1. Introduction

In many aspects mesoscale oceanic eddies, operating on the length scales of \(O(1–100)\) km are analogous to the cyclones and anticyclones that constitute the atmospheric weather phenomenon. The problem of resolving these eddies in a dynamically consistent way is very important for ocean modelling and, therefore, for global climate predictions. For achieving high Reynolds number \((Re)\) simulations, which are required for accurate modelling of the ocean, the models have to account for all important scales of motion.

Modern ocean models enter a new phase in which eddies will be, at least, permitted in the numerical simulation. For such models advection scheme is a very important component. A crucial element of numerical advection scheme is its ability to propagate finite-amplitude and -phase disturbances on a discrete grid either without generating spurious short-wave oscillations, because of not preserving the correct dispersion relation i.e., dispersion error, or any considerable damping of the amplitude i.e., dissipation error (e.g., Kravchenko and Moin, 1997; Pope, 2000). Note, that the generation of short-wave oscillations is particularly detrimental in case an inverse energy cascade takes place, and the small scales affect large scales (e.g., Tabeling, 2002; Vallis, 2006). In this paper, the effect of spurious small-scale dispersion and dissipation on important large-scale properties of the solution are captured by comparing numerical predictions obtained with two different advection methods implemented within the same quasigeostrophic ocean modelling code (Berloff et al., 2007), which solves for the classical double-gyre problem (Holland, 1978). The original model implements standard eddy viscosity for parameterising effects of the unresolved scales of turbulent diffusion and conservative second-order Arakawa scheme for advection (Arakawa, 1966). In this paper, we consider a few versions of the original code based on different advection methods. Comparisons with the converged fine-grid solutions are made to investigate effects of numerical advection schemes on the coarse-grid solutions.

Solving ‘convection-dominated’ problems is a long-standing challenge for computational fluid dynamics (e.g., Rozhdestvensky and Yanenko, 1978; Roache, 1982; Hirsch, 2007). One of the difficulties is that the conventional second-order finite-difference schemes have large dispersion errors, which generate spurious ripples in the solutions. To counterbalance this effect, a numerical dissipation, such as the classical von Neumann and Richtmyer (1950) artificial viscosity for compression-type pressure waves or such as the eddy viscosity in ocean circulation models, is added to the governing equations. However, a common negative side of this approach is the associated spurious dissipation that smears the large eddies along with the spurious ripples.

There are three general approaches for solving ‘convection-dominated’ problems: Eulerian, Lagrangian, and mixed Eulerian–Lagrangian. For the Eulerian methods, significant presence of both numerical dissipation and dispersion is common drawback. It is partially overcome in the Lagrangian and Eulerian–Lagrangian methods, which describe flow advection by following fluid particle coordinates, rather than by considering fixed coordinates on the...
Eulerian grid (e.g., Dritschel et al., 1999; Mohendarjoueh and Dritschel, 2004). A remarkable property of the Lagrangian methods is that they are exact for the linear-advection problem with a uniform velocity field, and in this case their accuracy is limited only by the accuracy of solving the corresponding Ordinary Differential Equations (ODEs), rather than by the accuracy of solving the full Partial Differential Equations (PDEs), as in the Eulerian case. The Lagrangian methods can be very efficient for simulations with multiple contact discontinuities and shocks, e.g., as those of multiphase flows (e.g., Margolin and Shashkov, 2004). However, for the problems dominated by chaotic folding and stretching of the material lines, the Lagrangian methods have to be complemented with ad hoc ‘repair’ (or ‘contour surgery’) procedures. They allow to remove redundant Lagrangian particles that aggregate in some locations and to add new particles to where they are needed. The ‘repair’ procedure can be viewed as a special kind of numerical dissipation that drains energy from the underestimated scales. The numerical effect of the ‘repair’ procedure on the numerical dissipation and dispersion error always limits Re for the resolved scales. Also, specifying standard physical boundary conditions (e.g., no-slip, free-slip, etc.) is problematic for the Lagrangian methods.

The implementation of Eulerian–Lagrangian methods, including semi-Lagrangian and vortex-in-a-cell methods is less intricate, since they employ an Eulerian interpolation/projection step after a Lagrangian step. However, they are no longer exact for the linear advection, and they are prone to the same dissipation and dispersion error problems as the Eulerian methods. For example, the mixed Eulerian–Lagrangian methods use artificial numerical dissipation for avoiding the ‘repair’ procedure and for ensuring a ‘non-oscillatory’ solution (e.g., Margolin, 1997). Finally, the Lagrangian-to-Eulerian grid interpolation is also a source of the numerical dissipation and dispersion errors.

Many Eulerian methods for ‘convection-dominated’ flows are based on emphasizing a particular property of the governing equations. Using such property as the basis for a “low-order” (first- or second-order) scheme, typically, further upgrades (e.g., high-order, non-oscillatory sequels, etc.) of the original scheme are developed. One such example is the family of optimised low-dissipation and low-dissipation finite-difference schemes, including implicit compact finite-difference schemes based on Pade-type approximations (Lele, 1992) and explicit schemes (Tam and Webb, 1993; Bogey and Bailly, 2004) that are popular in turbulence modelling. In particular, the family of explicit schemes was developed to overcome typical problems of the spectral and pseudo-spectral methods, such as handling non-periodic boundary conditions and suitability for parallel computations. The optimised explicit finite-difference schemes, typically, employ non-conservative forms of the governing equations, and use large computational stencils for more accurate approximation of the linear-wave dispersion relation in the physical domain. In order to cope with the large unresolved gradients emerging in nonlinear flows (e.g., shock waves) the high-order low-dissipative low-dispersive schemes use special filtering procedures (e.g., Robert, 1966; Asselin, 1972; Zhou and Wei, 2003). Such filtering is analogous to some form of artificial viscosity added to the scheme and is required, essentially, to reduce the second order in the vicinity of the unresolved solution gradients, in accordance with the Godunov Theorem (Godunov, 1959), that states that any monotone finite-difference scheme is first-order accurate.

Another example consists of the so-called “high-resolution schemes” (after A. Harten) that are based on Flux Corrected Transport (FCT), Total Variation Diminishing (TVD) solution ideas for quasi-linear hyperbolic conservation laws (e.g., Boris et al., 1975; Harten et al., 1987; LeVeque, 2002). In this approach a conservative approximation of the governing equations is used, that has several useful properties such as compact computational stencil, low CPU cost, and, often, the solution boundness, but have large dissipation and/or dispersion errors. Then such schemes are upgraded to a higher order, e.g., within the framework of ‘variabile-extrapolation’ and ‘flux-extrapolation’ techniques (e.g., van Leer, 1979; Woodward and Colella, 1984; Drikakis, 2003), by extending the spatial stencil. The upgrade is needed to enhance linear wave propagation properties of the solution away from the flow discontinuities. In the vicinity of high-gradients of the solution, either nonlinear flux limiter (FCT/TVD) functions are used (Boris et al., 1975; Harten et al., 1987; Pietrzak, 1998), or solution-adaptive stencils are used, as in WENO-type schemes (Shu and Osher, 1988). A significant improvement of the linear wave propagation properties can also be obtained by introducing additional degrees of freedom inside each computational cell, in the framework of Discontinuous Galerkin methods (Cockburn and Shu, 2001), which are less sensitive to spatial grid non-uniformity, relative to the high-resolution finite-difference schemes.

Standard second- and third-order non-oscillatory schemes are typically too dissipative for the problems in which the solution-front-sharpening mechanism is either absent or too weak to counterbalance the effect of the numerical dissipation. The higher cost of implementation and the extra CPU cost of a high-order scheme is accepted by the computational community for some situations, when the conventional methods require prohibitively fine grids. One may argue that high-order discretisation in space is the only way to improve the solution, but we take a complimentary view. We argue that in a number of situations it is more efficient to improve the properties of the underlying discretisation by addressing additional properties of the governing (hyperbolic) equations, within the class of second-order schemes. We argue that a hierarchy of high-order schemes is better to be developed from a “low-order” method that is as accurate and efficient as possible. Then the order of the method can be systematically increased, consistently in space and time, and the low-dispersive and low-dissipative properties of the original scheme can be improved. Finally, because of their robustness, the second-order methods are still widely used as a “working horse” in many hydrodynamics codes (e.g., for non-uniform grids, treatment of the shocks, ease with boundary conditions, etc.), and their improvement is important.

2. CABARET method

2.1. Linear advection scheme

To illustrate our ideas on a simple example, a scalar conservation law
\[ \partial_t u + \partial_x f(u) = 0 \] (1)

is considered on a finite-difference grid which is non-uniform in space \( x_{i+1} - x_i = h_{i+1/2} \) and time \( t^{n+1}_t - t^n_t = \tau^{n+1/2}_t \).

Fig. 1 shows the associated data structure on the non-uniform grid in space and time: the solid circles (labelled with letters) show the locations of conserved variable \( u \) with fractional \( i \)-indices and the open circles (labelled with numbers) show the locations of the flux variable \( f \) with whole \( i \)-indices. Our scheme assumes that all variables are known at the nth time level, and, then, it takes a half time step using forward-time central approximation,

\[ u^c_n - u^e_n + \frac{f^c_n - f^e_n}{h_{i+1/2}} = 0. \] (2)

It is continued with another half time step by using the backward-time central-space approximation,

\[ u^e_n - u^c_n + \frac{f^e_n - f^c_n}{h_{i+1/2}} = 0, \] (3)
where the fluxes $f_1$, $f_2$ are still to be determined. Note, that by adding these two equations, one obtains a conventional flux integral around the cell 1254, with a trapezoidal rule evaluation of the fluxes. By symmetry, the scheme will be second-order accurate for a sufficiently accurate evaluation of $f_1$ and $f_2$. We choose to evaluate these by a simple upwind extrapolation carried out in an upwind manner. Let’s assume for the moment that the sign of the wave speed, $\partial_u f(u)$, is positive everywhere. Then, we determine $f_1 = f(u_1)$ by assuming that
\[
u_1 = 2 u_c - u_s. \tag{4}\]

With this choice, the entire scheme (2)–(4) are time-reversible. It is also second order accurate, regardless of the grid non-uniformity in space and time, and it is non-dissipative.

The CABARET scheme (2)–(4) is an explicit single-temporal-stage method (note that the upwind extrapolation step for updating the fluxes is applied once per time step). The scheme is conservative and stable under the Courant (CFL) condition: $0 < \frac{cT}{h} \leq 1$. Due to its compact computational stencil, it remains second order accurate even on non-uniform spatial and temporal grids.

2.2. Treatment of unresolved short-wave scales in the solution

Regardless of how good the numerical scheme is, it is unlikely to correctly resolve all flow scales which can emerge in high-Re flow regimes. A feasible approach is to modify the original non-dissipative numerical method so that the small scales are harmlessly removed from the solution, without spurious backscatter from small to large scales (e.g., Pope, 2000; Grinstein et al., 2007). The discrete grid resolution limit is consistent with the Godunov Theorem (Godunov, 1959) which states that any monotone scheme is first order accurate. This implies for a numerical smoothing procedure to be used in the vicinity of unresolved solution gradients. Note that the resolution problem of Eulerian schemes can be directly related to their dispersion errors at high wavenumbers. The better numerical advection scheme is, the closer to the computational grid size is the properly resolved wave, and the better tailored is the use of the numerical smoothing.

For making the CABARET solution non-oscillatory, a simple tunable-parameter-free flux limiter is introduced through a non-linear correction procedure for the flux variables:
\[
u_1 = 2 u_c - u_s; \tag{4a}\]

if $(u_1 > \max(u_4, u_s, u_5)$),

if $(u_1 < \min(u_4, u_s, u_5))$, then
\[\nu_1 = \max(u_4, u_s, u_5); \quad \nu_1 = \min(u_4, u_s, u_5).\]

The above nonlinear correction procedure is based on the maximum principle (e.g., Boris et al., 1975; Harten et al., 1987) for characteristic wave that arrives at point 1 from the solution domain dependency $4 \rightarrow 5$, and that is approximated using the 3-point stencil $(u_4, u_s, u_5)$ within one cell in space. In comparison to the standard FCT/TVD schemes, the nonlinear correction algorithm is directly based on enforcing the maximum principle on flux variables, rather than limiting the slopes of conservation variables.

Despite the fact that the correction method (4a) does not explicitly enforce monotonicity on the conservation variables, the nonlinear flux correction allows the CABARET scheme to suppress non-physical oscillations in the solution. Numerical tests and theoretical investigations (e.g., Ostapenko, 2009) have confirmed that the solution of the CABARET method with flux correction (4a) remains strictly free both from spurious oscillations and “stair-casing” artifacts of the standard TVD schemes (e.g., Shchepetkin and McWilliams, 1998) for all Courant numbers $CFL \leq 0.5$. Fig. 2 shows the comparison of the CABARET solution for the linear advection of discontinuous and continuous initial data with the second-order van Leer TVD scheme based on MinMod and Superbee limiters (e.g., Hirsch, 2007).

It should be mentioned that there are more sophisticated ways of introducing conservative correction in the CABARET scheme, than the simple limiter (4a). For example, there are algorithms that enforce monotonicity simultaneously to the flux and conservation variables (e.g., Goloviznin et al., 2003). For simple cases, e.g., for linear advection with constant velocity, they produce more accurate results than the simple algorithms (2), (3) and (4a). A particular example is the so-called Digital Transport Algorithm (e.g., Karabasov and Goloviznin, 2004), which leads to the exact solution of 1-D and 2-D advection problems for any piece-wise constant initial data at $0 < CFL < 1$ ($0 < CFL < 0.5$ for 2-D problems) on Eulerian grids. Despite our interest in such generalizations of the CABARET method (2), (3) and (4a) they will not be discussed any further in the current publication, since their extension to the ocean modeling is in progress.

2.3. Comparison with other Leapfrog schemes

Schemes similar to (2)–(4) exist in the literature; they are the Upwind Leapfrog (UL) schemes first discussed by Iserles (1986) for linear advection, and later developed by Roe (1998), Tran and Scheurer (2002) and Kim (2004) for multidimensional wave propagation. However, these schemes were neither conservative, nor based on a compact one-space-cell one-time-step computational stencil. Also, they were not equipped with the limiters enabling them to overcome non-physical oscillations in the solution; hence, they were not robust enough for practical applications. In parallel to these efforts, Goloviznin and Samarskii (1998a,b) proposed an extension of the UL scheme to conservative methods. Later, Goloviznin and co-workers developed CABARET versions with the limiters (Goloviznin and Karabasov, 1998) and extended the method to multidimensional wave dispersion in fractured porous medium (Goloviznin et al., 2007) and gas dynamics (Goloviznin, 2005; Karabasov and Goloviznin, 2007).

To see that the present method is reduced to the UL for the linear advection ($f = c \cdot u$, $c = \text{const.}$) on a uniform grid in space and time, note that
\[
u_c - u_c = \nu(u_c - u_s) \quad \text{and} \quad u_s - u_c = \nu(u_s - u_c), \tag{5}\]

where $v = \frac{cT}{h}$ is the CFL number. Hence,
\[
u_c - u_c = \nu(u_c - u_s) = \frac{1}{2}(u_1 + u_5 - u_s - u_4) = \frac{1}{2}((u_1 - u_s) + (u_5 - u_4)), \tag{6}\]

which is the three-time-level UL method.
the Adams-type schemes, the CABARET scheme (2)–(4) is non-dis-

Note, that in comparison to the CABARET stencil (1-A-2-C-4-E-5) the UL method (1-4-5-8) uses a double-interval computational stencil in time, whereas the conventional Central Leapfrog (CL) method

\[ v(u_0 - u_4) = (u_2 - u_6) \]

uses computational stencil which is twice as wide as that of the UL scheme.

In addition to removing the defects of the UL, the CABARET scheme retains all of the UL good properties, such as ease of implementation near boundaries, zero numerical dissipation and very small dispersion error.

An important distinction between the implementations of the CABARET and UL schemes is that the UL scheme is three-time-level and needs to be properly initialized, otherwise a spurious mode can be excited. CABARET provides unambiguous prediction at level \((n + 1)\) from data at level \((n)\). However, the same spurious mode can be excited if at the initial time step the flux data are not consistent with the conservation variables.

An important feature of the CABARET/UL scheme is that it approximates the entire material derivative on the Eulerian grid:

\[ L = \langle \partial_t \hat{u} + \partial_x \hat{f}(u) \hat{c}_x \rangle \]

rather than splits it into independent, space and time components

\[ L_x = \langle \partial_x \hat{f}(u) \cdot \hat{c}_x \rangle, \quad L = \langle \hat{c}_t \rangle + L_x. \]

Note that separate space and time treatment (9) is used in many high-resolution schemes based on high-order spatial-flux reconstruction techniques, combined with time-integration methods based on low-dissipative multi-step Runge–Kutta techniques (e.g., Bogey and Bailly, 2004) or on single-step predictor–corrector methods of Adams-type (e.g., Shchepetkin and McWilliams, 1998). In particular, we argue that in comparison to the methods based on the Adams-type schemes, the CABARET scheme (2)–(4) is non-dissipative, which makes it more suitable for turbulent flow regimes. A comparison of the dispersion properties of the CABARET/UL scheme with those of the central finite differences, including optimised high-order schemes typically used with low-dissipative Runge–Kutta time-integration schemes, is in the next section.

2.4. Results of linear dispersion analysis

The non-staggered form of finite-difference schemes (without nonlinear flux correction) is well suited for analysis of the linear-wave dissipation and dispersion properties. In this analysis a travelling wave solution, \(q_m^e \sim \exp[\text{i}(\omega \cdot \tau - n \cdot \mathbf{k} \cdot \text{h} \cdot m)]\), is substituted into the scheme. The resulting algebraic equation is solved for the frequency, \(\omega = \omega(k \cdot \text{h})\), or \(\frac{m}{\text{CFL}} = f(k \cdot \text{h}, \text{CFL})\), where \(f\) is a function of the dimensionless wavenumber, \(k \cdot \text{h}\), and of the Courant number, \(\text{CFL} = |\text{c}| \cdot \text{h} / \text{CFL}\). The outcome is used to determine the dissipation error, \(3m(\omega(k \cdot \text{h}) \cdot |\tau|) \neq 0\), and the dispersion error, \(f - 1 \neq 0\), as functions of the wavenumber and the Courant number. In Fig. 3 we compare the dispersion error of the CABARET scheme at different Courant numbers against the errors of semi-discrete finite differences including two state-of-the art high-order low-dispersion schemes used in turbulence modelling (e.g., Lele, 1992). Note, that the dispersion properties of semi-discrete schemes correspond to exact integration in time, that neglects possible increase of the dispersion error due to inaccuracies in time marching, and that is equivalent to using a multi-stage Runge–Kutta method with a very small step for time integration. We find that for most Courant numbers and for a wide range of grid resolution (7–20 points per wavelength) the dispersion error of the CABARET scheme remains

![Fig. 2. 1-D linear advection test: (a) with a rectangular initial profile, (b) with a half-sine-wave initial profile after 250 time steps on a uniform grid with \(\text{CFL} = 0.45\); squares denote the solution of CABARET scheme, circles denote the solution of the second-order Van Leer scheme with TVD MinMod limiter, triangles denote the solution of the same scheme with TVD Superbee limiter.](image)

![Fig. 3. Phase errors of several spatial finite-difference schemes and of the CABARET scheme at different CFL numbers vs grid resolution (number of grid points per wavelength, \(N = \pi(k \cdot h)\)). E2, E4, E6 denote standard central differences of the second, fourth and sixth-order, respectively, DRP denotes the fourth order Dispersion Relation Preserving scheme (Tam and Webb, 1993); and LUI stands for a sixth order pentadiagonal compact scheme of Pade-type (Colonius and Lele, 2004). Results for all central schemes correspond to infinitesimally small Courant number, which corresponds to the highest accuracy for most of the central schemes, and CABARET0.x stands for the CABARET scheme at Courant number \(\text{CFL} = x\).](image)
below that of the conventional and optimised fourth-order central finite differences and close to that of the six-order central schemes. Away from the optimal Courant number range \((\text{CFL} = 0.1)\), the CABARET dispersion error is similar to that of the fourth-order scheme. Within this range of grid resolution the error decay rate (error slope) of the CABARET solution is in between the forth- and the second-order schemes.

Another important property pertaining to the numerical dispersion error is the numerical group speed (normalized by the advection speed), \(c_g = \frac{c}{c} \frac{h}{2}\). Any deviation of \(c_g\) from 1 corresponds to an error in physical wave propagation. In particular, negative values of \(c_g\) lead to spurious wave reflection. In this case, all waves starting from a certain frequency are “grid locked” and their energy is back-scattered (Colonius and Lele, 2004). Fig. 4 shows the numerical group speeds of the same finite-difference schemes as in Fig. 3. It can be seen that all central schemes, including the high-order compact Pade-type scheme, have \(c_g < 0\) for short enough wavelengths. In contrast, the CABARET group speed remains always positive, and the waves propagate at all frequencies. The notations are the same as in Fig. 1.

2.5. CABARET extension for linear advection in multiple dimensions

Let us consider a linear advection equation in multiple dimensions:

\[
\frac{\partial}{\partial t} u + \sum_k \frac{\partial}{\partial x_k} f_k = 0, \quad f_k = c_k u, \quad c_k = \text{const.} > 0.
\]

(10)

For solving this equation with the CABARET scheme on a quadrilateral/hexagonal, possibly multiple-block, Cartesian grid, first, we introduce the conservation variables, \(u\), referred to the cell centres, and the flux variables, \(f_k\), referred to the cell face centres in each coordinate direction, \(1 \leq k \leq d\), where \(d = 1/3\) is the dimension of the problem.

Fig. 5 shows the assumed data structure for case \(d = 2\), within one cell in space. Solid circles labelled with letters show locations of the conserved variable, and open circles labelled with numbers show locations of the flux variable, same as for the 1-D CABARET scheme shown in Fig. 1. Then, the CABARET algorithm becomes:

**Predictor step,**

\[
\frac{u_{C}}{2} \frac{\partial}{\partial t} u_{C} + \sum_k \frac{\partial}{\partial x_k} f_{k} = 0.
\]

(11)

**Extrapolation step,**

\[
u_{1,k} = 2u_{C} - u_{5,k}, \quad 1 \leq k \leq d.
\]

(12)

**Corrector step,**

\[
\frac{u_{C} - u_{C}}{2} \frac{\partial}{\partial t} u_{C} + \sum_k \frac{\partial}{\partial x_k} f_{k} = 0.
\]

(13)

Here, the divergence terms are approximated using the flux CABARET variables defined within the same cell, with the second-order accuracy in space. The multidimensional CABARET scheme (11)–(13) is second order accurate in time and space, similar to the 1-D CABARET scheme (7)–(9). The multidimensional CABARET scheme is stable for \(0 < \text{CFL} < 1/d\). For large-gradient flows, the nonlinear flux correction is applied at the extrapolation step (compare with (15)):

\[
u_{1,k} = 2u_{C} - u_{5,k};
\]

\[
M = \max(u_{4,k}, u_{k}, u_{5,k}) + \int_{t-n}^{t} Q(t) \, dt;
\]

\[
m = \min(u_{4,k}, u_{k}, u_{5,k}) + \int_{t-n}^{t} Q(t) \, dt;
\]

\[
Q(t) = -\sum_k \frac{\partial}{\partial x_k} f_{k};
\]

If \(u_{1,k} > M\), \(u_{1,k} = M\);

If \(u_{1,k} < m\), \(u_{1,k} = m\).

Thus, the multidimensional CABARET scheme updates each flux variable by performing 1-D extrapolation separately for each coordinate direction, but uses the same conservation variable for all directions. Thus, it combines the simplicity of the dimension-by-dimension splitting approach with genuinely multidimensional advection treatment.

For demonstration, the classical solid body rotation test (Crowley, 1968) is considered, in which the initial cone distribution is specified on a uniform Cartesian grid in the centre of 1/4 of square computational domain. The advection velocity field corresponds to a steady rotation around the computational domain centre. The
vertex of the cone is a singularity, and preserving it numerically is rather challenging. Fig. 6 shows that after one revolution on a coarse grid (20 points per the cone diameter), the cone symmetry is still well preserved. This solution corresponds to the relative cone vertex clipping of 16% which is twice smaller than 34% vertex reduction reported by Hundsdorfer et al. (1995) for the third-order MUSCL TVD scheme.

2.6. Example of CABARET scheme for the system of hyperbolic equations

The CABARET method has been extended to the compressible Euler equations with the use of characteristic decomposition. Standard characteristic Eulerian methods are based on flow field decomposition into simple characteristic waves of constant amplitudes obtained from the solution of 1-D Riemann problem at each cell face (e.g., LeVeque, 2002; Hirsch, 2007). In contrast to those, the CABARET extension to the equations in primitive variables is based on the extrapolation of groups of characteristic variables, i.e., local Riemann invariants. In particular, the solution algorithm for the 1-D gas dynamics is outlined below.

The components of the conservative variables are updated at the predictor and corrector step (Fig. 1):

$$\begin{align*}
U_i - U_{i-1/2} + \frac{F(U_i) - F(U_{i-1/2})}{h_{i-1/2}} &= 0; \\
U_i - U_{i+1/2} + \frac{F(U_i) - F(U_{i+1/2})}{h_{i+1/2}} &= 0; \\
U &= (\rho, \rho u, E)^T, \quad F = (\rho u, \rho u^2 + p, p(E + p))^T,
\end{align*}$$

(15)

where $\rho$, $\rho u$, $E$ are density, momentum and energy, respectively.

At the extrapolation step the flux variables are computed at cell faces. In order to update the flux variable on the new time level at cell face $(i)$, we use local Riemann invariants that are referred to each cell centre, left, $(i - 1/2)$, and right, $(i + 1/2)$, from the computational node $(i)$. For example, ideal gas flows the following form of the local Riemann invariants referred to the space-time cell $(i + 1/2, n + 1/2)$ can be used:

$$R(\rho, \rho u, p|S_C) = (u + Ap^\mu, u - Ap^\mu, \ln(p \cdot \rho^{\gamma^2}))^T, \quad A = \frac{2\sqrt{\gamma}}{\gamma - 1}, \quad \mu = \frac{\gamma - 1}{2\gamma},$$

(16)

where $S_C = p_C^{\mu} / (\rho C)^{\gamma^2}$ is the cell-based entropy variable. The three components of the characteristic vector correspond to eigenvalues $\lambda(C)_C = (u + a, u - a, u_C). q = \nabla q$, where $a$ is sound speed and $\gamma$ is specific heat ratio. For locally isentropic flows, this choice of characteristic invariants allows increasing the solution accuracy during the extrapolation stage. For each computational cell face $(i)$, we compute its neighbouring local Riemann invariants (16) defined in cells $(i - 1/2)$ and $(i + 1/2)$, according to the CABARET data structure shown in Fig. 1. For example, for point 2 this involves computation of the local Riemann invariants in points 4, 5, 6 and C.D.E.F. The extrapolation of local Riemann invariants from cells (1-2-4-5) and (2-3-5-6) to point 2 results in two sets of local Riemann invariants. Once the local Riemann invariants are obtained, their values are subject to the conservative flux correction (4a). From the two vectors of characteristic variables a single conservative flux vector is reconstructed by using only those components of each vector that correspond to incoming characteristics to the cell face. Note that the use of a double set of local Riemann invariants is analogous to that based on using a double characteristic Jacobian matrix that is known to increase the robustness of characteristic decomposition techniques (Donat and Marquina, 1996).

As a demonstration of the CABARET approach capabilities, we consider the so-called Shu–Osher problem (Shu and Osher, 1988), in which a shock wave of Mach 3 interacts with a stationary entropy wave:

$$(\rho, u, p) = \begin{cases} 
(3.857143, 2.629369, 10.333333) & \text{for } x < 2; \\
(1 + 0.2 \sin(5x), 0, 1) & \text{for } x \geq 2;
\end{cases}$$

(17)

$x \in [0, 10]$.

The test is challenging for it requires from the numerical method not only to be shock-capturing but also to preserve the rich solution structure behind the shock front. Table 1 shows the comparison of the CABARET solution with other high-resolution methods considered by Woodward and Colella (1984). Fig. 7 shows the instantaneous density field for the CABARET solution at a longer time $t = 1.872$ that corresponds to the grid resolution of 400 cells.

The numerical errors and total number of time iterations are summarized in Table 1. The total number of iterations corresponds to the number of time steps $(NT)$ required to reach the control time moment, $t = 1.8$. The numerical error is defined as a relative error based on the following norms:

$$e_t = \frac{1}{N + 1} \sum_{i=0}^{N} \left| \frac{\rho - \rho_{exact}}{\rho_{exact}} \right|, \quad e_c = \max \left( \frac{|\rho - \rho_{exact}|}{\rho_{exact}} \right).$$

(18)

Table 1

| $NT$ | PLIMDE $e_t$ | WENO $e_t$ | CABARET $e_t$
|------|-------------|------------|-------------
| 200  | 0.0185      | 0.429      | 0.0214      | 0.902      | 0.229      | 0.1211      |
| 400  | 0.0183      | 0.188      | 0.0195      | 0.667      | 0.462      | 0.1467      |
| 800  | 0.0166      | 0.290      | 0.0162      | 0.369      | 0.927      | 0.0058      | 0.2400      |

WENO is the third-order Weighted Essentially Non-Oscillatory Scheme and PLIMDE is the second-order Parabolic Interpolation Method.

Fig. 6. Solid body rotation test: cone profile after one full revolution on Cartesian grid with resolution of 20 grid cells per cone diameter.

Fig. 7. Density solution profile for the Shu–Osher test for $t = 1.872$. 

---

As follows from the table, the maximum error in the $g_1$-norm remains smallest for the CABARET scheme, among all three schemes. In contrast to the other schemes, in the $g_1$-norm, the CABARET solution error leads to the convergence rate which is higher than 1 for all grids, including the coarsest one.

### 3. Application to the double-gyre circulation

In this section we apply the CABARET approach to a classical ocean circulation problem.

#### 3.1. Ocean model

We keep in mind ultimately applying the CABARET approach to a fully comprehensive, primitive-equation, eddy-resolving general circulation model of the ocean, but, for the purpose of this paper, we focus on the more simple QG ocean dynamics. The QG model of the wind-driven double-gyre circulation is considered in a mid-latitude closed basin, which is in the shape of a square with north-south and east-west rigid walls (L x L = 3840 3840 km). This model simulates not only the subpolar and subtropical ocean gyres but also the nonlinear western boundary currents, such as the Gulf Stream or Kuroshio, and their eastward jet extensions (e.g., Holton, 1978). The model stratification is represented by three stacked isopycnal layers that are dynamically coupled through pressure fluctuations. The governing equations constitute the system of material conservation laws for potential vorticity (PV) anomaly, $\zeta$, with a source term due to the meridional gradient of the Coriolis parameter and with the additional source terms due to the lateral viscosity, bottom friction, and the wind forcing. The system of the quasilinear hyperbolic-type (e.g., Rozhdestvensky and Yanenko, 1978) equations for PV anomaly and the associated elliptic equations for velocity streamfunctions are:

$$
\frac{\partial}{\partial t} \zeta + \frac{\partial}{\partial x} (u \zeta) + \frac{\partial}{\partial y} (v \zeta) = F_q, \quad q = \nabla \zeta.
$$

$$
F_q = \delta_{1g} F_{\text{wind}} - \beta \cdot v_q + \delta_{3g} \cdot \mu_{\text{bot}} \nabla^2 \psi_q + \mu_{\text{eddy}} \nabla^2 D_q,
$$

$$
u_q = \frac{\partial \psi_q}{\partial x}, \quad v_q = \frac{\partial \psi_q}{\partial y},
$$

$$
\zeta_1 = \nabla^2 \psi_1 - s_1 (\psi_1 - \psi_2), \quad D_1 = \zeta_1 + s_1 (\psi_1 - \psi_2),
$$

$$
\zeta_2 = \nabla^2 \psi_2 - s_2 (\psi_2 - \psi_1) - s_2 (\psi_2 - \psi_3), \quad D_2 = \zeta_2 + s_2 (\psi_2 - \psi_1) + s_2 (\psi_2 - \psi_3),
$$

$$
\zeta_3 = \nabla^2 \psi_3 - s_3 (\psi_3 - \psi_2), \quad D_3 = \zeta_3 + s_3 (\psi_3 - \psi_2),
$$

where $\beta$, $\mu_{\text{bot}}$, $\mu_{\text{eddy}}$, $s_1$, $s_2$, $s_3$ are the viscosity coefficients of the bottom and eddy viscosity, respectively.

The partial-slip condition is imposed:

$$
(u_1 n_x + v_q n_y) \big|_{r} = 0, \quad \left( \frac{\partial}{\partial \nu_q} (u_1 n_x + v_q n_y) - \alpha^{-1} (u_1 n_x + v_q n_y) \right) \big|_{r} = 0, \quad \alpha = \text{const. > 0}, \quad q = \nabla \zeta.
$$

where the notations are the same as in (19). This form emphasises the characteristic behaviour of the PV anomaly, as if it is a scalar transported tracer in a non-uniform divergence-free velocity field.

At the closed-basin boundary, $\Gamma$, the normal derivative is:

$$
\left( \psi_q - \frac{\partial \psi_q}{\partial x} \right) \big|_{\Gamma} = 0, \quad q = \nabla \zeta.
$$

and we use $\alpha = 120$ km.

Together with the integral mass conservation constraints (McWilliams, 1977).

$$
\frac{\partial}{\partial \nu} \int (\psi_1 - \psi_2) \, dx \, dy = 0 \quad \text{and} \quad \frac{\partial}{\partial \nu} \int (\psi_2 - \psi_3) \, dx \, dy = 0.
$$

The boundary condition (22) is a part of the elliptic-problem inversion of PV anomaly, $\psi_q$, into the velocity streamfunctions.

#### 3.2. Numerical method based on the CABARET scheme

In this section we introduce CABARET method to the ocean model described in Section 3.1.
Let us consider a uniform Cartesian grid, \( x_{i+1/2} - x_i = h, \) \( y_{j+1/2} - y_j = h, \) \( 1 \leq i, j \leq N, \) which covers the entire computational domain. In addition to this grid, we consider the nodes staggered from it by \( h/2: \)

\[
x_{i+1/2,j+1/2} = x_j + h/2; \quad y_{i+1,j+1/2} = y_{i+1,j} + h/2; \quad (24)
\]

\[
x_{i+1/2,j} = x_j; \quad y_{i+1,j+1/2} = y_{i+1,j} + h/2; \quad (25)
\]

\[
x_{i+1/2,j} = x_j + h/2; \quad y_{i+1,j} = y_{i+1,j+1/2}; \quad (26)
\]

The conservation variables of the CABARET scheme are referred to the cell centre points (fractional \( i \) and \( j \) indices); the flux variables in the \( x \)-direction are referred to the cell face centres (whole \( i \)-indices); and the flux variables in the \( y \)-direction are referred to the cell face centres (whole \( j \)-indices).

The extension of the 1-D CABARET algorithms (7)–(9) for the conservation law (19) is the following. At the predictor step, the PV anomaly value is computed at the mid-time level:

\[
\left( \frac{\zeta^n}{\nu_{i+1/2,j+1/2}} - \frac{\zeta^n}{\nu_{i,j+1/2}} \right) - \frac{(u^n)_{i+1/2,j+1/2} - (u^n)_{i,j+1/2}}{h} + \frac{(v^n)_{i+1/2,j+1/2} - (v^n)_{i,j+1/2}}{h} = (F^n)_{i+1/2,j+1/2} - q = \frac{1}{3}, \quad (27)
\]

where \((F^n)_{i+1/2,j+1/2}\) denotes the second-order accurate approximation (discussed later in this section) of the source function at the mid-time level; and the minimal local step, according to the stability constraint, is

\[
\tau^{n+1/2} = \min_{i,j} \left[ \frac{CFL \cdot h}{\max \left( \left( u^n \right)_{i+1/2,j+1/2}, \left( v^n \right)_{i+1/2,j+1/2} \right) } \right], \quad (28)
\]

where \(0 < CFL < 0.5\). The updated values of PV anomaly are used to compute the mid-time-level velocity streamfunction and its Laplacian, by solving the system of the elliptic equations,

\[
\begin{align*}
(\zeta^n)_{i+1/2,j+1/2} &= \nabla^2 \psi^n_{i+1/2,j+1/2} = s_1 \left( \psi^n_{i+1/2,j+1/2} - \psi^n_{i+1/2,j+1/2} \right) , \\
(\zeta^n)_{i+1/2,j+2} &= \nabla^2 \psi^n_{i+1/2,j+2} = s_2 \left( \psi^n_{i+1/2,j+2} - \psi^n_{i+1/2,j+2} \right) , \\
(\zeta^n)_{i+1/2,j+1} &= \nabla^2 \psi^n_{i+1/2,j+1} = s_3 \left( \psi^n_{i+1/2,j+1} - \psi^n_{i+1/2,j+1} \right) ,
\end{align*}
\]

complemented by the partial-slip boundary condition (22) approximated with the second-order one-sided finite-differences and by the imposed mass conservation constraint (23).

Once \(\psi^n_{i+1,2,j+1/2}\) is calculated, it is used for updating the velocity components at the cell faces:

\[
\begin{align*}
(\psi^n_{i+1,2,j+1/2})_{i+1/2} &= \frac{1}{4} \left[ (\psi^n_{i+1,2,j+1/2})_{i+1/2,j+1/2} + (\psi^n_{i+1,2,j+1/2})_{i+1/2,j+1/2} - (\psi^n_{i+1,2,j+1/2})_{i+1/2,j+1/2} \right] , \\
(u^n)_{i+1/2,j+1/2} &= \frac{(\psi^n_{i+1,2,j+1/2})_{i+1/2} - (\psi^n_{i+1,2,j+1/2})_{i+1/2}}{h} , \\
(v^n)_{i+1/2,j+1/2} &= \frac{(\psi^n_{i+1,2,j+1/2})_{i+1/2} - (\psi^n_{i+1,2,j+1/2})_{i+1/2}}{h} .
\end{align*}
\]

Then, each velocity component is extrapolated to the new time level, \( n + 1 \), with the second-order accuracy:

\[
\begin{align*}
(u^n)_{i+1/2,j+1/2} &= (1 + \delta) (u^n)_{i+1/2,j+1/2} - \delta (u^n)_{i+1/2,j+1/2} , \\
(v^n)_{i+1/2,j+1/2} &= (1 + \delta) (v^n)_{i+1/2,j+1/2} - \delta (v^n)_{i+1/2,j+1/2} ,
\end{align*}
\]

\[
\delta = \frac{\tau^n}{\tau^{n+1/2} + \tau^{n-1/2}}
\]

Once the velocity field is updated, the extrapolation step yields the new values of the flux variables:

\[
\begin{align*}
(\zeta^n)_{i+1/2,j+1/2} &= (1 + \delta) (\zeta^n)_{i+1/2,j+1/2} - \delta (\zeta^n)_{i+1/2,j+1/2} , \\
(\zeta^n)_{i+1/2,j+2} &= (1 + \delta) (\zeta^n)_{i+1/2,j+2} - \delta (\zeta^n)_{i+1/2,j+2} , \\
(\zeta^n)_{i+1/2,j+1} &= (1 + \delta) (\zeta^n)_{i+1/2,j+1} - \delta (\zeta^n)_{i+1/2,j+1} ,
\end{align*}
\]

\[
\delta = \frac{\tau^n}{\tau^{n+1/2} + \tau^{n-1/2}}
\]

At the lateral boundary cells, the upstream cell-face values are not available, because the partial-slip boundary condition is imposed on the corresponding centre-cell variables. Therefore, we use the first-order upwind approximation, e.g.:

\[
(\zeta^n)_{i+1,2,j+1/2} = (\zeta^n)_{i+1,2,j+1/2} , \quad (24)
\]

The computed cell-face values of the PV anomaly are corrected, if they are outside the limits given by the maximum principle:

\[
(\zeta^n)_{i+1,2,j+1/2} < (M^n)_{i+1,2,j+1/2} , \quad (\zeta^n)_{i+1,2,j+1/2} = (M^n)_{i+1,2,j+1/2} , \quad (35)
\]

\[
(\zeta^n)_{i+1,2,j+1/2} > (M^n)_{i+1,2,j+1/2} , \quad (\zeta^n)_{i+1,2,j+1/2} = (M^n)_{i+1,2,j+1/2} , \quad (36)
\]

Here, the minimum and maximum limit values account not only for the solution at the previous time step, as in (15), but also for the non-zero right-hand side in (19a). Thus, for a cell facing in the \( x \)-direction, we have:

\[
\begin{align*}
(\zeta^n)_{i+1,2,j+1/2} &= \min \left( (\zeta^n)_{i+1,2,j+1/2}, (\zeta^n)_{i+1,2,j+1/2} \right) \quad (37)
\]

\[
\begin{align*}
(u^n)_{i+1,2,j+1/2} &= \min \left( (u^n)_{i+1,2,j+1/2}, (u^n)_{i+1,2,j+1/2} \right) \quad (37)
\]

\[
\begin{align*}
(m^n)_{i+1,2,j+1/2} &= \min \left( (m^n)_{i+1,2,j+1/2}, (m^n)_{i+1,2,j+1/2} \right) \quad (37)
\]

\[
\begin{align*}
(q^n)_{i+1,2,j+1/2} &= \min \left( (q^n)_{i+1,2,j+1/2}, (q^n)_{i+1,2,j+1/2} \right) \quad (37)
\]

1 Note that without the flux corrections (35) and (36) the CABARET solution of the double-gyre problem diverges, same as the Arakawa method without the Robert-Asselin filtering in time.
where the source term is evaluated by approximating (19a) with the first-order accuracy in time and with the second-order accuracy in space:
\[
\left(Q_q\right)^{n+1/2}_{i,j+1/2,j-1/2} = \frac{\left(\psi_q\right)^{n+1/2}_{i,j+1/2,j-1/2} - \left(\psi_q\right)^n_{i,j+1/2,j-1/2}}{\tau_{n+1/2}} + \frac{\left(\psi_q\right)^{n+1/2}_{i,j+1/2,j-1/2} - \left(\psi_q\right)^n_{i,j+1/2,j}}{h}.
\]
(39)

Here, \((u_q)_{i,j+1/2,j-1/2} = \frac{1}{2} \left( (u_q)_{i+1,j+1/2}^{n+1} + (u_q)_{i-1,j+1/2}^{n+1} \right)\) in the domain interior, and \((u_q)_{i,j+1/2,j-1/2}^0 = 0\) on the boundary. In (39), the first-order accuracy in time cannot affect the overall order of accuracy of the algorithm, because approximation (39) is used only in the non-differential constraint (36).

Once the flux variables are updated at the new time step, the new values of the centre-cell conservation variables are computed at the corrector step:
\[
\left(\psi_q\right)^{n+1/2}_{i,j+1/2,j-1/2} = \frac{\left(\psi_q\right)^{n+1/2}_{i,j+1/2,j-1/2} - \left(\psi_q\right)^n_{i,j+1/2,j-1/2}}{\tau_{n+1/2}} + \frac{\left(\psi_q\right)^{n+1/2}_{i,j+1/2,j-1/2} - \left(\psi_q\right)^n_{i,j+1/2,j}}{h} + \frac{\left(\psi_q\right)^{n+1/2}_{i,j+1/2,j} - \left(\psi_q\right)^n_{i,j+1/2,j}}{h} = (F_q)^{n+1/2}_{i,j+1/2,j-1/2}.
\]
\(q = \psi_q\).
(40)

For well-resolved solution regions, where the corrections (34) and (35) remain inactive, the CABARET scheme (27)-(40) approximates the conservation laws (19) to the second order of accuracy in both space and time, provided that the approximation of the full source term, \((F_q)^{n+1/2}_{i,j+1/2,j-1/2}\), is second-order accurate.

In our algorithm, the full source term is separated into the combined wind and viscous component and into the \(\beta\)-term component:
\[
(F_q)^{n+1/2}_{i,j+1/2,j-1/2} = \left(F_{q_{\text{wind}}}ight)^{n+1/2}_{i,j+1/2,j-1/2} + \left(F_{q_{\text{Corrds}}}ight)^{n+1/2}_{i,j+1/2,j-1/2}.
\]
(41)

We find that in the ocean simulations the \(\beta\)-term component is numerically stiff, and, therefore, it can excite spurious basin-scale barotropic modes. In order to avoid this problem, a separate second order accurate predictor-corrector Adams-type scheme is used in order to increase the stability of the approximation in time:
\[
\left(F_{q_{\text{Corrds}}}^{n+1/2}ight)^{i+1/2}_{i,j} = \left(1 + \delta\right)\left(R_q\right)^n_{i,j+1/2,j-1/2} - \delta\left(R_q\right)^{n+1/2}_{i,j+1/2,j-1/2} - \frac{\beta}{2}\tau_{n+1/2}.
\]
(42)

where the residual is
\[
\left(R_q\right)^n_{i,j+1/2,j-1/2} = -\frac{1}{2} \beta \left( (\psi_q)_{i+1,j+1/2}^n + (\psi_q)_{i,j+1/2}^n \right).
\]
(43)

In our numerical algorithm, the \(\beta\)-term component (42) is added to the equations at the predictor time step (27).

Since, unlike in the Arakawa method, in the absence of viscous forces our method is equivalent to the conservation law of PV anomaly plus the \(\beta\)-term, rather than to the conservation law of PV itself, \(\int (\zeta + \beta \cdot \mathbf{v}) \, dx \, dy = \text{const.} \), \(q = \psi_q\), we need to make sure that our scheme is compatible with the discrete conservation law. This is indeed the case, since (43) is equivalent to the following conservative approximation of the \(\beta\)-term component:
\[
\left(F_q^{n+1/2}_{i+1/2,j},j-1/2 = \text{Flux}_{y,j-1} - \text{Flux}_{y,j+1} - \text{Flux}_{x} + O(\text{div}),
\]
\[
\text{Flux}_{y} = -\beta (\psi_q_{i+1/2,j}) / h,
\text{Flux}_{x} = -\beta (u_q_{i+1/2,j}) / h,
\]
(43a)

\[
y_{j+1/2} = \frac{1}{2} \left( (q_{j} + y_{j}) \right).
\]

and \(O(\text{div}) = \beta y_{j+1} \left[ \left( \psi_q^n_{i+1/2,j-1/2} - \psi_q^n_{i-1/2,j+1/2} - \psi_q^n_{i-1/2,j-1/2} + \psi_q^n_{i+1/2,j+1/2} \right) / h \right] = 0\), because the velocity field obtained from (29) and (30) is divergence-free. We have checked that the velocity divergence term is indeed of the order of round off error in all our calculations.

In contrast to the \(\beta\)-term component, the other component of the source term is numerically benign, and, therefore, it is approximated at the mid-time level by the second-order differences of both the velocity streamfunction and its Laplacian. The corresponding term is added to the equations after the elliptic problem (29) is solved, and after the velocity streamfunction at the mid-time level is updated:
\[
(F_{q_{\text{wind}}})^{n+1/2}_{i,j+1/2,j-1/2} = \delta t_{\text{wind}} \cdot \text{Flux}_{y} + \delta t_{\text{wind}} \cdot \text{Flux}_{x} \cdot \mu_{\text{hol}} \left( \nabla^2 \psi_q \right)^{n+1/2}_{i,j+1/2,j-1/2} + \mu_{\text{hol}} \left( \nabla^2 D_q \right)^{n+1/2}_{i,j+1/2,j-1/2}.
\]
(44)

Here, the Laplacians are evaluated directly from the solution of elliptic problems at the mid-time level:
\[
\nabla^2 \psi_q^{n+1/2}_{i+1/2,j} = \left( \nabla^2 \psi_q^{n+1/2}_{i+1/2,j} + S_{\text{wind}} \left( \psi_q^{n+1/2}_{i+1/2,j} - \psi_q^{n+1/2}_{i-1/2,j} \right) \right),
\]
\[
\nabla^2 \psi_q^{n+1/2}_{i,j+1/2} = \left( \nabla^2 \psi_q^{n+1/2}_{i,j+1/2} + S_{\text{wind}} \left( \psi_q^{n+1/2}_{i,j+1/2} - \psi_q^{n+1/2}_{i,j-1/2} \right) \right),
\]
\[
\nabla^2 \psi_q^{n+1/2}_{i,j} = \left( \nabla^2 \psi_q^{n+1/2}_{i,j} + S_{\text{wind}} \left( \psi_q^{n+1/2}_{i,j} - \psi_q^{n+1/2}_{i-1/2,j-1/2} \right) \right).
\]
(45)

In order to untangle the variables in the elliptical problem (45), the vertical normal-mode variables are employed. They allow to diagonalise the elliptic problem into the three independent Helmholtz problems (for the individual vertical modes), that are solved with the combined Fast Fourier Transform (FFT) and cyclic reduction methods.

4. Ocean modelling results

In this section, we discuss the ocean circulation solutions obtained with the CABARET algorithm and compare the results with those obtained by using the conventional second-order Arakawa method. The Arakawa method is based on the CL advection scheme and Robert–Asselin filtering in time (with the relaxation constant equal to 0.1). We have actually checked that the results reported in this section for the Arakawa method are virtually insensitive to the relaxation constant in the range 0.01–0.3. Both methods are implemented in the same quasigeostrophic ocean modelling code that (with the Arakawa method) was previously used in double-gyre simulations (e.g., Berloff et al., 2007). We also checked that, in comparison with the conventional Arakawa method, the CABARET method (24)-(42) is about 1.2–2 times slower. However, this is a small drawback relative to the advantage of the greatly increased accuracy of the model.

4.1. Moderate Reynolds numbers

First, we compare the CABARET method with the conventional second-order method and with a low-dispersion first-order upwind method. We use \(\mu_{\text{hol}} = 50 \text{ m}^2 \text{ s}^{-1}\) that corresponds to a moderate-Re flow regime, \(Re = 3200\). We use four uniform Cartesian grids with 129 × 129, 257 × 257, 513 × 513, and 1025 × 1025 nodes. For the moderate-Re case the solution convergence is confirmed against the finest grid, 2059 × 2059 simulation, which is almost indistinguishable from the fine 1025-grid solution (e.g., the difference of maximum r.m.s. fluctuations of the eastward jet is less than 0.1%). Each simulation starts from the state of rest, and...
each physical run time is $10^4$ days, which corresponds to about 0.5–1 million time steps on the fine, 1025 grid. The spin-up time, during which all solutions reach statistically stationary states, is 8000 days. The data for further processing are stored for the remaining physical time of 2000 days, which is long relative to most of the characteristic time scales of the mesoscale eddies. For the purpose of the grid convergence study, we consider the time averages of the solutions and their r.m.s. fluctuations. At the moderate $Re = 3200$, the Arakawa scheme begins to converge starting from the 513-grid (Figs. 8 and 9). On the 257-grid, the western boundary current eastward jet (EJ) separates below the correct latitude (as given by the converged finest-grid solution). We suggest that this spurious behaviour of the EJ is an artifact of the numerical dissipation and dispersion errors that can be significantly corrected with the CABARET algorithm. Overall, correct modelling of the EJ is one of the key targets that we seek to achieve on relatively coarse grids.

The convergence of the CABARET and Arakawa methods have been checked in three error norms and by the location of the separation point of the jet on the EJ (Table 2). The first two errors correspond to the standard maximum norm of the time-averaged upper-ocean streamfunction solution and to that of the r.m.s. fluctuations of the upper-ocean streamfunction solution, respectively. The third error corresponds to the absolute value of the relative error in predicting the maximum of the r.m.s. fluctuations of the upper-ocean streamfunction solution that characterises the strength of the EJ. As seen from the table, the CABARET scheme shows significantly smaller errors and faster convergence rate for all error norms in comparison to the Arakawa scheme, which for max $\|\nu_{ij}\|$ norm even fails to converge. The CABARET convergence rate is approximately linear in the maximum norms. For both methods, the relative error of the maximum r.m.s. jet fluctuations decays faster than the quadratic rate, with the CABARET method being still superior. The convergence of the EJ separation point is in Table 3, and it also shows a superiority of the CABARET solution.

4.2. High Reynolds numbers

The advantages of the CABARET method over the conventional second-order methods are more pronounced at high $Re$. To demonstrate this, we computed the double-gyre problem for a large $Re \approx 13,300 \ (\mu_{edd} = 12 \ m^2 \ s^{-1})$.

With our grid resolutions, and, unlike in the moderate-Re case (Section 4.1), there is no evidence of the solution convergence for either of the schemes except for the jet separation point location on the western boundary (Table 4).

Figs. 10 and 11 compare time averaged and r.m.s. fluctuations of the upper-ocean velocity streamfunction for the coarse 257-grid Arakawa and CABARET solutions with that of the fine 1025-grid Arakawa solution. We find that in contrast to the coarse-grid Arakawa solution the other two solutions are very similar. This similarity is further examined by plotting typical instantaneous snapshots of the upper-ocean streamfunction and PV anomaly for the same solutions (Figs. 12 and 13). In all figures the same contour levels of the streamfunction and PV anomaly are plotted. While the 257-grid Arakawa solution is strongly contaminated by spurious short-wave oscillations which completely obliterate the EJ structure, the 257-grid CABARET solution is remarkably close to the reference fine 1025-grid solution.

Fig. 8. Convergence of the Arakawa and CABARET methods for the moderate-Re case. Time-averaged, upper-ocean velocity streamfunction solutions: (a) $257 \times 257$ grid, (b) $513 \times 513$ grid, and (c) $1025 \times 1025$ grid.
To further validate this qualitative picture, we have checked that the underprediction of the maximum r.m.s. fluctuations of the EJ of the 257-grid Arakawa solution, in comparison to the reference 1025 solution, is 65%, thus indicating a total failure of the solution. In contrast to this, the CABARET solution on the 257 grid results in only 19% error in the maximum r.m.s. fluctuations of the EJ. We find this qualitative agreement with the 1025-grid solution remarkable, given that computation with the coarse 257-grid CABARET is more than 30 times faster than that with the fine 1025-grid Arakawa scheme.

5. Conclusions and discussion

The novel Compact Accurately Boundary Adjusting high-Resolution Technique (CABARET) works remarkably well for the eddy-resolving quasigeostrophic double-gyre ocean circulation problem. In comparison with the conventional second-order method, it allows to increase Reynolds number (Re) of the simulations by an order of magnitude - this opens exciting perspectives for exploring more realistic flow regimes.

The CABARET approach is based on several important ideas, which have been circulating in the computational physics community:

Table 2
Error convergence for the moderate-Re case.

<table>
<thead>
<tr>
<th>Grid</th>
<th>CABARET</th>
<th>Arakawa</th>
<th>CABARET</th>
<th>Arakawa</th>
<th>CABARET</th>
<th>Arakawa</th>
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<td>max</td>
<td></td>
<td>\epsilon_{\text{e}}</td>
<td></td>
<td>&amp; max</td>
<td></td>
</tr>
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<td>138.48</td>
<td>99.39</td>
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<td>132.66</td>
<td>189.07</td>
<td>56.58</td>
<td>86.87</td>
<td>0.071</td>
<td>0.098</td>
</tr>
<tr>
<td>513</td>
<td>50.98</td>
<td>199.44</td>
<td>0.003</td>
<td>71.29</td>
<td>0.003</td>
<td>0.012</td>
</tr>
</tbody>
</table>

Table 3
Convergence of the location of the separation point, y/L.

<table>
<thead>
<tr>
<th>Grid</th>
<th>257</th>
<th>513</th>
<th>1025</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arakawa</td>
<td>0.4</td>
<td>0.5</td>
<td>0.55</td>
</tr>
<tr>
<td>CABARET</td>
<td>0.5</td>
<td>0.55</td>
<td>0.55</td>
</tr>
</tbody>
</table>

The confidence interval for the error in separation point location, which was estimated by graphical data post-processing in Tecplot, is about ±0.01.

Table 4
Location of the separation point, y/L.

<table>
<thead>
<tr>
<th>Grid</th>
<th>257</th>
<th>513</th>
<th>1025</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arakawa</td>
<td>N/A</td>
<td>0.3</td>
<td>0.5</td>
</tr>
<tr>
<td>CABARET</td>
<td>0.5</td>
<td>0.55</td>
<td>0.5</td>
</tr>
</tbody>
</table>

See the footnotes of Table 3.

Fig. 9. Convergence of the Arakawa and CABARET methods for the moderate-Re case. The r.m.s. fluctuation of upper-ocean velocity streamfunction solutions: (a) 257 × 257 grid, (b) 513 × 513 grid, and (c) 1025 × 1025 grid.
(i) **Fully discrete/Lagrangian property:** optimal approximation of the entire material derivative rather than optimisation of the time and space discretisations separately, as in the standard Eulerian schemes.

(ii) **Low dispersion and no-dissipation error** of the underlining finite-difference scheme.

(iii) **Non-oscillatory property:** enforcing the maximum principle on the solution as the means for efficient treatment of the underresolved scales.

(iv) **Ease of implementation for complicated boundary conditions and non-uniform grids due to the compactness of stencil in space and time.**

(v) **Reduced CPU cost due to the small computational stencil.**

The novelty of our approach is in achieving good dispersion and dissipation properties of the underlying advection scheme that are crucial in large-Re eddy-resolving simulations. We deal with these properties in the situation when the regularising
effect of the explicit eddy viscosity is small relative to the dissipation and dispersion errors of the scheme. In order to prove this point, we presented a systematic comparison of the new CABARET method with popular high-resolution methods such as MUSCL, PPM and WENO, for several linear-advection and gas dynamics problems. Finally, we contrasted the performance of the CABARET scheme with that of the standard second-order Arakawa method for the classical ocean circulation problem. In particular we show that the numerical dispersion and dissipation error of the latter can be responsible for largely incorrect eastward jet structure.

For the double-gyre circulation, the CABARET method allows one to obtain high-quality solutions on the grid that has half a point over the Munk lengthscale, \((\frac{L_{M}}{\eta})^{1/3}\), which characterises the viscous western boundary layer. This is a very significant upgrade relative to the conventional scheme that requires 2–3 grid points over this lengthscale (e.g., Berloff et al., 2007).

In the current publication we have not attempted to implement other high-resolution schemes, such as those quoted in Section 2, for the double-gyre problem and to compare them with the CABARET method. This is because of the complexity of implementation of the former near the boundaries (e.g., the western boundary treatment is a crucial part of the double-gyre solution), since many such methods have a larger computational stencil in comparison to the CABARET or Arakawa scheme.

Potential future directions include adaptation of the CABARET method for non-uniform grids, which can be achieved by capitalising on the remarkable compactness of its computational stencil, and extension of the method to the primitive equations for use in the comprehensive ocean general circulation models.

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References


