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2 Cosmological simulations on a grid of supercomputers

Based on:
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High performance cosmological simulations on a grid of supercomputers
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We present results from our cosmological $N$-body simulation which consisted of $2048^3$ particles and ran distributed across three supercomputers throughout Europe. The run, which was performed as the concluding phase of the GBBP DEISA project, integrated a 30Mpc box of dark matter using an optimized TreePM $N$-body integrator. We ran the simulation up to the present day ($z=0$), and obtained an efficiency of about 0.93 over 2048 cores compared to a single supercomputer run. In addition, we share our experiences on using multiple supercomputers for high performance computing and provide several recommendations for future projects.
2.1 Introduction

Cosmological simulations are an efficient method to gain understanding of the formation of large-scale structures in the Universe. Large simulations were previously applied to model the evolution of dark matter in the Universe (Springel et al. 2005), and to investigate the properties of Milky-Way sized dark matter halos (Springel et al. 2008; Ishiyama et al. 2009a). However, these simulations are computationally demanding, and are best run on large production infrastructures. We have previously run a cosmological simulation using two supercomputers across the globe (Portegies Zwart et al. 2010a) with the GreeM integrator, and presented the SUSHI N-body integrator, which we used to run simulations across up to four supercomputers. The simulations we ran in the GBBP produced over 110 TB of data, which we have used to characterize the properties of ultra-faint dwarf galaxies (Ishiyama et al. 2013), and to compare the halo mass function in our runs to analytical formulae for the mass function. Among other things, we found that the halo mass function in our runs shows good agreement with the Sheth and Tormen (1999) mass function down to $\sim 10^7 M_\odot$.

Here we present the performance results of a production simulation across three supercomputers, as well as several other runs which all use an enhanced version of SUSHI. The production simulation ran continuously for $\sim 8$ hours, using 2048 cores in total for calculations as well as 4 additional cores for communications. We achieved a peak performance of $3.31 \times 10^{11}$ tree force interactions per second, a sustained performance of $2.19 \times 10^{11}$ tree force interactions per second and a wide area communication overhead of less than 10% overall.

We briefly reflect on the improvements made to SUSHI for this work in section 2.2, while we report on tests performed on a single supercomputer in section 2.3. In section 2.4 we describe our experiments across three supercomputers and present our performance results. We reflect on our experiences on using multiple supercomputers for distributed supercomputing simulations, and provide several recommendations for users and resource providers in section 2.5 and present our conclusions in section 2.6.

2.1.1 Related work

There are several other projects which have run high performance computing applications across multiple supercomputers. These include simulations of a galaxy collision (Norman et al. 1996), a materials science problem (Pratt et al. 1997) as well as an analysis application for arthropod evolution (Stewart et al. 2004). A larger number of groups performed distributed computing across sites of PCs rather than supercomputers (e.g. Gualandris et al. 2007; Bal and Verstoep 2008;
Bar et al. 2009). Several software tools have been developed to facilitate high performance computing across sites of PCs (e.g. Karonis et al. 2003; Gabriel et al. 1998, 2004; Manos et al. 2008; Sundari M. et al. 2010)) and within volatile computing environments (Rood et al. 2010). The recently launched MAPPER EU-FP7 project (map 2012) seeks to run multi-scale applications across a distributed supercomputing environment, where individual subcodes periodically exchange information and (in some cases) run concurrently on different supercomputing architectures.

2.2 Improvements to SUSHI

Based on results of our earlier simulations and in preparation for the production run across three supercomputers we made several modifications to the SUSHI distributed $N$-body integrator. In our previous experiments a relatively large amount of computation and communication time was spent on (non-parallelised) particle-mesh integration. To reduce this bottleneck we now parallelised the particle-mesh integration routines using the parallel FFTW2 library and a one dimensional slab decomposition. We also optimized the communications of the particle-mesh integration by introducing a scheme where sites only broadcast those mesh cells which have actual particle content. This optimization reduced the size of the mesh communications by a factor roughly equal to the number of sites used, in the case of an equal domain distribution.

In some of the larger previous runs we also observed load imbalances if the code was run across two machines with different architectures, despite the presence of a load balancing scheme. This result has led us to further optimize the load balancing in SUSHI, taking into account not only the force integration time, but also the number of particles stored on each node. In addition to these changes, we also seized the opportunity to plug in a more recent MPWide version into SUSHI. This newer version contains several optimizations to improve the wide area communication over networks with a high latency.

2.3 Tests on a single site

2.3.1 Setup

We performed a number of runs on the Huygens supercomputer to validate the scalability of our new implementation, and to provide performance measurements against which we can compare our results using multiple sites. More information on the Huygens machine can be found in the second column of Table 2.3. The ini-
**Table 2.1**: Initial condition and accuracy parameters used for our simulations with 2048$^3$ particles.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Matter density parameter ($\omega_0$)</td>
<td>0.3</td>
</tr>
<tr>
<td>Cosmological constant ($\Omega_\Lambda$)</td>
<td>0.7</td>
</tr>
<tr>
<td>Hubble constant ($H_0$)</td>
<td>70.0 km/s/Mpc</td>
</tr>
<tr>
<td>Mass fluctuation parameter ($\sigma_8$)</td>
<td>0.8</td>
</tr>
<tr>
<td>Box size</td>
<td>(30Mpc)$^3$</td>
</tr>
<tr>
<td>Softening for 2048$^3$ particle run.</td>
<td>175 pc</td>
</tr>
<tr>
<td>Sampling rate for 2048$^3$ particle run.</td>
<td>20000</td>
</tr>
</tbody>
</table>

tial conditions for this simulation is the snapshot at redshift $z = 0.0026$ from the CosmoGrid simulation (described in Portegies Zwart et al. 2010a). We also use the simulation parameters chosen for the CosmoGrid simulation, which are summarized in Table 2.1. Here the first four parameters are constants, which are derived from WMAP observations (with a slight round-off), and the physical size of our simulated system is given by the fifth parameter (Box size). The softening in our simulation (i.e. a length value added to reduce the intensity of close interactions) and the sampling rate are given by the last two parameters. The sampling rate is the ratio of particles in the simulation divided by the number of particles sampled by the load balancing scheme. Our simulation used a mesh size of 512$^3$ cells. We ran the simulation using respectively 512 cores and 1024 cores until $z = 0.0024$, and using 2048 cores until the simulation completed (at $z = 0$). The number of force calculations per step in the simulation varies for different $z$ values, though these variations are negligible for $z < 0.01$.

### 2.3.2 Results

The performance results of our runs are shown in Table 2.2. In addition, the total runtime of the run using 2048 cores is given by the light blue line in Figure 2.1 (top). The overall performance of the code is dominated by calculations, with the communication overhead ranging from $\sim$5% for 512 cores to $\sim$10-15% for 2048 cores. During the run using 2048 cores, several snapshots were written. This resulted in a greatly increased execution time during two steps of the run.
2.4 Tests across three sites

We performed our main run using a total of 2048 cores across three supercomputers, which are listed in Table 2.3. These machines include Huygens in the Netherlands (1024 cores), Louhi in Finland (512 cores), and HECToR in Scotland (512 cores). The sites are connected to the DEISA shared network with either a 1Gbps interface (HECToR) or a 10Gbps interface (Huygens, Louhi). The initial conditions and simulation parameters chosen are identical to those of the runs using 1 supercomputer, although we use a mesh of $256^3$ cells. The use of a smaller mesh size results in a slightly higher calculation time as tree interactions are calculated over a longer range, but a somewhat lower time spent on intra-site communications. We configured MPWide to use 64 parallel TCP streams per path for the wide area communication channels, each with a TCP buffer size set at 768 kB and packet-pacing set at 10 MB/s maximum. We enabled some load balancing during the run, though we had to limit the boundary moving length per step to 0.00001 of the box length due to memory constraints on our communication nodes and the presence of dense halos in our initial condition.

In addition to the main run, we also performed three smaller runs using the same code across the same three supercomputers. These include one run with $1024^3$ particles using 80 cores per supercomputer, and two runs with $512^3$ particles using 40 cores per supercomputer. These runs also used a mesh size of $256^3$, though we did reduce the sampling rate to respectively 10000 and 5000 for the runs with $1024^3$ and $512^3$ particles. The force softening used for these runs were respectively 1.25kpc and 2.5kpc, and we set the boundary moving length limit to 0.01 of the box length. Some of the measurements were made using an opening angle $\theta$ of 0.3, rather than 0.5. Using a smaller opening angle results a higher accuracy of the

### Table 2.2: Overview of experiments performed with the enhanced SUSHI code on the Huygens supercomputer. The time spent on communication is given in the fourth column, while the total runtime is given in the fifth column. All times are measured per step, averaged over steps 1-11. In addition we included the timing results of the last 10 steps of the simulation running on 2048 cores (bottom row).

<table>
<thead>
<tr>
<th>$N$ $p$</th>
<th>$\theta$</th>
<th>comm. t [s]</th>
<th>runtime [s]</th>
<th>$z$ range $\times 10^{-3}$</th>
<th>speed-up</th>
</tr>
</thead>
<tbody>
<tr>
<td>$2048^3$</td>
<td>512</td>
<td>0.5</td>
<td>19.18</td>
<td>501.3</td>
<td>2.5-2.4</td>
</tr>
<tr>
<td>$2048^3$</td>
<td>1024</td>
<td>0.5</td>
<td>13.96</td>
<td>258.2</td>
<td>2.5-2.4</td>
</tr>
<tr>
<td>$2048^3$</td>
<td>2048</td>
<td>0.5</td>
<td>22.34</td>
<td>151.0</td>
<td>2.5-2.4</td>
</tr>
<tr>
<td>$2048^3$</td>
<td>2048</td>
<td>0.5</td>
<td>16.22</td>
<td>143.7</td>
<td>0.1-0.0</td>
</tr>
</tbody>
</table>
force integration on close range, but also results in a higher force calculation and tree structure communication time per step.

### 2.4.2 Results

The timing results of our production run are shown in Figure 2.1. Here, we also added the wall-clock time results of the simulation run using 2048 cores on Huygens as reference. The simulation run across three sites is only $\sim 9\%$ slower per step than the single-site run, despite the slightly higher force calculation time due to the lower number of mesh cells. The peaks in wall-clock time of the single site run are caused by the writing of snapshots during those steps (we only wrote one snapshot at the end of the three site run). The total wide area communication overhead of our run is $\lesssim 10\%$ at about 15s per step. Most of this time is required to exchange the tree structures between sites, though the communications for the parallelised particle-mesh require an additional $\sim 2.5s$ per step. Despite the use of a shared wide area network, the communication performance of our run shows very little jitter and no large slowdowns. We provide a snapshot of the final state of the simulation (at $z = 0$), distributed across the three supercomputers, in Figure 2.2.

We also provide a numerical overview of the production run performance, as well as that of several other runs which use the new code, in Table 2.4. The communication overhead for the runs with $512^3$ particles is less than 20%, while the overhead for the run with $1024^3$ particles is just 6.5%. The parallelisation of the particle-mesh integration and the enhanced load balancing greatly improved the performance of these runs, especially in the case with $1024^3$ particles. Here, the communication overhead was reduced by $\sim 60\%$ and the overall runtime by more than 25% compared to the previous version (Groen et al. 2011).
2.5 User Experiences

<table>
<thead>
<tr>
<th>$N$</th>
<th>$p$</th>
<th>$\theta$</th>
<th>comm. time</th>
<th>runtime</th>
<th>$z$ range</th>
</tr>
</thead>
<tbody>
<tr>
<td>512(^3)</td>
<td>120</td>
<td>0.3</td>
<td>6.925</td>
<td>7.312</td>
<td>39.70</td>
</tr>
<tr>
<td>512(^3)</td>
<td>120</td>
<td>0.5</td>
<td>5.982</td>
<td>6.335</td>
<td>24.60</td>
</tr>
<tr>
<td>1024(^3)</td>
<td>240</td>
<td>0.3</td>
<td>12.09</td>
<td>14.04</td>
<td>214.5</td>
</tr>
<tr>
<td>2048(^3)</td>
<td>2048</td>
<td>0.5</td>
<td>15.40</td>
<td>24.77</td>
<td>167.7</td>
</tr>
<tr>
<td>2048(^3)</td>
<td>2048</td>
<td>0.5</td>
<td>14.62</td>
<td>23.13</td>
<td>155.2</td>
</tr>
</tbody>
</table>

Table 2.4: Overview of experiments performed with the enhanced SUSHI code across all three supercomputers. All times are measured per step, averaged over 10 steps.

2.5 User Experiences

We have presented results from several cosmological simulations which run across three supercomputers, including a production run lasting for 8 hours. In the process of seeking a solution for wide area message passing between supercomputers, requesting allocations, arranging network paths and preparing for the execution of these simulations, we have learned a number of valuable lessons.

Primarily, we found that it is structurally possible to do high performance computing across multiple supercomputers. During the GBBP project we have run a considerable number of large-scale simulations using two or more supercomputers, with results improving as we were able to further enhance the $N$-body integrator and optimize the MPWide communication library for the wide area networks that we used.

The cooperation of the resource providers was particularly crucial in this project, as they enabled previously unavailable network paths and provided us with means to initiate simulations concurrently at the different sites. However, reserving networks and orchestrating concurrent supercomputer runs currently does require a disproportionate amount of time and effort, which makes performance optimization and debugging a challenging task. The effort required to run applications across supercomputers can be greatly reduced if resource providers were to adopt automated resource reservation systems for their supercomputers, and maintain shared high-bandwidth networking between sites. The persistent DEISA shared network connections helped greatly in our case, as we could use it at will without prior network reservations.

The software environment across different supercomputers, even within the same distributed infrastructure, is very heterogeneous. This made it unattractive to use existing middleware or message passing implementations to make different sites interoperable. We chose to use a modular approach where we connected platform-specific optimized versions of the SUSHI code with the MPWide com-
munication library. With MPWide being a user-space tool that requires no external libraries or administrative privileges, we are able to install and run the simulation code in the locally preferred software environments on each site without needing any additional (grid) middleware. We recommend adopting a similar modular software approach in future distributed supercomputing efforts for its ease of installation and optimization, at least until resource providers present a homogeneous and interoperable software environment for distributed supercomputing.

This paper focuses on the calculation and communication performance aspects of a single application run across supercomputers. However, the methods presented here can be applied for several other purposes. During this project we were confronted with additional overhead introduced by disk I/O, as can be observed in Figure 2.1. With supercomputer disk performance and capacity improving at a much slower rate than the compute power, the deployment of an application across sites may help to eliminate a disk I/O performance bottleneck, though a detailed investigation will be needed to quantify such potential benefit. Additionally, the communication technique could be used to facilitate periodic exchanges between different simulation codes, each of which runs on a different site and tackles a different aspect of a complex multi-scale or multi-physics problem.

2.6 Conclusion

Our results show that cosmological production simulations run efficiently across supercomputers for a prolonged time. The political effort required to arrange cross-supercomputer runs is considerable, and is an important reason why few people have attempted to run production simulations across supercomputers. We have shown that the added overhead of using a network of supercomputers is rather marginal for at least one optimized production application and that given the right (political) environment, supercomputers can be conveniently connected to form even larger high performance computing resources.

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2.6 Conclusion

**Figure 2.1:** Performance results of the production simulation across three sites. In the top figure we provide the total time spent on calculation per step in red, and on communication per step in blue. Here, the total wall-clock time of an identical simulation using 2048 processes only on Huygens is given by the light blue line. Time spent on the four communication phases is given in the bottom figure. These phases are (from top to bottom) the migration of particles between sites, the exchanges of sample particles for determining the site boundaries, the local essential tree exchanges (PP) and the mesh cell exchanges (PM). See Groen et al. (2011) for full details on the communication routines of the code.

**Figure 2.2:** The final snapshot of the production simulation across three supercomputers, taken at \( z = 0 \). The size of the box is 30x30x30 Mpc and the contents are coloured to match the particles residing in Espoo (green, left), Edinburgh (blue, centre) and Amsterdam (red, right) respectively.