

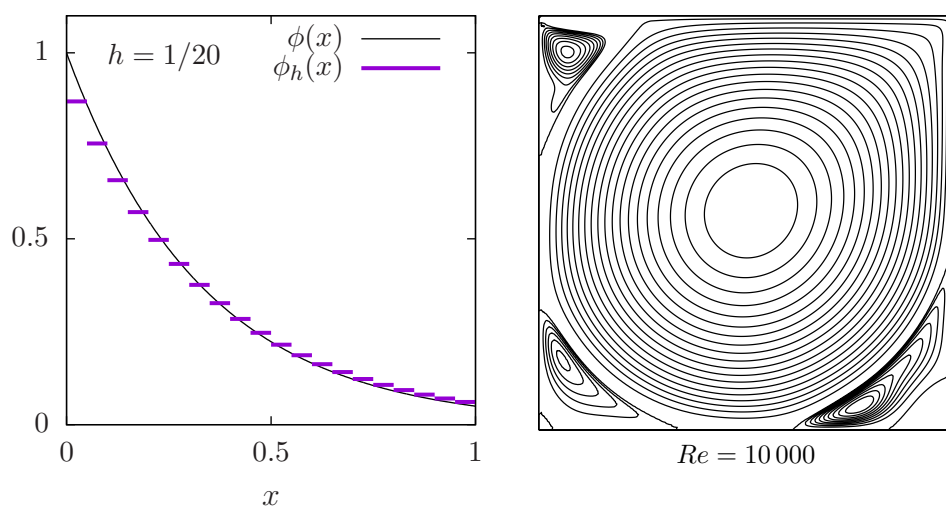
Efficient C++ finite element computing with Rheolef

volume 2:

discontinuous Galerkin methods

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version 6.7 update 24 March 2016



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Notations

Rheolef	mathematics	description
d	$d \in \{1, 2, 3\}$	dimension of the physical space
dot(u,v)	$\mathbf{u} \cdot \mathbf{v} = \sum_{i=0}^{d-1} u_i v_i$	vector scalar product
ddot(sigma,tau)	$\sigma : \tau = \sum_{i,j=0}^{d-1} \sigma_{i,j} \tau_{i,j}$	tensor scalar product
tr(sigma)	$\text{tr}(\sigma) = \sum_{i=0}^{d-1} \sigma_{i,i}$	trace of a tensor
trans(sigma)	σ^T	tensor transposition
sqr(phi) norm2(phi)	ϕ^2	square of a scalar
norm2(u)	$ \mathbf{u} ^2 = \sum_{i=0}^{d-1} u_i^2$	square of the vector norm
norm2(sigma)	$ \sigma ^2 = \sum_{i,j=0}^{d-1} \sigma_{i,j}^2$	square of the tensor norm
abs(phi) norm(phi)	$ \phi $	absolute value of a scalar
norm(u)	$ \mathbf{u} = \left(\sum_{i=0}^{d-1} u_i^2 \right)^{1/2}$	vector norm
norm(sigma)	$ \sigma = \left(\sum_{i,j=0}^{d-1} \sigma_{i,j}^2 \right)^{1/2}$	tensor norm
grad(phi)	$\nabla \phi = \left(\frac{\partial \phi}{\partial x_i} \right)_{0 \leq i < d}$	gradient of a scalar field in $\Omega \subset \mathbb{R}^d$
grad(u)	$\nabla \mathbf{u} = \left(\frac{\partial u_i}{\partial x_j} \right)_{0 \leq i,j < d}$	gradient of a vector field
div(u)	$\text{div}(\mathbf{u}) = \text{tr}(\nabla \mathbf{u}) = \sum_{i=0}^{d-1} \frac{\partial u_i}{\partial x_i}$	divergence of a vector field
D(u)	$D(\mathbf{u}) = (\nabla \mathbf{u} + \nabla \mathbf{u}^T) / 2$	symmetric part of the gradient of a vector field
curl(u)	$\mathbf{curl}(\mathbf{u}) = \nabla \wedge \mathbf{u}$	curl of a vector field, when $d = 3$
curl(phi)	$\mathbf{curl}(\phi) = \left(\frac{\partial \phi}{\partial x_1}, -\frac{\partial \phi}{\partial x_0} \right)$	curl of a scalar field, when $d = 2$
curl(u)	$\mathbf{curl}(\mathbf{u}) = \frac{\partial u_1}{\partial x_0} - \frac{\partial u_0}{\partial x_1}$	curl of a vector field, when $d = 2$
grad_s(phi)	$\nabla_s \phi = P \nabla \phi$ where $P = I - \mathbf{n} \otimes \mathbf{n}$	tangential gradient of a scalar

Rheolef	mathematics	description
<code>grad_s(u)</code>	$\nabla_s \mathbf{u} = \nabla \mathbf{u} P$	tangential gradient of a vector
<code>Ds(u)</code>	$D_s(\mathbf{u}) = P D(\mathbf{u}) P$	symmetrized tangential gradient
<code>div_s(u)</code>	$\operatorname{div}_s(\mathbf{u}) = \operatorname{tr}(D_s(\mathbf{u}))$	tangential divergence
<code>normal()</code>	\mathbf{n}	unit outward normal on $\Gamma = \partial\Omega$ or on an oriented surface Ω or on an internal oriented side S
<code>jump(phi)</code>	$[\![\phi]\!] = \phi _{K_0} - \phi _{K_1}$	jump accross inter-element side $S = \partial K_0 \cap K_1$
<code>average(phi)</code>	$\{\!\!\{\phi\}\!\!\} = (\phi _{K_0} + \phi _{K_1})/2$	average accross S
<code>inner(phi)</code>	$\phi _{K_0}$	inner trace on S
<code>outer(phi)</code>	$\phi _{K_1}$	outer trace on S
<code>h_local()</code>	$h_K = \operatorname{meas}(K)^{1/d}$	length scale on an element K
<code>penalty()</code>	$\varpi_s = \max\left(\frac{\operatorname{meas}(\partial K_0)}{\operatorname{meas}(K_0)}, \frac{\operatorname{meas}(\partial K_1)}{\operatorname{meas}(K_1)}\right)$	penalty coefficient on S
<code>grad_h(phi)</code>	$(\nabla_h \phi) _K = \nabla(\phi _K), \forall K \in \mathcal{T}_h$	broken gradient
<code>div_h(u)</code>	$(\operatorname{div}_h \mathbf{u}) _K = \operatorname{div}(\mathbf{u} _K), \forall K \in \mathcal{T}_h$	broken divergence of a vector field
<code>Dh(u)</code>	$(D_h(\mathbf{u})) _K = D(\mathbf{u} _K), \forall K \in \mathcal{T}_h$	broken symmetric part of the gradient of a vector field
<code>sin(phi)</code> <code>cos(phi)</code> <code>tan(phi)</code> <code>acos(phi)</code> <code>asin(phi)</code> <code>atan(phi)</code> <code>cosh(phi)</code> <code>sinh(phi)</code> <code>tanh(phi)</code> <code>exp(phi)</code> <code>log(phi)</code> <code>log10(phi)</code> <code>floor(phi)</code> <code>ceil(phi)</code> <code>min(phi,psi)</code>	$\sin(\phi)$ $\cos(\phi)$ $\tan(\phi)$ $\cos^{-1}(\phi)$ $\sin^{-1}(\phi)$ $\tan^{-1}(\phi)$ $\cosh(\phi)$ $\sinh(\phi)$ $\tanh(\phi)$ $\exp(\phi)$ $\log(\phi)$ $\log 10(\phi)$ $\lfloor \phi \rfloor$ $\lceil \phi \rceil$ $\min(\phi, \psi)$	standard mathematical functions extended to scalar fields largest integral not greater than ϕ smallest integral not less than ϕ

Rheolef	mathematics	description
<code>max(phi,psi)</code> <code>pow(phi,psi)</code> <code>atan2(phi,psi)</code> <code>fmod(phi,psi)</code>	$\max(\phi, \psi)$ ϕ^ψ $\tan^{-1}(\psi/\phi)$ $\phi - \lfloor \phi/\psi + 1/2 \rfloor \psi$	floating point remainder
<code>compose(f,phi)</code>	$f \circ \phi = f(\phi)$	applies an unary function f
<code>compose(f,phi1,...,phin)</code>	$f(\phi_1, \dots, \phi_n)$	applies a n -ary function f , $n \geq 1$
<code>compose(phi,X)</code>	$\phi \circ X, \quad X(x) = x + \mathbf{d}(x)$	composition with a characteristic

Part I

Getting started with simple problems

Chapter 1

Scalar first-order problems

The aim of this chapter is to introduce to discontinuous Galerkin methods within the **Rheolef** environment. For some recent presentations of discontinuous Galerkin methods, see [11] for theoretical aspects and [16] for algorithmic and implementation. The discontinuous Galerkin methods are in active development in **Rheolef**, and new features will appear soon.

1.1 The transport equation

The steady scalar transport problem writes:

(P): *find ϕ , defined in Ω , such that*

$$\begin{aligned} \mathbf{u} \cdot \nabla \phi + \sigma \phi &= f \text{ in } \Omega \\ \phi &= \phi_\Gamma \text{ on } \partial\Omega_- \end{aligned}$$

where \mathbf{u} , $\sigma > 0$, f and ϕ_Γ being known. Notice that this is the steady version of the unsteady diffusion-convection problem previously introduced in section 6.2, page 86 and when the diffusion coefficient ν vanishes. Here, the $\partial\Omega_-$ notation is the *upstream* boundary part, defined by

$$\partial\Omega_- = \{x \in \partial\Omega; \mathbf{u}(x) \cdot \mathbf{n}(x) < 0\}$$

Let us suppose that $\mathbf{u} \in W^{1,\infty}(\Omega)^d$ and introduce the space:

$$X = \{\varphi \in L^2(\Omega); (\mathbf{u} \cdot \nabla) \varphi \in L^2(\Omega)^d\}$$

and, for all $\phi, \varphi \in X$

$$\begin{aligned} a(\phi, \varphi) &= \int_{\Omega} (\mathbf{u} \cdot \nabla \phi \varphi + \sigma \phi \varphi) \, dx + \int_{\partial\Omega} \max(0, -\mathbf{u} \cdot \mathbf{n}) \phi \varphi \, ds \\ l(\varphi) &= \int_{\Omega} f \varphi \, dx + \int_{\partial\Omega} \max(0, -\mathbf{u} \cdot \mathbf{n}) \phi_\Gamma \varphi \, ds \end{aligned}$$

Then, the variational formulation writes:

(FV): *find $\phi \in X$ such that*

$$a(\phi, \varphi) = l(\varphi), \quad \forall \varphi \in X$$

Notice that the term $\max(0, -\mathbf{u} \cdot \mathbf{n}) = (|\mathbf{u} \cdot \mathbf{n}| - \mathbf{u} \cdot \mathbf{n})/2$ is positive and vanishes everywhere except on $\partial\Omega_-$. Thus, the boundary condition $\phi = \phi_\Gamma$ is weakly imposed on $\partial\Omega_-$ via the integrals on the boundary. The *discontinuous* finite element space is defined by:

$$X_h = \{\varphi_h \in L^2(\Omega); \varphi_h|_K \in P_k, \quad \forall K \in \mathcal{T}_h\}$$

where $k \geq 0$ is the polynomial degree. Notice that $X_h \not\subset X$ and that the $\nabla \phi_h$ term has no more sense for discontinuous functions $\phi_h \in X_h$. Following [11, p. 14], we introduce the *broken gradient* ∇_h as a convenient notation:

$$(\nabla_h \phi_h)|_K = \nabla(\phi_h|_K), \quad \forall K \in \mathcal{T}_h$$

Thus

$$\int_{\Omega} \mathbf{u} \cdot \nabla_h \phi_h \varphi_h \, dx = \sum_{K \in \mathcal{T}_h} \int_K \mathbf{u} \cdot \nabla \phi_h \varphi_h \, dx, \quad \forall \phi_h, \varphi_h \in X_h$$

This leads to a discrete version a_h of the bilinear form a , defined for all $\phi_h, \varphi_h \in X_h$ by (see e.g. [11, p. 57], eqn. (2.34)):

$$\begin{aligned} a_h(\phi_h, \varphi_h) &= \int_{\Omega} (\mathbf{u} \cdot \nabla_h \phi_h \varphi_h + \sigma \phi_h \varphi_h) \, dx + \int_{\partial\Omega} \max(0, -\mathbf{u} \cdot \mathbf{n}) \phi_h \varphi_h \, ds \\ &\quad + \sum_{S \in \mathcal{S}_h^{(i)}} \int_S \left(-\mathbf{u} \cdot \mathbf{n} \llbracket \phi_h \rrbracket \llbracket \varphi_h \rrbracket + \frac{\alpha}{2} |\mathbf{u} \cdot \mathbf{n}| \llbracket \phi_h \rrbracket \llbracket \varphi_h \rrbracket \right) \, ds \end{aligned}$$

The last term involves a sum over $\mathcal{S}_h^{(i)}$, the set of *internal sides* of the mesh \mathcal{T}_h . Each internal side

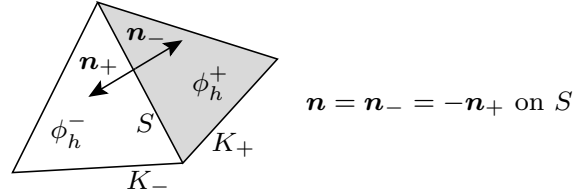


Figure 1.1: Discontinuous Galerkin method: an internal side, its two neighbor elements and their opposite normals.

$S \in \mathcal{S}_h^{(i)}$ has two possible orientations: one is chosen definitively. In practice, this orientation is defined in the ‘.geo’ file containing the mesh, where all sides are listed, together with their orientation. Let \mathbf{n} the normal to the oriented side S : as S is an internal side, there exists two elements K_- and K_+ such that $S = \partial K_- \cap \partial K_+$ and \mathbf{n} is the outward unit normal of K_- on $\partial K_- \cap S$ and the inward unit normal of K_+ on $\partial K_+ \cap S$, as shown on Fig. 1.1. For all $\phi_h \in X_h$, recall that ϕ_h is in general discontinuous across the internal side S . We define on S the *inner value* $\phi_h^- = \phi_h|_{K_-}$ of ϕ_h as the restriction $\phi_h|_{K_-}$ of ϕ_h in K_- along $\partial K_- \cap S$. Conversely, we define the *outer value* $\phi_h^+ = \phi_h|_{K_+}$. We also denote on S the *jump* $\llbracket \phi_h \rrbracket = \phi_h^- - \phi_h^+$ and the *average* $\{\!\{ \phi_h \}\!\} = (\phi_h^- + \phi_h^+)/2$. The last term in the definition of a_h is ponderated by a coefficient $\alpha \geq 0$. Choosing $\alpha = 0$ correspond to the so-called *centered flux* approximation, while $\alpha = 1$ is the *upwinding flux* approximation. The case $\alpha = 1$ and $k = 0$ (piecewise constant approximation) leads to the popular upwinding finite volume scheme. Finally, the discrete variational formulation writes:

$(FV)_h$: find $\phi_h \in X_h$ such that

$$a_h(\phi_h, \varphi_h) = l(\varphi_h), \quad \forall \varphi_h \in X_h$$

The following code implement this problem in the **Rheolef** environment.

Example file 1.1: transport_dg.cc

```

1 #include "rheolef.h"
2 using namespace rheolef;
3 using namespace std;
4 int main(int argc, char**argv) {
5     environment rheolef (argc, argv);
6     geo omega (argv[1]);
7     space Xh (omega, argv[2]);
8     Float alpha = (argc > 3) ? atof(argv[3]) : 1;
9     Float sigma = (argc > 4) ? atof(argv[4]) : 3;
10    point u (1,0,0);
11    trial phi (Xh); test psi (Xh);
12    form ah = integrate (dot(u,grad_h(phi))*psi + sigma*phi*psi)
13              + integrate ("boundary", max(0, -dot(u,normal()))*phi*psi)
14              + integrate ("internal_sides",
15                          - dot(u,normal())*jump(phi)*average(psi)
16                          + 0.5*alpha*abs(dot(u,normal()))*jump(phi)*jump(psi));
17    field lh = integrate ("boundary", max(0, -dot(u,normal()))*psi);
18    solver sah (ah.uu());
19    field phi_h(Xh);
20    phi_h.set_u() = sah.solve(lh.u());
21    dout << catchmark("sigma") << sigma << endl
22          << catchmark("phi") << phi_h;
23 }

```

Comments

The data are $\phi_\gamma = 1$ and $\mathbf{u} = (1, 0, 0)$, and then the exact solution is known: $\phi(x) = \exp(-\sigma x_0)$. The numerical tests are running with $\sigma = 3$ by default. The one-dimensional case writes:

```

make transport_dg
mkgeo_grid -e 10 > line.geo
./transport_dg line P0 | field -
./transport_dg line P1d | field -
./transport_dg line P2d | field -

```

Observe the jumps accross elements: these jumps decreases with mesh refinement or when polynomial approximation increases. The two-dimensional case writes:

```

mkgeo_grid -t 10 > square.geo
./transport_dg square P0 | field -elevation -
./transport_dg square P1d | field -elevation -
./transport_dg square P2d | field -elevation -

```

The elevation view shows details on inter-element jumps. Finally, the three-dimensional case writes:

```

mkgeo_grid -T 5 > cube.geo
./transport_dg cube P0 | field -
./transport_dg cube P1d | field -
./transport_dg cube P2d | field -

```

Fig. 1.2 plots the solution when $d = 1$ and $k = 0$: observe that the boundary condition $\phi = 1$ at $x_0 = 0$ is only weakly satisfied. It means that the approximation $\phi_h(0)$ tends to 1 when h tends to zero. Fig. 1.3 plots the error $\phi - \phi_h$ in L^2 and L^∞ norms: these errors behave as $\mathcal{O}(h^{k+1})$ for all $k \geq 0$, which is optimal. A theoretical $\mathcal{O}(h^{k+1/2})$ error bound was shown in [17]. The present numerical results confirm that these theoretical error bounds can be improved for some families of meshes, as pointed out by Richter [19], that showed a $\mathcal{O}(h^{k+1})$ optimal bound for the transport problem. This result was recently extended by Cockburn *et al.* [6], while Peterson [18] showed that the estimate $\mathcal{O}(h^{k+1/2})$ is sharp for general families of quasi-uniform meshes.

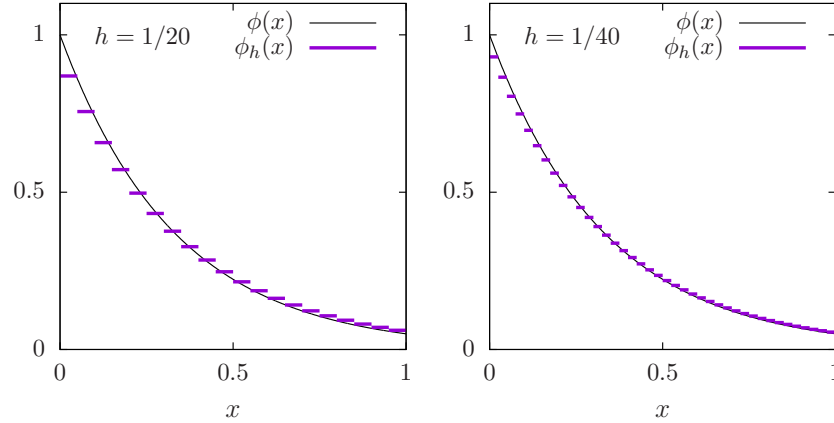


Figure 1.2: The discontinuous Galerkin method for the transport problem when $k = 0$ and $d = 1$.

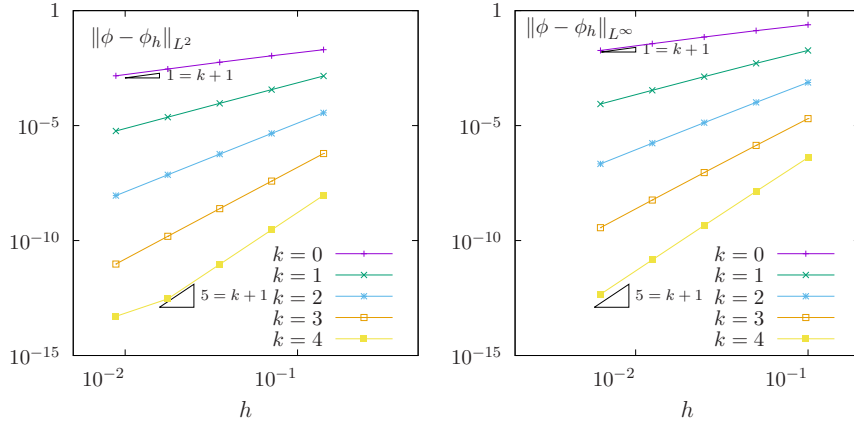


Figure 1.3: The discontinuous Galerkin method for the transport problem: convergence when $d = 2$.

1.2 Nonlinear scalar hyperbolic problems

The aim of this paragraph is to study the discontinuous Galerkin discretization of scalar nonlinear hyperbolic equations. This section presents the general framework and discretization tools while the next section illustrates the method for the Burgers equation.

1.2.1 Problem setting

A time-dependent nonlinear hyperbolic problem writes in general form [11, p. 99]:

(P): find u , defined in $]0, T[\times \Omega$, such that

$$\frac{\partial u}{\partial t} + \operatorname{div} \mathbf{f}(u) = 0 \quad \text{in }]0, T[\times \Omega \quad (1.1a)$$

$$u(t=0) = u_0 \quad \text{in } \Omega \quad (1.1b)$$

$$\mathbf{f}(u) \cdot \mathbf{n} = \Phi(\mathbf{n}; u, g) \quad \text{on }]0, T[\times \partial\Omega \quad (1.1c)$$

where $T > 0$, $\Omega \subset \mathbb{R}^d$, $d = 1, 2, 3$ and the initial condition \mathbf{u}_0 being known. As usual, \mathbf{n} denotes the outward unit normal on the boundary $\partial\Omega$. The function $\mathbf{f} : \mathbb{R} \rightarrow \mathbb{R}^d$ is also known and supposed

to be continuously differentiable. The initial data u_0 , defined in Ω , and the boundary one, g , defined on $\partial\Omega$ are given. The function Φ , called the Godunov flux associated to \mathbf{f} , is defined, for all $\nu \in \mathbb{R}^d$ and $a, b \in \mathbb{R}$, by

$$\Phi(\nu; a, b) = \begin{cases} \min_{v \in [a, b]} \mathbf{f}(v) \cdot \nu & \text{when } a \leq b \\ \max_{v \in [b, a]} \mathbf{f}(v) \cdot \nu & \text{otherwise} \end{cases} \quad (1.1d)$$

1.2.2 Space discretization

In this section, we consider first the semi-discretization with respect to space while the problem remains continuous with respect to time. The semi-discrete problem writes in variational form [11, p. 100]:

$(P)_h$: find $u_h \in C^1([0, T], X_h)$ such that

$$\begin{aligned} \int_{\Omega} \frac{\partial u_h}{\partial t} v_h \, dx - \int_{\Omega} \mathbf{f}(u_h) \cdot \nabla_h v_h \, dx + \sum_{S \in \mathcal{S}_h^{(i)}} \Phi(\mathbf{n}; u_h^-, u_h^+) \llbracket v_h \rrbracket \, ds + \int_{\partial\Omega} \Phi(\mathbf{n}; u_h, g) v_h \, ds &= 0, \quad \forall v_h \in X_h \\ u_h(t=0) &= \pi_h(u_0) \end{aligned}$$

where π_h denotes the Lagrange interpolation operator on X_h and others notations has been introduced in the previous section.

For convenience, we introduce the discrete operator G_h , defined for all $u_h, v_h \in X_h$ by

$$\int_{\Omega} G_h(u_h) v_h \, dx = - \int_{\Omega} \mathbf{f}(u_h) \cdot \nabla_h v_h \, dx + \sum_{S \in \mathcal{S}_h^{(i)}} \int_S \Phi(\mathbf{n}; u_h^-, u_h^+) \llbracket v_h \rrbracket \, ds + \int_{\partial\Omega} \Phi(\mathbf{n}; u_h, g) v_h \, ds \quad (1.2)$$

For a given $u_h \in X_h$, we also define the linear form g_h as

$$g(v_h) = \int_{\Omega} G_h(u_h) v_h \, dx$$

As the matrix M , representing the L^2 scalar product in X_h , is block-diagonal, it can be easily inverted at the element level, and for a given $u_h \in X_h$, we have $G(u_h) = M^{-1}g_h$. Then, the problem writes equivalently as a set of coupled nonlinear ordinary differential equations.

$(P)_h$: find $u_h \in C^1([0, T], X_h)$ such that

$$\frac{\partial u_h}{\partial t} + G_h(u_h) = 0$$

1.2.3 Time discretization

Let $\Delta t > 0$ be the time step. The previous nonlinear ordinary differential equations are discretized by using a specific explicit Runge-Kutta with intermediates states [13, 14, 22]. This specific variant of the usual Runge-Kutta scheme, called *strong stability preserving*, is suitable for avoiding possible spurious oscillations of the approximate solution when the exact solution has a poor regularity. Let u_h^n denotes the approximation of u_h at time $t_n = n\Delta t$, $n \geq 0$. Then u_h^{n+1} is defined by recurrence:

$$\begin{aligned} u_h^{n,0} &= u_h^n \\ u_h^{n,i} &= \sum_{j=0}^{i-1} \alpha_{i,j} u_h^{n,j} - \Delta t \beta_{i,j} G_h(u_h^{n,j}), \quad 1 \leq i \leq p \\ u_h^{n+1} &= u_h^{n,p} \end{aligned}$$

where the coefficients satisfy $\alpha_{i,j} \geq 0$ and $\beta_{i,j} \geq 0$ for all $1 \leq i \leq p$ and $0 \leq j \leq i-1$, and $\sum_{j=0}^{i-1} \alpha_{i,j} = 1$ for all $1 \leq i \leq p$. Notice that when $p = 1$ this scheme coincides with the usual explicit Euler scheme. For a general $p \geq 1$ order scheme, there are $p-1$ intermediate states $u_h^{n,i}$, $i = 1 \dots p-1$. Computation of the coefficients $\alpha_{i,j}$ and $\beta_{i,j}$ can be founded in [13, 14, 22] and are grouped in file ‘runge_kutta_ssp.icc’ of the examples directory for convenience.

1.2.4 Slope limiters

Slope limiters are required when the solution develops discontinuities: this is a very classical feature of most solutions of nonlinear hyperbolic equations. A preliminary version of the slope limiter proposed by Cocburn et al. [5, p. 208] is implemented in **Rheolef**: this preliminary version only supports the $d = 1$ dimension and $k = 1$ polynomial degree. Recall that the $k = 0$ case do not need any limiter. More general implementation will support the $d = 2, 3$ and $k \geq 2$ cases in the future. The details of the limiter implementation is presented in this section: the impatient reader, who is interested by applications, can jump to the next section, devoted to the Burgers equation.

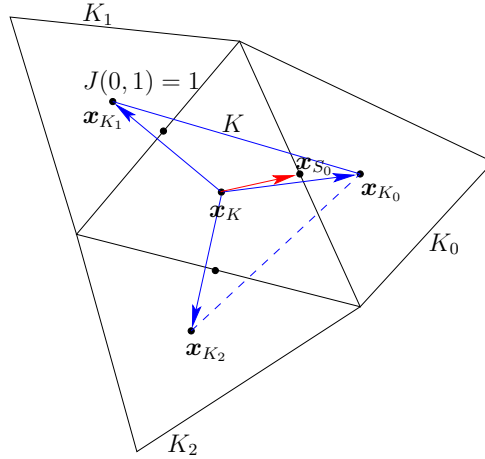


Figure 1.4: Limiter: the neighbors elements and the middle edge points.

Fig. 1.4 shows the $d+1$ neighbor elements K_i , $i = 0 \dots d$ around an element d . Let $S_i = \partial K \cap \partial K_i$, $i = 0 \dots d$ be the i -th side of K . We denote by \mathbf{x}_K , \mathbf{x}_{K_i} and \mathbf{x}_{S_i} the barycenters of these elements and sides, $i = 0 \dots d$. When $d = 2$, the barycenter \mathbf{x}_{S_i} of the edge belongs to the interior of a triangle $(\mathbf{x}_K, \mathbf{x}_{K_i}, \mathbf{x}_{K_{J_{i,1}}})$ for exactly one of the two possible $J_{i,1} \neq i$ and $0 \leq J_{i,1} \leq d$. When $d = 3$, the barycenter \mathbf{x}_{S_i} of the face belongs to the interior of a tetrahedron $(\mathbf{x}_K, \mathbf{x}_{K_i}, \mathbf{x}_{K_{J_{i,1}}}, \mathbf{x}_{K_{J_{i,2}}})$ for exactly one pair $(J_{i,1}, J_{i,2})$, up to a permutation, of the three possible pairs $J_{i,1}, J_{i,2} \neq i$ and $0 \leq J_{i,1}, J_{i,2} \leq d$. Let us denote $J_{i,0} = i$. Then, the vector $\overrightarrow{\mathbf{x}_K \mathbf{x}_{S_i}}$ decompose on the basis $(\overrightarrow{\mathbf{x}_K \mathbf{x}_{K_{J_{i,k}}}})_{0 \leq k \leq d-1}$ as

$$\overrightarrow{\mathbf{x}_K \mathbf{x}_{S_i}} = \sum_{k=0}^{d-1} \alpha_{i,k} \overrightarrow{\mathbf{x}_K \mathbf{x}_{K_{J_{i,k}}}} \quad (1.3)$$

where $\alpha_{i,k} \geq 0$, $k = 0 \dots d-1$. Let us consider now the patch ω_K composed of K and its d neighbors:

$$\omega_K = K \cup K_0 \cup \dots \cup K_d$$

For any affine function $\xi \in P_1(\omega_K)$ over this patch, let us denote

$$\begin{aligned}\delta_{K,i}(\xi) &= \sum_{k=0}^{d-1} \alpha_{i,k} \left(\xi(\mathbf{x}_{K_{J_{i,k}}}) - \xi(\mathbf{x}_K) \right), \quad i = 0 \dots d-1 \\ &= \xi(\mathbf{x}_{S_i}) - \xi(\mathbf{x}_K) \quad \text{from (1.3)}\end{aligned}$$

In other terms, $\delta_{K,i}(\xi)$ represents the departure of the value of ξ at \mathbf{x}_{S_i} from its average $\xi(\mathbf{x}_K)$ on the element K .

Let now $(\varphi_i)_{0 \leq i \leq d-1}$ denote the Lagrangian basis in K associated to the set of nodes $(\mathbf{x}_{S_i})_{0 \leq i \leq d-1}$:

$$\begin{aligned}\varphi_i(\mathbf{x}_{S_j}) &= \delta_{i,j}, \quad 0 \leq i, j \leq d-1 \\ \sum_{i=0}^{d-1} \varphi_i(\mathbf{x}) &= 1, \quad \forall \mathbf{x} \in K\end{aligned}$$

The affine function $\xi \in P_1(\omega_K)$ expresses on this basis as

$$\xi(\mathbf{x}) = \xi(\mathbf{x}_K) + \sum_{i=0}^{d-1} \delta_{K,i}(\xi) \varphi_i(\mathbf{x}), \quad \forall \mathbf{x} \in K$$

Let now $u_h \in \mathbb{P}_{1d}(\mathcal{T}_h)$. On any element $K \in \mathcal{T}_h$, let us introduce its average value:

$$\bar{u}_K = \frac{1}{\text{meas}(K)} \int_K u_h(\mathbf{x}) \, dx$$

and its departure from its average value:

$$\tilde{u}_K(\mathbf{x}) = u_h|_K(\mathbf{x}) - \bar{u}_K, \quad \forall \mathbf{x} \in K$$

Notice that $u_h \notin P_1(\omega_K)$. Let us extends $\delta_{K,i}$ to u_h as

$$\delta_{K,i}(u_h) = \sum_{k=0}^{d-1} \alpha_{i,k} \left(\bar{u}_{K_{J_{i,k}}} - \bar{u}_K \right), \quad i = 0 \dots d-1$$

Since $u_h \notin P_1(\omega_K)$, we have $\tilde{u}_K(\mathbf{x}_{K_{J_{i,k}}}) \neq \delta_{K,i}(u_h)$ in general. The idea is then to capture oscillations by controlling the departure of the values $\tilde{u}_K(\mathbf{x}_{K_{J_{i,k}}})$ from the values $\delta_{K,i}(u_h)$. Thus, associate to $u_h \in \mathbb{P}_{1d}(\mathcal{T}_h)$ the quantities

$$\Delta_{K,i}(u_h) = \text{minmod}_{\text{TVB}} \left(\tilde{u}_K(\mathbf{x}_{K_{J_{i,k}}}), \theta \delta_{K,i}(u_h) \right)$$

for all $i = 0 \dots d-1$ and where $\theta \geq 1$ is a parameter of the limiter and

$$\text{minmod}_{\text{TVB}}(a, b) = \begin{cases} a & \text{when } |a| \leq Mh^2 \\ \text{minmod}(a, b) & \text{otherwise} \end{cases}$$

where $M > 0$ is a tunable parameter which can be evaluated from the curvature of the initial datum at its extrema by setting

$$M = \sup_{\mathbf{x} \in \Omega, \nabla u_0(\mathbf{x})=0} |\nabla \otimes \nabla u_0| \quad (1.4)$$

Introduced in [21], the basic idea is to deactivate the limiter when space derivatives are of order h^2 . This improves the limiter behavior near smooth local extrema. The minmod function is defined by

$$\text{minmod}(a, b) = \begin{cases} \text{sgn}(a) \min(|a|, |b|) & \text{when } \text{sgn}(a) = \text{sgn}(b) \\ 0 & \text{otherwise} \end{cases}$$

Then, for all $i = 0 \dots d-1$ we define

$$r_K(u_h) = \frac{\sum_{j=0}^{d-1} \max(0, -\Delta_{K,j}(u_h))}{\sum_{j=0}^{d-1} \max(0, \Delta_{K,j}(u_h))} \geq 0$$

$$\hat{\Delta}_{K,i}(u_h) = \min(1, r_K(u_h)) \max(0, \Delta_{K,i}(u_h)) - \min(1, 1/r_K(u_h)) \max(0, -\Delta_{K,i}(u_h)), \quad i = 0 \dots d-1 \text{ when } r_K(u_h) \neq 0$$

Finally, the limited function $\Lambda_h(u_h)$ is defined element by element for all element $K \in \mathcal{T}_h$ for all $\mathbf{x} \in K$ by

$$\Lambda_h(u_h)|_K(\mathbf{x}) = \begin{cases} \bar{u}_K + \sum_{i=1}^{d-1} \Delta_{K,i}(u_h) \varphi_i(\mathbf{x}) & \text{when } r_K(u_h) = 0 \\ \bar{u}_K + \sum_{i=1}^{d-1} \hat{\Delta}_{K,i}(u_h) \varphi_i(\mathbf{x}) & \text{otherwise} \end{cases}$$

Notice that there are two types of computations involved in the limiter: one part is independent of u_h and depends only upon the mesh: $J_{i,k}$ and $\alpha_{i,k}$ on each element. It can be computed one time for all. The other part depends upon the values of u_h .

Notice that the limiter preserves the average value of u_h on each element K and also the functions that are globally affine on the patch ω_K . Also we have, inside each element K and for all side index $i = 0 \dots d-1$:

$$|\Lambda_h(u_h)|_K(\mathbf{x}_{S_i}) - \bar{u}_K| \leq \max(|\Delta_{K,i}(u_h)|, |\hat{\Delta}_{K,i}(u_h)|) \leq |\Delta_{K,i}(u_h)| \leq |u_h|_K(\mathbf{x}_{S_i}) - \bar{u}_K|$$

It means that, inside each element, the gradient of the P_1 limited function is no larger than that of the original one.

The limiter on an element close to the boundary should takes into account the inflow condition. In [7], the modifications are described.

1.3 Example: the Burgers equation

As an illustration, let us consider now the test with the one-dimensional ($d = 1$) Burgers equation for a propagating slant step (see e.g. [3, p. 87]) in $\Omega =]0, 1[$. We have $\mathbf{f}(u) = u^2/2$, for all $u \in \mathbb{R}$. In that case, the Godunov flux (1.1d), introduced page 11, can be computed explicitly for any $\nu = (\nu_0) \in \mathbb{R}^d$ and $a, b \in \mathbb{R}$:

$$\Phi(\nu; a, b) = \begin{cases} \nu_0 \min(a^2, b^2) / 2 & \text{when } \nu_0 \geq 0 \text{ and } a \leq b \text{ or } \nu_0 \leq 0 \text{ and } a \geq b \\ \nu_0 \max(a^2, b^2) / 2 & \text{otherwise} \end{cases}$$

Example file 1.2: burgers.icc

```
1 point f (const Float& u) { return point (sqr(u)/2); }
```

Example file 1.3: burgers_flux_godunov.icc

```
1 Float phi (const point& nu, Float a, Float b) {
2   if ((nu[0] >= 0 && a <= b) || (nu[0] <= 0 && a >= b))
3     return nu[0]*min(sqr(a),sqr(b))/2;
4   else
5     return nu[0]*max(sqr(a),sqr(b))/2;
6 }
```

1.3.1 Computing an exact solution

An exact solution is useful for testing numerical methods. The computation of such an exact solution for the one dimensional Burgers equation is described by Hartens *et al.* [15]. The authors consider first the problem with a periodic boundary condition:

(P): find $u :]0, T[\times]-1, 1[\rightarrow \mathbb{R}$ such that

$$\begin{aligned} \frac{\partial u}{\partial t} + \frac{\partial}{\partial x} \left(\frac{u^2}{2} \right) &= 0 \quad \text{in }]0, T[\times]-1, 1[\\ u(t=0, x) &= \alpha + \beta \sin(\pi x + \gamma), \quad \text{a.e. } x \in]-1, 1[\\ u(t, x=-1) &= u(t, x=1) \quad \text{a.e. } t \in]0, T[\end{aligned}$$

where α, β and γ are real parameters. Let us denote w the solution of the problem when $\beta = 1$ and $\alpha = \gamma = 0$; i.e. with the initial condition $w(t=0, x) = \sin(\pi x)$, a.e. $x \in]-1, 1[$. For any $x \in]0, 1[$ and $t > 0$, the solution $\bar{w} = w(t, x)$ satisfies the characteristic relation

$$\bar{w} = \sin(\pi(x - \bar{w}t))$$

This nonlinear relation can be solved by a Newton algorithm. Then, for $x \in]-1, 0[$, the solution is completed by symmetry: $w(t, x) = -w(t, -x)$. Finally, the general solution for any α, β and $\gamma = 0$ writes $u(t, x) = \alpha + w(\beta t, x - \alpha t + \gamma)$. File ‘harten.icc’ implements this approach.

Example file 1.4: harten.icc

```

1 #include "harten0.icc"
2 struct harten {
3     Float operator() (const point& x) const {
4         Float x0 = x[0] - a*t + c;
5         Float shift = -2*floor((x0+1)/2);
6         Float xs = x0 + shift;
7         check_macro (xs >= -1 && xs <= 1, "invalid xs=" << xs);
8         return a + b*h0 (point(xs));
9     }
10     harten (Float t1=0, Float a1=1, Float b1=0.5, Float c1=0):
11         h0(b1*t1), t(t1), a(a1), b(b1), c(c1) {}
12     Float M() const { Float pi = acos(-1.0); return sqr(pi)*b; }
13     Float min() const { return a-b; }
14     Float max() const { return a+b; }
15 protected:
16     harten0 h0;
17     Float t, a, b, c;
18 };
19 using u_init = harten;
20 using g = harten;
```

Example file 1.5: harten_show.cc

```

1 #include "rheolef.h"
2 using namespace rheolef;
3 using namespace std;
4 #include "harten.icc"
5 int main(int argc, char**argv) {
6     environment rheolef (argc, argv);
7     geo omega (argv[1]);
8     space Xh (omega, argv[2]);
9     size_t nmax = (argc > 3) ? atoi(argv[3]) : 1000;
10    Float tf = (argc > 4) ? atof(argv[4]) : 2.5;
11    Float a = (argc > 5) ? atof(argv[5]) : 1;
12    Float b = (argc > 6) ? atof(argv[6]) : 0.5;
13    Float c = (argc > 7) ? atof(argv[7]) : 0;
14    branch even("t", "u");
15    for (size_t n = 0; n <= nmax; ++n) {
16        Float t = n*tf/nmax;
17        field pi_h_u = interpolate (Xh, harten(t, a, b, c));
18        dout << even(t, pi_h_u);
19    }
20 }
```

The included file ‘`harten0.icc`’ is not shown here by is available in the example directory.

Comments

Notice that the constant M , used by the limiter in (1.4), can be explicitly computed for this solution: $M = \beta\pi^2$.

The animation of this exact solution is performed by the following commands:

```
make harten_show
mkgeo_grid -e 2000 -a -1 -b 1 > line2.geo
./harten_show line2 P1 1000 2.5 > line2-exact.branch
branch line2-exact -gnuplot
```

Fig. 1.5 shows the solution u for $\alpha = 1, \beta = 1/2$ and $\gamma = 0$. It is regular until $t = 2/\pi$ (Fig. 1.5.c) and then develops a chock for $t > 2/\pi$ (Fig. 1.5.d). After its apparition, this chock interacts with the expansion wave in $] -1, 1[$: this brings about a fast decay of the solution (Figs. 1.5.e and f). Fig. 1.5 plots also a numerical solution: its computation is the aim of the next section.

1.3.2 Numerical resolution

When replacing the periodic boundary condition with a inflow one, associated with the boundary data g , we choose g to be the value of the exact solution of the problem with periodic boundary conditions: $g(t, x) = \alpha + w(\beta t, x - \alpha t)$ for $x \in \{-1, 1\}$.

Example file 1.6: burgers_dg.cc

```

1 #include "rheolef.h"
2 using namespace rheolef;
3 using namespace std;
4 #include "harten.icc"
5 #include "burgers.icc"
6 #include "burgers_flux_godunov.icc"
7 #include "runge_kutta_ssp.icc"
8 int main(int argc, char**argv) {
9     environment rheolef (argc, argv);
10    geo omega (argv[1]);
11    space Xh (omega, argv[2]);
12    Float cfl = 1;
13    limiter_option_type lopt;
14    size_t nmax = (argc > 3) ? atoi(argv[3]) : numeric_limits<size_t>::max();
15    Float tf = (argc > 4) ? atof(argv[4]) : 2.5;
16    size_t p = (argc > 5) ? atoi(argv[5]) : pmax;
17    lopt.M = (argc > 6) ? atoi(argv[6]) : u_init().M();
18    if (nmax == numeric_limits<size_t>::max()) {
19        nmax = floor(1+tf/(cfl*omega.hmin()));
20    }
21    Float delta_t = tf/nmax;
22    form_option_type fopt;
23    fopt.invert = true;
24    trial u (Xh); test v (Xh);
25    form inv_m = integrate (u*v, fopt);
26    vector<field> uh(p+1, field(Xh,0));
27    uh[0] = interpolate (Xh, u_init());
28    branch even("t","u");
29    dout << catchmark("delta_t") << delta_t << endl
30        << even(0,uh[0]);
31    for (size_t n = 1; n <= nmax; ++n) {
32        for (size_t i = 1; i <= p; ++i) {
33            uh[i] = 0;
34            for (size_t j = 0; j < i; ++j) {
35                field lh =
36                    - integrate (dot(compose(f,uh[j]),grad_h(v)))
37                    + integrate ("internal_sides",
38                                compose (phi, normal(), inner(uh[j]), outer(uh[j]))*jump(v))
39                    + integrate ("boundary",
40                                compose (phi, normal(), uh[j], g(n*delta_t))*v);
41                uh[i] += alpha[p][i][j]*uh[j] - delta_t*beta[p][i][j]*(inv_m*lh);
42            }
43            uh[i] = limiter(uh[i], g(n*delta_t)(point(-1)), lopt);
44        }
45        uh[0] = uh[p];
46        dout << even(n*delta_t,uh[0]);
47    }
48 }

```

Comments

The Runge-Kutta time discretization combined with the discontinuous Galerkin space discretization is implemented for this test case.

The P_0 approximation is performed by the following commands:

```

make burgers_dg
mkgeo_grid -e 200 -a -1 -b 1 > line2-200.geo
./burgers_dg line2-200.geo P0 1000 2.5 > line2-200-P0.branch
branch line2-200-P0.branch -gnuplot

```

The two last commands compute the P_0 approximation of the solution, as shown on Fig. 1.5. Observe the robust behavior of the solution at the vicinity of the chock. By replacing P_0 by P_1 in the previous commands, we obtain the P_1 -discontinuous approximation.

```
./burgers_dg line2-200.geo P1d 1000 2.5 > line2-200-P1d.branch
branch line2-200-P1d.branch -gnuplot
```

Fig. 1.6 plots the error vs h for $k = 0$ and $k = 1$. Fig. 1.6.a plots in a time interval $[0, T]$ with $T = 1/\pi$, before the chock that occurs at $t = 2/\pi$. In that interval, the solution is regular and the error approximation behaves as $\mathcal{O}(h^{k+1})$. The time interval has been chosen sufficiently small for the error to depend only upon h . Fig. 1.6.b plots in a larger time interval $[0, T]$ with $T = 5/2$, that includes the chock. Observe that the error behaves as $\mathcal{O}(h)$ for both $k = 0$ and 1 . This is optimal when $k = 0$ but not when $k = 1$. This is due to the loss of regularity of the exact solution that presents a discontinuity; A similar observation can be found in [26], table 4.1.

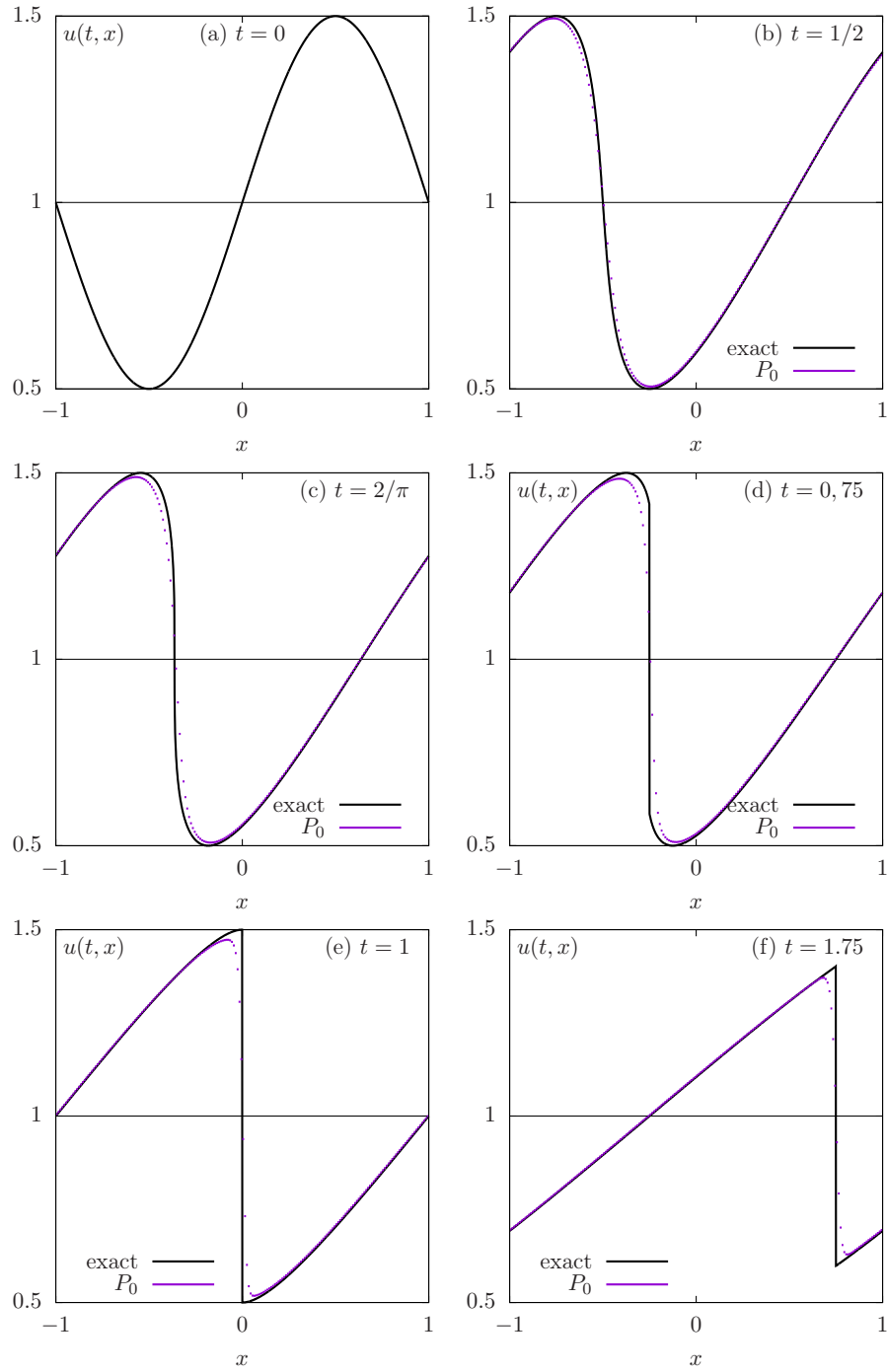


Figure 1.5: Harten's exact solution of the Burgers equation ($\alpha = 1, \beta = 1/2, \gamma = 0$). Comparison with the P_0 approximation ($h = 1/100$, RK-SSP(3)).

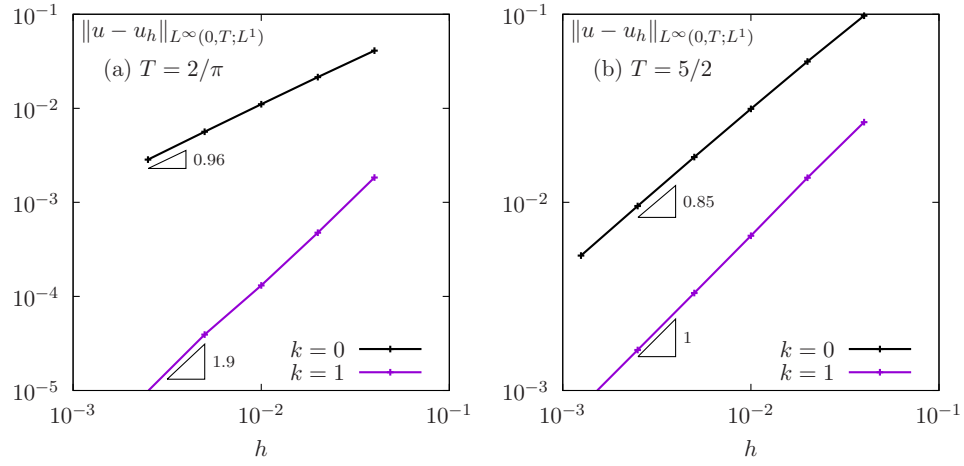


Figure 1.6: Burgers equation: error between the P_0 approximation and the exact solution of the Harten's problem ($\alpha = 1, \beta = 1/2, \gamma = 0$): (a) before chock, with $T = 1/\pi$; (b) after chock, with $T = 5/2$.

Chapter 2

Scalar second-order problems

2.1 The Poisson problem with Dirichlet boundary conditions

The Poisson problem with non-homogeneous Dirichlet boundary conditions has been already introduced in volume 1, section 2.1, page 27:

(P): find u , defined in Ω , such that

$$\begin{aligned} -\Delta u &= f \text{ in } \Omega \\ u &= g \text{ on } \partial\Omega \end{aligned}$$

where f and g are given.

The *discontinuous* finite element space is defined by:

$$X_h = \{v_h \in L^2(\Omega); v_h|_K \in P_k, \forall K \in \mathcal{T}_h\}$$

where $k \geq 1$ is the polynomial degree. As elements of X_h do not belongs to $H^1(\Omega)$, due to discontinuities at inter-elements, we introduce the broken Sobolev space:

$$H^1(\mathcal{T}_h) = \{v \in L^2(\Omega); v|_K \in H^1(K), \forall K \in \mathcal{T}_h\}$$

such that $X_h \subset H^1(\mathcal{T}_h)$. We introduce the following bilinear form $a_h(.,.)$ and linear form $l_h(.,.)$, defined for all $u, v \in H^1(\mathcal{T}_h)$ by (see e.g. [11, p. 125 and 127], eqn. (4.12)):

$$a_h(u, v) = \int_{\Omega} \nabla_h u \cdot \nabla_h v \, dx + \sum_{S \in \mathcal{S}_h} \int_S (\eta_s \llbracket u \rrbracket \llbracket v \rrbracket - \{\!\!\{ \nabla_h u \cdot \mathbf{n} \}\!\!\} \llbracket v \rrbracket - \llbracket u \rrbracket \{\!\!\{ \nabla_h v \cdot \mathbf{n} \}\!\!\}) \, ds \quad (2.1)$$

$$l_h(v) = \int_{\Omega} f u \, dx + \int_{\partial\Omega} (\eta_s g v - g \nabla_h v \cdot \mathbf{n}) \, ds \quad (2.2)$$

The last term involves a sum over \mathcal{S}_h , the set of *all sides* of the mesh \mathcal{T}_h , i.e. the internal sides and the boundary sides. By convenience, the definition of the jump and average are extended to all boundary sides as $\llbracket u \rrbracket = \{\!\!\{ u \}\!\!\} = u$. Notice that, as for the previous transport problem, the Dirichlet boundary condition $u = g$ is weakly imposed on $\partial\Omega$ via the integrals on the boundary. Finally, $\eta_s > 0$ is a stabilization parameter on a side S . The stabilization term associated to η_s is present in order to achieve coercivity: it penalize interface and boundary jumps. A common choice is $\eta_s = \beta h_s^{-1}$ where $\beta > 0$ is a constant and h_s is a local length scale associated to the current side S . One drawback to this choice is that it requires the end user to specify the numerical constant β . From one hand, if the value of this parameter is not sufficiently large, the form $a_h(.,.)$ is not coercive and the approximate solution develops instabilities and do not converge [12]. From other hand, if the value of this parameter is too large, it affects the overall efficiency of the iterative solver of the linear system: the spectral condition number of the matrix associated to $a_h(.,.)$

grows linearly with this parameter [4]. An explicit choice of penalty parameter is proposed in [20]: $\eta_s = \beta \varpi_s$ where $\beta = (k+1)(k+d)/d$ and

$$\varpi_s = \begin{cases} \frac{\text{meas}(\partial K)}{\text{meas}(K)} & \text{when } S = K \cap \partial\Omega \text{ is a boundary side} \\ \max\left(\frac{\text{meas}(\partial K_0)}{\text{meas}(K_0)}, \frac{\text{meas}(\partial K_1)}{\text{meas}(K_1)}\right) & \text{when } S = K_0 \cap K_1 \text{ is an internal side} \end{cases}$$

Notice that ϖ_s scales as h_s^{-1} . Now, the computation of the penalty parameter is fully automatic and the convergence of the method is always guaranted to converge. Moreover, this choice has been founded to be sharp and it preserves the optimal efficiency of the iterative solvers. Finally, the discrete variational formulation writes:

$(FV)_h$: find $u_h \in X_h$ such that

$$a_h(u_h, v_h) = l_h(v_h), \forall v_h \in X_h$$

The following code implement this problem in the **Rheolef** environment.

Example file 2.1: dirichlet_dg.cc

```

1 #include "rheolef.h"
2 using namespace rheolef;
3 using namespace std;
4 #include "cosinusprod_laplace.icc"
5 int main(int argc, char**argv) {
6     environment rheolef (argc, argv);
7     geo omega (argv[1]);
8     space Xh (omega, argv[2]);
9     size_t d = omega.dimension();
10    size_t k = Xh.degree();
11    Float beta = (k+1)*(k+d)/d;
12    trial u (Xh); test v (Xh);
13    form a = integrate (dot(grad_h(u), grad_h(v)))
14              + integrate ("sides", beta*penalty()*jump(u)*jump(v)
15                          - jump(u)*average(dot(grad_h(v), normal()))
16                          - jump(v)*average(dot(grad_h(u), normal())));
17    field lh = integrate (f(d)*v)
18                  + integrate ("boundary", beta*penalty()*g(d)*v
19                              - g(d)*dot(grad_h(v), normal()));
20    solver sa (a.uu());
21    field uh(Xh);
22    uh.set_u() = sa.solve(lh.u());
23    dout << uh;
24 }

```

Comments

The `penalty()` pseudo-function implements the computation of ϖ_s in **Rheolef**. The right-hand side f and g are given by (2.1), volume 1, page 28. In that case, the exact solution is known. Running the one-dimensional case writes:

```

make dirichlet_dg
mkgeo_grid -e 10 > line.geo
./dirichlet_dg line P1d | field -
./dirichlet_dg line P2d | field -

```

Fig. 2.1 plots the one-dimensional solution when $k = 1$ for two meshes. Observe that the jumps at inter-element nodes decreases very fast with mesh refinement and are no more perceptible on the plots. Recall that the Dirichlet boundary conditions at $x = 0$ and $x = 1$ is only weakly imposed: the corresponding jump at the boundary is also not perceptible.

The two-dimensional case writes:

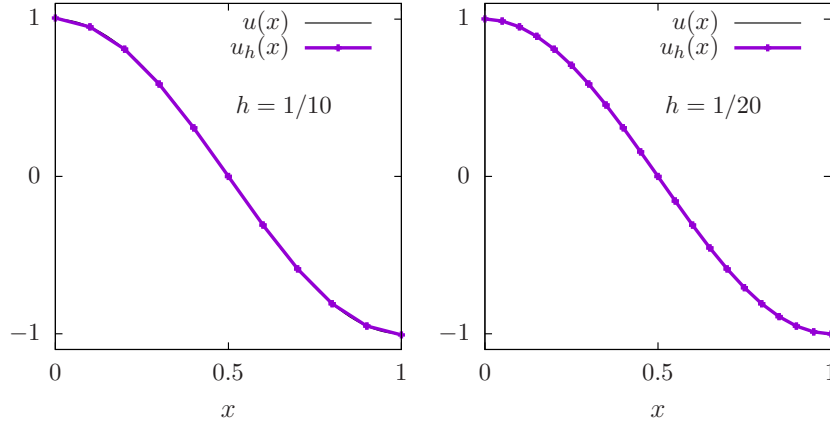


Figure 2.1: The discontinuous Galerkin method for the Poisson problem when $k = 1$ and $d = 1$.

```
mkgeo_grid -t 10 > square.geo
./dirichlet_dg square P1d | field -elevation -
./dirichlet_dg square P2d | field -elevation -
```

and the three-dimensional one

```
mkgeo_grid -T 10 > cube.geo
./dirichlet_dg cube P1d | field -
./dirichlet_dg cube P2d | field -
```

Error analysis

The space $H^1(\mathcal{T}_h)$ is equipped with the norm $\|\cdot\|_{1,h}$, defined for all $v \in H^1(\mathcal{T}_h)$ by [11, p. 128]:

$$\|v\|_{1,h}^2 = \|\nabla_h v\|_{0,\Omega}^2 + \sum_{S \in \mathcal{S}_h} \int_S h_s^{-1} \llbracket v \rrbracket^2 ds$$

The code `cosinusprod_error_dg.cc` compute the error in these norms. This code it is not listed here but is available in the **Rheolef** example directory. The computation of the error writes:

```
make cosinusprod_error_dg
./dirichlet_dg square P1d | cosinusprod_error_dg
```

Fig. 2.2 plots the error $u - u_h$ in L^2 , L^∞ and the $\|\cdot\|_{1,h}$ norms. The L^2 and L^∞ error norms behave as $\mathcal{O}(h^{k+1})$ for all $k \geq 0$, while the $\|\cdot\|_{1,h}$ one behaves as $\mathcal{O}(h^k)$, which is optimal.

2.2 The Helmholtz problem with Neumann boundary conditions

The Poisson problem with non-homogeneous Neumann boundary conditions has been already introduced in volume 1, section 2.2, page 35:

(P): find u , defined in Ω , such that

$$\begin{aligned} u - \Delta u &= f \text{ in } \Omega \\ \frac{\partial u}{\partial n} &= g \text{ on } \partial\Omega \end{aligned}$$

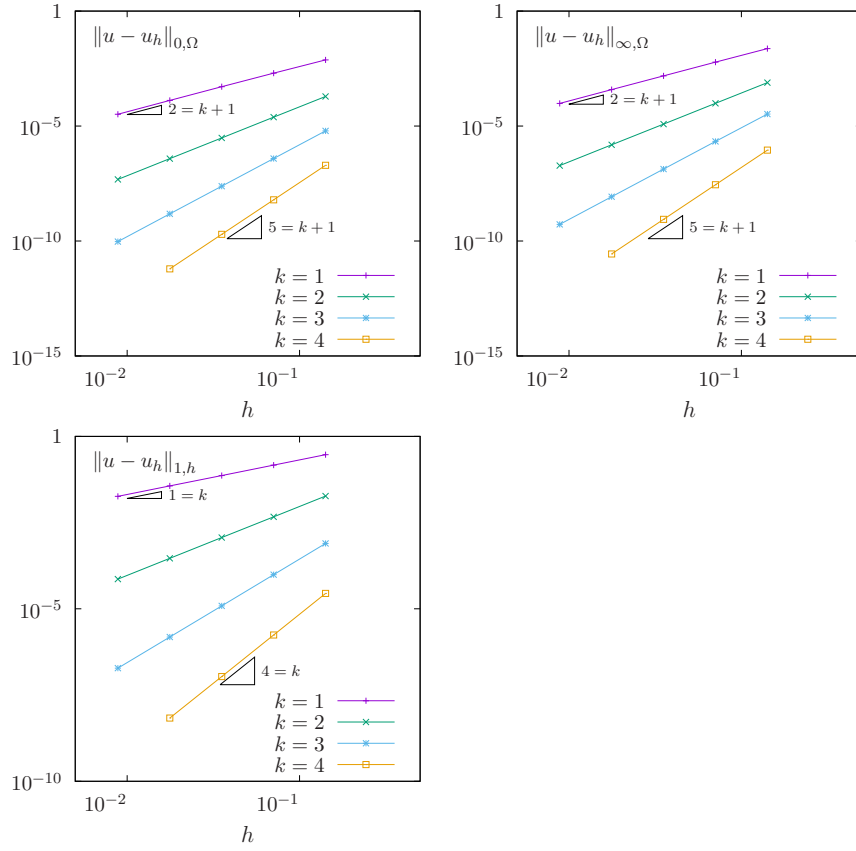


Figure 2.2: The discontinuous Galerkin method for the Poisson problem: convergence when $d = 2$.

where f and g are given. We introduce the following bilinear form $a_h(.,.)$ and linear for $l_h(.,.)$, defined for all $u, v \in H^1(\mathcal{T}_h)$ by (see e.g. [11, p. 127], eqn. (4.16)):

$$a_h(u, v) = \int_{\Omega} (u v + \nabla_h u \cdot \nabla_h v) \, dx \quad (2.3)$$

$$+ \sum_{S \in \mathcal{S}_h^{(i)}} \int_S (\beta \varpi_s \llbracket u \rrbracket \llbracket v \rrbracket - \{\{\nabla_h u \cdot \mathbf{n}\}\} \llbracket v \rrbracket - \llbracket u \rrbracket \{\{\nabla_h v \cdot \mathbf{n}\}\}) \, ds \quad (2.4)$$

$$l_h(v) = \int_{\Omega} f u \, dx + \int_{\partial\Omega} g v \, ds \quad (2.5)$$

Let us comment the changes between these forms and those used for the Poisson problem with Dirichlet boundary conditions. The Poisson operator $-\Delta$ has been replaced by the Helmholtz one $I - \Delta$ in order to have an unique solution. Remark also that the sum is performed in (2.1) for all internal sides in $\mathcal{S}_h^{(i)}$, while, in (2.1), for Dirichlet boundary conditions, it was for all sides in \mathcal{S}_h , i.e. for both boundary and internal sides. Also, the right-hand-side linear form $l_h(.,.)$ do no more involves any sum over sides.

Finally, the discrete variational formulation writes:

$(FV)_h$: find $u_h \in X_h$ such that

$$a_h(u_h, v_h) = l_h(v_h), \quad \forall v_h \in X_h$$

The following code implement this problem in the **Rheolef** environment.

Example file 2.2: neumann_dg.cc

```

1 #include "rheolef.h"
2 using namespace rheolef;
3 using namespace std;
4 #include "sinusprod_helmholtz.icc"
5 int main(int argc, char**argv) {
6     environment rheolef (argc, argv);
7     geo omega (argv[1]);
8     space Xh (omega, argv[2]);
9     size_t d = omega.dimension();
10    size_t k = Xh.degree();
11    Float beta = (k+1)*(k+d)/d;
12    trial u (Xh); test v (Xh);
13    form a = integrate (u*v + dot(grad_h(u),grad_h(v)))
14             + integrate ("internal_sides",
15                         beta*penalty()*jump(u)*jump(v)
16                         - jump(u)*average(dot(grad_h(v),normal()))
17                         - jump(v)*average(dot(grad_h(u),normal())));
18    field lh = integrate (f(d)*v) + integrate ("boundary", g(d)*v);
19    solver sa (a.uu());
20    field uh (Xh);
21    uh.set_u() = sa.solve(lh.u());
22    dout << uh;
23 }

```

Comments

The right-hand side f and g are given by (2.2), volume 1, page 28. In that case, the exact solution is known. Running the program is obtained from the non-homogeneous Dirichlet case, by replacing `dirichlet_dg` by `neumann_dg`.

2.3 Nonlinear scalar hyperbolic problems with diffusion

A time-dependent nonlinear second order problem with nonlinear first order dominated terms problem writes:

(P): find u , defined in $]0, T[\times \Omega$, such that

$$\frac{\partial u}{\partial t} + \operatorname{div} \mathbf{f}(u) - \varepsilon \Delta u = 0 \quad \text{in }]0, T[\times \Omega \quad (2.6a)$$

$$u(t=0) = u_0 \quad \text{in } \Omega \quad (2.6b)$$

$$u = g \quad \text{on }]0, T[\times \partial\Omega \quad (2.6c)$$

where $\varepsilon > 0$, $T > 0$, $\Omega \subset \mathbb{R}^d$, $d = 1, 2, 3$ and the initial condition \mathbf{u}_0 being known. The function $\mathbf{f} : \mathbb{R} \rightarrow \mathbb{R}^d$ is also known and supposed to be continuously differentiable. The initial data u_0 , defined in Ω , and the boundary one, g , defined on $\partial\Omega$ are given.

Comparing (2.6a)-(2.6c) with the non-diffusive case (1.1a)-(1.1c) page 10, the second order term has been added in (2.6a) and the upstream boundary condition has been replaced by a Dirichlet one (2.6c).

2.4 Example: the Burgers equation with diffusion

2.4.1 Problem statement and its exact solution

Our model problem in this chapter is the one-dimensional Burgers equation. It was introduced in section 1.3, page 14 with the choice $\mathbf{f}(u) = u^2/2$, for all $u \in \mathbb{R}$. Equation (2.6a) admits an exact

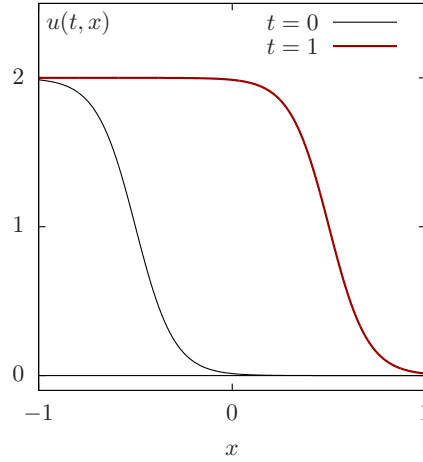


Figure 2.3: An exact solution for the Burgers equation with diffusion ($\varepsilon = 10^{-1}$, $x_0 = -1/2$).

solution

$$u(t, x) = 1 - \tanh\left(\frac{x - x_0 - t}{2\varepsilon}\right) \quad (2.7)$$

Example file 2.3: burgers_diffusion_exact.icc

```

1 struct u_exact {
2   Float operator() (const point& x) const {
3     return 1 - tanh((x[0]-x0-t)/(2*epsilon)); }
4   u_exact (Float e1, Float t1=0) : epsilon(e1), t(t1), x0(-0.5) {}
5   Float M() const { return 0; }
6   Float epsilon, t, x0;
7 };
8 using u_init = u_exact;
9 using g = u_exact;

```

The solution is represented on Fig. 2.3. Here x_0 represents the position of the front at $t = 0$ and ε is a characteristic width of the front. The initial and boundary conditions are chosen such that $u(t, x)$ is the solution of (2.6a)-(2.6c).

Fig. 2.4.a plots the error versus Δt for the semi-implicit scheme when $k = 1$ and 2, and for $h = 2/50$. The time step for which the error becomes independent upon Δt and depends only upon h is of about $\Delta t = 10^{-3}$ when $k = 1$ and of about 10^{-5} when $k = 2$. This approach is clearly inefficient for high order polynomial k and a higher order time scheme is required.

Fig. 2.4.b plots the error versus Δt for the Runge-Kutta semi-implicit scheme with $p = 3$, $k = 1$ and $h = 2/200$. The scheme is clearly only first-order, which is still unexpected. More work is required...

2.4.2 Space discretization

The *discontinuous* finite element space is defined by:

$$X_h = \{v_h \in L^2(\Omega); v_h|_K \in P_k, \forall K \in \mathcal{T}_h\}$$

where $k \geq 1$ is the polynomial degree. As elements of X_h do not belong to $H^1(\Omega)$, due to discontinuities at inter-elements, we introduce the broken Sobolev space:

$$H^1(\mathcal{T}_h) = \{v \in L^2(\Omega); v|_K \in H^1(K), \forall K \in \mathcal{T}_h\}$$

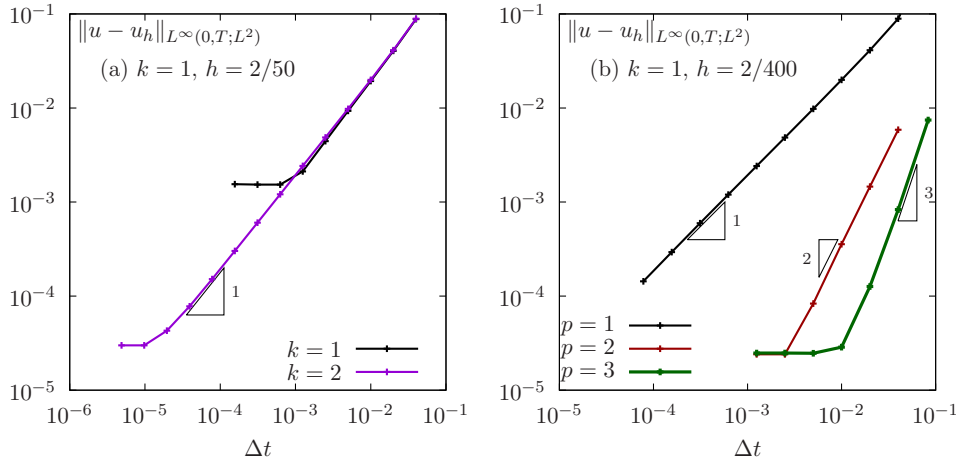


Figure 2.4: Convergence of the first order semi-implicit scheme for the Burgers equation with diffusion ($\epsilon = 0.1, T = 1$). (a) first order semi-implicit scheme ; (b) Runge-Kutta semi-implicit scheme with $p = 3$.

such that $X_h \subset H^1(\mathcal{T}_h)$. As for the Dirichlet problem, introduce the following bilinear form $a_h(.,.)$ and linear for $\ell_h(.,.)$, defined for all $u, v \in H^1(\mathcal{T}_h)$ by (see e.g. [11, p. 125 and 127], eqn. (4.12)):

$$a_h(u, v) = \int_{\Omega} \nabla_h u \cdot \nabla_h v \, dx + \sum_{S \in \mathcal{S}_h} \int_S (\eta_s \llbracket u \rrbracket \llbracket v \rrbracket - \llbracket \nabla_h u \cdot \mathbf{n} \rrbracket \llbracket v \rrbracket - \llbracket u \rrbracket \llbracket \nabla_h v \cdot \mathbf{n} \rrbracket) \, ds \quad (2.8)$$

$$\ell_h(v) = \int_{\partial\Omega} (\eta_s g v - g \nabla_h v \cdot \mathbf{n}) \, ds \quad (2.9)$$

The semi-discrete problem writes in variational form [11, p. 100]:

$(P)_h$: find $u_h \in C^1([0, T], X_h)$ such that

$$\begin{aligned} \int_{\Omega} \frac{\partial u_h}{\partial t} v_h \, dx + \int_{\Omega} G_h(u_h) v_h \, dx + \varepsilon a_h(u_h, v_h) &= \varepsilon \ell_h(v_h), \quad \forall v_h \in X_h \\ u_h(t=0) &= \pi_h(u_0) \end{aligned}$$

where G_h has been introduced in (1.2), page 11.

2.4.3 Time discretization

Explicit Runge-Kutta scheme is possible for this problem but it leads to an excessive Courant-Friedrichs-Levy condition for the time step Δt , that is required to be lower than an upper bound that varies in $\mathcal{O}(h^2)$. The idea here is to continue to explicit the first order nonlinear terms and implicit the linear second order terms. Semi-implicit second order Runge-Kutta scheme was first introduced in 1997 by Ascher, Ruuth and Spiteri [1] and then extended in 2001 to third and fourth order by Calvo, de Frutos and Novo [2]. In 2015, Wang, Shu and Zhang [24, 25] applied it in the context of the discontinuous Galerkin method. The finite dimensional problem can be rewritten as

$(P)_h$: find $u_h \in C^1([0, T], X_h)$ such that

$$\begin{aligned} \frac{\partial u_h}{\partial t} + G_h(t, u_h) + A_h(t, u_h) &= 0, \quad \forall t \in]0, T[\\ u_h(t=0) &= \pi_h(u_0) \end{aligned}$$

where G_h has been introduced in (1.2), page 11 and A_h denotes the diffusive term. The semi-implicit Runge-Kutta scheme with $p \geq 0$ intermediate steps writes at time step t_n :

$$u_h^{n,0} = u_h^n \quad (2.10a)$$

$$u_h^{n,i} = u_h^n - \Delta t \sum_{j=1}^i \alpha_{i,j} A_h(t_{n,j}, u_h^{n,j}) - \Delta t \sum_{j=0}^{i-1} \tilde{\alpha}_{i,j} G_h(t_{n,j}, u_h^{n,j}), \quad i = 1, \dots, p \quad (2.10b)$$

$$u_h^{n+1} = u_h^n - \Delta t \sum_{i=1}^p \beta_i A_h(t_{n,i}, u_h^{n,i}) - \Delta t \sum_{i=0}^p \tilde{\beta}_i G_h(t_{n,i}, u_h^{n,i}) \quad (2.10c)$$

where $(u_h^{n,i})_{1 \leq i \leq p}$ are the $p \geq 1$ intermediate states, $t_{n,i} = t_n + \gamma_i \Delta t$, $\gamma_i = \sum_{j=1}^i \alpha_{i,j} = \sum_{j=0}^{i-1} \tilde{\alpha}_{i,j}$, and $(\alpha_{i,j})_{0 \leq i, j \leq p}$, $(\tilde{\alpha}_{i,j})_{0 \leq i, j \leq p}$, $(\beta_i)_{0 \leq i \leq p}$ and $(\tilde{\beta}_i)_{0 \leq i \leq p}$ are the coefficients of the scheme [1, 2, 25]. At each time step, have to solve p linear systems. From (2.10b) we get for all $i = 1, \dots, p$:

$$(I + \Delta t \alpha_{i,i} A_h(t_{n,i})) u_h^{n,i} = u_h^n - \Delta t \sum_{j=1}^{i-1} \alpha_{i,j} A_h(t_{n,j}, u_h^{n,j}) - \Delta t \sum_{j=0}^{i-1} \tilde{\alpha}_{i,j} G_h(t_{n,j}, u_h^{n,j})$$

Notice that when the matrix coefficients of $A_h(t, \cdot)$ are independent of t , the matrix involved on the right-hand-side of the previous equation can be factored one time for all.

Example file 2.4: burgers_diffusion_dg.cc

```

1 #include "rheolef.h"
2 using namespace rheolef;
3 using namespace std;
4 #include "burgers.icc"
5 #include "burgers_flux_godunov.icc"
6 #include "runge_kutta_semiimplicit.icc"
7 #include "burgers_diffusion_exact.icc"
8 #undef NEUMANN
9 #include "burgers_diffusion_operators.icc"
10 int main(int argc, char**argv) {
11     environment rheolef (argc, argv);
12     geo omega (argv[1]);
13     space Xh (omega, argv[2]);
14     size_t k = Xh.degree();
15     Float epsilon = (argc > 3) ? atof(argv[3]) : 0.1;
16     size_t nmax = (argc > 4) ? atoi(argv[4]) : 500;
17     Float tf = (argc > 5) ? atof(argv[5]) : 1;
18     size_t p = (argc > 6) ? atoi(argv[6]) : min(k+1, rk::pmax);
19     Float delta_t = tf/nmax;
20     size_t d = omega.dimension();
21     Float beta = (k+1)*(k+d)/d;
22     trial u (Xh); test v (Xh);
23     form m = integrate (u*v);
24     form_option_type fopt;
25     fopt.invert = true;
26     form inv_m = integrate (u*v, fopt);
27     form a = epsilon*(
28         integrate (dot(grad_h(u), grad_h(v)))
29 #ifdef NEUMANN
30         + integrate ("internal_sides",
31 #else // NEUMANN
32         + integrate ("sides",
33 #endif // NEUMANN
34         beta*penalty()*jump(u)*jump(v)
35         - jump(u)*average(dot(grad_h(v), normal()))
36         - jump(v)*average(dot(grad_h(u), normal()))));
37     vector<solver> sc (p+1);
38     for (size_t i = 1; i <= p; ++i) {
39         form ci = m + delta_t*rk::alpha[p][i][i]*a;
40         sc[i] = solver(ci.uu());
41     }
42     vector<field> uh(p+1, field(Xh,0));
43     uh[0] = interpolate (Xh, u_init(epsilon));
44     branch even("t", "u");
45     dout << catchmark("epsilon") << epsilon << endl
46         << even(0, uh[0]);
47     for (size_t n = 0; n < nmax; ++n) {
48         Float tn = n*delta_t;
49         Float t = tn + delta_t;
50         field uh_next = uh[0] - delta_t*rk::tilde_beta[p][0]*(inv_m*gh(epsilon, tn, uh[0], v));
51         for (size_t i = 1; i <= p; ++i) {
52             Float ti = tn + rk::gamma[p][i]*delta_t;
53             field rhs = m*uh[0] - delta_t*rk::tilde_alpha[p][i][0]*gh(epsilon, tn, uh[0], v);
54             for (size_t j = 1; j <= i-1; ++j) {
55                 Float tj = tn + rk::gamma[p][j]*delta_t;
56                 rhs -= delta_t*( rk::alpha[p][i][j]*(a*uh[j] - lh(epsilon, tj, v))
57                     + rk::tilde_alpha[p][i][j]*gh(epsilon, tj, uh[j], v));
58             }
59             rhs += delta_t*rk::alpha[p][i][i]*lh (epsilon, ti, v);
60             uh[i].set_u() = sc[i].solve (rhs.u());
61             uh_next -= delta_t*(inv_m*( rk::beta[p][i]*(a*uh[i] - lh(epsilon, ti, v))
62                 + rk::tilde_beta[p][i]*gh(epsilon, ti, uh[i], v)));
63         }
64         uh_next = limiter(uh_next);
65         dout << even(tn+delta_t, uh_next);
66         uh[0] = uh_next;
67     }
68 }

```

Example file 2.5: burgers_diffusion_operators.icc

```

1 field lh (Float epsilon, Float t, const test& v) {
2 #ifdef NEUMANN
3     return field (v.get_vf_space(), 0);
4 #else // NEUMANN
5     size_t d = v.get_vf_space().get_geo().dimension();
6     size_t k = v.get_vf_space().degree();
7     Float beta = (k+1)*(k+d)/d;
8     return epsilon*integrate ("boundary",
9         beta*penalty()*g(epsilon,t)*v
10        - g(epsilon,t)*dot(grad_h(v),normal()));
11 #endif // NEUMANN
12 }
13 field gh (Float epsilon, Float t, const field& uh, const test& v) {
14     return - integrate (dot(compose(f,uh),grad_h(v)))
15        + integrate ("internal_sides",
16            compose (phi, normal(), inner(uh), outer(uh))*jump(v))
17        + integrate ("boundary",
18            compose (phi, normal(), uh, g(epsilon,t))*v);
19 }

```

Running the program

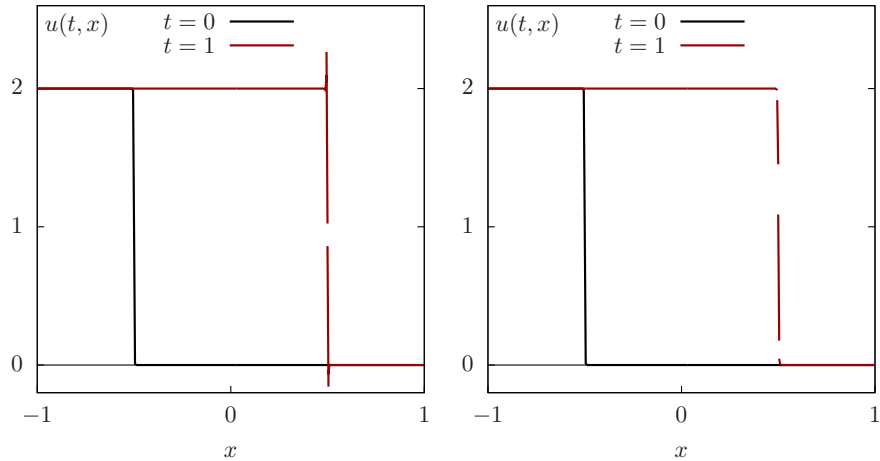


Figure 2.5: Burgers equation with a small diffusion ($\varepsilon = 10^{-3}$). Third order in time semi-implicit scheme with P_{1d} element. (left) without limiter ; (right) with limiter.

Running the program writes with $h = 2/400$ and $\varepsilon = 10^{-2}$ writes:

```

make burgers_diffusion_dg
mkgeo_grid -e 400 -a -1 -b 1 > line.geo
./burgers_diffusion_dg line P1d 0.01 1000 1 3 > line.branch
branch -gnuplot line.branch -umin -0.1 -umax 2.1

```

Decreasing $\varepsilon = 10^{-3}$ leads to a sharper solution:

```

./burgers_diffusion_dg line P1d 0.001 1000 1 3 > line.branch
branch -gnuplot line.branch -umin -0.1 -umax 2.1

```

As mentioned in [25], the time step should be chosen smaller when ε decreases. The result is shown on Fig. 2.5.left. Observe the oscillations near the smoothed shock when there is no limiter while the value goes outside $[0, 2]$. Conversely, with a limiter (see Fig. 2.5.right) the approximate solution is decreasing and there is no more oscillations: the values remains in the range $[0 : 2]$.

Part II

Fluids and solids computations

Chapter 3

The linear elasticity and the Stokes problems

3.1 The linear elasticity problem

The elasticity problem (4.2) has been introduced in volume 1, section 4.1, page 51.

(P): find \mathbf{u} such that

$$\begin{aligned} -\operatorname{div}(\lambda \operatorname{div}(\mathbf{u}).I + 2D(\mathbf{u})) &= \mathbf{f} \text{ in } \Omega \\ \mathbf{u} &= \mathbf{g} \text{ on } \partial\Omega \end{aligned}$$

where $\lambda \geq -1$ is a constant and \mathbf{f}, \mathbf{g} given. This problem is a natural extension to vector-valued field of the Poisson problem with Dirichlet boundary conditions.

The variational formulation writes:

(FV)_h: find $\mathbf{u} \in \mathbf{V}(\mathbf{g})$ such that

$$a(\mathbf{u}, \mathbf{v}) = l_h(\mathbf{v}), \quad \forall \mathbf{v} \in \mathbf{V}(0)$$

where

$$\begin{aligned} \mathbf{V}(\mathbf{g}) &= \{\mathbf{v} \in H^1(\Omega)^d; \mathbf{v} = \mathbf{g} \text{ on } \partial\Omega\} \\ a(\mathbf{u}, \mathbf{v}) &= \int_{\Omega} (\lambda \operatorname{div}(\mathbf{u}) \operatorname{div}(\mathbf{v}) + 2D(\mathbf{u}) : D(\mathbf{v})) \, dx \\ l(\mathbf{v}) &= \int_{\Omega} \mathbf{f} \cdot \mathbf{v} \, dx \end{aligned}$$

The discrete variational formulation writes:

(FV)_h: find $u_h \in \mathbf{X}_h$ such that

$$a_h(u_h, v_h) = l_h(v_h), \quad \forall v_h \in \mathbf{X}_h$$

where

$$\begin{aligned} \mathbf{X}_h &= \{\mathbf{v}_h \in L^2(\Omega)^d; \mathbf{v}_h|_K \in P_K^d, \forall K \in \mathcal{T}_h\} \\ a_h(\mathbf{u}, \mathbf{v}) &= \int_{\Omega} (\lambda \operatorname{div}_h(\mathbf{u}) \operatorname{div}_h(\mathbf{v}) + 2D_h(\mathbf{u}) : D_h(\mathbf{v})) \, dx \\ &\quad + \sum_{S \in \mathcal{S}_h} \int_S (\beta \varpi_s \llbracket \mathbf{u} \rrbracket \cdot \llbracket \mathbf{v} \rrbracket - \llbracket \mathbf{u} \rrbracket \cdot \llbracket \lambda \operatorname{div}_h(\mathbf{v}) \mathbf{n} + 2D_h(\mathbf{v}) \mathbf{n} \rrbracket - \llbracket \mathbf{v} \rrbracket \cdot \llbracket \lambda \operatorname{div}_h(\mathbf{u}) \mathbf{n} + 2D_h(\mathbf{u}) \mathbf{n} \rrbracket) \, ds \\ l_h(\mathbf{v}) &= \int_{\Omega} \mathbf{f} \cdot \mathbf{v} \, dx + \int_{\partial\Omega} \mathbf{g} \cdot (\beta \varpi_s \mathbf{v} - \lambda \operatorname{div}_h(\mathbf{v}) \mathbf{n} - 2D_h(\mathbf{v}) \mathbf{n}) \, ds \end{aligned}$$

where $k \geq 1$ is the polynomial degree in \mathbf{X}_h .

Example file 3.1: elasticity_taylor_dg.cc

```

1 #include "rheolef.h"
2 using namespace rheolef;
3 using namespace std;
4 #include "taylor.icc"
5 int main(int argc, char**argv) {
6     environment rheolef (argc, argv);
7     geo omega (argv[1]);
8     space Xh (omega, argv[2], "vector");
9     Float lambda = (argc > 3) ? atof(argv[3]) : 1;
10    size_t d = omega.dimension();
11    size_t k = Xh.degree();
12    Float beta = (k+1)*(k+d)/d;
13    trial u (Xh); test v (Xh);
14    form a = integrate (lambda*div_h(u)*div_h(v) + 2*ddot(Dh(u),Dh(v)))
15            + integrate (omega.sides(),
16                        beta*penalty()*dot(jump(u),jump(v))
17                        - lambda*dot(jump(u),average(div_h(v)*normal()))
18                        - lambda*dot(jump(v),average(div_h(u)*normal()))
19                        - 2*dot(jump(u),average(Dh(v)*normal()))
20                        - 2*dot(jump(v),average(Dh(u)*normal())));
21    field lh = integrate (dot(f(),v))
22                + integrate (omega.boundary(),
23                            beta*penalty()*dot(g(),jump(v))
24                            - lambda*dot(g(),average(div_h(v)*normal()))
25                            - 2*dot(g(),average(Dh(v)*normal())));
26    solver sa (a.uu());
27    field uh(Xh);
28    uh.set_u() = sa.solve(lh.u());
29    dout << uh;
30 }

```

Comments

The data are given when $d = 2$ by:

$$\mathbf{g}(x) = \begin{pmatrix} -\cos(\pi x_0) \sin(\pi x_1) \\ \sin(\pi x_0) \cos(\pi x_1) \end{pmatrix} \quad \text{and} \quad \mathbf{f} = 2\pi^2 \mathbf{g} \quad (3.1)$$

This choice is convenient since the exact solution is known $\mathbf{u} = \mathbf{g}$. This benchmark solution was proposed in 1923 by Taylor [23] in the context of the Stokes problem. Notice that the solution is independent of λ since $\text{div}(\mathbf{u}) = 0$.

Example file 3.2: taylor.icc

```

1 struct g {
2     point operator() (const point& x) const {
3         return point(-cos(pi*x[0])*sin(pi*x[1]),
4                     sin(pi*x[0])*cos(pi*x[1])); }
5     g() : pi(acos(Float(-1.0))) {}
6     const Float pi;
7 };
8 struct f {
9     point operator() (const point& x) const { return 2*sqr(pi)*_g(x); }
10    f() : pi(acos(Float(-1.0))), _g() {}
11    const Float pi; g _g;
12 };

```

As the exact solution is known, the error can be computed. The code elasticity_taylor_error_dg.cc compute the error in L^2 , L^∞ and energy norms. This code it is not listed here but is available in the **Rheolef** example directory. The computation writes:

```
make elasticity_taylor_dg elasticity_taylor_error_dg
```

```
mkgeo_grid -t 10 > square.geo
./elasticity_taylor_dg square P1d | ./elasticity_taylor_error_dg
./elasticity_taylor_dg square P2d | ./elasticity_taylor_error_dg
```

3.2 The Stokes problem

Let us consider the Stokes problem for the driven cavity in $\Omega =]0, 1[^d$, $d = 2, 3$. The problem has been introduced in volume 1, section 4.4, page 62.

(P): find \mathbf{u} and p , defined in Ω , such that

$$\begin{aligned} -\operatorname{div}(2D(\mathbf{u})) + \nabla p &= \mathbf{f} \text{ in } \Omega, \\ -\operatorname{div} \mathbf{u} &= 0 \text{ in } \Omega, \\ \mathbf{u} &= \mathbf{g} \text{ on } \partial\Omega \end{aligned}$$

where \mathbf{f} and \mathbf{g} are given. This problem is the extension to divergence free vector fields of the elasticity problem. The variational formulation writes:

$(VF)_h$ find $\mathbf{u} \in \mathbf{V}(\mathbf{g})$ and $p \in L^2(\Omega)$ such that:

$$\begin{aligned} a(\mathbf{u}, \mathbf{v}) + b(\mathbf{v}, p) &= l(\mathbf{v}), \quad \forall \mathbf{v} \in \mathbf{V}(0), \\ b(\mathbf{u}, q) &= 0, \quad \forall q \in L^2(\Omega) \end{aligned} \tag{3.2}$$

where

$$\begin{aligned} \mathbf{V}(\mathbf{g}) &= \{\mathbf{v} \in H^1(\Omega)^d; \mathbf{v} = \mathbf{g} \text{ on } \partial\Omega\} \\ a(\mathbf{u}, \mathbf{v}) &= \int_{\Omega} 2D(\mathbf{u}) : D(\mathbf{v}) \, dx \\ b(\mathbf{u}, q) &= - \int_{\Omega} \operatorname{div}(\mathbf{u}) q \, dx \\ l(\mathbf{v}) &= \int_{\Omega} \mathbf{f} \cdot \mathbf{v} \, dx \end{aligned}$$

The discrete variational formulation writes:

$(VF)_h$ find $\mathbf{u}_h \in \mathbf{X}_h$ and $p_h \in Q_h$ such that:

$$\begin{aligned} a_h(\mathbf{u}_h, \mathbf{v}_h) + b_h(\mathbf{v}_h, p_h) &= l_h(\mathbf{v}_h), \quad \forall \mathbf{v}_h \in \mathbf{X}_h, \\ b_h(\mathbf{u}_h, q_h) - c_h(p_h, q_h) &= k_h(q), \quad \forall q_h \in Q_h. \end{aligned} \tag{3.3}$$

The discontinuous finite element spaces are defined by:

$$\begin{aligned} \mathbf{X}_h &= \{\mathbf{v}_h \in L^2(\Omega)^d; \mathbf{v}_h|_K \in P_k^d, \quad \forall K \in \mathcal{T}_h\} \\ Q_h &= \{q_h \in L^2(\Omega)^d; q_h|_K \in P_k^d, \quad \forall K \in \mathcal{T}_h\} \end{aligned}$$

where $k \geq 1$ is the polynomial degree. Notice that velocity and pressure are approximated by the same polynomial order. This method was introduced by [9] and some recent theoretical results can be founded in [10]. The forms are defined for all $u, v \in H^1(\mathcal{T}_h)^d$ and $q \in L^2(\Omega)$ by (see

e.g. [11, p. 249]):

$$\begin{aligned}
a_h(\mathbf{u}, \mathbf{v}) &= \int_{\Omega} 2D_h(\mathbf{u}) : D_h(\mathbf{v}) \, dx \\
&\quad + \sum_{S \in \mathcal{S}_h} \int_S (\beta \varpi_s [\![\mathbf{u}]\!] \cdot [\![\mathbf{v}]\!] - [\![\mathbf{u}]\!] \cdot \{\{2D_h(\mathbf{v})\mathbf{n}\}\} - [\![\mathbf{v}]\!] \cdot \{\{2D_h(\mathbf{u})\mathbf{n}\}\}) \, ds \\
b_h(\mathbf{u}, q) &= \int_{\Omega} \mathbf{u} \cdot \nabla_h q \, dx - \sum_{S \in \mathcal{S}_h^{(i)}} \int_S \{\{\mathbf{u}\}\} \cdot \mathbf{n} [q] \, ds \\
c_h(p, q) &= \sum_{S \in \mathcal{S}_h^{(i)}} \int_S h_s [p] [q] \, ds \\
l_h(\mathbf{v}) &= \int_{\Omega} \mathbf{f} \cdot \mathbf{v} \, dx + \int_{\partial\Omega} \mathbf{g} \cdot (\beta \varpi_s \mathbf{v} - 2D_h(\mathbf{v})\mathbf{n}) \, ds \\
k_h(q) &= \int_{\partial\Omega} \mathbf{g} \cdot \mathbf{n} q \, ds
\end{aligned}$$

The stabilization form c_h controls the pressure jump accross internal sides. This stabilization term is necessary when using equal order polynomial approximation for velocity and pressure. The definition of the forms is grouped in a subroutine: it will be reused later for the Navier-Stokes problem.

Example file 3.3: stokes_dirichlet_dg.icc

```

1 void stokes_dirichlet_dg (const space& Xh, const space& Qh,
2   form& a, form& b, form& c, form& mp, field& lh, field& kh,
3   quadrature_option_type qopt = quadrature_option_type())
4 {
5   size_t k = Xh.degree();
6   size_t d = Xh.get_geo().dimension();
7   Float beta = (k+1)*(k+d)/d;
8   trial u (Xh), p (Qh);
9   test v (Xh), q (Qh);
10  a = integrate (2*ddot(Dh(u), Dh(v)), qopt)
11    + integrate ("sides", beta*penalty()*dot(jump(u), jump(v))
12      - 2*dot(jump(u), average(Dh(v)*normal()))
13      - 2*dot(jump(v), average(Dh(u)*normal())), qopt);
14  lh = integrate (dot(f(), v), qopt)
15    + integrate ("boundary", beta*penalty()*dot(g(), v)
16      - 2*dot(g(), Dh(v)*normal()), qopt);
17  b = integrate (dot(u, grad_h(q)), qopt)
18    + integrate ("internal_sides", - dot(average(u), normal())*jump(q), qopt);
19  kh = integrate ("boundary", dot(g(), normal())*q, qopt);
20  c = integrate ("internal_sides", h_local()*jump(p)*jump(q), qopt);
21  mp = integrate (p*q, qopt);
22 }

```

A simple test program writes:

Example file 3.4: stokes_taylor_dg.cc

```

1 #include "rheolef.h"
2 using namespace rheolef;
3 using namespace std;
4 #include "taylor.icc"
5 #include "stokes_dirichlet_dg.icc"
6 int main(int argc, char**argv) {
7     environment rheolef (argc, argv);
8     geo omega (argv[1]);
9     space Xh (omega, argv[2], "vector");
10    space Qh (omega, argv[2]);
11    form a, b, c, mp;
12    field lh, kh;
13    stokes_dirichlet_dg (Xh, Qh, a, b, c, mp, lh, kh);
14    field uh (Xh, 0), ph (Qh, 0);
15    solver_abtb stokes (a.uu(), b.uu(), c.uu(), mp.uu());
16    stokes.solve (lh.u(), kh.u(), uh.set_u(), ph.set_u());
17    dout << catchmark("u") << uh
18         << catchmark("p") << ph;
19 }

```

Comments

The data are given when $d = 2$ by (3.1). This choice is convenient since the exact solution is known $\mathbf{u} = \mathbf{g}$ and $p = 0$. The code `stokes_taylor_error_dg.cc` compute the error in L^2 , L^∞ and energy norms. This code it is not listed here but is available in the **Rheolef** example directory. The computation writes:

```

make stokes_taylor_dg stokes_taylor_error_dg
mkgeo_grid -t 10 > square.geo
./stokes_taylor_dg square P1d | ./stokes_taylor_error_dg
./stokes_taylor_dg square P2d | ./stokes_taylor_error_dg

```

3.3 The stationnary Navier-Stokes problem

3.3.1 Problem statement

The Navier-Stokes problem has been already introduced in volume 1, section 3.3 page 37. Here we consider the stationnary version of this problem. Let $Re \geq 0$ be the Reynolds number. The problem writes:

(P): find \mathbf{u} and p , defined in Ω , such that

$$\begin{aligned}
 Re(\mathbf{u} \cdot \nabla) \mathbf{u} - \operatorname{div}(2D(\mathbf{u})) + \nabla p &= \mathbf{f} \text{ in } \Omega, \\
 -\operatorname{div} \mathbf{u} &= 0 \text{ in } \Omega, \\
 \mathbf{u} &= \mathbf{g} \text{ on } \partial\Omega
 \end{aligned}$$

Notice that, when $Re > 0$, the problem is nonlinear, due to the inertia term $\mathbf{u} \cdot \nabla \mathbf{u}$. When $Re = 0$ the problem reduces to the linear Stokes problem, presented in the previous section/

The variationnall formulation of this nonlinear problem writes:

(FV): find $\mathbf{u} \in \mathbf{V}(\mathbf{g})$ and $p \in L^2(\Omega)$ such that

$$\begin{aligned}
 Ret(\mathbf{u}; \mathbf{u}, \mathbf{v}) + a(\mathbf{u}, \mathbf{v}) + b(\mathbf{v}, p) &= l(\mathbf{v}), \forall \mathbf{v} \in \mathbf{V}(0), \\
 b(\mathbf{u}, q) &= 0, \forall q \in L^2(\Omega)
 \end{aligned}$$

where the space $\mathbf{V}(\mathbf{g})$ and forms a , b and l are given as in the previous section 3.2 for the Stokes problem and the trilinear form $t(\cdot; \cdot, \cdot)$ is given by:

$$t(\mathbf{w}; \mathbf{u}, \mathbf{v}) = \int_{\Omega} ((\mathbf{w} \cdot \nabla) \mathbf{u}) \cdot \mathbf{v} \, dx$$

3.3.2 The discrete problem

Let

$$t(\mathbf{w}; \mathbf{u}, \mathbf{u}) = \int_{\Omega} (\mathbf{w} \cdot \nabla \mathbf{u}) \cdot \mathbf{u} \, dx$$

Observe that, for all $\mathbf{u}, \mathbf{w} \in H^1(\Omega)^d$ we have

$$\begin{aligned} \int_{\Omega} (\mathbf{w} \cdot \nabla \mathbf{u}) \cdot \mathbf{u} \, dx &= \sum_{i,j=0}^{d-1} \int_{\Omega} u_i w_j \partial_j(u_i) \, dx \\ &= \sum_{i,j=0}^{d-1} - \int_{\Omega} u_i \partial_j(u_i w_j) \, dx + \int_{\partial\Omega} u_i^2 w_j n_j \, ds \\ &= \sum_{i,j=0}^{d-1} - \int_{\Omega} u_i \partial_j(u_i) w_j \, dx - \int_{\Omega} u_i^2 \partial_j(w_j) \, dx + \int_{\partial\Omega} u_i^2 w_j n_j \, ds \\ &= - \int_{\Omega} (\mathbf{w} \cdot \nabla \mathbf{u}) \cdot \mathbf{u} \, dx - \int_{\Omega} \operatorname{div}(\mathbf{w}) |\mathbf{u}|^2 \, dx + \int_{\partial\Omega} \mathbf{w} \cdot \mathbf{n} |\mathbf{u}|^2 \, ds \end{aligned} \quad (3.4)$$

Thus

$$t(\mathbf{w}; \mathbf{u}, \mathbf{u}) = \int_{\Omega} (\mathbf{w} \cdot \nabla \mathbf{u}) \cdot \mathbf{u} \, dx = -\frac{1}{2} \int_{\Omega} \operatorname{div}(\mathbf{w}) |\mathbf{u}|^2 \, dx + \frac{1}{2} \int_{\partial\Omega} \mathbf{w} \cdot \mathbf{n} |\mathbf{u}|^2 \, ds$$

When $\operatorname{div}(\mathbf{w}) = 0$, the trilinear form $t(\cdot, \cdot, \cdot)$ reduces to a boundary term: it is formally skew-symmetric. The skew-symmetry of t is an important property: let $(\mathbf{v}, q) = (\mathbf{u}, p)$ as test functions in (FV) . We obtain:

$$a(\mathbf{u}, \mathbf{u}) = l(\mathbf{u})$$

In other words, we obtain the same energy balance as for the Stokes flow and inertia do not contribute to the energy balance. This is an important property and we aim at obtaining the same one at the discrete level. As the discrete solution \mathbf{u}_h is not exactly divergence free, following Temam, we introduce the following modified trilinear form:

$$t^*(\mathbf{w}; \mathbf{u}, \mathbf{v}) = \int_{\Omega} \left((\mathbf{w} \cdot \nabla \mathbf{u}) \cdot \mathbf{v} + \frac{1}{2} \operatorname{div}(\mathbf{w}) \mathbf{u} \cdot \mathbf{v} \right) dx - \frac{1}{2} \int_{\partial\Omega} (\mathbf{w} \cdot \mathbf{n}) \mathbf{u} \cdot \mathbf{v} \, ds, \quad \forall \mathbf{u}, \mathbf{v}, \mathbf{w} \in H^1(\Omega)^d$$

This form integrates the non-vanishing terms and we have:

$$t^*(\mathbf{w}; \mathbf{u}, \mathbf{u}) = 0, \quad \forall \mathbf{u}, \mathbf{w} \in H^1(\Omega)^d$$

When the discrete solution is not exactly divergence free, it is better to use t^* than t .

The discontinuous finite element spaces \mathbf{X}_h and Q_h and forms a_h, b_h, c_h, l_h and k_h are defined as in the previous section. Let us introduce t_h^* , the following discrete trilinear form, defined for all $\mathbf{u}_h, \mathbf{v}_h, \mathbf{w}_h \in \mathbf{X}_h$:

$$t_h^*(\mathbf{w}_h; \mathbf{u}_h, \mathbf{v}_h) = \int_{\Omega} \left((\mathbf{w}_h \cdot \nabla_h \mathbf{u}_h) \cdot \mathbf{v}_h + \frac{1}{2} \operatorname{div}_h(\mathbf{w}_h) \mathbf{u}_h \cdot \mathbf{v}_h \right) dx - \frac{1}{2} \int_{\partial\Omega} (\mathbf{w}_h \cdot \mathbf{n}) \mathbf{u}_h \cdot \mathbf{v}_h \, ds$$

Notice that t_h^* is similar to t^* : the gradient and divergence has been replaced by their broken counterpart in the first term. As $\mathbf{X}_h \not\subset H^1(\Omega)^d$, the skew-symmetry property is not expected to be true at the discrete level. Then

$$t_h^*(\mathbf{w}_h; \mathbf{u}_h, \mathbf{u}_h) = \sum_{K \in \mathcal{T}_h} \int_K \left((\mathbf{w}_h \cdot \nabla \mathbf{u}_h) \cdot \mathbf{u}_h + \frac{1}{2} \operatorname{div}(\mathbf{w}_h) |\mathbf{u}_h|^2 \right) dx - \frac{1}{2} \int_{\partial\Omega} (\mathbf{w}_h \cdot \mathbf{n}) |\mathbf{u}_h|^2 \, ds$$

As the restriction of \mathbf{u}_h and \mathbf{w}_h to each $K \in \mathcal{T}_h$ belongs to $H^1(K)^d$, we have, using a similar integration by part:

$$\int_K (\mathbf{w}_h \cdot \nabla \mathbf{u}_h) \cdot \mathbf{u}_h \, dx = -\frac{1}{2} \int_K \operatorname{div}(\mathbf{w}_h) |\mathbf{u}_h|^2 \, dx + \frac{1}{2} \int_{\partial K} (\mathbf{w}_h \cdot \mathbf{n}) |\mathbf{u}_h|^2 \, ds$$

Thus

$$t_h^*(\mathbf{w}_h; \mathbf{u}_h, \mathbf{u}_h) = \frac{1}{2} \sum_{K \in \mathcal{T}_h} \int_{\partial K} (\mathbf{w}_h \cdot \mathbf{n}) |\mathbf{u}_h|^2 \, ds - \frac{1}{2} \int_{\partial \Omega} (\mathbf{w}_h \cdot \mathbf{n}) |\mathbf{u}_h|^2 \, ds$$

The terms on boundary sides vanish while those on internal sides can be grouped:

$$t_h^*(\mathbf{w}_h; \mathbf{u}_h, \mathbf{u}_h) = \frac{1}{2} \sum_{S \in \mathcal{S}_h^{(i)}} \int_S \llbracket |\mathbf{u}_h|^2 \mathbf{w}_h \rrbracket \cdot \mathbf{n} \, ds$$

The jump term $\llbracket (\mathbf{u}_h \cdot \mathbf{v}_h) \mathbf{w}_h \rrbracket \cdot \mathbf{n}$ is not easily manageable and could be developed. A short computation shows that, for all scalar fields ϕ, φ we have on any internal side:

$$\llbracket \phi \varphi \rrbracket = \llbracket \phi \rrbracket \{\varphi\} + \{\phi\} \llbracket \varphi \rrbracket \quad (3.5)$$

$$\{\phi \varphi\} = \{\phi\} \{\varphi\} + \frac{1}{4} \llbracket \phi \rrbracket \llbracket \varphi \rrbracket \quad (3.6)$$

Then

$$\begin{aligned} t_h^*(\mathbf{w}_h; \mathbf{u}_h, \mathbf{u}_h) &= \frac{1}{2} \sum_{S \in \mathcal{S}_h^{(i)}} \int_S (\{\mathbf{w}_h\} \cdot \mathbf{n} \llbracket |\mathbf{u}_h|^2 \rrbracket + \llbracket \mathbf{w}_h \rrbracket \cdot \mathbf{n} \{\|\mathbf{u}_h\|^2\}) \, ds \\ &= \sum_{S \in \mathcal{S}_h^{(i)}} \int_S \left(\{\mathbf{w}_h\} \cdot \mathbf{n} (\llbracket \mathbf{u}_h \rrbracket \cdot \{\mathbf{u}_h\}) + \frac{1}{2} \llbracket \mathbf{w}_h \rrbracket \cdot \mathbf{n} \{\|\mathbf{u}_h\|^2\} \right) \, ds \end{aligned}$$

Thus, as expected, the skew-symmetry property is no more satisfied at the discrete level, due to the jumps of the fields at the inter-element boundaries. Following the previous idea, we introduce the following modified discrete trilinear form:

$$\begin{aligned} t_h(\mathbf{w}_h; \mathbf{u}_h, \mathbf{v}_h) &= t_h^*(\mathbf{w}_h; \mathbf{u}_h, \mathbf{v}_h) - \sum_{S \in \mathcal{S}_h^{(i)}} \int_S \left(\{\mathbf{w}_h\} \cdot \mathbf{n} (\llbracket \mathbf{u}_h \rrbracket \cdot \{\mathbf{v}_h\}) + \frac{1}{2} \llbracket \mathbf{w}_h \rrbracket \cdot \mathbf{n} \{\mathbf{u}_h \cdot \mathbf{v}_h\} \right) \, ds \\ &= \int_{\Omega} \left((\mathbf{w}_h \cdot \nabla \mathbf{u}_h) \cdot \mathbf{v}_h + \frac{1}{2} \operatorname{div}_h(\mathbf{w}_h) \mathbf{u}_h \cdot \mathbf{v}_h \right) \, dx - \frac{1}{2} \int_{\partial \Omega} (\mathbf{w}_h \cdot \mathbf{n}) \mathbf{u}_h \cdot \mathbf{v}_h \, ds \\ &\quad - \sum_{S \in \mathcal{S}_h^{(i)}} \int_S \left(\{\mathbf{w}_h\} \cdot \mathbf{n} (\llbracket \mathbf{u}_h \rrbracket \cdot \{\mathbf{v}_h\}) + \frac{1}{2} \llbracket \mathbf{w}_h \rrbracket \cdot \mathbf{n} \{\mathbf{u}_h \cdot \mathbf{v}_h\} \right) \, ds \quad (3.7) \end{aligned}$$

This expression has been proposed by Pietro and Ern [10, p. 22], eqn (72) (see also [11, p. 272], eqn (6.57)). The boundary term introduced in t_h may be compensated in the right-hand side:

$$l_h^*(\mathbf{v}) := l_h(\mathbf{v}) - \frac{Re}{2} \int_{\partial \Omega} (\mathbf{g} \cdot \mathbf{n}) \mathbf{g} \cdot \mathbf{v}_h \, ds$$

Notice that the boundary term introduced in t_h is compensated in the right-hand side l_h^* .

Example file 3.5: inertia.icc

```

1  template<class W, class U, class V>
2  form inertia (W w, U u, V v,
3    quadrature_option_type qopt = quadrature_option_type())
4  {
5    return
6      integrate (dot(grad_h(u)*w,v) + 0.5*div_h(w)*dot(u,v), qopt)
7    + integrate ("boundary", - 0.5*dot(w,normal())*dot(u,v), qopt)
8    + integrate ("internal_sides",
9      - dot(average(w),normal())*dot(jump(u),average(v))
10     - 0.5*dot(jump(w),normal())
11       *(dot(average(u),average(v)) + 0.25*dot(jump(u),jump(v))), qopt);
12 }
13 field inertia_fix_rhs (test v,
14   quadrature_option_type qopt = quadrature_option_type())
15 {
16   return integrate("boundary", - 0.5*dot(g(),normal())*dot(g(),v), qopt);
17 }

```

The discrete problem is

$(FV)_h$: find $\mathbf{u}_h \in \mathbf{X}_h$ and $p \in Q_h$ such that

$$\begin{aligned}
 Ret_h(\mathbf{u}_h; \mathbf{u}_h, \mathbf{v}_h) + a_h(\mathbf{u}_h, \mathbf{v}_h) + b_h(\mathbf{v}_h, p_h) &= l_h^*(\mathbf{v}_h), \quad \forall \mathbf{v}_h \in \mathbf{X}_h, \\
 b_h(\mathbf{u}_h, q_h) - c_h(p_h, q_h) &= k_h(q), \quad \forall q_h \in Q_h
 \end{aligned} \tag{3.8}$$

The simplest approach for solving the discrete problem is to consider a fixed-point algorithm. The sequence $(\mathbf{u}_h^{(k)})_{k \geq 0}$ is defined by recurrence as:

- $k = 0$: let $\mathbf{u}_h^{(0)} \in \mathbf{X}_h$ being known.

- $k \geq 0$: let $\mathbf{u}_h^{(k-1)} \in \mathbf{X}_h$ given. Find $\mathbf{u}_h^{(k)} \in \mathbf{X}_h$ and $p_h^{(k)} \in Q_h$ such that

$$\begin{aligned}
 Ret_h(\mathbf{u}_h^{(k-1)}; \mathbf{u}_h^{(k)}, \mathbf{v}_h) + a_h(\mathbf{u}_h^{(k)}, \mathbf{v}_h) + b_h(\mathbf{v}_h, p_h^{(k)}) &= l_h^*(\mathbf{v}_h), \quad \forall \mathbf{v}_h \in \mathbf{X}_h, \\
 b_h(\mathbf{u}_h^{(k)}, q_h) - c_h(p_h^{(k)}, q_h) &= k_h(q), \quad \forall q_h \in Q_h.
 \end{aligned}$$

At each step $k \geq 0$, this algorithm involves a linear subproblem of Stokes-type.

Example file 3.6: navier_stokes_taylor_dg.cc

```

1 #include "rheolef.h"
2 using namespace rheolef;
3 using namespace std;
4 #include "taylor.icc"
5 #include "stokes_dirichlet_dg.icc"
6 #include "inertia.icc"
7 int main(int argc, char**argv) {
8     environment rheolef (argc, argv);
9     geo omega (argv[1]);
10    space Xh (omega, argv[2], "vector");
11    space Qh (omega, argv[2]);
12    Float Re = (argc > 3) ? atof(argv[3]) : 1;
13    size_t max_iter = (argc > 4) ? atoi(argv[4]) : 1;
14    form a, b, c, mp;
15    field lh, kh;
16    stokes_dirichlet_dg (Xh, Qh, a, b, c, mp, lh, kh);
17    field uh (Xh, 0), ph (Qh, 0);
18    solver_abtb stokes (a.uu(), b.uu(), c.uu(), mp.uu());
19    stokes.solve (lh.u(), kh.u(), uh.set_u(), ph.set_u());
20    trial u (Xh); test v (Xh);
21    form a1 = a + Re*inertia (uh, u, v);
22    lh += Re*inertia_fix_rhs (v);
23    derr << "#k r as" << endl;
24    for (size_t k = 0; k < max_iter; ++k) {
25        solver_abtb stokes (a1.uu(), b.uu(), c.uu(), mp.uu());
26        stokes.solve (lh.u(), kh.u(), uh.set_u(), ph.set_u());
27        form th = inertia (uh, u, v);
28        a1 = a + Re*th;
29        field rh = a1*uh + b.trans_mult(ph) - lh;
30        derr << k << " " << rh.max_abs() << " " << th(uh,uh) << endl;
31    }
32    dout << catchmark("Re") << Re << endl
33         << catchmark("u") << uh
34         << catchmark("p") << ph;
35 }

```

Comments

The data are given when $d = 2$ by (3.1). This choice is convenient since the exact solution is known $\mathbf{u} = \mathbf{g}$ and $p = -(Re/4)(\cos(2\pi x_0) + \cos(2\pi x_1))$. The code `navier_stokes_taylor_error_dg.cc` compute the error in L^2 , L^∞ and energy norms. This code it is not listed here but is available in the **Rheolef** example directory. The computation writes:

```

make navier_stokes_taylor_dg navier_stokes_taylor_error_dg
./navier_stokes_taylor_dg square P1d 10 10 | ./navier_stokes_taylor_error_dg
./navier_stokes_taylor_dg square P2d 10 10 | ./navier_stokes_taylor_error_dg

```

3.3.3 A conservative variant

Remark the identity

$$\operatorname{div}(\mathbf{u} \otimes \mathbf{u}) = (\mathbf{u} \cdot \nabla) \mathbf{u} + \operatorname{div}(\mathbf{u}) \mathbf{u}$$

The momentum conservation can be rewritten in conservative form and the problem writes:

(\tilde{P}): find \mathbf{u} and p , defined in Ω , such that

$$\begin{aligned}
 \operatorname{div}(Re \mathbf{u} \otimes \mathbf{u} - 2D(\mathbf{u})) + \nabla p &= \mathbf{f} \text{ in } \Omega, \\
 -\operatorname{div} \mathbf{u} &= 0 \text{ in } \Omega, \\
 \mathbf{u} &= \mathbf{g} \text{ on } \partial\Omega
 \end{aligned}$$

Notice the Green formulae (see volume 1, appendix A.2, page 147):

$$\int_{\Omega} \mathbf{div}(\mathbf{u} \otimes \mathbf{u}) \cdot \mathbf{v} \, dx = - \int_{\Omega} (\mathbf{u} \otimes \mathbf{u}) : \nabla \mathbf{v} \, dx + \int_{\partial\Omega} (\mathbf{u} \cdot \mathbf{n}) (\mathbf{u} \cdot \mathbf{v}) \, ds$$

The variationnal formulation is:

(\widetilde{FV}) : find $\mathbf{u} \in \mathbf{V}(\mathbf{g})$ and $p \in L^2(\Omega)$ such that

$$\begin{aligned} Re \tilde{t}(\mathbf{u}; \mathbf{u}, \mathbf{v}) + a(\mathbf{u}, \mathbf{v}) + b(\mathbf{v}, p) &= \tilde{l}(\mathbf{v}), \quad \forall \mathbf{v} \in \mathbf{V}(0), \\ b(\mathbf{u}, q) &= 0, \quad \forall q \in L^2(\Omega) \end{aligned}$$

where the forms \tilde{t} and \tilde{l}_h are given by:

$$\begin{aligned} \tilde{t}(\mathbf{w}; \mathbf{u}, \mathbf{v}) &= - \int_{\Omega} (\mathbf{w} \otimes \mathbf{u}) : \nabla \mathbf{v} \, dx \\ \tilde{l}(\mathbf{v}) &= l(\mathbf{v}) - Re \int_{\partial\Omega} (\mathbf{g} \cdot \mathbf{n}) (\mathbf{g} \cdot \mathbf{v}) \, ds \end{aligned}$$

Notice that the right-hand side \tilde{l} contains an additional term that compensates those coming from the integration by parts. Then, with $\mathbf{v} = \mathbf{u}$:

$$\begin{aligned} \tilde{t}(\mathbf{w}; \mathbf{u}, \mathbf{u}) &= - \int_{\Omega} (\mathbf{w} \otimes \mathbf{u}) : \nabla \mathbf{u} \, dx \\ &= \int_{\Omega} \mathbf{div}(\mathbf{w} \otimes \mathbf{u}) \cdot \mathbf{u} \, dx - \int_{\partial\Omega} (\mathbf{w} \otimes \mathbf{u}) : (\mathbf{u} \otimes \mathbf{n}) \, dx \\ &= \int_{\Omega} (((\mathbf{u} \cdot \nabla) \mathbf{w}) \cdot \mathbf{u} + \mathbf{div}(\mathbf{u}) (\mathbf{u} \cdot \mathbf{w})) \, dx - \int_{\partial\Omega} (\mathbf{u} \cdot \mathbf{n}) (\mathbf{u} \cdot \mathbf{w}) \, dx \end{aligned}$$

From an integration by part similar to (3.4):

$$\int_{\Omega} (\mathbf{u} \cdot \nabla \mathbf{w}) \cdot \mathbf{u} \, dx = - \int_{\Omega} (\mathbf{u} \cdot \nabla \mathbf{u}) \cdot \mathbf{w} \, dx - \int_{\Omega} \mathbf{div}(\mathbf{u}) (\mathbf{u} \cdot \mathbf{w}) \, dx + \int_{\partial\Omega} (\mathbf{u} \cdot \mathbf{n}) (\mathbf{u} \cdot \mathbf{w}) \, ds$$

The term $(\mathbf{u} \cdot \nabla \mathbf{w}) \cdot \mathbf{u}$ do not reapper after the integration by parts: instead, it appears $(\mathbf{u} \cdot \nabla \mathbf{u}) \cdot \mathbf{w}$. Thus, the structure of the \tilde{t} trilinear form do not permit a general skew-symmetry property as it was the case for t . It requires the three arguments to be the same:

$$\tilde{t}(\mathbf{u}; \mathbf{u}, \mathbf{u}) = \int_{\Omega} (((\mathbf{u} \cdot \nabla) \mathbf{u}) \cdot \mathbf{u} + \mathbf{div}(\mathbf{u}) |\mathbf{u}|^2) \, dx - \int_{\partial\Omega} (\mathbf{u} \cdot \mathbf{n}) |\mathbf{u}|^2 \, ds$$

Using (3.4) with $\mathbf{w} = \mathbf{u}$ leads to:

$$\int_{\Omega} ((\mathbf{u} \cdot \nabla) \mathbf{u}) \cdot \mathbf{u} \, dx = - \frac{1}{2} \int_{\Omega} \mathbf{div}(\mathbf{u}) |\mathbf{u}|^2 \, dx + \frac{1}{2} \int_{\partial\Omega} (\mathbf{u} \cdot \mathbf{n}) |\mathbf{u}|^2 \, ds \quad (3.9)$$

Then

$$\tilde{t}(\mathbf{u}; \mathbf{u}, \mathbf{u}) = \frac{1}{2} \int_{\Omega} \mathbf{div}(\mathbf{u}) |\mathbf{u}|^2 \, dx - \frac{1}{2} \int_{\partial\Omega} (\mathbf{u} \cdot \mathbf{n}) |\mathbf{u}|^2 \, ds$$

When working with velocities that are not divergence-free, a possible modification of the trilinear form \tilde{t} is to consider

$$\begin{aligned} \tilde{t}^*(\mathbf{w}; \mathbf{u}, \mathbf{v}) &= \tilde{t}(\mathbf{w}; \mathbf{u}, \mathbf{v}) - \frac{1}{2} \int_{\Omega} \mathbf{div}(\mathbf{v}) (\mathbf{u} \cdot \mathbf{w}) \, dx + \frac{1}{2} \int_{\partial\Omega} (\mathbf{v} \cdot \mathbf{n}) (\mathbf{u} \cdot \mathbf{w}) \, ds \\ &= - \int_{\Omega} \left((\mathbf{w} \otimes \mathbf{u}) : D(\mathbf{v}) + \frac{1}{2} \mathbf{div}(\mathbf{v}) (\mathbf{u} \cdot \mathbf{w}) \right) \, dx + \frac{1}{2} \int_{\partial\Omega} (\mathbf{v} \cdot \mathbf{n}) (\mathbf{u} \cdot \mathbf{w}) \, ds \end{aligned}$$

Then we have

$$\tilde{t}^*(\mathbf{u}; \mathbf{u}, \mathbf{u}) = 0, \quad \forall \mathbf{u} \in H^1(\Omega)^d$$

The new variationnal formulation is:

$(\widetilde{FV})^*$: find $\mathbf{u} \in \mathbf{V}(\mathbf{g})$ and $\tilde{p} \in L^2(\Omega)$ such that

$$\begin{aligned} Re \tilde{t}^*(\mathbf{u}; \mathbf{u}, \mathbf{v}) + a(\mathbf{u}, \mathbf{v}) + b(\mathbf{v}, \tilde{p}) &= \tilde{l}(\mathbf{v}), \quad \forall \mathbf{v} \in \mathbf{V}(0), \\ b(\mathbf{u}, q) &= 0, \quad \forall q \in L^2(\Omega) \end{aligned}$$

One can easily check that when (\mathbf{u}, \tilde{p}) is a solution of $(\widetilde{FV})^*$, then (\mathbf{u}, p) is a solution of (\widetilde{FV}) with $p = \tilde{p} + Re|\mathbf{u}|/2$. The apparition of the kinetic energy term $Re|\mathbf{u}|/2$ in the modified pressure field \tilde{p} is due to the introduction of the $\text{div}(\mathbf{v})(\mathbf{u} \cdot \mathbf{w})$ term in the trilinear form \tilde{t}^* .

At the discrete level, let us define for all $\mathbf{u}_h, \mathbf{v}_h, \mathbf{w}_h \in \mathbf{X}_h$:

$$\begin{aligned} \tilde{t}_h^*(\mathbf{w}_h; \mathbf{u}_h, \mathbf{v}_h) &= - \int_{\Omega} \left((\mathbf{w}_h \otimes \mathbf{u}_h) : \nabla_h \mathbf{v}_h + \frac{1}{2} \text{div}_h(\mathbf{v}_h)(\mathbf{u}_h \cdot \mathbf{w}_h) \right) dx \\ &\quad + \frac{1}{2} \int_{\partial\Omega} (\mathbf{v}_h \cdot \mathbf{n})(\mathbf{u}_h \cdot \mathbf{w}_h) ds \end{aligned}$$

Notice that \tilde{t}_h^* is similar to \tilde{t}^* : the gradient and divergence has been replaced by their broken counterpart in the first term. As $\mathbf{X}_h \not\subset H^1(\Omega)^d$, the skew-symmetry property is not expected to be true at the discrete level. Then

$$\tilde{t}_h^*(\mathbf{u}_h; \mathbf{u}_h, \mathbf{u}_h) = - \int_{\Omega} \left((\mathbf{u}_h \otimes \mathbf{u}_h) : \nabla_h \mathbf{u}_h + \frac{1}{2} \text{div}_h(\mathbf{u}_h) |\mathbf{u}_h|^2 \right) dx + \frac{1}{2} \int_{\partial\Omega} (\mathbf{u}_h \cdot \mathbf{n}) |\mathbf{u}_h|^2 ds$$

Next, using (3.9) in each K , and then developing thanks to (3.5)-(3.6), we get

$$\begin{aligned} \tilde{t}_h^*(\mathbf{u}_h; \mathbf{u}_h, \mathbf{u}_h) &= \frac{1}{2} \int_{\partial\Omega} (\mathbf{u}_h \cdot \mathbf{n}) |\mathbf{u}_h|^2 ds - \frac{1}{2} \sum_{K \in \mathcal{T}_h} \int_{\partial K} (\mathbf{u}_h \cdot \mathbf{n}) |\mathbf{u}_h|^2 ds \\ &= -\frac{1}{2} \sum_{S \in \mathcal{S}_h^{(i)}} \int_S \llbracket (\mathbf{u}_h \cdot \mathbf{n}) |\mathbf{u}_h|^2 \rrbracket ds \\ &= -\frac{1}{2} \sum_{S \in \mathcal{S}_h^{(i)}} \int_S \left((\llbracket \mathbf{u}_h \rrbracket \cdot \mathbf{n}) \llbracket |\mathbf{u}_h|^2 \rrbracket + (\llbracket \mathbf{u}_h \rrbracket \cdot \mathbf{n}) \llbracket |\mathbf{u}_h|^2 \rrbracket \right) ds \\ &= - \sum_{S \in \mathcal{S}_h^{(i)}} \int_S \left((\llbracket \mathbf{u}_h \rrbracket \cdot \mathbf{n}) (\llbracket \mathbf{u}_h \rrbracket \cdot \llbracket \mathbf{u}_h \rrbracket) + \frac{1}{2} (\llbracket \mathbf{u}_h \rrbracket \cdot \mathbf{n}) \llbracket |\mathbf{u}_h|^2 \rrbracket \right) ds \end{aligned}$$

The idea is to integrate this term in the definition of a discrete \tilde{t}_h . One of the possibilities is

$$\begin{aligned} \tilde{t}_h(\mathbf{w}_h; \mathbf{u}_h, \mathbf{v}_h) &= \tilde{t}_h^*(\mathbf{w}_h; \mathbf{u}_h, \mathbf{v}_h) + \sum_{S \in \mathcal{S}_h^{(i)}} \int_S \left((\llbracket \mathbf{u}_h \rrbracket \cdot \mathbf{n}) (\llbracket \mathbf{w}_h \rrbracket \cdot \llbracket \mathbf{v}_h \rrbracket) + \frac{1}{2} \llbracket \mathbf{u}_h \cdot \mathbf{w}_h \rrbracket (\llbracket \mathbf{v}_h \rrbracket \cdot \mathbf{n}) \right) ds \\ &= - \int_{\Omega} \left((\mathbf{w}_h \otimes \mathbf{u}_h) : \nabla_h \mathbf{v}_h + \frac{1}{2} \text{div}_h(\mathbf{v}_h)(\mathbf{u}_h \cdot \mathbf{w}_h) \right) dx \\ &\quad + \frac{1}{2} \int_{\partial\Omega} (\mathbf{v}_h \cdot \mathbf{n})(\mathbf{u}_h \cdot \mathbf{w}_h) ds \\ &\quad + \sum_{S \in \mathcal{S}_h^{(i)}} \int_S \left((\llbracket \mathbf{u}_h \rrbracket \cdot \mathbf{n}) (\llbracket \mathbf{w}_h \rrbracket \cdot \llbracket \mathbf{v}_h \rrbracket) + \frac{1}{2} \llbracket \mathbf{u}_h \cdot \mathbf{w}_h \rrbracket (\llbracket \mathbf{v}_h \rrbracket \cdot \mathbf{n}) \right) ds \end{aligned} \quad (3.10)$$

This expression was proposed by [10, p. 21], eqn (73) (see also [11, p. 282]) follong and original idea introduced in [8].

Example file 3.7: inertia_cks.icc

```

1 form inertia (field w, trial u, test v,
2   quadrature_option_type qopt = quadrature_option_type())
3 {
4   return
5     integrate (- dot(trans(grad_h(v))*w,u) - 0.5*div_h(v)*dot(u,w), qopt)
6   + integrate ("internal_sides",
7     dot(average(u),normal())*dot(jump(v),average(w))
8     + 0.5*dot(jump(v),normal())
9     *(dot(average(u),average(w)) + 0.25*dot(jump(u),jump(w))), qopt)
10  + integrate ("boundary", 0.5*dot(v,normal())*dot(u,w), qopt);
11 }
12 field inertia_fix_rhs (test v,
13   quadrature_option_type qopt = quadrature_option_type())
14 {
15   return integrate("boundary", -dot(g(),normal())*dot(g(),v), qopt);
16 }

```

The discrete problem is

$(\widetilde{FV})_h$: find $\mathbf{u}_h \in \mathbf{X}_h$ and $\tilde{p} \in Q_h$ such that

$$\begin{aligned} Re \tilde{t}_h(\mathbf{u}_h; \mathbf{u}_h, \mathbf{v}_h) + a_h(\mathbf{u}_h, \mathbf{v}_h) + b_h(\mathbf{v}_h, \tilde{p}_h) &= \tilde{l}_h^*(\mathbf{v}_h), \quad \forall \mathbf{v}_h \in \mathbf{X}_h, \\ b_h(\mathbf{u}_h, q_h) - c_h(p_h, q_h) &= k_h(q), \quad \forall q_h \in Q_h \end{aligned}$$

A simple test program is obtained by replacing in `navier_stokes_taylor_dg.cc` the include `inertia.icc` by `inertia_cks.icc`. The compilation and run are similar.

3.3.4 A Newton solver

The discrete problems $(FV)_h$ can be put in a compact form:

$$F(\mathbf{u}_h, p_h) = 0$$

where F is defined in variationnal form:

$$\langle F(\mathbf{u}_h, p_h), (\mathbf{v}_h, q_h) \rangle = \left(\begin{aligned} &Re t_h(\mathbf{u}_h; \mathbf{u}_h, \mathbf{v}_h) + a_h(\mathbf{u}_h, \mathbf{v}_h) + b_h(\mathbf{v}_h, p_h) - l_h^*(\mathbf{v}_h) \\ &b_h(\mathbf{u}_h, q_h) - c_h(p_h, q_h) - k_h(q) \end{aligned} \right)$$

for all $(\mathbf{v}_h, q_h) \in \mathbf{X}_h \times Q_h$. Notices that, after some minor modifications in the definition of F , this method could also applies for the locally conservative formulation $(\widetilde{FV})_h$. The previous formulation is simply the variationnal expression of $F(\mathbf{u}_h, p_h) = 0$. The Newton method defines the sequence $(\mathbf{u}_h^{(k)})_{k \geq 0}$ by recurrence as:

- $k = 0$: let $\mathbf{u}_h^{(0)} \in \mathbf{X}_h$ being known.
- $k \geq 0$: let $\mathbf{u}_h^{(k-1)} \in \mathbf{X}_h$ given. Find $\delta \mathbf{u}_h \in \mathbf{X}_h$ and $\delta p_h \in Q_h$ such that

$$F'(\mathbf{u}_h^{(k-1)}, p_h^{(k-1)}) . (\delta \mathbf{u}_h, \delta p_h) = -F(\mathbf{u}_h^{(k-1)}, p_h^{(k-1)})$$

and then defines

$$\mathbf{u}_h^{(k)} = \mathbf{u}_h^{(k-1)} + \delta \mathbf{u}_h \quad \text{and} \quad p_h^{(k)} = p_h^{(k-1)} + \delta p_h$$

At each step $k \geq 0$, this algorithm involves a linear subproblem involving the jacobian F' that is defined by its variationnal form:

$$\begin{aligned} &\langle F'(\mathbf{u}_h^{(k-1)}, p_h^{(k-1)}) . (\delta \mathbf{u}_h, \delta p_h), (\mathbf{v}_h, q_h) \rangle \\ &= \left(\begin{aligned} &Re (t_h(\delta \mathbf{u}_h; \mathbf{u}_h, \mathbf{v}_h) + t_h(\mathbf{u}_h; \delta \mathbf{u}_h, \mathbf{v}_h)) + a_h(\delta \mathbf{u}_h, \mathbf{v}_h) + b_h(\mathbf{v}_h, \delta p_h) \\ &b_h(\delta \mathbf{u}_h, q_h) - c_h(\delta p_h, q_h) \end{aligned} \right) \end{aligned}$$

Example file 3.8: navier_stokes_taylor_newton_dg.cc

```

1 #include "rheolef.h"
2 using namespace rheolef;
3 using namespace std;
4 #include "taylor.icc"
5 #include "stokes_dirichlet_dg.icc"
6 #include "inertia.icc"
7 #include "navier_stokes_dg.h"
8 int main(int argc, char**argv) {
9     environment rheolef (argc, argv);
10    Float eps = numeric_limits<Float>::epsilon();
11    geo omega (argv[1]);
12    string approx = (argc > 2) ? argv[2] : "P1d";
13    Float Re = (argc > 3) ? atof(argv[3]) : 100;
14    Float tol = (argc > 4) ? atof(argv[4]) : eps;
15    size_t max_iter = (argc > 5) ? atoi(argv[5]) : 100;
16    string restart = (argc > 6) ? argv[6] : "";
17    navier_stokes_dg F (Re, omega, approx);
18    navier_stokes_dg::value_type xh = F.initial (restart);
19    int status = damped_newton (F, xh, tol, max_iter, &derr);
20    dout << catchmark("Re") << Re << endl
21         << catchmark("u") << xh[0]
22         << catchmark("p") << xh[1];
23    return status;
24 }

```

Comments

The implementation of the Newton method follows the generic approach introduced in volume 1, section 8.3, page 132. For that purpose we define a class `navier_stokes_dg`.

Example file 3.9: navier_stokes_dg.h

```

1 struct navier_stokes_dg {
2     typedef valarray<field> value_type;
3     typedef Float float_type;
4     navier_stokes_dg (Float Re, const geo& omega, string approx);
5     value_type initial (string restart) const;
6     value_type residue (const value_type& uh) const;
7     void update_derivative (const value_type& uh) const;
8     value_type derivative_solve (const value_type& mrh) const;
9     value_type derivative_trans_mult (const value_type& mrh) const;
10    Float space_norm (const value_type& uh) const;
11    Float dual_space_norm (const value_type& mrh) const;
12    Float Re;
13    space Xh, Qh;
14    quadrature_option_type qopt;
15    form a0, b, c, mu, mp;
16    field lh0, lh, kh;
17    solver smu, smp;
18    mutable form a1;
19    mutable solver_abtb stokes1;
20 };
21 #include "navier_stokes_dg1.icc"
22 #include "navier_stokes_dg2.icc"

```

The member functions of the class are defined in two separate files.

Example file 3.10: navier_stokes_dg1.icc

```

1  navier_stokes_dg::navier_stokes_dg (
2    Float Re1, const geo& omega, string approx)
3    : Re(Re1), Xh(), Qh(), qopt(), a0(), b(), c(), mu(), mp(), lh0(), lh(), kh(),
4      smu(), smp(), a1(), stokes1()
5  {
6    Xh = space (omega, approx, "vector");
7    Qh = space (omega, approx);
8    qopt.set_family(quadrature_option_type::gauss);
9    qopt.set_order(2*Xh.degree()+1);
10   stokes_dirichlet_dg (Xh, Qh, a0, b, c, mp, lh0, kh, qopt);
11   trial u (Xh); test v (Xh);
12   lh = lh0 + Re*inertia_fix_rhs (v, qopt);
13   mu = integrate (dot(u,v), qopt);
14   smu = solver(mu.uu());
15   smp = solver(mp.uu());
16 }
17 navier_stokes_dg::value_type
18 navier_stokes_dg::initial (string restart) const {
19   value_type xh(2);
20   xh[0] = field (Xh, 0);
21   xh[1] = field (Qh, 0);
22   Float Re0 = 0;
23   if (restart == "") {
24     solver_abtb stokes0 (a0.uu(), b.uu(), c.uu(), mp.uu());
25     stokes0.solve (lh0.u(), kh.u(), xh[0].set_u(), xh[1].set_u());
26   } else {
27     idiststream in (restart);
28     in >> catchmark("Re") >> Re0
29         >> catchmark("u") >> xh[0]
30         >> catchmark("p") >> xh[1];
31     check_macro (xh[1].get_space() == Qh, "unexpected " << xh[0].get_space().stamp()
32               << " approximation in file \"" << restart << "\" (" << Xh.stamp() << " expected)");
33   }
34   derr << "# continuation: from Re=" << Re0 << " to " << Re << endl;
35   return xh;
36 }
37 navier_stokes_dg::value_type
38 navier_stokes_dg::residue (const value_type& xh) const {
39   trial u (Xh); test v (Xh);
40   form a = a0 + Re*inertia(xh[0], u, v, qopt);
41   value_type mrh(2);
42   mrh[0] = a*xh[0] + b.trans_mult(xh[1]) - lh;
43   mrh[1] = b*xh[0] - c*xh[1] - kh;
44   return mrh;
45 }
46 void navier_stokes_dg::update_derivative (const value_type& xh) const {
47   trial u (Xh); test v (Xh);
48   a1 = a0 + Re*(inertia(xh[0], u, v, qopt) + inertia(u, xh[0], v, qopt));
49   stokes1 = solver_abtb (a1.uu(), b.uu(), c.uu(), mp.uu());
50 }
51 navier_stokes_dg::value_type
52 navier_stokes_dg::derivative_solve (const value_type& mrh) const {
53   value_type delta_xh(2);
54   delta_xh[0] = field (Xh, 0);
55   delta_xh[1] = field (Qh, 0);
56   stokes1.solve (mrh[0].u(), mrh[1].u(),
57                 delta_xh[0].set_u(), delta_xh[1].set_u());
58   return delta_xh;
59 }
60 navier_stokes_dg::value_type
61 navier_stokes_dg::derivative_trans_mult (const value_type& mrh) const {
62   value_type rh(2);
63   rh[0] = field (Xh);
64   rh[1] = field (Qh);
65   rh[0].set_u() = smu.solve(mrh[0].u());
66   rh[1].set_u() = smp.solve(mrh[1].u());
67   value_type mgh(2);
68   mgh[0] = a1.trans_mult(rh[0]) + b.trans_mult(rh[1]);
69   mgh[1] = b*rh[0] - c*rh[1];
70   return mgh;
71 }

```

Example file 3.11: navier_stokes_dg2.icc

```

1 Float navier_stokes_dg::space_norm (const value_type& xh) const {
2   return sqrt (mu(xh[0],xh[0]) + mp(xh[1],xh[1]));
3 }
4 Float navier_stokes_dg::dual_space_norm (const value_type& mrh) const {
5   value_type rh(2);
6   rh[0] = field (Xh,0);
7   rh[1] = field (Qh,0);
8   rh[0].set_u() = smu.solve(mrh[0].u());
9   rh[1].set_u() = smp.solve(mrh[1].u());
10  return sqrt (dual(rh[0],mrh[0]) + dual(rh[1],mrh[1]));
11 }

```

```

make navier_stokes_taylor_newton_dg navier_stokes_taylor_error_dg
./navier_stokes_taylor_newton_dg square P2d 1000 | ./navier_stokes_taylor_error_dg

```

3.3.5 Application to the driven cavity benchmark

Example file 3.12: cavity_dg.icc

```

1 struct g {
2   point operator() (const point& x) const {
3     return point((abs(1-x[1]) < 1e-7) ? 1 : 0, 0, 0); }
4 };
5 struct f {
6   point operator() (const point& x) const { return point(0,0,0); }
7 };

```

The program `navier_stokes_cavity_newton_dg.cc` is obtained by replacing in `navier_stokes_taylor_newton_dg.cc` the include `taylor.icc` by `cavity_dg.icc` that defines the boundary conditions. The compilation and run are similar.

```

make navier_stokes_cavity_newton_dg streamf_cavity
./navier_stokes_cavity_newton_dg square P1d 500 > square.field
field -proj square.field -field | ./streamf_cavity | \
    field -bw -n-iso-negative 10 -

```

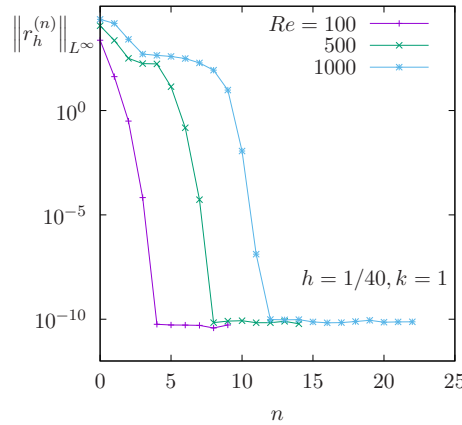


Figure 3.1: The discontinuous Galerkin method for the Navier-Stokes problem on the driven cavity benchmark when $k = 1$ and $d = 2$: convergence of the damped Newton algorithm.

3.3.6 Upwinding

The skew symmetry property is generalized to the requirement that t_h be non-dissipative (see [11, p. 282], eqn (6.68)):

$$t_h(\mathbf{w}_h; \mathbf{u}_h, \mathbf{u}_h) \geq 0, \forall \mathbf{w}_h, \mathbf{u}_h \in \mathbf{X}_h$$

A way to satisfy this property is to add an *upwinding* term in t_h :

$$\check{t}_h(\mathbf{w}_h; \mathbf{u}_h, \mathbf{v}_h) := t_h(\mathbf{w}_h; \mathbf{u}_h, \mathbf{v}_h) + s_h(\mathbf{w}_h; \mathbf{u}_h, \mathbf{v}_h)$$

with

$$s_h(\mathbf{w}_h; \mathbf{u}_h, \mathbf{v}_h) = \frac{1}{2} \sum_{S \in \mathcal{S}_h^{(i)}} \int_S |\llbracket \mathbf{w}_h \rrbracket \cdot \mathbf{n}| (\llbracket \mathbf{u}_h \rrbracket \cdot \llbracket \mathbf{v}_h \rrbracket) \, ds$$

We aim at using a Newton method. We replace t_h by its extension \check{t}_h containing the upwind terms in the definition of F , and then we compute its jacobian F' . As the absolute value is not differentiable, the functions s_h , \check{t}_h and then F are also not differentiable with respect to \mathbf{w}_h . Nevertheless, the absolute value is convex and we can use some concets of the sudifferential calculus. Let us introduce the multi-valued sign function:

$$\text{sgn}(x) = \begin{cases} \{1\} & \text{when } x > 0 \\ [-1, 1] & \text{when } x = 0 \\ \{-1\} & \text{when } x < 0 \end{cases}$$

Then, the subdifferential of the absolute value function is $\text{sgn}(x)$ and for all $\delta \mathbf{w}_h, \mathbf{w}_h, \mathbf{u}_h, \mathbf{v}_h \in \mathbf{X}_h$, we define a generalization of the partial derivative as

$$\frac{\partial s_h}{\partial \mathbf{w}_h}(\mathbf{w}_h; \mathbf{u}_h, \mathbf{v}_h) \cdot (\delta \mathbf{w}_h) = \frac{1}{2} \sum_{S \in \mathcal{S}_h^{(i)}} \int_S \text{sgn}(\llbracket \mathbf{w}_h \rrbracket \cdot \mathbf{n}) (\llbracket \delta \mathbf{w}_h \rrbracket \cdot \mathbf{n}) (\llbracket \mathbf{u}_h \rrbracket \cdot \llbracket \mathbf{v}_h \rrbracket) \, ds$$

Example file 3.13: inertia_upw.icc

```

1 #include "sgn.icc"
2 form inertia_upw (field w, trial u, test v,
3   quadrature_option_type qopt = quadrature_option_type())
4 {
5   return integrate ("internal_sides",
6     0.5*abs(dot(average(w), normal()))*dot(jump(u), jump(v)));
7 }
8 form d_inertia_upw (field w, trial dw, field u, test v,
9   quadrature_option_type qopt = quadrature_option_type())
10 {
11   return integrate ("internal_sides",
12     0.5*compose (sgn, dot(average(w), normal()))
13     *dot(average(dw), normal())*dot(jump(u), jump(v)));
14 }
```

A multi-valued jacobian F' is then defined:

$$\begin{aligned}
& \langle F' \left(\mathbf{u}_h^{(k-1)}, p_h^{(k-1)} \right) \cdot (\delta \mathbf{u}_h, \delta p_h), (\mathbf{v}_h, q_h) \rangle \\
&= \text{Re} \left(\begin{aligned} & t_h(\delta \mathbf{u}_h; \mathbf{u}_h, \mathbf{v}_h) + t_h(\mathbf{u}_h; \delta \mathbf{u}_h, \mathbf{v}_h) + \frac{\partial s_h}{\partial \mathbf{w}_h}(\mathbf{u}_h; \mathbf{u}_h, \mathbf{v}_h) \cdot (\delta \mathbf{u}_h) + s_h(\mathbf{u}_h; \delta \mathbf{u}_h, \mathbf{v}_h) \\ & 0 \end{aligned} \right) \\
&+ \begin{pmatrix} a_h(\delta \mathbf{u}_h, \mathbf{v}_h) & + & b_h(\mathbf{v}_h, \delta p_h) \\ b_h(\delta \mathbf{u}_h, q_h) & - & c_h(\delta p_h, q_h) \end{pmatrix}
\end{aligned}$$

We are able to extend the Newton method to the F function that allows a multi-valued subdifferential F' . During iterations, we can choose any of the available directions in the subdifferential. One the possibilities is then to replace the multi-valued sign function by a single-value one:

$$\widetilde{\text{sgn}}(x) = \begin{cases} 1 & \text{when } x \geq 0 \\ -1 & \text{when } x < 0 \end{cases}$$

Example file 3.14: `sgn.icc`

```
1 Float sgn (Float x) { return (x >= 0) ? 1 : -1; }
```

Example file 3.15: `navier_stokes_upw_dg.h`

```
1 #include "navier_stokes_dg.h"
2 struct navier_stokes_upw_dg: navier_stokes_dg {
3     typedef valarray<field> value_type;
4     typedef Float float_type;
5     navier_stokes_upw_dg (Float Re, const geo& omega, string approx);
6     value_type residue (const value_type& uh) const;
7     void update_derivative (const value_type& uh) const;
8 };
9 #include "navier_stokes_upw_dg.icc"
```

Example file 3.16: `navier_stokes_upw_dg.icc`

```
1 #include "inertia_upw.icc"
2 navier_stokes_upw_dg::navier_stokes_upw_dg (
3     Float Re1, const geo& omega, string approx)
4     : navier_stokes_dg (Re1, omega, approx) {}
5
6 navier_stokes_upw_dg::value_type
7 navier_stokes_upw_dg::residue (const value_type& xh) const {
8     trial u (Xh); test v (Xh);
9     form a = a0 + Re*( inertia (xh[0], u, v, qopt)
10                        + inertia_upw (xh[0], u, v, qopt));
11     value_type mrh(2);
12     mrh[0] = a*xh[0] + b.trans_mult(xh[1]) - lh;
13     mrh[1] = b*xh[0] - c*xh[1] - kh;
14     return mrh;
15 }
16 void navier_stokes_upw_dg::update_derivative (const value_type& xh) const {
17     trial du (Xh); test v (Xh);
18     a1 = a0 + Re*( inertia (xh[0], du, v, qopt)
19                  + inertia_upw (xh[0], du, v, qopt)
20                  + inertia (du, xh[0], v, qopt)
21                  + d_inertia_upw (xh[0], du, xh[0], v, qopt));
22     stokes1 = solver_abtb (a1.uu(), b.uu(), c.uu(), mp.uu());
23 }
```

The program `navier_stokes_cavity_newton_upw_dg.cc` is obtained by replacing in `navier_stokes_taylor_newton_dg.cc` the string `navier_stokes_dg` by `navier_stokes_upw_dg` (two occurrences: in the includes and then in the definition of F). Also replace the include `taylor.icc` by `cavity_dg.icc` that defines the boundary conditions. The compilation and run are similar.

```
make navier_stokes_cavity_newton_upw_dg stream_cavity
mkgeo_grid -t 80 > square.geo
./navier_stokes_cavity_newton_upw_dg square P1d 500 1e-15 100 > square-500.field
field -proj square-500.field -field | ./streamf_cavity | \
    field -bw -n-iso 30 -n-iso-negative 20 -
```

Computations for higher Renolds numbers are performed by continuation, starting from a previous computation at lower Re :

```
./navier_stokes_cavity_newton_upw_dg square P1d 1000 1e-15 100 square-500.field > square-1000.fi
```

```
./navier_stokes_cavity_newton_upw_dg square P1d 1500 1e-15 100 square-1000.field > square-1500.fi
```

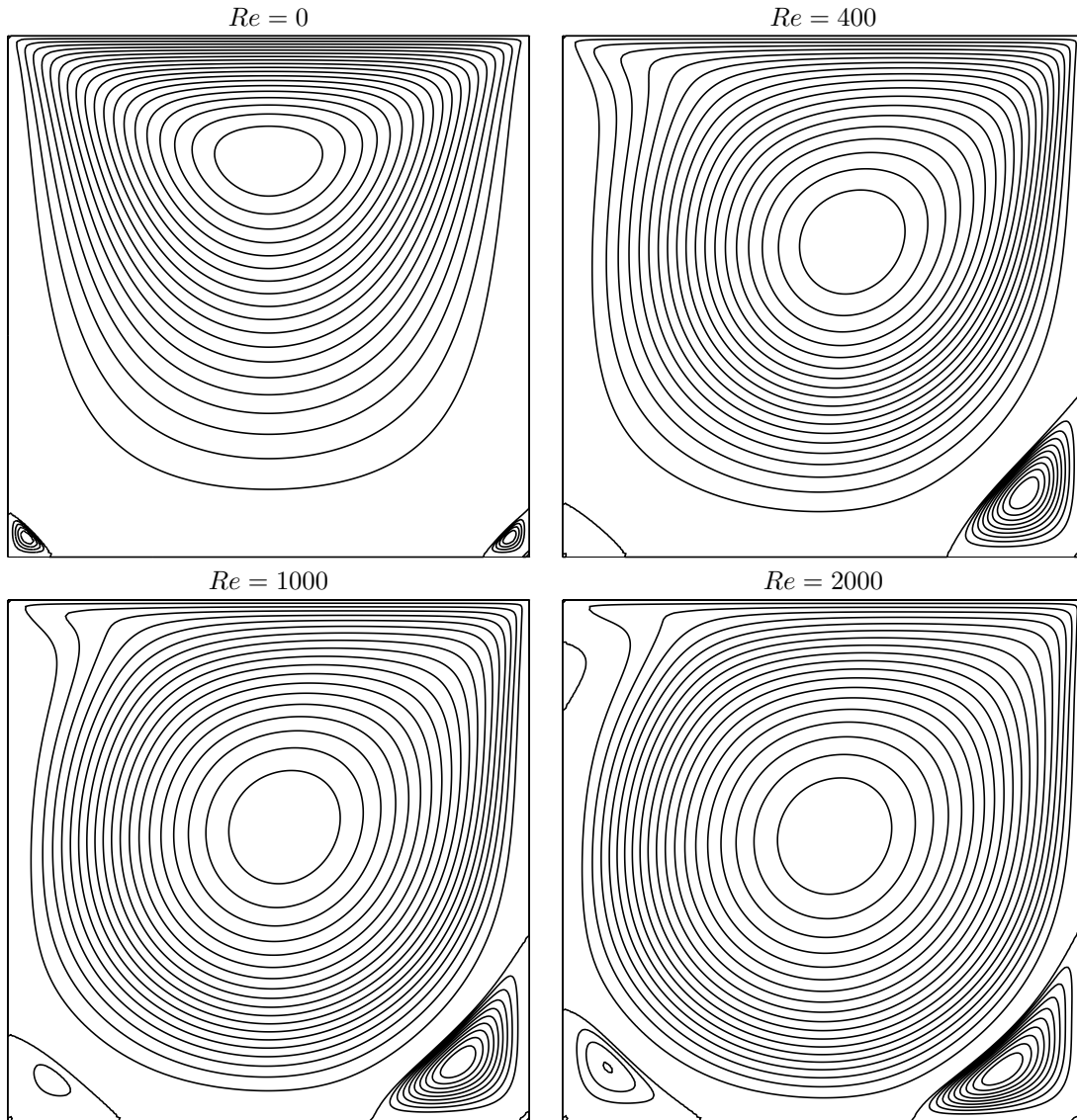


Figure 3.2: The discontinuous Galerkin method for the Navier-Stokes problem on the driven cavity benchmark when $k = 1$ (80×80 grid): stream function isovalues for various Re .

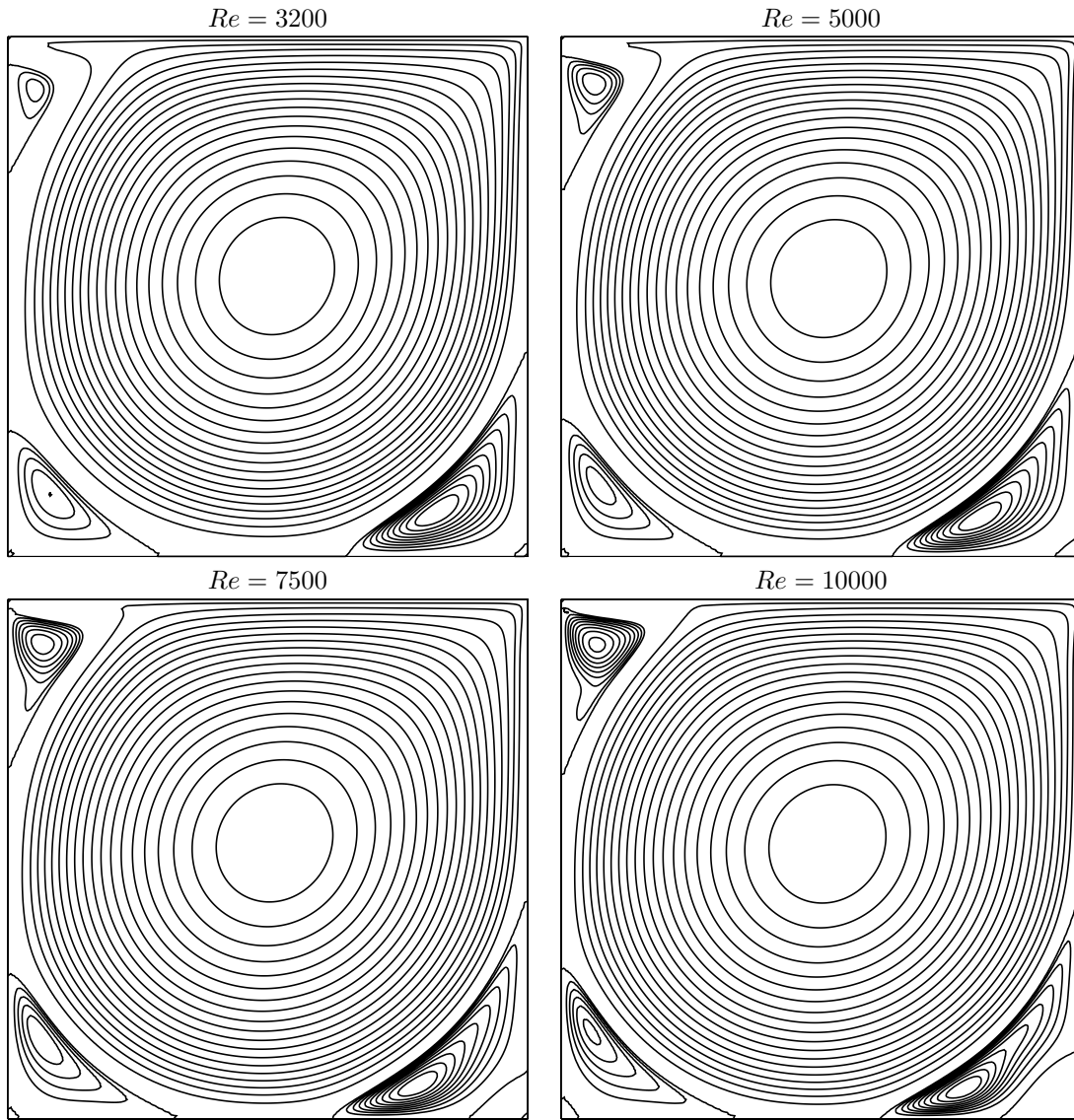


Figure 3.3: The discontinuous Galerkin method for the Navier-Stokes problem on the driven cavity benchmark when $k = 1$ (80×80 grid): stream function isovalues for various Re (cont.).

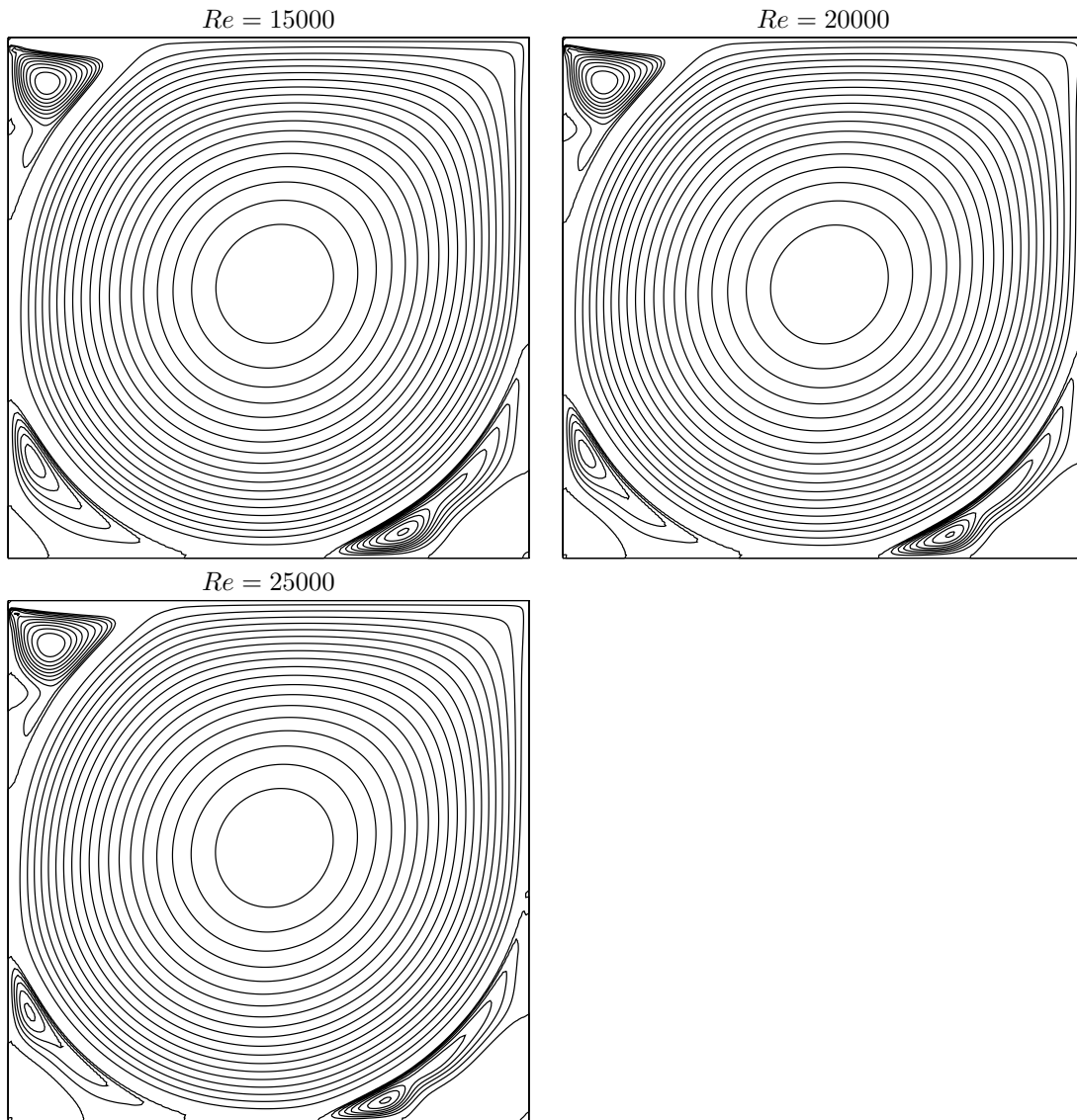


Figure 3.4: The discontinuous Galerkin method for the Navier-Stokes problem on the driven cavity benchmark when $k = 1$ (80×80 grid): stream function isovalues for various Re (cont.).

Part III

Technical appendices

Appendix A

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