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AN ADAPTIVE QUADRILATERAL MESH IN CURVED DOMAINS

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ABSTRACT. An nonlinear elliptic system for generating adaptive quadrilateral meshes in curved domains is presented. The presented technique has been implemented in the C++ language with the help of the standard template library. The software package writes the converged meshes in the GMV and the Matlab formats. Grid generation is the first very important step for numerically solving partial differential equations. Thus, the presented C++ grid generator is extremely important to the computational science community.

1. Introduction. Quadrilateral meshes are used for visualization, interpolation and numerically solving partial differential equations. The accuracy of a numerical solution of partial differential equation strongly depends on the quality of the underlying mesh [22, 2, 3, 4, 7, 5]. Here, quality means orthogonality at the boundaries and quasi-orthogonality within the critical regions, smoothness, bounded aspect ratios and solution adaptive behavior.

ACM Computing Classification System (1998): G.1.0, G.4, G.1.7.

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Grid adaptation is used for increasing the efficiency of numerical schemes by focusing the computational effort where it is needed [19, 18]. In this work, we present the elliptic grid generation system for generating adaptive quadrilateral meshes. The presented method has been implemented in the C++ language. The presented method generates adaptive meshes without destroying the structured nature of the mesh. It is easier to develop solvers based on a structured mesh than on an unstructured mesh [20]. There are various software packages available for generating adaptive triangular meshes, but we do not know of any software package that can be used for generating adapting quadrilateral meshes. Thus, the presented method and its C++ implementation are very useful for solving partial differential equations.

For meshing a domain into non-simplex elements (quadrilaterals in 2D and hexahedra in 3D), we seek a mapping from a reference square or cube to the physical domain. This mapping can be algebraic in nature such as Transfinite Interpolation, or it can be expressed by a system of nonlinear partial differential equations such as an elliptic system [6, 8, and references therein]. We are looking for a vector mapping, $\mathcal{F}_k(\hat{k}) = (x, y)^t$, from a unit square in the reference space ($\hat{k} = [0, 1] \times [0, 1]$) to a physical space (k), that is $\mathcal{F}_k: \hat{k} \mapsto k$. See Figure 1. Mapping \mathcal{F}_k gives the position of a point in the physical space corresponding to a point in the computational or reference space. Let the physical space be given by the x and y coordinates, and the computational space be given by the ξ and η coordinates. Here, $\xi \in [0, 1]$ and $\eta \in [0, 1]$. We are using the following elliptic system for defining the mapping $\mathcal{F}_k = (x, y)^t$

$$(1) \quad g_{22} \frac{\partial^2 x}{\partial \xi^2} - 2 g_{12} \frac{\partial^2 x}{\partial \xi \partial \eta} + g_{11} \frac{\partial^2 x}{\partial \eta^2} + P x_\xi + Q x_\eta = 0,$$

$$(2) \quad g_{22} \frac{\partial^2 y}{\partial \xi^2} - 2 g_{12} \frac{\partial^2 y}{\partial \xi \partial \eta} + g_{11} \frac{\partial^2 y}{\partial \eta^2} + P y_\xi + Q y_\eta = 0.$$

Here, terms P and Q are used for grid adaptation and are given as

$$(3) \quad P = g_{22} P_{11}^1 - 2 g_{12} P_{12}^1 + g_{11} P_{22}^1,$$

$$(4) \quad Q = g_{22} P_{11}^2 - 2 g_{12} P_{12}^2 + g_{11} P_{22}^2.$$

Equations (1–2) are non-linear and are coupled through metric coefficients g_{ij} (coefficients of the metric tensor). Metric coefficients are given as

$$(5) \quad g_{11} = x_\xi^2 + y_\xi^2, \quad g_{22} = x_\eta^2 + y_\eta^2 \quad \text{and} \quad g_{12} = x_\xi x_\eta + y_\xi y_\eta.$$

For generating grids in the physical space, elliptic system (1–2) is solved for coordinates (x, y) on a unit square in the computational space by the method of Finite Differences. The boundary of the physical domain is specified as the Dirichlet boundary condition on the unit square in the computational space. In Figure 1, $\mathbf{g}_1 (= \mathbf{r}_\xi)$ and $\mathbf{g}_2 (= \mathbf{r}_\eta)$ are the covariant base vectors at point (x_i, y_j) . Figure 2 shows a finite difference stencil around point (ξ_i, η_j) in the computational space. A finite difference approximation of x_ξ and x_η at point (i, j) (see Figure 2) is

$$x_\xi = \frac{[x(i + 1, j) - x(i - 1, j)]}{2 \Delta\xi} \quad \text{and} \quad x_\eta = \frac{[x(i, j + 1) - x(i, j - 1)]}{2 \Delta\eta}.$$

Similarly, y_ξ and y_η can be defined. Here, we are assuming that the grid in the computational space is uniform. However, the grid in the physical space can be compressed or stretched.

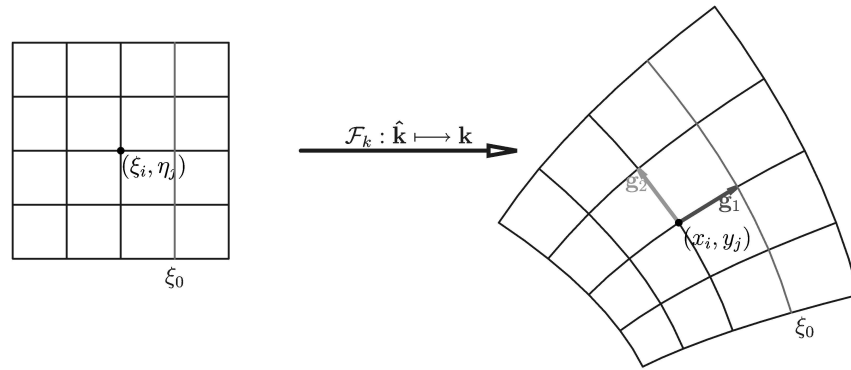


Fig. 1. Mapping \mathcal{F}_k from a reference unit square (\hat{k}) on the left to a physical domain (k)

Terms P_{ij}^k in equations (1–2) are determined through another mapping \mathcal{F}_1 . Here, $i = 1, 2; j = 1, 2; k = 1, 2$ and $P_{12}^k = P_{21}^k$. Mapping \mathcal{F}_1 is shown in Figure 3. This mapping maps a unit square in the computational space to a unit square in the parameter space. For defining mapping $\mathcal{F}_1: \hat{k} \rightarrow k_1$, the boundary and internal grid points of the parameter space are mapped to the reference space. The Jacobian matrix \mathbf{T} of mapping \mathcal{F}_1 and vectors \mathbf{P}_{11} , \mathbf{P}_{12} and \mathbf{P}_{22} are given as follows

$$(6) \quad \mathbf{T} = \begin{pmatrix} s_\xi & s_\eta \\ t_\xi & t_\eta \end{pmatrix}, \quad \mathbf{P}_{11} = -\mathbf{T}^{-1} \begin{pmatrix} s_{\xi\xi} \\ t_{\xi\xi} \end{pmatrix},$$

$$(7) \quad \mathbf{P}_{22} = -\mathbf{T}^{-1} \begin{pmatrix} s_{\eta\eta} \\ t_{\eta\eta} \end{pmatrix}, \quad \mathbf{P}_{12} = -\mathbf{T}^{-1} \begin{pmatrix} s_{\xi\eta} \\ t_{\xi\eta} \end{pmatrix}.$$

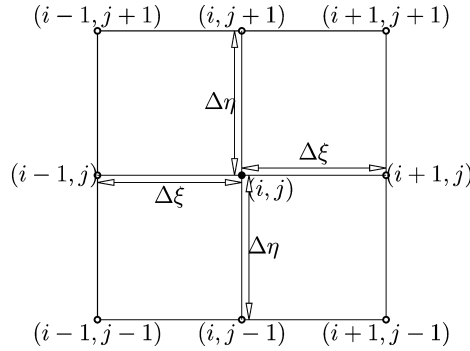


Fig. 2. Finite difference stencil in the ξ - η computational space

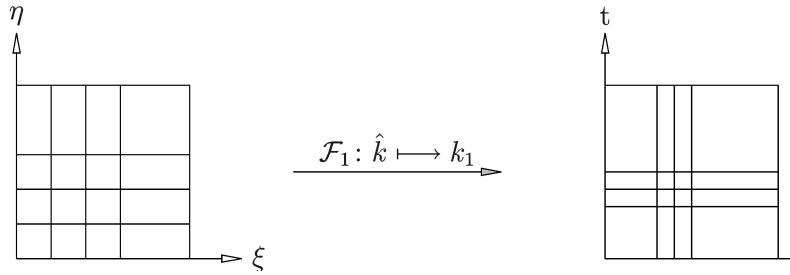


Fig. 3. Mapping \mathcal{F}_1 from a unit square (\hat{k}) in the reference space to a unit square in the parameter space (k_1)

Terms P_{ij}^1 ($i, j = 1, 2$) are the first component of vector \mathbf{P}_{ij} , and terms P_{ij}^2 are the second component of the vector \mathbf{P}_{ij} . It should be noted that vectors \mathbf{P}_{11} , \mathbf{P}_{12} and \mathbf{P}_{22} can be computed a priori for clustering the grid points in the physical space. A second-order finite difference approximation of different operators required for computing vectors \mathbf{P}_{11} , \mathbf{P}_{22} , \mathbf{P}_{12} and the Jacobian \mathbf{T} are given in Table 1. We are using the stencil shown in Figure 2.

2. C++ Implementation. We have implemented the presented technique in the C++ language for generating adaptive grids. The software package can write meshes in the Matlab and GMV [21] formats. It consists of one Domain class. See subsections 2.2 and 2.3. Domain class is used for representing the unit square in the computational space, parameter space and the physical domain. See line numbers **44**, **22** and **37** in subsection 2.1.

The physical domain is defined in the file functions.h in subsection 2.6. For clustering grids in the parameter space different functions are defined in the domain class. See line numbers **25**, **26**, **28**, **29**, **31**, **32** in subsection 2.2.

Table 1. Finite difference approximation of continuous operators

$$\begin{aligned}
s_{\xi} &= \frac{s(i+1, j) - s(i-1, j)}{2 \Delta \xi}, & s_{\xi\xi} &= \frac{s(i+1, j) - 2s(i, j) + s(i-1, j)}{\Delta \xi^2} \\
t_{\xi} &= \frac{t(i, j+1) - t(i, j-1)}{2 \Delta \eta}, & t_{\eta\eta} &= \frac{t(i, j+1) - 2t(i, j) + t(i, j-1)}{\Delta \eta^2} \\
s_{\eta\eta} &= \frac{s(i, j+1) - 2s(i, j) + s(i, j-1)}{\Delta \eta^2}, & t_{\xi\xi} &= \frac{t(i+1, j) - 2t(i, j) + t(i-1, j)}{\Delta \xi^2} \\
s_{\xi\eta} &= \frac{s(i+1, j+1) + s(i-1, j-1) - s(i-1, j+1) - s(i+1, j-1)}{4 \Delta \xi \Delta \eta} \\
t_{\xi\eta} &= \frac{t(i+1, j+1) + t(i-1, j-1) - t(i-1, j+1) - t(i+1, j-1)}{4 \Delta \xi \Delta \eta}
\end{aligned}$$

The coupled elliptic system are linearised by the method of Finite Differences, and the resulting system is solved by the SOR relaxation (see subsection 2.5). The SOR algorithm consists of three loops: the outer loop (see line number **42** in subsection 2.5) and two inner for loops (see line numbers **45** and **46** in subsection 2.5). Each iteration of an inner loop provides a new mesh by the SOR relaxation. The outer loop is controlled by the maximum number of SOR iterations (see line number **28** in subsection 2.5) and a given tolerance (see line number **26** in subsection 2.5).

The overall algorithm proceeds as follows. Generate grids in the computational and parameter spaces. Compute matrix \mathbf{T} and vectors \mathbf{P}_{ij} for defining mapping \mathcal{F}_1 (from computational space to parameter space). An initial grid, \mathbf{r}_{old} , in the physical region is generated by Transfinite Interpolation. This information is then passed to the SOR solver (see line number **42** in subsection 2.1).

2.1. main.cpp

```

1 //+++++
2 #include <iostream>
3 #include <iomanip>
4 #include <vector>
5 #include <iterator>
6 #include <fstream>
7 #include <sstream>
8 #include <map>
9 #include "domain.h"
10 #include "matrix.h"
11 #include "sor_solver.cpp"
12 //+++++
13 int main(){

```

```

15     bool grid_dist = true;
16     bool run_ellip = true;
17     unsigned xdim, ydim ;
18     xdim = 31, ydim = 31;
19     double del_xi = 1.0/double(xdim-1.0);
20     double del_eta = 1.0/double(ydim-1.0);
21     //Parameter Space ref(xdim, ydim);
22     Domain_parm(xdim, ydim);
23     //Meshing the Parameter
24     parm.Grid_Gen();
25     //Clustering the Mesh
26     //Example 1
27     //parm.Cluster_X_Near(0.5);
28     //parm.Cluster_Y_Near(0.5);
29     //Example 2
30     //parm.Cluster_Two_Lines_X(0.25,0.750);
31     //parm.Cluster_Two_Lines_Y(0.25,0.750);
32     //Example 3
33     parm.Bound_Clust_X(0.5);
34     parm.Bound_Clust_Y(0.5);
35
36     parm.Fill_del_xi_eta(del_xi, del_eta);
37
38     Domain_physical(xdim, ydim);
39     physical.Read_Bd();
40     physical.Fill_del_xi_eta(del_xi, del_eta);
41     unsigned max_iter = 100;
42     double w = 1.90;
43     SORSOLVER(physical, parm, xdim, ydim);
44     //Reference or computational space
45     Domain_ref(xdim, ydim);
46
47     //Writing the mesh in the physical space (GMV)
48     std::ofstream outPhy("gmv_Physical.dat", std::ios::out);
49     if (!(outPhy)) std::cerr << "ERROR: UNABLE TO OPEN "outPhy"\n";
50     physical.GMV_Writer(outPhy);
51     if (outPhy.is_open()) outPhy.close();
52
53     //writing mesh in the parameter space (GMV)
54     std::ofstream outParm("gmv_Para.dat", std::ios::out);
55     if (!(outParm)) std::cerr << "ERROR: UNABLE TO OPEN "outParm"\n";
56     parm.GMV_Writer(outParm);
57     if (outParm.is_open()) outParm.close();
58     parm.Matlab_Writer();
59
60     return EXIT_SUCCESS;
61 }

```

2.2. domain.h

```

1 #ifndef PARAMETER_SPACE
2 #define PARAMETER_SPACE
3 //+++++
4 #include<iostream>
5 #include<iomanip>
6 #include<vector>
7 #include<iterator>
8 #include<fstream>
9 #include<sstream>
10 #include<map>
11 //+++++
12 #include "matrix.h"
13 //+++++
14 class Domain{
15 public:
16     Domain();
17     Domain(unsigned int xdim1, unsigned ydim1);
18     Domain(const Domain & org);
19     unsigned int XDIM() const ;
20     unsigned int YDIM() const ;
21     void Grid_Gen();

```

```

22     std::vector<double> XCOORDS();
23     std::vector<double> YCOORDS();
24     double Eriksson_1(double eta);

26     void Cluster_X_Near(double eta0);
27     void Cluster_Y_Near(double eta0);

28
29     void Cluster_Two_Lines_X(double eta1,double eta2);
30     void Cluster_Two_Lines_Y(double eta1,double eta2);

32     void Bound_Clust_X(double eta1);
33     void Bound_Clust_Y(double eta1);

34
35     double& XCOORD(unsigned int i , unsigned int j);
36     double& YCOORD(unsigned int i , unsigned int j);
37     void Read_Bd();
38     void Matlab_Writer();
39     void GMV_Writer(std::ofstream & outFile);
40     void Fill_del_xi_eta(double xi,double eta);
41     //void Call_Grid_Adapter();

42
43     //+++++++
44     std::vector<double> P11(unsigned int i , unsigned int j);
45     std::vector<double> P22(unsigned int i , unsigned int j);
46     std::vector<double> P12(unsigned int i , unsigned int j);
47     //+++++++
48     Matrix MeshX();
49     Matrix MeshY();
50     //+++++++
51     double G22(unsigned int i , unsigned int j);
52     double G11(unsigned int i , unsigned int j);
53     double X_xi(unsigned int i , unsigned int j);
54     double Y_xi(unsigned int i , unsigned int j);
55     double X_eta(unsigned int i , unsigned int j);
56     double Y_eta(unsigned int i , unsigned int j);
57     double X_xieta(unsigned int i , unsigned int j);
58     double Y_xieta(unsigned int i , unsigned int j);

60 private:
61     double del_eta , del_xi;
62     unsigned xdim , ydim;
63     Matrix x , y;
64     std::vector<double> xcoords , ycoords;
65 };
66 #endif

```

2.3. domain.cpp

```

#include "domain.h"
2
#ifdef _FUNCTIONS_
4 #include "functions.h"
5 #endif
6
7 #include <cassert>
8
9 Domain::Domain(){
10     xdim = 0 ; ydim = 0;
11 }
12 Domain::Domain(unsigned int xdim1,unsigned int ydim1){
13     xdim = xdim1 ; ydim = ydim1;
14 }
15 Domain::Domain(const Domain & org){
16     xdim = org.XDIM();
17     ydim = org.YDIM();
18     Grid_Gen();
19 }
20 unsigned int Domain::XDIM() const{
21     return xdim;
22 }
23 unsigned int Domain::YDIM() const{

```

```

24     return ydim;
25 }
26 void Domain::Grid_Gen(){
27     Matrix xt(xdim, ydim), yt(xdim, ydim);
28     assert(0 != xdim && 0 != ydim);
29     for (unsigned int j = 0 ; j < ydim ; ++j){
30         for (unsigned int i = 0 ; i < xdim ; ++i){
31             double t_x = double(i)/double(xdim-1.0);
32             double t_y = double(j)/double(ydim-1.0);
33             xt(i, j) = t_x ; yt(i, j) = t_y;
34         }
35     }
36     x = xt ; y = yt;
37 }
38 std::vector<double> Domain::XCOORDS(){
39     xcoords.resize(xdim*ydim); ycoords.resize(xdim*ydim);
40     for (int j = 0 ; j < ydim ; ++j){
41         for (int i = 0 ; i < xdim ; ++i){
42             int no = i+j*xdim;
43             xcoords[no] = x(i, j);
44             ycoords[no] = y(i, j);
45         }
46     }
47     return xcoords;
48 }
49 std::vector<double> Domain::YCOORDS(){
50     xcoords.resize(xdim*ydim); ycoords.resize(xdim*ydim);
51     for (int j = 0 ; j < ydim ; ++j){
52         for (int i = 0 ; i < xdim ; ++i){
53             int no = i+j*xdim;
54             xcoords[no] = x(i, j);
55             ycoords[no] = y(i, j);
56         }
57     }
58     return ycoords;
59 }
60 void Domain::Bound_Clust_X(double eta1){
61
62     double alpha = 4.0;
63     double h = 1.0;
64     double h2 = 1.0;
65     double h1 = 0.0;
66
67     for (int j = 0 ; j < ydim ; ++j){
68         for (int i = 0 ; i < xdim ; ++i){
69
70             if (x(i, j) <= eta1 && 0 <= x(i, j)){
71                 double eta = x(i, j);
72                 x(i, j) = (h2-h1)* eta1*(std::exp(alpha*eta/eta1)-1.0)/(std::exp(alpha)-1.0)+h1;
73             }
74
75             if (x(i, j) >= eta1 && x(i, j) <= 1.0){
76                 double eta = x(i, j);
77                 x(i, j) = (h2-h1)*(1.0-(1.0-eta1)*(((std::exp(alpha*(1.0-eta)/
78                 (1.0-eta1)))-1.0)/(std::exp(alpha)-1.0))); }
79
80         }
81     }
82 }
83
84 }
85
86 void Domain::Bound_Clust_Y(double eta1){
87
88     double alpha = 4.0;
89     double h = 1.0;
90     double h2 = 1.0;
91     double h1 = 0.0;
92
93     for (int j = 0 ; j < ydim ; ++j){
94         for (int i = 0 ; i < xdim ; ++i){
95
96             if (y(i, j) <= eta1 && 0 <= y(i, j)){
97                 double eta = y(i, j);

```



```

98         }   y(i,j) =(h2-h1)* eta1*(std::exp(alpha*eta/eta1)-1.0)/(std::exp(alpha)-1.0)+h1;
100     }
101     if (y(i,j) >= eta1 && y(i,j) <= 1.0){
102         double eta = y(i,j);
103         y(i,j) = (h2-h1)*(1.0-(1.0-eta1)*(((std::exp(alpha*(1.0-eta)/
104             (1.0-eta1)))-1.0)/(std::exp(alpha)-1.0)));}
105     }
106 }
107 }
108 }
109 //=====
110 double Domain::Eriksson_1(double eta){
111     double h = 1.0;
112     double alpha = 3.0;
113
114     return h*((std::exp(alpha*eta)-1.0)/(std::exp(alpha)-1.0));
115 }
116
117 void Domain::Cluster_Two_Lines_X(double eta1,double eta2){
118     double alpha = 5.0;
119     double h = 1.0;
120     double eta0 = (eta1+eta2)*0.5;
121
122     for (int j = 0 ; j < ydim ; ++j){
123         for (int i = 0 ; i < xdim ; ++i){
124
125             if (x(i,j) <= eta1 && 0 <= x(i,j)){
126                 double eta = x(i,j);
127                 x(i,j) = eta1*(h-Eriksson_1(1-eta/eta1));
128             }
129
130             if (x(i,j) >= eta1 && x(i,j) <= eta0 ){
131                 double eta = x(i,j);
132                 x(i,j) = h*eta1+ (eta0-eta1)*Eriksson_1((eta-eta1)/(eta0-eta1));
133             }
134
135             if (x(i,j) >= eta0 && x(i,j) <= eta2){
136                 double eta = x(i,j);
137                 x(i,j) = h*eta0 + (eta2-eta0)*(h-Eriksson_1((eta2-eta)/(eta2-eta0)));
138             }
139
140             if (x(i,j) >= eta2 && x(i,j) <= 1.0){
141                 double eta = x(i,j);
142                 x(i,j) = h*eta2 + (1.0-eta2)*Eriksson_1((eta-eta2)/(1.0-eta2));
143             }
144         }
145     }
146 }
147 }
148 }
149
150 void Domain::Cluster_Two_Lines_Y(double eta1,double eta2){
151     double alpha = 5.0;
152     double h = 1.0;
153     double eta0 = (eta1+eta2)*0.5;
154
155     for (int j = 0 ; j < ydim ; ++j){
156         for (int i = 0 ; i < xdim ; ++i){
157
158             if (y(i,j) <= eta1 && 0 <= y(i,j)){
159                 double eta = y(i,j);
160                 y(i,j) = eta1*(h-Eriksson_1(1-eta/eta1));
161             }
162
163             if (y(i,j) >= eta1 && y(i,j) <= eta0 ){
164                 double eta = y(i,j);
165                 y(i,j) = h*eta1+ (eta0-eta1)*Eriksson_1((eta-eta1)/(eta0-eta1));
166             }
167
168             if (y(i,j) >= eta0 && y(i,j) <= 1.0){
169                 double eta = y(i,j);
170                 y(i,j) = h*eta0 + (1.0-eta0)*(h-Eriksson_1((eta-eta0)/(1.0-eta0)));
171             }
172         }
173     }
174 }

```

```

172         y(i,j) = h*eta1+ (eta0-eta1)*Eriksson_1((eta-eta1)/(eta0-eta1));
174     }
176     if (y(i,j) >= eta0 && y(i,j) <= eta2){
177         double eta = y(i,j);
178         y(i,j) = h*eta0 + (eta2-eta0)*(h-Eriksson_1((eta2-eta)/(eta2-eta0)));
179     }
180     if (y(i,j) >= eta2 && y(i,j) <= 1.0){
181         double eta = y(i,j);
182         y(i,j) = h*eta2 + (1.0-eta2)*Eriksson_1((eta-eta2)/(1.0-eta2));
183     }
184 }
185 }
186 }
187 }
188 }
189 }
190 //=====
191 void Domain::Cluster_X_Near(double eta0){
192     double alpha = 3.0;
193     for (int j = 0 ; j < ydim ; ++j){
194         for (int i = 0 ; i < xdim ; ++i){
195             if (x(i,j) < eta0){
196                 double eta = x(i,j);
197                 x(i,j) = (double) eta0*(exp(alpha)-exp(alpha*(double)(1-eta/eta0)))/
198                     (exp(alpha)-0.1e1);
199             }
200             if (x(i,j) > eta0){
201                 double eta = x(i,j);
202                 x(i,j) = (double) eta0+(double)(1-eta0)*(exp((double)(alpha*(eta-eta0)/
203                     (1-eta0)))-0.1e1)/(exp((double) alpha)-0.1e1);
204             }
205         }
206     }
207 }
208 void Domain::Cluster_Y_Near(double eta0){
209     double alpha = 3.0;
210     for (int j = 0 ; j < ydim ; ++j){
211         for (int i = 0 ; i < xdim ; ++i){
212             if (y(i,j) < eta0){
213                 double eta = y(i,j);
214                 y(i,j) = (double)eta0*(exp(alpha)-exp(alpha*(double)(1-eta/eta0)))/
215                     (exp(alpha)-0.1e1);
216             }
217             if (y(i,j) > eta0){
218                 double eta = y(i,j);
219                 y(i,j) = (double)eta0+(double)(1-eta0)*(exp((double)(alpha*(eta-eta0)/
220                     (1-eta0)))-0.1e1)/(exp((double) alpha)-0.1e1);
221             }
222         }
223     }
224 }
225 double& Domain::XCOORD(unsigned int i , unsigned int j) {
226     if (i >= xdim || j >= ydim || i < 0 || j < 0){
227         std::cerr << "In_XCOORDS(...)_dim_mismatch\n";
228     }
229     return x(i,j);
230 }
231 double& Domain::YCOORD(unsigned int i , unsigned int j){
232     if (i >= xdim || j >= ydim || i < 0 || j < 0){
233         std::cerr << "In_YCOORDS(...)_dim_mismatch\n";
234     }
235     return y(i,j);
236 }
237 void Domain::Read_Bd(){
238     //Read the boundary of the physical domain
239     Matrix xt(xdim,ydim),yt(xdim,ydim);
240     for (int j = 0 ; j < ydim ; ++j){
241         for (int i = 0 ; i < xdim ; ++i){
242             if (0 == i || xdim-1 == i || 0 == j || ydim-1 == j){
243                 double zeta1 = double(i)/double(xdim-1.0);
244                 double eta1 = double(j)/double(ydim-1.0);
245                 xt(i,j) = zeta1; yt(i,j) = eta1;

```

```

246         xt(i,j) = XYcircle(zetal,etal,1);
248         yt(i,j) = XYcircle(zetal,etal,2);
250     }
252 }
//Create grid by the TFI
254 for (int j = 1 ; j < ydim-1 ; ++j){
    for (int i = 1 ; i < xdim-1 ; ++i){
256         double zetal = double(i)/double(xdim-1.0);
                double etal = double(j)/double(ydim-1.0);
258         xt(i,j)=(1.0-zetal)*xt(0,j)+zetal*xt(xdim-1,j)+(1.0-etal)*xt(i,0)+etal*xt(i,ydim-1)-
                ((1.0-zetal)*(1.0-etal)*xt(0,0) + (zetal)*(1.0-etal)*xt(xdim-1,0)
260                 + (zetal)*(etal)*xt(xdim-1,ydim-1) + (1.0-zetal)*etal*xt(0,ydim-1));
                yt(i,j)=(1-zetal)*yt(0,j)+zetal*yt(xdim-1,j) + (1.0-etal)*yt(i,0)+etal*yt(i,ydim-1)-
262                 ((1-zetal)*(1-etal)*yt(0,0) + (zetal)*(1-etal)*yt(xdim-1,0)
                + (zetal)*(etal)*yt(xdim-1,ydim-1)+ (1-zetal)*etal*yt(0,ydim-1));
264     }
    }
266     x = xt ; y = yt ;
}
//=====
268 void Domain::Matlab_Writer(){
270     std::vector<double> x1 = XCOORDS();
272     std::vector<double> y1 = YCOORDS();
    std::vector<double>::const_iterator viter;
274     std::ofstream outfile("matlab_out.m",std::ios::out);
    if (!outfile) std::cerr << "Unable to open the matlab outfile\n";
276     outfile << "clear;\n";
    outfile << "holdon=ishold;\n";
278     for (int j = 0 ; j < ydim ; ++j){
280         for (int i = 0 ; i < xdim ; ++i){
                int no = i + j*xdim;
282                 outfile << "x1(" << i+1 << ", " << j+1 << ")=" << x1[no] << ";\n";
                << "y1(" << i+1 << ", " << j+1 << ")=" << y1[no] << ";\n";
284         }
    }
286     outfile << "m=\n" << xdim << std::endl;
288     outfile << "n=\n" << ydim << std::endl;
290     outfile << "plot(x1(1,:),y1(1,:),'r');\nholdon" << std::endl;
    outfile << "plot(x1(m,:),y1(m,:),'r');" << std::endl;
292     outfile << "plot(x1(:,1),y1(:,1),'r');" << std::endl;
    outfile << "plot(x1(:,n),y1(:,n),'r');" << std::endl;
294     outfile << "%Plot internal grid lines\n";
296     outfile << "for i=2:m-1, plot(x1(i,:),y1(i,:),'b');\nend\n";
    outfile << "for j=2:n-1, plot(x1(:,j),y1(:,j),'b');\nend\n";
298     outfile << "if(~holdon), hold off, end" << std::endl;
300     outfile << "axis off;\n";
302     outfile.close();
304 }
306 void Domain::GMV_Writer(std::ofstream & outFile){
308     std::vector<double> xcoords = XCOORDS();
    std::vector<double> ycoords = YCOORDS();
310     outFile << "gmvinput_ascii\n";
    outFile << "nodes\n" << xdim*ydim << std::endl;
312     for (int j = 0 ; j < ydim ; ++j){
        for (int i = 0 ; i < xdim ; ++i){
314             int no = i + j*xdim;
            outFile << xcoords[no] << ";\n";
316         }
    }
318 }

```

```

320     }
321     outFile << std::endl << std::endl;
322     //writing y coord
323     for (int j = 0 ; j < ydim ; ++j){
324         for (int i = 0 ; i < xdim ; ++i){
325             int no = i + j*xdim;
326             outFile << ycoords[no] << "UUUUUUUU" ;
327         }
328     }
329     outFile << std::endl << std::endl;
330     //forming cells
331     outFile << "cellsUU" << (xdim-1)*(ydim-1) << std::endl;
332     for (int j = 0 ; j < (ydim-1) ; ++j){
333         for (int i = 0 ; i < (xdim-1) ; ++i){
334             int no = ( i + j*(xdim) )+1;
335             int no1 = ( i + (j+1)*(xdim)+1);
336             outFile << "quadUU4UU" << std::endl;
337             outFile << no << "UUU" << no+1 << "UUU"
338                 << no+1 << "UU" << no1 << std::endl;
339         }
340     }
341     outFile << std::endl;
342
343     outFile << std::endl << "endgmv\n";
344     outFile.close();
345 }
346 //+++++
347 Matrix Domain::MeshX(){
348     return x;
349 }
350 Matrix Domain::MeshY(){
351     return y;
352 }
353 //+++++
354 double Domain::G22(unsigned int i , unsigned int j){
355     double x1;double x2;double y1;double y2;
356     x1 = x(i,j-1); x2 = x(i,j+1) ;
357     y1 = y(i,j-1); y2 = y(i,j+1);
358     double g22 = std::pow((x2-x1)/(2.0*del_eta),2) +
359                 std::pow((y2-y1)/(2.0*del_eta),2);
360     return g22;
361 }
362 double Domain::G11(unsigned int i , unsigned int j){
363     double x1 = x(i-1,j); double x2 = x(i+1,j);
364     double y1 = y(i-1,j); double y2 = y(i+1,j);
365     double g11 = std::pow((x2-x1)/(2.0*del_xi),2) +
366                 std::pow((y2-y1)/(2.0*del_xi),2);
367     return g11;
368 }
369 double Domain::X_xi(unsigned int i , unsigned int j){
370     double x_xi;
371     x_xi = (x(i+1,j)-x(i-1,j))/(2.0*del_xi);
372     return x_xi;
373 }
374 double Domain::X_eta(unsigned int i , unsigned int j){
375     double x_eta;
376     x_eta = (x(i,j+1)-x(i,j-1))/(2.0*del_eta);
377     return x_eta;
378 }
379 double Domain::Y_xi(unsigned int i , unsigned int j){
380     double y_xi;
381     y_xi = (y(i+1,j)-y(i-1,j))/(2.0*del_xi);
382     return y_xi;
383 }
384 double Domain::Y_eta(unsigned int i , unsigned int j){
385     double y_eta;
386     y_eta = (y(i,j+1)-y(i,j-1))/(2.0*del_eta);
387     return y_eta;
388 }
389 double Domain::X_xieta(unsigned int i , unsigned int j){
390     return (x(i+1,j+1)+x(i-1,j-1)-x(i-1,j+1)-x(i+1,j-1))/(4.0*del_xi*del_eta);
391 }

```

```

394 double Domain::Y_xieta(unsigned int i , unsigned int j){
    return (y(i+1,j+1)+y(i-1,j-1)-y(i-1,j+1)-y(i+1,j-1))/(4.0*del_xi*del_eta);
396 }
void Domain::Fill_del_xi_eta(double xi,double eta){
398     del_xi = xi; del_eta = eta;
    }
400 std::vector<double> Domain::P11(unsigned int i , unsigned int j){
    //x-t coordinate
402     //compute the jacobian at the point
    double s_xi = (x(i+1,j)-x(i-1,j))/(2.0*del_xi);
404     double s_eta = (x(i,j+1)-x(i,j-1))/(2.0*del_eta);
    double t_xi = (y(i+1,j)-y(i-1,j))/(2.0*del_xi);
406     double t_eta = (y(i,j+1)-y(i,j-1))/(2.0*del_eta);
    double det = s_xi*t_eta-t_xi*s_eta;
408     double Tl_11 = t_eta/det; double Tl_12 = -t_xi/det;
    double Tl_21 = -s_eta/det; double Tl_22 = s_xi/det;
410     double s_xixi = (x(i+1,j)-2.0*x(i,j)+x(i-1,j))/(del_xi*del_xi);
    double s_etaeta = (x(i,j+1)-2.0*x(i,j)+x(i,j-1))/(del_eta*del_eta);
412     double s_xieta = (x(i+1,j+1)+x(i-1,j-1)-x(i-1,j+1)-x(i+1,j-1))/(4.0*del_xi*del_eta);
    double t_xieta = (y(i+1,j+1)+y(i-1,j-1)-y(i-1,j+1)-y(i+1,j-1))/(4.0*del_xi*del_eta);
414     double t_xixi = (y(i+1,j)-2.0*y(i,j)+y(i-1,j))/(del_xi*del_xi);
    double t_etaeta = (y(i,j+1)-2.0*y(i,j)+y(i,j-1))/(del_eta*del_eta);
416     std::vector<double> P11(2);

    P11[0] = -(s_xixi*Tl_11+t_xixi*Tl_12);
    P11[1] = -(s_xixi*Tl_21+t_xixi*Tl_22);
420     return P11;
422 }
std::vector<double> Domain::P22(unsigned int i , unsigned int j){
424     //x-t coordinate
    //compute the jacobian at the point
426     double s_xi = (x(i+1,j)-x(i-1,j))/(2.0*del_xi);
    double s_eta = (x(i,j+1)-x(i,j-1))/(2.0*del_eta);
428     double t_xi = (y(i+1,j)-y(i-1,j))/(2.0*del_xi);
    double t_eta = (y(i,j+1)-y(i,j-1))/(2.0*del_eta);
430     double det = s_xi*t_eta-t_xi*s_eta;
    double Tl_11 = t_eta/det; double Tl_12 = -t_xi/det;
432     double Tl_21 = -s_eta/det; double Tl_22 = s_xi/det;
    double s_xixi = (x(i+1,j)-2.0*x(i,j)+x(i-1,j))/(del_xi*del_xi);
434     double s_etaeta = (x(i,j+1)-2.0*x(i,j)+x(i,j-1))/(del_eta*del_eta);
    double s_xieta = (x(i+1,j+1)+x(i-1,j-1)-x(i-1,j+1)-x(i+1,j-1))/(4.0*del_xi*del_eta);
436     double t_xieta = (y(i+1,j+1)+y(i-1,j-1)-y(i-1,j+1)-y(i+1,j-1))/(4.0*del_xi*del_eta);
    double t_xixi = (y(i+1,j)-2.0*y(i,j)+y(i-1,j))/(del_xi*del_xi);
438     double t_etaeta = (y(i,j+1)-2.0*y(i,j)+y(i,j-1))/(del_eta*del_eta);
    std::vector<double> P22(2);
440     P22[0] = -(s_etaeta*Tl_11+t_etaeta*Tl_12);
    P22[1] = -(s_etaeta*Tl_21+t_etaeta*Tl_22);
442     return P22;
    }
444 std::vector<double> Domain::P12(unsigned int i , unsigned int j){
    //x-t coordinate
446     //compute the jacobian at the point
    double s_xi = (x(i+1,j)-x(i-1,j))/(2.0*del_xi);
448     double s_eta = (x(i,j+1)-x(i,j-1))/(2.0*del_eta);
    double t_xi = (y(i+1,j)-y(i-1,j))/(2.0*del_xi);
450     double t_eta = (y(i,j+1)-y(i,j-1))/(2.0*del_eta);
    double det = s_xi*t_eta-t_xi*s_eta;
452     double Tl_11 = t_eta/det; double Tl_12 = -t_xi/det;
    double Tl_21 = -s_eta/det; double Tl_22 = s_xi/det;
454     double s_xixi = (x(i+1,j)-2.0*x(i,j)+x(i-1,j))/(del_xi*del_xi);
    double s_etaeta = (x(i,j+1)-2.0*x(i,j)+x(i,j-1))/(del_eta*del_eta);
456     double s_xieta = (x(i+1,j+1)+x(i-1,j-1)-x(i-1,j+1)-x(i+1,j-1))/(4.0*del_xi*del_eta);
    double t_xieta = (y(i+1,j+1)+y(i-1,j-1)-y(i-1,j+1)-y(i+1,j-1))/(4.0*del_xi*del_eta);
458     double t_xixi = (y(i+1,j)-2.0*y(i,j)+y(i-1,j))/(del_xi*del_xi);
    double t_etaeta = (y(i,j+1)-2.0*y(i,j)+y(i,j-1))/(del_eta*del_eta);
460     std::vector<double> P12(2);
    P12[0] = -(s_xieta*Tl_11+t_xieta*Tl_12);
462     P12[1] = -(s_xieta*Tl_21+t_xieta*Tl_22);
    return P12;
464 }

```

2.4. matrix.h

```

1 #ifndef MATRIX_H
2 #define MATRIX_H
3 class Matrix{
4 public:
5     unsigned int nx,ny;
6     std::vector<std::vector<double>> Elements;
7     Matrix(){
8     }
9
10    Matrix(unsigned int nx1,unsigned int ny1){
11        nx = nx1 ; ny = ny1;
12        //number of rows
13        Elements.resize(nx);
14        //fill each rows
15        for (unsigned int i = 0 ; i < nx ; ++i)
16            Elements[i].resize(ny,99.0);
17    }
18
19    void Clear(){
20        for (unsigned int i = 0 ; i < nx ; ++i)
21            Elements[i].clear();
22    }
23
24    double& operator()(unsigned int ix , unsigned int iy){
25        return Elements[ix][iy];
26    }
27 };
28 #endif

```

2.5. sor_solver.cpp

```

1 #ifndef SOR_SOLVER
2 #define SOR_SOLVER
3
4 #include <vector>
5
6 #include "domain.h"
7 #include "matrix.h"
8
9 double Mesh_Residual(Matrix x, Matrix y ,
10 Matrix x_old, Matrix y_old,
11 unsigned int xdim, unsigned int ydim){
12     double resid = 0 ;
13     for (int j = 0 ; j < ydim ; ++j){
14         for (int i = 0 ; i < xdim ; ++i){
15             double x_resd = (x(i,j)-x_old(i,j));
16             double y_resd = (y(i,j)-y_old(i,j));
17             resid += (x_resd*x_resd+y_resd*y_resd);
18         }
19     }
20     return std::sqrt(resid);
21 }
22
23 bool SORSOLVER( Domain& physical , Domain & parm, unsigned int xdim1,unsigned int ydim1){
24     bool grid_dist = true;
25     bool run_ellip = true;
26
27     double tolerance = 1.0e-4;
28     unsigned max_iter = 100;
29     double w = 1.90;
30     double residual = 10.0;
31     unsigned int iter = 0;
32
33     unsigned int xdim =xdim1; unsigned int ydim = ydim1;
34     Matrix x_old(xdim,ydim), y_old(xdim,ydim);
35
36     double del_xi = 1.0/double(xdim-1.0);
37     double del_eta = 1.0/double(ydim-1.0);
38

```

```

40     std::ofstream fout("gauss.dat",std::ios::out);
41     if (!(fout.is_open())) std::cerr << "ERROR:UNABLE TO OPEN THE FILE \"gauss.dat\"\\n";
42     if (run_ellip){
43         while (iter < max_iter & residual > tolerance){
44             iter++;
45             x_old = physical.MeshX() ; y_old = physical.MeshY();
46             for (unsigned int j = 1 ; j < ydim-1 ; ++j){
47                 for (unsigned int i = 1 ; i < xdim-1 ; ++i){
48
49                     double g22 = physical.G22(i, j);
50                     double g11 = physical.G11(i, j);
51                     double x_xi = physical.X_xi(i, j);
52                     double x_eta = physical.X_eta(i, j);
53                     double y_xi = physical.Y_xi(i, j);
54                     double y_eta = physical.Y_eta(i, j);
55                     double x_xieta = physical.X_xieta(i, j);
56                     double y_xieta = physical.Y_xieta(i, j);
57                     double g12 = x_xi*x_eta+y_xi*y_eta;
58                     double g = std::pow(x_xi*y_eta-y_xi*x_eta, 2);
59
60                     std::vector<double> P11 , P22, P12;
61
62                     if (grid_dist){
63                         P11 = parm.P11(i, j);
64                         P22 = parm.P22(i, j);
65                         P12 = parm.P12(i, j);
66                     }else{
67                         P11.push_back(0);P11.push_back(0);
68                         P22.push_back(0);P22.push_back(0);
69                         P12.push_back(0);P12.push_back(0);
70                     }
71
72                     double tmpx, tmpy;
73
74                     tmpx = (g22*P11[0]-2.0*g12*P12[0]+g11*P22[0])*x_xi+
75                             (g22*P11[1]-2.0*g12*P12[1]+g11*P22[1])*x_eta;
76
77                     tmpy = (g22*P11[0]-2.0*g12*P12[0]+g11*P22[0])*y_xi+
78                             (g22*P11[1]-2.0*g12*P12[1]+g11*P22[1])*y_eta;
79
80                     double lhsx = 2.0*(g22/(del_xi*del_xi)+g11/(del_eta*del_eta));
81                     double rhsx = g22*(physical.XCOORD(i+1,j)+physical.XCOORD(i-1,j))/
82                             (del_xi*del_xi)+g11*(physical.XCOORD(i, j+1)+physical.XCOORD(i, j-1))/
83                             (del_eta*del_eta)-2.0*g12*x_xieta+tmpx;
84
85                     physical.XCOORD(i, j)=physical.XCOORD(i, j)+w*(rhsx/lhsx-physical.XCOORD(i, j))
86
87                     double lhsy = lhsx;
88                     double rhsy = g22*(physical.YCOORD(i+1,j)+physical.YCOORD(i-1,j))/
89                             (del_xi*del_xi)+g11*(physical.YCOORD(i, j+1)+physical.YCOORD(i, j-1))/
90                             (del_eta*del_eta)-2.0*g12*y_xieta+ tmpy;
91
92                     physical.YCOORD(i, j)=physical.YCOORD(i, j)+w*(rhsy/lhsy-physical.YCOORD(i, j))
93                 }
94             }
95
96             double mesh_resid = Mesh_Residual(physical.MeshX(), physical.MeshY(), x_old,
97                 y_old, xdim, ydim)/((xdim-2)*(ydim-2));
98             std::cout << "Iteration_====" << iter << ", Residual_=" <<
99                 << mesh_resid << std::endl;
100
101             fout << iter << "====" << mesh_resid << std::endl;
102             residual = mesh_resid;
103         }
104     }
105     return true;
106 }
107 #endif

```

2.6. functions.h

```

1 #ifndef _FUNCTIONS_
2 #define _FUNCTIONS_

4 #include <vector>
5 #include <cmath>
6 #include <iomanip>
7 #include <iostream>
8
9 double XYcircle(double x, double y, unsigned int x_or_y){
10     double r = 1.0;
11     double theta = 0.0;
12     double Pi = 4.0*atan(1.0);
13     if (0==y){
14         theta = Pi/2.0*x;
15         if (x_or_y == 1)
16             return r*cos(theta);
17         else
18             return r*sin(theta);
19     }
20     if (1==x){
21         theta = Pi/2+Pi/2*y;
22         if (x_or_y == 1)
23             return r*cos(theta);
24         else
25             return r*sin(theta);
26     }
27     if (1==y){
28         theta=Pi+Pi/2*(1.0-x);
29         if (x_or_y == 1)
30             return r*cos(theta);
31         else
32             return r*sin(theta);
33     }
34     if (0==x){
35         theta = 3.0*Pi/2.0+Pi/2.0*(1.0-y);
36         if (x_or_y == 1)
37             return r*cos(theta);
38         else
39             return r*sin(theta);
40     }
41 }
42 #endif

```

2.7. makefile

```

1 # Generic make file for LaTeX: requires GNU make
2 #
3 # This makefile provides four targets: dvi, ps, pdf and clean.
4 # The default is "pdf".
5 # To make a dvi file, type "make_dvi"
6 # To make a ps file, type "make_ps".
7 # To make a pdf file, type "make_pdf" or simply "make".
8 # To remove all files generated by make, type "make_clean".
9 #
10 #
11 #
12 TEXFILE = Main_MS.tex
13
14
15
16 .PHONY: dvi ps pdf clean
17
18 pdf: $(TEXFILE:.tex=.pdf)
19 ps: $(TEXFILE:.tex=.ps)
20 dvi: $(TEXFILE:.tex=.dvi)
21
22 %.dvi: %.tex
23     ( \
24         \latex $< \

```



```

26 while \grep -q "Rerun\to\get\cross-references\right."
    do \
28     \latex $<; \
    done \
30 )
32 %.ps: %.dvi
    \dvips -q -t a4 $<
34 %.pdf: %.ps
36 \ps2pdf -dPDFSETTINGS=/prepress $<
38 clean:
    @\rm -f \
40     $(TEXFILE:.tex=.aux) \
42     $(TEXFILE:.tex=.log) \
44     $(TEXFILE:.tex=.out) \
    $(TEXFILE:.tex=.dvi) \
    $(TEXFILE:.tex=.ps)

```

3. Numerical Examples.

3.1. Example 1. In this example, we cluster the grids along the centre lines of a circular physical domain. Figure 4 shows the grid in the parameter space for concentrating grids at the centre lines of the physical space. Grid density in the physical space is determined by grid density in the parameter space. Figure 5 shows the converged grid in the physical space. For generating this grid lines 26 and 27 in Subsection 2.1 are used.

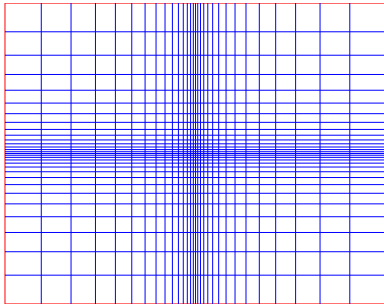


Fig. 4. A grid in the parameter space

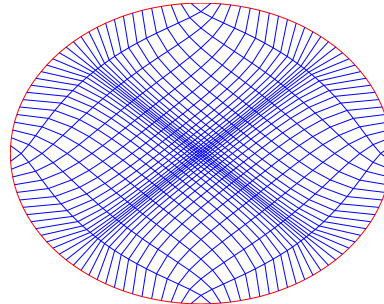


Fig. 5. Adapted grid by elliptic system

3.2. Example 2. See Figure 6 for grids in the parameter space and Figure 7 for the converged grids in the physical space. For generating the grids lines 29 and 30 of subsection 2.1 are used.

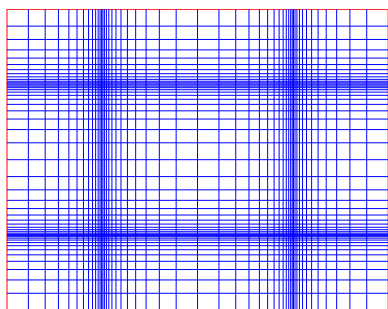


Fig. 6. A grid in the parameter space

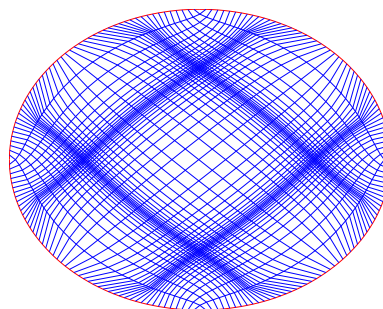


Fig. 7. Adapted grid by elliptic system

3.3. Example 3. In this example, we are interested in concentrating grids at the boundary of the physical space. Figure 8 shows the grid in the parameter space for concentrating grids at the boundary of the physical domain. The converged grids in the physical space is shown in Figure 9. For generating the grids, lines **32** and **33** of subsection 2.1 are used.

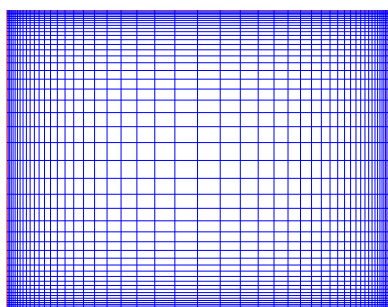


Fig. 8. A grid in the parameter space

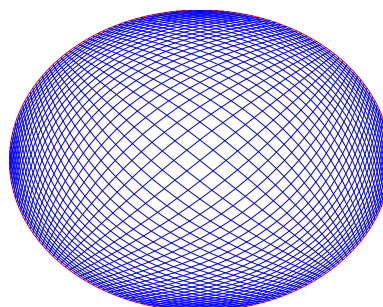


Fig. 9. Adapted grid by elliptic system

4. Conclusions. An elliptic system for generating adaptive quadrilateral meshes in curved domains has been presented. A C++ implementation of the presented technique is also given. Three examples are reported for demonstrating the effectiveness of the technique and the implementation. Since the quadrilateral meshes are very extensively used for numerical simulations, and

grid adaptation is required for capturing many important phenomenon such as the boundary layers, this software package is a very useful tool.

The software package can also be downloaded from the author's web-site.

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