Simulation of the Poisson problem on the 3D Torus

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1 Introduction

We consider the torus defined as follow

$$\Gamma = \{(x, y, z) \in \mathbb{R}^3, (\sqrt{x^2 + y^2} - R)^2 + z^2 - r^2 = 0\},\$$

where r > 0 is the minor radius and R > r is the major radius.

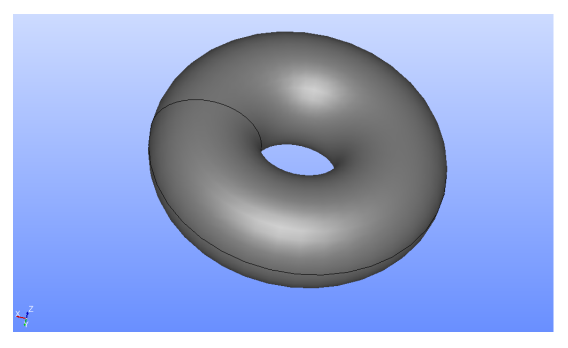


Figure 1: The torus in SALOME CAO module

The torus is a C^{∞} manifold of dimension 2 embedded in \mathbb{R}^3 . The **Laplace-Beltrami operator** \triangle_{Γ} , a generalisation of the euclidean laplacean, can be defined on the torus as the combination of a surface divergence ∇_{Γ} , and of a surface gradient $\vec{\nabla}_{\Gamma}$ (see for instance [3, 6, 12]).

Using the torus coordinates (θ, ϕ) where $\theta, \phi \in [0, 2\pi]$, the Laplace-Beltrami operator takes the following form on the torus

$$\Delta_{\Gamma} f = \frac{1}{r^2 (R + r \cos \theta)} \frac{\partial}{\partial \theta} \left((R + r \cos \theta) \frac{\partial f}{\partial \theta} \right) + \frac{1}{(R + r \cos \theta)^2} \frac{\partial^2 f}{\partial \phi^2}.$$

We consider the following **Poisson problem** on the torus

$$\begin{cases}
-\Delta_{\Gamma} u = f \text{ on } \Gamma \\
\int_{\Gamma} u = 0
\end{cases} , \tag{1}$$

where the right hand side $f \in L^2(\Gamma)$ and the unknown $u \in H^1(\Gamma)$ are **zero** mean functions.

For the following choice of $u(\theta, \phi) = \sin(3\phi)\cos(3\theta + \phi)$ (see [5]), the right hand side is given by

$$f = 9r^{-2}\sin(3\phi)\cos(3\theta + \phi) - (10\sin(3\phi)\cos(3\theta + \phi) + 6\cos(3\phi)\sin(3\theta + \phi))((R + r\cos(\theta))^{-2} - 3\sin(\theta)\sin(3\phi)\sin(3\theta + \phi)(r(R + r\cos(\theta)))^{-1}.$$

Our objective is to solve numerically the Poissson problem (1) using the finite element method described in [1, 4, 2, 3].

2 Finite elements method for 3D Poisson problem

Since Γ is closed (no boundary), we have to impose the global condition $\int_{\Gamma} u = 0$ to guarantee the uniqueness of solution. For this reason we define the following Lebesgue space

$$L^{2}_{\#}(\Gamma) = \{ w \in L^{2}(\Gamma) : \int_{\Gamma} w = 0 \},$$

and the following Sobolev space

$$H^1_{\#}(\Gamma) = \{ w \in H^1(\Gamma) : \int_{\Gamma} w = 0 \}.$$

2.1 Well-posedness of the problem

2.1.1 Variational formulation and Poincaré inequality

In this section we are going to recall two important properties of the surface gradient operator $\vec{\nabla}_{\Gamma}$: the Green-Ostrogradski formula (i.e. integration by part) and Poincaré inequality.

Thanks to the **Green-Ostrograski** theorem, the variational formulation of (1) is:

Find
$$u \in H^1_{\#}(\Gamma)$$
 such that $\forall v \in H^1_{\#}(\Gamma), \int_{\Gamma} \overrightarrow{\nabla}_{\Gamma} u \cdot \overrightarrow{\nabla}_{\Gamma} v = \int_{\Gamma} f v.$ (2)

As for the classical gradient, there is a **Poincaré's inequality** involving the surface gradient (see theorem 2.12 in [3]).

Theorem 1 (Poincaré's inequality).

Assume that Γ is an embedded C^3 hypersurface. There exists a constant c such that, for every function $f \in H^1(\Gamma)$ with $\int_{\Gamma} f = 0$, we have the inequality

$$||f||_{L^2(\Gamma)} \le c||\nabla_{\Gamma} f||_{L^2(\Gamma)}.$$

2.1.2 Existence of a unique weak solution

The bilinear form

$$a(u,v) = \int_{\Gamma} \overrightarrow{\nabla}_{\Gamma} u \cdot \overrightarrow{\nabla}_{\Gamma} v$$

is continuous and coercive thanks to Poincaré inequality.

The linear form

$$b(v) = \int_{\Gamma} fv$$

is continuous.

By application of the **Lax-Milgram theorem**, the variational formulation (2) of problem (1) admits a unique weak solution, which depends continuously on the data f (see Theorem 3.1 in [3]).

Theorem 2 (Well-posedness). Let Γ , be a compact C^2 hypersurface in \mathbb{R}^3 and assume that $f \in L^2(\Gamma)$ and $\int_{\Gamma} f = 0$. Then there exists a unique solution $u \in H^1(\Gamma)$ of (2) with $\int_{\Gamma} u = 0$.

2.1.3 Regularity of the solution

The regularity of the solution requires the regularity of both the right hand side and the manifold. The following theorem is taken from [3] theorem 3.3.

Theorem 3. Let Γ , be a compact C^2 hypersurface in \mathbb{R}^3 and assume that $f \in L^2(\Gamma)$ and $\int_{\Gamma} f = 0$. Then, the unique weak solution of (2) satisfies $u \in H^2_{\#}(\Gamma)$, and there exists a constant C > 0 such that

$$||u||_{H^2_{\#}(\Gamma)} \le C||f||_{L^2_{\#}(\Gamma)}.$$

For more details see [1] for euclidian case and [2, 3] for the case of curved surfaces.

2.2 The P1 finite elements

Following [3], we first approximate the torus Γ by a polyhedral surface Γ_h with triangular faces $(\mathcal{T}_k)_{k\geq 1}$ called elements having their nodes on Γ . We approximate functions $f\in H^1_\#(\Gamma)$ by functions $f_h\in H^1_\#(\Gamma_h)$ via the lift operator (10).

We consider u_h the weak solution of the following Poisson problem on the piecewise linear manifold Γ_h :

$$\begin{cases}
-\Delta_{\Gamma_h} u_h = f_h \text{ on } \Gamma_h \\
u_h \in H^1_\#(\Gamma_h)
\end{cases} , \tag{3}$$

and its variational formulation, analog to (2) is

Find
$$u_h \in H^1_\#(\Gamma_h)$$
 such that $\forall v_h \in H^1_\#(\Gamma_h), \int_{\Gamma_h} \overrightarrow{\nabla}_{\Gamma_h} u_h \cdot \overrightarrow{\nabla}_{\Gamma_h} v_h = \int_{\Gamma_h} f_h v_h.$

(4)

We look for \tilde{u}_h the projection of the solution u_h of (4) on the space $V_0(\Gamma_h)$ of continuous piecewise affine functions with zero mean on Γ_h . The discrete form of the variational formulation (4) is then given by.

Find
$$\tilde{u}_h \in V_0(\Gamma_h)$$
 such that $\forall \tilde{v}_h \in V_0(\Gamma_h), \int_{\Gamma_h} \overrightarrow{\nabla}_{\Gamma_h} \tilde{u}_h \cdot \overrightarrow{\nabla}_{\Gamma_h} \tilde{v}_h = \int_{\Gamma_h} \tilde{f}_h \tilde{v}_h$, (5)

where \tilde{f}_h is the projection of f_h on $V_0(\Gamma_h)$.

2.3 The linear system to be solved

Since $V_0(\Gamma_h)$ is generated by the nodal functions $\phi_i : \Gamma_h \to \mathbb{R}$, i = 1, ..., n such that $\phi_i(x_j) = \delta_{ij}$, (5) takes the following algebraic form

$$A_{\triangle_{\Gamma_h}} X = b_h, \tag{6}$$

where

$$\tilde{u}_h = \sum_{i=1}^n u_i \phi_i,\tag{7}$$

 $A_{\triangle_{\Gamma_h}} = (a_{ij})_{i,j=1,...,n}, X = {}^t(u_1,...,u_n) \text{ and } b_h = {}^t(b_1,...,b_n) \text{ with}$

$$a_{ij} = \int_{\Gamma_h} \overrightarrow{\nabla}_{\Gamma_h} \phi_i \cdot \overrightarrow{\nabla}_{\Gamma_h} \phi_j = \sum_{k=1}^n \int_{\mathcal{T}_k} \overrightarrow{\nabla}_{\Gamma_h} \phi_i \cdot \overrightarrow{\nabla}_{\Gamma_h} \phi_j,$$

$$b_j = \int_{\Gamma_h} f \phi_j = \sum_{k=1}^n \int_{\mathcal{T}_k} f \phi_j.$$

 $A_{\triangle_{\Gamma_h}}$ is symmetric positive and sparse but not invertible since constants are in its kernel, hence the linear system (6) is singular. However it admits a unique solution with zero mean provided the right hand side has zero mean (see [3]).

2.4 Convergence of the numerical method

2.4.1 Fermi coordinates and lift operator

A function u defined on Γ can be extended to a neighborhood of Γ in \mathbb{R}^3 using a lift operator based on the Fermi coordinates around Γ . Following [3], we define the δ -strip around Γ as

$$U_{\delta,\Gamma} = \{ x \in \mathbb{R}^3, dist(x,\Gamma) < \delta \}. \tag{8}$$

For δ small enough it is possible to define the projection $a: U_{\delta} \to \Gamma$ onto Γ and the distance function $d: U_{\delta} \to \mathbb{R}_+$ to Γ . a(x) and d(x) are called the **Fermi coordinates** of x and their existence is given by the following theorem (see Lemma 2.8 in [3] for the proof).

Theorem 4 (Fermi coordinates).

Let Γ be an embedded C^2 hypersurface. There exists $\delta_{Fermi} > 0$ such that for every point $x \in U_{\delta_{Fermi},\Gamma}$, there exists a unique point $a(x) \in \Gamma$, and a function $d \in C^2(U_{\delta_{Fermi},\Gamma})$ such that

$$\forall x \in U_{\delta_{Fermi}, \Gamma}, \quad x = a(x) + d(x) \overrightarrow{n}(x),$$
 (9)

where $\overrightarrow{n}(x)$ is the unit normal vector to Γ at x.

Thanks to the **Fermi coordinates** defined in theorem 4, we can define as in [3] (equation 4.2) a **lift operator** L such that

$$L: C(\Gamma_h) \to C(\Gamma) u_h \to u_h \circ a^{-1} ,$$
 (10)

provided

$$\Gamma_h \subset U_{\delta_{Fermi},\Gamma}.$$
 (11)

2.4.2 Convergence theorems

In order to study the convergence of the finite element approximation, we need to compare $u \in H^1(\Gamma)$ with $\tilde{u}_h \in H^1(\Gamma_h)$ but don't share the same support. Hence we need to use the lift operator (10) which requires the assumption (11) that the triangulated surface Γ_h is close enough to Γ .

As the parameter h goes to zero the distance between Γ_h and Γ converges to zero as expressed in the following theorem taken from [3] Lemma 4.1.

Theorem 5 (Convergence of Γ_h towards Γ). Let $\Gamma \in \mathbb{R}^3$ be an embedded C^2 hypersurface and $\Gamma_h \subset U_{\delta_{Fermi},\Gamma}$ a piecewise linear surface. Let h be the largest diameter of trianges in Γ_h . There exists a constant c such that

$$\forall x \in \Gamma_h, \quad dist(x, \Gamma) \le ch^2.$$

Once proven that Γ_h converges towards Γ , we can prove that \tilde{u}_h converges to u using the lift operator (10). The following convergence theorem is taken from [2] Theorem 8, Lemma 6 and Lemma 7.

Theorem 6 (Convergence of \tilde{u}_h towards u). Let $\Gamma \in \mathbb{R}^3$ be an embedded C^2 hypersurface and $\Gamma_h \subset U_{\delta_{Fermi},\Gamma}$ a piecewise linear surface. Let h be the largest diameter of trianges in Γ_h .

If u is a continuous solution of the Poisson problem (1) and \tilde{u}_h is the discrete solution of (5), then there exists c > 0 such that

$$||u - \tilde{u}_h \circ a^{-1}||_{L^2(\Gamma)} \le ch^2, \quad ||\nabla_{\Gamma}(u - \tilde{u}_h \circ a^{-1})||_{L^2(\Gamma)} \le ch.$$
 (12)

3 Numerical results for Laplace-Beltrami operator on Torus

For the coding the finite element method, we use the Python language and the open-source Linux based library CDMATH [11] which is very simple for the manipulation of large matrices, vectors, meshes and fields. It (CDMATH) can handle finite element and finite volume discretizations, read general 3D geometries and meshes generated by SALOME.

3.1 Meshing of the domain

For the design and meshing of the domain we use GEOMETRY and MESH modules of the software SALOME 9.5 (see [8, 10, 9]).

Below are the meshes used in our convergence analysis.

meshTorus 1	meshTorus 2	meshTorus 3	meshTorus 4
·			
1022 cells	6461 cells	20006 cells	43910 cells

Figure 2: Mesh of domain

3.2 Visualization of the results

For the numerical resolution of our discrete problem, we use an iterative solver because the stiffness matrix A_{\triangle_R} is large and sparse (see [7]).

because the stiffness matrix $A_{\triangle_{\Gamma_h}}$ is large and sparse (see [7]) . For the visualization of the result, we use the PARAVIS module included in SALOME (see [9]).

Below are visualizations of the numerical results obtained on the different meshes of picture 2.

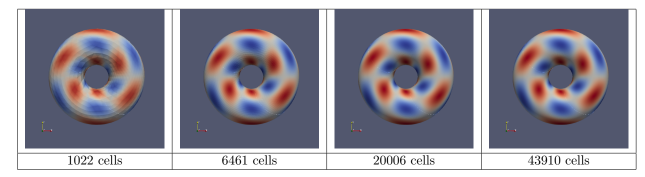


Figure 3: Numerical results of the finite elements on the torus

Below are clipings of the previous numerical results.

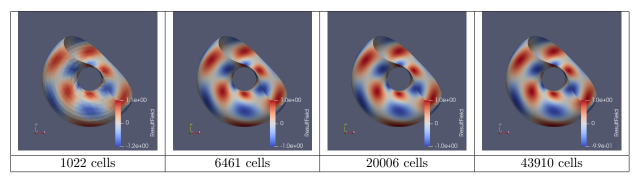


Figure 4: Cliping of the numerical result on the torus

3.3 Numerical convergence of the finite element method

Convergence of finite elements for Laplace operator on 3D torus triangular meshes log(|numerical-exact|) -0.6 least square slope : -1.655 -0.8 -1.0log(error) -1.2 -1.4-1.6-1.8-2.01.8 1.6 1.7 1.9 2.0 2.1 2.2 1.5 2.3 log(sqrt(number of nodes))

Figure 5: Convergence of the finite element method on the torus

The method converges with a numerical order of approximately 1.65.

3.4 Computational time of the finite element method

Computational time of finite elements for Laplace operator on 3D torus triangular meshes 2.0 log(cpu time)) 1.5 1.0 log(cpu time) 0.5 0.0 -0.5 1.7 1.9 2.0 2.1 2.2 1.5 1.6 1.8 2.3 log(sqrt(number of nodes)

Figure 6: Computational time of the finite element method on the torus

3.5 Ploting over slice circle

Here we have drawn a circle on each torus to extract the values. This circle is visible on the torus in Figure 3.

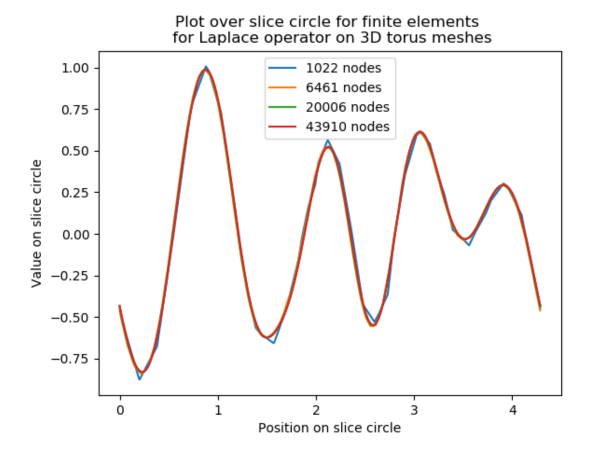


Figure 7: Convergence of the data ploted over a circle drawn on the torus

4 The script

```
Solution exacte = f/12 : il s'agit d'un vecteur
8 #
      propre du laplacien sur le tore
                  Résolution d'un système linéaire à matrice singuliè
9 #
      re : les vecteurs constants sont dans le noyau
10 #
11
12 import cdmath
13 import time, json
14 from math import sin, cos, atan2, sqrt
15 import PV_routines
16 import VTK_routines
17 import paraview.simple as pvs
19 test_desc={}
20 test_desc["Initial_data"]="No"
21 test_desc["Boundary_conditions"]="Dirichlet"
22 test_desc["Global_name"]="FE simulation of the Poisson equation on
      a torus"
23 test_desc["Global_comment"]="Triangular mesh, compact surface (no
      boundary) "
24 test_desc["PDE_model"]="Poisson-Beltrami"
25 test_desc["PDE_is_stationary"]=True
26 test_desc["PDE_search_for_stationary_solution"]=False
27 test_desc["Numerical_method_name"]="P1 FE"
28 test_desc["Numerical_method_space_discretization"]="Finite elements
29 test_desc["Numerical_method_time_discretization"]="None"
30 test_desc["Mesh_is_unstructured"]=True
31 test_desc["Geometry"]="Square"
32 test_desc["Part_of_mesh_convergence_analysis"]=True
33
def solve(filename,resolution,meshType, testColor):
      start = time.time()
35
      test_desc["Mesh_type"]=meshType
36
37
      test_desc["Test_color"]=testColor
38
      # Torus radii (calculation will fail if the mesh is not
39
      correct)
      R=1 #Grand rayon
40
      r=0.6 #Petit rayon
41
42
      #Chargement du maillage triangulaire du tore
43
44
      my_mesh = cdmath.Mesh(filename+".med")
45
      if(not my_mesh.isTriangular()) :
46
          raise ValueError("Wrong cell types : mesh is not made of
      triangles")
      if (my_mesh.getMeshDimension()!=2) :
48
          raise ValueError("Wrong mesh dimension : expected a surface
49
       of dimension 2")
       if (my_mesh.getSpaceDimension()!=3) :
50
          raise ValueError("Wrong space dimension : expected a space
      of dimension 3")
      nbNodes = my_mesh.getNumberOfNodes()
53
54
      nbCells = my_mesh.getNumberOfCells()
55
      test_desc["Space_dimension"] = my_mesh.getSpaceDimension()
56
```

```
test_desc["Mesh_dimension"] = my_mesh.getMeshDimension()
57
             test_desc["Mesh_number_of_elements"] = my_mesh.getNumberOfNodes()
58
            test_desc["Mesh_cell_type"] = my_mesh.getElementTypesNames()
59
60
            print("Mesh building/loading done")
61
            print("nb of nodes=", nbNodes)
print("nb of cells=", nbCells)
62
63
64
             #Discrétisation du second membre et détermination des noeuds
65
            intérieurs
66
             -----
            my_RHSfield = cdmath.Field("RHS_field", cdmath.NODES, my_mesh,
67
            1)
             exactSolField = cdmath.Field("Exact solution field", cdmath.
68
            NODES, my_mesh, 1)
            {\tt maxNbNeighbours} = 0#This is to determine the number of non zero
              coefficients in the sparse finite element rigidity matrix
            #parcours des noeuds pour discrétisation du second membre et
71
            extraction du nb max voisins d'un noeud
             for i in range(nbNodes):
                    Ni=my_mesh.getNode(i)
                    x = Ni.x()
74
                    y = Ni.y()
75
                    z = Ni.z()
76
77
                     theta=atan2(z,sqrt(x*x+y*y)-R)
                    phi=atan2(y,x)
79
80
                    exactSolField[i] = sin(3*phi)*cos(3*theta+ phi) # for the
81
             exact solution we use the funtion given in the article of
            Olshanskii, Reusken 2009, page 19
                    my_RHSfield[i] = 9*sin(3*phi)*cos(3*theta+ phi)/(r*r) +
82
             (10*\sin(3*phi)*\cos(3*theta+phi) + 6*\cos(3*phi)*\sin(3*theta+phi)
            phi))/((R+r*cos(theta))*(R+r*cos(theta))) - 3*sin(theta)*sin(3*sin(theta))*(R+r*cos(theta))) - 3*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*sin(theta)*
            side we use the function given in the article of Olshanskii,
            Reusken 2009, page 19
                    if my_mesh.isBorderNode(i): # Détection des noeuds frontiè
83
                             raise ValueError("Mesh should not contain borders")
84
85
                     else:
86
                             maxNbNeighbours = max(1+Ni.getNumberOfCells(),
             maxNbNeighbours)
87
             test_desc["Mesh_max_number_of_neighbours"]=maxNbNeighbours
88
89
             print("Right hand side discretisation done")
90
             print("Max nb of neighbours=", maxNbNeighbours)
91
             print("Integral of the RHS", my_RHSfield.integral(0))
92
93
            # Construction de la matrice de rigidité et du vecteur second
94
            membre du système linéaire
            Rigidite = cdmath.SparseMatrixPetsc(nbNodes, nbNodes,
96
            \verb|maxNbNeighbours|| \verb|# warning|| : third argument is number of non
             zero coefficients per line
        RHS=cdmath.Vector(nbNodes)
```

```
98
       # Vecteurs gradient de la fonction de forme associée à chaque
       noeud d'un triangle
100
       GradShapeFunc0=cdmath.Vector(3)
       GradShapeFunc1=cdmath.Vector(3)
101
       GradShapeFunc2=cdmath.Vector(3)
102
103
       normalFace0=cdmath.Vector(3)
       normalFace1=cdmath.Vector(3)
105
       #On parcourt les triangles du domaine
108
       for i in range(nbCells):
109
           Ci=my_mesh.getCell(i)
110
111
           #Contribution à la matrice de rigidité
112
           nodeId0=Ci.getNodeId(0)
           nodeId1=Ci.getNodeId(1)
114
           nodeId2=Ci.getNodeId(2)
116
           NO=my_mesh.getNode(nodeIdO)
            N1=my_mesh.getNode(nodeId1)
117
           N2=my_mesh.getNode(nodeId2)
118
119
           #Build normal to cell Ci
120
           normalFace0[0] = Ci.getNormalVector(0,0)
            normalFace0[1]=Ci.getNormalVector(0,1)
           normalFace0[2]=Ci.getNormalVector(0,2)
123
124
           normalFace1[0]=Ci.getNormalVector(1,0)
            normalFace1[1]=Ci.getNormalVector(1,1)
           normalFace1[2]=Ci.getNormalVector(1,2)
126
127
           normalCell = normalFace0.crossProduct(normalFace1)
128
           normalCell = normalCell/normalCell.norm()
129
130
           cellMat=cdmath.Matrix(4)
           cellMat[0,0]=N0.x()
132
            cellMat[0,1]=N0.y()
           cellMat[0,2]=N0.z()
134
135
            cellMat[1,0]=N1.x()
           cellMat[1,1]=N1.y()
136
137
            cellMat[1,2]=N1.z()
            cellMat[2,0]=N2.x()
138
           cellMat[2,1]=N2.y()
139
140
            cellMat[2,2] = N2.z()
141
            cellMat[3,0]=normalCell[0]
           cellMat[3,1]=normalCell[1]
142
            cellMat[3,2]=normalCell[2]
143
            cellMat[0,3]=1
144
           cellMat[1,3]=1
145
            cellMat[2,3]=1
146
           cellMat[3,3]=0
147
148
            #Formule des gradients voir EF P1 -> calcul déterminants
149
            GradShapeFunc0[0] = cellMat.partMatrix(0,0).determinant()/2
150
            GradShapeFunc0[1] = - cellMat.partMatrix(0,1).determinant()/2
151
            GradShapeFunc0[2] = cellMat.partMatrix(0,2).determinant()/2
            GradShapeFunc1[0] = - cellMat.partMatrix(1,0).determinant()/2
153
            GradShapeFunc1[1] = cellMat.partMatrix(1,1).determinant()/2
154
            GradShapeFunc1[2] = - cellMat.partMatrix(1,2).determinant()/2
            GradShapeFunc2[0] = cellMat.partMatrix(2,0).determinant()/2
156
            GradShapeFunc2[1] = - cellMat.partMatrix(2,1).determinant()/2
           GradShapeFunc2[2] = cellMat.partMatrix(2,2).determinant()/2
158
```

```
159
           #Création d'un tableau (numéro du noeud, gradient de la
       fonction de forme
161
           GradShapeFuncs={nodeId0 : GradShapeFunc0}
           GradShapeFuncs[nodeId1]=GradShapeFunc1
162
           GradShapeFuncs[nodeId2]=GradShapeFunc2
163
           # Remplissage de la matrice de rigidité et du second
       membre
           for j in [nodeId0, nodeId1, nodeId2] :
               #Ajout de la contribution de la cellule triangulaire i
167
       au second membre du noeud j
               RHS[j]=Ci.getMeasure()/3*my_RHSfield[j]+RHS[j] # inté
168
       grale dans le triangle du produit f x fonction de base
               \#Contribution de la cellule triangulaire i à la ligne j
        du système linéaire
               for k in [nodeId0, nodeId1, nodeId2] :
                   Rigidite.addValue(j,k,GradShapeFuncs[j]*
       GradShapeFuncs[k]/Ci.getMeasure())
172
173
       print("Linear system matrix building done")
174
       # Résolution du système linéaire
176
       LS = \texttt{cdmath.LinearSolver(Rigidite,RHS,100,1.E-6,"CG","CHOLESKY")}
177
       LS.setMatrixIsSingular()#En raison de l'absence de bord
       LS.setComputeConditionNumber()
179
180
       SolSyst=LS.solve()
181
       print( "Preconditioner used : ", LS.getNameOfPc() )
182
       print( "Number of iterations used : ", LS.getNumberOfIter() )
       print( "Final residual : ", LS.getResidu() )
184
       print("Linear system solved")
185
186
       test_desc["Linear_solver_algorithm"]=LS.getNameOfMethod()
187
       test_desc["Linear_solver_preconditioner"]=LS.getNameOfPc()
188
       test_desc["Linear_solver_precision"]=LS.getTolerance()
189
       test_desc["Linear_solver_maximum_iterations"]=LS.
190
       getNumberMaxOfIter()
       test_desc["Linear_system_max_actual_iterations_number"]=LS.
191
       getNumberOfIter()
       test_desc["Linear_system_max_actual_error"]=LS.getResidu()
       test_desc["Linear_system_max_actual_condition number"]=LS.
       getConditionNumber()
       # Création du champ résultat
196
       my_ResultField = cdmath.Field("ResultField", cdmath.NODES,
       my_mesh, 1)
       for j in range(nbNodes):
           my_ResultField[j]=SolSyst[j];#remplissage des valeurs pour
199
       les noeuds intérieurs
       #sauvegarde sur le disque dur du résultat dans un fichier
200
       paraview
       my_ResultField.writeVTK("FiniteElementsOnTorusPoisson_"+
       meshType+str(nbNodes))
202
       end = time.time()
203
204
205
       print("Integral of the numerical solution", my_ResultField.
       integral(0))
```

```
print("Numerical solution of poisson equation on a torus using
206
       finite elements done")
207
       #Calcul de l'erreur commise par rapport à la solution exacte
208
209
       max_abs_sol_exacte=exactSolField.getNormEuclidean().max()
210
       erreur_abs=(exactSolField - my_ResultField).getNormEuclidean().
       max()
       max_sol_num=my_ResultField.max()
212
       min_sol_num=my_ResultField.min()
214
       print("Absolute error = max(| exact solution - numerical
215
       solution |)/max(| exact solution |) = ",erreur_abs/
       max_abs_sol_exacte)
       print("Maximum numerical solution = ", max_sol_num, " Minimum
       numerical solution = ", min_sol_num)
       print("Maximum exact solution = ", exactSolField.max(), "
       Minimum exact solution = ", exactSolField.min())
218
219
       assert erreur_abs/max_abs_sol_exacte <1.
220
       test desc["Computational time taken by run"] = end-start
221
       test_desc["Absolute_error"] = erreur_abs
       test_desc["Relative_error"] = erreur_abs/max_abs_sol_exacte
223
224
       #Postprocessing:
226
       # save 3D picture
227
       PV_routines.Save_PV_data_to_picture_file("
       FiniteElementsOnTorusPoisson_"+meshType+str(nbNodes)+'_0.vtu',"
       ResultField",'NODES', "FiniteElementsOnTorusPoisson_"+meshType+
       str(nbNodes))
       # save 3D clip
229
       VTK_routines.Clip_VTK_data_to_VTK("
       FiniteElementsOnTorusPoisson_"+meshType+str(nbNodes)+'_0.vtu',"
       Clip_VTK_data_to_VTK_"+ "FiniteElementsOnTorusPoisson_"+
       meshType+str(nbNodes)+'_0.vtu',[0.25,0.25,0.25],
       [-0.5, -0.5, -0.5], resolution)
231
       PV_routines.Save_PV_data_to_picture_file("Clip_VTK_data_to_VTK_
       "+"FiniteElementsOnTorusPoisson_"+meshType+str(nbNodes)+',_0.vtu
       ', "ResultField", 'NODES', "Clip_VTK_data_to_VTK_"+'
       FiniteElementsOnTorusPoisson_"+meshType+str(nbNodes))
       # save plot around circumference
232
       finite Elements On Torus\_Ovtu \ = \ pvs. XMLUnstructured Grid Reader (
233
       FileName = ["FiniteElementsOnTorusPoisson_"+meshType+str(nbNodes)
       +'_0.vtu'])
       slice1 = pvs.Slice(Input=finiteElementsOnTorus_Ovtu)
234
       slice1.SliceType.Normal = [0.5, 0.5, 0.5]
       renderView1 = pvs.GetActiveViewOrCreate('RenderView')
236
       finiteElementsOnTorus_OvtuDisplay = pvs.Show(
       finiteElementsOnTorus_Ovtu , renderView1)
       pvs.ColorBy(finiteElementsOnTorus_OvtuDisplay, ('POINTS', '
238
       ResultField'))
       slice1Display = pvs.Show(slice1, renderView1)
239
       pvs.SaveScreenshot("./FiniteElementsOnTorusPoisson"+"_Slice_"+
       meshType+str(nbNodes)+'.png', magnification=1, quality=100,
       view=renderView1)
       plotOnSortedLines1 = pvs.PlotOnSortedLines(Input=slice1)
       pvs.SaveData('./FiniteElementsOnTorusPoisson_PlotOnSortedLines')
242
       +meshType+str(nbNodes)+'.csv', proxy=plotOnSortedLines1)
```

```
with open('test_Poisson'+str(my_mesh.getMeshDimension())+'D_EF_
'+meshType+str(nbCells)+ "Cells.json", 'w') as outfile:
    json.dump(test_desc, outfile)

return erreur_abs/max_abs_sol_exacte, nbNodes, min_sol_num,
    max_sol_num, end - start

if __name__ == """__main__""":
    solve("meshTorus",100,"Unstructured_3D_triangles","Green")
```

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