

WAVETRISK-1.0: an adaptive wavelet hydrostatic dynamical core

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Abstract. This paper presents the new adaptive dynamical core wavetrisk. The fundamental features of the waveletbased adaptivity were developed for the shallow water equation on the β plane and extended to the icosahedral grid on the sphere in previous work by the authors. The threedimensional dynamical core solves the compressible hydrostatic multilayer rotating shallow water equations on a multiscale dynamically adapted grid. The equations are discretized using a Lagrangian vertical coordinate version of the dynamico model. The horizontal computational grid is adapted at each time step to ensure a user-specified relative error in either the tendencies or the solution. The Lagrangian vertical grid is remapped using an arbitrary Lagrangian-Eulerian (ALE) algorithm onto the initial hybrid σ -pressurebased coordinates as necessary. The resulting grid is adapted horizontally but uniform over all vertical layers. Thus, the three-dimensional grid is a set of columns of varying sizes. The code is parallelized by domain decomposition using mpi, and the variables are stored in a hybrid data structure of dyadic quad trees and patches. A low-storage explicit fourthorder Runge-Kutta scheme is used for time integration. Validation results are presented for three standard dynamical core test cases: mountain-induced Rossby wave train, baroclinic instability of a jet stream and the Held and Suarez simplified general circulation model. The results confirm good strong parallel scaling and demonstrate that wavetrisk can achieve grid compression ratios of several hundred times compared with an equivalent static grid model.

1 Introduction

Atmospheric flows are intrinsically non-stationary and multiscale. They are characterized by length scales varying from millimetres to thousands of kilometres and timescales from seconds to decades. Effective climate modelling requires long runs (typically decades) that resolve the dynamically significant scales of motion. However, the smallest significant scales of motion are highly intermittent in space and non-stationary in time. Especially at higher resolutions, the portion of the flow with active small scales is relatively small. Attempting to resolve these important small scales with a uniform grid is necessarily inefficient. Either we need to choose a very fine grid to resolve the smallest scales of interest, which limits simulation times, or we use a coarser grid that filters out these small scales, potentially losing important dynamics. In addition, a uniform resolution does not allow effective control of numerical error. Some regions will be under-resolved and others will be over-resolved. A static computational grid also uses computational resources inefficiently by spending CPU time computing dynamically inactive regions. Ideally, the numerical error should scale predictably with the number of computational elements (e.g. nodes) which should, in turn, be proportional to CPU time. This paper introduces a new wavelet-based adaptive dynamical core, wavetrisk, that attempts to achieve these goals.

Behrens (2009) provides a good overview of early developments in adaptive atmospheric modelling, including early work by Skamarock et al. (1989) and previous work using moving grids to model tropical cyclones (Harrison, 1973; Jones, 1977). Since this early work, adaptive mesh refinement (AMR) has found many applications, e.g. fluid dynamics (Popinet and Rickard, 2007) and astrophysics (Mignone et al., 2012; Bryan et al., 2014). In such applications, the main added value of AMR is its ability to accurately capture events that are highly localized in space and time at a relatively low computational cost. Jablonowski et al. (2009) evaluated a block-structured AMR method for scalar twodimensional transport on the sphere. Refinement and coarsening levels are constrained so that there is a uniform 2:1 mesh ratio at all fine-grid/coarse-grid interfaces. In a series of test cases, they find that the additional resolution helps preserve the shape and amplitude of the transported tracer while saving computing resources in comparison to uniform-grid model runs. Kopera and Giraldo (2014) evaluated the performance of AMR in a discontinuous Galerkin-based implicitexplicit (IMEX) model on a planar two-dimensional grid. They found that AMR could provide up to a $15 \times$ speedup with minimal overhead. Ferguson et al. (2016) analyzed the performance of an AMR model for the shallow water equations on the cubed sphere. Their model is built using the general purpose Chombo-AMR toolbox of finite difference and finite volume methods for the solution of partial differential equations on block-structured adaptively refined rectangular grids. They test the model for one or two levels of refinement with refinement ratios of $\times 2$, $\times 4$ and $\times 8$ between each level. In comparison, wavetrisk currently uses a fixed refinement ratio of $\times 2$, but has been tested for up to six levels of refinement. Ferguson et al. (2016) conclude that AMR can be effective provided that a sufficiently fine coarsest grid is selected.

Beyond its ability to accurately capture events that are quite localized in space and time at a low cost, several additional properties are required for AMR to be an attractive option for atmospheric applications. Firstly, AMR must demonstrate its ability to accurately simulate complex threedimensional flows, where a large number of important features such as cyclones and waves continuously appear, move and disappear. Wavetrisk has already demonstrated this ability for (two-dimensional) shallow water flows. Furthermore, especially for applications to climate modelling, the robustness of the method over long timescales and its ability to capture accurate statistics should be shown. Finally, an efficient parallel implementation must be developed in order to compete with state-of-the art operational models. Skamarock and Klemp (1993) presented pioneering initial results from area-limited two- and three-dimensional AMR codes. Popinet et al. (2012) reported results for a threedimensional AMR code on the cubed sphere at a conference, and Ferguson (2018)'s thesis also describes the development of a global three-dimensional AMR code (based on the CHOMBO library). However, although these groups made significant progress in developing global three-dimensional AMR, to the best of our knowledge, no AMR for complex three-dimensional atmospheric flows has been fully documented and analyzed in a peer-reviewed publication.

A key element to the robustness and accuracy of any AMR method is its refinement criterion, which decides when and where to coarsen or refine the computational grid. Ferguson et al. (2016) evaluate several ad hoc criteria and identify appropriate ones for their numerical experiments. However, they do not find any "clear strategy for establishing the best general refinement criteria". In contrast, wavetrisk uses objective and clearly defined refinement criteria which control the multiscale relative error of the solution or of its tendencies as measured directly by the wavelet coefficients. Although this strategy is objective and clearly defined, since it is based solely on the normalized local approximation error of the solution or tendencies, it is not sufficient to control more qualitative measures of error, such as the location of a vortex. In particular, this approach would need to be augmented when subgrid parameterizations are included, such as for cloud formation or turbulence. Other refinement indicators would be appropriate when considering p refinement (Naddei et al., 2019).

In the following section, we review briefly the basic properties and computational features of the wavelet-based dynamical adaptivity laid out in Dubos and Kevlahan (2013) and Aechtner et al. (2015). Section 3 summarizes the discrete equations solved by the model and presents the arbitrary Lagrangian–Eulerian (ALE) vertical coordinate. Section 4 applies the principle of wavelet-based adaptivity to the three-dimensional model. In Sect. 5, implementation details are given and the adaptive solver is evaluated using three test cases of increasing complexity. Section 6 concludes the paper.

2 The adaptive wavelet method

2.1 Wavelet adaptivity on the plane

The foundations for wavetrisk were set out in Dubos and Kevlahan (2013) for the shallow water equations on the β plane. We used the TRiSK discretization scheme on the hexagonal-triangular C grid proposed by Ringler et al. (2010) because of its excellent mimetic properties. This scheme is second order in space in the L^2 norm. The principal goal of our approach was that the adaptivity should be an overlay on the flux-based discretization of any order. Mimetic properties (e.g. mass conservation) should be preserved by the adaptivity and the discretizations should be unchanged. The building blocks of the method are one-scale operators (in this case, the TRiSK discretization) and twoscale operators between a fine scale j + 1 and a coarse scale *j*. To conserve the mimetic properties of the TRiSK scheme, the restriction operators from scale j + 1 to scale j and the discrete differential operators (div, grad, curl) must satisfy the following commutation properties:



Figure 1. The regular C grid on the plane. Vorticity is located on the primal grid of triangles at points x_v and mass located on the dual grid of hexagons at nodes x_i . The velocities/fluxes are located on edges x_e . A multiscale hierarchy of nested refined C grids is generated by bisecting the triangle edges.

$$R^{j}_{\mu} \circ \operatorname{div}^{j+1} = \operatorname{div}^{j} \circ R^{j}_{\mathrm{F}} \text{ conserves mass,}$$
(1)

)

 $\operatorname{curl}^{j} \circ R_{u}^{j} = R_{\zeta}^{j} \circ \operatorname{curl}^{j+1}$ conserves circulation, (2)

$$\operatorname{grad}^{j} \circ R_{\mathrm{B}}^{j} = R_{\boldsymbol{u}}^{j} \circ \operatorname{grad}^{j+1}$$
 no spurious vorticity, (3)

where R_{μ} is mass density (or height) restriction, $R_{\rm F}$ is the flux restriction, R_u is the velocity restriction, R_{ζ} is the circulation (vorticity) restriction, and $R_{\rm B}$ is the Bernoulli function restriction. The third commutation relation ensures that a flow with uniform potential vorticity remains uniform under the advection by an arbitrary velocity field (i.e. vorticity is advected like a tracer).

The C grid is a staggered grid where vorticity is located on triangles (the primal grid), and mass is located on the hexagons formed from the bisectors of the triangle edges (the dual grid). Velocity is located on the edges of triangles, which are also the perpendicular bisectors of the hexagonal edges. (Note that we have chosen the opposite notation to Ringler et al. (2010) and Dubos et al. (2015) since in the multiscale case the triangle grid is generated first by repeated bisection from the icosahedron and the hexagon grid is generated as the dual grid of the triangles.) The regular C grid is shown in Fig. 1. Starting with a coarsest grid, a multiscale hierarchy of primal grids is constructed by bisection of the triangle edges. The dual grid of hexagons is constructed from the perpendicular edge bisectors of the primal grid. On the plane, both the hexagons and triangles are nested and regular. We will see below that this is not the case for the sphere.

This hierarchy of computational grids leads to a so-called wavelet multiresolution analysis (MRA), i.e. the nested sequence of approximation subspaces used to construct the second-generation biorthogonal wavelets (Sweldens, 1998) that are the basis of the grid adaptivity algorithm. The MRA provides a sequence of smooth approximations of a function f(x) on each grid level j and an associated sequence of "details" which give the differences between the approximation at a fine level j + 1 and a coarse level j. The details are effectively the interpolation errors between scales j + 1 and j. The smooth approximation at scale j has a basis of scaling functions $\{\phi_k^j(x)\}$, while the details between scales j + 1 and j have a basis of wavelets $\{\psi_k^j(x)\}$.

If the wavelet coefficient at a particular position k and scale j is sufficiently small, i.e. $|\psi_k^j(x)| < \varepsilon$, then the value of f(x) is well approximated by the function $f_{\geq}(x)$ interpolated from neighbouring scaling function values at the same scale. If the wavelet coefficient is large, the value of the wavelet coefficient (i.e. the details) must be retained and added to the interpolated value to obtain an sufficiently accurate approximation. Neglecting the small wavelet coefficients (and associated grid points) generates a multiscale hierarchy of adapted grids. The commutation relations (Eqs. 1–3) ensure that the mimetic properties of the discretization are also satisfied on the adapted grid.

One can prove that this nonlinear wavelet filtering provides error control:

$$||f(x) - f_{\geq}(x)||_{\infty} = O(\varepsilon), \tag{4}$$

$$\mathcal{N} = O\left(\varepsilon^{-1/2N}\right),\tag{5}$$

$$||f(x) - f_{\geq}(x)||_{\infty} = O\left(\mathcal{N}^{-2N}\right),$$
 (6)

where \mathcal{N} is the number of grid points retained on the adapted grid and N is the order of interpolation. Thus, we can also estimate how many grid points (i.e. computational elements) are required to obtain an approximation with a desired error ε . Setting ε therefore determines the numerical error of the approximation (the tolerance) and also determines the number of grid points in the adapted grid. In principle, it is not necessary to set a maximum resolution scale J since it is determined automatically by ε . Note that an approximation $f_{\geq}^{j}(x)$ at each scale j is provided by the set of scaling functions $\{\phi_{k}^{j}\}_{>}$ and their coefficients.

In previous work, we have shown that the CPU time τ_n per time step is directly proportional to the number of active grid points \mathcal{N} . Combined with the error estimates (Eq. 6), this shows that adaptive wavelet method is efficient in the sense mentioned in the introduction: the CPU time scales with the specified error and the error is controlled regardless of the structure of the flow. This is not true for non-adaptive methods where there is no explicit link between the actual numerical error and the number of grid points. At best, the approximation error of the discretization on a particular static grid gives only an upper bound on the actual error. Even this upper bound may not be satisfied if the a priori estimate of the smallest structures (e.g. strongest gradients) that determined the chosen grid resolution did not properly take into account intermittency and rare events. For example, Yakhot and Sreenivasan (2005) argue that temporal and spatial intermittency mean that turbulent flows require far higher resolution than that found using the usual Kolmogorov-scale-based estimate, $\Delta x \sim Re^{-3/4}$.

The above adaptive algorithm controls the error of the solution at each time step, but it does not allow for dynamics. The solution can change over one time step from t^n to $t^n + \Delta t$ by translating at the same scale, coarsening (gradient weakens) or refining (gradient strengthens). If the Courant–Friedrichs–Lewy (CFL) criterion is 1, i.e. $\Delta t <$ $\Delta x_{\min}/||u||_{\infty}$, the solution can translate by a maximum of one grid point at the same scale over one time step. If the nonlinearities in the governing equations are quadratic, the active scale can increase by a maximum of a factor of 2 from *i* to i + 1 over one time step. To allow for these changes, an "adjacent zone" is added to the set of active wavelet coefficients $\{\psi_k^J\}_{\geq}$ that includes its nearest neighbours in both position and scale (Liandrat and Tchamitchian, 1990). The adaptive grid must also satisfy the perfect reconstruction criterion: there must be sufficient scaling functions (grid points) to construct the wavelets. There must also be sufficient grid points present at each scale *j* to construct the required TRiSK differential operators (by interpolation, if their associated wavelets are inactive). Once the new adapted grid has been constructed, the prognostic variables are interpolated onto the new grid by performing an inverse wavelet transform. After the solution is advanced, the wavelets on the union of the adapted grid and adjacent zone are again filtered using the threshold ε to obtain a new set of active wavelets at time $t^n + \Delta t$.

Because we use a staggered grid, the adaptive wavelet algorithm described above differs fundamentally from previous adaptive wavelet collocation methods (e.g. Mehra and Kevlahan, 2008; Kevlahan and Vasilyev, 2005; Roussel and Schneider, 2010; Schneider and Vasilyev, 2010). Since mass and velocity are located at different points, we must construct two distinct wavelet transforms: a scalar-valued wavelet transform for mass density μ and a vector-valued wavelet transform for velocity u. To control the errors in the tendencies, the corresponding thresholds ε_{μ} and ε_{u} must be properly scaled. The details of these scalings in the inertia–gravity wave and geostrophic regimes are given in Dubos and Kevlahan (2013).

Finally, note that ideally the time step should also adapt to the local grid scale. In other words, the solution at each scale should be advanced on the time step Δt^{j} appropriate for that scale. Although this scale-dependent time stepping is optimal, in practice it does not provide much advantage unless only a small portion of the total active grid points is at the finest scale, which is not usually the case. Domingues et al. (2008) developed a second-order Runge–Kutta scaledependent time stepping and McCorquodale et al. (2015) extended scale-dependent time stepping to arbitrary order. For simplicity, we have decided not to implement scaledependent time stepping in wavetrisk, although it could be added in the future.

The ability of the adaptive wavelet method to control the errors of the tendencies was verified in Dubos and Kevlahan (2013), and the computational performance and parallel efficiency of the method on the sphere were confirmed in Aechtner et al. (2015). Since the three-dimensional hydrostatic code uses the multilayer shallow equations and horizontal adaptivity only, these properties are inherited by wavetrisk.

The prototype matlab serial implementation of the adaptive algorithm in the plane is relatively straightforward because the computational grid is uniform and we did not parallelize the algorithm. In the following section, we review the extension of the algorithm to the sphere.

2.2 Wavelet adaptivity on the sphere

We have reviewed the essential elements of the adaptive wavelet method in the previous section; however, to be practically useful for a dynamical core, the method must be extended to the sphere and parallelized. The extension to the sphere presents numerous technical challenges. First, the grid is non-uniform, each computational element is geometrically unique, and the icosahedral grid on the sphere has 12 pentagonal dual grid cells (a sphere cannot be tiled uniformly). Secondly, the repeated bisection of the primal triangular grid used to generate the multiscale grid structure produces a grid that is increasingly distorted near the edges of the original icosahedron. Finally, the dual hexagonal grid is no longer nested between successive scales because the triangles used to generate it are not equilateral. This makes it complicated to construct the flux restriction from fine to coarse scales because we must keep track of small overlapping regions. In the following, we focus on the most important modifications necessary to deal with the non-uniform geometry and to parallelize the code in the following.

The wavelet transform on the non-adaptive primal (triangular) icosahedral grid is shown in Fig. 2. Table 1 gives the properties of the multiscale grids for each level of refinement J.

The basic principles of the adaptive transforms are the same as on the plane; however, we must take account of the fact that the geometry of the grid is irregular. The TRiSK discretization (Ringler et al., 2010) is second-order accurate for equilateral triangles, but drops to first-order accurate when the triangles are far from equilateral. This problem is not restricted to adaptive methods but is an issue for all methods on fine icosahedral grids. To deal with this problem, we first generate a coarsest grid (e.g. $J_{min} = 5$ levels of bisection) and then optimize its geometry to ensure that the bisection of primal and dual edges is as close as possible. This optimized primal grid is then bisected as needed to generate the multiscale grids required for the wavelet transform. The default method is to read in optimized grids provided by Heikes

Table 1. Hierarchy of multiscale primal (triangle) grids derived by edge bisection from the icosahedron at scale J = 0. $N = 10 \times 4^J + 2$ is the number of computational elements (lozenges), $\overline{\Delta x} = (4/\sqrt{3}4\pi^2a^2/20/4^J)^{1/2}$ is the average triangle edge length on Earth. Each computational element is made up of one node (for scalars), three edges (for velocities) and two triangles (circulation). Therefore, the total number of data elements is 5N per vertical level. Note that there are two exceptional points to deal with the poles when the icosahedron is unfolded into 10 lozenges. Typically, the coarsest (optimized) level is $4 \le J \le 9$ with three to five levels of refinement.

J	Ν	$\overline{\Delta x}$ (°)	$\overline{\Delta x}$ (km)
0	12	65.9	7317
1	42	34.1	3790
2	162	17.2	1913
3	642	8.6	960
4	2562	4.3	480
5	10242	2.2	240
6	40 962	1.1	120
7	163 842	0.54	60
8	655 362	0.27	30
9	2 621 442	0.14	15
10	10485762	0.068	7.5

et al. (2013). As an alternative, we can also use the grid optimization proposed by Xu (2006), although it produces less optimal grids.

A more fundamental challenge particular to this adaptive flux-based method on staggered grids is that the dual hexagonal grids on two successive scales are no longer nested as they are on the plane. This means that we must keep track of the various configurations of small overlapping hexagons when computing the flux restriction (see Sect. 4.3 and Fig. 4 of Aechtner et al., 2015, for more details). Note that the primal grid of triangles remains nested on the sphere, which means that the restrictions of velocity, Bernoulli and circulation are straightforward.

2.3 Parallelization and data structure

The parallelization and data structure must take account both the icosahedral geometry and topology of the spherical discretization and the fact that the grid is adaptive and multiscale. The C grid is stored as a regular data structure by grouping one node (mass and other scalars), two triangles (circulations) and three edges (velocities) into one computational element, a "lozenge" as shown in Fig. 3. The icosahedron is composed of 20 triangles grouped into 10 lozenges. Therefore, a grid resulting from refining an icosahedron can be divided into 10 subgrids, each of which can be stored and accessed in a regular fashion. Note that at the edges of the lozenges the two adjacent regular grids of the original icosahedron are rotated with respect to each other. This is dealt with by surrounding the 10 lozenge subdomains by ghost/halo cells. The halo cells are also used for parallel communication during boundary updates between cores.

The most natural way to store two-dimensional data with a dyadic multiscale structure is to use a quad tree, where each branching represents a new finer scale. However, in an adaptive method, some branches are pruned and accessing neighbours would require wasteful traversing of the tree structure. We decided to use a hybrid data structure where each quad tree terminates in a patch. The patches are small 4×4 or 8×8 regular grids. This structure reduces the number of levels in the quad tree and makes it more computationally efficient to find neighbours. A similar hybrid approach was used by Behrens (2009) and Hejazialhosseini et al. (2010). Using patches increases memory slightly because inactive elements are stored. It is also possible to try to optimize the patch size for best computational performance for a particular problem, although 8×8 appears optimal in most cases. Note that the patches do not affect the results of the computation; they just improve computational efficiency.

The domain decomposition used for parallelizing the computation is based on distributing the lozenge subdomains on the coarsest level J_{min} (and their associated children) to different cores. Each core can compute several subdomains and having several small subdomains per core can improve cache efficiency. Note that there are $10 \times 4^{J_{min}-(P+1)}$ subdomains at the coarsest level with a patch size of $2^P \times 2^P$. For example, if $J_{min} = 7$ and P = 2, the coarsest scale contains $N_D = 2560$ subdomains. Because the number of cores $N_{core} \le N_D$, large numbers of cores are usable only by large J_{min} .

In an adaptive simulation, each subdomain typically has a different number of active computational elements and thus generates different loads for communication and computation. This load imbalance between cores can seriously degrade parallel performance. To remedy this, we use a simple rebalancing algorithm to redistribute subdomains amongst the cores to produce a more balanced load. This rebalancing is done at each checkpoint save. The checkpoint saves the current load for each coarse subdomain, which is simply the number of active columns in that subdomain. This load information is then read when the code restarts. The rebalancing algorithm starts by trying to sequentially distribute the $N_{\rm D}$ coarse subdomains over the available ranks to satisfy a user-specified maximum load imbalance (i.e. (maximum load - average load) / average load - 1, currently set to 0.1)while ensuring that every rank has at least one domain and every domain is assigned to a rank. When a rank is "full" (i.e. the load is greater than the average load), the algorithm starts assigning subdomains to the next rank. If not all subdomains can been assigned to a rank, the acceptable maximum acceptable load imbalance is increased by a factor of 2 and a new attempt is made to distribute the subdomains to the available ranks. This maximum acceptable load imbalance is increased iteratively until all domains have been assigned to a rank.



Figure 2. Wavelet transform on a non-adaptive icosahedral grid with three scales.



Figure 3. Lozenge: basic computational element containing one node (for mass and other scalars), three edges (for velocities) and two triangles (for circulation). Separate wavelet transforms are provided for the nodes (scalar valued) and edges (vector valued). The adaptive grid consists of the significant nodes and edges, together with nearest neighbours in position and scale necessary for dynamics.

Every subdomain is extended to hold as many ghost/halo cells as necessary for the various required operators. The values at the halo cells are communicated as needed. Intracore communication is done by copying and inter-core communication is done using mpi. During grid adaption, new patches are added and removed as required and grid connectivity between domains is updated (via mpi as necessary). Critical communications are carried out locally point to point rather than using global communication where possible. Where possible, communication is non-blocking so that the computations can continue while communication is taking place in the background.

This parallelization algorithm is reasonably efficient for at least several hundred or a few thousand cores. Dubos and Kevlahan (2013) found that good weak parallel efficiency is possible with as few as 1300 computational elements per core in adaptive runs. The three-dimensional code has better parallel efficiency because the column structure of the data produces a higher computational load for each active grid element. We present some representative strong and weak parallel scaling results in Sect. 5.1.

3 Hydrostatic dynamical equations and ALE vertical coordinates

As mentioned in the introduction, we use the dynamico discretization of the three-dimensional hydrostatic multilayer shallow water equations (Dubos et al., 2015) in compressible form. The discrete dynamical equations are derived from the discrete Hamiltonian, which allows the construction of energy or potential enstrophy conserving equations. The prognostic variables are m_{ik} (mass), Θ_{ik} (mass-weighted potential temperature) and v_{ek} (velocity), where k labels a full vertical level, l an interface (half level) between full vertical levels, i an hexagonal or pentagonal cell, v a triangle and e an edge. In terms of the Hamiltonian, their evolution equations are

$$\frac{\partial m_{ik}}{\partial t} + \delta_i \frac{\partial H}{\partial v_{ek}} = 0, \quad \frac{\partial \Theta_{ik}}{\partial t} + \delta_i \left(\theta_{ek}^* \frac{\partial H}{\partial v_{ek}} \right) = 0, \tag{7}$$

$$\frac{\partial v_{ek}}{\partial t} + \left(\frac{f_v + \delta_v v_k}{\overline{m_{ik}}^v} \frac{\partial H}{\partial v_{ek}} \right)^\perp + \delta_e \frac{\partial H}{\partial m_{ik}} + \theta_{ek}^* \delta_e \frac{\partial H}{\partial \Theta_{ik}} = 0, \tag{8}$$

where δ_i , δ_e and δ_v are discrete divergence, gradient and curl operators yielding values at cells, edges and triangles, respectively, f_v is the Coriolis parameter, and θ_{ek}^* , $\overline{m_{ik}}^v$ are values of θ and *m* reconstructed at edges and triangles, respectively, by appropriate averaging. Indices than can be inferred may be omitted, as in $\delta_v v_k \equiv \delta_v v_{ek}$. Evaluating the Hamiltonian terms and discretizing, we obtain the inviscid discrete dynamical equations:

$$\frac{\partial \mu_{ik}}{\partial t} + \delta_i U_k = 0, \tag{9}$$

$$\frac{\partial \Theta_{ik}}{\partial t} + \delta_i (\theta_{ek}^* U_k) = 0, \tag{10}$$

$$\frac{\partial v_{ek}}{\partial t} + \delta_e B_k + \theta_{ek}^* \delta_e \pi_k + (q_k U_k)_e^{\perp} = 0,$$
(11)

where $\mu_{ik} = \rho_{ik} \Delta z_{ik}$, and we have assumed Lagrangian vertical coordinates (so the vertical mass fluxes are not present). Potential temperature $\theta_{ik} = \Theta_{ik}/\mu_{ik}$ and $\theta_{ek} = \overline{\theta_{ik}}^e$. μ_{ik} is a pseudo mass density (equivalent to $\rho_{ik} \Delta z_{ik}$), U_{ek} is the horizontal mass flux, B_{ik} is the Bernoulli function, $\pi_{ik} = \pi(\alpha_{ik}, \theta_{ik})$ is the Exner function, α_{ik} is the specific volume, and q_{vk} is the potential vorticity. In the compressible case we consider here, the Bernoulli function is given by

$$B_{ik} = K_{ik} + \overline{\Phi_{il}}^k, \tag{12}$$

where K_{ik} is the discrete kinetic energy computed from u_{ek}^2 using appropriate averaging, and Φ_{il} is the geopotential at vertical layer interfaces l. The discrete operators δ_i (divergence with result at a node), δ_e (gradient with result at an edge) and $(\cdot)^{\perp}$ (perpendicular flux) are defined as in Ringler et al. (2010) and the shallow water equation version of wavetrisk.

Each evaluation of the trend requires integrating vertically down to find the surface pressure and then integrating up to find the Exner function and geopotential (and hence the Bernoulli function).

The choice of Lagrangian vertical coordinates (rather than mass based) is simple, computationally efficient and especially well suited for ocean modelling since it virtually eliminates numerical vertical diapycnal diffusion, unlike a z coordinate. This will become important when we develop the ocean version of wavetrisk (see Kevlahan et al., 2015). The vertical coordinates are pressure based and may either be evenly spaced or hybrid (σ). As the flow develops, the vertical levels expand and contract, which can lead to loss of accuracy or even negative mass (if a layer collapses to zero thickness). To avoid this problem, we remap the vertical coordinates back to the original coordinates either every time step or every 10 time steps or so. The remapping takes about 7 % of CPU time if done every time step.

Remapping Lagrangian vertical coordinates is a common technique in ocean modelling (e.g. Petersen et al., 2015), where the target grid is updated at each time step to optimize numerical accuracy and stability (e.g. the target grid may be based on approximately isopycnal or isentropic coordinates) and is referred to as an ALE coordinate system. Optimizing the target grid in wavetrisk would add a sort of vertical r adaptivity in addition to the wavelet-based horizontal hadaptivity, where the number of vertical levels would remain constant but their locations would be chosen optimally. For example, Kavcic and Thuburn (2018) choose the target grid levels at each time step to keep the vertical levels close to isentropic. This requires solving a small elliptic optimization problem at each grid level. In principle, it is also possible to include vertical *h* adaptivity by locally de-activating certain vertical layers if the vertical interpolation errors are small. We have tested wavetrisk with a variety of piecewise constant, piecewise linear, piecewise parabolic and piecewise quartic remapping schemes, modified from packages supplied by Shchepetkin (2001) and Engwirda and Kelley (2016, https://github.com/dengwirda/PPR, last access: 25 November 2019). The user can select no limiter, a monotone limiter or a weighted essentially non-oscillatory (WENO) limiter. Any of these remapping algorithms can be selected simply by changing a parameter.

Mass density μ , potential temperature θ and velocities u, v, w are remapped onto the original hybrid pressure coordinates. This remapping scheme conserves mass, potential temperature and divergence and appears to perform well. We also tested Lin (2004)'s scheme, which remaps momentum and total energy, but it proved to be less stable.

We found that simple piecewise constant remapping gives qualitatively incorrect results for zonally averaged statistics in the Held and Suarez (1994) test case, but that it is sufficient for the mountain-induced Rossby wave and baroclinic instability test cases. Piecewise linear and piecewise parabolic remapping give qualitatively accurate results. The target grid could be optimized at each remap, as in ocean models and as explored by Kavcic and Thuburn (2018) for the endgame general circulation model (GCM). However, the current procedure gives good results in the test cases we have examined.

The dynamical equations are advanced in time using the fourth-order four-stage low-storage Runge-Kutta routine used in dynamico (Dubos et al., 2015). Various strong stability-preserving schemes (e.g. the third-order three-stage scheme RK33ssp, RK45ssp) are also available as options (Spiteri and Ruuth, 2002).

In many cases, the code runs stably without any additional diffusion or filtering of small scales. However, in rare cases, the code crashes due to numerical instability. In order to improve stability, a regular or second-order hyperdiffusion term is added to the dynamical equations for the prognostic variables to damp the largest wavenumbers (both the divergent and vortical modes of the momentum equation are damped). Using the discrete operator notation of Dubos et al. (2015), the diffusion terms for scalars, divergence and vorticity are, respectively,

$$D_{\phi} = K_{\phi} \delta_i \left[\frac{l_e}{d_e} \delta_e \left(\frac{\phi}{A_i} \right) \right], \tag{13}$$

$$D_{\delta} = K_{\delta} \delta_e \left[\frac{1}{A_i} \delta_i \left(\frac{l_e}{l_d} v_e \right) \right], \tag{14}$$

$$D_{\omega} = K_{\omega} \delta_{e} \left[\frac{1}{A_{v}} \delta_{v} \left(v_{e} \right) \right], \tag{15}$$

where d_e is a triangle edge length (primal grid), l_e is a hexagon edge length (dual grid), A_i is a hexagon area, A_v is a triangle area. These discretizations correspond to the continuous differential operators $\nabla \cdot \nabla(\phi)$, $\nabla(\nabla \cdot u)$ and $\nabla \times (\nabla \times u)$, respectively.

A general *p*th-order hyperdiffusion operator is defined as an iterated Laplacian operator Δ^p corresponding to D_{ϕ}^p , D_{δ}^p , D_{ω}^p . We choose either p = 1 (regular diffusion) or p = 2 hyperdiffusion. Diffusion may be applied at each time step or every $N_{\text{diff}} > 1$ time steps (where N_{diff} is limited by viscous stability in time). The diffusion coefficient *K* is chosen to give the same amount of damping over a time step:

$$K = \frac{\overline{\Delta x}^{2p}}{\Delta t} C N_{\text{diff}},\tag{16}$$

where $\overline{\Delta x} = (4\pi a^2/(10 \ 4^J + 2))^{1/2}$ is the average grid scale and *C* is an empirical constant chosen to ensure stability, which may be different for different variables.

Note that in addition to ensuring stability, adding diffusion can improve the efficiency of the adaptivity by damping out small fluctuations that might otherwise produce some local grid refinement. This effect is especially significant when adapting on the wavelets of the tendencies. We present results both with and without explicit diffusion.

4 Adaptivity

To extend the adaptive algorithm for the two-dimensional shallow water equations reviewed in Sect. 2.1 to the threedimensional case, we simply apply the two-dimensional algorithm to each vertical layer in turn and then define the adapted grid to be the union of the adapted grids over all vertical layers k = 1, ..., N.

After the time step and vertical regridding have been completed, the wavelet transforms of mass density μ , massweighted potential temperature Θ and velocity u are calculated for each vertical level. Then, grid points are labelled active if their associated wavelet coefficient has a magnitude greater than or equal to the appropriate relative error threshold. Nearest neighbours are added to the adjacent zone in position and scale. Note that a node is labelled active if any of its associated scalar wavelets (i.e. mass wavelet or mass-weighted potential temperature wavelet) is significant in any vertical layer. An edge is labelled active if its associated vector wavelet (i.e. velocity wavelet) is active in any vertical layer. Grid points are added as necessary to satisfy the perfect reconstruction criterion (so the grid points necessary to compute the wavelet coefficients are present). Finally, grid points required for the TRiSK operators are labelled. An inverse wavelet transform interpolates the solution conservatively onto the new adapted grid. The adaptivity algorithm is summarized in Algorithm 1.

Algorithm 1 Adaptive grid algorithm. Executed after the time step and vertical grid remapping is complete. Note that wavelets are calculated and filtered at all vertical levels so that the final adapted grid is the union of adapted grids over all vertical levels.

After time step and vertical grid remapping have been completed **if** adapt on trend **then**

Compute trend

Compute trend wavelets

end if

for k = 1to N do {loop over all vertical levels}

Update relative error thresholds ε_{μ} , ε_{Θ} , ε_{u} by computing appropriate norms at each vertical level

end for

Label all active nodes and edges at coarsest scale $j = J_{\min}$ as adjacent zone

Label all nodes and edges for scales $j > J_{\min+1}$ as **inactive** for k = 1 to N do {check all vertical levels}

for j = J - 1 to J_{\min} do {check wavelets at all scales} label as active all nodes with an associated wavelet coefficients $\geq \varepsilon_{\mu}$ OR $\geq \varepsilon_{\Theta}$ label as active all edges with an associated wavelet coefficients $\geq \varepsilon_{\mu}$

end for

end for

Add **nearest neighbours** of active grid points at **coarser scale** to **adjacent zone**

Add nearest neighbours of active grid points at same scale to adjacent zone

Add/remove **patches** as required for neighbours at finer scales for j = J - 1 to J_{min} do

add neighbours of active grid points at **finer scales** end for

Perfect reconstruction criterion: add grid points required to compute active and adjacent zone wavelets

Label grid points required by **TRiSK operators** to compute trend at active and adjacent zone wavelets

Set all **wavelets** not active or in the adjacent zone to zero **Inverse wavelet transform** of solution onto new adapted grid

The adaptive algorithm produces an adapted grid consisting of vertical columns of varying horizontal size. In practice, a maximum scale J is usually set based on available computational resources and user requirements. This "column adaptivity" approach is not optimal for vertically tilted structures, but it provides much better load balancing and is far simpler than dealing with fully three-dimensional adaptivity. In addition, we can take advantage of an ALE formulation for the vertical coordinate which is often more flexible and accurate than a z-coordinate system. We show in the results section that column adaptivity provides accurate results and good grid compression ratios. Note that the vertical grid is remapped (if necessary) before adapting the horizontal grid.

The normalizations for the absolute tolerances $\varepsilon_{\mu} = \varepsilon ||\mu||_{\infty}$, $\varepsilon_{\Theta} = \varepsilon ||\Theta||_{\infty}$ and $\varepsilon_{u} = \varepsilon ||u||_{\infty}$ determining the grid adaptation may be estimated a priori by dimensional analysis and knowledge of the flow being simulated, or they

are determined dynamically by calculating the appropriate norms separately for each vertical layer at each time step.

The choice of normalization for the absolute thresholds ε_{μ} , ε_{Θ} and ε_{u} is a crucial and sensitive part of the algorithm since they guarantee the relative accuracy and efficiency of the method. In the shallow water case, we determined the tolerances for mass and velocity based on dimensional analysis of the tendencies in the inertia–gravity and quasi-geostrophic regimes. This sort of dimensional analysis is not feasible for the three-dimensional equations, so we have developed two strategies to ensure uniform control of the relative error in the tendencies of the prognostic variables in each vertical layer.

As we mentioned earlier, in the first approach, these tolerances are set by dimensional analysis using a knowledge of the appropriate scales for the test problem under consideration. This gives reasonable results for problems with fairly stationary evolution but can lead to less accurate results if the dynamics change significantly during the course of the simulation. It also generally means that the tolerances are set to the same value for all vertical levels.

In the second approach, the relevant norms are calculated dynamically at each time step and at each vertical level to ensure they are consistent with the actual state of the flow. This approach is more generally applicable and does not require the user to have any a priori knowledge of the solution.

In addition to determining how to calculate the normalization for the thresholds, we also need to decide which wavelets we are filtering. In the shallow water case, we always directly filtered the wavelets of the solution, which measure the interpolation errors of the variables themselves between two grid levels at a given time step. Although this does not directly control the tendency error, analysis of a linearized problem suggests that it should provide some control of the tendency error provided the solution is smooth.

Consider a set of n coupled linear ordinary differential equations:

$$\frac{\mathrm{d}\boldsymbol{u}}{\mathrm{d}t} = \mathbf{A}(\boldsymbol{u}), \, \boldsymbol{u}(0) = \boldsymbol{u}_0, \tag{17}$$

where **A** is an $(n \times n)$ constant coefficient matrix. Applying an Euler scheme gives the time step $u^{n+1} = u^n + \Delta t A u^n$. The error (wavelet coefficient) \tilde{u}^n satisfies the same equation, so we have

$$\widetilde{\boldsymbol{u}}^{n+1} = (\mathbf{I} + \Delta t \mathbf{A}) \widetilde{\boldsymbol{u}}^n,$$

$$\frac{||\widetilde{\boldsymbol{u}}^{n+1}||}{||\boldsymbol{u}^n||} \le (1 + \Delta t ||\mathbf{A}||) \frac{||\widetilde{\boldsymbol{u}}^n||}{||\boldsymbol{u}^n||},$$

where **I** is the identity matrix and we have used the triangle inequality. Now, if we assume that $||\tilde{u}^n||/||u^n|| \le \varepsilon$ by wavelet filtering of u^n , we have the following bound on the relative error:

$$\frac{||\widetilde{\boldsymbol{u}}^{n+1}||}{||\boldsymbol{u}^{n}||} \le \varepsilon (1 + \Delta t ||\mathbf{A}||).$$
(18)

If **A** is symmetric, then $||\mathbf{A}||_2 = \rho(\mathbf{A})$, where $\rho(\mathbf{A})$ is the spectral radius of **A** (largest magnitude eigenvalue of **A**, $|\lambda|_{\text{max}}$). Thus, we have that

$$\frac{||\widetilde{\boldsymbol{u}}^{n+1}||_2}{||\boldsymbol{u}^n||_2} \le \varepsilon (1 + \Delta t \,\rho(\mathbf{A})). \tag{19}$$

However, a necessary condition for stability of the Euler method is that

$$\Delta t \,\rho(\mathbf{A}) \le 2,\tag{20}$$

which gives

$$\frac{||\widetilde{\boldsymbol{u}}^{n+1}||_2}{||\boldsymbol{u}^n||_2} \le 3\varepsilon.$$

$$(21)$$

We have therefore shown that wavelet filtering of the solution itself provides control of the relative error of the solution over one time step for symmetric discretizations. In general, for non-symmetric \mathbf{A} , we have

$$\frac{||\widetilde{\boldsymbol{u}}^{n+1}||_2}{||\boldsymbol{u}^n||_2} \le \varepsilon (1 + \Delta t \sigma(\mathbf{A})), \tag{22}$$

where $\sigma(\mathbf{A})$ is the largest singular value of \mathbf{A} . However, there exists an $\epsilon > 0$ such that $||\mathbf{A}|| \le \rho(\mathbf{A}) + \epsilon$ for any matrix norm $||\cdot||$. Thus, we can expect similar relation to Eq. (21) to hold for more non-symmetric discretizations. Although we have only considered the linear constant coefficient case, the results suggest that dynamic wavelet filtering of the solution, together with dynamic calculation of the normalization of the thresholds, should control the relative error of the solution over one time step.

Now, consider filtering on the wavelets of the tendencies T themselves at the previous time step so that $||\tilde{T}^n||/||T^n|| \leq \varepsilon$, where \tilde{T} is now the error (wavelet coefficient) of the tendency. To first order in Δt , we have

$$\widetilde{\boldsymbol{T}}^{n+1} = \widetilde{\boldsymbol{T}}^n + \Delta t \left(\mathbf{J}_T(\widetilde{\boldsymbol{u}}^n, t^n) \widetilde{\boldsymbol{T}}^n + \frac{\partial \boldsymbol{T}}{\partial t}(\widetilde{\boldsymbol{u}}^n, t^n) \right),$$
(23)

where $\mathbf{J}_T(\tilde{\boldsymbol{u}}^n, t^n)$ is the Jacobian matrix of the tendency evaluated at the previous time step. In the case of an Euler method applied to a linear constant coefficient system of ordinary differential equations as we considered above, we have $T = A\boldsymbol{u}$ and the errors in the tendency \tilde{T} (wavelets of the tendency) satisfy

$$\widetilde{\boldsymbol{T}}^{n+1} = (\mathbf{I} + \mathbf{A})\widetilde{\boldsymbol{T}}^{n},$$

$$||\widetilde{\boldsymbol{T}}^{n+1}|| \le ||\widetilde{\boldsymbol{T}}^{n}|| + ||\mathbf{A}\widetilde{\boldsymbol{T}}^{n}||,$$

$$||\widetilde{\boldsymbol{T}}^{n+1}|| \le ||\widetilde{\boldsymbol{T}}^{n}|| + ||\mathbf{A}||||\widetilde{\boldsymbol{T}}^{n}||$$

where we have used the triangle and Schwartz inequalities. Normalizing by $||T^n||$ and recalling that wavelet filtering ensures that $||\tilde{T}^n||/||T^n|| \le \varepsilon$ gives the result

$$\frac{||\tilde{\boldsymbol{T}}^{n+1}||}{||\boldsymbol{T}^{n}||} \le (1+||\mathbf{A}||)\varepsilon.$$
(24)

www.geosci-model-dev.net/12/4901/2019/



Figure 4. Strong scaling of wavetrisk on the Compute Canada machine niagara for a simulation of the Held and Suarez (1994) general circulation experiment for a perfectly balanced (non-adaptive) run at a resolution of J = 8 (1/4°) and for a strongly unbalanced (dynamically adaptive) run at a maximum resolution of J = 9 (1/8°) resolution with trend-based error tolerance $\varepsilon = 0.08$. (a) Speedup compared with perfect (linear) scaling. The non-adaptive case has perfect linear scaling for more than eight cores, while the adaptive case has power law scaling of approximately 0.78. (b) Absolute strong scaling performance in milliseconds (ms) (wall-clock time per time step multiplied by the number of cores, i.e. CPU hours per time step, divided by the average number of active nodes over all vertical levels and all scales). \overline{N} is the average number of active nodes over all scales. Note that the absolute times shown in panel (b) are slower than equivalent times reported in Table 2 because we used a RK45ssp (with one additional trend evaluation) and code was compiled with gfortran rather than ifort for the scaling runs.

Again, if **A** is symmetric, we have $||\mathbf{A}||_2 = \rho(\mathbf{A})$, and using $\rho(\mathbf{A}) < 1/\Delta t$ for the Euler method we find

$$\frac{||\widetilde{\boldsymbol{T}}^{n+1}||}{||\boldsymbol{T}^{n}||} \le \left(1 + \frac{1}{\Delta t}\right)\varepsilon \le C\varepsilon.$$
(25)

Inequality (25) shows that filtering the wavelets of the tendencies \tilde{T}^n from the previous time step effectively controls the relative tendency error in the next time step up to a constant factor depending on the discretized system. Note that filtering the wavelets of the tendencies is slightly more expensive since it requires an additional evaluation of the tendencies (or an additional inverse wavelet transform).

In the following section, we apply wavetrisk to solve three standard test problems for hydrostatic dynamical cores. The emphasis is on evaluating the adaptivity in the threedimensional hydrostatic version, rather the basic accuracy of the method since the underlying discretization is the same as dynamico and we have already assessed the basic features of the error control in previous work on the shallow water equations. We directly compare the options of nonlinear filtering the wavelets of the solution and wavelet filtering the wavelets of the tendencies. Filtering the tendencies is more precise but more sensitive to the choice of threshold.

5 Validation test case results

5.1 Parallel and computational efficiency

The adaptive wavetrisk code has significant computational overhead compared to dynamico, which solves the same discretized equations. This overhead is required to deal with the local geometry of the grid (which is not precomputed), the multiscale grid structure and the parallel communication on the hybrid data structure. This overhead increases with the number of refinement levels and decreases for larger patch sizes. A lower bound on the overhead can be estimated by directly comparing wall-clock time for wavetrisk and dynamico since both codes are based on the same TRiSK discretization. The codes solved Dynamical Core Model Intercomparison Project (DCMIP) 2008 test case 8 (see Sect. 5.2) on the uniform grid J = 6 (1° degree) with 27 Lagrangian vertical levels and 8×8 patches for wavetrisk. The codes were run on 160 cores. This limited test suggests that wavetrisk is approximately 50% slower than dynamico per active grid point. The actual overhead depends on the number of refinement levels and the patch size (larger patch sizes are more computational efficient but limit the number of cores and increase memory overhead). Note that dynamico usually runs with massbased vertical coordinates, which adds an additional 15 % to its CPU time.

We have also directly estimated the overhead due to multiscale adaptivity for wavetrisk by comparing single-scale runs (with the largest possible patch sizes for the given resolution for best performance) and multiple-scale runs with minimum resolution J = 4 and maximum resolutions J =6, 7, 8, 9 on 40 cores with minimum scale J = 4. The tolerance was set to zero to give an upper bound on the cost of the multiscale runs. The code solved the Held and Suarez (1994) test case. We found that the overhead due to the multiscale adaptivity is about a factor of 3. We therefore conclude that wavetrisk is about 4 times slower per active grid point than dynamico, and therefore a grid compression factor greater than 4 is required for wavetrisk to be faster than dynamico. Remapping accounts for about 7%



Figure 5. Adaptive Held and Suarez (1994) general circulation experiment at t = 500 h with maximum resolution of $1/8^{\circ}$ used for the strong parallel scaling results shown in Fig. 4. Results are shown at the vertical level corresponding to 850 hPa. The rightmost figure is the adaptive grid with a grid compression ratio of about 4.5.

of the cost, so remapping every 10 time steps decreases the overhead difference to about a factor of 3.7. Using RK33ssp instead of RK4 further reduces the overhead to about a factor of 2.8. Note that, unlike dynamico, we have not made a serious effort to optimize the performance of wavetrisk and it is certainly possible to reduce this overhead significantly. Nevertheless, we will show that compression ratios of up to 1000 times are achievable using the adaptive code and it is certainly much faster than the equivalent non-adaptive code for high-resolution intermittent problems.

In order to estimate strong parallel scaling performance, we ran two Held and Suarez (1994) general circulation experiments. The first run was non-adaptive at horizontal resolution $1/4^{\circ}$ with 30 vertical levels for 10 time steps. This case is balanced and Fig. 4a shows that it has linear speedup from about 8 to at least 2560 cores (the lack of linear scaling for four or fewer cores is due to the intrinsic overhead of parallel computations). Although speedup is a commonly used measure of parallel scaling, a more sensitive measure is CPU time per time step divided by the number of vertical columns per core. Perfect scaling is then a constant, and its value gives an indication of absolute performance. Figure 4b shows that this measure is indeed approximately constant from 4 to 2560 cores.

The second trial was fully adaptive using trend filtering, with a minimum horizontal resolution of $1/2^{\circ}$ and three levels of refinement to $1/8^{\circ}$ and 18 vertical levels computed with a relative error tolerance $\varepsilon = 0.08$. This simulation was first run for 500 h to allow the climate dynamics to develop. Figure 5 shows the solution and adaptive grid at t = 500 h. The grid compression ratio is a relatively modest 4.5. The simulation was then restarted and run for another 4 h (about 300 time steps) to estimate strong parallel scaling and timing. The computational load was rebalanced on restart, but there was no further rebalancing. The load imbalance (ratio

of highest to average load) varied from 3 to 8 during this simulation. This run therefore represents a very unbalanced case. Nevertheless, Fig. 4a shows close to linear speedup (power law exponent of 0.78) to at least 640 cores. The more precise measure of strong scaling in Fig. 4b is further from a constant (with a maximum variation of 3 times), although there is not a definite trend with increasing numbers of cores. Since weak scaling performance is a better indication of parallel performance than the strong scaling, these results suggest weak scaling should be good to much larger numbers of cores. In fact, the shallow water code on which wavetrisk is based showed 70 % weak scaling efficiency for as few as 1300 computational elements per core (Aechtner et al., 2015). The current three-dimensional code has better parallel performance since the subdomains are distributed to the cores as complete vertical columns, which means each core has a larger load.

In summary, wavetrisk has a minimum overhead of about 50% compared with dynamico associated with the management of the adaptive grids and local geometry. This overhead increases with the number of refinement levels and decreases with patch size. However, the cost per active node is independent of the grid compression ratio. We also show that strong parallel performance is good for $O(10^3)$ cores. This suggests that wavetrisk requires a grid compression ratio of at least 2 to be more computationally efficient than an equivalent non-adaptive code. As we show in the following sections, grid compression ratios of 4 to 200 or more are achievable depending on the flow (see Fig. 11, for example). In general, higher grid compression ratios are achievable at higher resolutions and in more turbulent (intermittent) cases.

Table 2 summarizes absolute computational performance for the test cases considered in the following sections.

Table 2. Summary of actual computational performance for each of the test cases considered here. All runs were done on the Compute Canada machine niagara and the values shown are averages over the whole simulation. CPU is wall-clock time, \overline{N} is the average number of active nodes (over all vertical levels and all scales), total cost is wall-clock time per time step multiplied by the number of cores (i.e. CPU hours per active node per vertical level. (Note that total cost does not take into account the speedup due to parallelism or adaptivity: it measures CPU hours per active node.) The Rossby wave run is more expensive because it uses a smaller patch size (4 × 4 rather than 8 × 8) in order to run on 160 cores with $J_{\min} = 5$. Please see the discussion at the beginning of Sect. 5.1 for an explanation of the trade-offs involved in patch size versus number of domains.

Test case	Adaptivity	Cores	$\overline{\Delta t}$	$CPU d^{-1}$	$CPU/\Delta t$	$\overline{\mathcal{N}}\times 10^4$	Compression	Total cost
Rossby wave	tendency	160	170 s	203 s	0.40 s	4.85	4.43	0.0508 ms
Baroclinic instability	variables	40	240 s	310 s	0.40 s	4.10	5.22	0.0152 ms
Baroclinic instability	tendency	40	237 s	411 s	1.13 s	10.2	2.11	0.0173 ms
Held–Suarez (1°)	variables	40	287 s	81 s	0.27 s	2.78	1.93	0.0216 ms
Held–Suarez (1/4°)	variables	320	66.5 s	375 s	0.29 s	11.4	7.58	0.0456 ms



Figure 6. Results of the adaptive simulation of the mountain-induced Rossby wave case shown at 700 hPa at day 25.

5.2 DCMIP 2008 test case 5: mountain-induced Rossby wave train

The first validation we consider is the mountain-induced Rossby wave train used in DCMIP in 2008 (Jablonowski et al., 2008). This test case is relatively smooth and does not produce much small-scale structure. It is, however, a good test of the ability of the adaptive algorithm to track developing wave instabilities. It is similar to the case described in Tomita and Sato (2004) but with a hydrostatic surface pressure. The simulation starts from smooth, balanced isothermal initial conditions and a Rossby wave train instability is generated by an isolated Gaussian mountain over the course of the first 15 days.

The coarsest scale in the simulation is J = 5 (2°) and the finest scale is J = 7 (1/2°) with 27 vertical hybrid σ pressure levels. The vertical grid is remapped when the thickness of a vertical level has dropped to 30% of its initial value. There is no diffusion. The grid is adapted on the tendency wavelets. We run the simulation for a total of 30 days. The results are shown at day 25 on the sphere in Fig. 6. Figure 7 shows equidistant cylindrical rectangular latitude–longitude projections of the results of the adaptive simulation at 700 hPa at day 25. The adaptive data are first interpolated to a uniform J = 7 (1/2°) grid, then interpolated to the desired pressure level and finally projected onto the plane. The results are in good qualitative agreement with those shown in the DCMIP 2008 report (Jablonowski et al., 2008). There is some difference in the weaker structures, which is inevitable given that the adaptive simulation necessarily resolves the more intense structures more highly than the weaker ones.

The mountain-induced Rossby wave train is a good first validation of the wavetrisk. However, it is relatively smooth, it does not test the ability of the code to deal with intense instabilities, and it has no "physics" (e.g. cooling or Rayleigh drag). The following baroclinic instability test case is much more intense and generates small-scale vorticity filaments.



Figure 7. Latitude–longitude projections at 700 hPa of the adaptive simulation of the mountain-induced Rossby wave test case at day 25.

5.3 DCMIP 2012 test case 4: baroclinic instability of jet stream

Jablonowski and Williamson (2006) proposed a deterministic test case for dry dynamical cores of atmospheric general circulation models that simulates the evolution of a baroclinic wave in the Northern Hemisphere. Perturbation of an analytic steady state solution triggers a baroclinic instability that generates a series of vortices. These vortices grow and interact, eventually producing a two-dimensional turbulencelike vorticity field of intense filaments and vortex cores. The rapid development of the vortical instability and its subsequent evolution is a challenging case for an adaptive method. This baroclinic instability was test case 4 of DCMIP 2012.

With a sufficiently low relative tolerance (e.g. $\varepsilon = 0.03$ when adapting on the solution), wavetrisk successfully captures the explosive cyclogenesis around day 8 and subsequent breaking of the wave train around day 12 to produce a series of intense filamentary vortices (see Sect. 10). This calculation uses quadratic piecewise parabolic remapping at each time step Engwirda and Kelley (2016, https://github. com/dengwirda/PPR, last access: 25 November 2019).

Figure 8 shows the surface pressure and temperature at day 9, which agree reasonably well with the equivalent results in Fig. 6 from Jablonowski and Williamson (2006). Note that we do not expect exact agreement since the adaptive simulation does not resolve all regions uniformly. Beyond day 12, the vortices interact to produce turbulence-like flow in both hemispheres. These results therefore validate the ability of the model to capture suddenly developing instabilities and track their evolution to a turbulence-like state.

Next, we compare the characteristics of the solutionfiltered and trend-filtered variants of wavetrisk. In both cases, the maximum scale is $J = 7 (1/2^{\circ})$ and we use 27 vertical hybrid σ pressure levels as specified in Jablonowski and



Figure 8. Latitude–longitude projections of surface pressure and temperature at vertical level 4 (about 867 hPa) of the adaptive simulation of the baroclinic instability test case at day 9.

Williamson (2006). There is no explicit diffusion added to stabilize the dynamics or to damp out grid-scale oscillations and both simulations using RK45ssp time integration (Spiteri and Ruuth, 2002). It is important to note that, although this is a deterministic test case when simulated non-adaptively, adaptivity necessarily neglects some less dynamically important structures and so the details of flows with different tolerances eventually diverge once they become turbulent (after roughly 12 days). This is, however, an optimal test case for adaptive methods, since at early times, only a small portion of the flow is active, which allows high compression ratios. In fact, until the instability develops, a very coarse grid and large time step are sufficient.

Figure 9 compares the grid compression ratios for the baroclinic instability when adapting on the solution and on the trend. The tolerances were set to achieve similar com-



Figure 9. Comparison of the number of active nodes \mathcal{N} for the baroclinic instability when adapting on the trend or adapting on the solution. The relative error threshold was set to $\varepsilon = 0.04$ when adapting on the solution and $\varepsilon = 2.6$ when adapting on the trend. In both cases, the active scales are j = 5, 6, 7 and there is no additional diffusion. The tolerances were set to achieve similar compression ratios at 9 days. The total number of available grid points is 8.6×10^5 , and so the minimum grid compression ratio is 3.6 when adapting on the solution and 1.5 when adapting on the trend. The average compression ratio is 5.2 when adapting on the solution and 2.1 when adapting on the trend.

pression ratios at day 9 and were set based on dimensional analysis (i.e. they are fixed in time). Both options use similar numbers of grid points until about day 12 when the flow becomes more turbulent. Once the flow is turbulent, adapting on the trend uses significantly more grid points. Although the number of grid points used by both options is similar until day 12, the methods distribute the same number of grid points quite differently.

Figure 10 compares the vorticity fields and active grids at days 9 and 20. At day 9, adapting on the solution distributes the available grid points to track all the developing vortices. In contrast, adapting on the trend concentrates all grid points on the two strongest vortices. In addition, the peak vorticity when adapting on the solution, $5.2 \times 10^{-4} \text{ s}^{-1}$, is higher than when adapting on the trend, $1.9 \times 10^{-4} \text{ s}^{-1}$. Once the flow is turbulent, at day 20, adapting on the trend uses about 2.8 times more grid points than adapting on the solution.

Adapting on the solution appears to be more efficient since adapting on the trend is sensitive to grid-scale noise in the trend (the solution is smoother than the trend). Adapting on the trend uses fewer overall grid points when the tolerances are rescaled dynamically (i.e. when the trend norms are recomputed at each time step), but the resulting adapted grid is quite sensitive to local fluctuations in the trend. This effect is much less pronounced when diffusion is added. Based on this and other examples we have investigated, adapting on the solution appears to be preferable when there is little or no diffusion added to damp out grid-scale noise.

It is interesting to investigate how the grid compression ratios change as we increase the maximum allowed resolution. The main advantage of adaptive methods is for intermittent problems that require very high local resolution not attainable using static grid methods. This requires that the grid compression ratio increases significantly with the maximum allowed resolution. Figure 11 compares grid compression ratios for four different maximum resolutions ranging from $J = 7 (1/2^{\circ})$ to $J = 10 (1/16^{\circ})$ from day 8 to day 9 at the onset of the instability. The simulations have no diffusion and are adapted on the trends. Unsurprisingly, the compression is very high at the onset of the instability at day 8, over 1300 at J = 10. At day 9, once the initial vortices have developed, the compression ranges from about 20 at J = 7 to about 200 at J = 10. Note that we have found that we can also use a higher relative error threshold at higher resolutions which further increases the compression.

This scaling test suggests that the adaptive method should be especially advantageous at high maximum resolutions for vorticity-dominated flows. Although in this flow the instability is highly localized, Aechtner et al. (2015) found compression ratios of about 20 for a homogeneous isotropic shallow water turbulence case on the sphere with scales J =5, ..., 10.

5.4 Held and Suarez general circulation experiment

Held and Suarez (1994) is a classic test case for dynamical cores of atmospheric general circulation models. It uses simplified "physics" (i.e. radiation and friction/drag models) that nevertheless produce realistic general circulation over relatively short timescales of $O(10^2)$ days. For example, Liu and Schneider (2010) have used simple Held-Suarez-type physics to realistically model the general atmospheric circulation of Saturn. This model uses height-dependent Rayleigh damping to represent boundary-layer friction and a heightand latitude-dependent Newton cooling relaxation of potential temperature to a prescribed radiative-convective equilibrium. The temperature relaxation includes parameters accounting for cooling at the surface and top of the atmosphere as well as a tropopause. The Held-Suarez general circulation experiment adds a qualitatively new aspect to the Rossby wave and baroclinic instability tests we considered above: it includes physics source terms for the temperature and momentum equations. None of the previous simulations have included these sorts of physics terms, either in three dimensions or in two dimensions on the sphere or on the plane. Nevertheless, we expect the grid adaptation to track the effect of the source terms either directly through their effect on the trend or indirectly through their effect on the prognostic variables.

We compare low- and high-resolution runs since previous work has suggested that the choice of coarsest can affect the result. We also want to explore how grid compression changes with resolution (as in the baroclinic test case). The



Figure 10. Comparison at days 9 and 20 of the grid compression ratios for the baroclinic instability when adapting on the trend or adapting on the solution. The relative error threshold was set to $\varepsilon = 0.04$ (fixed) when adapting on the solution and $\varepsilon = 2.6$ when adapting on the trend. In both cases, the resolution is j = 5, 6, 7 and there is no additional diffusion. The tolerances were set to achieve similar compression ratios at 9 days. The vorticity field is shown at hybrid vertical level 4 (about 870 hPa).

low-resolution run has coarsest scale $J_{\min} = 4$ and finest resolution J = 6 (about 120 km or 1°) with tolerance $\varepsilon = 0.04$. The high-resolution run has coarsest scale $J_{\min} = 6$ and finest resolution J = 8 (about 30 km or 1/4°) with tolerance $\varepsilon = 0.02$. The low-resolution case is run on 40 cores and the high-resolution case is run on 320 cores. As in Wan et al. (2013), both cases start from the Jablonowski and Williamson (2006) zonally symmetric initial condition with random noise of magnitude 1 m s⁻¹ added to the zonal wind. In both cases, the simulations are first run non-adaptively for 200 days at the coarsest resolution. The time step is adaptive with that of the CFL criterion. The maximum possible grid compression ratio for both cases is 21.

We use 19 vertical σ -pressure levels concentrated at the top and bottom of the atmosphere. The vertical grid is

remapped using a piecewise parabolic method with WENO limiting every 10 time steps.

Small-scale noise is damped with p = 2 hyperdiffusion with diffusion constant $K_{\phi} = 3.48 \times 10^{15} \text{ m}^4 \text{ s}^{-1}$, $K_{\delta} = 3.48 \times 10^{16} \text{ m}^4 \text{ s}^{-1}$ and $K_{\omega} = 2.17 \times 10^{14} \text{ m}^4 \text{ s}^{-1}$ for the low-resolution run and $K_{\phi} = 2.33 \times 10^{14} \text{ m}^4 \text{ s}^{-1}$, $K_{\delta} = 5.98 \times 10^{14} \text{ m}^4 \text{ s}^{-1}$ and $K_{\omega} = 1.46 \times 10^{13} \text{ m}^4 \text{ s}^{-1}$. The diffusion is applied each time step in the main trend routine. The source terms for the potential temperature and the velocity representing cooling and Rayleigh damping are implemented as a separate Euler step.

Mean and variance statistics are calculated using a parallel version of Welford's inline algorithm (Chan et al., 1983) by first interpolating the solution to the finest grid from checkpoints saved every 24 h. The second-order statistics are essentially converged after 200 days (the first-order statistics converge much more quickly).



Figure 11. Grid compression ratios at the beginning of the baroclinic instability for maximum resolutions J = 7 (1/2°), J = 8(1/4°), J = 9 (1/8°) and J = 10 (1/16°). In all cases, the coarsest resolution is J = 5 (2°). The grids are adapted on the trend and there is no diffusion. Note that the J = 10 uses a higher relative error threshold $\varepsilon = 4$ than the others, which use $\varepsilon = 2$. Compression ratios increase significantly with increasing resolution, reaching as high as 200 times at day 9 for the maximum resolution case. Even at J = 7, the code achieves a compression ratio of about 20 at day 9.

A typical low-resolution result is shown in Fig. 12 (top). The average grid compression ratio at this low resolution is only 1.9 ± 0.1 with $\varepsilon = 0.04$. The adaptive algorithm is able to track the development and evolution of the fine-scale filamentary vortex structures over long times. Note that since the adapted grid is the union of adapted grids over all vertical levels, the adapted grid does not necessarily correspond exactly to the structures at the vertical level 11 (about 250 hPa) at the level of the upper atmosphere jets shown in the figure.

Figure 13 shows standard first- and second-order statistics averaged zonally and over time for the low-resolution run. These statistics are qualitatively similar to those of Dubos et al. (2015), Wan et al. (2013) and Lin (2004). (Lin, 2004 is the only one that uses Lagrangian vertical coordinates.) The main quantitative difference is in the slightly lower magnitude of the eddy kinetic energy. This difference is due partly to the use of Lagrangian vertical coordinates, where the remapping introduces additional dissipation, and partly to the additional dissipation generated by the adaptivity at these relatively low resolutions. Lin (2004) did not show the eddy kinetic energy, but he reported a maximum variance of zonal wind of about $300 \text{ m}^2 \text{ s}^{-2}$, slightly larger than the $284 \text{ m}^2 \text{ s}^{-2}$ we find here (not shown), and similar to the 301 m² s⁻² we find for the high-resolution case discussed below. Note that the original Held and Suarez (1994) paper did not present results for eddy kinetic energy.

The choice of remapping has a significant influence on the eddy kinetic energy. For example, we found that changing from a piecewise linear remapping to a piecewise parabolic remapping increases the maximum eddy kinetic energy by $53 \text{ m}^2 \text{ s}^{-2}$ for a non-adaptive simulation with resolution J =

5 (240 km). Similarly, Lin (2004) found that including a monotonicity constraint in the remapping lowered the maximum variance of the zonal velocity by about $20 \text{ m}^2 \text{ s}^{-2}$. We would also like to emphasize that piecewise constant remapping gives qualitatively incorrect results (e.g. zonal jets are too high), presumably because it is too dissipative and is not a good choice for this test case.

Larger tolerances ε effectively increase dissipation, which decreases maximum eddy kinetic energy, while choosing a smaller tolerance leads to essentially no compression in the low-resolution run. (The other statistics are much less sensitive to ε .) Note that a fixed (non-adaptive) resolution $J_{\min} = 4$ (4°) gives a maximum eddy kinetic energy of only $280 \text{ m}^2 \text{ s}^{-2}$, much less than the published values of over $400 \text{ m}^2 \text{ s}^{-2}$ at higher resolutions of 2 and 1°. This suggests that the coarsest grid may be the main factor behind the lower eddy kinetic energy we observe here at higher compression ratios.

In contrast, Fig. 12 (bottom) shows typical results from the high-resolution simulation at $\varepsilon = 0.02$. The grid compression is clearly much higher than in the low-resolution run, and the fine scales are limited to a small neighbourhood of the high-intensity vorticity filaments. The average grid compression ratio for the high-resolution simulation is 7.6 ± 0.5 , exactly 4 times as large as in the low-resolution case. Both the compression and the fluctuations in the compression are much higher than in the low-resolution run. These results suggest that the adaptive method is useful primarily at higher resolutions and for more turbulent flows. This confirms our earlier observations for two-dimensional shallow water turbulence on the sphere (see Figs. 17 and 18 of Aechtner et al., 2015). The advantages of the adaptive method should be even greater for maximum resolutions higher than $J = 8 (1/4^{\circ})$.

Figure 14 shows standard first- and second-order statistics averaged zonally and over time for the high-resolution run. The results are qualitatively very similar to the lowresolution run, with the main differences being more intense eddy momentum and eddy kinetic energy. The negative highaltitude zonal jet is also a bit stronger.

The results presented here suggest that the eddy kinetic energy is quite sensitive to the coarsest resolution and relative error tolerance chosen and that, not surprisingly, the adaptive method does not provide significant benefits for lowresolution simulations. However, dynamic adaptivity makes much-higher-resolution runs feasible at a significantly reduced computational cost (both in terms of memory and CPU time). This advantage should increase for larger maximum resolutions unattainable using non-adaptive codes. The ability to restart a simulation adaptively at a higher resolution (e.g. add three levels of refinement) allows events of interest to be explored at very high local resolutions, even if it is only for relatively short times.



Figure 12. Typical results for the low-resolution (top) and high-resolution (bottom) Held and Suarez general circulation test case at 250 hPa. The grid is adapted on the solution with relative error tolerance $\varepsilon = 0.04$ in the low-resolution case and $\varepsilon = 0.02$ in the high-resolution case. The grid compression ratio is 2.0 for the low-resolution case and 7.5 for the high-resolution case.

6 Conclusions

This paper introduces wavetrisk: a new adaptive dynamical core. wavetrisk uses the hydrostatic dynamico discretization of Dubos et al. (2015) with a Lagrangian vertical coordinate. Second-generation discrete wavelet transforms provide control of the relative error of the solution in each vertical layer at each time step. The adaptive grid is uniform vertically and is composed of columns of varying horizontal sizes. In addition to the horizontal adaptivity, the vertical coordinates are remapped onto a hybrid σ -pressure coordinates by optimizing the target grid at each remap. *h* adaptivity may also be possible in the future by deactivating some vertical layers (the so-called "dormant layers" used in ocean modelling).

The code is parallelized via domain decomposition using mpi and the data are stored in a hybrid quad tree-patch data structure. The computational load is rebalanced at each checkpoint save. We demonstrate excellent strong parallel scaling up to at least 2560 cores in the perfectly balanced case and close to linear scaling (exponent 0.78) up to at least 640 cores in an unbalanced case. wavetrisk is 3–4 times slower per active grid point than the non-adaptive dynamico code, which suggests that we require a grid compression ratio of more than 4 for adaptivity to provide an advantage in CPU time.

The grid may be adapted based on the wavelet coefficients (interpolation errors) of either the prognostic variables themselves (mass density, mass-weighted potential temperature and velocity) or of their associated tendencies. The desired relative error tolerance is multiplied by the appropriate norm (of the variables or tendencies) computed at each time step to control the relative error. Note that the adaptivity is necessarily dissipative compared to an equivalent simulation with uniform resolution on the finest grid since some energy is lost when computational elements are coarsened. However, by construction, this dissipation is controlled and may be reduced by decreasing the tolerance or increasing the maximum allowed resolution. In previous work (Aechtner et al., 2015), we explored how the effective Reynolds number of a



Figure 13. Time–zonal statistics for the low-resolution (Held and Suarez, 1994) test case with scales J = 4, 5, 6 (maximum resolution of 1°). The grid is adapted on the solution error with tolerance $\varepsilon = 0.04$. Statistics are averaged over 400 days after day 200 by interpolating saved data to the finest grid.



Figure 14. Time–zonal statistics for the high-resolution (Held and Suarez, 1994) test case with scales J = 6, 7, 8 (maximum resolution $1/4^{\circ}$). The grid is adapted on the solution error with tolerance $\varepsilon = 0.02$. Statistics are averaged over 200 days after day 700 by interpolating saved data to the finest grid.

turbulent flow depends on the choice of error tolerance parameter.

We have validated the code on three standard benchmarks: a mountain-induced Rossby wave train, baroclinic instability of a jet stream and the Held and Suarez general circulation experiment. These tests show that wavetrisk correctly captures the dynamics, including rapidly developing instabilities, with only a small portion of the total grid points available on a similar non-adaptive grid. The grid compression ratio can reach over 200 in ideal cases (e.g. the start of the baroclinic instability with five levels of refinement) and is advantageous at sufficiently high resolutions even in more homogeneous flows like Held and Suarez.

Because adaptive climate simulation is a new field, we have deliberately included many options in our adaptive algorithm. wavetrisk can adapt on errors in the solution or on errors in the trend; it can run with no diffusion, Laplacian diffusion or hyperdiffusion. The vertical grid can be regridded (using a large selection of remapping schemes) each time step or only when a level becomes too narrow. And, of course, we can choose different relative tolerances ε and maximum and minimum grid resolutions. In many cases, the code is stable without diffusion and with grids adapted either on the solution or the trend. However, our test cases suggest that adapting on the solution (rather than the trend) generally gives more accurate and faster solutions for a given number of grid points. Including a small amount of diffusion stabilizes the code and reduces the number of active grid points by reducing grid-level noise.

It is clear that the main application of wavetrisk is for simulations at maximum resolutions unattainable by non-adaptive dynamical cores. In future work, we will use wavetrisk to simulate simple Held–Suarez-type climates at much higher resolutions and for longer times and investigate the behaviour of more sophisticated physics parameterizations in adaptive simulations. We are also developing an ocean variant of wavetrisk that will improve the ALE formulation of the vertical coordinate and use penalization for bathymetry and coastlines. This work builds on the shallow water ocean model we presented in Kevlahan et al. (2015).

Code availability. WAVETRISK-1.0 is published under the Creative Commons 4.0 license at https://doi.org/10.5281/zenodo.3459710 (Kevlahan et al., 2019). The latest version of the code is WAVETRISK-1.1, which includes some bug fixes and two incompressible cases.

Author contributions. Both NKRK and TD have contributed to the research and paper preparation.

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