Abstract

We present Sapporo, a library for performing high-precision gravitational N-body simulations on NVIDIA Graphical Processing Units (GPUs). Our library mimics the GRAPE-6 library, and N-body codes currently running on GRAPE-6 can switch to Sapporo by a simple relinking of the library. The precision of our library is comparable to that of GRAPE-6, even though internally the GPU hardware is limited to single precision arithmetics. This limitation is effectively overcome by emulating double precision for calculating the distance between particles. The performance loss of this operation is small (∼20%) compared to the advantage of being able to run at high precision. We tested the library using several GRAPE-6-enabled N-body codes, in particular with Starlab and phiGRAPE. We measured peak performance of 800 Gflop/s for running with 10^6 particles on a PC with four commercial G92 architecture GPUs (two GeForce 9800GX2). As a production test, we simulated a 32k Plummer model with equal mass stars well beyond core collapse. The simulation took 41 days, during which the mean performance was 113 Gflop/s. The GPU did not show any problems from running in a production environment for such an extended period of time.

1. Introduction

Graphical processing units (GPUs) are quickly becoming mainstream in computational science. The introduction of Compute Unified Device Architecture (CUDA, Fernando, 2004), in which GPUs can be programmed effectively, has generated a paradigm shift in scientific computing (Hoekstra et al., 2007). Modern GPUs are greener in terms of CO$_2$ production, have a smaller footprint, are cheaper, and as easy to program as traditional parallel computers. In addition, you will not have a waiting queue when running large simulations on your local GPU-equipped workstation.

Newtonian stellar dynamics is traditionally on the forefront of high-performance computing. The first dedicated Newtonian solver (Applegate et al., 1986) was used to study the stability of the solar system (Sussman and Wisdom, 1992). And soon even faster specialized hardware was introduced by the inauguration of the GRAPE family of computers, which have an impressive history of breaking computing speed records (Makino and Taiji, 1998).

Nowadays, the GPUs are being used in various scientific areas, such as molecular dynamics (Anderson et al., 2008; van Meel et al., 2008), solving Kepler’s equations (Ford, 2009) and Newtonian N-body simulations. Solving the Newtonian N-body problem with GPUs started in the early 2000s by adopting a shared time step algorithm with a 2nd order integrator (Nyland et al., 2004). A few years later this algorithm was improved to include individual time steps and a higher order integrator (Portegies Zwart et al., 2007), in a code that was written in the device specific language Cg (Fernando and Kilgard, 2003). The performance was still relatively low compared to later implementations in CUDA via the Cunbody package (Hamada and Iitaka, 2007), Kirin library (Belleman et al., 2008), and the Yebisu N-body code (Nitadori and Makino, 2008; Nitadori, 2009). The main problem with the two former implementations was the limited accuracy of the GPU, which only enabled single precision. In part this problem was solved with the introduction of the double precision GPU (GTX280), but at a dramatic performance-hit as only a limited number of processor pipelines supported double-precision calculation.

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1http://www.nvidia.com/object/product_tesla_s1070_us.html
In this paper we introduce Sapporo\textsuperscript{2}, a library, which is written in CUDA, for running gravitational N-body simulations on NVIDIA GPUs. The Sapporo library closely matches the calling sequence of the GRAPE-6 library [Makino et al. 2003], and codes which are already running on a GRAPE-6 can be immediately be run on GPUs without any modifications to their source code.

Here, we describe the implementation of our library, and present both accuracy and performance measurements using Sapporo. For the latter, we use two direct N-body simulations environment, one being Starlab (Portegies Zwart et al. 2001) and the other is phiGRAPE (Harfst et al. 2007) as it is implemented in the Multi Scale Software Environment (MUSE, Portegies Zwart et al. 2009).

Our new implementation of the N-body force calculation in CUDA is an important step towards high-precision direct N-body simulations using GPUs, as the library is more flexible and more general than previous implementations. In addition we identify the operations in the code where double precision accuracy is most important and implement double precision arithmetic in those locations. The cost for doing this is limited to a $\sim 20\%$ loss in performance (but overall performance is still very high).

2. Implementation

We have designed a library to calculate gravitational forces on a GPU in order to accelerate N-body simulations. The library can be used in combination with a standard 4th-order predictor-corrector Hermite integration scheme (Makino & Aarseth 1992), either with shared or block time steps. Such a scheme consists of three essential steps: predictor, force calculation, and corrector. In the predictor step, the positions and velocities of all particles are predicted for the next time step. Then, the gravitational accelerations and their first derivatives (jerks) are calculated using the predicted positions and velocities. Finally, the predicted positions and velocities are corrected using the newly computed accelerations and jerks. In the case of a block time step scheme, the last two steps are only executed on a block of active particles. In the following, we will focus on block time steps, since shared time step scheme can be considered a variant of the block time step scheme.

In the case of the block time steps, the system is divided into $i$-particles and $j$-particles, which are the sinks and sources of the gravitational forces\textsuperscript{3}, respectively. Within the Hermite scheme, we take the following subsequent steps: we predict the positions and velocities of the $i$- and $j$-particles (predictor step); calculate the gravitational forces exerted by the $j$-particles on the $i$-particles (calculation step); correct the positions and velocities of the $i$-particles (corrector step).

The actual parallelisation of the force calculation is not too difficult and many examples for parallel N-body algorithms exist (Gualandris et al. 2007; Dorband et al. 2003). Implementing the algorithm on a GPU is a little more challenging due to the specific design of GPUs. For example, a G80/G92 NVIDIA GPU consist of 16 multi-processors (MPs) each being able to execute up to 768 threads in parallel. A parallel, SIMD (single instruction multiple data), element on such a GPU is called a warp. A warp is a set of 32 threads which execute the same instruction but operate on different data, and up to 24 warps can be executed in parallel on each MP. In total, a single G80/G92 GPU is able to execute up to 12288 threads in parallel, which means that a program which runs on a GPU, called a kernel, should be able to efficiently exploit such a high degree of parallelism.

In addition, the GPU design impose a strict memory access pattern on such a kernel. For example, a GPU has two types of memory: global memory, which is an equivalent of RAM on the CPU, and the shared memory, which is equivalent to L1-cache on the CPU. The global memory is further subdivided into local memory which is allocated on a per thread basis, and both texture and constant cached read-only memory. The major difference between the shared and the global memory is the access latency. Access to an element of the shared memory is as fast as access to a register, whereas access to an element in the global memory has a latency of 400-600 cycles. Since the data initially reside in the global memory, each of the parallel thread should cooperatively load the data to the shared memory in a specific pattern in order to reduce the access latency. The threads can then efficiently operate on the data in the shared memory.

\textsuperscript{2}The Sapporo library is free and can be downloaded from http://modesta.science.uva.nl.

\textsuperscript{3}In the following, by gravitational forces we imply gravitational accelerations, potentials and jerks.
Figure 1: Decomposition of the tasks between the CPU and the GPU. The CPU is responsible for the communication, prediction of the \(i\)-particles and the correction step, whereas the prediction of the \(j\)-particles and the force calculation step are carried out on the GPU.

2.1. Task decomposition between the CPU and the GPU

At first, we estimate the complexity of each of the steps in the Hermite scheme. We assume that the number of \(j\)-particles is equal to \(n\), and the number of \(i\)-particles is equal to \(m\). The time complexities of the predictor, the calculation, and the corrector steps are \(O(n)\), \(O(nm)\), and \(O(m)\), respectively. It is important to note, that the \(n^2\)-scaling of the force calculation has been reduced to \(nm\) by the introduction of block time steps. As a result, the time contribution of the calculation and of the corrector steps decreases with block size, whereas that of the prediction step remains constant. This is important as block sizes are typically small on average in direct N-body simulations.

Motivated by this, we implemented both the prediction of \(j\)-particles and the force calculation on the GPU. The prediction of \(i\)-particles and the corrector step are carried out on the CPU (Fig. 1); in fact, GRAPE-enabled N-body codes have exactly the same decomposition. Another motivation for this decomposition is the communication overhead: if the predictor step would be carried out on the CPU, all \(n\) particles would have to be communicated to the GPU at every time-step. In this case, communication would become a bottle-neck for the calculation. In our chosen implementation, only \(i\)-particles need to be communicated, and this therefore decreases the communication overhead.

2.2. Predictor step

In this step, the GPU’s task is to predict positions and velocities of the \(j\)-particles. It is a rather trivial step to parallelise since a particle does not require information from other particles. Hence, we assign a particle to each of the parallel threads on the GPU. Each thread reads for its particle position \((x)\), velocity \((v)\), acceleration \((a)\), jerk \((j)\), and time step \((dt)\) from global memory into registers. This data is required to execute the predictor step:

\[
x_{\text{pred}} = x_0 + v_0 dt + a_0 \frac{dt^2}{2} + j_0 \frac{dt^3}{6},
\]

\[
v_{\text{pred}} = v_0 + a_0 dt + j_0 \frac{dt^2}{2},
\]

where subscripts “0” and “pred” refer to the initial and predicted values, respectively; both the predicted position and velocity are stored in the global memory for later use. The prediction of the \(i\)-particles is similarly implemented on the CPU.

Prediction of particle positions must be computed in double precision (DP) arithmetics because the integration scheme is most sensitive to round-off errors in positions (Aarseth, 1985). This creates difficulties for implementing the predictor on the GPU due to the following reason: previous generations of NVIDIA GPUs (G80/G92) do not natively support DP arithmetics, and the current generation (GT200) executes DP instructions an order of magnitude slower than their single precision (SP) equivalents. However, not all operations in Eq. 1 require double precision. Only the position is stored in DP, whereas the rest of the terms are stored in SP. Consequently, the multiplications and divisions can be carried out in SP, but the summation should be carried out in DP. This can be done most efficiently by emulating DP only where it is necessary, instead of implementing all of Eq. 1 in DP. As a result, the loss in performance is acceptable and the implementation is not dependent on hardware supporting DP arithmetics.

The memory storage requirement for DP float (64 bit) is twice that of a SP float (32 bit). Therefore, it is natural to store DP numbers in two SP numbers: one of the SP numbers stores the most significant
Figure 2: An illustration of the decomposition between multiprocessors (MP) on the GPU. The $n$ $j$-particles are equally distributed between all $P$ MPs, where $P$ is the number of MPs. An identical copy of all $m$ $i$-particles is distributed to each of the MPs. As a result, every MP computes gravitational forces from $n/P$ bodies on the same set of $i$-particles in parallel.

digits, the other stores the least significant digits. Such a representation of DP is known as a double-single (DS)\(^4\). In this representation, the number of significant digits is 14 compared to 16 in DP\(^5\), but nevertheless it is a factor of two larger than in SP.

In the following, we will use DS to emulate DP where it is required. In Eq. 1 this only needs to be done for the positions $x_0$ and $x_P$. All multiplications and divisions are carried out and have their results stored in SP. These SP numbers are then added together and the resulting number is added to the DS number\(^6\) $x_0$. The result is stored again in DS as $x_P$. Therefore, the floating point operation (FLOP) count for Eq. 1 is 1 DS + 10 SP operations or 11 FLOP per particle; the Eq. 2 requires only 6 FLOP per particle.

2.3. The calculation step

We use predicted positions and velocities of the $j$-particles to calculate gravitational forces that the $j$-particles exert on the $i$-particles. We schematically depict the parallelisation strategy for such problem in Fig. 2. As before, we assume that the total number of the $j$-particles is equal to $n$ and the total number of the $i$-particles is equal to $m$.

We split the problem in $P$ parallel blocks, where $P$ is the number of parallel multiprocessors (MP) on the GPU. The $j$-particles are distributed evenly among the $P$ MP's. Each of the MP's then computes the

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4. \url{http://crd.lbl.gov/~dhbailey/mpdist/}
5. The mantissa of DP consists of 53 bits, compared to 48 bits in DS because a SP number has a mantissa of 24 bits.
6. Addition of DS with SP operation requires 10 SP floating point operations (FLOP)
Figure 3: Illustration of the decomposition on one multiprocessor. 1) Each thread reads in parallel one of the \( n/P \) particles to shared memory; the total number of shared-particles is therefore equals to \( m \). 2) These shared particles are processed sequentially. Steps 1) and 2) are repeated until all the \( n/P \) particles have been processed.

Figure 4: Illustration of the decomposition on a multiprocessor in the case when the number of \( i \)-particles is equal to half the number of parallel threads that can be executed simultaneously. In this case, every particle is processed by two threads in order to guarantee that all threads are active. However, every thread processes half of the \( j \)-particles that are associated with this MP.

partial gravitational forces, exerted by the \( n/P \ j \)-particles assigned to that MPs, on all the \( i \)-particles in parallel. This is accomplished by assigning each of the \( m \) parallel threads one of the \( i \)-particles in the block. Here, we assume that the number of \( i \)-particles \( m \), is smaller than or equal to the maximal number of parallel threads that a block is able to execute. Should \( m \) be larger, we split the \( i \)-particles in segments such that each of the segments contains the maximal number of \( i \)-particles that can be processed in parallel; the segments themselves are processed in a serial manner.

The \( P \) parallel blocks have no means of communication with each other, whereas the threads within a multiprocessor are able to exchange information via shared memory, which can be considered an equivalent of an on-chip cache memory. Therefore, once all the \( i \)-particles are processed, partial accelerations from each of the blocks must be accumulated.

The parallelisation method that we use in each block is illustrated in Fig. 3. Each thread in a block loads one particle from the global memory to the shared memory, and therefore the total number of particles stored in shared memory (shared particles) equals to the number of threads in a block. Afterwards, each thread sequentially calculates and sums the partial gravitational forces from the shared particles. Once completed, this loop is repeated until all of \( n/P \)-particles in the block have been processed.

If the number of the \( i \)-particles is smaller than the maximal number of threads that a block can execute, it is desirable to further parallelise the algorithm; we show this schematically in Fig. 4. Here, we assume that the number of \( i \)-particles \( m \), is smaller than or equal to the maximal number of \( i \)-particles in parallel. Thus, we split the \( n/P \) \( j \)-particles in two equal parts, such that each thread processes \( n/(2P) \) particles. In this way, the load per thread decreases by a factor of two, but since the total number of the \( i \)-particles that are processed concurrently doubles, the performance is unaffected. If we were to refrain from this parallelisation step, the performance would suffer since there would not be enough \( i \)-particles to fully occupy the multiprocessor, and half of the parallel threads would remain idle.

The final step is to sum up the partial forces computed by each of the \( P \) parallel blocks. A simple approach, in which all the data from the GPU memory is first copied to the host and then reduced using the CPU, is sufficiently efficient (Nitadori, 2009). Here, we have chosen to reduce the partial forces directly in the GPU memory and only copy back the final result.

The gravitational accelerations and jerks are sensitive to a round-off errors which occur during the
calculation of distances between two particles. We decrease these round-off errors by calculating this distances using DS arithmetics. This can be understood considering the following example: if the first four significant digits in the position of the two particles are the same and the positions are stored in SP, the resulting distance has only three significant digits. The use of such low accuracy result degrades the quality of the accelerations and jerks, and the accuracy of the simulation overall. If, however, the positions are stored in DS, the number of significant digits in the distance is nine instead of three, and this is sufficient for an accurate integration. Since the rest of the calculations is carried out in SP, the significant digits beyond the seventh can be discarded, implying that the separation, even though computed in DS, can be stored in SP.

The actual implementation is quite simple. One may bluntly apply formulae to calculate the difference between two DS numbers. Such method is expensive as it requires approximately ten SP floating point operations to subtract two DS. However, such operation takes proper care of both the most and the least significant digits in the result. However, for our purpose, we can simplify the subtraction by discarding the least significant digits. In this way, we reduce the number of FLOPs. If the positions of two particles are stored in DS format, \( x_i = \{x_i,hi; x_i,lo\} \) and \( x_j = \{x_j,hi; x_j,lo\} \), the separation \( \Delta x_{ij} \) is

\[
\Delta x_{ij} = (x_j,hi - x_i,hi) + (x_j,lo - x_i,lo).
\]

(3)

The number of FLOPs required to carry out this difference is equal to 3. With this implementation, we will be able to resolve particles separations to one part in ten millions or better.

This is the only operation that is carried out in DS arithmetics. Since it is usually assumed that it takes 60 FLOPs to calculate gravitational accelerations and jerks, the number of FLOPs that our method requires is therefore equals to 66. Here, we substitute each of the three SP subtraction operations with Eq.3.

2.4. Parallelization over multiple GPUs

Our library has a built-in support for multiple GPUs installed on a single host PC. The support of multiple GPUs is done with the help of the GPUWorker library which is a part of the HOOMD molecular dynamics GPU-code (Anderson et al., 2008). The parallelisation is carried out automatically, based on the availability of multiple GPUs and the user request given in a configuration file. The application that uses the library is unaware of this, meaning that no modifications to source code are required for using multiple GPUs on one host. The parallelisation strategy in Sapporo is rather straightforward: given a number \( Q \) of GPUs, the library distributes \( n \) of the \( j \)-particles equally between all of the GPUs. As a result, each of the GPUs processes \( n/Q \) of \( j \)-particles, but the same set of \( i \)-particles. Using multiple GPUs results in a speed up for sufficiently large \( n \), otherwise the smaller number of \( j \)-particles per GPU may result in a performance loss.

2.5. Interface

We designed the library with the same application interface (API) as the standard GRAPE6 library. To replicate the GRAPE-6 functionality, Sapporo supports softening per block of \( i \)-particles and searches for nearest neighbours for each of the \( i \)-particles. This makes Sapporo suitable for high-precision collisional \( N \)-body simulations. The swapping of libraries during the linking process allows existing applications which already support the GRAPE6 API to be used directly with Sapporo without any changes to the software. Compilation of Sapporo does require the presence of a CUDA-enabled GPU on the host computer as well as the CUDA run-time libraries. From our experience, GRAPE6-enabled \( N \)-body codes, such as Starlab (Portegies Zwart et al., 2001), phiGRAPE or phiGRAPEch (Harfst et al., 2008), and NBODY4 (Aarseth, 1999) can be used with the Sapporo library without any modifications to their source code.

3. Results

We measure the performance of Sapporo by integrating equal mass Plummer (Plummer, 1915) spheres with a various number of particles, \( N \), for a fraction of an \( N \)-body time unit (Heggie and Mathieu, 7

\[7\] The number of pipelines in Sapporo is 256 instead of 48 for the GRAPE6, which may require resizing some arrays if an \( N \)-body code is written in a programming language with static memory allocation.
These experiments were performed using two different codes for high precision collisional N-body simulations, Starlab (Portegies Zwart et al., 2001) and phiGRAPE (Harfst et al., 2007). We also performed an integration of a $N = 32^k$ Plummer sphere well beyond core collapse.

The host computer used for these tests is equipped with one Intel Core2 Quad CPU operating at 2.5GHz, 2GB of RAM, and two NVIDIA GeForce 9800GX2 graphics cards. Each of these cards is equipped with two independent G92 GPU chips. For the user the system appears to be equipped with four independent GPUs. We used the NVIDIA CUDA driver v177.67, CUDA Toolkit v1.1 and CUDA SDK v1.1 and the installed operating system is Debian GNU/Linux 4.0 with the 2.6.21 SMP x86_64 kernel. The library is also compatible with CUDA Toolkit v2.1.

For comparison, some of the calculations were repeated on GRAPE-6A, for which we used eight nodes of the Rochester Institute of Technology cluster (Harfst et al., 2007). Each of these nodes is equipped with a GRAPE-6A. The code phiGRAPE runs in parallel over several nodes on this cluster, whereas in the GPU setup several GPUs are hosted by a single PC. Some difference in performance can therefore be attributed to the inter-node communication on the GRAPE cluster, which is negligible on the single GPU-equipped PC. For comparing the accuracy, we have also done a number of calculations on a single PC with GRAPE-6A, which we refer to as GRAPE PC. This PC is equipped with a 2.8GHz Pentium D processor with 1GB RAM and a GRAPE-6A PCI card, and running Debian 4.0 GNU/Linux with the 2.6.18 SMP x86_64 kernel.

3.1. Performance

The results of the performance tests are presented in Fig. 5, where we show the wall-clock time as a function of the number of particles, $N$, for 1, 2 and 4 GPUs. In all the cases, the wall-clock time scales as expected, with $N^2$ and the wall-clock time decreases as the number of GPUs increases. Simulations with $N \leq 64k$ on 4 GPUs show a degradation in the speed-up, which is caused by the smaller number of particles in each block per GPU, see (2.4) and we notice similar behaviour for the GRAPE-6A cluster (Harfst et al., 2007). The performance of two GRAPE-6A-nodes is comparable to that of a single GPU.

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Figure 5: The wall-clock time as function of the number of particles using GPUs (solid line with bullets) or GRAPE-6A (dotted line with stars). The bullets and stars are measured from integrating a Plummer models with $N$ particles using phiGRAPE. From top to bottom, one, two, or four GPUs and one, two, four and eight GRAPEs were used. The dashed-dotted line shows the expected $N^2$-scaling, offset not to overlap with the measurements.

1986[5].

http://en.wikipedia.org/wiki/Natural_units#N-body_units
In Fig. 6 we show the sustained performance in GFlop/s for a range of \( N \). The performance of the GPU is better than that of the GRAPE-cluster by about a factor of two. The relatively slow PCI-bus used in the GRAPE nodes causes the performance for very small \( N \) to be even worse compared to the GPU-equipped PC.

The average number of particles in a block time-step can also be used as a diagnostic tool. In phiGRAPE the block size is controlled by the standard time step criterion (Aarseth, 1985). This criterion includes a term, that depends on the difference in force on a the particle at the beginning and end of a time step. This can have an effect on time steps (and therefore block size) if the forces are calculated with low precision. We found that small time steps tend to become much smaller in this case, as the relative error in the force difference becomes larger (when time steps are small, the force is usually large but it changes very little over this time step, resulting in the same problem as discussed before for calculating the distance of two particles). Smaller time steps generally result in smaller block sizes, which in turn results in an inefficient use of the GPU and also increases the number of integration steps needed.

In Fig. 7 we present the average block size as a function of \( N \) for GRAPE-6 as well as for GPU. The difference between the block size for GRAPE and GPU is small for the adopted standard time-step criterion, which indicates that the accuracy in the force calculation on the GPU is comparable to that of GRAPE. It is important to note, that the emulation of double precision is the key for achieving this accuracy.

### 3.2. Accuracy

The accuracy of the force evaluation and integration of the equations of motion is crucial for a reliable \( N \)-body simulation. We therefore test in this section the adopted method for achieving high accuracy on the GPU hardware and compare this with GRAPE-6. There are various sources of error in any \( N \)-body integration: the most straightforward comes from the integration scheme itself and depends on the order of the integrator; another error is caused by the limited (single) precision of the GPU. The error in the integration is controlled via the accuracy parameter \( \eta \) (Aarseth, 1985), such that smaller \( \eta \) results in a

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9The exact number of floating point operations required for one force calculation varies in the literature. Here we assumed that a single force calculation requires 60 operations, which was also used for determining the theoretical peak performance of GRAPE (Fukushige et al., 2003)
lower energy error. For a sufficiently small $\eta$ the integration errors are dominated by the limited precision of the hardware.

To test accuracy of Sapporo we integrated an equal mass Plummer models with various $N$. For every $N$ we studied the dependence of the relative energy error, $dE/E = (E_1 - E_0)/E_0$, as a function of the accuracy parameter, $\eta$, and the number of particles, $N$. For this purpose, we varied $\eta$ and $N$ between $10^{-4}$ to 0.3 and 16k to 256k respectively. Our results show that the error is strongly dependent on $\eta$, whereas dependence on $N$ is rather weak. In Fig. 8 we show the results for $N = 32k$ and $N = 128k$.

For $\eta \gtrsim 3 \cdot 10^{-3}$, the relative errors $dE/E$ decreases with $\eta$ and the difference between GRAPE and GPU is indistinguishable independent of $N$ (see Fig. 8). This behaviour is expected, since for relatively high $\eta$ the energy error is dominated by the integration errors rather than by the errors in the forces evaluation. For $\eta \lesssim 3 \cdot 10^{-3}$, however, the integration error saturates at the point where the hardware limits the accuracy of the force evaluation. As expected for $\eta \lesssim 3 \cdot 10^{-3}$ the error $dE/E$ does not depend on $\eta$, although the saturation level is weakly dependent on $N$. On the GPU, the relative error is $dE/E \simeq 10^{-9}$ due to the limited precision, which is of order of the minimal relative error that can be achieved with single precision. The errors produced by the GRAPE-6 saturates at a somewhat lower level $dE/E \simeq 10^{-11}$, which is expected based on the higher accuracy of the internal GRAPE hardware \cite{Makino and Taiji, 1998}. Note, that typical values for $\eta$ are between $10^{-2}$ and $10^{-1}$, and therefore well within the range where Sapporo is not limited by the single precision accuracy of the GPU.

### 3.3. Star cluster evolution and core collapse

We tested Sapporo in a production environment by calculating the time evolution of an equal-mass Plummer model with $N=32k$ particles through core collapse. The integration was performed using Starlab N-body simulation environment, and we used only one GPU for this calculation.

The results of this calculation are presented in Fig. 9 where we show time evolution of some cluster parameters. In particular, it is evident that the cluster reaches core collapse at $t \simeq 6440$ N-body time units. The wall-clock time it took to reach core collapse was about 24 days, whereas it took the simulation only 9 days to reach the point half-way to core collapse. The whole simulation, which we terminated at $t \simeq 9200$ N-body time units, was completed in about 41 days. We note, that a relatively inexpensive commercial GPU, which we used for this experiment, proved to be stable during the entire simulation.

The sustained performance up to $t \simeq 3000$ N-body time units was about 140 Gflop/s, which reduced to about 132 Gflop/s by the time core collapse was reached, and to drop even further to about 88 Gflop/s in the post core collapse phase. The lower performance after core collapse is mainly caused by the
Figure 8: The relative error in total energy as a function of the time-step accuracy parameter $\eta$ for a Plummer model with 32k (left panel) and 128k (right panel) particles were integrated for $1/4 N$-body time units using phiGRAPE. The dotted line with open squares shows the energy error for the GPU with Sapporo, the dashed line are from GRAPE-6A.

Figure 9: The left panel shows time evolution of the core radius (black line) and 1, 2, 5, 10, 25, 50, 75, 90% lagrangian radii (green lines, from bottom to top). The right panel shows the time evolution of both the number of stars in the core (top) and core density (bottom). The initial condition was a equal-mass Plummer distribution with $N = 32768$ particles. The core collapse is reached at $t \simeq 6440$. The calculation was terminated at $t = 9200$. The average sustained speed for the entire calculation was about 113 Gflop/s. The sudden decrease of lagrangian radii at $t \simeq 6800$ is caused by the removal of the escaped stars.
presence of a dense core and hard binaries, which formed during core collapse and which require additional calculations performed on the host. The mean sustained performance for the entire simulation was about 113 Gflop/s.

4. Summary

The integration of a large self gravitating system for many relaxation times is a computationally demanding procedure. Currently, the GRAPE6 special-purpose hardware is the state-of-the-art for carrying out such simulations. In this paper, we have presented a library named Sapporo, which emulates the GRAPE6 functionality on NVIDIA GPUs using the CUDA programming environment. Our library is publicly available and we emulate GRAPE6 interface as closely as possible (with the exception of some rarely used functions). The library is designed in such a way that it can be simply swapped with the original GRAPE6 library and, as a result, a wide range of current N-body programs, that have been written using the GRAPE6 interface, can now be used on GPUs using Sapporo without any modification of the source code.

We have carried out a number of basic N-body experiments in order to test the efficiency and accuracy of the library and we compared our results with the GRAPE6. We found that our library is twice as fast as commonly used GRAPE6A/GRAPE6-BLX cards, and it is as accurate as GRAPE with standard integration parameters. In particular, the performance of a PC equipped with two NVIDIA GeForce 9800GX2 cards is on par with that of a 8-node GRAPE6A cluster or a 32-chip GRAPE6 board.

We have also carried out a production run to test the ability of the library to handle a real astrophysical problem. For such stress-test, we have chosen to integrate an equal-mass Plummer model beyond core collapse. As expected, the cluster reaches core collapse in about 15 initial half-mass relaxation times and afterwards enters the phase of gravothermal oscillations. Our results are consistent with those previously published in the literature [Heggie and Hut, 2003]. The ability of Sapporo to handle such a demanding problem demonstrates that the library can be safely used for realistic astrophysical problems.

The high performance of the Sapporo library is achieved by splitting the steps of the Hermite integration scheme between CPU and GPU in the same way as it is done when using the GRAPE6. In particular, the prediction and calculation steps are carried out on the GPU, whereas the corrector is computed on the CPU. In this way, the force calculation is always a dominant part of the integration. If the prediction for all particles is carried out on the host, as it is done in the kirin library [Belleman et al., 2008], the integration would be dominated by the prediction step and communication, and thus degrading the performance.

It is common nowadays, that a single production node is equipped with at least two GPUs. Therefore, we designed the library to make an efficient use of such a configuration, and the user can configure the library to use any combination of available GPUs. The application which makes use of the library is unaware that multiple GPUs are utilised, i.e. no modification to its source code is required.

We also implemented a neighbour lists functionality in Sapporo. The maximum number of neighbours that a particle in a block can store is configured at compile time (default is 256), and it cannot be changed during runtime. Calculating and storing neighbour lists has an impact on performance of a few tens of percent, depending on the maximum number of neighbours allowed. On the other hand, it is also possible to disable neighbour lists all together, and by this gain up to 50% in speed.

An important part of Sapporo is the emulation of double precision using single precision numbers by using standard methods from the literature. Similar techniques were implemented in the Yebisu N-body code which also runs on NVIDIA GPUs [Nitadori, 2009]. The partial emulation of double precision has a minimal impact on performance, and places current and future NVIDIA GPUs on a very competitive ground with current and future GRAPE hardware.

The emulation of double precision was required because the previous generation of NVIDIA GPUs (based on G80/G92 architecture) does not support it. Current NVIDIA GPUs (based on GT200 architecture) have native double precision support, however, only for a limited number of pipelines. This enables in principle the use of double precision, but the performance is an order of magnitude smaller in that case. The GT200 architecture also has almost three times as many parallel elements and twice the theoretical peak-performance of the previous G80/G92 architectures. Using a GT200 GPU, we were able

10http://www.nvidia.com/object/product_tesla_s1070_us.html
to get a noticeable speedup of about 50%, which is smaller than the factor of two in peak performance. The reason is, that Sapporo was designed for the older GPU architectures, and therefore does not in the current version efficiently utilise all the parallel elements of a GT200 GPU. However, we are currently working on a new version of Sapporo which will be scalable across future generations of NVIDIA GPUs, including the GT200.

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