# Asynchronous OpenCL/MPI Discontinous Galerkin Solver for Conservation laws

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#### Outline

- ▶ The DG method
  - Macrocell/subcell formulation
  - Hexahedral elements
  - ► Gauss-Legendre vs. Gauss-Lobatto points
- Implementation
  - ► The OpenCL programming language
  - ► Performance analysis
- Example simulation results

## **Evolution Equations**

We consider the general hyperbolic equation

$$\partial_t w + \sum_{k=1}^{k=d} \partial_k F^k(w) = S, \tag{1}$$

in d = dimensions. F is the flux and S the source term.

#### Examples:

- Navier–Stokes equations
- Maxwell's equations
- ► MHD
- Vlasov equations

We would like to numerically solve such equations in complex geometries with as general boundary conditions as possible.

#### Discontinuous Galerkin Method

The physical domain is divided into cells.

In each cell L, w is projected onto a finite set of basis functions  $\psi_i^L(x)$ :

$$w(x,t) \approx \sum_{i \in L} w_L^i(t) \psi_i^L(x). \tag{2}$$

The evolution equation is approximated by

$$\int_{L} \partial_{t} w \psi_{i}^{L} - \int_{L} F(w, w, \nabla \psi_{i}^{L}) + \int_{\partial L} F(w_{L}, w_{R}, \boldsymbol{n}_{LR}) \psi_{i}^{L} = S_{i}^{L},$$
(3)

where  $n_{LR}$  is the normal vector from cell L to cell R. The DG formulation is good for conserving invariants. Elements can be curved and meshes can be non-conformal.

## Macrocell/subcell

The physical domain is divided into an unstructured mesh of macrocells.

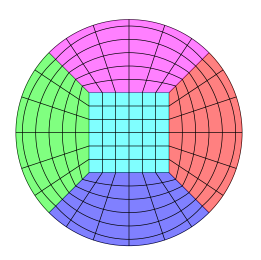
The macrocells are divided into subcells.

- ▶ The curvature is constant within a macrocell.
- ► The subcells are arranged in a Cartesian grid in the reference space.

#### Advantages:

- ▶ We are able consider complex geometries.
- Memory access is coalescent when looping over subcells in a macrocell.
- Macrocells provide coarse-grained parallelism.

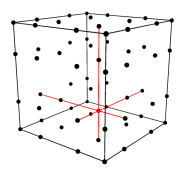
## ${\sf Macrocell/subcell}$



#### Hexahedral elements

We make use of hexahedral elements.

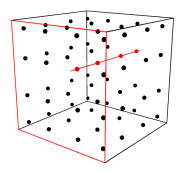
- ▶ Memory access is coalescent within an element.
- ▶ The volumic term is non-zero on a subdimensional set.



## Gauss Legendre Quadrature

Gauss Legendre collocation points gives convergence

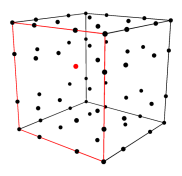
- ▶ degree n gives  $\mathcal{O}(dx^{n+1})$  spatial convergence,
- ▶ but the flux terms are expensive.



## Gauss Lobatto Quadrature

Gauss Lobatto collocation points gives convergence

- ▶ degree n gives  $\mathcal{O}(dx^n)$  spatial convergence,
- ▶ but the flux terms are cheaper.



## Source-term computation

The computation of the source term is divided into the following steps:

- Volume terms
- Subcell interface flux terms
- Macrocell interface flux terms
- Mass division
- Computation of source terms

We time-step using either RK2 or RK4.

### **Implementation**

We use OpenCL for our implementation.

- ► OpenCL is a C-like programming language.
- ▶ It works on CPUs, GPUs, and MICs.
- ► No restriction to a specific vendor (unlike CUDA).

GPUs and MICs are interesting computing platforms:

- ▶ Many cores (fine-grained parallelism).
- ► Fast memory access on board (less bottleneck).
- CPU-to-GPU transfers are slow (so stay on the board if possible).

## Implementation

We would like to use multiple GPUs.

- ► Macrocells allow for coarse-grained parallelism
- Inter-macrocell flux computation doesn't need a lot of bandwidth.
- ▶ We can use MPI between nodes with different macrocells.

The StarPU runtime environment is an interesting possibility for managing data transfers and a heterogeneous computing environment.

## Performance analysis

How do we compare performance between platforms?

Roofline: Count FLOPs, count i/o, compare with manufacturer specs.

#### Questions:

- 1. is a \* x + b two FLOPs or one FLOP?
- 2. Do multiplication, division, and addition count equally?
- 3. What about integer operations?
- 4. Is the listed speed a specification or an advertisement?

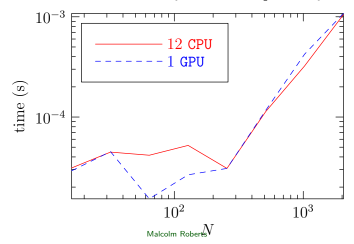
#### Comparison is difficult

► CPU manufacturers do not publish GFLOP/s We get around 200 GFLOP/s on a 1TFLOP/s GPU.

## Comparison of CPU and GPU performance

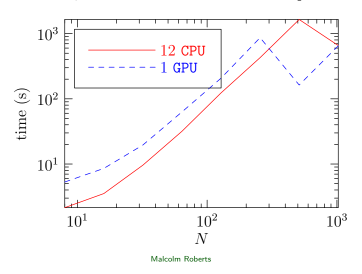
We can compare platforms by comparing execution speed.

We use clffT, an FFT library written in OpenCL by AMD.



## Comparison of CPU and GPU performance

#### We perform the same test with schnaps



## Comparison of CPU and GPU performance

#### schnaps:

- ► Fully uses all 12 CPUs.
- ▶ Is faster than our C implementation.
- ► Has similar performance on 12 CPU or on GPU.

So we claim that schnaps is well-optimized for both for CPUs and GPUs

#### Who's on the MIC?

MIC: Many integrated cores, Intel's Xeon Phi.

- ▶ A combination of a GPU and a CPU
- ► Same advertised performance as Tahiti GPU we're using.
- ▶ Works with OpenCL.

Requires optimization:

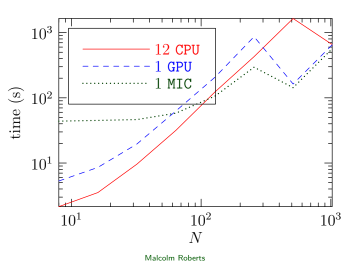
"There is no free lunch."

Evaluating kernels on Xeon Phi to accelerate Gysela application by G Latu, M Haefele, J Bigot, V Grandgirard, T Cartier-Michaud, F Rozar

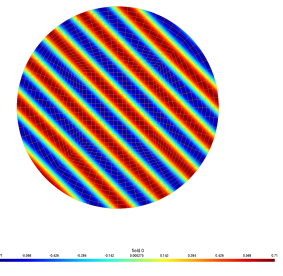
"There may be no lunch even if you pay."

#### Who's on the MIC?

schnaps works well on the Xeon Phi.



## Example simulation: Maxwell's equations



#### Conclusion

schnaps is a new general-purpose DG implementation in OpenCL.

- ▶ For the DG method we use:
  - An unstructured mesh of macrocells.
  - ▶ A structured mesh of subcells within each macrocell.
  - hexahedral elements.
  - Gauss-Lobatto collocation points.
- ▶ We can run on CPUs, GPUs, and MICs
  - and we can do so efficiently.
  - Performance metric is based on comparison with established libraries.

schnaps is available at schnaps.gforge.inria.fr
This talk is available at malcolmiwroberts.com

Thank you for your attention!

Merci pour votre attention!