

Chapter 1

Example problem: Solution of the 2D unsteady heat equation with restarts

Simulations of time-dependent problem can be very time consuming and it is important to be able to restart simulations, e.g. to continue a run after a system crash, etc. We shall illustrate `oomph-lib`'s dump/restart capabilities by re-visiting the 2D unsteady heat equation discussed in a [previous example](#).

The two-dimensional unsteady heat equation in a square domain.

Solve

$$\sum_{i=1}^2 \frac{\partial^2 u}{\partial x_i^2} = \frac{\partial u}{\partial t} + f(x_1, x_2, t), \quad (1)$$

in the square domain $D = \{x_i \in [0, 1]; i = 1, 2\}$, subject to the Dirichlet boundary conditions

$$u|_{\partial D} = g_0 \quad (2)$$

and initial conditions

$$u(x_1, x_2, t = 0) = h_0(x_1, x_2), \quad (3)$$

where the functions g_0 and h_0 are given.

As before, we consider the unforced case, $f = 0$, and choose boundary and initial conditions that are consistent with the exact solution

$$u_0(x_1, x_2, t) = e^{-Kt} \sin(\sqrt{K}(x_1 \cos \Phi + x_2 \sin \Phi)), \quad (4)$$

where K and Φ are constants, controlling the decay rate of the solution and its spatial orientation, respectively.

The figure below shows a plot of computed and exact solution at a control node as a function of time. The solid lines represent quantities computed during the original simulation; the dashed line shows the corresponding data from a second simulation that was restarted with the restart file generated at timestep 23 of the original simulation.

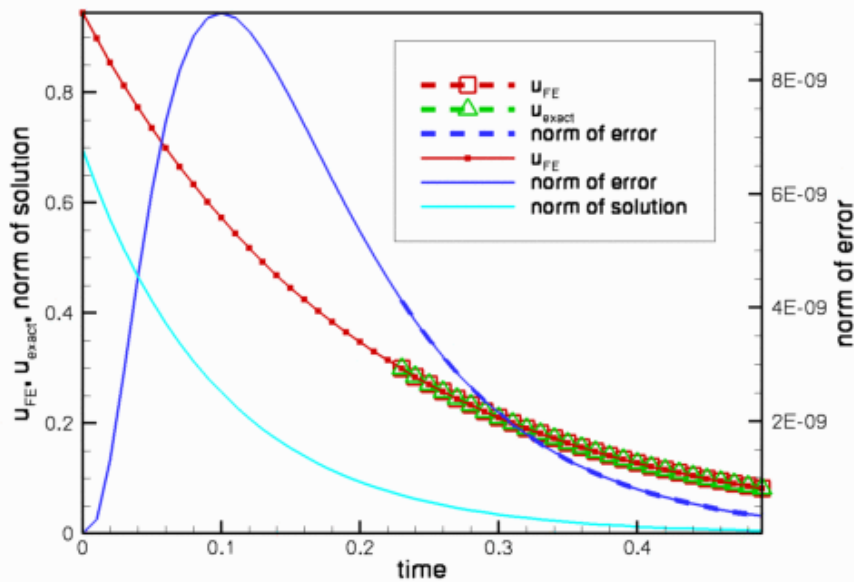


Figure 1.1 Time evolution of the computed and exact solutions at a control node, the global error norm and the norm of the solution. Solid lines: original simulation; dashed lines: restarted simulation.

Most of the driver code for this example is identical to that discussed in the [previous example](#), therefore we only discuss the modifications required to enable the dump and restart operations:

- We use optional command line arguments to specify the name of the restart file. If the code is run without any command line arguments, we start the simulation at time $t = 0$ and generate the initial conditions as in the [previous example](#)
- We add dump and restart functions to the Problem class and call the dump function when post-processing the solution in `doc_solution(...)`.
- We modify the Problem member function `set_initial_condition()` so that the initial conditions are read from the restart file if a restart file was specified on the command line.

1.1 Global parameters and functions

The namespace `ExactSolnForUnsteadyHeat` that stores the problem parameters is identical to that in the [previous example](#).

1.2 The driver code

The only change to the main function is that we record the command line arguments and store them in the namespace `CommandLineArgs`

```
=====start_of_main=====
/// Driver code for unsteady heat equation with option for
/// restart from disk: Only a single command line argument is allowed.
/// If specified it is interpreted as the name of the restart file.
=====
int main(int argc, char* argv[])
{
    // Store command line arguments
    CommandLineArgs::setup(argc, argv);
}
```

The rest of the main function is identical to that in the [previous example](#).

1.3 The problem class

The problem class contains the two additional member functions

```
// Dump problem to disk to allow for restart.  
void dump_it(ofstream& dump_file);  
  
// Read problem for restart from specified restart file.  
void restart(ifstream& restart_file);
```

1.4 The problem constructor

The problem constructor is identical to that in the [previous example](#).

1.5 The problem destructor

The problem destructor is identical to that in the [previous example](#).

1.6 Actions before timestep

This function is identical to that in the [previous example](#).

1.7 Set initial condition

We start by checking the validity of the command line arguments, accessible via the namespace `CommandLineArgs`, as only a single command line argument is allowed. If a command line argument is provided, it is interpreted as the name of the restart file. We try to open the file and, if successful, pass the input stream to the `restart(...)` function, discussed below. If no command line arguments are specified, we generate the initial conditions, essentially as in the

[previous example](#). The only difference is that in the current version of the code, we moved the specification and initialisation of the timestep from the `main` function into `set_initial_condition()`. This is because in a restarted simulation, the value of `dt` must be consistent with that used in the original simulation. If the simulation is restarted, the generic `Problem::read(...)` function, called by `restart(...)`, automatically initialises the previous timestep; otherwise we have to perform the initialisation ourselves.

```
=====start_of_set_initial_condition=====  
// Set initial condition: Assign previous and current values  
// from exact solution or from restart file.  
=====  
template<class ELEMENT>  
void UnsteadyHeatProblem<ELEMENT>::set_initial_condition()  
{  
  
    // Pointer to restart file  
    ifstream* restart_file_pt=0;  
  
    // Restart?  
    //-----  
    // Restart file specified via command line [all programs have at least  
    // a single command line argument: their name. Ignore this here.]  
    if (CommandLineArgs::Argc==1)  
    {  
        cout << "No restart -- setting IC from exact solution" << std::endl;  
    }  
    else if (CommandLineArgs::Argc==2)  
    {  
        // Open restart file  
        restart_file_pt= new ifstream(CommandLineArgs::Argv[1],ios_base::in);  
        if (restart_file_pt!=0)  
        {  
            cout << "Have opened " << CommandLineArgs::Argv[1] <<  
            " for restart. " << std::endl;  
        }  
        else  
        {  
            std::ostringstream error_stream;  
            error_stream  
            << "ERROR while trying to open " << CommandLineArgs::Argv[1] <<  
            " for restart." << std::endl;  
  
            throw OomphLibError(  
                error_stream.str(),  
                OOMPH_CURRENT_FUNCTION,  
                OOMPH_EXCEPTION_LOCATION);  
        }  
    }  
}
```

```

        }

    // More than one command line argument?
else
{
    std::ostringstream error_stream;
    error_stream << "Can only specify one input file\n"
                  << "You specified the following command line arguments:\n";
    //Fix this
    CommandLineArgs::output();

    throw OomphLibError(
        error_stream.str(),
        OOMPH_CURRENT_FUNCTION,
        OOMPH_EXCEPTION_LOCATION);
}

// Read restart data:
//-----
if (restart_file_pt!=0)
{
    // Read the problem data from the restart file
    restart(*restart_file_pt);

}

// Assign initial condition from exact solution
//-----
else
{
    // Choose timestep
    double dt=0.01;

    // Initialise timestep -- also sets the weights for all timesteppers
    // in the problem.
    initialise_dt(dt);

    // Backup time in global Time object
    double backed_up_time=time_pt()->time();

    // Past history fo needs to be established for t=time0-deltat, ...
    // Then provide current values (at t=time0) which will also form
    // the initial guess for the first solve at t=time0+deltat

    // Vector of exact solution value
    Vector<double> soln(1);
    Vector<double> x(2);

    //Find number of nodes in mesh
    unsigned num_nod = mesh_pt()->nnode();

    // Set continuous times at previous timesteps
    int nprev_steps=time stepper_pt()->nprev_values();
    Vector<double> prev_time(nprev_steps+1);
    for (int itime=nprev_steps;itime>=0;itime--)
    {
        prev_time[itime]=time_pt()->time(unsigned(itime));
    }

    // Loop over current & previous timesteps
    for (int itime=nprev_steps;itime>=0;itime--)
    {
        // Continuous time
        double time=prev_time[itime];
        cout << "setting IC at time =" << time << std::endl;

        // Loop over the nodes to set initial guess everywhere
        for (unsigned n=0;n<num_nod;n++)
        {
            // Get nodal coordinates
            x[0]=mesh_pt()->node_pt(n)->x(0);
            x[1]=mesh_pt()->node_pt(n)->x(1);

            // Get intial solution
            ExactSolnForUnsteadyHeat::get_exact_u(time,x,soln);

            // Assign solution
            mesh_pt()->node_pt(n)->set_value(itime,0,soln[0]);

            // Loop over coordinate directions: Previous position = present position
            for (unsigned i=0;i<2;i++)
            {
                mesh_pt()->node_pt(n)->x(itime,i)=x[i];
            }
        }
    }
}

```

```

}

// Reset backed up time for global timestepper
time_pt() ->time() =backed_up_time;

}

} // end of set_initial_condition

```

1.8 Post-processing

The Problem member function `doc_solution(...)` is identical to that in the [previous example](#), apart from the addition of a call to the new `dump_it(...)` function, discussed below.

```

// Write restart file
snprintf(filename, sizeof(filename), "%s/restart%i.dat", doc_info.directory().c_str(),
         doc_info.number());
some_file.open(filename);
dump_it(some_file);
some_file.close();

```

1.9 Dumping the solution

The `Problem::dump(...)` function writes the generic `Problem` data in ASCII format to the specified output file. The content of the file can therefore be inspected and, if necessary, manipulated before a restart. However, the specific content of the file is generally of little interest – it is written in a format that can be read by the corresponding function `Problem::read(...)`.

Briefly, the dump file contains:

- A flag that indicates if the data was produced by a time-dependent simulation.
- The current value of the "continuous" time, i.e. the value returned by `Problem::time_pt() ->time()`;
- The number of previous timesteps, `dt`, stored in the `Problem`'s `Time` object, and their values.
- The values and history values for all `Data` objects in the `Problem`, as well as the present and previous coordinates of all `Nodes`.

The "raw data" is augmented by brief comments that facilitate the identification of individual entries.

In the present problem, the generic `Problem::dump(...)` function is sufficient to record the current state of the simulation, therefore no additional information needs to be added to the dump file. The section [Comments and Exercises](#) below contains an exercise that illustrates how to customise the dump function to record additional parameters; the [demo code with spatial adaptivity](#) provides another example of a customised dump/restart function.

```

=====start_of_dump_it=====
/// Dump the solution to disk to allow for restart
=====template<class ELEMENT>
void UnsteadyHeatProblem<ELEMENT>::dump_it(ofstream& dump_file)
{
    // Call generic dump()
    Problem::dump(dump_file);

} // end of dump_it

```

1.10 Reading a solution from disk

Since the restart file was written by the generic `Problem::dump(...)` function, it can be read back with the generic `Problem::read(...)` function. If any additional data had been recorded in the restart file, additional `read` statements could be added here; see [Comments and Exercises](#).

```

=====start_of_restart=====
/// Read solution from disk for restart
=====template<class ELEMENT>
void UnsteadyHeatProblem<ELEMENT>::restart(ifstream& restart_file)
{
    // Read the generic problem data from restart file
    Problem::read(restart_file);

} // end of restart

```

1.11 Comments and Exercises

The `Problem::dump(...)` and `Problem::read(...)` functions write/read the generic data that is common to all `oomph-lib` problems. Occasionally, it is necessary to record additional data to re-create the system's state when the simulations is restarted. We will explore this in more detail in [another example](#). Here we provide a brief exercise that illustrates the general idea and addresses a shortcoming of the driver code: Currently the program computes the same number of timesteps, regardless of whether or not the simulation was restarted. If a simulation is restarted, the computation therefore continues past $t = t_{max}$.

1.11.1 Exercises

- Change the `for` loop over the fixed number of timesteps in the `main` function to a `while` loop that checks if the continuous time, accessible via `Problem::time_pt() -> time()`, has reached or exceeded t_{max} . This works because the `Problem::dump(...)` and `Problem::read(...)` functions dump and restore the value of the continuous time.
- Following this trivial change, the restarted simulation is terminated at the appropriate point but the numbering of the output files begins at 0, making it difficult to merge the results from the original and the restarted simulations. Modify the functions `dump_it(...)` and `restart(...)` so that they write/read the current label for the output files to/from the restart file.

Hints: (i) You can write to/read from the restart file before calling `Problem::dump(...)` and `Problem::read(...)`; just make sure you do it in the same order in both functions. (ii) The easiest way to make the label, currently stored in the `DocInfo` object in the `main` function, accessible to all member functions in the `Problem` is to make the `DocInfo` object a private data member of the `Problem` class.

1.12 Source files for this tutorial

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

```
demo_drivers/unsteady_heat/two_d_unsteady_heat/
```

- The driver code is:

```
demo_drivers/unsteady_heat/two_d_unsteady_heat/two_d_unsteady_heat_←
restarted.cc
```

1.13 PDF file

A [pdf version](#) of this document is available. \