Chapter 1

Example problem: The spatially-adaptive solution of the azimuthally Fourier-decomposed 3D Helmholtz equation

In this document we discuss the spatially-adaptive finite-element-based solution of the 3D Helmholtz equation in cylindrical polar coordinates, using a Fourier-decomposition of the solution in the azimuthal direction.

The driver code is very similar to the one discussed in <u>another tutorial</u> – the main purpose of the current tutorial is to demonstrate the use of spatial adaptivity on unstructured meshes.

1.1 A specific example

We will solve the azimuthally Fourier-decomposed Helmholtz equation

$$\nabla^2 u_N(r,z) + \left(k^2 - \frac{N^2}{r^2}\right) u_N(r,z) = 0, \tag{1}$$

where N is the azimuthal wavenumber, in the finite domain $1<\sqrt{r^2+z^2}<3$. We impose the Sommerfeld radiation condition at the outer boundary of the computational domain at $\sqrt{r^2+z^2}=3$, using a Dirichlet-to- \leftarrow Neumann mapping, and apply flux boundary condition on the surface of the unit-sphere (where $\sqrt{r^2+z^2}=1$) such that the exact solution is given by

$$u_N(r,z) = u_N^{[exact]}(r,z) = \sum_{l=N}^{N_{\mathrm{terms}}} h_l^{(1)}(k\sqrt{r^2 + z^2}) P_l^N\left(\frac{z}{\sqrt{r^2 + z^2}}\right).$$

This solution corresponds to the superposition of several outgoing waves that emerge from the unit sphere. The two plots below show a comparison between the exact and computed solutions for $N_{\rm terms}=6$, a Fourier wavenumber of N=1, and a (squared) Helmholtz wavenumber of $k^2=10$.

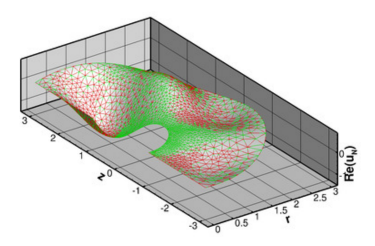


Figure 1.1 Plot of the exact (green) and computed (red) real parts of the solution of the Fourier-decomposed Helmholtz equation for N=1 and a wavenumber of $k^2 = 10$.

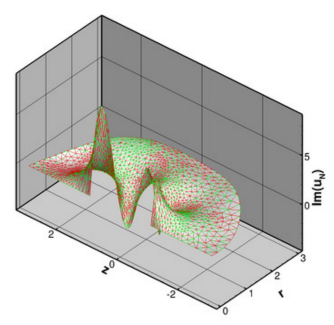


Figure 1.2 Plot of the exact (green) and computed (red) imaginary parts of the solution of the Fourier-decomposed Helmholtz equation for N=1 and a wavenumber of $k^2 = 10$.

1.2 The numerical solution

The driver code for this problem is very similar to the one discussed in $\mbox{another tutorial}$. Running \mbox{sdiff} on the driver codes

and

```
demo_drivers/fourier_decomposed_helmholtz/sphere_scattering/unstructured_←
                           sphere_scattering.cc
```

shows the main differences required to discretise the computational domain with an adaptive, unstructured mesh:

- The provision of the functions actions_before/after_adapt() to detach/re-attach the Face← Elements that are used to enforce the Neumann boundary conditions before and after every spatial adaptation, and to pass the physical parameters to the newly created bulk elements.
- The generation of an unstructured mesh whose curvilinear boundaries are represented by GeomObjects - this ensures that the domain boundaries become increasingly well resolved under mesh refinement.

That's all!

Code listing 1.3

Here's a listing of the complete driver code:

```
//LIC// This file forms part of oomph-lib, the object-oriented,
//LIC// multi-physics finite-element library, available
//LIC// at http://www.oomph-lib.org.
//LIC//
//LIC//
          Version 1.0; svn revision $LastChangedRevision$
//IJTC//
//LIC// $LastChangedDate$
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//LIC// Foundation, Inc., 51 Franklin Street, Fifth Floor, Boston, MA
//LIC// 02110-1301 USA.
//LIC//
//LIC// The authors may be contacted at oomph-lib@maths.man.ac.uk.
//LIC//
//IJTC//==
//Driver for Fourier-decomposed Helmholtz problem
#include <complex>
#include <cmath>
//Generic routines
#include "generic.h"
// The Helmholtz equations
#include "fourier_decomposed_helmholtz.h"
// The mesh
#include "meshes/triangle mesh.h"
// Get the Bessel functions
#include "oomph_crbond_bessel.h"
using namespace oomph;
using namespace std;
//---- start_of_namespace_planar_wave-----
/// Namespace to test representation of planar wave in spherical
/// polars
//-----
namespace PlanarWave
/// Number of terms in series
unsigned N_terms=100;
```

```
/// Wave number double K=3.0*MathematicalConstants::Pi;
/// Imaginary unit
std::complex<double> I(0.0,1.0);
/// Exact solution as a Vector of size 2, containing real and imag parts void get\_exact\_u (const Vector<double>& x, Vector<double>& u)
 // Switch to spherical coordinates
 double R=sqrt(x[0]*x[0]+x[1]*x[1]);
 double theta;
 theta=atan2(x[0],x[1]);
 // Argument for Bessel/Hankel functions
 // Need half-order Bessel functions
 double bessel_offset=0.5;
 // Evaluate Bessel/Hankel functions
 Vector<double> jv(N_terms);
Vector<double> yv(N_terms);
Vector<double> djv(N_terms);
 Vector<double> dyv(N_terms);
 double order_max_in=double(N_terms-1)+bessel_offset;
 double order_max_out=0;
 // This function returns vectors containing
 // J_k(x), Y_k(x) and their derivatives
 // up to k=order_max, with k increasing in
 // integer increments starting with smallest
 // positive value. So, e.g. for order_max=3.5
// jv[0] contains J_{1/2}(x),
// jv[1] contains J_{3/2}(x),
 // jv[2] contains J_{5/2}(x),
 // jv[3] contains J_{7/2}(x).
 CRBond_Bessel::bessjyv(order_max_in,
                            kr,
                            order max out,
                            &jv[0],&yv[0],
                            &djv[0],&dyv[0]);
 // Assemble exact solution (actually no need to add terms
 // below i=N_fourier as Legendre polynomial would be zero anyway)
 complex<double> u_ex(0.0,0.0);
 for(unsigned i=0;i<N_terms;i++)</pre>
   //Associated_legendre_functions
   double p=Legendre_functions_helper::plgndr2(i,0,cos(theta));
   // Set exact solution
   u_ex+=(2.0*i+1.0)*pow(I,i)*
    sqrt (MathematicalConstants::Pi/(2.0*kr))*jv[i]*p;
 // Get the real & imaginary part of the result
 u[0]=u ex.real();
 u[1]=u_ex.imag();
}//end of get_exact_u
/// Plot
void plot()
 unsigned nr=20;
 unsigned nz=100;
 unsigned nt=40;
ofstream some_file("planar_wave.dat");
for (unsigned i_t=0;i_t<nt;i_t++)</pre>
   double t=2.0*MathematicalConstants::Pi*double(i_t)/double(nt-1);
   some_file « "ZONE I="« nz « ", J="« nr « std::endl;
   Vector<double> x(2);
   Vector<double> u(2);
   for (unsigned i=0;i<nr;i++)</pre>
      x[0]=0.001+double(i)/double(nr-1);
      for (unsigned j=0;j<nz;j++)</pre>
        x[1]=double(j)/double(nz-1);
        get_exact_u(x,u);
complex<double> uu=complex<double>(u[0],u[1])*exp(-I*t);
        some_file « x[0] « " " « x[1] « "
```

```
« uu.real() « " " « uu.imag() « "\n";
      }
    }
  }
}
/// Namespace for the Fourier decomposed Helmholtz problem parameters
namespace ProblemParameters
 /// \short Square of the wavenumber
double K_squared=10.0;
 /// Fourier wave number
int N fourier=3;
 /// Number of terms in computation of DtN boundary condition
unsigned Nterms_for_DtN=6;
 /// Number of terms in the exact solution
unsigned N terms=6:
 /// Coefficients in the exact solution
Vector<double> Coeff(N_terms, 1.0);
 /// Imaginary unit
 std::complex<double> I(0.0,1.0);
 /// Exact solution as a Vector of size 2, containing real and imag parts
 void get_exact_u(const Vector<double>& x, Vector<double>& u)
  \ensuremath{//} Switch to spherical coordinates
  double R=sqrt(x[0]*x[0]+x[1]*x[1]);
  double theta;
  theta=atan2(x[0],x[1]);
 // Argument for Bessel/Hankel functions
double kr = sqrt(K_squared) *R;
  // Need half-order Bessel functions
  double bessel_offset=0.5;
  // Evaluate Bessel/Hankel functions
  Vector<double> jv(N_terms);
  Vector<double> yv(N_terms);
Vector<double> djv(N_terms);
  Vector<double> dyv(N_terms);
  double order_max_in=double(N_terms-1)+bessel_offset;
  double order_max_out=0;
  // This function returns vectors containing
  // J_k(x), Y_k(x) and their derivatives // up to k=order_max, with k increasing in
  // integer increments starting with smallest
  // positive value. So, e.g. for order_max=3.5
  // jv[0] contains J_{1/2}(x),
// jv[1] contains J_{3/2}(x),
// jv[2] contains J_{5/2}(x),
  // jv[3] contains J_{7/2}(x).
  CRBond_Bessel::bessjyv(order_max_in,
                          order_max_out,
                          &jv[0],&yv[0],
                          &djv[0],&dyv[0]);
  // Assemble exact solution (actually no need to add terms
  // below i=N_fourier as Legendre polynomial would be zero anyway)
  complex<double> u_ex(0.0,0.0);
  for(unsigned i=N_fourier;i<N_terms;i++)</pre>
    //Associated_legendre_functions
   double p=Legendre_functions_helper::plgndr2(i, N_fourier,
                                                 cos(theta));
    // Set exact solution
     u_ex + = Coeff[i] * sqrt (MathematicalConstants:: Pi/(2.0*kr)) * (jv[i] + I*yv[i]) * p; \\
  // Get the real & imaginary part of the result
  u[0]=u ex.real();
```

```
u[1]=u_ex.imag();
 }//end of get exact u
 /// \short Get -du/dr (spherical r) for exact solution. Equal to prescribed
 /// flux on inner boundary.
 void exact_minus_dudr(const Vector<double>& x, std::complex<double>& flux)
  // Initialise flux
  flux=std::complex<double>(0.0,0.0);
  // Switch to spherical coordinates
  double R=sqrt(x[0]*x[0]+x[1]*x[1]);
  double theta:
  theta=atan2(x[0],x[1]);
  // Argument for Bessel/Hankel functions
  double kr=sqrt(K_squared)*R;
  // Helmholtz wavenumber
  double k=sqrt(K_squared);
  // Need half-order Bessel functions
  double bessel_offset=0.5;
  // Evaluate Bessel/Hankel functions
  Vector<double> jv(N_terms);
  Vector<double> yv(N_terms);
Vector<double> djv(N_terms);
  Vector<double> dyv(N_terms);
  double order_max_in=double(N_terms-1)+bessel_offset;
  double order_max_out=0;
  // This function returns vectors containing
  // J_k(x), Y_k(x) and their derivatives // up to k=order_max, with k increasing in
  // integer increments starting with smallest
  // positive value. So, e.g. for order_max=3.5
    jv[0] contains J_{1/2}(x),
  // jv[1] contains J_{3/2}(x),
  // jv[2] contains J_{5/2}(x), // jv[3] contains J_{7/2}(x).
  CRBond_Bessel::bessjyv(order_max_in,
                         kr,
                         order_max_out,
                         &jv[0],&yv[0],
                         &djv[0],&dyv[0]);
  // Assemble exact solution (actually no need to add terms
  // below i=N_fourier as Legendre polynomial would be zero anyway)
  complex<double> u_ex(0.0,0.0);
  for(unsigned i=N_fourier;i<N_terms;i++)</pre>
    //Associated_legendre_functions
   double p=Legendre_functions_helper::plgndr2(i, N_fourier,
                                                 cos(theta));
    // Set flux of exact solution
    \texttt{flux-=Coeff[i]} * \texttt{sqrt} \texttt{ (MathematicalConstants::Pi/(2.0*kr))} * \texttt{p*}
     ( k*(djv[i]+I*dyv[i]) - (0.5*(jv[i]+I*yv[i])/R) );
}// end of exact_normal_derivative
} // end of namespace
//===== start_of_problem_class===============================
/// Problem class
                   ______
template<class ELEMENT>
class FourierDecomposedHelmholtzProblem : public Problem
public:
 /// Constructor
 FourierDecomposedHelmholtzProblem();
 /// Destructor (empty)
 ~FourierDecomposedHelmholtzProblem(){}
 /// Update the problem specs before solve (empty)
 void actions_before_newton_solve(){}
```

```
/// Update the problem after solve (empty)
void actions_after_newton_solve(){}
 /// \short Doc the solution. DocInfo object stores flags/labels for where the
 /// output gets written to
 void doc_solution(DocInfo& doc_info);
 /// Recompute gamma integral before checking Newton residuals
 void actions_before_newton_convergence_check()
   if (!CommandLineArgs::command_line_flag_has_been_set("--square_domain"))
     Helmholtz_outer_boundary_mesh_pt->setup_gamma();
   }
 /// Actions before adapt: Wipe the mesh of prescribed flux elements
 void actions_before_adapt();
 /// Actions after adapt: Rebuild the mesh of prescribed flux elements
void actions_after_adapt();
 /// Check gamma computation
void check_gamma(DocInfo& doc_info);
private:
 /// \ Short Create BC elements on outer boundary
void create_outer_bc_elements();
 /// Create flux elements on inner boundary
void create_flux_elements_on_inner_boundary();
 /// \short Delete boundary face elements and wipe the surface mesh
 void delete_face_elements( Mesh* const & boundary_mesh_pt)
   \ensuremath{//} Loop over the surface elements
   unsigned n_element = boundary_mesh_pt->nelement();
   for(unsigned e=0;e<n_element;e++)</pre>
     // Kill surface element
    delete boundary_mesh_pt->element_pt(e);
   // Wipe the mesh
  boundary_mesh_pt->flush_element_and_node_storage();
#ifdef ADAPTIVE
 /// Pointer to the "bulk" mesh
RefineableTriangleMesh<ELEMENT>* Bulk_mesh_pt;
#else
 /// Pointer to the "bulk" mesh
TriangleMesh<ELEMENT>* Bulk_mesh_pt;
#endif
 /// \short Pointer to mesh containing the DtN boundary
FourierDecomposedHelmholtzDtNMesh<ELEMENT>* Helmholtz_outer_boundary_mesh_pt;
 /// on the inner boundary
Mesh* Helmholtz inner boundary mesh pt;
 /// Trace file
 ofstream Trace_file;
}; // end of problem class
              -----start_of_actions_before_adapt-----start_of_actions_before_adapt
/// Actions before adapt: Wipe the mesh of face elements
template<class ELEMENT>
void FourierDecomposedHelmholtzProblem<ELEMENT>::actions_before_adapt()
 \ensuremath{//} Kill the flux elements and wipe the boundary meshs
if (!CommandLineArgs::command_line_flag_has_been_set("--square_domain"))
  delete_face_elements(Helmholtz_outer_boundary_mesh_pt);
 delete_face_elements(Helmholtz_inner_boundary_mesh_pt);
 // Rebuild the Problem's global mesh from its various sub-meshes
 rebuild_global_mesh();
}// end of actions_before_adapt
             =======start of actions after adapt======
/// Actions after adapt: Rebuild the face element meshes
```

```
template<class ELEMENT>
void FourierDecomposedHelmholtzProblem<ELEMENT>::actions_after_adapt()
// Complete the build of all elements so they are fully functional
 // Loop over the Helmholtz bulk elements to set up element-specific
 // things that cannot be handled by constructor: Pass pointer to
 // wave number squared
unsigned n_element = Bulk_mesh_pt->nelement();
 for (unsigned e=0;e<n_element;e++)</pre>
  {
   // Upcast from GeneralisedElement to Helmholtz bulk element
   ELEMENT *el_pt = dynamic_cast<ELEMENT*>(Bulk_mesh_pt->element_pt(e));
   //Set the k_squared pointer
   el_pt->k_squared_pt() = &ProblemParameters::K_squared;
   // Set pointer to Fourier wave number
   el_pt->fourier_wavenumber_pt() =&ProblemParameters::N_fourier;
 // Create prescribed-flux elements and BC elements
 \ensuremath{//} from all elements that are adjacent to the boundaries and add them to
 // Helmholtz_boundary_meshes
create_flux_elements_on_inner_boundary();
if (!CommandLineArgs::command_line_flag_has_been_set("--square_domain"))
   create_outer_bc_elements();
 // Rebuild the Problem's global mesh from its various sub-meshes
rebuild_global_mesh();
}// end of actions_after_adapt
/// Constructor for Fourier-decomposed Helmholtz problem
template<class ELEMENT>
FourierDecomposedHelmholtzProblem<ELEMENT>::
FourierDecomposedHelmholtzProblem()
// Open trace file
Trace_file.open("RESLT/trace.dat");
 // Create circles representing inner and outer boundary
double x_c=0.0;
double y_c=0.0;
double r_min=1.0;
double r_max=3.0;
Circle* inner_circle_pt=new Circle(x_c,y_c,r_min);
Circle* outer_circle_pt=new Circle(x_c,y_c,r_max);
 // Edges/boundary segments making up outer boundary
Vector<TriangleMeshCurveSection*> outer_boundary_line_pt(4);
 // Number of segments used for representing the curvilinear boundaries
unsigned n_segments = 20;
 // All poly boundaries are defined by two vertices
Vector<Vector<double> > boundary_vertices(2);
 // Bottom straight boundary on symmetry line
boundary_vertices[0].resize(2);
boundary_vertices[0][0]=0.0;
boundary_vertices[0][1]=-r_min;
boundary_vertices[1].resize(2);
boundary_vertices[1][0]=0.0;
boundary_vertices[1][1]=-r_max;
 unsigned boundary_id=0;
outer_boundary_line_pt[0]=
 new TriangleMeshPolyLine(boundary_vertices,boundary_id);
 if (CommandLineArgs::command_line_flag_has_been_set("--square_domain"))
   // Square outer boundary:
   Vector<Vector<double> > boundary_vertices(4);
   boundary_vertices[0].resize(2);
   boundary_vertices[0][0]=0.0;
   boundary_vertices[0][1]=-r_max;
boundary_vertices[1].resize(2);
boundary_vertices[1][0]=r_max;
   boundary_vertices[1][1]=-r_max;
   boundary_vertices[2].resize(2);
   boundary_vertices[2][0]=r_max;
boundary_vertices[2][1]=r_max;
   boundary_vertices[3].resize(2);
boundary_vertices[3][0]=0.0;
   boundary_vertices[3][1]=r_max;
```

```
boundary_id=1;
  outer_boundary_line_pt[1]=
   new TriangleMeshPolyLine(boundary_vertices, boundary_id);
else
   // Outer circular boundary:
  // The intrinsic coordinates for the beginning and end of the curve
  double s_start = -0.5*MathematicalConstants::Pi;
  double s_end
                 = 0.5*MathematicalConstants::Pi;
  boundary_id = 1;
  outer_boundary_line_pt[1]=
   new TriangleMeshCurviLine(outer_circle_pt,
                               s start.
                               s end,
                               n_segments,
                               boundary_id);
// Top straight boundary on symmetry line
boundary_vertices[0][0]=0.0;
boundary_vertices[0][1]=r_max;
boundary_vertices[1][0]=0.0;
boundary_vertices[1][1]=r_min;
boundary_id=2;
outer boundary line pt[2]=
 new TriangleMeshPolyLine (boundary_vertices, boundary_id);
// Inner circular boundary:
// The intrinsic coordinates for the beginning and end of the curve
double s_start = 0.5*MathematicalConstants::Pi;
double s_end = -0.5*MathematicalConstants::Pi;
boundary_id = 3;
outer_boundary_line_pt[3]=
 new TriangleMeshCurviLine(inner_circle_pt,
                             s start,
                             s_end,
                             n_segments,
                             boundary_id);
// Create closed curve that defines outer boundary
TriangleMeshClosedCurve *outer_boundary_pt =
 new TriangleMeshClosedCurve(outer_boundary_line_pt);
// Use the TriangleMeshParameters object for helping on the manage of the // TriangleMesh parameters. The only parameter that needs to take is the
 // outer boundary.
TriangleMeshParameters triangle_mesh_parameters(outer_boundary_pt);
// Specify maximum element area
double element_area = 0.1;
triangle_mesh_parameters.element_area() = element_area;
#ifdef ADAPTIVE
 // Build "bulk" mesh
Bulk_mesh_pt=new RefineableTriangleMesh<ELEMENT>(triangle_mesh_parameters);
// Create/set error estimator
Bulk_mesh_pt->spatial_error_estimator_pt() = new Z2ErrorEstimator;
// Choose error tolerances to force some uniform refinement
Bulk_mesh_pt->min_permitted_error()=0.00004;
Bulk_mesh_pt->max_permitted_error()=0.0001;
#else
// Pass the TriangleMeshParameters object to the TriangleMesh one
Bulk_mesh_pt= new TriangleMesh<ELEMENT>(triangle_mesh_parameters);
#endif
// Check what we've built so far...
Bulk_mesh_pt->output("mesh.dat");
Bulk_mesh_pt->output_boundaries("boundaries.dat");
if (!CommandLineArgs::command_line_flag_has_been_set("--square_domain"))
   // Create mesh for DtN elements on outer boundary
  Helmholtz_outer_boundary_mesh_pt=
   new FourierDecomposedHelmholtzDtNMesh<ELEMENT>(
    r_max, ProblemParameters::Nterms_for_DtN);
   // Populate it with elements
   create_outer_bc_elements();
```

```
// Create flux elements on inner boundary
Helmholtz inner boundary mesh pt=new Mesh;
create_flux_elements_on_inner_boundary();
 // Add the several sub meshes to the problem
add_sub_mesh(Bulk_mesh_pt);
add_sub_mesh(Helmholtz_inner_boundary_mesh_pt);
 if (!CommandLineArgs::command_line_flag_has_been_set("--square_domain"))
  add_sub_mesh(Helmholtz_outer_boundary_mesh_pt);
 // Build the Problem's global mesh from its various sub-meshes
build_global_mesh();
 // Complete the build of all elements so they are fully functional
 unsigned n_element = Bulk_mesh_pt->nelement();
 for (unsigned i=0;i<n_element;i++)</pre>
   // Upcast from GeneralsedElement to the present element
  ELEMENT *el_pt = dynamic_cast<ELEMENT*>(Bulk_mesh_pt->element_pt(i));
   //Set the k_squared pointer
  el_pt->k_squared_pt()=&ProblemParameters::K_squared;
   // Set pointer to Fourier wave number
  el_pt->fourier_wavenumber_pt() =&ProblemParameters::N_fourier;
// Setup equation numbering scheme
cout «"Number of equations: " « assign_eqn_numbers() « std::endl;
} // end of constructor
/// Check gamma computation: f \gamma = -du/dn \f$
//==========
template<class ELEMENT>
void FourierDecomposedHelmholtzProblem<ELEMENT>::check_gamma(DocInfo& doc_info)
 // Compute gamma stuff
Helmholtz_outer_boundary_mesh_pt->setup_gamma();
ofstream some_file;
char filename[100];
 sprintf(filename, "%s/gamma_test%i.dat", doc_info.directory().c_str(),
        doc_info.number());
 some_file.open(filename);
 //first loop over elements e
 unsigned nel=Helmholtz_outer_boundary_mesh_pt->nelement();
 for (unsigned e=0;e<nel;e++)</pre>
   // Get a pointer to element
  FourierDecomposedHelmholtzDtNBoundaryElement<ELEMENT>* el_pt=
   dynamic_cast<FourierDecomposedHelmholtzDtNBoundaryElement<ELEMENT>*>
   (Helmholtz_outer_boundary_mesh_pt->element_pt(e));
   //Set the value of n_intpt
  const unsigned n_intpt =el_pt->integral_pt()->nweight();
   // Get gamma at all gauss points in element
   Vector<std::complex<double> > gamma(
   Helmholtz_outer_boundary_mesh_pt->gamma_at_gauss_point(el_pt));
   //Loop over the integration points
   for(unsigned ipt=0;ipt<n_intpt;ipt++)</pre>
   {
     //Allocate and initialise coordiante
     Vector<double> x(el_pt->dim()+1,0.0);
     //Set the Vector to hold local coordinates
    unsigned n=el_pt->dim();
Vector<double> s(n,0.0);
     for (unsigned i=0;i<n;i++)</pre>
      s[i]=el_pt->integral_pt()->knot(ipt,i);
     //Get the coordinates of the integration point
     el_pt->interpolated_x(s,x);
     complex<double> flux;
     ProblemParameters::exact_minus_dudr(x,flux);
     some\_file \ll atan2(x[0],x[1]) \ll ""
              « gamma[ipt].real() « " "
              « gamma[ipt].imag() « " "
              « flux.real() «
               « flux.imag() « " "
```

```
« std::endl;
    }// end of loop over integration points
  }// end of loop over elements
some_file.close();
}//end of output_gamma
//========start_of_doc=======
/// Doc the solution: doc_info contains labels/output directory etc.
template<class ELEMENT>
void FourierDecomposedHelmholtzProblem<ELEMENT>::doc_solution(DocInfo& doc_info)
ofstream some file:
char filename[100];
 // Number of plot points: npts x npts
unsigned npts=5;
 // Output solution
sprintf(filename, "%s/soln%i.dat", doc_info.directory().c_str(),
         doc_info.number());
 some_file.open(filename);
Bulk_mesh_pt->output(some_file,npts);
 some file.close();
 // Output exact solution
 sprintf(filename, "%s/exact_soln%i.dat", doc_info.directory().c_str(),
         doc_info.number());
 some_file.open(filename);
Bulk_mesh_pt->output_fct(some_file,npts,ProblemParameters::get_exact_u);
some file.close();
 // Doc error and return of the square of the L2 error
double error, norm;
sprintf(filename, "%s/error%i.dat", doc_info.directory().c_str(),
         doc info.number());
 some_file.open(filename);
Bulk_mesh_pt->compute_error(some_file,ProblemParameters::get_exact_u,
                              error, norm);
some_file.close();
// Doc L2 error and norm of solution
cout « "\nNorm of error : " « sqrt(error) « std::endl;
cout « "Norm of solution: " « sqrt(norm) « std::endl « std::endl;
 // Write norm of solution to trace file
Bulk_mesh_pt->compute_norm(norm);
Trace file « norm « std::endl;
 if (!CommandLineArgs::command_line_flag_has_been_set("--square_domain"))
   // Check gamma computation
   check_gamma(doc_info);
} // end of doc
//======start of create outer bc elements==
/// Create BC elements on outer boundary
template<class ELEMENT>
void FourierDecomposedHelmholtzProblem<ELEMENT>::create_outer_bc_elements()
 // Outer boundary is boundary 1:
unsigned b=1;
 // Loop over the bulk elements adjacent to boundary b?
 unsigned n_element = Bulk_mesh_pt->nboundary_element(b);
 for(unsigned e=0;e<n_element;e++)</pre>
   // Get pointer to the bulk element that is adjacent to boundary b
   ELEMENT* bulk_elem_pt = dynamic_cast<ELEMENT*>(
   Bulk_mesh_pt->boundary_element_pt(b,e));
   //Find the index of the face of element {\tt e} along boundary {\tt b}
   int face_index = Bulk_mesh_pt->face_index_at_boundary(b,e);
   // Build the corresponding DtN element
   FourierDecomposedHelmholtzDtNBoundaryElement<ELEMENT>* flux_element_pt = new
   FourierDecomposedHelmholtzDtNBoundaryElement<ELEMENT>(bulk_elem_pt,
   //Add the flux boundary element to the helmholtz outer boundary mesh
   Helmholtz_outer_boundary_mesh_pt->add_element_pt(flux_element_pt);
// Set pointer to the mesh that contains all the boundary condition
   // elements on this boundary
```

```
flux_element_pt->
    set_outer_boundary_mesh_pt(Helmholtz_outer_boundary_mesh_pt);
} // end of create_outer_bc_elements
//======start_of_create_flux_elements=========
/// Create flux elements on inner boundary
template<class ELEMENT>
void FourierDecomposedHelmholtzProblem<ELEMENT>::
create_flux_elements_on_inner_boundary()
// Apply flux bc on inner boundary (boundary 3)
unsigned b=3;
// Loop over the bulk elements adjacent to boundary b
unsigned n_element = Bulk_mesh_pt->nboundary_element(b);
 for (unsigned e=0;e<n_element;e++)</pre>
   // Get pointer to the bulk element that is adjacent to boundary b
  ELEMENT* bulk_elem_pt = dynamic_cast<ELEMENT*>(
   Bulk_mesh_pt->boundary_element_pt(b,e));
   //Find the index of the face of element e along boundary b
   int face_index = Bulk_mesh_pt->face_index_at_boundary(b,e);
   // Build the corresponding prescribed incoming-flux element
  FourierDecomposedHelmholtzFluxElement<ELEMENT>* flux_element_pt = new
   FourierDecomposedHelmholtzFluxElement<ELEMENT>(bulk_elem_pt,face_index);
   //Add the prescribed incoming-flux element to the surface mesh
  Helmholtz_inner_boundary_mesh_pt->add_element_pt(flux_element_pt);
   // Set the pointer to the prescribed flux function
  flux_element_pt->flux_fct_pt() = &ProblemParameters::exact_minus_dudr;
  } //end of loop over bulk elements adjacent to boundary b
} // end of create flux elements on inner boundary
//==== start_of_main======
/// Driver code for Fourier decomposed Helmholtz problem
int main(int argc, char **argv)
 // Store command line arguments
CommandLineArgs::setup(argc,argv);
 // Define possible command line arguments and parse the ones that
 // were actually specified
 // Square domain without DtN
CommandLineArgs::specify_command_line_flag("--square_domain");
 // Parse command line
CommandLineArgs::parse_and_assign();
 // Doc what has actually been specified on the command line
CommandLineArgs::doc_specified_flags();
 // Check if the claimed representation of a planar wave in
 // the tutorial is correct -- of course it is!
 //PlanarWave::plot();
 // Test Bessel/Hankel functions
 // Number of Bessel functions to be computed
 unsigned n=3;
  // Offset of Bessel function order (less than 1!)
 double bessel_offset=0.5;
 ofstream bessely_file("bessely.dat");
 ofstream bessely_deriv_file("dbesselY.dat");
  ofstream besselj_file("besselJ.dat");
  ofstream besselj_deriv_file("dbesselJ.dat");
  // Evaluate Bessel/Hankel functions
  Vector<double> jv(n+1);
  Vector<double> yv(n+1);
  Vector<double> djv(n+1);
  Vector<double> dyv(n+1);
 double x_min=0.5;
 double x_max=5.0;
unsigned nplot=100;
  for (unsigned i=0;i<nplot;i++)</pre>
   double x=x_{min}+(x_{max}-x_{min})*double(i)/double(nplot-1);
    double order_max_in=double(n)+bessel_offset;
   double order_max_out=0;
    // This function returns vectors containing
    // J_k(x), Y_k(x) and their derivatives
```

```
// up to k=\operatorname{order\_max}, with k increasing in
    \ensuremath{//} integer increments starting with smallest
    // positive value. So, e.g. for order_max=3.5
// jv[0] contains J_{1/2}(x),
    // jv[1] contains J_{3/2}(x),
     // jv[2] contains J_{5/2}(x),
     // jv[3] contains J_{7/2}(x).
    CRBond_Bessel::bessjyv(order_max_in,x,
                                order_max_out,
                                &jv[0],&yv[0],
                                &djv[0],&dyv[0]);
    bessely_file « x « " ";
    for (unsigned j=0; j<=n; j++)</pre>
       bessely_file « yv[j] « " ";
    bessely_file « std::endl;
    besselj_file « x « " ";
    for (unsigned j=0;j<=n;j++)</pre>
       besselj_file « jv[j] « " ";
    besselj_file « std::endl;
    bessely_deriv_file « x « " ";
    for (unsigned j=0;j<=n;j++)</pre>
       bessely_deriv_file « dyv[j] « " ";
    bessely_deriv_file « std::endl;
    besselj_deriv_file « x « " ";
    for (unsigned j=0; j<=n; j++)</pre>
       besselj_deriv_file « djv[j] « " ";
    besselj_deriv_file « std::endl;
 bessely_file.close();
besselj_file.close();
  bessely_deriv_file.close();
 besselj_deriv_file.close();
 // Test Legrendre Polynomials
  // Number of lower indices
  unsigned n=3;
  ofstream some_file("legendre3.dat");
  unsigned nplot=100;
  for (unsigned i=0;i<nplot;i++)</pre>
    double x=double(i)/double(nplot-1);
some_file « x « " ";
for (unsigned j=0;j<=n;j++)</pre>
       some_file « Legendre_functions_helper::plgndr2(n,j,x) « " ";
    some_file « std::endl;
  some_file.close();
#ifdef ADAPTIVE
 // Create the problem with 2D six-node elements from the \,
// TFourierDecomposedHelmholtzElement family.
FourierDecomposedHelmholtzProblem<ProjectableFourierDecomposedHelmholtzElement<
  TFourierDecomposedHelmholtzElement<3> > problem;
#else
// Create the problem with 2D six-node elements from the
// TFourierDecomposedHelmholtzElement family.
FourierDecomposedHelmholtzProblem<TFourierDecomposedHelmholtzElement<3> >
 problem;
 // Create label for output
DocInfo doc info;
 // Set output directory
doc_info.set_directory("RESLT");
```

```
// Solve for a few Fourier wavenumbers
for (ProblemParameters::N_fourier=0;ProblemParameters::N_fourier<4;
        ProblemParameters::N_fourier++)
{
        // Step number
        doc_info.number()=ProblemParameters::N_fourier;

#ifdef ADAPTIVE
        // Max. number of adaptations
        unsigned max_adapt=1;

        // Solve the problem with Newton's method, allowing
        // up to max_adapt mesh adaptations after every solve.
        problem.newton_solve(max_adapt);

#else
        // Solve the problem
        problem.newton_solve();
#endif

        //Output the solution
        problem.doc_solution(doc_info);
}</pre>
```

//end of main

1.4 Source files for this tutorial

• The source files for this tutorial are located in the directory:

```
demo_drivers/fourier_decomposed_helmholtz/sphere_scattering/
```

· The driver code is:

demo_drivers/fourier_decomposed_helmholtz/sphere_
 scattering/unstructured_sphere_scattering.cc

1.5 PDF file

A pdf version of this document is available.