of finite element having its own class.

For instance, the class `FiniteElement::Lagrange_Q1` is a class derived from `FiniteElement::withMaster`. It will only `dock_on` quadrilaterals (two-dimensional `Cells` with four sides). Recall that in *manil* the vertices do not have necessarily any number attached. When we create a `FiniteElement::Lagrange_Q1` object, it will enumerate vertices of the mesh if a mesh exists already, and it will prepare the ground for vertices created in the future to be enumerated, too. This enumeration is useful to identify a vector of real numbers with the values (at each vertex) of a function defined on the mesh. When docking on a cell, the `FiniteElement::Lagrange_Q1` object will build four "shape functions", a transformation map (from a master element occupying the square  $[-1, 1]^2$  to the current cell) and the jacobian of this transformation.

## 6. Technical details

The present section is meant for those interested in developing and extending *maniℰℛ*. Of course the ultimate documentation is the source code; this section can be used as a guide through the source code.

### 6.1. General considerations

Designing and implementing *maniℰℛ* has certainly been fun (and we have learned a lot along the process), but it was difficult.

Take cells and meshes, for instance. As explained in paragraph 1.2, at the conceptual level meshes are roughly collections of cells of the same dimension and cells are essentially defined by their boundary which is a lower-dimensional mesh. This conceptual simplicity does not survive to the demands of an efficient code (which doesn't mean it has not been extremely useful in the process of designing *maniℰℛ*; it should also be useful for learning and using it).

Vertices (zero-dimensional cells) have no boundary at all. One-dimensional cells (segments) have all the same shape and have a rudimentary boundary (a zero-dimensional mesh consisting of a **base** and a **tip**). In order to save space in the computer's memory, specific classes have been created for vertices and for segments.

Also, there are positive cells and negative cells, positive meshes and negative meshes. A positive cell and its negative counterpart (its **reverse**) share some information. The dimension is the same. The boundary of each one is the reverse of the boundary of the other. Again, to save memory space, negative cells are implemented in different classes from positive cells, thus avoiding the storage of redundant information.

We want, however, to manipulate all these different cells through a uniform interface, which is where C++'s inheritance and polymorphism mechanisms come handy. Thus, there are classes `PositiveVertex`, `PositiveSegment` and `PositiveCell`, all derived from `PositiveBaseCell`. And we have `NegativeVertex`, `NegativeSegment` and `NegativeCell`, all derived from `NegativeBaseCell`. Both `PositiveBaseCell` and `NegativeBaseCell` are derived from `BaseCell`. And we do not want to bloat the user's namespace with too many names, so all these class names are **hidden** (see paragraph 6.2).

On the other hand, we want cells and meshes to be persistent objects (not subject to syntactic scope). We create them within some function and we want them to remain alive after returning to the main program. Also, they are unique entities, it does not make sense to copy them. This is why we have implemented `Cell` as a thin wrapper around `BaseCell` with most methods of `Cell` being delegated to `BaseCell`. When it goes out of its syntactic scope, the wrapper is destroyed but the `BaseCell` object inside (its **core**) remains intact. Also, you can copy the wrapper as in `Cell A = B` or `Mesh BA = AB.reverse()` but these operations do not create new core objects, they just give new names to already existing cells or meshes. See paragraph ... for details. You can think of `Cells` and `Meshes` as customized pointers towards `BaseCells` and `PositiveMeshes`, respectively.

Negative meshes contain no useful information (they only appear as boundaries of negative cells) so there is no such class as `NegativeMesh`. The wrapper class `Mesh` contains

a pointer to a `PositiveMesh` and a flag telling it to reverse everything if the mesh is considered negative. Zero-dimensional meshes are not stored at all. One-dimensional meshes have a specific structure (they are open or closed chains of segments) and will receive in the future a specific treatment (a specialized class).

To save memory, we don't even keep the dimension of a cell as an attribute, it's a static property for vertices and segments, while for higher-dimensional cells it's obtained from the boundary's dimension. The dimension of a mesh is not kept as an attribute either; it's computed on-the-fly by counting the levels of collections of cells the mesh is made of (and then subtracting one). For instance, the mesh in paragraph 1.1 has three layers of cells : points, segments and squares.

Iterators ...

The same goes for manifolds, for instance. There are Euclidian manifolds, cartesian products, implicitly defined submanifolds and parametrized submanifolds. Distinct notions with very different internal implementations. But we would like to offer the user the possibility of declaring any of these as "a `Manifold`"-

## 6.2. Hidden, private, public

In *manifem*, there are no `private` or `protected` class members or methods. Everything is `public`; the user can make use of any class member if he or she so chooses. This can be considered poor design; we endorse this criticism with no further comments.

However, some class members and methods are intended to be used by the final user, while others are used in the internal implementation of the former. Classes not intended for the final user have been gathered under the namespace `hidden` (see paragraph 6.8). Also, in the source code there are comments like

```
// do not use directly, let [some other method] do the job
```

As a rule of thumb, there is no reason for you to use in your programs any name beginning with `hidden`, nor any class members or methods not mentioned in this manual. On the other hand, you are encouraged to read them, understand them and hopefully improve them.

## 6.3. Tags

We use extensively `tags`. These are structures gathered in the namespace `tag`. Most of them contain no data; only their type is useful, at compile time.

A `tag` is used to clearly distinguish between functions with the same name (overloaded functions) :

```
// example please
```

The above could be achieved by giving longer names to the functions, but we believe `tags` make the code more readable. Besides that, in the case of constructors, we do not have the choice of the name of the function (a constructor has the same name as its class). *Manifem* uses `tags` extensively for constructors.

In *manifem*, namespaces and class names begin with capital letter (see paragraph 6.8) The namespace `tag` (or `ManiFEM::tag` if you are not using namespace `ManiFEM`) is an

exception to the above rule. We prefer its name to have only lower case letters because we want it to be discrete. For instance, if we wrote `Tag::reverse_of`, the reader's eye would catch `Tag` much before `reverse_of`; thus, `tag::reverse_of` is more readable. Of course, the final user has the choice of using namespace `tag`.

## 6.4. Orientation of cells and meshes

Both classes `Cell` and `Mesh` have an attribute `hidden_reverse` which is `NULL` if the cell or mesh has no reverse, otherwise points to the reverse object. Note that negative cells and meshes must have a reverse, which is a positive cell or mesh. Positive cells and meshes may have no reverse. In the code below, `c11.reverse` simply returns `c11.hidden_reverse` :

```
Cell & c11 = ...; // some factory function
... more code ...
Cell * r_c11_p = c11.reverse ( tag::may_not_exist );
```

Note that this is quite different from the code below where a reverse cell is built on-the-fly if needed (also, a reference is returned rather than a pointer) :

```
Cell & c11 = ...; // some factory function
... more code ...
Cell & r_c11 = c11.reverse();
```

Although the most natural choice for implementing the orientation may seem to be a `sign` attribute of classes `Cell` and `Mesh`, we have chosen a different internal implementation. There is an attribute `positive` in both classes which points to the object itself if the cell or mesh is positive, and points to the reverse object if the cell or mesh is negative. In the former case, `positive == this`; in the latter, `positive == hidden_reverse`. The method `is_positive` hides these details from the user (it simply returns the result of `this == positive`).

## 6.5. Forgetting meshes

In paragraph 1.3 and others we have shown how to join meshes. The question arises, do we need to keep the intermediate meshes or just the final, big, one ? That's up to the user to decide. If we prefer to keep only the final mesh, we can "get rid" of the meshes we don't need anymore by forgetting them.

Note that it is not enough for the respective C++ object to go out of scope. In *manikern*, cells and meshes are created on the free store and survive outside their syntactic scope (see paragraph 6.1). The user must release them explicitly.

For instance, in paragraph 1.3, after building `L_shaped`, we may add the following lines of code :

```
BC.forget(); // BC.reverse() will be discarded, too
CD.forget(); // CD.reverse() will be discarded, too
ABCD.forget(); CEFD.forget(); BGHC.forget();
```

The `forget` operation will not discard the component cells of that mesh. If we want that, we must use the `shread` method, but the need for such should be rare.



## 6.6. About `init_cell`

The `Cell` class has static attributes `init_cell`, `init_cell_r`, `data_for_init`, `data_for_init_r`. The attribute `init_cell` is a list of pointers to functions to be called by the constructor of a positive cell, while `data_for_init` is a void pointer which can be used to pass supplementary information to `init_cell`. Attributes `init_cell_r` and `data_for_init_r` fulfill a similar task when building negative cells.

For instance, `Manifold::coordinate_system` inserts into the list `Cell::init_cell[0]` a call to `Mesh::prescribe_on` which in turn calls `FunctionOnMesh::prescribe_on` in order to prepare the ground for instructions like `x == 1.0` to produce the desired effect after the creation of each point.

Another example is the constructor `FiniteElement::Lagrange::Q1` which adds a call to `FiniteElement::Lagrange::Q1::enumerate_new_vertex` to the list `Cell::init_cell[0]`. This way, future vertices will receive automatically a `size_t` label, to be used by Lagrange finite elements.

## 6.7. About `hook`

Both `Cell` and `Mesh` classes have an attribute called `hook` which allows the user to attach supplementary data to a cell without having to declare a derived class. The `hook` attribute is a map from strings to void pointers. The user may insert (pointers to) data of any type into this map by converting them to `void*` through explicit casting. In order to be used later, this data must be converted back to the original type. The authors of *manikern* are aware that this approach is prone to errors. The programmer must be very careful when casting back and forth between types, and must keep record of keys of the `hook` map which are already in use (recall that these keys are strings).

Below is a list of keys already used in maps `Cell::hook` and `Mesh::hook`. Note that they exist in positive cells and meshes, not in negative ones.

The factory function `Mesh::segment` inserts pointers towards the first and last nodes as `hook["first point"]` and `hook["last point"]`. The method `Mesh::join` honors this convention by setting these pointers as appropriate, if applied to Meshes having these attributes. These attributes are used in the `reset` method of iterators over cells of open chains `msh.iter_over ( tag::vertices, tag::along )`, `msh.iter_over ( tag::segments, tag::along )`, `msh.iter_over ( tag::vertices, tag::along, tag::reverse )`, `msh.iter_over ( tag::segments, tag::along, tag::reverse )` (described in paragraph 3.5).

If a one-dimensional mesh is built by some means other than `Mesh::segment` and `Mesh::join`, the `hook` map may have no keys "first point" and "last point". A safe way of getting the first and last vertices is to use an iterator like in

```
CellIterator it1 = msh.iter_over ( tag::vertices, tag::along );
it1.reset(); Cell & first_ver = *it1;
CellIterator it2 = msh.iter_over ( tag::vertices, tag::along, tag::reverse );
it2.reset(); Cell & last_ver = *it2;
```

This will always provide the correct vertices because the `reset` method looks for the key "first point" or "last point" and, if it doesn't find it, performs a direct search to find the first or last vertex. Beware that, if the mesh is not an open chain (if it's a loop),

this search may last forever. The above method has the advantage of working correctly even for a negative mesh.

The factory function `Mesh::rectangle` inserts pointers to two maps `hook["sides"]` and `hook["corners"]`. The first one of these maps holds pointers to the four sides of the rectangle under the keys "south", "east", "north" and "west". The second map holds pointers to the four corners of the rectangle under the keys "SW", "SE", "NE" and "NW".

## 6.8. Namespaces and class names

All names in *manifem* are wrapped into the namespace `ManiFEM`. We recommend to be using `namespace ManiFEM` in your code, otherwise the text will become cumbersome. For instance, you will have to write `ManiFEM::Mesh::rectangle` instead of `Mesh::rectangle`, and so on. We are using `namespace ManiFEM` in the examples of this manual.

As a general rule, namespaces and class names are written with capital initial letter : `Cell`, `Mesh`, `Integrator`, `FiniteElement`, `VariationalFormulation`. In contrast, factory functions (see paragraph 3.1) are named with lower case letters, e.g. `Mesh::rectangle` or `msh.iter_over`.

Namespaces `hidden` (see paragraph 6.2) and `tag` (see paragraph 6.3) are exceptions to the above rule.

## 6.9. Frequent errors at compile time

```
use of deleted function 'ManiFEM::Cell::Cell(const ManiFEM::Cell&)\n-- or --\nuse of deleted function 'ManiFEM::Mesh::Mesh(const ManiFEM::Mesh&)
```

Cells and meshes cannot be copied. You cannot ask for things like `cell_2 = cell_1` or `mesh_2 = mesh_1`. Also, when initializing a cell or a mesh through a factory function, you should use a reference; `Cell tri = Cell::triangle (...)` is illegal, as well as `auto tri = Cell::triangle (...)`; use `Cell & tri = Cell::triangle (...)` or `auto & tri = Cell::triangle (...)`.

## 6.10. Frequent errors at run time

When you think your program is ready for shipping, you may want to optimize it by adding the `-dNDEBUG` option to your compilation command (check your `Makefile`). Remember to `make clean` before re-building your application. Optimizing your code should increase its execution speed.

If your program produces unpredictable, random errors at run-time, e.g. throws `Segmentation fault`, try to run it in debug mode. To achieve this, simply remove any `-dNDEBUG` option from your compilation command (check your `Makefile`). Remember to `make clean` before re-building your application.

Errors described below are produced by assertions and thus will only show up in debug mode.

```
ManiFEM::Cell& ManiFEM::Cell::base(): Assertion 'dim == 1' failed.\n-- or --\nManiFEM::Cell& ManiFEM::Cell::tip(): Assertion 'dim == 1' failed.
```

It seems you are trying to get the base or the tip of a cell of dimension different from 1. This does not make sense.

```
double& ManiFEM::OneDimField::operator()(ManiFEM::Cell&) const:
Assertion 'c11.real_heap_size() > index_min' failed.
```

You probably tried to access a coordinate (or some other value) at a cell to which no value has been associated. Either you picked a cell of a different dimension (e.g. a segment instead of a vertex) or you are looking at a negative cell. Values are usually stored at positive cells; negative cells have no information attached. For any cell, you may use the `positive` attribute which is a pointer equal to `this` if the cell is positive or points to the reverse if the cell is negative (so the reverse is positive). Or you may check if a certain cell is positive or negative by using the method `is_positive` (which simply returns the result of the comparison `this == positive`). See paragraphs 3.6 and 6.4 for more details about orientation of cells and meshes.

Note that, if `seg` is a segment (a one-dimensional cell), then `seg.tip()` is a positive cell but `seg.base()` is a negative cell. So, you probably need to use `seg.base().reverse()` instead. Some iterators have versions ending in `tag::not_oriented` which produce positive cells. Iterators with `tag::vertices_along` or `tag::vertices_around` produce positive points, while iterators with `tag::segments_along` or `tag::segments_around` produce oriented segments (which may be positive or negative), see paragraph 3.5.

```
ManiFEM::Cell::Cell(short int, bool): Assertion 'is_segment or ( d != 1 )' failed.
```

Did you try to build a one-dimensional cell by using directly the `Cell` constructor ? Don't. Always use the factory function `Cell::segment`. When you build a cell whose dimension `d` is unknown at compile time, use an if statement to give special treatment to the case `d == 1`. Segments belong to a separate class, derived from `Cell` (see paragraph 3.4). The fact that *manikern* stores them as `Cell` objects, rather than as `Segment` objects, can be considered poor design. We endorse this criticism with no further comments.

```
ManiFEM::Cell& ManiFEM::Mesh::cell_in_front_of(ManiFEM::Cell&,
const ManiFEM::tag::SurelyExists&): Assertion 'c11 != NULL' failed.
-- or --
ManiFEM::Cell& ManiFEM::Mesh::cell_behind(ManiFEM::Cell&,
const ManiFEM::tag::SurelyExists&): Assertion 'c11 != NULL' failed.
```

You are navigating dangerously close to the boundary of a mesh. See paragraph 3.8.

```
ManiFEM::Mesh::diff(size_t, size_t): Assertion 'a >= b' failed.
```

You probably defined the maximum topological dimension using `Mesh::set_max_dim` (and the `DECLARE_MAX_DIM` compiler directive is in effect). And you are probably trying to build a mesh of higher dimension than the one defined. Along your program, you may have different maximum topological dimensions (that is, you may use `Mesh::set_max_dim` several times) but, at each moment, you can only build meshes of dimension up to the value most recently defined.

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