A scalable RBF–FD method for atmospheric flow

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Abstract

Radial basis function-generated finite difference (RBF–FD) methods have recently been proposed as very interesting for global scale geophysical simulations, and have been shown to outperform established pseudo-spectral and discontinuous Galerkin methods for shallow water test problems. In order to be competitive for very large scale simulations, the RBF–FD methods needs to be efficiently implemented for modern multicore based computer architectures. This is a challenging assignment, because the main computational operations are unstructured sparse matrix-vector multiplications, which in general scale poorly on multicore computers due to bandwidth limitations. However, with the task parallel implementation described here we achieve 60–100\% of theoretical speedup within a shared memory node, and 80–100\% of linear speedup across nodes. We present results for global shallow water benchmark problems with a 30 km resolution.

Keywords: shallow water, scattered node, task parallel, distributed memory, multicore, radial basis function, RBF–FD
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1. Introduction

In computational geoscience, models are often global in the sense that the considered domain is the atmosphere around the earth, the surface of the earth, or the mantle of the earth. Furthermore, there may also be a variety of coupled physical processes as in global climate simulations, where different models for the atmosphere, the land surface,
the oceans, the large ice sheets, and others all contribute to the global results. These simulations are both time consuming and produce large volumes of data. Hence, they cannot be performed without the use of parallel processing. Some of the largest computer systems in the world are dedicated to weather and climate simulations.

Existing community models such as EC-Earth connected with the European Center for Medium-Range Weather Forecasts (ECMWF), and the Community Earth System Model (CESM) connected with the National Center for Atmospheric Research (NCAR), USA, use grid based numerical methods for the simulations. This makes it costly to resolve local solution features, because typically a larger region needs to be resolved in order to preserve grid integrity. Reported resolutions in current simulations are at a level of 30–100 km, but processes that are of importance for the global simulation result may occur on even finer scales.

An interesting alternative to using grid based methods are meshfree methods based on radial basis function (RBF) approximation. These methods work with scattered nodes and easily allow for local refinement of the numerical solutions. Global RBF approximation methods have been shown to be competitive compared with other commonly used numerical approaches in computational geosciences in a series of papers [17, 18, 44, 15]. However, for large node sets, it becomes expensive in terms of computational time and storage to work with the dense linear systems that arise from the global approximations.

By replacing the global RBF approximations with local RBF approximations, we can reduce the computational cost while retaining the flexibility with respect to geometry and the ability to work with scattered nodes. RBF-generated finite differences (RBF–FD) are similar in nature to finite difference methods, but with stencils that are built on scattered nodes instead of nodes in a grid, and an underlying interpolation that is RBF based instead of polynomial. In the paper [16], RBF–FD methods were compared with, and outperformed state-of-the-art methods for fluid flow test cases over the sphere. These comparisons were done with a sequential implementation. However, in order to demonstrate the viability of the RBF–FD method for large scale computational geoscience, we also need to show that it scales to large problem sizes and can be efficiently parallelized for large computer systems.

The RBF–FD method is relatively new, with an early mentioning in the conference paper [39] from 2000, and further development and analysis for example in [33, 43, 10, 45, 3, 9, 8, 5, 4]. There is still not much research done on parallelization of these methods. However, a parallel implementation of an RBF–FD method for the shallow water equations on the sphere for multiple CPUs and GPUs can be found in [6]. Each CPU runs a Message Passing Interface (MPI) process and handles the necessary communication. Local computations are off-loaded to the GPU attached to the CPU. Speedups of up to 7 compared with execution on one CPU are reported. An OpenMP parallelization of an RBF–FD method for a coupled thermal-fluid flow problem can be found in [25]. The stencils and problem sizes are small, and only two cores are used, but the results are good. The parallel method is further developed in [26] with experiments for up to 16 cores. An MPI implementation of a similar application problem and similar RBF–FD approach is presented in [11], where experiments are performed on a 36-node cluster. However, no performance results are reported. In the recent conference paper [14], parallel sparse matrix-vector products for applications such as the RBF–FD method for the shallow water equations are implemented for the Intel Many Integrated Core (MIC) architecture. The conclusions made there are similar to the ones we make in the present paper, even
if the implementation method is different. A preliminary version of the shared memory part of this paper can be found in [38].

In the MPI/OpenCL and MPI approaches used for parallelization in [6, 11], the programmer is responsible for explicitly expressing all data transfers and decide where in the system each computation takes place. Also the implementation for the MIC in [14] involves some rather technical operations in order to exploit the vector processing units. With the OpenMP approach of [25, 26] the programmer works with pragma directives to the compiler inserted into the sequential code. The latter is significantly easier to implement, however the performance may be suboptimal due to the global synchronization points introduced by the fork-join style programming model.

In this paper, we employ a task-based parallel programming model. These models are emerging as one of the most promising approaches to achieve high performance in scientific applications at a reasonable programming effort. By a task we mean an object that can be called asynchronously, that is, a piece of code together with its required data. The programmer writes a sequential code in terms of tasks, while a run-time system provided by a programming framework is responsible for scheduling and executing tasks in parallel. There are several widely spread frameworks of this type such as OmpSs [13] (from the StarsSs family of programming models) and StarPU [1]. In this work we use the SuperGlue framework [36, 34, 35], developed by the first author. Its performance with respect to other frameworks was tested in [35], and SuperGlue was shown to be highly competitive.

The RBF–FD discretization of the shallow water equations on the sphere generates sparse unstructured matrices, where the size of the matrix corresponds to the total number of node points (degrees of freedom) and the number of non-zero elements per row is determined by the stencil size. The main operations of the solver are sparse matrix-vector multiplications. Due to bandwidth limitations, this type of operation does not perform well on multicore architectures in general. However, for each evaluation of the spatial differential operator, 16 sparse matrix-vector multiplications are required in order to form the necessary derivatives. We show that by reusing the sparsity pattern across these multiplications, we increase the ratio of computations per memory access and regain the scalability. In this way, we address our first objective to establish that RBF–FD methods on unstructured grids can be efficiently implemented for modern multicore based computer architectures.

The second objective of the paper is to demonstrate that task-based parallel programming is a tool that provides high programmer productivity and significantly facilitates the writing of scientific software without sacrificing absolute performance. Samples from the implementation are included for reference.

Finally, we show that the RBF–FD method can produce highly resolved and very sharp solutions to challenging shallow water benchmark problems. A series of illustration for two test cases, flow over an isolated mountain and evolution of a highly non-linear way are shown. Future work involves how to implement adaptive node refinement. This is not pursued here.

The paper is organized as follows: In Section 2 we define the shallow water problems that we use as benchmarks. Then in Section 3, we discuss the RBF–FD method. Section 4 contains the discrete formulation of the problems. The task-based programming model is explained in Section 5, and the parallel implementation of the solver is described in Section 6. Finally, Section 7 provides performance experiments and solution examples,
and the conclusions are given in Section 8.

2. The shallow water equations on the sphere

For a new method to be adopted in any field it must first be shown to perform well for certain benchmark problems. In computational geosciences, and especially for global atmospheric simulations, the shallow water equations (SWE) constitute an important problem class for numerical testing. The SWE are a set of non-linear partial differential equations (PDE) which capture the main features of the horizontal dynamics of atmospheric flow around the earth.

In this paper, we will use two of the benchmarks that were used in [16] to evaluate the parallel performance of RBF–FD methods for the SWE. The first test case that we use is called “flow over an isolated mountain”. The initial condition is perfectly laminar flow in the easterly direction, but due to the presence of a very large cone shaped mountain, the flow develops a wave pattern with time. A full description can be found in [40]. The second test case, described in [24], represents the evolution of a highly non-linear wave. The solution contains high frequency components and sharp gradients. Figure 1 shows the solutions of the two test cases at different times.

In combination with grid based methods on the sphere, the SWE are often formulated in spherical coordinates. However, this introduces unphysical singularities at the poles. RBF based methods are oblivious to the coordinate system used and its orientation. Hence, we choose to work with the Cartesian formulation of the SWE, which is singularity free and given by

\[
\frac{\partial u}{\partial t} = -(u \cdot \nabla)u - f(x \times u) - g\nabla h,
\]

\[
\frac{\partial h}{\partial t} = -\nabla \cdot (hu),
\]

where \( u = (u, v, w) \) is the wind field, \( x = (x, y, z) \) is the location, \( f = 2\Omega z \) is the Coriolis force, where \( \Omega \) is the angular velocity of the earth, \( h \) is the geopotential height, and \( g \) is the gravity.

Using the Cartesian formulation, we instead need to project the operators onto the curved surface of the earth. This approach was introduced in [18], and is applied also in [16]. For practical purposes, we first scale the problem to the unit sphere. The projection operator onto the unit sphere is defined as \( P = I - xx^T \). The projected form of the SWE then becomes

\[
\frac{\partial u}{\partial t} = -P[(u \cdot P\nabla)u + f(x \times u) + gP\nabla h],
\]

\[
\frac{\partial h}{\partial t} = -P\nabla \cdot (hu).
\]

3. The RBF–FD method and its properties

Here we will discuss the RBF–FD method. We start with the general framework, and then discuss different properties that are important for the approximations. For
Figure 1: RBF–FD solutions to the shallow water test case “flow over an isolated mountain” using 655,532 nodes and showing the geopotential height (top), and the “evolution of a highly non-linear wave” using 612,346 nodes and showing relative vorticity (bottom) at different times. The black circle shows the location of the cone shaped mountain. In both cases, a perturbation of an initially laminar flow develops over time to affect the flow on a global scale. The equator and the 0°/180° meridian are indicated by solid grid lines, and north is upward.

Each property, we will in particular discuss aspects that may influence the ability of the method to scale to large problem sizes.

We start by defining the RBFs that we will use in the underlying approximations. A radial basis function $\phi(r)$ is a function in $d$ dimensions whose value depends only on the distance $r = \|x - c\|$ from its center point $c$. Many commonly used RBFs are also equipped with a so called shape parameter $\varepsilon$ that is applied to the argument to influence the flatness of the RBF. Figure 2 shows the effect of scaling on a Gaussian RBF, $\phi(r) = e^{-\varepsilon r^2}$. 

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Figure 2: The shape of a Gaussian RBF for different values of the shape parameter $\varepsilon$.

Assume there are $N$ scattered nodes on the surface of the sphere (or in any other geometry of choice). A differential operator $\mathcal{D}$ is approximated at the location $x_c$ by using a weighted combination of the function values $f_k$, $k = 1, \ldots, n$ at the $n \ll N$ nearest neighboring nodes. That is,

$$\mathcal{D}f(x_c) \approx \sum_{k=1}^{n} w_k f_k,$$

where the weights $w_k$ are determined by requiring the approximation to be exact when the solution can be exactly represented by the basis underlying the approximation, which here consists of radial basis functions centered at the scattered nodes. Assuming that $f(x) = \sum_{k=1}^{n} \lambda_k \phi(||x - x_k||) \equiv \sum_{k=1}^{n} \lambda_k \phi_k(x)$ yields the following linear system of equations for the weights

$$\begin{bmatrix}
\phi_1(x_1) & \phi_1(x_2) & \cdots & \phi_1(x_n) \\
\phi_2(x_1) & \phi_2(x_2) & \cdots & \phi_2(x_n) \\
\vdots & \vdots & \ddots & \vdots \\
\phi_n(x_1) & \phi_n(x_2) & \cdots & \phi_n(x_n) \\
\end{bmatrix} \begin{bmatrix}
w_1 \\
w_2 \\
\vdots \\
w_n \\
\end{bmatrix} = \begin{bmatrix}
\mathcal{D}\phi_1(x_c) \\
\mathcal{D}\phi_2(x_c) \\
\vdots \\
\mathcal{D}\phi_n(x_c) \\
\end{bmatrix}.$$

Figure 3 provides a graphical representation of a differentiation stencil on the sphere. The exemplified stencil is quite large with $n = 75$. Because the sphere is a periodic domain (no boundaries) we can use large stencils without running in to the particular problems that are associated with boundaries and high-order stencil approximations. It has been shown that RBF approximations in the limit where the basis functions become flat are polynomials \[12, 28, 32, 30\]. Therefore, at least for small values of $\varepsilon$, the approximation error for a stencil for a first-order derivative with $n$ points behaves as $O(h^k)$, where

$$ (k + 1)^2 \leq n < (k + 2)^2, $$

and $(k + 1)^2$ is the number of degrees of freedom in a polynomial of degree $k$ restricted to the sphere.
Figure 3: Left: This is an illustration of a 75 node stencil on the sphere. The differential operator is evaluated at the node marked with a square. The size of the markers indicates the magnitude of the stencil weights and the color indicates the signs. The nodes marked with green rings (further away) are not included in the stencil. Right: The structure of a global differentiation matrix for $N = 6400$ nodes. A bandwidth minimization algorithm has been applied to the matrix.

To assemble a global differentiation matrix that approximates the operator $D$ at all node points given the function values at these points, we let $x_{c}$ traverse all node points, and then we enter the generated weights into the corresponding row in the global sparse differentiation matrix. A typical sparsity pattern for a differential operator approximation on the sphere is shown in the right part of Figure 3.

3.1. Stability for time-dependent PDEs

The most critical issue in scaling the RBF–FD method to large problem sizes is stability, in particular for hyperbolic PDEs (lacking a diffusion term) like the SWE. The discretization of the spatial operator may, as illustrated in [16], exhibit eigenvalues with a positive real part, leading to instability. A remedy for this was given in [20] in the form of hyperviscosity operators that can be added to the discrete operator in order to suppress the unstable modes. The hyperviscosity operator is of the type $\Delta^{m}$, where $m$ needs to be higher for larger stencil sizes (corresponding to higher order methods).

To ensure convergence to the inviscid solution, the hyperviscosity must be inversely scaled with respect to the resolution, and to obtain optimal accuracy, the order of the hyperviscosity operator should match the numerical order of the advection discretization.

Using Gaussian RBFs, the hyperviscosity operators can be efficiently implemented [20], see also [29] for the flat limit version. Therefore, we will use Gaussian RBFs throughout this paper.

3.2. Stencil sizes

The stencil size has already been discussed above and how the order of accuracy $k$ relates to the stencil size $n$ is given by equation (5). A larger stencil leads to higher accuracy, but more stability problems.
The first test case, flow over an isolated mountain, has a non-smooth forcing term representing the conical mountain. Therefore, the accuracy is not improved by increasing the stencil size beyond a certain point. For this problem we will use the size $n = 31$ (corresponding to a fourth order approximation) chosen in [16] for all experiments. For the second test case, we use a larger stencil with $n \approx 75$. This corresponds to a seventh order approximation.

Following equation (5), it might seem reasonable to choose the stencil size equal to $(k + 1)^2$. However, it has been observed in experiments that certain stencil sizes have better stability properties than others. These special numbers are typically $n = 17, 31, 50, 75,$ and 101. We do not know yet why these sizes are better, but one theory is that they have symmetry properties, depending on the layout of the nodes.

### 3.3. Node sets on the sphere

There are different approaches to create quasi uniform node distributions on the sphere. Minimum energy (ME) nodes [42] are of high quality, but only relatively small node sets can be generated. Maximum determinant (MD) nodes [41, 42] are also high quality and somewhat larger node sets can be generated. These are used for the smaller problem sizes in our numerical experiments. For the larger problem sizes, we have tried icosahedral (ICO) node sets [2] that can be easily generated for large problem sizes. However, the quality of the icosahedral node sets is a little bit worse. The number of close neighbors to a node point depends on the location in the original icosahedron. This does not matter very much for small problem sizes, but for the large problem sizes, it is one of the factors that influence stability. Therefore, we have instead used DistMesh [31] (DM) to generate a first version of the node set, and then smoothed it by application of an approximate electrostatic repulsion. Local node refinement is a possibility that should be further explored. For a problem with localized solution features, this allows for a smaller total number of nodes than with a uniform distribution, while providing appropriate resolution in each part of the domain. However, moving to non-uniform node sets would require better stencil selection algorithms in order to attain stability. The node sets must also be smoothly varying to avoid large dispersion and aliasing errors.

### 3.4. Scaling of stencils and RBFs

Stencil weights are computed by solving a linear system $Aw = b$. If we fix the node configuration, i.e., the number of stencil nodes and their relative location up to a scaling and translation, then we can focus on the effects of the scaling of the coordinates and the scaling of the RBFs. Define the stencil size $h$ as the largest distance between two nodes in the stencil. We use the notation $A = A(h, \varepsilon)$ and $b = b(h, \varepsilon)$ in order to facilitate the discussion of the scaling issues. Due to the interchangeability of the scaling of the distances $r$ and the shape parameter $\varepsilon$ in the argument of the RBFs, we have the following relations

$$A(\alpha h_0, \varepsilon_0) = A(h_0, \alpha \varepsilon_0), \quad (6)$$

$$b(\alpha h_0, \varepsilon_0) = \alpha^q b(h_0, \alpha \varepsilon_0), \quad (7)$$

where $q$ is the order of the differential operator $\mathcal{D}$.

We will discuss two main strategies for choosing the scaling parameters: (i) keeping the shape parameter constant and (ii) stationary scaling. With the first strategy, theory
tells us that the error will go to zero as $h \to 0$ in exact arithmetic. However, the numerical stability of the computation of the stencil weights depends on the conditioning of the matrices $A$. Equation (6) tells us that decreasing $h$ with a fixed $\varepsilon$ is numerically equivalent to decreasing $\varepsilon$ for a fixed $h$. For a fixed node set, the condition number of $A$ goes to infinity as $\varepsilon^{-2k}$ when $\varepsilon \to 0$ as shown in [28] and [32]. Hence, if the weights are computed directly from (4), we get numerical stability problems for small $\varepsilon$ and/or small $h$.

A number of numerically stable methods for evaluating RBF interpolants or approximants in the flat $\varepsilon \to 0$ limit have been developed [23, 19, 21], see also [22] for global RBF approximations on the sphere. Common for these are that the cost for evaluating the stencil weights becomes higher (by a constant factor), but computations can be stably performed for any small value of $\varepsilon$. Some problems may arise when the stencils become very small in relation to the size of the spherical surface. The surface is then numerically flat, but we are trying to generate a stencil in a three-dimensional space. A work-around is to project the nodes onto the tangent plane and compute the weights in the two-dimensional plane instead (personal communication Bengt Fornberg).

The second strategy, stationary scaling, is commonly adopted in practical applications, as, e.g. in [16]. By employing equation (6) for both arguments, we have that $A(h_0, \varepsilon_0) = A(\alpha h_0, \varepsilon_0/\alpha)$. That is, if we decrease $h$ while at the same time increasing $\varepsilon$, the condition number stays constant. This is of course an advantage, but as $h \to 0$ we will run into a saturation error at some point, beyond which accuracy is not further improved. The stencil can be modified by inclusion of lower order polynomial terms, by which some order of convergence can be retained. How the stencil approximation errors depend on $\varepsilon$ and $h$ is discussed in [3] and explicit formulas are given for low order stencils.

In the article [29], the two strategies for stencil scaling are compared with each other as well as with stencils that include polynomial terms. It can be seen that up to the point where the saturation error enters, the performance is similar.

In [16], RBF–FD approximations were computed for $N$ up to $O(10^5)$ nodes with maintained convergence for the flow over an isolated mountain test case. However, a slight change in the convergence trend for the largest problem size could indicate that the saturation error occurs in that regime. This might put in question the need for higher resolutions and parallel implementations. We will try to put this issue to rest by a hypothetical example. Assume that we want to approximate two different functions $f(x)$ and $g(x) = f(\beta x)$. Then any derivative of order $q$ is scaled by $\beta^q$ for $g(x)$ compared with the same derivative of $f(x)$ for the corresponding location. If we apply a stencil with the combination $(h_0, \varepsilon_0)$ to $f(x)$, we should apply a stencil with $(h_0/\beta, \beta \varepsilon_0)$ to $g(x)$. Then the function values at the node points for the two functions will be identical and the stencils will be identical up to a scaling factor of $\beta^q$ (see eq. (7)), corresponding to the higher derivative of $g(x)$. The conclusion is that if saturation strikes at a certain $h$ for $f(x)$, then it strikes at $h/\beta$ for $g(x)$. Hence, a function with variations at shorter scales ($\beta > 1$) can be resolved more before reaching the saturation error. Furthermore, by using the stable methods, we can use scaling approach (i), which does not suffer from saturation errors.
4. The discrete formulation of the shallow water equations

To obtain a semi-discretization of the PDE, the RBF–FD stencils are used for approximation of the spatial PDE operators at each node point and the results are assembled into one sparse global differentiation matrix per operator involved. Here the spatial operators are the different components of $P \nabla$, and we denote the differentiation matrices by $D_N^x$, $D_N^y$, and $D_N^z$. For details on how to apply the projected operators to the RBFs for the right hand side in the stencil weight computation (4), see [16].

We adopt a tensor notation, where each element of a matrix or a vector in turn is a matrix or a vector. An underlined quantity such as $x = (x_1, \ldots, x_n)^T$ denotes a node function with a (matrix, vector or scalar) value at each node point. Operations that involve node functions are interpreted as being applied element wise for each node. The discrete gradient operator applied to a vector or a scalar node function yields

$$D_N u = \begin{bmatrix} D_N^x u & D_N^y v & D_N^z w \\ D_N^x u & D_N^y v & D_N^z w \\ D_N^x u & D_N^y v & D_N^z w \end{bmatrix}, \quad D_N h = \begin{bmatrix} D_N^x h \\ D_N^y h \\ D_N^z h \end{bmatrix},$$

(8)

and the discrete divergence operator similarly becomes

$$D_N \cdot u = D_N^x u + D_N^y v + D_N^z w.$$

(9)

With this notation, the semi-discrete form of the shallow water equations on the sphere becomes

$$\frac{\partial u}{\partial t} = -P [u \cdot D_N u + f(x \times u) + gD_N h],$$

(10)

$$\frac{\partial h}{\partial t} = - (u \cdot D_N h + hD_N \cdot u).$$

(11)

For the time-derivative, the classical fourth order Runge-Kutta method is used. The SWE are of hyperbolic type, which motivates the choice of an explicit time-stepping method. To ensure stability, small hyperviscosity terms $\gamma D_N^x u$ and $\gamma D_N^z h$ respectively, are incorporated into the scheme. These represent a small amount of diffusion, which acts as stabilizer without significantly altering the solution values. As in [16], all operators are discretized using Gaussian RBFs, which allows for efficient evaluation of the hyperviscosity operator [20].

The computationally most expensive operations in evaluating the right hand side functions in the system of ODEs (10)–(11) resulting from the spatial discretization of the PDEs (1)–(2) are the applications of differentiation matrices to node functions. There are four solution components $u, v, w, \text{ and } h$, and each is differentiated with respect to the three coordinate directions. Furthermore, we apply the hyperviscosity once to each component. This results in a total of 16 sparse matrix-vector multiplications for each evaluation of the right hand side functions.

5. Task based parallel programming

We have chosen to use a task-based programming model to parallelize the SWE solver. There are two main reasons for this choice. The first is the programmer productivity
aspect. A key idea in task parallel programming is that it allows the programmer to write a sequential code in terms of computational tasks, and then a run-time system handles the parallel execution of the tasks. In scientific applications, tasks typically share some data and this leads to dependencies between tasks in terms of constraints on the order in which they can execute. The run-time system needs to respect these constraints when scheduling tasks for execution. Figure 4 shows a schematic representation of an algorithm in terms of dependent tasks.

The second reason to choose a task-based programming model is performance. Tasks are scheduled dynamically at run-time to the cores, processors, and computational nodes. Load balancing, so far only in the shared memory setting, is achieved at run-time by task stealing. No particular assumptions are made about the hardware beforehand except for the distinction between shared memory and distributed memory. The execution instead adapts to the conditions of the system where the software is running. Another important aspect is that in the task model, global barriers, which typically do not scale well, can be avoided. Synchronization is instead fine grained between different tasks. In [27] and [7] the suitability of different programming models and scheduling approaches was investigated for linear algebra problems on multicore based architectures, and dynamic scheduling was found to be beneficial for performance.

In this work, we use the SuperGlue\textsuperscript{1} library, developed by the first author [36, 34, 35], for dependency-aware task-based parallel programming. The performance of SuperGlue has been compared with that of related efforts and it was shown that SuperGlue has comparatively very low run-time overhead. In SuperGlue, the programmer defines tasks and also registers what type of access to shared data that each task performs. The types can be for example read, write, and add, where the latter is used for accesses that can be performed in any order, but not at the same time. The tasks are submitted to the run-time system, which infers the dependencies from the data accesses and schedules the tasks accordingly.

An important principle in our view of task parallelism is that tasks depend on data, not on other tasks. This allows us to add dependencies without having to synchronize with other tasks. We use a system with data versioning, where tasks depend on a particular version of the data. When the required version becomes available, the task can execute. It should be noted that we do not duplicate data. The versions replace

\textsuperscript{1}https://github.com/tillenius/superglue
each other as they become available. The versions are handled by data objects that we call (data) handles. This gives additional flexibility and allows us to construct handles not only for consecutive blocks of data, but for arbitrary data structures, as well as for logical resources such as, for example, disk I/O.

6. The sequential and parallel implementations of the solver

In this section, we will describe the implementation, optimization, and parallelization of the RBF–FD method for the SWE step by step, starting with the sequential code and leading up to the parallel distributed code.

6.1. The sequential MATLAB code

The first implementation of the RBF–FD SWE solver was done in MATLAB. However, as reported in [16], even the pilot implementation was 4–10 times faster, when comparing serial execution times, than the latest C++ discontinuous Galerkin solver developed at the National Center for Atmospheric Research (NCAR), Boulder, CO, USA. In the comparison, the discontinuous Galerkin model was running on a cubed sphere grid utilizing spatially uniform $h$- and $p$-refinement. The performance difference can in part be attributed to the restrictive time step condition of high order spectral elements.

The main computational tasks in the code are the initialization part where the weights in the differentiation matrices are computed, and the Runge-Kutta time-stepping loop. An excerpt from the MATLAB-code containing these two steps is shown in Appendix A.

Profiling of the MATLAB program shows that the majority of the time (74% for the tested problem size) is spent in the evaluation of the right hand side of the SWE. Therefore, this is the part of the program that we target initially. Note that the initial computation of the different stencil weights is trivially parallelizable in the sense that for each operator, the $N$ linear systems of size $(n \times n)$ are independent of each other.

Further examining the computations in the right hand side function, we find that over 90% of the time is spent in sparse matrix-vector multiplications between differentiation matrices and intermediate solution vectors. Hence, parallelizing this code efficiently amounts to handling sparse unstructured matrix-vector multiplications, which is a classical example of a bandwidth-bound operation that is hard to get to scale on multicore based computer systems.

6.2. The sequential C++ code

We consider the time-stepping loop only and start by performing optimizations and adjustments of the sequential code. The particular characteristic of sparse matrix-vector multiplications that makes them scale poorly is the low ratio of computations to memory accesses. A common feature of this and other RBF–FD methods is that differentiation matrices for different operators have identical sparsity patterns, since they use the same stencil sizes. This can be exploited to increase the computational intensity. First, we store the solution variables in the $N \times 4$ matrix $H$, where each row contains the values of $(u, v, w, h)$ at one point $(x, y, z)$. Then, we also store the differentiation matrices, denoted by $DPx$, $DPy$, $DPz$, and $L$ (for the hyperviscosity) together in the aggregated operator $D$, reusing the common sparsity pattern. Instead of performing four separate
Table 1: The execution time in millions of cycles and the speedup of (the different parts of) the right hand side computation when the sequential C++ code is compared with the sequential MATLAB code.

<table>
<thead>
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<th></th>
<th>MATLAB</th>
<th>C++</th>
<th>Speedup</th>
</tr>
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<tbody>
<tr>
<td>Differentiation</td>
<td>8186</td>
<td>1441</td>
<td>5.7</td>
</tr>
<tr>
<td>Hyperviscosity</td>
<td>2606</td>
<td>679</td>
<td>3.8</td>
</tr>
<tr>
<td>Other rhs ops</td>
<td>790</td>
<td>200</td>
<td>4.0</td>
</tr>
<tr>
<td>Total</td>
<td>12062</td>
<td>2402</td>
<td>5.0</td>
</tr>
</tbody>
</table>

matrix-vector multiplications with $H$, we compute them all at the same time as $T = D \cdot H$, where $T$ has 16 values per node point.

The computations can be further optimized by noting that each instance of the solution value $(u, v, w, h)$ is of length four. This allows us to employ AVX (Advanced Vector Extensions to the x86 instruction set) SIMD instructions when multiplying the differentiation matrices with $H$. The SIMD instructions are significantly more efficient than performing four single instruction operations. We have also changed from column major storage as used by MATLAB to row-major compressed sparse row (CSR) storage of the matrices.

The overall speedup from the MATLAB implementation to the sequential optimized C++ code was 5.0 times. For a breakdown into different parts, see Table 1. Note that much of the gain was achieved by exploiting the structure of this specific problem. We could not have achieved the same speedup by using a standard optimized library routine for sparse matrix-vector multiplication.

### 6.3. The task parallel shared memory C++ code

As discussed in Section 5, we use the task parallel framework SuperGlue for the implementation of the parallel code. The first step is a shared memory parallel implementation, and the problem we will discuss is how to do the sparse matrix-vector multiplications efficiently using SuperGlue.

We need to formulate the algorithm in terms of tasks. This can be done in different ways for any given problem and it does have implications for performance. Task size is one important parameter. Too small tasks make the overhead from scheduling visible, whereas too large tasks are hard to schedule efficiently and may lead to reduced parallelism. Figure 5 shows how the matrix is blocked to allow parallelism, and the resulting access pattern of a single task of the sparse matrix-vector multiplication.

![Figure 5: The partitioning of the matrix and vectors, and the responsibility of a single task of the matrix-vector multiplication. The data accessed by the task is shaded.](image)

The SuperGlue implementation of the blocked version is included in Appendix A. We would like to emphasize that the code complexity is quite similar to that of the original MATLAB implementation. We need to write a wrapper for each computational kernel.
to make it into a task, but it is still easy to follow the code and recognize the different steps in the algorithm.

6.4. The task parallel distributed memory C++ code

The management of the distributed environment and the communication between different computational nodes is implemented as a layer on top of SuperGlue, using MPI for the message passing calls. Because the distributed framework builds on the SuperGlue programming model, the user code for the distributed case is almost identical to the shared memory code. The main difference from the user perspective is that the handles and tasks now are upgraded to MPI_Handle and MPI_Task with some additional functionality. Therefore, when an MPI_Handle is created as opposed to the basic handle, the user must in addition specify the data owner process rank and the associated memory block. Each computational node submits the same sequence of tasks to the run-time system, which then decides which tasks will be executed locally and which to discard. The decision is based on the location of the output data from the task. The run-time system also detects for which tasks data transfers from other nodes are needed and inserts the appropriate communication tasks. All communication is point-to-point, and the implementation cannot currently take advantage of collective communication. This is not an issue for the SWE solver, as its communication pattern is primarily point-to-point. We are also developing another version of the distributed task-based framework, DuctTeip [46], with the same programming model, and also built on SuperGlue, but with slightly different design choices.

Figure 6 shows a simplified example of what the run-time system does in a situation when a transfer is needed. The transfer is detected when the Copy task is submitted. For the process with rank 0, the owner of the data x, a task that will send the data is submitted. When the send is completed, a FinishedSendTask updates the version number of the data. At rank 1, when the data has been received, a PublishTask is submitted that copies the data to the handle. In addition to keeping track of version numbers, the handles also register who last wrote to a certain data, and which nodes already have a copy of the latest version. When scheduling tasks, the run-time system gives priority to tasks that work on data that will be communicated.

```cpp
x.set_owner_rank(0);
y.set_owner_rank(1);
submit(new Write(x));
submit(new Copy(x, y));
submit(new Write(x));
```

Figure 6: The code to the left shows the tasks that the programmer submits, and the figure to the right shows also the additional tasks that are submitted by the run-time system and how they interact with the MPI process thread.
7. Experimental results

The numerical experiments are performed on the Tintin cluster at the UPPMAX facilities at Uppsala University. The cluster has 160 computational nodes equipped with dual AMD Opteron 6220 Bulldozer CPUs. Formally, this gives us 16 cores per node. However, it is important to note that each floating point unit is shared between two cores. Hence, for the type of floating point heavy simulations performed here, we can not expect a speedup of more than 8 times on one computational node.

The result section consists of two main parts, first we analyze the performance of the parallel implementation, and then we discuss the SWE solutions computed with the parallel RBF–FD method. The code that we used can be downloaded from GitHub\(^2\).

The convergence of the RBF–FD method is not the focus of this paper. However, we did perform some experiments for the second test case, which has a smooth solution, using the solution with \(N = 155\,718\) nodes as reference. With a stencil size \(n = 31\), we achieve almost the expected fourth order convergence. The larger stencil size \(n = 75 \pm 2\) leads to approximately fifth order convergence. This is lower than the expected seventh order, but the actual errors are about 8 times smaller than for \(n = 31\) and a given \(N\). More extensive experimental results on the convergence of RBF–FD methods applied to convective problems formulated over the sphere are reported in [20, 16].

7.1. Parameter values in the experiments

For the “flow over an isolated mountain” test case, we use stencil size \(n = 31\), hyperviscosity of order 4, and the amount of hyperviscosity \(\gamma = -0.05 \cdot N^{-4}\). For the “evolution of a highly non-linear wave” test case, we instead use stencil size \(n = 75\), hyperviscosity of order 8, and the amount of hyperviscosity \(\gamma = -0.3 \cdot N^{-8}\). Even at coarse resolutions, these terms are very small, and do not qualitatively affect the solution apart from ensuring stability.

The node sets, shape parameters, time steps, and block sizes used for the first test case are listed in Table 2. These are the values that were also used in [16]. The shape parameter values are chosen close to the ill-conditioning limit for optimal accuracy. However, for the largest problem size, due to the lack of uniformity of the icosahedral node set, we needed to increase the shape parameter value in order to have stability. This decreases the accuracy of this solution compared to having the expected value \(\varepsilon = 28\). The last column in the table shows how this affects the product \(\varepsilon h\) (which relates to the stationary scaling), assuming that \(h\) is proportional to \(1/\sqrt{N}\).

For the second test case we have also increased the shape parameter value for the largest problem. We had problems with the ICO node sets, and therefore we have used the smoothed DM nodes. Furthermore, we included a stencil selection procedure, where stencils of sizes \(n = 75 \pm 2\) are compared with respect to the total magnitude of the weights \(\sum |w_i|/\sqrt{n}\), and the one with smallest result is used. The rationale behind this procedure is that stencils with weights of large magnitude have in experiments been found to produce local instabilities. The method parameters that were used are given in Table 3.

\(^2\)https://github.com/tillenius/rgba-sw
Table 2: The node sets, shape parameter values, time steps, and block sizes used for the numerical simulations of the “flow over an isolated mountain” test case. The last column indicates how the relation between the shape parameter and the stencil size changes.

<table>
<thead>
<tr>
<th>N</th>
<th>Type</th>
<th>( \varepsilon )</th>
<th>( \Delta t )</th>
<th>( n_b )</th>
<th>( \varepsilon / \sqrt{N} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>6 400</td>
<td>MD</td>
<td>2.7</td>
<td>900</td>
<td>400</td>
<td>0.0338</td>
</tr>
<tr>
<td>25 600</td>
<td>MD</td>
<td>5.5</td>
<td>300</td>
<td>1280</td>
<td>0.0344</td>
</tr>
<tr>
<td>40 962</td>
<td>ICO</td>
<td>7.0</td>
<td>180</td>
<td>1280</td>
<td>0.0346</td>
</tr>
<tr>
<td>163 842</td>
<td>ICO</td>
<td>14.1</td>
<td>60</td>
<td>1600</td>
<td>0.0348</td>
</tr>
<tr>
<td>655 362</td>
<td>ICO</td>
<td>40.0</td>
<td>60</td>
<td>2048</td>
<td>0.0494</td>
</tr>
</tbody>
</table>

Table 3: The parameter values for the numerical simulations of the second test case. Note that the stencil size (and the shape parameter values) are larger than for the first test case.

<table>
<thead>
<tr>
<th>N</th>
<th>Type</th>
<th>( \varepsilon )</th>
<th>( \Delta t )</th>
<th>( \varepsilon / \sqrt{N} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>42 768</td>
<td>DM</td>
<td>12.6</td>
<td>180</td>
<td>0.0609</td>
</tr>
<tr>
<td>86 111</td>
<td>DM</td>
<td>18.0</td>
<td>90</td>
<td>0.0613</td>
</tr>
<tr>
<td>155 718</td>
<td>DM</td>
<td>24.3</td>
<td>60</td>
<td>0.0616</td>
</tr>
<tr>
<td>344 444</td>
<td>DM</td>
<td>36.2</td>
<td>60</td>
<td>0.0617</td>
</tr>
<tr>
<td>612 346</td>
<td>DM</td>
<td>60.0</td>
<td>30</td>
<td>0.0767</td>
</tr>
</tbody>
</table>

The time step was investigated in the previous paper [16], and similar time steps are used here for the smaller problem sizes. The proportionally larger time steps for the highest resolution runs were chosen close to the stability limit, in order to reduce the total run time.

7.2. Performance analysis

All experiments in this section are performed for the “flow over an isolated mountain” test case. In each experiment, we run the code for 100 time-steps, which is enough to compute the speedup, but does not correspond to a complete run. For the largest problem with time step \( \Delta t = 60s \), the total simulation time corresponding to 15 days requires 21 600 time steps. The sparse matrices are divided into blocks of different sizes depending on the problem size. These block sizes work well, but are not completely optimized. For the largest problem we use a block size of \( 2048 \times 2048 \), resulting in \( 320 \times 320 \) blocks. Blocks that are empty are discarded.

We start with experiments for the shared memory case as this is the basis for the distributed memory code and the performance properties will partly be inherited. Figure 7 visualizes an execution trace for the largest problem size considered, with \( N = 655 362 \) nodes. The tasks from different time steps do not need to synchronize globally, and the scheduler can take advantage of this to schedule any tasks that are ready to run, in parallel. The relatively small block size implies a large number of tasks, which means more time is spent on task generation and task management. In this case, 2% was spent on task generation, and 5% was idle time, which includes task management. When the block size is increased to \( 4096 \times 4096 \) elements, the task generation time is decreased to 0.5% of the run time, and the idle time is reduced to only 1.36%. In spite of this, the overall computation time is about 5% slower. The reason is that SuperGlue schedules...
Figure 7: Part of an execution trace for $N = 655,362$ nodes and 16 shared memory cores. Each triangle represents a task with start time at the base and finish time at the tip. Tasks of the same color belong to the same time-step. The longer tasks are the ones that submit the computational tasks.

Tasks by locality, and smaller working sets means that more data can be reused between tasks. Increasing the block size increased the number of cache misses with almost 20%.

Table 4 shows the resulting speedups for different problem sizes. The parallel code with execution time $T_p$ is compared both against the best serial (unblocked) code with execution time $T_{best}$, and against the parallel (blocked) code running on 1 core with execution time $T_1$. The estimated total execution time of the whole simulation $T_p^{tot}$ is also given as this is the time that the end user will experience.

<table>
<thead>
<tr>
<th>Size N</th>
<th>$T_p$ (s)</th>
<th>Speedup $T_{best}/T_p$</th>
<th>Speedup $T_1/T_p$</th>
<th>$T_p^{tot}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>6 400</td>
<td>0.1</td>
<td>7.0</td>
<td>8.7 (109%)</td>
<td>1s</td>
</tr>
<tr>
<td>25 600</td>
<td>0.5</td>
<td>6.0</td>
<td>6.8  (84%)</td>
<td>21s</td>
</tr>
<tr>
<td>40 962</td>
<td>0.9</td>
<td>5.3</td>
<td>6.0  (75%)</td>
<td>1m 3s</td>
</tr>
<tr>
<td>163 842</td>
<td>4.3</td>
<td>4.3</td>
<td>5.1  (64%)</td>
<td>15m 27s</td>
</tr>
<tr>
<td>655 362</td>
<td>18.3</td>
<td>4.1</td>
<td>4.9  (61%)</td>
<td>1h 5m 51s</td>
</tr>
</tbody>
</table>

The theoretical optimal speedup is 8 on the system we are using, and with the very low idle time seen in the execution trace, we might expect a nearly perfect speedup. However, this is where the bandwidth-bound characteristics of the sparse matrix-vector multiplication comes in. In current multicore processors, resources such as caches and memory bandwidth are shared between several cores. This may result in contention between the cores and consequently performance degradation when the resource is over-subscribed. The efficiency decreases as the problem size increases, because of the larger working set. Comparing the memory behavior when solving the largest and the smallest problems, the larger problem generated over two times as many last-level cache misses per instruction.

To evaluate the contention hypothesis, we compute the total time spent in the sparse
matrix-vector products in the sequential case and compare with the time in the parallel case (sum over the cores). We then find that the time in the parallel case is 152% of the sequential. The slowdown that we experience indicates that the cores are waiting for data to be delivered from memory. The problem is a combined effect of having few computations per data access, and not accessing data consecutively in memory.

How to schedule to decrease the resource contention is discussed in [37]. However, this approach only improves the situation if there are tasks that do not need the resource in question that can be interleaved with the resource bound tasks. This is not the case here, since all the heavy tasks are similar in nature.

Figure 8: The execution trace for $N = 655,362$ nodes using a distributed system with 10 computational nodes, each with 16 shared memory cores. The top subfigure shows the full execution trace for 100 time steps. There are 10 bands, each representing one computational node. The lower subfigure shows a part of the trace only for the first computational node (the lower band in the top subfigure). The colors represent the different time steps.

Figure 8 shows execution traces for the distributed memory code, running 100 time-steps of the problem with $N = 655,363$ nodes and block size $4096 \times 4096$. Thread 0 on each node is responsible for communication, which is why it executes fewer tasks. All MPI calls are non-blocking to overlap with task execution, and are not shown in the execution trace. Even when excluding thread 0, the idle time is here 20.40%, which is larger than in the shared memory case.

The distribution of data between the computational nodes, and consequently the distribution of work is currently static (determined before the execution starts). By implementing a load-balancing strategy at the node level, this can be improved. We are currently investigating mechanisms for automatic load-balancing over the nodes.
Table 5: Speedups for 100 time steps using the distributed memory code. We compare against the best shared memory code, $T_{\text{shared}}$, and execution on one node with the parallel code, $T_1$.

<table>
<thead>
<tr>
<th>Nodes</th>
<th>Block size</th>
<th>Execution time</th>
<th>Speedup</th>
<th>Efficiency</th>
</tr>
</thead>
<tbody>
<tr>
<td>$q$</td>
<td>$n_b$</td>
<td>$T_q$ (s)</td>
<td>$T_{\text{tot}}$</td>
<td>$T_{\text{shared}}/T_q$</td>
</tr>
<tr>
<td>1</td>
<td>2048</td>
<td>18.76</td>
<td>1h 7m 32s</td>
<td>0.98</td>
</tr>
<tr>
<td>2</td>
<td>2048</td>
<td>9.22</td>
<td>33m 11s</td>
<td>1.98</td>
</tr>
<tr>
<td>4</td>
<td>2048</td>
<td>4.48</td>
<td>16m 8s</td>
<td>4.08</td>
</tr>
<tr>
<td>6</td>
<td>2048</td>
<td>2.99</td>
<td>10m 45s</td>
<td>6.12</td>
</tr>
<tr>
<td>8</td>
<td>2048</td>
<td>2.22</td>
<td>8m 0s</td>
<td>8.23</td>
</tr>
<tr>
<td>10</td>
<td>2048</td>
<td>2.32</td>
<td>8m 21s</td>
<td>7.89</td>
</tr>
<tr>
<td>10</td>
<td>4096</td>
<td>1.83</td>
<td>6m 35s</td>
<td>10.00</td>
</tr>
</tbody>
</table>

Table 5 shows the speedup of the distributed code for different numbers of computational nodes. The execution time $T_q$ for running on $q$ computational nodes is compared both against the best shared memory code (without MPI) with execution time denoted by $T_{\text{shared}}$, and against the distributed code (with MPI) running on one node, $T_1$. We have also computed the total parallel efficiency as the speedup against the best serial code $T_{\text{best}}/T_q$, divided by the total (effective) number of cores, which is $8q$.

Looking at these numbers, we find that the scaling of the parallel code is very close to the theoretical best speedup of $q$ up until 10 nodes. This means that the total efficiency is more or less constant for all numbers of nodes. What happens at 10 nodes is that generating the tasks starts taking longer time than executing them. However, by increasing the block size, we can regain perfect scalability.

If we now look at the overall speedup that has been achieved, the distributed parallel code running on 10 computational nodes is currently 205 times faster than the original MATLAB implementation, 41 times faster than the optimized serial C++ code, and 10 times faster than the parallel shared memory code. Clearly, this enables both larger and faster SWE simulations using the RBF–FD method.

7.3. Shallow water equation simulation results

The “flow over an isolated mountain” test case is comparatively easy to solve, and visually, the solutions appear identical for all problem sizes. Figure 9 shows the solution for the lowest resolution $N = 6400$, which corresponds to 300 km, and the error measured against the solution with $N = 163842$, which corresponds to around 60 km. In [16], the finer solution was compared against a highly resolved discontinuous Galerkin solution showing that the error in the geopotential height was less than 1 m. As mentioned before, we do not carry out a full convergence study here, instead we focus on finding out if we can solve the larger problems with a qualitatively well behaved solution.

For the “evolution of a highly non-linear wave” test case it is much harder to get a physically correct solution. It also takes longer to solve the problem because we use a larger stencil and a smaller time step. In Figure 10, a number of solutions are displayed with a resolution of about 30 km for the largest problem. Solutions for smaller node numbers can be found in [16] for comparison. All the solutions have the expected qualitative behavior, but we can see that the features of the solution become sharper and better resolved with each refinement. Hence, the higher resolution that we could reach with the parallel code really has an effect.
We believe that the behavior can be improved by employing one of the stable methods, RBF-QR [19] or RBF-GA [21], to compute stencils for smaller \( \varepsilon \)-values. This is something that we will implement and test.

8. Conclusions

Our three main objectives in this paper were to show that the RBF–FD method can be efficiently parallelized, that task-based parallel programming is a suitable approach both for productivity and performance, and that the method can produce high quality solutions for the shallow water benchmark problems.

Starting from the original MATLAB implementation, which was shown to perform well sequentially compared with a discontinuous Galerkin solver [16], we have achieved 205 times speedup. Compared with the best sequential C++ code, we have a speedup of 41 using 10 computational nodes (160 cores, 80 FPUs) in a cluster. Part of this speedup comes from specific optimizations of the sparse matrix-vector multiplications. These are method and problem specific and can not be found in standard library routines. However, similar opportunities would arise in other RBF–FD solvers as well. Judging from the performance experiments, the code can scale to even larger computer systems, if needed.

The parallelization was done using the SuperGlue library for task-based parallel programming. The resulting code is quite similar to the original code except that the operations on matrices and vectors are blocked. The programmer does not need to pay any particular attention to the parallel aspects, except that in the distributed code, the data ownership is specified. The block size does affect performance to some extent. Overall, the productivity goal is met compared with the effort it would take to hardcode all interactions and schedules.

The task-based approach resulted in a very efficient code. There is very little idle time, and the performance losses that are observed are due to resource sharing, which cannot be completely avoided for a multicore based system. We have no reason to believe that another parallel approach would be more efficient.
We were able to compute highly resolved solutions to the shallow water benchmark problems. The second test case with the highly non-linear wave is quite sensitive to the method parameters and to the node layout. We believe that local node refinement could prove beneficial for stability, since all features would be resolved properly. However, this still needs to be investigated.
Appendix A. Sample MATLAB and C++ code

Here we show the main function of the original MATLAB program together with the shared memory C++ implementation, to illustrate that the difference in code complexity is relatively small.

Listing 1 shows the main computational steps of the MATLAB implementation. First the necessary RBF–FD matrices are set up, and then the time stepping is performed in a loop. The variable \( H \) is an \((N \times 4)\) matrix with the node functions of the four solution components at the current time step as its columns.

```matlab
% Build differentiation matrices
% and hyperviscosity operator
[DPx, DPy, DPz, L] = rbfmatrix_fd();

for i=1:timesteps
% Runge-Kutta
F1 = dt*rhs(H);
F2 = dt*rhs(H + 0.5*F1);
F3 = dt*rhs(H + 0.5*F2);
F4 = dt*rhs(H + F3);
H = H + (1/6)*(F1 + 2*F2 + 2*F3 + F4);
end
```

Listing 1: Excerpt from the main program of the MATLAB implementation.

Listing 2 shows the SuperGlue user code for the blocked matrix-vector multiplication. First, the involved blocked data structures with \( nb \) blocks in each dimension are created. Each block of these data structures contains a SuperGlue handle. Then the task to multiply a single block is defined. The data accesses are registered in the constructor and the run method (which is not displayed here) implements the computational kernel. Finally, a helper function is defined, which submits the block multiplication tasks in a loop.

```matlab
BlockedMatrix D(nb,nb);
BlockedVector H(nb);
BlockedVector T(nb);

struct MultTask : public Task {
    mult(VectorBlock &T, MatrixBlock &D, VectorBlock &H) {
        registerAccess(add, T.handle);
        registerAccess(read, D.handle);
        registerAccess(read, H.handle);
    }
    void run() { /* T += D*H */ }
};

void mult(BlockedMatrix &D, BlockedVector &H, BlockedVector &T) {
    for (int i = 0; i < nb; ++i)
        for (int j = 0; j < nb; ++j)
            submit(new MultTask(T(i), D(i,j), H(j)));
}
```

Listing 2: Blocked matrix-vector multiplication using SuperGlue. Note that the actual multiplication kernel is not shown.
To see how we can build a more complete application, we also show the code for the whole time-stepping loop in Listing 3. GenTasks is a task that submits other tasks. It calls a number of subroutines, which are listed below, that in turn submits the computational tasks. When all task for one time-step have been submitted, a new instance of GenTasks is added.

```c++
// Runge-Kutta step
void GenTasks::run() {
    f(F1, H); // F1 = f(H)
    add(H1, H, 0.5*dt, F1); f(F2, H1); // F2 = f(H + 0.5*dt*F1)
    add(H2, H, 0.5*dt, F2); f(F3, H2); // F3 = f(H + 0.5*dt*F2)
    add(H3, H, dt, F3); f(F4, H3); // F4 = f(H + dt*F3)

    step(H, F1, F2, F3, F4); // H = H + dt/6*(F1+2*F2+2*F3+F4)
    submit(new GenTasks(H)); // Generate new tasks for next step
}

// evaluate \partial H/\partial t
void f(dH, H) {
    mult(T, D, H); // T = D*H
    rhs(dH, H, T); // dH = ...
}
```

Listing 3: The main loop in the task parallel code. Note that all the subroutines here submit tasks.

The implementation of the `mult` subroutine was indicated in Listing 2. The other helper routines are displayed in Listing 4.

```c++
void add(Htmp, a, H) {
    for (int r = 0; r < nb; ++r)
        submit(new AddTask(Htmp(r), a, H(r)));
}

void rhs(dH, H, T) {
    for (int r = 0; r < nb; ++r)
        submit(new RHSTask(H(r), T(r)));
}

void step(H, F1, F2, F3, F4) {
    for (int r = 0; r < nb; ++r)
        submit(new StepTask(H(r), F1(r), F2(r), F3(r), F4(r)));
}
```

Listing 4: The helper tasks that submit the computational tasks for each block of the matrix H.


URL http://doi.acm.org/10.1145/2597652.2597656
[30] Y. J. Lee, G. J. Yoon, J. Yoon, Convergence of increasingly flat radial basis interpolants to poly-


