Sailfish: a flexible multi-GPU implementation of the lattice Boltzmann method

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Abstract

We present Sailfish, an open source fluid simulation package implementing the lattice Boltzmann method (LBM) on modern Graphics Processing Units (GPUs) using CUDA/OpenCL. We take a novel approach to GPU code implementation and use run-time code generation techniques and a high level programming language (Python) to achieve state of the art performance, while allowing easy experimentation with different LBM models and tuning for various types of hardware. We discuss the general design principles of the code, scaling to multiple GPUs in a distributed environment, as well as the GPU implementation and optimization of many different LBM models, both single component (BGK, MRT, ELBM) and multicomponent (Shan-Chen, free energy). The paper also presents results of performance benchmarks spanning the last three NVIDIA GPU generations (Tesla, Fermi, Kepler), which we hope will be useful for researchers working with this type of hardware and similar codes.

Keywords: lattice Boltzmann, LBM, computational fluid dynamics, graphics processing unit, GPU, CUDA

PROGRAM SUMMARY

Manuscript Title: Sailfish: a flexible multi-GPU implementation of the lattice Boltzmann method
Authors: Michal Januszewski, Marcin Kostur
Program Title: Sailfish
Code Repository: https://github.com/sailfish-team/sailfish
Journal Reference:
Catalogue identifier:
Licensing provisions: LGPLv3
Programming language: Python, CUDA C, OpenCL
Computer: any with an OpenCL or CUDA-compliant GPU

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Operating system: no limits (tested on Linux and Mac OS X)
RAM: Hundreds of megabytes to tens of gigabytes for typical cases.
Keywords: lattice Boltzmann, LBM, CUDA, OpenCL, GPU, computational fluid dynamics, Python.
Classification: 12, 6.5
External routines/libraries: PyCUDA/PyOpenCL, Numpy, Mako, ZeroMQ (for multi-GPU simulations)
Nature of problem:
GPU-accelerated simulation of single- and multi-component fluid flows.
Solution method:
A wide range of relaxation models (LBGK, MRT, regularized LB, ELBM, Shan-Chen, free energy, free surface) and boundary conditions within the lattice Boltzmann method framework. Simulations can be run in single or double precision using one or more GPUs.
Restrictions:
The lattice Boltzmann method works for low Mach number flows only.
Unusual features:
The actual numerical calculations run exclusively on GPUs. The numerical code is built dynamically at run-time in CUDA C or OpenCL, using templates and symbolic formulas. The high-level control of the simulation is maintained by a Python process.
Running time: problem-dependent, typically minutes (for small cases or short simulations) to hours (large cases or long simulations)

1. Introduction

Merely a few years ago the computational power on the order of TFLOPS was an attribute of a supercomputer. Today, the latest commodity Graphics Processing Units (GPU) are capable of more than 5 TFLOPS and have much lower energy requirements than a set of Central Processing Units (CPUs) of comparable performance – a result achieved by employing a massively parallel architecture with hundreds of cores on a single device.

In order to take advantage of the parallel hardware, appropriate algorithms have to be developed, and this generally needs to be done on a problem-specific basis. Ideally, the problem naturally decomposes into a large number of (semi-)independent tasks whose results can be combined in a simple way, as in e.g. numerical Monte-Carlo solution of a stochastic differential equation [1], parameter space studies of dynamical systems, or event simulation and reconstruction in particle physics [2].

In recent years, the lattice Boltzmann method (LBM) emerged as an interesting alternative to more established methods for fluid flow simulations. Originally developed as an extension of lattice gas automata, nowadays LBM stands on well-established theoretical foundations and serves as a method of choice for many researchers in various fields [3 4], while still retaining its relative simplicity. Perhaps its most important feature in practice is its suitability
for parallel architectures – the algorithm is executed on a regular lattice and
typically only interactions between nearest neighbor nodes are necessary.

This paper discusses various issues related to the implementation of the LBM on GPUs, and presents a concrete and flexible solution, taking a new overall approach to the problem of LB software design. Our Sailfish code has been under development since April 2009 and is publicly available under an open source license. With support for both single and binary fluid simulations, a wide variety of boundary conditions, and calculations on both a single GPU and multiple GPUs, it is, to the best of our knowledge, the most comprehensive open source LBM code for GPUs.

The text is organized as follows. First, a short overview of the discussed lattice Boltzmann models is presented. Then, the primary design principles of the Sailfish project are laid out, and various LB GPU implementations are compared. This is followed by performance measurements on diverse GPU hardware, both on single machines and in clusters. In the next section, the code is validated on 4 standard test cases. The final section concludes the paper and provides some directions for future work.

2. Lattice Boltzmann methods overview

In this section, all models implemented in Sailfish as of September 2013 will be briefly overviewed. We start with the basic concepts and nomenclature of LBM. We then describe single fluid models: single-relaxation time LBGK and regularized dynamics, multiple relaxation times and the entropic LBM, and multi-component models: the Shan-Chen model and the free energy model. Finally we give a short overview of our implementation of body forces and various boundary conditions. We refer the reader to the reviews [3, 4] for a more detailed discussion of the LBM.

In an LB simulation space is discretized into a regular Cartesian grid, where each node represents a small pocket of fluid. The state of the fluid is encoded by a particle distribution function $f_i$ where $i$ spans a set of discrete velocity vectors $\{ \vec{e}_i \}$ indicating the allowed directions of mass movement between the nodes of the lattice. $f_i$ are often also called mass fractions. LB lattices are typically named using the DxQy scheme, where $x$ indicates the dimensionality of the lattice, and $y$ is the number of discrete velocity vectors. Figure 1 shows some common LB lattices.

Macroscopic fluid fields, such as density ($\rho$) or velocity ($\vec{u}$) are defined as moments of the distribution function:

\[
\rho = \sum_i f_i, \quad \rho \vec{u} = \sum_i f_i \vec{e}_i
\]  

(1)

In the simplest case the system dynamics is described by:

\[
f_i(\vec{x} + \vec{e}_i, t + 1) - f_i(\vec{x}, t) = \frac{1}{\tau} (f_i - f_i^{eq})(\vec{x}, t)
\]  

(2)
where \( \{ f_i^\text{eq}(\vec{x}, t) \} \) is a set of equilibrium distributions which are functions of the macroscopic fields \((\rho, \vec{u})\) at node \( \vec{x} \) at time \( t \), \( \tau \) is a relaxation time related to the kinematic viscosity \( \nu \) via \( \tau = (1 + 6\nu)/2 \), and where the positions and time are expressed in the lattice unit system, i.e. the time unit represents a single simulation step and the lattice nodes are separated by 1 distance unit along the primary axes. This LB model is often referred to as Lattice Bhatnagar-Gross-Krook (LBGK), which owes its name to the BGK operator from kinetic theory \[6\], and by which the collision operator on the right hand side of (2) is inspired.

When \( f_i^\text{eq} \) takes the form:

\[
    f_i^\text{eq} = \rho w_i \left( 1 + 3 \vec{e}_i \cdot \vec{u} + \frac{9}{2} (\vec{e}_i \cdot \vec{u})^2 - \frac{3}{2} u^2 \right)
\]  

(3)

where \( \{ w_i \} \) is a set of weights (see Table 1), the Navier-Stokes equations can be recovered from (2) in the limit of low Mach numbers through Chapman-Enskog expansion \[3\] or expansion in a Hermite basis \[7\]. It can be shown that (2) is second-order accurate in space and time in the bulk of the fluid \[3\].

It should be noted that (1) and the right hand side of (2) are fully local, i.e. only information from node \( \vec{x} \) is necessary to calculate the system state at
the next step of the simulation. When data is exchanged between nodes, the process is limited to the nearest neighbors (left hand side of 2). This makes the LB method ideally suited for implementation on massively parallel computer architectures.

Conceptually, 2 is often decomposed into two steps: relaxation (evaluation of the collision operator, right hand side) and propagation (movement of mass fractions to neighboring nodes, left hand side). While such a decomposition is often suboptimal from the point of view of a software implementation, it still provides a useful mental framework to discuss LB algorithms, and we will use these two names throughout the text.

Many extended LB models have been proposed to (a) improve numerical stability, or (b) simulate more complex systems. A comprehensive discussion of such models would be prohibitively long, so here we limit ourselves to a short overview of models that are currently implemented within the Sailfish framework. In the first group, we discuss the multiple-relaxation times (MRT) model, the entropic LB model (ELBM) and the Smagorinsky large eddy model. In the second group, Sailfish implements two multicomponent fluid models: the Shan-Chen model [8], and the free energy model [9].

2.1. Single fluid models with enhanced stability

One practical problem with the LBGK approximation is the limited range of viscosities at which the model remains stable, which directly translates into a limited range of Reynolds numbers Re = Lu/ν at which fluid flows can be simulated (L being the spatial extent of the domain expressed as a number of nodes). With u limited to about 0.05 by the low Mach number requirement for most LB models where the speed of sound c_s = 1/√3 and the stable range of numerical viscosities at these speeds being about (10^{-3}, 1/6), the highest attainable Reynolds number assuming L = 300 is on the order of Re = 15000. In practice, the range of Reynolds numbers and viscosities is often even narrower due to instabilities introduced by complex geometry or boundary conditions. To address this problem, a number of models exhibiting enhanced numerical stability have been developed.

2.1.1. Regularized LBM

Perhaps the simplest modification of the LBGK model aiming at improving its stability is the regularized model [10]. The basic idea of this approach is to replace the nonequilibrium part of the distributions \( f_{\text{neq}} = f_i - f_{\text{eq}}^i \) with a new first order regularized value \( f_{\text{reg}} \) derived from the non-equilibrium stress tensor \( \Pi_{\text{neq}}^{\alpha\beta} = \sum_i f_{\text{neq}}^i (e_{i\alpha} e_{i\beta} - \delta_{\alpha\beta} c_s^2) \) as \( f_{\text{reg}} = \frac{\mu_i}{2c_s^4} (e_{i\alpha} e_{i\beta} - c_s^2 \delta_{\alpha\beta}) \Pi_{\text{neq}}^{\alpha\beta} \). After this regularization step, the collision and propagation proceed as in the standard LBGK scheme.

2.1.2. Multiple relaxation times

If the particle distributions \( f_i \) are considered to form vectors \( |f_i| \), the collision operator in the LBGK approximation (2) can be interpreted as a diagonal matrix
$S = \omega I$, with $\omega = 1/\tau$ acting on the vector difference $|f^{eq}(\vec{x}, t)| - |f(\vec{x}, t)|$. The basis of the distribution vector can be changed using an invertible matrix $M$: $|m\rangle = M |f\rangle$. In particular, the matrix $M$ can be chosen such that $|m\rangle$ is a vector of moments of the particle distributions $f_i$.

The idea of MRT is to perform the collision in the moment basis, and then change the basis to recover the original form of the particle distributions for the streaming step [11]. The MRT collision operator can be written as:

$$M^{-1}\hat{S}(|m^{eq}(\vec{x}, t)) - |m(\vec{x}, t))|$$

where the collision matrix $\hat{S} = MSM^{-1}$ is diagonal with its elements $s_i = 1/\tau_i$ representing the inverses of relaxation times for individual moments. Collision matrix elements corresponding to conserved moments, such as density or momentum are set to 0. Some of the remaining elements are related to kinematic and bulk viscosities, but the other ones can be tuned to increase the numerical stability of the simulation without changing its physical results. We refer the reader to the paper of D’Humieres et al. [11] for a detailed discussion of optimal choices for relaxation times, and the form of the matrix $M$.

2.1.3. Entropic LBM

In the entropic LB model, a discrete H-function $H(f) = \sum_i f_i \ln(f_i/w_i)$ is defined and the equilibrium distribution function is defined as the extremum of $H(f)$ under constraints of mass and momentum conservation:

$$f_i^{eq} = \rho w_i \prod_{\alpha=1}^{D} \left( 2 - \sqrt{1 + u_i^2} \right) \left( \frac{2u_{\alpha} + \sqrt{1 + 3u_i^2}}{1 - u_{\alpha}} \right) e^{\epsilon_{i\alpha}}$$

where $D$ is the dimensionality of the lattice. The relaxation process is also modified to include a new dynamically adjusted parameter $\alpha$:

$$f_i(\vec{x} + \vec{e}_i, t + 1) - f_i(\vec{x}, t) = \omega_0 \frac{\alpha}{2} (f_i - f_i^{eq})(\vec{x}, t)$$

with $\omega_0 = 1/\tau$. $\alpha$ is evaluated at every simulation step as the solution of the $H$-function monotonicity constraint:

$$H(f) = H(f - \alpha(f - f^{eq}))$$

This procedure guarantees unconditional stability of the numerical scheme, as the $H(f)$ function can be proven to be a Lyapunov function for the system. When $\alpha = 2$ as is the case when the system is close to equilibrium, [9] has the same form as [2]. The corrections resulting from H-theorem compliance can be both positive and negative, temporarily increasing and decreasing the effective local viscosity of the fluid [12, 13].
2.1.4. Smagorinsky subgrid model

The idea behind the Smagorinsky subgrid model is to locally modify the fluid viscosity by adding an eddy viscosity term dependent on the magnitude of the strain rate tensor $S$:

$$\nu = \nu_0 + \nu_t$$

where $\nu_0$ is the baseline viscosity of the fluid and $\nu_t = \tau_t / 3$ is an eddy viscosity, the form of which depends on the subgrid model used in the simulation. In the Smagorinsky model, the relaxation time $\tau_t$ can be calculated using the momentum flux tensor $Q_{\alpha\beta} = \sum_i e_i e_i (f_i - f_i^{eq})$:

$$\tau_t = \frac{1}{2} \left( \sqrt{\tau_0^2 + 4c_s^{-4}C_s^2(Q_{\alpha\beta}Q_{\alpha\beta})^{1/2}} - \tau_0 \right)$$

where $C_s$ is the Smagorinsky constant, which for LB models can be taken to be 0.1. This effectively abandons the single relaxation time approximation by making it a spatially and temporally varying quantity dependent on the local gradients of the fluid velocity [14].

2.2. Multi-fluid models

The multi-fluid models discussed here are all diffuse interface models, without any explicit interface tracking – the dynamics of the interface emerges naturally from the interactions between the fluid components. All models in this class use an additional lattice (whose mass fractions we will call $g_i$) to represent the second fluid species and nonlocal interactions between the lattices in the collision process.

2.2.1. Shan-Chen

In the Shan-Chen model, both lattices are used to represent fluid components, with the 0-th moment of $f_i$ and $g_i$ representing the density of the first (A) and second fluid components (B), respectively. The equilibrium function and relaxation schemes remain unchanged for both lattices, but an additional coupling term in the form of a body force:

$$\vec{F}_A(x) = G\psi_A(x) \sum_i w_i \psi_B(x + \vec{e}_i)\vec{e}_i$$

is introduced into (2) (see Section 2.4 for a short overview of ways of adding body forces to an LB simulation), where $\psi_B(x)$ is a pseudopotential function, dependent on the density $\rho_B$ at node $x$ and $G$ is a coupling constant. A similar term $\vec{F}_B$ of the form (10) with $\psi_A$ replaced by $\psi_B$ and vice versa is added to the collision operator for the second component.

A commonly used pseudopotential function is $\psi(\rho) = 1 - e^{-\rho}$. The velocity of the fluid becomes a weighted average of the first moments of the distribution functions: $\vec{u} = \frac{\rho_A \vec{u}_A + \rho_B \vec{u}_B}{\rho_A + \rho_B + \rho_0}$, where $\vec{u}_A$, $\vec{u}_B$ and $\rho_A$, $\rho_B$ are velocities and densities computed respectively from $f_i$ and $g_i$ using (1), and $\tau_A$, $\tau_B$ are relaxation times for the two fluid components.
2.2.2. The free energy model

The free energy model is based on a Landau free energy functional for a binary fluid [15]. In the LB realization, the 0-th moment of $f_i$ represents the density $\rho$ of both components, while the 0-th moment of $g_i$ is a smoothly varying order parameter $\phi$, with $-1$ indicating a pure first component and $1$ indicating a pure second component:

$$
\rho = \sum_i f_i \quad \phi = \sum_i g_i \quad \mu = \sum_i e_i f_i.
$$

(11)

The macroscopic dynamics of the fluid is described by the system of equations

$$
\begin{align*}
\partial_t \rho &+ \partial_\alpha (\rho u_\alpha) = 0 \\
\partial_t (\rho u_\alpha) &+ \partial_\alpha (\rho u_\alpha u_\beta) = -\partial_\alpha P_{\alpha\beta} + \partial_\alpha \{\nu \rho (\partial_\beta u_\alpha + \partial_\alpha u_\beta)\} \\
\partial_t \phi &+ \partial_\alpha (\phi u_\alpha) = M \nabla^2 \mu
\end{align*}
$$

(12)

where $M$ is a mobility parameter, $\mu$ is the chemical potential and the pressure tensor $P_{\alpha\beta}$ is defined as:

$$
P_{\alpha\beta} = \left( p_0 - \kappa \phi \nabla^2 \phi - \frac{\kappa}{2} |\nabla \phi|^2 \right) \delta_{\alpha\beta} + \kappa \partial_\alpha \phi \partial_\beta \phi,
$$

(13)

and $p_0 = \rho/3 + a \left( \phi^2/2 + 3\phi^4/4 \right)$ is the bulk pressure.

In the corresponding LB scheme, distributions on both lattices are relaxed and streamed using the standard LBGK scheme (2) using the relaxation times $\tau_\rho = \tau_B + \frac{\phi+1}{2} (\tau_A - \tau_B)$ and $\tau_\phi$, respectively. The following equilibrium functions [15] are used to recover (12) in the macroscopic limit:

$$
f_{eq}^i = w_i \left( p_0 - \kappa \phi \nabla^2 \phi + e_{ia} u_\alpha \rho + \frac{3}{2} \left[ e_{ia} e_{i\beta} - \frac{\delta_{i\beta}}{3} \right] \rho u_\alpha u_\beta \right) + \kappa w_i^\alpha \beta \partial_\alpha \phi \partial_\beta \phi
$$

(14)

$$
g_{eq}^i = w_i \left( \Gamma \mu + e_{ia} u_\alpha \phi + \frac{3}{2} \left[ e_{ia} e_{i\beta} - \frac{\delta_{i\beta}}{3} \right] \phi u_\alpha u_\beta \right)
$$

(15)

where $\Gamma$ is a tunable parameter related to mobility via $M = \Gamma(\tau_\phi - 1/2)$, and the chemical potential $\mu = a(-\phi + \phi^3) - \kappa \nabla^2 \phi$. $\kappa$ and $a$ are constant parameters related to the surface tension $\gamma = \sqrt{8ka/9}$ and interface width $\xi = 2\sqrt{2\kappa/a}$. The values of the weights $w_i$ and $w_i^\alpha \beta$ can be found in the paper by Kusumaatmaja and Yeomans [16].

In order to minimize spurious currents at the interface between the two fluids, we use optimized gradient and Laplacian stencils [17].

2.3. Boundary conditions

Sailfish implements various boundary conditions, which taken together make it possible to model a wide range of physical situations:
Figure 2: The bounce-back scheme illustrated for the D2Q9 lattice. (1) full-way bounce-back, (2) half-way bounce-back. The bottom row of nodes within each group represents the bounce-back nodes, while the top row represents normal fluid. Black nodes are "dry" (do not represent fluid, do not undergo relaxation), while grey nodes are "wet". The dashed line represents the effective location of the no-slip boundary condition. For simplicity, only distributions originating from the top left node are shown in subsequent time steps. The steps are: (a) initial state, time $t$ (b) streaming, time $t+1$ (c) relaxation, time $t+1$ (arrows ending with disks represent the post-relaxation state), (d) streaming, time $t+2$.

- periodic boundary conditions (used to simulate a system spatially infinite along a given axis),
- no-slip (solid walls): half-way bounce-back, full-way bounce-back (both with configurable wetting when used with the free energy model), Tamm-Moth-Smith [18],
- density/pressure: Guo’s method, Zou-He’s method, equilibrium distribution, regularized [19],
- velocity: Zou-He’s method, equilibrium distribution, regularized [19],
- outflow: Grad’s approximation [20], Yu’s method [21], Neumann’s, nearest-neighbor copy, "do nothing" [22].

The bounce-back method is a simple idea originating from lattice gas automatons – a particle distribution is streamed to the wall node, and scattered back to the node it came from (see Figure 2.3). In the full-way bounce-back scheme the wall nodes do not represent any fluid ("dry" nodes) and are only used to temporarily store distributions before scattering. This approach is slightly easier to implement, but has the downside of the distributions needing two time steps to reach back to the originating fluid node (one step to reach the wall node, then another to be reflected back and streamed to the fluid node). The physical wall is effectively located between the last fluid node and the wall node. In contrast, in the half-way bounce-back the wall nodes do represent fluid, the wall is located beyond the wall node (away from the fluid domain), and the distributions are reflected in the same time step in which they reach the wall node [23].
For other types of boundary conditions, the problem is more complex, as the mass fractions pointing into the fluid domain are undefined (there are no nodes streaming to them in these directions). Various schemes have been proposed to work around this problem. In the equilibrium approach, the missing incoming distributions are assumed to have the same values as those opposite to them (bounce-back), and the post-collision distributions are defined to be \( f_i^\text{eq}(\rho, \vec{u}) \) with one of \( \{\rho, \vec{u}\} \) being specified as part of the boundary condition, and the other calculated from the local distribution function. Zou’s and He’s method \cite{24} uses the idea of the bounce-back of the non-equilibrium parts of the distributions to calculate the missing mass fractions. The regularized boundary conditions also use the non-equilibrium bounce-back idea, but only for an intermediate step to calculate the 2-nd moment of the non-equilibrium distribution, which is then used to reset all mass fractions at the boundary node via a regularization procedure. We refer the reader to the excellent paper by Latt et al. \cite{19} for a detailed overview of these and other boundary conditions.

The Grad’s approximation method uses a formula dependent on the density, velocity and the pressure tensor \( P_{\alpha\beta} = \sum_i e_{i\alpha} e_{i\beta} f_i \) to replace all distributions on the boundary node. The three macroscopic quantities can be taken to be those from the boundary node at the previous time step (less precise, fully local) or extrapolated from neighboring fluid nodes (more precise, nonlocal). Other outflow methods, such as the Neumann’s boundary implementation or Yu’s method, are also nonlocal and use a simple extrapolation scheme. The simplest methods (nearest-neighbor copy, do nothing) do not require extrapolation, but also give less control over the resulting macroscopic fields.

### 2.4. Body forces

Sailfish implements two popular ways of adding a body force \( \vec{F} \) to the simulation: Guo’s method \cite{25}, and the Exact Difference Method (EDM) \cite{26}. In both schemes, the actual fluid velocity \( \vec{v} \) can be computed as

\[
\rho \vec{v} = \sum_i \vec{e}_i f_i + \vec{F}/2 = \rho \vec{u} + \vec{F}/2,
\]

and a force term \( F_i \) is added to the right hand side of \( (16) \). Guo et al. analyzed a number of popular schemes and concluded that the optimal choice for \( F_i \) in an LBGK simulation is

\[
F_i = \left(1 - \frac{1}{2\tau}\right) w_i \left[ \frac{\vec{e}_i - \vec{v}}{c_s^2} + \frac{\vec{e}_i \cdot \vec{v}}{c_s^4} \right] \cdot \vec{F}.
\]

In the EDM, the body force term added to the collision operator is equal to the difference of equilibrium distribution functions computed using momentum after and before the action of the force \( \vec{F} \):

\[
F_i = f_i^\text{eq}(\rho, \vec{u} + \Delta \vec{u}) - f_i^\text{eq}(\rho, \vec{u})
\]

with \( \Delta \vec{u} = \vec{F}/\rho \). The advantages of the EDM are its lack of any spurious terms.
in the macroscopic Navier-Stokes equations and its applicability to any collision operator (not only LBGK).

3. Software design and algorithms

Computational software is traditionally developed in one of two different paradigms. Prototype code is written with relatively little effort in a high level environment such as Matlab or Python for exploratory or teaching purposes. Due to the relatively low performance of these environments, this type of code is unsuitable for large scale problems. In contrast, production code is written in lower level languages such as C++ or Fortran in a higher effort process, which can sometimes span many years and involve teams of developers. The resulting programs are more efficient, but also longer, less readable, and potentially difficult to maintain.

In our implementation of the lattice Boltzmann method for GPUs released under the name project Sailfish, we took a hybrid approach. We use the Python programming language with a template-based code generation module (using the Mako language) and a computer algebra system (Sympy) to generate code in CUDA C or OpenCL. We chose Python because it is a very expressive language, with bindings to many system libraries and great support for GPU programming via the PyCUDA and PyOpenCL packages [27]. In Sailfish, Python is used for setting up the simulation (initial conditions, boundary conditions, selecting an LB model), for simulation control, communication (e.g. between compute nodes in a cluster) and input/output (loading geometry, saving simulation results). We also employ the NumPy package to perform matrix operations efficiently.

```python
from sympy import Rational
def bgk_equilibrium(grid):
    out = []
    for ei, weight in zip(grid.basis, grid.weights):
        out.append(weight * (S.rho + S.rho * (3 * ei.dot(grid.v) +
                              Rational(9, 2) * (ei.dot(grid.v))**2 - Rational(3, 2) *
                              grid.v.dot(grid.v)))
    return out
```

Listing 1: BGK equilibrium function defined using Sympy expressions. $S$ is a class with predefined symbols. $grid$ is a class representing the lattice used for the simulation. Note that this code works for all lattices and dimensions. The discrete velocity vectors are stored in symbolic form in the grid.basis list. This allows for e.g. simple computation of dot products. Numerical coefficients in the formulas are also stored in symbolic form as rational expressions (e.g. Rational(3, 2)) instead of floating point values. This makes them exact and allows further symbolic simplification.

Mathematical formulas, such as (1), (3), (15) are stored in the form of Sympy expressions. We decided to do this after noticing that computer code for numerical calculations is often very repetitive, e.g. (3), when evaluated on a D3Q19 lattice, would expand to 19 long lines of code, even though the initial formula easily fits in one line (see Listing 1). This approach makes the code easier to read (the formulas can also be automatically converted to LaTeX expressions),
allows for automated consistency checks (e.g. one can easily, in a single line of code, verify that the 0th moment of $f^{eq}$ is $\rho$), enables easy experimentation with code optimization techniques (formulas can be reorganized e.g. for speed or precision of floating-point operations), and enables code reuse (e.g. the formula for equilibrium or the bounce-back rule is written down once, but can generate code for various types of lattices, both in 2D and 3D, see Listing 2 for an example).

Finally, the Mako template language is used to render the formulas into low-level CUDA/OpenCL code. The generated code is fairly small, contains comments, and is automatically formatted, making it suitable for instructional purposes as well as for one-off, ad-hoc modifications. This is in stark contrast to large, multimodular codes where the architecture of the system can be difficult to understand without extensive documentation.

Sailfish was designed from scratch and optimized for modern massively parallel architectures, such as GPUs. We wanted to achieve high flexibility of the code, ease of use when running or defining new simulations, and not take any performance hits compared to code written directly in CUDA C or OpenCL. In order to achieve that, we decided to use run-time code generation techniques. This provides some isolation from low-level hardware details, important in the case of GPUs which are still evolving rapidly and changing with every new GPU architecture. It also makes it possible to generate optimized code on a case-by-case basis and to automatically explore parameter spaces to find optimal solutions, thus saving programmer time and increasing their productivity. With many details, such as the number of compute units, size of on-chip memory, speed of access patterns to on-chip and off-chip memory, host-device latency and bandwidth, memory bandwidth to computational performance ratio directly impacting the performance of the code, experimentation and microbenchmarking are necessary to find combinations of parameters that work well.

We also considered other metaprogramming approaches to code generation, such as domain-specific languages, and templates in C++. We deemed the first solution to have too much overhead, and decided against the latter one since the expanded code cannot be saved for inspection and modification, and there were no open source computer algebra libraries providing a level of flexibility and sophistication comparable to Sympy.

```c
${device_func} inline void
bounce_back(Dist *fi)
{
    float t;

    for i in sym.bb_swap_pairs(grid):
        a = grid.idx_name[i]
        opp_i =
            grid.idx_opposite[i]
        b = grid.idx_name[opp_i]
        t = fi->$a;
        fi->$a = fi->$b;
        fi->$b = t;
}
```
__device__ inline void bounce_back(Dist * fi) {
    float t;
    t = fi->fE;
    fi->fE = fi->fW;
    fi->fW = t;
    t = fi->fN;
    fi->fN = fi->fS;
    fi->fS = t;
    t = fi->fNE;
    fi->fNE = fi->fSW;
    fi->fSW = t;
    t = fi->fNW;
    fi->fNW = fi->fSE;
    fi->fSE = t;
}

Listing 2: Full-way bounce-back rule. Left column: source code in Mako. Right column: generated CUDA C code for the D2Q9 lattice.

3.1. High-level simulation architecture

Sailfish takes extensive advantage of object-oriented programming techniques for modularization and code reuse purposes. Each simulation is defined using two classes – a simulation class, and a geometry class (see Listing 3). The simulation domain can be divided into cuboid subdomains, which do not need to fill the parts of the domain that do not contain any fluid (this makes it possible to handle complex geometries). The geometry class defines initial and boundary conditions for a single subdomain. The simulation class derives from a base class specific to the LB model used such as LBFluidSim (single component simulations) or LBBinaryFluidFreeEnergy (binary fluids using the free energy model). The simulation class can optionally also add body forces or define custom code to be run after selected steps of the simulation (e.g. to display a status update, check whether the steady state has been reached, etc).

The base simulation class specifies the details of the used LB model, such as the form of the equilibrium distribution function (in symbolic form, stored as a Sympy expression), number and names of macroscopic fields (density, velocity, order parameter, etc), and names of GPU code functions that need to be called at every simulation step.

When executed, every Sailfish simulation starts a controller process which parses any command line parameters and reads configuration files, decides how many computational nodes and GPUs are to be used (in case of distributed simulations), and how the subdomains are going to be assigned to them. It then starts a master process on every computational node (see Figure 3), which in turn starts a subdomain handler process for every subdomain assigned to the computational node. We use subprocesses instead of threads in order to avoid limitations of the Python runtime in this area (the global interpreter lock preventing true multithreading), as well as to simplify GPU programming. The subdomain handlers instantiate the simulation and geometry classes for their respective subdomains. The system state is initialized in the form of Numpy arrays describing the initial macroscopic fields, and these are then used to initialize the distribution functions on the GPU (using the equilibrium distribution function). Optionally, a self-consistent initialization procedure [28] can be performed to compute the density field from a known velocity field. Once the system is fully initialized, the simulation starts running, with the subdomain handlers exchanging information in a peer-to-peer fashion as determined by the connectivity of the global geometry.
Figure 3: High-level architecture of a distributed Sailfish simulation. The simulation is divided into 4 subdomains. The controller process uses the execnet Python library to start machine master processes on two computational nodes, which then spawn children processes to handle individual subdomains. The master and subdomain handlers use the ZeroMQ library to communicate.

```python
from sailfish.subdomain import Subdomain2D
from sailfish.node_type import NTFullBBWall, NTZouHeVelocity
from sailfish.controller import LBSimulationController
from sailfish.lb_single import LBFluidSim

class LDCSubdomain(Subdomain2D):
    max_v = 0.1

    def boundary_conditions(self, hx, hy):
        wall_map = (hx == self.gx-1) | (hx == 0) | (hy == 0)
        self.set_node((hy == self.gy-1) & (hx > 0) & (hx < self.gx-1),
                          NTZouHeVelocity((self.max_v, 0.0)))
        self.set_node(wall_map, NTFullBBWall)

    def initial_conditions(self, sim, hx, hy):
        sim.rho[:] = 1.0
        sim.vx[hy == self.gy-1] = self.max_v

class LDCSim(LBFluidSim):
    subdomain = LDCSubdomain

if __name__ == '__main__':
    LBSimulationController(LDCSim).run()
```

Listing 3: 2D Lid-driven cavity example in Sailfish.

3.2. GPU architecture overview

Modern GPUs are massively parallel computational devices, capable of performing trillions of floating-point operations per second. We will now briefly present the architecture of CUDA-compatible devices as a representative example of the hardware architecture targeted by Sailfish. Other devices not supporting CUDA but supporting OpenCL are based on the same core concepts. CUDA
devices can be grouped into three generations, called Tesla, Fermi, and Kepler—each one offering progressively more advanced features and better performance.

The CUDA GPU is organized around the concept of a streaming multiprocessor (MP). Such a multiprocessor consists of several scalar processors (SPs), each of which is capable of executing a thread in a SIMT (Single Instruction, Multiple Threads) manner. Each MP also has a limited amount of specialized on-chip memory: a set of 32-bit registers, a shared memory block and L1 cache, a constant cache, and a texture cache. The registers are logically local to the scalar processor, but the other types of memory are shared between all SPs in a MP, which allows data sharing between threads.

Perhaps the most salient feature of the CUDA architecture is the memory hierarchy with 1-2 orders of magnitude differences between access times at each successive level. The slowest kind of memory is the host memory (RAM). While the RAM can nowadays be quite large, it is separated from the GPU by the PCIe bus, with a maximum theoretical throughput in one direction of 16 GB/s (PCI Express 3.0, x16 link).

Next in line is the global device memory of the GPU, which is currently limited to several gigabytes and which has a bandwidth of about 100-200 GB/s. Global memory accesses are however high-latency operations, taking several hundred clock cycles of the GPU to complete.

The fastest kind of memory currently available on GPUs is the shared memory block residing on MPs. It is currently limited in size to just 48 kB (16 kB on Tesla devices), but has a bandwidth of ca 1.3 TB/s and a latency usually no higher than that of a SP register access.

The above description readily suggests an optimization strategy which we will generally follow in the next section and which can be summarized as: move as much data as possible to the fastest kind of memory available and keep it there as long as possible, while minimizing accesses to slower kinds of memory. When memory accesses are necessary, it also makes sense to try to overlap them with independent computation, which can then be executed in parallel effectively hiding the memory latency.

From the programmer’s point of view, CUDA programs are organized into kernels. A kernel is a function that is executed multiple times simultaneously on different MPs. Each instance of this function is called a thread, and is assigned to a single scalar processor. Threads are then grouped in one-, two- or three-dimensional blocks assigned to multiprocessors in an 1-1 manner (1 block - 1 MP). The blocks are organized into a one- or two-dimensional grid. The size and dimensionality of the grid and blocks is determined by the programmer at the time of kernel invocation. Knowledge of the grid position, and the in-block position makes it possible to calculate a thread ID that is unique during the kernel execution. Within a single block threads can synchronize their execution and share information through the on-chip shared memory. Synchronization between blocks is not supported in any way other than serializing the execution of kernels, and through atomic operations on global memory.
3.3. Low-level LB algorithms and data structures on GPUs

The great potential of modern GPUs as a hardware platform for simulating both two-dimensional [29] and three-dimensional flows [30] with the lattice Boltzmann method was quickly realized after the initial release of the CUDA programming environment in 2007. Recently, GPUs have also been used to investigate more complex, two-dimensional multicomponent flows [31].

There are three main factors that need to be carefully optimized in order to fully utilize the computational power of modern GPUs: memory access patterns, register utilization, and overlap of memory transfers and arithmetic operations. The first factor has received the most attention in the literature, but in this work we show that the remaining two are also important.

The most important data structure used in an LB simulation are the particle distributions \( f_i \). They are stored in the global GPU memory in a Structure of Arrays (SoA) fashion, effectively forming a 4D or 3D array (for 3D and 2D simulations, respectively), with the following index ordering: \((q,z,y,x)\), where \(q\) is the number of discrete velocities in the lattice. An Array of Structures (AoS) approach, while elegant from the point of view of object-oriented programming, is completely unsuitable for GPU architectures due to how global reads and writes are performed by the hardware.

Global memory accesses are performed by thread warps (32 threads) in transactions of 32, 64, or 128 bytes. On Fermi devices, accesses cached in the L1 cache are serviced with 128-byte transactions (the size of a full L1 cache line), while those cached in the L2 cache are serviced with 32-byte transactions. In order to attain good bandwidth utilization, the memory has to be accessed in contiguous blocks so that all bytes in a transaction are used for meaningful data. The memory location also has to be naturally aligned, i.e. the first address in the transferred segment must by a multiple of the segment’s size.

In Sailfish, we run all kernels in 1-dimensional thread blocks spanning the X axis of the subdomain, with a typical block size being 64, 128 or 192 nodes. Each thread handles a single node of the subdomain. Due to the layout of the distributions in global memory, we issue \(q\) read requests to load a full set of mass fractions in a thread block. Each request results in a fully utilized memory transaction. To ensure natural alignment of transactions, the X dimension of the distributions array in global memory is padded with unused nodes so that \(x\) is a multiple of 32 or 16, for single and double precision floating point numbers, respectively.

In addition to the distributions array, we also store a node type map (see Section 3.4), and macroscopic field arrays in global memory, all following the same memory layout and padding as described above.

A simple LB simulation in Sailfish repeatedly calls a single CUDA kernel called `CollideAndPropagate`, which implements both the collision and propagation step (see e.g. [2]), as well as any boundary conditions. A high-level summary of this kernel is presented in Algorithm 1.

Two basic patterns of accessing the distributions have been described in the literature, commonly called AB and AA [32]. In the AB access pattern, there
Algorithm 1 A GPU kernel to perform one step of a lattice Boltzmann simulation, using shared memory for propagation in the $x$ direction. offset($x, y, z$) is a function computing a linear index in a 1D array corresponding to a shift in the 3D subdomain space. $i$ is a thread index within the local block of threads ($threadIdx.x$ in CUDA C). $e_{km}$ is the $m$-th component of the $k$-th discrete velocity vector.

1: allocate shared memory array $buf_k[blockSize]$
2: compute global array index $i_G$ for the current node $i$
3: load and decode node type from global memory to $nt$
4: load distributions from global memory to $f$
5: compute macroscopic variables $\rho$ and $\vec{v}$
6: if $nt$ is a boundary node then
7: apply boundary conditions
8: end if
9: if output requested then
10: save $\rho$ and $\vec{v}$ in global memory arrays
11: end if
12: compute the equilibrium distribution $f^{eq}$ using $\rho$ and $\vec{v}$
13: relaxation: $f \leftarrow f + \frac{1}{\tau}(f^{eq} - f)$
14: for $k$ such that $e_{kx} = 0$ do
15:      $buf_k[i - 1] \leftarrow f_k$
16: end for
17: if $i \geq -1$ then
18:      $buf_k[i + 1] \leftarrow f_k$
19: end if
20: if $i < blockSize - 1$ then
21:      for $k$ such that $e_{kx} = -1$ do
22:          write $f_k$ to global memory at $i_G + offset(e_{ky}, e_kz)$
23:      end for
24: else
25:      for $k$ such that $e_{kx} = -1$ do
26:          write $f_k$ to global memory at $i_G + offset(e_{ky}, e_kz)$
27:      end for
28: end if
29: synchronize threads within the block
30: if $i > 0$ then
31:      for $k$ such that $e_{kx} = 1$ do
32:          write $buf_k[i]$ to global memory at $i_G + offset(0, e_ky, e_kz)$
33:      end for
34: else
35:      for $k$ such that $e_{kx} = 1$ do
36:          write $f_k$ to global memory at $i_G + offset(e_{ky}, e_kz)$
37:      end for
38: end if
39: synchronize threads within the block
40: if $i > 0$ then
41:      for $k$ such that $e_{kx} = 1$ do
42:          write $buf_k[i]$ to global memory at $i_G + offset(0, e_ky, e_kz)$
43:      end for
44: end if
are two copies of the distributions in global memory (A and B). The simulation step alternates between reading from A and writing to B, and vice versa. In the AA access pattern, only a single copy of the distributions array is stored in global memory. Since there are no guarantees about the order of execution of individual threads, care has to be taken that a thread reads from and writes to exactly the same locations in memory. This is illustrated in Figure 3.3. A third access pattern that is used in practice is the so-called indirect addressing. In this mode, in order to access data for a node, its address has to be read first. This causes overhead both in storage (need to store the addresses) and in memory accesses, but can be very useful for complex geometries where the active nodes are only a tiny fraction of the volume of the bounding box. Indirect addressing can be combined with both AA and AB access patterns. For a dense subdomain using indirect addressing, the performance can be 5-25% lower than when using direct addressing, with C2050 showing better results with the AA access pattern than in AB, and the GTX 680 exhibiting the opposite tendency. The exact performance is however necessarily geometry-dependent, and as such it is not discussed further here.

The propagation step of the LB algorithm shifts the distributions by $\pm 1$ node in all directions – when this happens for the X axis, it results in misaligned reads/writes. To reduce the impact of misaligned writes, Sailfish utilizes shared memory to shift data in the X direction within the thread block (see Algorithm 1).

We performed numerical tests to evaluate the impact of various data access

Figure 4: Memory layouts supported by Sailfish. In (a)-(d), dark arrows represent distributions used for calculations on the central node (middle square in every panel). In the AB layout, there are two copies of the lattice stored in memory. Data is read from the first lattice (a), and propagated into the second lattice (c). In the AA layout, there is only one lattice copy. Data is read from (a), stored as (b), and in the following step, read from (d) and stored as (c). In indirect addressing, there is a dense array of pointers to the actual distributions, which are stored in a continuous array (e). Dark nodes with black circles represent active (fluid) nodes. White nodes represent areas outside of the simulation domain which do not have corresponding distributions.
patterns on the overall performance of the simulation. Earlier works [33, 34] indicate that the cost of unaligned writes is significantly higher than the cost of unaligned reads, and that a propagate-on-read strategy results in up to 15% performance gain. Our experiments confirmed this on older GT200 hardware (GTX 285), however we failed to replicate this effect on Fermi and Kepler devices (Tesla C2050, K10, K20), where the performance of both implementations was nearly identical (typically, with a few % loss for propagate-on-read). This is most likely caused by hardware improvements in the Fermi and later architectures.

We also compared the performance for the AB and AA access patterns. On Tesla-generation devices (GTX 285, Tesla C1060), the AA memory layout results in a slightly higher performance, and is therefore clearly preferred. On newer devices the AB scheme is typically a little faster, but the performance gains over the AA scheme are minor (<10%), and as such the AB scheme is only preferable when ample GPU memory is available.

3.4. Boundary condition handling

Boundary conditions are handled in Sailfish with the help of a node type map – an unsigned 32-bit integer array stored in the global GPU memory. Each entry in this array contains encoded information about the node type (fluid, unused, ghost, type of boundary condition to use), orientation (vector normal to the boundary, pointing into the fluid) and a parameter ID. The parameter ID is an index to a global array of values used by boundary conditions (e.g. densities, velocities). The encoding scheme uses variable-size bitfields, which are dynamically chosen for every simulation depending on the usage of different boundary conditions in a subdomain (see Figure 5).

Time-dependence is supported for all types of boundary conditions. When a value changing in time is required, it is typically specified in the form of a Sympy expression. This expression is then transformed into a GPU function and assigned an ID, analogous to the parameter ID for static boundary conditions.

![Figure 5: Node parameter and type encoding into a single 32-bit unsigned integer to be used in the node type map. Bitfields (node type, node parameter index, ...) have adjustable size which can vary between simulations.](image)
3.5. Multicomponent models

Models with more than one distribution function, such as the Shan-Chen model or the free energy model introduce a nonlocal coupling via one or more macroscopic fields. To minimize the amount of data read from global memory, we split the simulation step into two logical parts, implemented as two GPU kernels. In the first kernel, ComputeMacroFields, we load the distributions and compute the macroscopic field value for every field which needs to be accessed in a nonlocal manner in the collision kernel. The collision step is implemented similarly to single fluid models, and the nonlocal quantities (Shan-Chen force, gradient and Laplacian of the order parameter in the free energy model) are computed by directly accessing the macroscopic field values in global memory.

We considered three approaches to realizing the collision and propagation part of the algorithm for multifluid models. In the first variant, we used a single collision kernel, which loaded both distributions into registers and ran the collision and propagation step for both lattices. In the second variant, we tried to speed-up the calculations of nonlocal values by binding the macroscopic fields to GPU textures and accessing them through these bindings. In the last variant, we split the collision process into two separate kernels, one for every lattice.

The second variant yielded minimal speed-ups (on the order of a few percent) on old Tesla-generation devices, which however did not carry over to Fermi and Kepler ones. We abandoned the idea as it introduced unnecessary complexity to the code. The third approach using split kernels proved to be the most efficient one. We were able to obtain 5.4% (D2Q9, free energy) – 53% (D3Q19, free energy) speed-ups as compared to a monolithic kernel, mainly due to the kernels using fewer registers and being able to achieve higher occupancy, which hid the minor overhead introduced by a slightly increased number of global memory accesses. As a side benefit, this approach also resulted in simpler code, so we decided to standardize on it in the Sailfish framework. We also expect it to scale well to models requiring more than 2 lattices.

3.6. Distributed simulations

In Sailfish, the mechanisms that support distributed simulations are very similar to those supporting multiple subdomains on a single GPU or many GPUs on one computational node. Every subdomain has a layer of ghost nodes around it. These nodes do not participate in the simulation, but are used for data storage for mass fractions leaving the subdomain or for macroscopic fields of neighboring nodes located in remote subdomains. Once mass fractions are streamed into the ghost nodes, we run additional CUDA kernels to collect the data leaving the subdomain into a linear buffer in global memory, which is then transferred back to the host and sent to remote nodes using a network link.

The subdomain is also split into two areas called bulk and boundary, which are simulated via two separate kernel calls. The boundary area is defined as all nodes belonging to CUDA thread blocks where at least one node touches the subdomain boundary. The simulation step is first performed for nodes in
the boundary area, so that communication with remote subdomain runners can start as soon as possible and can overlap in time with the simulation of the bulk of the subdomain. As a further optimization, we limit the data sent between computational nodes exactly to the mass fractions that actually need to be sent (e.g. if two nodes are connected along the X axis, then only the mass fractions corresponding to discrete velocities with a positive X component will be transferred from the left subdomain to the right one). We also make it possible to optionally compress the data before sending it to the remote node, which can be helpful if the simulation is run on multiple machines with slow network links between them, or when only a small fraction of nodes on the subdomain interface plane is active.

3.7. Impact of single precision

Many GPUs are significantly faster when calculations are done in single precision as opposed to the standard double precision which is typically used in scientific and engineering applications. The speed-up factor can vary between 10 and 2, depending on the device model and generation. The performance gap is smaller in newer devices (e.g. Kepler generation). Earlier works [35, 36] used the lid-driven cavity benchmark to verify that single precision calculations produce satisfactory results.

Here, we use the 2D Taylor-Green decaying vortex flow – a benchmark problem with known analytical solution – to study the accuracy of our LB implementation in both single (SP) and double (DP) precision. The simulations are done using the LBGK relaxation model on a D2Q9 lattice. For single precision, both the standard formulation and the round-off minimizing formulation [11, 21] (SRO) are tested. The Taylor-Green vortex can be described by the following system of equations:

\[
\begin{align*}
    u_x(t) &= -u_0 \cos(x) \sin(y)e^{-2\nu t} \\
    u_y(t) &= u_0 \sin(x) \cos(y)e^{-2\nu t} \\
    \rho(t) &= \frac{\rho}{4} (\cos(2x) + \cos(2y)) e^{-4\nu t}
\end{align*}
\]

(19)

where \( u_0 \) is a velocity constant and \( x, y \in [0; 2\pi) \). (19) can be easily verified to satisfy the incompressible Navier-Stokes equations. We performed multiple runs of the simulation on a 256^2 lattice with periodic boundary conditions in both directions, varying the viscosity \( \nu \) but keeping the Reynolds number constant at \( Re = 1000 \). Each simulation was run until it reached a specific point in physical time, corresponding to \( 10^6 \) iterations at \( u_0 = 0.0005 \). We use the L2 norm of the velocity and density field difference to characterize the deviation of the numerical solution from the analytical one:

\[
\epsilon = \sum_{\text{nodes}} \sqrt{\frac{(u_x - \hat{u}_x)^2 + (u_y - \hat{u}_y)^2}{u_x^2 + u_y^2}}
\]

(20)

where \( \hat{u}_x, \hat{u}_y \) is the numerical solution.
The results presented in Figure 6 illustrate interesting differences between double precision and single precision calculations. In double precision, the error stays constant until \( u_0 \approx 0.02 \) and raises quadratically for higher values of velocity, as could be expected from the \( O(\text{Ma}^2) \) accuracy of the LBGK model. In single precision however, lower speeds lead to higher error values. The interplay between model accuracy and numerical round-off leads to a sweet spot around \( u_0 \approx 0.05 \), where the errors are minimized. For higher speeds, there is no difference between double and single precision. This result shows that the conventional wisdom that lower speeds always lead to better precision is not true when applied to single precision simulations. Since many practical simulations are run at velocities 0.05 and higher for reasons of efficiency, it also explains why in many cases no differences are observed between single and double precision results. The density error shows that the round-off minimization model generates significantly better results than the standard single precision implementation, and as such should be preferred in cases where the accuracy in the density field is important.

4. Performance

All performance figures in this section are quoted in MLUPS (Millions of Lattice-site Updates per Second). The tests were run using CUDA Toolkit 5.0, PyCUDA 2013.1.1 on 64-bit Linux systems.

4.1. Comparison of different models

Single fluid models were benchmarked using a lid-driven cavity test case, with a \( 254^3 \) lattice at \( \text{Re} = 1000 \), using full bounce-back walls and equilibrium velocity boundary conditions for the lid. To illustrate the impact of the entropy-stabilized time-stepping, the ELBM solver was also tested at \( \text{Re} = 10000 \).
Figure 7: Performance comparison of different models using the D3Q19 lattice, AB memory access pattern. Used acronyms: ELBM: entropic LBM, SC: Shan-Chen, FE: free energy. All GPUs had ECC disabled. ELBM used intrinsic functions as described in section 4.4. Both logical GPUs were used for the K10 tests. Left panel: single precision. Right panel: double precision.

Figure 8: Performance of an LBGK simulation with the AA memory access pattern as a function of lattice type. The lower performance for the K10 simulation for D3Q13 and D3Q15 lattices is caused by overhead in copying the data between the two GPUs through the host (both logical GPUs were used for the K10 tests). Test case: Kida vortex (see text). Left panel: single precision. Right panel: double precision.
binary fluid models were benchmarked using a simple spinodal decomposition test case, where a uniform mix of both fluid components fills the whole simulation domain (254 × 100 × 100) and periodic boundary conditions are enabled in all directions. Whenever the domain size was too large to fit on a specific GPU we tested, we reduced the size in the Z direction until it fit.

We find our results to be comparable or better to those reported by Habich et al. [37], who used hardware very similar to ours (Tesla C2050 and C2070 differ only in memory size, according to NVIDIA specifications). The performance of single precision simulations can be shown to be limited by global memory bandwidth. We find that our code runs at \( \sim 80\% \) of the theoretical bandwidth as reported in NVIDIA whitepapers, and close to 100\% of the real bandwidth measured by sample code from the NVIDIA SDK. Double precision simulations run at \( \sim 60\% - 80\% \) of the theoretical maximum. They are limited by the double precision instruction throughput on the GPU on Fermi-class hardware and by memory bandwidth in Kepler hardware. Overall, we find that the memory bandwidth is a reliable indicator of expected performance for single precision simulations across all three generations of NVIDIA devices (see also Figure 8, which shows that the simulation performance is inversely proportional to the lattice connectivity \( Q \) in single precision).

For double precision simulations on Fermi hardware, we have found increasing the L1 cache size to 48 kB, disabling L1 cache for global memory accesses and replacing division operations by equivalent multiplication operations to have a large positive impact on the performance of the code (\( \sim 3.5 \) speed-up in total). The impact of the L1 cache is understandable if one considers the fact that double precision codes use significantly more registers. This causes the compiler to spill some of them over to local memory, accesses to which always go via the L1 cache. The larger the size of the unused part of that cache, the more operations can execute without actually accessing the global memory.

This optimization strategy does not apply to Kepler-class devices, where both in single and double precision we found that disabling the preference for L1 cache for the main computational kernel had a positive impact on performance.

We also tested our code on lower end GPUs (mobile versions used in laptops) with compute capability 3.0, where we found a significant speed up (up to 40\%) by using shuffle operations for in-warp propagation, and limiting shared memory for data exchange between warps only. This optimization strategy does not work with the higher end GPUs discussed in this paper, where the performance is limited by global memory bandwidth already.

Overall, we unsurprisingly find that more recent GPUs perform noticeably better. The K10 is an interesting option if only single precision is required as it delivers the highest overall performance at a level higher than 1.3 GLUPS per board with D3Q19 and simple fluid models. For double precision, the K20x card being the most advanced NVIDIA GPU available on the market when this paper is written, is a clear winner performance-wise.
4.2. **Scaling on GPU clusters**

While the computational power provided by a single GPU is impressive, practical simulations often require large domains, and for these the total size of GPU memory (a few GBs) is an important limitation. The natural solution of this problem is to run a distributed simulation using multiple GPUs, which can be physically located in a single computer or multiple computers in a network.

In order to measure performance of distributed simulations, we ran a 3D duct flow test case (periodic boundary conditions in the streamwise $Z$ direction, bounce-back walls at other boundaries) on the Zeus cluster (part of the PLGRID infrastructure), consisting of computational nodes with 8 M2090 GPUs and interconnected with an Infiniband QDR network.

4.2.1. **Weak scaling**

The first test we performed measured weak scaling, i.e. code performance as a function of increasing size of the computational domain. The domain size was $254 \times 127 \times 512 \times N$, where $N$ is the number of GPUs. We used the D3Q19 lattice, the AA memory access pattern, a CUDA block size of 128, and single precision. Figure 9 shows excellent scaling up to 64 GPUs, which was the largest job size we were able to run on the cluster. The 1.5% efficiency loss takes place as soon as more than 1 subdomain is used, and does not degrade noticeably as the domain size is increased. This small efficiency loss could be further minimized by employing additional optimization techniques, such as using peer-to-peer copies when the GPUs are located on the same host.

4.3. **Strong scaling**

The second test we ran measured strong scaling, i.e. code performance with a constant domain size, as a function of increasing number of GPUs. We used a $254 \times 127 \times 1664$ geometry (largest domain size that fit within the memory of a
single M2090 GPU) and other settings as in the weak scaling test, and divided the domain into equal-length chunks along the Z axis as more GPUs were used for the simulation. The results of this test are presented in Figure 10. A slightly worse performance is visible in comparison to the weak scaling test, but even when the simulation is expanded to run on 8 GPUs, only a 3.5% efficiency loss can be observed.

It should also be noted, that there is a minimum domain size below which performance quickly degrades due to the overhead of starting kernels on the GPUs. This size can be seen in Figure 11 to be about 14% of the GPU memory or 8.2 M lattice nodes.

### 4.4. Further optimization with intrinsic functions

CUDA GPUs provide an alternative hardware implementation of various transcendental functions such as the exponent, logarithm or trigonometric functions. These functions, known as intrinsic functions, are faster but less precise than their normal counterparts, especially when their arguments do not fall within a narrow range specified for each function.

We analyzed the impact of these functions on the performance and precision of the LB models that can take advantage of them, namely the Shan-Chen model with a non-linear pseudopotential and the entropic LBM. With ELBM the use of intrinsic functions, together with the FMAD (fused multiply-add) instruction yields a speed-up of $\sim 43\%$ without any noticeable impact on the correctness of the results (in terms of global metrics such as the total kinetic energy, enstrophy and the kinetic energy spectrum). While testing these optimizations, we also found that the FTZ (denormalize to 0) option of the CUDA compiler causes the ELBM simulation to crash.

With the proposed optimizations, the performance of ELBM is at 72% of the LBGK performance, making it a very interesting alternative for some simulations.
Figure 11: Duct flow simulation performance as a function of used memory fraction of a single M2090 GPU. The vertical line shows the point at which the computational capabilities of the GPU are saturated. The geometry was the same as in the strong scaling test and the memory fill fraction was controlled by varying the extent of the domain in the Z direction. Very similar behavior is observed for K10 and K20 GPUs (not shown here).

For the Shan-Chen model with a nonlinear pseudopotential we saw 17-20% speed-ups in 2D and 3D, with relative changes in the density fields smaller than 1.5% after 10000 steps.

Unfortunately, the same approach does not yield speed-ups in double precision, as most intrinsic functions are available in single precision only.

5. Validation

In order to validate our implementation of the LBM, we performed simulations for four classical computational fluid dynamics test cases and compared our results with those published in the literature.

5.1. Lid-driven cavity

The lid-driven cavity geometry consists of a cube cavity with a face length $L$. The geometric center of the cavity is located at the origin of the coordinate system. The cube face at $x = -L/2$ moves tangentially in the $y$-direction with a constant velocity $v$, while all other faces are no-slip walls. We carried out simulations of this problem at $Re = 1000$ with various LB models (BGK, MRT, regularized BGK, ELBM) using a $201^3$ D3Q19 lattice, full-way bounce-back for no-slip walls, and the regularized velocity boundary condition with $u_y = 0.05$ for the moving wall.

Our results (both in single and double precision) agree with those published by Albensoeder et al. (see Figure 12).
Figure 12: Lid-driven cavity velocity profiles $u_y(x,0,0)$ and $u_x(0,y,0)$. Round dots: data from Tables 5 and 6 in Albensoeder et al. [38]. Solid line: results from Sailfish simulations on a 201$^3$ lattice, after $2 \cdot 10^5$ steps using LBGK, MRT, regularized BGK and ELBM in single and double precision. The results from all Sailfish simulations are in agreement to within the width of the line on the plot. LB results are rescaled using: $\vec{u} = \vec{u}_{LB}/0.05$.

5.2. Kida vortex

The Kida vortex flow is a free decay from the initial conditions [39]:

$$u_x(\vec{x}, 0) = u_0 \sin x \cos y \cos z - \cos y \cos 3z$$
$$u_y(\vec{x}, 0) = u_0 \sin y \cos 3z \cos x - \cos z \cos 3x$$
$$u_z(\vec{x}, 0) = u_0 \sin z \cos 3x \cos y - \cos 3y \cos x$$

defined on a cubic domain with face length $2\pi$, with periodic boundary conditions in all directions. To validate our code, we performed simulations for $Re = Nu_0/\nu = 4000$, $1.28 \cdot 10^4$, and $1.28 \cdot 10^5$ and compared them with results published in [40] and [41], respectively. The simulations were run using $u_0 = 0.05$ on a 700$^3$ grid (350$^3$ for $Re = 4000$), using both single and double precision (with no noticeable difference between them). The $Re = 4000$ and $Re = 1.28 \cdot 10^4$ cases were investigated using the LBGK, MRT, regularized LBGK, Smagorinsky-LES and entropic models. At $Re = 1.28 \cdot 10^5$, only simulations using the entropic model and the Smagorinsky subgrid model (with $C_S = 0.1$) remained stable. During the simulation time kinetic energy $E = \frac{1}{2V} \int d^3x \vec{v}^2$ and enstrophy $\Omega = \frac{1}{2V} \int d^3x (\vec{\nabla} \times \vec{v})^2$, where $V$ is the volume of the simulation domain, were tracked directly on the GPU for optimal efficiency. Vorticity was computed using the first order central difference scheme in the bulk of the fluid and forward/backward differences at subdomain boundaries.

For $Re = 4000$ all four models gave the same results (Figure 13). At $Re = 1.28 \cdot 10^4$ some minor differences are visible, particularly in the evolution of enstrophy. Its peak value is slightly underpredicted by both models that locally modify effective viscosity (Smagorinsky, ELBM) (see Figure 13(c)). At $Re = 1.28 \cdot 10^5$, the differences are more pronounced and we observe that the Smagorinsky model underpredicts the absolute value of peak enstrophy. The kinetic energy spectrum shown on Figure 13(d) was computed as $E(k) = \sum_{k \leq k' < k+1} \hat{u}(k)^2$, for $k = 0, 1, 2, \ldots$. A good agreement is visible in comparison
Figure 13: (a) evolution of normalized kinetic energy, (b) evolution of normalized enstrophy, (c) evolution of normalized enstrophy at Re = 12800 for various collision models, (d) kinetic energy spectrum for selected collision models and Reynolds numbers. For panels (a)-(c) time is rescaled assuming a domain size of \((2\pi)^3\) and \(u_0 = 1\).
to the Kolmogorov scaling $k^{-5/3}$, especially for the high Reynolds number cases. All collision models lead to similar spectra, with ELBM at Re = 4000 predicting a slightly higher value around $k = 10$ than LBGK or other models, and with ELBM keeping a slightly flatter spectrum for high $k$ values at Re = $1.28 \cdot 10^5$. In all cases the simulation results show the same features as those discussed in previous papers on this topic [39, 40, 41].

5.3. Binary Poiseuille flow

To verify the binary fluid models we consider a 2D Poiseuille flow in the $x$-direction. No-slip walls are imposed at $y = \pm L$ using the half-way bounce-back boundary conditions, and periodic boundary conditions are used in the $x$-direction. A body force $G$ drives the flow. In the core flow region ($|y| \leq L/2$) a fluid of viscosity $\nu_1$ is placed, while in the boundary flow region ($L/2 < |y| < L$) the viscosity of the fluid is $\nu_0$. The analytical solution for this case can be expressed as:

$$u_x(y) = \begin{cases} \frac{G}{2\rho_0} \left( L^2 - y^2 \right) & \text{if } L/2 < |y| < L \\ \frac{GL^2}{8} \left( \frac{3}{\rho_0} \rho_1 + \frac{1}{\rho_0} \left( 1 - 4\frac{y^2}{L^2} \right) \right) & \text{if } |y| \leq L/2 \end{cases}$$  \hspace{1cm} (21)

We run the simulation on a $64 \times 256$ grid with $\nu_0 = 1/6$, and $\nu_1 = 1/24$. The simulation starts with $u_x = 0$ in the whole domain and we let the flow reach the stationary state on its own. The free energy simulation was run with $\Gamma = 25$, $\kappa = 10^{-4}$, and $A = 32 \cdot 10^{-4}$, while for the Shan-Chen model, $G = 1.5$ was used. The parameters were chosen to ensure that the interface remains as sharp as possible without destabilizing the simulation. The Exact Difference Method was used to introduce body forces. The Shan-Chen model permits residual mixing between the fluid components, so the effective density and viscosity were calculated as $\rho = \rho_0 + \rho_1$ and $\nu\rho = \rho_0\nu_0 + \rho_1\nu_1$, respectively. Figure 14 illustrates the good agreement of the simulation results with (21).

5.4. Capillary waves

In order to verify the binary fluid models also in a dynamic case, we simulate the decay of the amplitude of a capillary wave. The boundary conditions of the system are the same as in the binary Poiseuille flow case, but we now use a larger $512 \times 512$ lattice in order to accommodate higher frequency waves. The region $y > 0$ is filled with one component (A) and the region $y < 0$ is filled with another component (B). For simplicity, we choose both components to have the same viscosities $\nu = 1/18$. The interface between the two fluids is initialized to a sinusoidal curve of wavelength $\lambda$ chosen such that an integer number of wavelengths fits exactly in the simulation domain. The interface is then allowed to relax naturally with no external forces, resulting in a damped capillary wave. At each timestep of the simulation, the height of the interface at $x = L - \lambda/4$ is recorded. In order to recover the frequency of the wave, an exponentially damped sine function is fit to the interface height data. For the Shan-Chen model, we used $G = 0.9$ and for the free energy model we used $A = 0.02$. 

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Figure 14: Comparison between simulated and theoretical velocity profiles for the binary Poiseuille flow. Left panel: Results for the free energy model. Right panel: Shan-Chen results. The effective viscosity ratio was 3.7985 due to mixing between the two fluid components. The free energy model shows better agreement with the theoretical profile due to a thinner interface.

Figure 15: Theoretical and measured dispersion relation $\omega(k)$ for the capillary wave for the free energy and Shan-Chen models.
\( \kappa = 0.04 \), and \( \Gamma = 0.8 \). As expected [42, 43], the dispersion relation shows a power law form \( \omega \propto k^{3/2} \) for both the Shan-Chen and free energy models (see Figure 15).

6. Conclusions

In the previous sections we have demonstrated our Sailfish code as a flexible framework for implementing lattice Boltzmann models on modern graphics processing units. With novel optimization techniques for complex models it provides a very efficient tool for a wide range of simulations. We hope that our observations collected while running the presented benchmarks will serve as a guideline in the choice of both LB models and computational hardware for users of Sailfish and of other similar codes.

For single precision simulations, we advocate a careful choice of parameters and correctness testing via comparisons to similar test cases in double precision. While all of our benchmark problems did not show any noticeable differences between single and double precision, the Taylor-Green test case clearly demonstrates that these do exist and can significantly impact the results if the simulation is in the slow velocity regime. Whenever possible, the round-off minimizing model should be used to reduce precision losses without any impact on performance.

While the capabilities of Sailfish are already quite extensive, much work remains to be done. Among the most important remaining tasks, we mention ongoing efforts to implement a hierarchical lattice, allowing for local grid refinement and a fluid-structure interaction model based on the immersed boundary method. Since the code is freely available under an open source license, we would like to invite the reader to participate in its development and contribute new enhancements according to their interests.

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