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Multi-GPU Implementation of the Lattice Boltzmann Method

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Abstract

The lattice Boltzmann method (LBM) is an increasingly popular approach for solving fluid flows in a wide range of applications. The LBM yields regular, data-parallel computations; hence, it is especially well fitted to massively parallel hardware such as graphics processing units (GPU). Up to now, though, single-GPU implementations of the LBM are of moderate practical interest since the on-board memory of GPU based computing devices is too scarce for large scale simulations.

In this paper, we present a multi-GPU LBM solver based on the well-known D3Q19 MRT model. Using appropriate hardware, we managed to run our program on six Tesla C1060 computing devices in parallel. We observed up to $2.15 \times 10^9$ node updates per second for the lid-driven cubic cavity test case. It is worth mentioning that such performance is comparable to the one obtained with large high performance clusters or massively parallel supercomputers.

Our solver enabled us to perform high resolution simulations for large Reynolds numbers without facing numerical instabilities. Though, we could observe symmetry breaking effects for long-extended simulations of unsteady flows. We describe the different levels of precision we implemented, showing that these effects are due to round off errors, and we discuss their relative impact on performance.

Keywords: GPU programming, CUDA, Lattice Boltzmann method, TheLMA project

1. Introduction

Although the original Moore’s law \cite{15}, i.e. the exponential growth of transistor count on processors is still valid nowadays, the advances in computing performance are less straightforward. During the last decade, graphics processing units (GPU) have gradually outrun CPUs in terms of raw computational power. Using nVidia’s CUDA technology \cite{16}, GPUs have proven to be effective platforms to implement various high performance computing applications, ranging from linear algebra \cite{2} to CFD \cite{6} and PDE solvers \cite{14}.

The lattice Boltzmann method (LBM) is a novel approach in computational fluid dynamics. It appears to be an interesting alternative to the solving of the Navier-Stockes equations for various applications such as multiphase flows or porous media. As other CFD methods, the LBM is very demanding from a computational standpoint. High performance parallel implementations are therefore necessary for the LBM to be of practical interest.

Several successful implementations for the GPU are described in literature \cite{8, 21, 22}. Nonetheless, single-GPU implementations are bound by the device memory. The maximum available amount, when using GT200 GPUs, is 4 GB, which enables to handle at most about $2.83 \times 10^7$ nodes in single precision using the three-dimensional D3Q19 stencil. Multi-GPU implementations are therefore mandatory to run large scale LBM simulations, but are still a pioneering field and performance is often below what is expected from such hardware \cite{20}.

Recently released motherboards are able to manage up to eight GPU based computing devices. Although a MPI based multi-GPU LBM solver would be of great interest to run on hybrid clusters, we chose as a first step to implement a simpler POSIX thread based solver. The remaining of the paper is organized as follows. We first briefly re-
view the LBM and the CUDA technology. Then we give some general guidelines for implementing the LBM on GPUs and describe our multi-GPU implementation. Last, we discuss numerical issues we could observe running large scale simulations at high Reynolds numbers.

2. Multiple-Relaxation-Time LBM

Although originating from the lattice-gas automata theory [7], the lattice Boltzmann method is now generally interpreted as a way to solve the linearised Boltzmann equation [13]. In our work, we used the multiple-relaxation-time (MRT) approach [4] instead of the more popular Bhatnagar-Gross-Krook (BGK) version of the LBM [19]. In this section we shall briefly describe the MRT LBM.

With the Boltzmann equation, a fluid is described using a single-particle distribution function \( f \) depending on space and particular velocity, i.e. phase space, and on time. In the LBM, space is usually represented by a regular orthogonal mesh of resolution \( \delta x \) and time is split in constant steps \( \delta t \). The discrete counterpart of the continuous velocity space is a finite set of velocities \( \xi_i \), carefully chosen in order to ensure sufficient isotropy. Usually, vectors \( \delta \xi_i \) link nodes to only some of their nearest neighbours. As an example, fig. 1 shows the D3Q19 stencil we used in our computations.

![D3Q19 stencil](image)

**Figure 1**: D3Q19 stencil

Let us denote: \( |a_i| = (a_0, \ldots, a_N)^T \), \( T \) being the transpose operator. The lattice Boltzmann equation (LBE) writes:

\[
|f_i(x + \delta t \xi_i, t + \delta t) - |f_i(x, t)| = \Omega\left[|f_i(x, t)|\right]
\]

(1)

where \( \{f_i| i = 0, \ldots, N\} \) is the discrete equivalent of \( f \), and \( \Omega \) is the collision operator.

In the MRT approach, collision is performed in moment space. The particle distribution is mapped to a set of moments \( \{m_i| i = 0, \ldots, N\} \) by an orthogonal matrix \( M \):

\[
|f_i(x, t)| = M^{-1}|m_i(x, t)|
\]

(2)

where \( |m(x, t)| \) is the moment vector. Matrix \( M \) for the D3Q19 stencil can be found in appendix A of [5]. The corresponding moment vector is:

\[
|m_i(x, t)| = (\rho, \varepsilon, j_x, j_y, j_z, q_x, q_y, q_z, 3p_{xx}, 3p_{yx}, p_{xy}, p_{yz}, p_{pz}, m_x, m_y, m_z)^T
\]

(3)

where \( \rho \) is the mass density, \( \varepsilon \) is energy, \( \varepsilon \) is energy square, \( j = (j_x, j_y, j_z) \) is the momentum, \( q = (q_x, q_y, q_z) \) is the heat flux, \( p_{xx}, p_{yx}, p_{yz}, p_{pz} \) are related to the components of the stress tensor, \( \pi_{xx}, \pi_{yy}, \pi_{zz} \) are third order moments, \( m_x, m_y, m_z \) are fourth order moments. The mass density and the momentum are the conserved moments.

The LBE thus writes:

\[
|f_i(x + \delta t \xi_i, t + \delta t) - |f_i(x, t)| = -M^{-1}S\left[|m_i(x, t)| - |m_i^{eq}(x, t)|\right]
\]

(4)

where \( S \) is a diagonal collision matrix and the \( m_i^{eq} \) are the equilibrium values of the moments. For the sake of isotropy, \( S \) obeys:

\[
S = \text{diag}(0, s_1, s_2, 0, s_4, 0, s_4, 0, s_4, s_9, s_10, s_9, s_1,s_{12}, s_{13}, s_{13}, s_{15}, s_{16}, s_{16})
\]

(5)

We additionally set \( s_9 = s_{13} \), the initial density \( \rho_0 = 1 \), and the speed of sound \( c_s = 1/\sqrt{3} \), the unit of speed being \( \delta x/\delta t \). The equilibrium values of the non-conserved moments are thus given by:

\[
\varepsilon^{eq} = -11\rho + 19j^2
\]

(6)

\[
\varepsilon^{eq} = -\frac{475}{63}j^2
\]

(7)

\[
q_x^{eq} = -\frac{2}{3}j
\]

(8)

\[
3p_{xx}^{eq} = 3j^2_x - j^2
\]

(9)

\[
p_{xy}^{eq} = j^2_y - j^2_z
\]

(10)

\[
p_{yz}^{eq} = j^2_x - j^2_z
\]

(11)
The kinematic viscosity $\nu$ of the model is related to relaxation rate $s_9$ by:

$$\nu = \frac{1}{3} \left( \frac{1}{s_9} - \frac{1}{2} \right)$$

(14)

The other rates are set according to [9]. Namely: $s_1 = 1.19$, $s_2 = s_{10} = 1.4$, $s_4 = 1.2$, and $s_{16} = 1.98$.

3. Overview of the CUDA Technology

The Compute Unified Device Architecture (CUDA) is nowadays leading technology for general purpose computations on GPUs. Initiated in late 2007 by the nVidia company, CUDA defines both a programming model and general hardware specifications. CUDA capable GPUs consist in a set of streaming multiprocessors (SM), each containing several scalar processors (SP) as outlined in fig. 2. The SPs within a SM follow a single-instruction multiple-data (SIMD) execution scheme. Yet, SMs are not globally synchronised, thus the overall execution scheme may be described as single-instruction multiple-thread (SIMT).

CUDA computing devices show a complex memory hierarchy. The main storage consists in a rather large off-chip device memory. This memory is not cached except for specific read-only data (i.e. constants and textures); hence it suffers of high latency which has to be properly hidden. Each SM provides its SPs with non-addressable registers and some addressable shared memory which allows inter-SP communication.

The CUDA programming language is an extension to C/C++ (with some restrictions). A CUDA program basically consists in CPU code and (at least) one kernel, i.e. a void returning function to be executed by the GPU. Kernels are executed in several threads with private local variables. Threads are grouped in identical blocks which may have up to three dimensions. During execution, a block cannot be partitioned and therefore must fit into a single SM. Nonetheless, a SM may execute several blocks concurrently. Threads within a block may be synchronised and have access to a shared memory space. Yet, no protection mechanism, e.g. mutexes, is available: it is up to the programmer to manage this aspect.

Blocks are grouped into a one or two-dimensional execution grid, specified at launch time. Blocks are executed asynchronously and there is no efficient dedicated mechanism to ensure global synchronisation. All threads within a grid have access to a global memory space which is hosted in the device memory and is persistent along the application life cycle. Global synchronisation is therefore achieved by performing multiple kernel launches.

$$\begin{align*}
(\mathbf{f}_i(x, t)) &= M^{-1} \left( |m_i(x, t)| \right) \\
&\quad - S \left( |m_i(x, t)| - |m_i^{\mathrm{eq}}(x, t)| \right) \\
&\quad \left| \mathbf{f}_i(x + \delta t \xi_i, t + \delta t) = |\mathbf{f}_i(x, t)| \right.
\end{align*}$$

(15)

(16)
where eq. 15 describes the collision step and eq. 16 the propagation step. Thus, the LBM described in sec. 2, may be summarised by the following pseudo-code:

1. **for each** time step \( t \) **do**
2. **for each** lattice node \( x \) **do**
3. read velocity distribution \( f_i(x, t) \)
4. **if** node \( x \) is on boundaries **then**
5. apply boundary conditions
6. **end if**
7. compute moments \( m_i(x, t) \)
8. compute equilibrium values \( m_i^{(eq)}(x, t) \)
9. compute updated distribution \( \tilde{f}_i(x, t) \)
10. propagate to neighboring nodes \( x + \delta \xi_i \)
11. **end for**
12. **end for**

The most convenient approach to take advantage of the massive parallelism of GPUs is to assign one thread to each node. Threads within a block are executed in groups of 32 threads named **warps**. Global memory transactions are issued by half-warp. Best performance is achieved when these operations may be coalesced into single transactions of 32 B, 64 B, or 128 B. Yet, segment transactions face the important restriction that the segment’s offset has to be a multiple of its size.

Optimised CPU implementations of the LBM generally store the particle distribution in an array of structures, which improves data locality. In order to allow coalescing, GPU implementations must adopt a reverse approach. A simple and efficient solution is to use one dimensional blocks corresponding to a given spatial direction and to store the particle distribution in a multi-dimensional array. The minor dimension of the array is chosen such as contiguous threads access to contiguous memory locations.

Nonetheless, this approach is not sufficient to ensure optimal memory transactions. For most of the particle populations, the propagation step leads to one unit shifts in addresses as illustrated by fig. 4.

With the first generation of CUDA enabled GPUs, i.e. for compute capability up to 1.1, alignment is mandatory for coalescence to occur, hence misalignment has dramatic impact on performance. To address this problem, propagation within the blocks may be performed using shared memory as described in [21]. As of compute capability 1.2, misaligned memory accesses are issued in as few segment transactions as possible. As thoroughly shown in [18], misaligned reads are far less expensive than misaligned writes, hence a rather efficient way to perform propagation is to use the out-of-place propagation scheme [17], outlined in fig. 5.

5. **Multi-GPU Implementation of the LBM**

Developing libraries is a common and acknowledged practice in software engineering. The Palabos project [11] for instance, in the case of LBM, provides a wide set of generic functions which allows to efficiently implement a parallel CPU LBM solver with given geometry, boundary conditions, and lattice Boltzmann model.
Nevertheless, the CUDA technology has some inherent limitations which make difficult to follow the same path when developing GPU LBM solvers. The compilation tool chain, for instance, being unable to link GPU binaries forbids actual modular programming. Likewise, devices of compute capability up to 1.3 have limited support for functions. The so-called device functions, i.e. functions to be executed by the GPU, are mostly inlined at compile time, which restricts their use in practice.

In order to improve code reusability, we designed the TheLMA framework [1]. TheLMA stands for Thermal LBM on Many-core Architectures, thermal flow simulation being our main topic of interest. It provides a global template for multi-GPU LBM solvers on which we developed the present implementation. Figure 6 outlines the structure of the framework.

![Figure 6: The TheLMA framework](image)

The main.c file contains the main loop of the simulation and may access to a set of commodity functions in order to retrieve parameters, initialise variables, perform statistical calculations, and output simulation results in various formats. The thelma.cu file is a hub containing some general macros and including the CUDA components responsible for setting up the geometry, initializing, running the simulation and extracting results. Each of these component contains a launch function which is accessible to the C part of the program and handles the actual kernel invocation.

At initialisation, the program creates one POSIX thread for each requested computing device in order to hold the corresponding CUDA context. A sub-domain of the global lattice is assigned to each device. As for single-GPU implementation, synchronisation within the sub-domains is achieved by launching a kernel for each time step. Global synchronisation uses standard POSIX barriers. Inter-GPU communication is performed using page-locked CPU memory and zero-copy memory transactions.

As for global memory accesses, zero-copy transactions require coalescing to achieve optimal performance. This implies that the interfaces between sub-domains should be parallel to the direction associated with the minor dimension of the particle distribution array. For the sake of simplicity, we chose to split the lattice in rectangular cuboids along the direction corresponding to the major dimension. Figure 7 outlines the inter-GPU communication scheme.

![Figure 7: Inter-GPU communication scheme](image)

Each interface between sub-domains is associated to four buffers: two for incoming and two for outgoing populations. Pointers are switched after each time step. Maximal parallelization efficiency requires perfect overlapping of computations and communication. The zero-copy feature enables such overlapping, but the overlapping ratio depends on the scheduling of memory transactions at warp level. The execution grid set-up is therefore an important optimisation target.

Another problem arise when considering the configuration of the execution grid, since it may only have up to two dimensions. The simple solution of using one-dimensional blocks and a two-dimensional grid to span the three spatial dimensions does not apply to large lattices. On GT200 hardware, for instance, the resource requirements of a LBM kernel are likely to forbid the use of blocks greater than 256.

We therefore chose to use a two-dimensional grid of size \((\ell_x \times \ell_y \times \ell_z/2^m) \times (2^m - n)\) with one-dimensional blocks of size \(2^n\); \(\ell_x, \ell_y, \ell_z\) being the...
dimensions of the lattice, \( m \) and \( n \) being free parameters. Retrieval of coordinates is done using the following code:

\[
w = \text{blockIdx.x} \ll m \mid \text{blockIdx.y} \ll n \mid \text{threadIdx.x};
\]

\[
x = w \mod 1X;
\]

\[
y = (w/1X) \mod 1Y;
\]

\[
z = w/(1X*1Y);
\]

The optimal values for \( m \) and \( n \) are \( m = 15 \) and \( n = 7 \), which were determined empirically. To validate our code, we implemented the well-known lid-driven cubic cavity test case in which five walls have null velocity boundary conditions and the top lid has imposed constant velocity. In order to study the scalability of the program, we chose to run performance tests on a \( 192^3 \) lattice which may be handled by one single GPU or split in two, three, four, or six identical sub-domains as well. Figure 8 shows the obtained performance in million lattice node updates per second (MLUPS) for single precision with Tesla C1060 computing devices on a Tyant B7015 server. It is worth mentioning that the maximal performance is of the same order of magnitude than the one obtained with optimised double precision code on supercomputers (see [10], for instance).

![Figure 8: Performance on a 192^3 lattice](image)

Scalability is excellent with no less than 90% parallelization efficiency. Table 1 displays the required throughput for data exchange at 100% efficiency. With the Tyant S7015 motherboard of our server, the bandwidthTest program that comes with the CUDA development kit gives 2.78 GB/s host to device and 1.80 GB/s device to host maximum sustained throughput. The data exchange being symmetric, we may use the arithmetic mean of these values, i.e. 2.29 GB/s as a rough estimate of the available throughput for one PCI-E 16× slot. A comprehensive study of communications between computing devices and main memory is beyond the scope of this work and shall be given in future reference.

Table 1 shows that even with six GPUs, i.e. five sub-domain interfaces, the required throughput is comparable to the one achievable with a single PCI-E 16× slot, therefore data exchange is not likely to overflow the capacity of the PCI-E links. Furthermore, we see that the execution grid configuration we propose enables very satisfactory communication/computation overlapping.

According to the bandwidthTest program, the GPU to device memory maximum sustained throughput is 73.3 GB/s for the Tesla C1060. Performance in single precision using one GPU is 387 MLUPS which correspond to a data throughput of 80.4% of the maximum. We may therefore conclude that our single precision solver is memory bound and that performance is nearly optimal.

Performance for the double precision version of our solver ranges from 117 MLUPS using one GPU to 683 MLUPS using six, with similar scalability than for the single precision version. Considering one GPU, the corresponding data throughput is only 48.5% of the maximum, which implies that the double precision version is not memory bound but computation bound.

6. Numerical Issues

Although the lid-driven cubic cavity test case is well documented at low Reynolds numbers, there are to the best of our knowledge–very few references for \( \text{Re} \geq 12,000 \) [3, 12]. Using the six available Tesla C1060 cards, our solver is able to handle cubic lattices containing as much as \( 512^3 \) nodes for single precision D3Q19 and \( 384^3 \) nodes for double precision D3Q19. We could therefore run simulations for Reynolds numbers up to 30,000 without facing numerical instabilities.

According to nVidia, peak performance for the Tesla C1060 is 933 GFlops in single precision and 78 GFlops in double precision. As a matter of fact, GT200 GPUs are usually less efficient with double precision computations than with single precision. In our case, the performance ratio is about 3.2 to one. Nevertheless, the GT200 implementation of single precision is not fully IEEE 754 compliant. When first running our solver at \( \text{Re} = 30,000 \) in
single precision, we could see some numerical issue arise: the flow loses symmetry at a very early stage of simulation. Further investigation showed us that the average deviation from initial density decreases at a constant pace instead of fluctuating around zero.

To evaluate the impact of machine accuracy on our simulations, we experimented three levels of precision: single precision (sp), mixed precision (mp), i.e. double precision computations with single precision storage, and double precision (dp). It has been reported that using $\delta \rho = \rho - \rho_0$ instead of $\rho$ in moment space improves accuracy [5]. Thus we also experimented this approach for the three levels of precision: sp*, mp*, dp*.

Figures 9 and 10 show the average deviation from initial density when running a simulation at Re = 30,000, on a 384$^3$ lattice for the six levels of precision. We can see that, regarding conservation of mass, mixed precision does not provide significant improvement over single precision, and that sp*, mp*, and dp* perform better than dp by an order of magnitude. Furthermore, we may conclude that sp and mp should not be used when simulating unsteady flows.

In order to study the loss of symmetry from a quantitative standpoint, we used the following estimator:

$$L = \max_{x} \| u(x, t) - \bar{u}(\bar{x}, t) \|$$  (17)

where $x$ and $\bar{x}$, and $u$ and $\bar{u}$ are symmetric with respect of the symmetry plane of the cavity. Figure 11 shows the evolution of $L$ for the different precision levels running the same simulation than for mass conservation, i.e. Re = 30,000, on a 384$^3$ lattice. One can deduce from this diagram that the accumulation of round-off errors is the cause for the loss of symmetry. Past a certain threshold, due to the turbulent nature of the flow, the numerical perturbations are steeply amplified.

<table>
<thead>
<tr>
<th>Number of GPUs</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>6</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kernel duration (ms)</td>
<td>18.29</td>
<td>9.14</td>
<td>6.10</td>
<td>4.57</td>
<td>3.05</td>
</tr>
<tr>
<td>Exchanged data (MB)</td>
<td>0.00</td>
<td>1.47</td>
<td>2.95</td>
<td>4.42</td>
<td>7.37</td>
</tr>
<tr>
<td>Required throughput (GB/s)</td>
<td>0.00</td>
<td>0.16</td>
<td>0.48</td>
<td>0.97</td>
<td>2.42</td>
</tr>
</tbody>
</table>

Table 1: Required throughput for data exchange at 100% efficiency

Figure 9: Mass conservation issue (large scale)

Figure 10: Mass conservation issue (small scale)

Figure 11: Evolution of $L$ for the six precision levels

Figure 12 displays the evolution of $L$ at different
Reynolds numbers for the DP* precision level on a 384^3 lattice. This diagram shows that the more turbulent the flow pattern is, the sooner the symmetry breaking occurs, which corroborates the former point of view.

From a performance standpoint, SP, MP, and DP behave similarly than their stared counterparts, since the difference in implementation only affects the initialisation section. Using $\delta \rho$ instead of $\rho$ is therefore an advisable improvement. Mixed precision has almost identical performance than double precision. In this case, the gain in accuracy is not worth the performance trade-off.

7. Conclusion

In this contribution, we describe a multi-GPU implementation of the LBM, based on rather simple technical choices, i.e. POSIX threads and basic domain tilling. Nevertheless, performance is nearly optimal, rivaling with the one of supercomputer or large cluster implementations. Further investigations are needed to improve understanding of the inter-GPU communication potential. Moreover, work has to be done to design some execution grid layout and domain decomposition compatible with MPI parallelization.

Our multi-GPU LBM solvers enables the use of large lattices, thus allowing direct numerical simulation of unsteady flows. We describe some numerical issues that arise at high Reynolds numbers and investigate the impact of different precision levels both on accuracy and performance.

The TheLMA framework we designed to implement our flow solver is meant to improve code reusability. We are currently developing several applications based on TheLMA, including a hybrid thermal solver and a LES solver. In near future, we plan to extend this framework to generic multi-GPU parallelization.

References


