### Exercises for Tutorial05

# Adaptivity

### Exercise 1 Playing with the Parameters

The adaptive finite element algorithm is controlled by several parameters which can be set in the ini-file. Besides the polynomial degree these are:

- steps: how many times the solve—adapt cycle is executed.
- uniformlevel: how many times uniform mesh refinement is used before adaptive refinement starts.
- tol: tolerance where the adaptive algorithm is stopped.
- fraction: Remember that the global error is estimated from element-wise contributions:

$$\gamma^2(u_h) = \sum_{T \in \mathcal{T}_h} \gamma_T^2(u_h).$$

The fraction parameter controls which elements are marked for refinement. More precisely, the largest threshold  $\gamma^*$  is determined such that

$$\sum_{\{T \in \mathcal{T}_h : \gamma_T(u_h) > \gamma^*\}} \gamma_T^2(u_h) \ge \operatorname{fraction} \cdot \gamma^2(u_h).$$

If fraction is set to zero no elements need to be refined, i.e.  $\gamma^* = \infty$ . When fraction is set to one then all elements need to be refined, i.e.  $\gamma^* = 0$ . A smaller fraction parameter results in a more optimized mesh but needs more steps and thus might be more expensive. If fraction is chosen too large then almost all elements are refined which might also be expensive, so there exists an (unknown) optimal value for fraction.

Carry out the following experiments:

1. Choose polynomial degree 1 and fix a tolerance, e.g. tol=0.01. Set steps to a large number so that the algorithm is not stopped before the tolerance is reached. Set uniformlevel to zero. Now vary the fraction parameter and measure the execution time with

#### \$ time ./exercise05

2. Repeat the previous experiment with polynomial degrees two and three. Compare the optimal meshes generated for the different polynomial degrees.

3. The idea of the heuristic refinement algorithm refining only the elements with the largest contribution to the error is to equilibrate the error. To check how well this is achieved use the calculator filter in ParaView (use cell data) to plot  $\log(\gamma_T(u_h))$ . Note that the cell data exported by the (unchanged) code is  $\gamma_T^2(u_h)!$  Now check the minimum and maximum error contributions. What happens directly at the singularity, i.e. the point (0,0)?

## Exercise 2 Compute the True $L_2$ -error

Disclaimer: In this exercise we compute and evalute the  $L_2$ -norm of the error. The residual based error estimator implemented in the code, however, estimates the  $H^1$ -seminorm (which is equivalent to the  $H^1$ -norm here) and thus also optimizes the mesh with respect to that norm. This is done to make the exercise as simple as possible. It would be more appropriate to carry out the experiments either with the true  $H^1$ -norm or to change the estimator to the  $L_2$ -norm.

To do this exercise we need some more background on PDELab grid functions. The following code known from several tutorials by now constructs a PDELab grid function object g:

```
Problem <RF> problem(eta);
auto glambda =
[&](const auto& e, const auto& x){return problem.g(e,x);};
auto g = Dune::PDELab::makeGridFunctionFromCallable(gv,glambda);
```

Also the following code segment used in many tutorials constructs a PDELab grid function object zdgf from a grid function space and a coefficient vector:

```
typedef Dune::PDELab::DiscreteGridFunction < GFS, Z > ZDGF;
ZDGF zdgf(gfs,z);
```

PDELab grid functions can be evaluated in local coordinates given by an element and a local coordinate within the corresponding reference element. The result is stored in an additional argument passed by reference. Here is a code segment doing the job:

```
Dune::FieldVector < RF, 1 > truesolution(0.0);
g.evaluate(e,x,truesolution);
```

Here we assume that  $\mathbf{e}$  is an element and  $\mathbf{x}$  is a local coordinate. The result is stored in a Dune::FieldVector with one component of type RF. This allows also to return vector-valued results.

Now here are your tasks:

1. Extend the code in the file driver.hh in tutorial05/exercise/task to provide a grid function that provides the error  $u - u_h$ . Note that the method g in the parameter class already returns the true solution (for  $\eta = 0$ ). You can solve this task by writing a lambda function subtracting the values returned by g.evaluate and zdgf.evaluate.

- 2. Compute the  $L_2$ -norm of the error, i.e.  $\|u-u_h\|_0 = \sqrt{\int_{\Omega} (u(x)-u_h(x))^2 dx}$ . This can be done by creating a grid function with the squared error using Dune::PDELab::SqrGridFunctionAdapter from <dune/pdelab/function/sqr.hh> and the function Dune::PDELab::integrateGridFunction contained in the header file <dune/pdelab/common/functionutilities.hh>. Output the result to the console by writing in a single line a tag like L2ERROR, the number of degrees of freedom using gfs.globalSize() and the  $L_2$ -norm. This lets you grep the results later for post-processing.
- 3. Investigate the  $L_2$ -error for different polynomial degrees. Run the code using different polynomial degrees and also uniform and adaptive (use your optimal fraction) refinement. Use gnuplot to produce graphs such as those in Figure 1. An example gnuplot file plot.gp is given.
  - The graph shows that for uniform refinement polynomial degree 2 is better than polynomial degree 1 only by a constant factor. For adaptive refinement a better convergence rate can be achieved. The plot also shows that for adaptive refinement the optimal convergence rate can be recovered. For  $P_1$  finite elements and fully regular solution in  $H^2(\Omega)$  the a-priori estimate for uniform refinement yields  $||u-u_h||_0 \leq Ch^2$ . Expressing h in terms of the number of degrees of freedom N yields  $h \sim N^{-1/d}$ , so  $||u-u_h||_0 \leq Ch^2 \leq C'N^{-1}$  for d=2. The curve  $N^{-1}$  is shown for comparison in the plot. Clearly, the result for  $P_1$  is parallel to that line.
- 4. Optional: Also write the true error to the VTK output file.

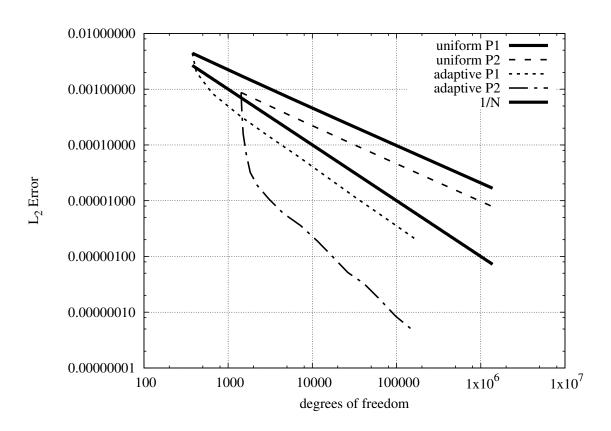


Figure 1:  $L_2$ -error versus number of degrees of freedom for different polynomial degrees.