University of Warsaw Lagrangian Cloud Model (UWLCM) 2.0: Adaptation of a mixed Eulerian-Lagrangian numerical model for heterogeneous computing clusters

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Abstract.

A numerical cloud model with Lagrangian particles coupled to an Eulerian flow is adapted for distributed memory systems. Eulerian and Lagrangian calculations can be done in parallel on CPUs and GPUs, respectively. The fraction of time when CPUs and GPUs work simultaneously is maximized at around 80% for an optimal ratio of CPU and GPU workloads. The optimal ratio of workloads is different for different systems, because it depends on the relation between computing performance of CPUs and GPUs. GPU workload can be adjusted by changing the number of Lagrangian particles, which is limited by device memory. Lagrangian computations scale with the number of nodes better than Eulerian computations, because the former do not require collective communications. This means that the ratio of CPU and GPU computation times also depends on the number of nodes. Therefore, for a fixed number of Lagrangian particles, there is an optimal number of nodes, for which the time CPUs and GPUs work simultaneously is maximized. Scaling efficiency up to this optimal number of nodes is close to 100%. Simulations that use both CPUs and GPUs take between 10 to 120 less time and use between 10 to 60 times less energy than simulations run on CPUs only. Simulations with Lagrangian microphysics take up to eight times longer to finish than simulations with Eulerian bulk microphysics, but the difference decreases as more nodes are used. Presented method of adaptation for computing clusters can be used in any numerical model with Lagrangian particles coupled to an Eulerian fluid flow.

1 Introduction

As CPU clock frequencies no longer stably increase over time and the cost per transistor increases, new modeling techniques are required to match the demand for more precise numerical simulations of physical processes (Bauer et al., 2021). We present an implementation of the cloud model UWLCM for distributed memory systems that uses some of the modeling techniques reviewed by Bauer et al. (2021): use of heterogeneous clusters (with parallel computations on CPU and GPU), mixed-precision computations, semi-implicit solvers, different time steps for different processes, portability to different hardware. Although we discuss a numerical cloud model, conclusions and the techniques used can be applied to modeling of other processes in which Lagrangian particles are coupled to an Eulerian field, such as the particle-in-cell method used in plasma physics (Hockney and Eastwood, 1988).
In numerical models of the atmosphere, clouds are represented using various approximations depending on resolution of the model. In large scale models, like global climate and weather models, clouds are described with a simplistic process, which is known as cloud parameterization. Cloud parameterizations are developed based on observations, theoretical insights and on fine scale numerical modeling. Therefore correct fine scale modeling is important for better understanding of Earth’s climate and for better weather prediction. Highest resolution numerical modeling is known as direct numerical simulation (DNS). In DNS, even the smallest turbulent eddies are resolved, which requires spatial resolution in the millimeter range. Largest current DNS simulations model a volume of the order of several cubic meters, not enough to capture many important cloud scale processes. Whole clouds and cloud fields can be modeled with the the Large Eddy Simulations (LES) technique. In LES, small scale eddies are parameterised, so that only large eddies, typically of the order of tens of meters, are resolved. Thanks to this, it is feasible to model a domain spanning tens of kilometers.

DNS and LES models of clouds need to resolve air flow, which is referred to as cloud dynamics, and evolution of cloud droplets, which is known as cloud microphysics. UWLCM is a tool for LES of clouds with a focus on detailed modeling of cloud microphysics. Dynamics are represented in an Eulerian manner. Cloud microphysics are modeled in a Lagrangian particle-based manner based on the Super-Droplet Method (SDM) (Shima et al., 2009). Lagrangian particle-based cloud microphysics models have gained popularity in the last decade (Shima et al., 2009; Andrejczuk et al., 2010; Riechelmann et al., 2012). These are very detailed models applicable both to DNS and LES. Their level of detail and computational cost are comparable to the more traditional Eulerian bin models, but Lagrangian methods have several advantages over bin methods (Grabowski et al., 2019). Simpler, Eulerian bulk microphysics schemes are also available in UWLCM.

We start with a brief presentation of the model, with particular attention given to the way the model was adapted to distributed memory systems. This was done using a mixed OpenMP + Message Passing Interface (MPI) approach. Next, model performance is tested on single-node systems followed by tests on a multi-node system. The main goals of these tests are to determine simulation parameters that give optimal use of computing hardware and check model scaling efficiency. Other discussed topics are the GPU vs CPU speedup and performance of different MPI implementations.

2 Model description

Full description of UWLCM can be found in Dziekan et al. (2019). Here, we briefly present key features. Cloud dynamics are modeled using an Eulerian approach. Eulerian variables are the flow velocity, potential temperature and water vapor content. Equations governing time evolution of these variables are based on the Lipps-Hemler anelastic approximation (Lipps and Hemler, 1982), which is used to filter acoustic waves. These equations are solved using a finite difference method. Spatial discretization of the Eulerian variables is done using the staggered Arakawa-C grid (Arakawa and Lamb, 1977). Integration of equations that govern transport of Eulerian variables is done with the multidimensional positive-definite advection transport algorithm (MPDATA) (Smolarkiewicz, 2006). Forcings are applied explicitly with the exception of buoyancy and pressure gradient, which are applied implicitly. Pressure perturbation is solved using the generalized conjugate residual solver (Smo-
Diffusion of Eulerian fields caused by subgrid-scale (SGS) turbulence can be modeled with a Smagorinsky-type model (Smagorinsky, 1963) or with the implicit LES approach (Grinstein et al., 2007).

Cloud microphysics can be modeled with a single- or double-moment bulk scheme, or with a Lagrangian particle-based model. Depending on the microphysics model, simulations are named UWLCM-B1M (single-moment bulk scheme), UWLCM-B2M (double-moment bulk scheme) or UWLCM-SDM (super-droplet method). Details of microphysics models can be found in Arabas et al. (2015). In both bulk schemes, cloud water and rain water mixing ratios are prognostic Eulerian variables. In the double-moment scheme, cloud droplet and rain drop concentrations are also prognostic Eulerian variables. In the Lagrangian particle-based scheme, all hydrometeors are modeled in a Lagrangian manner. The scheme is based on the super-droplet method (SDM) (Shima et al., 2009). In particular, it employs the the all-or-nothing coalescence algorithm (Schwenkel et al., 2018). In SDM, a relatively small number of computational particles, called super-droplets (SD), represent the vast population of all droplets. Equations that govern behaviour of SD are very similar to the well-known equations that govern behaviour of real droplets. The condensation equation includes the Maxwell-Mason approximation and the \( \kappa \)-Köhler parameterization of water activity (Petters and Kreidenweis, 2007). SD follow the resolved, large-scale flow and sediment at all times with the terminal velocity. Velocity of SD associated with SGS eddies can be modeled as an Ornstein-Uhlenbeck process (Grabowski and Abade, 2017). Collision-coalescence of SD is treated as a stochastic process in which probability of collision is proportional to the collision kernel. All particles, including humidified aerosols, are modeled in the same way. Therefore, particle activation is resolved explicitly, which often requires short time steps for solving the condensation equation. Short time steps are sometimes also required when solving collision-coalescence. To permit time steps for condensation and collision-coalescence shorter than for other processes, two separate substepping algorithms, one for condensation and one for collision-coalescence, are implemented.

Equations for the Eulerian variables, including cloud and rain water in bulk microphysics, are solved by a CPU. Lagrangian microphysics can be modeled either on a CPU or on a GPU. In the latter case, information about super-droplets is stored in device memory and GPU calculations can be done in parallel with the CPU calculations of Eulerian variables. Compared to UWLCM 1.0 described in Dziekan et al. (2019), order of operations has been changed to allow for GPU calculations to continue in parallel to the CPU calculations of the Eulerian SGS model. An updated UML sequence diagram is shown in fig. 1.

All CPU computations are done in double precision. Most of the GPU computations are done in single precision. The only exception are high order polynomials, e.g. in the equation for terminal velocity of droplets, that are done in double precision. So far, UWLCM has been used to model stratocumuli (Dziekan et al., 2019, 2021b), cumuli (Grabowski et al., 2019; Dziekan et al., 2021b) and raising thermals (Grabowski et al., 2018).

### 3 Adaptation to distributed memory systems

The strategy of adapting UWLCM to distributed memory systems was developed with a focus on UWLCM-SDM simulations with Lagrangian microphysics computed by GPUs. Therefore, this most complicated case is discussed first. Simpler cases with microphysics calculated by CPUs will be discussed afterwards.
The difficulty in designing a distributed memory implementation of code in which CPUs and GPUs simultaneously conduct different tasks is in obtaining a balanced workload distribution between different processing units. This is because GPUs have higher throughput than CPUs, but device memory is rather low, which puts an upper limit on the GPU workload. Taking this into account, we chose to use a domain decomposition approach that is visualized in fig. 2. The modeled domain is divided into equal slices along the horizontal axis $x$. Computations in each slice are done by a single MPI process, which can control multiple GPUs and CPU threads. Cloud microphysics within the slice are calculated on GPUs, with super-droplets residing in device memory. Eulerian fields in the slice reside in host memory and their evolution is calculated by CPU threads. Since the CPU and GPU data attributed to a process are colocated in the modeled space, all CPU-to-GPU and GPU-to-CPU communications happen via PCI-Express and do not require inter-node data transfer. The only inter-node communications are CPU-to-CPU and GPU-to-GPU. If an MPI process controls more than one GPU, computations within the subdomain of that process are divided among the GPUs also using domain decomposition along the $x$ axis. Intra-node communication between GPUs controlled by a single process makes use of the NVIDIA GPUDirect Peer to Peer technology, which allows direct transfers between memories of different devices. Intra-node and inter-node transfers between GPUs controlled by different processes are handled by the MPI implementation. If the MPI implementation uses the NVIDIA GPUDirect Remote Direct Memory Access (RMDA) technology, inter-node GPU-to-GPU transfers go directly from device memory to the interconnect, without host memory buffers.

Computations are divided between CPU threads of a process using domain decomposition of the process’ subdomain, but along the $y$ axis. The maximum number of GPUs that can be used in a simulation is equal to the number of cells in the $x$ direction. MPI communications are done using two communicators, one for the Eulerian data and one for the Lagrangian data. Transfers of the Eulerian data are handled simultaneously by two threads, one for each boundary that is perpendicular to the $x$ axis. This requires that the MPI implementation supports the MPI_THREAD_MULTIPLE thread level. Transfers of the Lagrangian data are handled by the thread that controls the GPU that is on the edge of the process’ subdomain. Collective MPI communication is done only on the Eulerian variables and most of it is associated with solving the pressure problem.

It is possible to run simulations with microphysics, either Lagrangian particle-based or bulk, computed by CPUs. In the case of bulk microphysics, microphysical properties are represented by Eulerian fields that are divided between processes and threads in the same manner as described in the previous paragraph, i.e. like the Eulerian fields in UWLCM-SDM. In UWLCM-SDM with microphysics computed by CPUs, all microphysical calculations in the subdomain belonging to a given MPI process are divided amongst the process’ threads by the NVIDIA Thrust library (Bell and Hoberock, 2012).

File output is done in parallel by all MPI processes using the parallel HDF5 C++ library (The HDF Group).

4 Performance tests

4.1 Simulation setup

Model performance is tested in simulations of a rising moist thermal (Grabowski et al., 2018). In this setup, an initial spherical perturbation is introduced to a neutrally stable atmosphere. Within the perturbation, water vapour content is increased to obtain
RH=100%. With time, the perturbation is lifted by buoyancy and water vapor condenses within it. We chose this setup, because it has significant differences in buoyancy and cloud formation already at the start of a simulation. This puts pressure solver and microphysics model to test without need of a spinup period.

Subgrid-scale diffusion of Eulerian fields is modeled with the Smagorinsky scheme. SGS motion of hydrometeors is modeled with a scheme described in (Grabowski & Abade 17). Model time step length is 0.5s. Substepping is done to achieve a time step of 0.1s for condensation and coalescence. These are values typically used when modeling clouds with UWLCM. No output of model data is done.

4.2 Computers used

Performance tests were ran on three systems: Rysy, a02 and Prometheus. Hardware and software of these systems is given in table 1 and table 2, respectively. Rysy and a02 were used only in the single-node tests, while Prometheus was used both in single- and multi-node tests. Prometheus has 72 GPU nodes connected with Infiniband. We chose to use the MVAPICH2 2.3.1 MPI implementation on Prometheus, because it supports the MPI_THREAD_MULTIPLE thread level, is CUDA-aware and is free to use. Other implementation that meets these criteria is OpenMPI, but it was found to give greater simulation wall time in scaling tests of libmpdata++ (appendix B). NVIDIA GPUDirect RDMA was not used by the MPI implementation, because it is not supported by MVAPICH2 for the type of interconnect used on Prometheus. MVAPICH2 does not allow more than one GPU per process. Therefore multi-node tests were done for 2 processes per node, each process controlling 1 GPU and 12 CPU threads.

4.3 Performance metrics

Wall time taken to complete one model time step $t_{tot}$ is divided into three parts, $t_{tot} = t_{CPU} + t_{GPU} + t_{CPU\&GPU}$, where:

- $t_{CPU}$ is the time when CPU is performing work and the GPU is idle,
- $t_{GPU}$ is the time when GPU is performing work and the CPU is idle,
- $t_{CPU\&GPU}$ is the time when CPU and GPU are performing work simultaneously.

The total time of CPU (GPU) computations is $t_{CPU}^{tot} = t_{CPU} + t_{CPU\&GPU}$ ($t_{GPU}^{tot} = t_{GPU} + t_{CPU\&GPU}$). The degree to which CPU and GPU computations are parallelized is measured with $t_{CPU\&GPU}/t_{tot}$. Timings $t_{CPU}$, $t_{GPU}$ and $t_{CPU\&GPU}$ are obtained using a built-in timing functionality of UWLCM that is enabled at compile-time by setting the UWLCM_TIMING CMake variable. The timing functionality does not have any noticeable effect on simulation wall time. Timer for GPU computations is started by a CPU thread just before a task is submitted to the GPU, and is stopped by a CPU thread when the GPU task returns. Therefore GPU timing in $t_{CPU\&GPU}$ and in $t_{GPU}$ includes time it takes to dispatch (and to return from) the GPU task.
4.4 Single-node performance

In this section we present tests of computational performance of UWLCM-SDM run on a single-node system. The goal is to determine how parallelization of CPU and GPU computations can be maximized. We also estimate the speedup achieved thanks to the use of GPUs. No MPI communications are done in these tests. Size of the Eulerian computational grid is 128x128x128 cells. In the super-droplet method, quality of microphysics solution depends on the number of super-droplets. We denote the initial number of super-droplets per cell by \( N_{SD} \). We perform test for different values of \( N_{SD} \). The maximum possible value of \( N_{SD} \) depends on available device memory.

The average wall time it takes to do one model time step is plotted in fig. 3. Time complexity of Eulerian computations depends on grid size and, ideally, does not depend on \( N_{SD} \). In reality, we see that \( t_{tot}^{CPU} \) slightly increases with \( N_{SD} \). Space and time complexity of Lagrangian computations increases linearly with \( N_{SD} \) (Shima et al., 2009). It is seen that \( t_{tot}^{GPU} \) in fact increases linearly with \( N_{SD} \), except for low values of \( N_{SD} \). For \( N_{SD} = 3 \), CPU computations take longer than GPU computations (\( t_{tot}^{CPU} > t_{tot}^{GPU} \)) and almost all GPU computations are done in parallel with CPU computations (\( t_{GPU} \approx 0 \)). As \( N_{SD} \) is increased, we observe that both \( t_{tot} \) and \( t_{CPU\&GPU} \) increase, with \( t_{CPU\&GPU} \) increasing faster than \( t_{tot} \), and that \( t_{CPU} \) decreases. This trend continues up to some value of \( N_{SD} \), for which \( t_{tot}^{CPU} \approx t_{tot}^{GPU} \). Parallelization of CPU and GPU computations (\( t_{CPU\&GPU}/t_{tot} \)) is highest for this value of \( N_{SD} \). If \( N_{SD} \) is increased above this value, GPU computations take longer than CPU computations, \( t_{tot} \) increases linearly and the parallelization of CPU and GPU computations decreases. The threshold value of \( N_{SD} \) depends on the system; it is 10 on *Prometheus*, 32 on *a02* and 64 on *Rysy*. This difference comes from differences in relative CPU to GPU computational power between these systems. In LES, \( N_{SD} \) is usually between 30 and 100.

The test shows that high parallelization of CPU and GPU computations, with \( t_{CPU\&GPU}/t_{tot} \) up to 80 %, can be obtained in typical cloud simulations.

In UWLCM-SDM, microphysical computations can also be done by the CPU. From the user perspective, all that needs to be done is to specify --backend=OpenMP at runtime. To investigate how much speedup is achieved by employing GPU resources, in fig. 4 we plot time step wall time of CPU-only simulations (with microphysics computed by the CPU) and of CPU+GPU simulations (with microphysics computed by the GPU). Estimated energy cost per time step is also compared in fig. 4. We find that simulations that use both CPUs and GPUs take between 10 to 130 times less time and use between 10 to 60 times less energy than simulations that use only CPUs. Speedup and energy savings increase with \( N_{SD} \) and depend on the number and type of CPUs and GPUs. It is important to note that microphysics computations in UWLCM-SDM are dispatched to CPU or GPU by the NVIDIA Thrust library. It is reasonable to expect that the library is better optimized for GPUs, because it is developed by the producer of the GPU.

4.5 Multi-node performance

Computational performance of UWLCM-SDM, UWLCM-B1M and UWLCM-B2M on distributed memory systems is discussed in this section. We consider four scenarios in which UWLCM is run on a distributed memory system for different reasons. Depending on the scenario and on the number of nodes used, number of Eulerian grid cells is between 0.5 and 18.5
million, and number of Lagrangian particles is between 40 million and 18.5 billion. Details of the simulation setup for each scenario are given in table 3. The scenarios are:

- **strong scaling** - More nodes are used in order to decrease the time it takes to complete the simulation.

- **SD scaling** - More nodes are used to increase the total device memory, allowing for more SD to be modeled, while grid size remains the same. This results in weak scaling of GPU workload and strong scaling of CPU workload. This test is applicable only to UWLCM-SDM.

- **2D grid scaling** - As more nodes are used, the number of grid cells in the horizontal directions is increased, while the number of cells in the vertical is constant. In UWLCM-SDM, number of SD per cell is constant. Therefore, as more cells are added, the total number of SD in the domain increases. This results in weak scaling of both CPU and GPU workloads. This test represents two use cases: domain size increase and horizontal resolution refinement. Typically in cloud modeling, domain size is increased only in the horizontal because clouds form only up to certain altitude.

- **3D grid scaling** - similar to **2D grid scaling**, but more cells are used in each dimension. This would typically be used to increase resolution of a simulation.

In each case, the maximum number of super-droplets that fit the device memory is used in UWLCM-SDM. The only exception is the **strong scaling** test in which, as more nodes are added, the number of SD per GPU decreases. Note how **SD scaling** is similar to **strong scaling**, but with more SDs added as more GPUs are added. Also note that the **2D grid scaling** and **3D grid scaling** tests are similar, but with differences in sizes of distributed memory data transfers.

UWLCM-SDM simulation time versus number of nodes used is plotted in fig. 5. First, we discuss the **strong scaling** scenario. We find that most of the time is spent on CPU-only computations. This is because in this scenario \( N_{SD} = 3 \), below the threshold value of \( N_{SD} = 10 \) determined by the single-node test (section 4.4). As more nodes are added, \( t_{CPU\&GPU} \) and \( t_{tot} \) decrease. Ratio of these two values, which describes the amount of parallelization of CPU and GPU computations, is low (30%) in a single-node run and further decreases, however slowly, as more nodes are used.

Better parallelization of CPU and GPU computations is seen in the **SD scaling** scenario. In this scenario, the CPU workload scales the same as in the **strong scaling** scenario, but the workload per GPU remains constant. Largest value of \( t_{CPU\&GPU}/t_{tot} \), approximately 80%, is found for \( N_{SD} = 10 \). The same value of \( N_{SD} \) was found to give highest \( t_{CPU\&GPU}/t_{tot} \) in the single-node tests section 4.4. We observe that \( t_{tot}^{GPU} \) is approximately constant. Given the weak scaling of GPU workload in this scenario, we conclude that the cost of GPU-GPU MPI communications is small. The small cost of GPU-GPU MPI communications, together with the fact that for \( N_{SD} > 10 \) the total time of computations is dominated by GPU computations, gives very high scaling efficiencies, around 100%.

**2D and 3D grid scaling** are weak scaling tests which differ in the way the size of CPU MPI communications scales. We find that wall time per time step \( t_{tot} \) scales very well (scaling efficiency exceeds 95%) and that \( t_{tot} \) is dominated by \( t_{tot}^{GPU} \) \( (t_{tot}^{GPU} > t_{tot}^{CPU}) \). The latter observations is consistent with the fact that the number of super-droplets \( (N_{SD} = 100) \) is larger than the threshold, \( N_{SD} = 10 \), determined in single-node tests. As in **SD scaling**, approximately constant \( t_{tot}^{GPU} \) indicates low cost
of MPI communications between GPUs. Contrary to $t_{\text{tot}}^{\text{GPU}}$, $t_{\text{tot}}^{\text{CPU}}$ clearly increases with the number of nodes. This shows that the cost of CPU-CPU MPI communications is non-negligible. Increase of $t_{\text{tot}}^{\text{GPU}}$ does not cause an increase of $t_{\text{tot}}$, because additional CPU computations are done simultaneously with GPU computations and $t_{\text{tot}}^{\text{GPU}} > t_{\text{tot}}^{\text{CPU}}$ in the studied range of the number of nodes. It is reasonable to expect that $t_{\text{tot}}^{\text{GPU}}$ scales better than $t_{\text{tot}}^{\text{CPU}}$ also for more nodes than used in this study. In that case, there should be some optimal number of nodes for which $t_{\text{tot}}^{\text{GPU}} \approx t_{\text{tot}}^{\text{CPU}}$. For this optimal number of nodes both scaling efficiency and parallelization of CPU and GPU computations are expected to be high.

Comparison of wall time scaling in UWLCM-B1M, UWLCM-B2M and UWLCM-SDM is shown in fig. 6. UWLCM-B1M and UWLCM-B2M use simple microphysics schemes that are computed by the CPU. UWLCM-B2M, which has four Eulerian prognostic variables for microphysics, is more complex than UWLCM-B1M, which has two. Regardless of this, wall time is very similar for UWLCM-B1M and UWLCM-B2M. Wall time of UWLCM-SDM, which uses much more complex microphysics scheme, is higher by a factor that depends on the number of SD. In 2D grid scaling and 3D grid scaling tests of UWLCM-SDM there are 100 SD per cell, which is a typical value used in LES. Then, on a single node, UWLCM-SDM simulations take approximately eight times longer than UWLCM-B1M or UWLCM-B2M simulations. However, UWLCM-SDM scales better than UWLCM-B1M and UWLCM-B2M, because scaling cost is associated with the Eulerian part of the model and in UWLCM-SDM this cost does not affect total wall time, as total wall time is dominated by Lagrangian computations. In result, the difference in wall time between UWLCM-SDM and UWLCM-B1M or UWLCM-B2M decreases with the number of nodes. For the largest number of nodes used in 2D grid scaling and 3D grid scaling, UWLCM-SDM simulations take approximately five times longer than UWLCM-B1M or UWLCM-B2M simulations. The strong scaling UWLCM-SDM test uses 3 SD per cell. For such low number of SD, time complexity of Lagrangian computations in UWLCM-SDM is low and we see that the wall time and it’s scaling are very similar to that of UWLCM-B1M and UWLCM-B2M.

5 Summary

A numerical model with Lagrangian particles embedded in an Eulerian fluid flow has been adapted to clusters equipped with GPU accelerators. On multi-node systems, computations are distributed among processes using static domain decomposition. The Eulerian and Lagrangian computations are done in parallel on CPUs and GPUs, respectively. We identified simulation parameters for which the amount of time during which CPUs and GPUs work in parallel is maximized.

Single-node performance tests were done on three different systems, each equipped with multiple GPUs. Percentage of time during which CPUs and GPUs simultaneously compute depends on the ratio of CPU to GPU workloads. GPU workload depends on the number of Lagrangian computational particles. For optimal workload ratio, parallel CPU and GPU computations can take more than 80% of wall time. This optimal workload ratio depends on the relative computational power of CPUs and GPUs. On all systems tested, workload ratio was optimal for between 10 and 64 Lagrangian particles per Eulerian cell. If only CPUs are used for computations, simulation take up to 120 times longer and consume up to 60 times more energy than simulations that use both CPUs and GPUs. We conclude that GPU accelerators enable running useful scientific simulation on single-node systems at a decreased energy cost.
Computational performance of the model on a distributed memory system was tested on the Prometheus cluster. We found that cost of communication between nodes slows down computations related to the Eulerian part of the model by a much higher factor than computations related to the Lagrangian part of the model. For example, in a weak scaling scenario (3D grid scaling) $t_{\text{CPU}}^{\text{tot}}$ is approximately three times larger on 27 nodes than on one node, while $t_{\text{GPU}}^{\text{tot}}$ is increased by only around 7% (fig. 5).

The reason why Eulerian computations scale worse than Lagrangian computation is that solving the pressure perturbation, which is done by the Eulerian component, requires collective communications, while the Lagrangian component requires peer-to-peer communications only. In single-node simulations on Prometheus an optimal ratio of CPU to GPU workloads is seen for 10 Lagrangian particles per Eulerian cell. In the literature, number of Lagrangian particles per Eulerian cell typically is higher, between 30 and 100 (Shima et al., 2009; Dziekan et al., 2019, 2021b). When such higher number of Lagrangian particles is used in single-nodes simulations on Prometheus, most of the time is spent on Lagrangian computations. However, in multi-node runs, Eulerian computation time scales worse than Lagrangian computation time. Since Eulerian and Lagrangian computations are done simultaneously, there is an optimal number of nodes for which the amount of time during which CPUs and GPUs work in parallel is maximized and the scaling efficiency is high. In scenarios in which GPU computations take most of the time, scaling efficiency exceeds 95% for up to 40 nodes. Fraction of time during which CPU and GPU work in parallel is between 20% and 50% for the largest number of nodes used. In weak scaling scenarios, the fraction of time during which both processing units work could be increased by using more nodes, but it was not possible due to the limited size of the cluster. Single-node simulations with Lagrangian microphysics computed by GPUs are around eight times slower than simulations with bulk microphysics computed by CPUs. However, the difference decreases with the number of nodes. For 36 nodes simulations with Lagrangian microphysics are five times slower, and the difference should be further reduced if more nodes were used.

Our approach of using CPUs for Eulerian calculations and GPUs for Lagrangian calculations results in CPUs and GPUs computing simultaneously for majority of the time step and gives good scaling on multi-node systems with several dozen nodes. The same approach can be used in other numerical models with Lagrangian particles embedded in an Eulerian flow.

**Code and data availability.** Source code of UWLCM, libmpdata++ and libcloudph++ is available at https://github.com/igfuw. In the study, the following code versions were used: UWLCM v2.0 (Dziekan and Waruszewski, 2021), libmpdata++ v2.0-beta (Arabas et al., 2021) and libcloudph++ v3.0 (Dziekan et al., 2021a). Dataset, run scripts and plotting scripts are available at Dziekan and Zmijewski (2021).

**Appendix A: Software**

UWLCM is written in C++14. It makes extensive use of two C++ libraries that are also developed at the Faculty of Physics of the University of Warsaw: libmpdata++ (Jaruga et al., 2015; Waruszewski et al., 2018) and libcloudph++ (Arabas et al., 2015; Jaruga and Pawlowska, 2018). Libmpdata++ is a collection of solvers for generalised transport equations that use the
Multidimensional Positive Definite Advection Transport Algorithm (MPDATA) algorithm. Libcloudph++ is a collection of cloud microphysics schemes.

In libcloudph++, the particle-based microphysics algorithm is implemented using the NVIDIA Thrust library. Thanks to that, the code can be run on GPUs as well as on CPUs. It is possible to use multiple GPUs on a single machine, without MPI. Then, each GPU is controlled by a separate thread and communications between GPUs are done with asynchronous cudaMemcpy. Libmpdata++ uses multidimensional array containers from the blitz++ library (Veldhuizen, 1995). Threading can be done either with OpenMP, Boost.Thread or std::thread. In UWLCM we use the OpenMP threading as it was found to be the most efficient. Output in UWLCM is done using the HDF5 output interface that is a part of libmpdata++. It is based on the thread safe version of the C++ HDF5 library. UWLCM, libcloudph++ and libmpdata++ make use of various components of the Boost C++ library (Koranne, 2011). In order to have parallel CPU and GPU computations in UWLCM, functions from libmpdata++ and from libcloudph++ are launched using std::async. UWLCM, libcloudph++ and libmpdata++ are open source software distributed via the Github repository https://github.com/igfuw/. They have test suits that are automatically run on Github Actions. To facilitate deployment, a Singularity container with all needed dependencies is included in UWLCM (https://cloud.sylabs.io/library/pdziekan/default/uwlcm).

Libcloudph++ and libmpdata++ have been adapted to work on distributed memory systems. This has been implemented using the C interface of MPI in libcloudph++ and the Boost.MPI library in libmpdata++. Tests of scalability of libmpdata++ are presented in appendix B.

Appendix B: Scalability of libmpdata++

UWLCM uses the libmpdata++ library for solving the equations that govern time evolution of Eulerian variables. The library had to be adapted for work on distributed memory systems. The domain decomposition strategy is as in fig. 2, but without GPUs. Here, we present strong scaling tests of standalone libmpdata++. The tests are done using a dry planetary boundary layer setup, which is a part of the libmpdata++ test suite. Grid size is 432x432x51. Tests were done on the Prometheus cluster. Note that all libmpdata++ calculations are done on CPUs. Two implementations of MPI are tested, OpenMPI v4.1.0 and MVAPICH2 v2.3.1. Note that Prometheus has 2 GPU per node, but MVAPICH2 does not support more than 1 GPU per process, so 2 processes per node would need to be run in UWLCM-SDM. OpenMPI does not have this limitation. For this reason in the libmpdata++ scalability tests we consider two scenarios, one with two processes per node and the other with one process per node. In the case with two processes per node, each process controls half of the available threads. Test results are shown in fig. A1. In general, better performance is seen with MVAPICH2 than with OpenMPI. Running two processes per node improves performance in MVAPICH2, but decreases performance in OpenMPI. In the best case, scaling efficiency exceeds 80% for up to 500 threads.
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Figure 1. UML sequence diagram showing the order of operations in the UWLCM 2.0 model. Right-hand-side terms are divided into condensational, non-condensational and subgrid-scale parts, $R = R_c + R_n + R_{SGS}$. Other notation follows Dziekan et al. (2019).
Figure 2. Visualization of the domain decomposition approach of UWLCM. Top-down view on a grid with 10 cells in each horizontal direction. Scalar variables are located at cell centers and vector variables are located at cell walls. Computations are divided among 5 MPI processes, each controlling 2 GPUs and 10 CPU threads. Local thread/GPU rank is the rank within respective MPI process. Dashed lines represent boundaries over which communications need to be done using MPI assuming periodic horizontal boundary conditions.
Figure 3. Single-node (no MPI) UWLCM-SDM performance for different hardware. Wall time per model time step averaged over 100 time steps. Results of LES of a raising thermal done on three different systems for varying number of super-droplets, $N_{SD}$. $t_{CPU}$, $t_{GPU}$ and $t_{CPU&GPU}$ are wall times of CPU-only, GPU-only and parallel CPU and GPU computations, respectively. These timings are presented as stacked areas of different color. Total wall time per time step $t_{tot}$ is the upper boundary of the green area. The dashed red line is the percentage of time spent on parallel CPU and GPU computations.
Figure 4. Wall time (top row) and energy usage (bottom row) per time step of CPU-only simulations (blue) and simulations utilizing both CPU and GPU (orange). In CPU-only simulations, energy usage is $t_{\text{tot}} \times P_{\text{CPU}}$, where $P_{\text{CPU}}$ is the sum of thermal design power of all CPUs. In CPU+GPU simulations, energy usage is $t_{\text{tot}}^{\text{CPU}} \times P_{\text{CPU}} + t_{\text{tot}}^{\text{GPU}} \times P_{\text{GPU}}$, where $P_{\text{GPU}}$ is the sum of thermal design power of all GPUs. $P_{\text{CPU}}$ and $P_{\text{GPU}}$ are listed in table 1. Results averaged over 100 time steps of UWLCM-SDM simulations on different single-node machines.
Figure 5. As in fig. 3, but for multi-node tests done on the Prometheus cluster for different scaling scenarios. The dotted black line is perfect scaling of $t_{\text{tot}}$. The solid red line is scaling efficiency, defined as $t_{\text{tot}}$ assuming perfect scaling divided by the actual $t_{\text{tot}}$. Perfect scaling is defined in table 3.
Figure 6. Multi-node model performance for different microphysics schemes. Wall time per model time step averaged over 100 time steps. Results of LES of a raising thermal done on the Prometheus cluster for different scaling scenarios.
Table 1. List of hardware on systems used. Computing performance and memory bandwidths are maximum values provided by the processing unit producer. Power usage of a processing unit is measured by the thermal design power (TDP).

|                | Rysy                                      | a02                                       | Prometheus<sup>a</sup>                          |
|----------------|-------------------------------------------|-------------------------------------------|-----------------------------------------------|
| CPUs           | 2x Xeon Gold 6154 @ 2.50 GHz              | 2x Xeon E5-2630 v3 @ 2.40GHz              | 2x Xeon E5-2680 v3 @ 2.50 GHz                |
| GPUs           | 4x Tesla V100                             | 2x Tesla K80                              | 2x Tesla K40 XL                              |
| CPU cores      | 2x18                                      | 2x8                                       | 2x12                                         |
| CPU performance| 2x1209.6 Gflops                           | 2x307.2 Gflops                            | 2x480 Gflops                                 |
| GPU performance<sup>b</sup> | 4x14.028 (7.014) Tflops | 2x8.73 (2.91) Tflops | 2x5.34 (1.78) Tflops | |
| CPU TDP        | 2x200 W                                   | 2x85 W                                    | 2x120 W                                      |
| GPU TDP        | 4x250 W                                   | 2x300 W                                   | 2x235 W                                      |
| host memory    | 384 GB                                    | 128 GB                                    | 128 GB                                       |
| device memory  | 4x32 GB                                   | 2x24 GB                                   | 2x12 GB                                      |
| host memory bandwidth | 2x128 GB/s                          | 2x68 GB/s                                | 2x68 GB/s                                   |
| device memory bandwidth | 4x900 GB/s                          | 2x480 GB/s                               | 2x288 GB/s                                  |
| host-device bandwidth (PCI-E) | 4x15.754 GB/s                          | 2x15.754 GB/s                            | 2x15.754 GB/s                               |
| interconnect   | n/a<sup>c</sup>                           | n/a<sup>c</sup>                           | Infiniband 56 Gb/s                           |

<sup>a</sup> The cluster has 72 such nodes.

<sup>b</sup> Single-precision performance. Double-precision performance is given in the brackets. Almost all GPU computations are done in single precision.

<sup>c</sup> Used in single-node tests only.
Table 2. List of software on systems used.

| Name      | CUDA | gcc  | Boost | HDF5  | Thrust | blitz++ |
|-----------|------|------|-------|-------|--------|---------|
| Rysy\(^a\) | 11.0 | 9.3.0| 1.71.0| 1.10.4| 1.9.5-1| 1.0.2   |
| a02       | 10.1 | 4.8.5| 1.60.0| 1.8.12| 1.9.7  | 0.10    |
| Prometheus| 11.2 | 9.3.0| 1.75.0| 1.10.7| 1.10.0 | 1.0.2   |

\(^a\) software from a Singularity container distributed with UWLCM.
Table 3. Details of multi-node scaling tests. $n_x$, $n_y$ and $n_z$ is the total number of Eulerian grid cells in the respective direction. $N_{SD}$ is the initial number of super-droplets per Eulerian grid cell. $N_{nodes}$ is the number of nodes used for the simulation. Number of Eulerian grid cells in the domain is equal to $n_x \times n_y \times n_z$. Number of superdroplets in the domain is equal to $n_x \times n_y \times n_z \times N_{SD}$. Workload per CPU is estimated assuming that it is proportional to the number of grid cells per CPU only. Workload per GPU is estimated assuming that it is proportional to the number of super-droplets per GPU only. MPI transfers, data transfers between host and device memories, and GPU handling of information about Eulerian cells are not included in these workload estimates. Data transfer sizes are for copies between different MPI processes, but do not include copies between host and device memories of the same process. Data transfer sizes are estimated assuming that time step length and air flow velocities do not change with grid size. $t^1$ is the wall time on a single node. $t^2_{GPU}$ is the wall time of GPU and CPU&GPU calculations in a simulation on two nodes.

|                  | strong scaling | SD scaling | 2D grid scaling | 3D grid scaling |
|------------------|----------------|------------|----------------|-----------------|
| $n_x$            | 240            | 240        | $\sqrt{N_{nodes}} \times 72$ | $\sqrt{N_{nodes}} \times 80$ |
| $n_y$            | 240            | 240        | $\sqrt{N_{nodes}} \times 72$ | $\sqrt{N_{nodes}} \times 80$ |
| $n_z$            | 240            | 240        | 100            | 100             |
| $N_{SD}$         | 3              | $N_{nodes} \times 3$ | 100            | 100             |
| Eulerian cells in domain [$10^3$] | 13824          | 13824      | $N_{nodes} \times 518.4$ | $N_{nodes} \times 512$ |
| superdroplets in domain [$10^6$]  | 41.472         | $N_{nodes} \times 41.472$ | $N_{nodes} \times 51.84$ | $N_{nodes} \times 51.2$ |
| workload per CPU | $\propto 1/N_{nodes}$ | $\propto 1/N_{nodes}$ | const.         | const.          |
| workload per GPU | $\propto 1/N_{nodes}$ | const.     | const.         | const.          |
| data transfer size per CPU | const.          | const.     | $\propto \sqrt{N_{nodes}}$ | $\propto N_{nodes}^{2/3}$ |
| data transfer size per GPU | const.          | $\propto N_{nodes}$ | $\propto N_{nodes}^{2/3}$ | $\propto N_{nodes}^{2/3}$ |
| time assuming perfect scaling | $t^1/N_{nodes}$ | $\max(t^1/N_{nodes}, t^2_{GPU})^b$ | $t^1$            | $t^1$            |

a Assuming that grid scaling is used to refine the resolution, as done in this paper. If it is done to increase the domain, data transfer size per GPU scales as the one per CPU.

b GPU time from two nodes simulation is taken as reference, because it is ca. 15% lower than on a single node. A plausible explanation for this is that, although the number of SD per GPU does not depend on the number of nodes, GPUs also store information about conditions in grid cells, and the amount of grid cells per GPU decreases as more nodes are used. For more than 2 nodes, GPU calculation time is approximately the same as for 2 nodes.
Figure A1. Strong scaling test of the libmpdata++ library. Wall time per time step of a dry planetary boundary layer simulation. The dotted black line shows perfect scaling.