

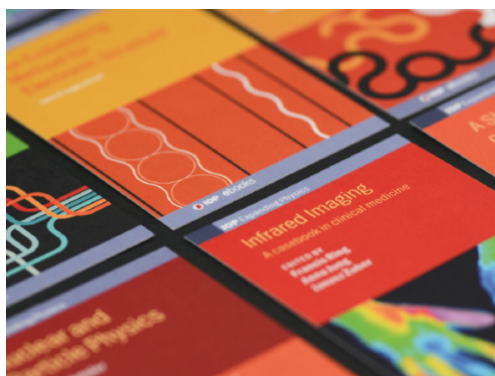
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Multi-block simulations in general relativity: high-order discretizations, numerical stability and applications

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Abstract

The need to smoothly cover a computational domain of interest generically requires the adoption of several grids. To solve a given problem under this grid structure, one must ensure the suitable transfer of information among the different grids involved. In this work, we discuss a technique that allows one to construct finite-difference schemes of arbitrary high order which are guaranteed to satisfy linear numerical and strict stability. The method relies on the use of difference operators satisfying summation by parts and *penalty terms* to transfer information between the grids. This allows the derivation of semi-discrete energy estimates for problems admitting such estimates at the continuum. We analyse several aspects of this technique when used in conjunction with high-order schemes and illustrate its use in one-, two- and three-dimensional numerical relativity model problems with non-trivial topologies, including truly spherical black hole excision.

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(Some figures in this article are in colour only in the electronic version)

1. Introduction

Many systems of interest have a non-trivial natural topology that a single cubical computational domain cannot accommodate in a smooth manner. Examples of these topologies in three-dimensional settings are $S^2 \times R$ (or a subset of it), encountered when dealing with spacetimes with smooth outer boundary and inner boundaries where required to excise singularities, and $S^2 \times S^1$ or S^3 topologies, commonly found in cosmological problems.

The need to treat these scenarios naturally leads one to consider multiple coordinate patches in order to cover the region of integration. These, in turn, translate into having to adopt multiple grids at the implementation level. Each one of these represents a region of discrete space, *a patch*, which might come equipped with a discrete Cartesian coordinate system or discrete charts; that is, invertible maps from discrete space to regions of Z^3 (i.e. the integers labelling coordinate grid points in each direction).

Covering a spacetime by charts is commonly done at the continuum level when considering the differential geometry of the spacetime (see for instance [1]). These charts are usually thought of as defining a map of a portion of the spacetime into a subset of R^3 , and the combination of charts (which usually overlap in some regions) covers the whole spacetime. Points belonging to an overlapping region are considered as belonging to any of the involved charts. Here a well-defined coordinate transformation between the charts is naturally defined by the combination of the maps and their inverse inbetween the spacetime and the charts.

At the discrete level, one can in principle adopt an analogue of the above construction. However, it is often the case that in the overlapping region a grid point in one of the charts does not have a corresponding one in the other. Consequently, the coordinate transformation is not defined. This presents a problem at the practical level as communication between patches must take place. This issue is commonly solved in two different ways: (i) by introducing further points via interpolation where needed; (ii) by considering patches that only *abut*, i.e. do not overlap.

In the first case—commonly referred to as the *overlapping-grids approach*—non-existent points in a given grid within the overlapping region are defined where needed by appropriate interpolations. Although this can be done in a straightforward manner with a relatively simple multiple-grid structure, the drawback of this approach is the introduction of a new ingredient—the interpolation—which does not have a counterpart at the continuum. This complicates the assessment of stability of even simple evolution problems as the details of the interpolation itself are intertwined with any attempt in this direction in an involved manner. As a consequence, there exist a few stability proofs for such evolution schemes and have so far been restricted to one-dimensional settings. Notwithstanding this point, a number of implementations in numerical relativity, with the addition of a certain amount of dissipation or filtering, make use of this approach with good results (see for instance [2–5]).

In the second case—commonly referred to as the *multi-block approach*—grids are defined in a way such that there is no overlap and only grid points at boundaries are common to different grids. This requirement translates into having to define the multiple grids with greater care than in the previous option. This extra effort, however, has as one pay-off that schemes preserving important continuum properties can be constructed. In particular, this allows the construction of stability analyses which are similar to those of a single grid. More explicitly, following Abarbanel, Carpenter, Nordström and Gottlieb [6–8], one can construct schemes of arbitrary high order for which semi-discrete energy estimates are straightforward to derive in a general way. The availability of stability results for this second approach makes it a very attractive option in involved problems—like those typically found when evolving Einstein equations—where schemes eliminating spurious sources of instabilities provide a strong starting point for a stable implementation of the problem.

In this paper, we discuss and analyse the use of this multi-block approach in the context of numerical relativity. At the core of the technique to treat outer and patch interfaces is the addition of suitable *penalty terms* to the evolution equations [6–8]. In the case of hyperbolic systems, these terms penalize the possible *mismatches* between the different values the characteristic fields take at the interface between several patches.

Not only does this method provide a consistent way to communicate information between the different patches but, more importantly, does so in a way which allows for the derivation of energy estimates at the semi-discrete level. Consequently, numerical stability can be ensured for a large set of problems. These estimates can be obtained with difference operators of any accuracy order, provided they satisfy the summation by parts (SBP) property and the penalty terms are constructed appropriately.

In this work, we discuss this technique in a context relevant to numerical relativity, analyse its properties and illustrate it in specific examples. In particular, we show results for the case of the $S^2 \times R$ topology used in black hole excision techniques.

This work is organized as follows. Section 2 includes a description of the numerical analysis needed to attain stability in the presence of multiple grids and summarizes how the penalty method of [6–8] allows for achieving this goal.

In section 3, we study some aspects of Strand’s [9] high-order operators satisfying SBP with respect to diagonal norms, when combined with the penalty technique. We find that in some cases, typically used operators that minimize the bandwidth have a very large spectral radius, with corresponding limitations in the Courant–Friedrich–Levy (CFL) factor when used in evolution equations. We therefore construct operators that minimize the spectral radius instead. Additionally, we examine the behaviour of the convergence rate and the propagation behaviour that different modes have when employing different higher order operators.

In section 4, we present and analyse different tests relevant to numerical relativity employing derivative operators of different orders of accuracy and the penalty technique to deal with multiple grids. These tests cover from linearized Einstein equations (in effectively one-dimensional scenarios) to propagation of three-dimensional fields in curved backgrounds.

We defer to appendices the discussion of several issues. Appendix A presents details of the higher derivative operators and diagonal norms which we employ in this work. Appendix B discusses our construction of high-order dissipative operators which are negative definite with respect to the corresponding SBP scalar product. Lastly, appendix C lists some useful properties that finite-difference derivative operator satisfy, which help in our construction of dissipative operators.

2. Interface treatment for symmetric hyperbolic problems in multiple blocks

As mentioned, we are interested in setting up a computational domain which consists of several grids which just abut. This domain provides the basic arena on which symmetric hyperbolic systems are to be numerically implemented. The basic strategy is to discretize the equations at each individual grid or block, treating boundary points in a suitable way. Boundary points at each grid either represent true boundary ones from a global perspective or lie at the interface between grids. In the latter case, since these points are common to more than one grid, the solution at them can be regarded as multi-valued. As we show below, this issue can be dealt with consistently and stably, ensuring that any possible mismatch converges to zero with resolution.

At the core of the technique is the appropriate communication of these possibly different values of the solution at the interfaces. Intuitively, since we are dealing with symmetric hyperbolic systems, a natural approach would be to communicate the characteristic variables from one domain to the other. However, this is not known to be numerically stable. There exists nonetheless a technique based on this strategy which does guarantee numerical stability [6–8]. This relies on adding *penalty terms* to the evolution equations of characteristic fields which penalize (a) in the interface case the mismatch between the different values each characteristic field takes at the interface of several grids, and (b) in the outer boundary case the difference

between each incoming characteristic field and the boundary conditions one wishes to impose on it.

These penalty terms are constructed so as to guarantee the stability of the whole composite grid if it can be guaranteed at each individual grid through the energy method. To this end, the use of schemes with difference operators satisfying SBP are employed. Hence, on each single grid there exists a family of natural semi-discrete energies, defined by both a symmetrizer of the continuum equations *and* a discrete scalar product with respect to which SBP holds⁵. One can then define an energy for the whole domain by simply adding the different energies of each grid. The use of operators satisfying SBP allows one to get an energy estimate, up to outer boundary and interface terms left after SBP. The penalties allow us to control their contribution, thus obtaining an estimate for the global grid. This is achieved if the contribution to the time derivative of the energy due to the interface and outer boundary terms (in the latter case when, say, homogenous maximally dissipative outer boundary conditions are imposed) left after SBP is non-positive. When these terms are exactly zero, the penalty treatment of [6–8] is, in a precise sense, ‘energy non-dissipative’. On the other hand, if these terms are negative, the scheme is numerically stable but at fixed resolution a damping of the energy (with respect to the growth one would obtain in the absence of an interface) in time arises. This damping is proportional to either the mismatch of a given characteristic variable at each interface or its failure to satisfy an outer boundary condition. As we describe below, these interface and boundary terms left after SBP are controlled precisely by the mentioned penalties, each of which depends on *the possible mismatch, the characteristic speeds, the corresponding SBP scalar product at the interface, the resolution at each intervening grid and a free parameter which regulates the strength of the penalties*.

Next, we explicitly describe how this penalty technique allows one to derive semi-discrete energy estimates. We first discuss in detail the one-dimensional example of an advection equation on a domain with a single interface. The more general case of systems of equations in several dimensions follows essentially the same principles, applying the 1D treatment to each characteristic field. We illustrate this by discussing a general constant-coefficient system in a given two-dimensional setting. From this, the generalization to the three-dimensional general case is straightforward, and we therefore highlight only its salient features.

2.1. A one-dimensional example

Consider a computational domain represented by a discrete grid consisting of points $i = i_{\min}, \dots, i_{\max}$ and grid spacing h covering $x \in [a, b]$. A 1D difference operator D on such a domain is said to satisfy SBP with respect to a positive definite scalar product (defined by its coefficients σ_{ij})

$$\langle u, v \rangle = h \sum_{i,j} u_i v_j \sigma_{ij}, \quad (1)$$

if the property

$$\langle u, Dv \rangle + \langle v, Du \rangle = (uv)|_a^b$$

holds for all grid functions u, v . The scalar product/norm is said to be *diagonal* if $\sigma_{ij} = \sigma_{ii} \delta_{i,j}$.⁶ One advantage of 1D difference operators satisfying SBP w.r.t. diagonal norms is that SBP is

⁵ Symmetrizers are not unique, already at the continuum level. In the case in which there is a preferred one, there is, similarly, a single, preferred semi-discrete energy on each grid. This is the case, for example, when one can derive a sharp energy estimate at the continuum, which gives rise to the construction of a strictly stable scheme; see [18] for a detailed discussion in the context of numerical relativity.

⁶ Here $\delta_{i,j}$ is the Kronecker delta ($\delta_{i,j} = 1$ if $i = j$ and zero otherwise).

guaranteed to hold in several dimensions if the 1D operator is used in each direction (which is not known to hold in the non-diagonal case in general) [10]. Even in 1D, in the variable coefficients and non-diagonal case, the commutator between D and the principal part might not be bounded for all resolutions (something that is *generically*⁷ needed for an energy estimate to hold) [11]. Another advantage is that the operators are, for a given order in the interior, simpler in their expressions. The disadvantage is that their order at and close to boundaries is half of that in the interior, while in the non-diagonal case the operators lose only one order with respect to the interior [9, 12]. Throughout this paper, we will mostly restrict our treatment to the use of diagonal norms.

As an example of how to impose interface or outer boundary conditions through penalty terms, we concentrate next on the advection equation for u propagating with speed Λ ,

$$\dot{u} = \Lambda \partial_x u. \quad (2)$$

2.1.1. A domain with an interface. Consider the interval $(-\infty, \infty)$ with appropriate fall-off conditions at infinity. We consider two grids: a *left* one covering $(-\infty, 0]$ and a *right* one covering $[0, \infty)$. We refer to the grid function u on each grid by u^l and u^r (corresponding to the left and right grids, respectively). Both of these grid functions have a point defined at the $x = 0$ interface and they need not coincide there, except at the initial time. Therefore, the numerical solution will in principle be multi-valued at $x = 0$, though, as we shall see, the penalty technique is designed to keep this difference small.

The problem is discretized using on the right and left grids, respectively, grid spacings h^l, h^r —not necessarily equal—and difference operators D^l, D^r satisfying SBP with respect to scalar products given by the weights σ^l, σ^r at their *individual grids*. That is, these scalar products are defined through

$$\langle u^l, v^l \rangle = h^l \sum_{i,j=-\infty}^0 \sigma_{ij}^l u_i^l v_j^l, \quad \langle u^r, v^r \rangle = h^r \sum_{i,j=0}^{\infty} \sigma_{ij}^r u_i^r v_j^r.$$

The semi-discrete equations are written as

$$\dot{u}_i^l = \Lambda D^l u_i^l + \frac{\delta_{i,0} S^l}{h^l \sigma_{00}^l} (u_0^r - u_0^l), \quad i = -\infty, \dots, 0, \quad (3)$$

$$\dot{u}_i^r = \Lambda D^r u_i^r + \frac{\delta_{i,0} S^r}{h^r \sigma_{00}^r} (u_0^l - u_0^r), \quad i = 0, \dots, \infty. \quad (4)$$

Note in the above equations the second term on each right-hand side, which constitutes the penalty added to the problem. They are defined by the possible mismatch, the grid spacing, the inner product employed and the free parameters $\{S^l, S^r\}$, which will be determined by requiring an energy estimate to hold.

We define a natural energy for the whole domain, which is the sum of the energies for each grid (for this simple example with a trivial symmetrizer),

$$E := \langle u^l, u^l \rangle + \langle u^r, u^r \rangle.$$

Taking a time derivative of this energy, using the semi-discrete equations (3) and (4) and the SBP property, one gets

$$E_t = (\Lambda - 2S^l)(u_0^l)^2 + (-\Lambda - 2S^r)(u_0^r)^2 + 2(S^l + S^r)u_0^l u_0^r. \quad (5)$$

In order to get an energy estimate, the above interface term (i.e., the right-hand side of equation (5)) must be non-positive for all u_0^l, u_0^r . It is straightforward to check that this is

⁷ That is, unless the scheme can be written in strictly stable form.

equivalent to the following three conditions holding:

$$\Lambda - 2S^l \leq 0, \quad (6)$$

$$-\Lambda - 2S^r \leq 0, \quad (7)$$

$$(\Lambda + S^r - S^l)^2 \leq 0. \quad (8)$$

From this, it is clear that we need $\Lambda + S^r - S^l = 0$. And with this condition, the other two become $S^l + S^r \geq 0$. There are three possibilities:

- *Positive Λ* : we can take

$$S^l = \Lambda + \delta, \quad S^r = \delta, \quad \text{with } \delta \geq -\frac{\Lambda}{2}. \quad (9)$$

The time derivative of the energy with this choice becomes

$$E_t = -(u_0^l - u_0^r)^2 (\Lambda + 2\delta) \leq 0.$$

- *Negative Λ* : similarly, we can take

$$S^r = -\Lambda + \delta, \quad S^l = \delta, \quad \text{with } \delta \geq \frac{\Lambda}{2}. \quad (10)$$

The corresponding time derivative of the energy with this choice becomes

$$E_t = (u_0^l - u_0^r)^2 (\Lambda - 2\delta) \leq 0.$$

- *Vanishing Λ* : this can be seen as the limiting case of any of the above two, and we can take

$$S^l = S^r = \delta, \quad \text{with } \delta \geq 0. \quad (11)$$

Hence,

$$E_t = -(u_0^l - u_0^r)^2 2\delta \leq 0.$$

The coefficients S^l and S^r need not be equal, but the following symmetry is important, under the change $\Lambda \rightarrow -\Lambda$ we should have $S^l \rightarrow S^r$ and vice versa, since it transforms incoming modes to outgoing ones. This is clearly satisfied by the above choice.

Summarizing, there is a freedom in the penalty factors of equations (3) and (4), encoded in the parameter δ , which has to satisfy $\delta \geq -|\Lambda|/2$. If $\delta = -|\Lambda|/2$, there is no interface term in the estimate (that is, exact energy conservation in the above model), while if $\delta > -|\Lambda|/2$ there is a negative definite interface term in the estimate (which represents a damping in the energy proportional to the mismatch).

As we will see below, one proceeds similarly in the more general case of systems of equations in several dimensions. The penalty terms are applied to the evolution equation of each characteristic mode, with factors given by equations (9)–(11) (where Λ in the general case is the corresponding characteristic speed).

2.1.2. A domain with an outer boundary. The penalty method also allows us to treat outer boundaries in a similar way. As an example, consider again the advective equation (2), but now on the domain $(-\infty, 0]$, with grid points $i = -\infty, \dots, 0$. Assume $\Lambda > 0$; boundary conditions therefore need to be given at $x = 0$, say $u(x = 0, t) = g$. The semi-discrete equations are written as

$$\dot{u}_i = \Lambda D u_i + \frac{\delta_{i,0} T}{h \sigma_{00}} (g - u_0). \quad (12)$$

Defining the energy to be

$$E = \langle u, u \rangle,$$

its time derivative is

$$\dot{E} = (\Lambda - 2T)u_0^2 + 2gu_0T, \quad (13)$$

$$\leq (\Lambda - T)u_0^2 + Tg^2. \quad (14)$$

As in the interface case with positive speed (cf equation (9)), we can therefore take $T = \Lambda + \delta$. For the homogeneous case, $g = 0$, the equality (13) holds and we have

$$\dot{E} = (\Lambda - 2T)u_0^2,$$

indicating that for $\delta \geq -\Lambda/2$ the energy will not increase. For the non-homogeneous case ($g \neq 0$), the inequality (14) yields

$$\dot{E} \leq \Lambda g^2 + \delta(g^2 - u_0^2).$$

Note that for $\delta = 0$ one trivially recovers the continuum estimate. For other values of δ , the consistency with the continuum estimate follows from the observation that u_0 converges to g to the $(s+1)$ th order if the SBP derivative operator has accuracy s at the boundary point⁸. This implies that both the numerical implementation and the corresponding energy estimate are consistent with those defined at the continuum level.

At this point, we find it important to make the following remark. Note that just having an energy inequality is not enough, as one further needs to ensure consistency of the discrete equations with respect to the continuum ones. In the case of the penalty approach, this is not straightforward as the penalty term diverges when the grid size decreases unless u converges to g sufficiently fast (as mentioned above, u does converge to g fast enough if u is an incoming mode). In fact, the penalty term can be viewed at the continuum as approximating the original equation and boundary conditions through the introduction of a suitable delta function at the boundary. The argument of the delta function must be consistent with the underlying problem. For instance, if one were to put a penalty term on a boundary where the mode is outgoing—and thus the value of the function there is determined by the evolution itself—the inconsistency would manifest itself through a lack of convergence. In the above example, this would be the case if we take $\Lambda < 0$ but insist on putting a penalty term at $x = 0$. Note that in such a situation the energy inequality would still hold if $T \geq \Lambda/2$. This would imply that the numerical solution is still bounded in the L^2 norm, but no more than that; indeed, numerical experiments show that a high-frequency solution travelling in the incoming direction (that is, with velocity opposite to that at the continuum, see section 3.3), whose amplitude depends on the size of the penalty term, is generated. Naturally, if $T < \Lambda/2$, the energy inequality is violated and the solution blows up exponentially with a rate increasing with the highest frequency that can be accommodated by the grid being employed.

2.2. The two-dimensional case

Consider now the system of equations

$$\dot{u} = A^\mu \partial_\mu u = A^x \partial_x u + A^y \partial_y u,$$

where u is a vector-valued function, A^x, A^y are symmetric and, for simplicity, constant-

⁸ We thank Olivier Sarbach for clarifying discussions on this point.

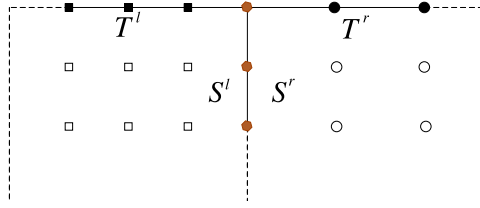


Figure 1. Example of a multi-block domain in two dimensions. Only the spacing in the vertical direction need be the same in both grids so as to ensure that boundary points—represented by grey hexagons—coincide.

coefficient matrices. The domain is composed of two grids: a *left* one covering $(-\infty, 0] \times (-\infty, 0]$ and a *right* one covering $[0, \infty) \times (-\infty, 0]$, with an interface at $x = 0$ and an outer boundary at $y = 0$ (see figure 1). At $y = 0$, we impose homogeneous maximally dissipative boundary conditions by setting to zero the incoming characteristic fields.

We assume that the scalar products associated with the different 1D difference operators are diagonal. This, as already mentioned, ensures certain properties that guarantee an energy estimate.

We denote the grid spacings, difference operators and associated scalar products corresponding to the x, y directions by h_x, h_y, D_x, D_y and $\sigma_{(x)}, \sigma_{(y)}$, respectively. These quantities need not coincide on the different subdomains, except for the grid spacing in the transversal direction at an interface (so that grid points belonging to different grids align with each other). To all quantities we add an l or r supraindex to denote quantities belonging to the left or right domain. Thus, the only condition we require is $h_y^l = h_y^r =: h_y$.

On each subdomain, the 2D scalar product is defined as the product of the scalar product in each direction,

$$\langle u, v \rangle = h_x h_y \sum_{i,j} (u_{ij}, v_{ij}) \sigma_{(x)i} \sigma_{(y)j}, \quad (15)$$

where (u, v) is the pointwise Euclidean scalar product of two vectors.

As in 1D, the total energy is defined as the sum of the energies on each subdomain (in this case with a trivial symmetrizer, as the equations are already in symmetric form):

$$E = E^l + E^r,$$

where

$$E^l = h_x^l h_y \sum_{i \leq 0} \sum_{j \leq 0} (u_{ij}^l, u_{ij}^l) \sigma_{(x)i}^l \sigma_{(y)j}^l, \quad (16)$$

$$E^r = h_x^r h_y \sum_{i \geq 0} \sum_{j \leq 0} (u_{ij}^r, u_{ij}^r) \sigma_{(x)i}^r \sigma_{(y)j}^r. \quad (17)$$

The evolution equations are a combination of equations (3), (4) and (12),

$$\dot{u}_{ij}^l = A^\mu D_\mu^l u_{ij}^l + \frac{\delta_{i,0} S^l}{h_x^l \sigma_{(x)i=0}^l} (u_{0j}^r - u_{0j}^l) - \frac{\delta_{j,0} T^l}{h_y \sigma_{(y)j=0}^l} u_{i0}^l, \quad (18)$$

$$\dot{u}_{ij}^r = A^\mu D_\mu^r u_{ij}^r + \frac{\delta_{i,0} S^r}{h_x^r \sigma_{(x)i=0}^r} (u_{0j}^l - u_{0j}^r) - \frac{\delta_{j,0} T^r}{h_y \sigma_{(y)j=0}^r} u_{i0}^r. \quad (19)$$

In the above expressions, S^l, S^r, T^l, T^r are operators (as opposed to scalars), since we are dealing with a system of equations. The first two correspond to penalty terms added to handle grid interfaces and the latter two for imposing outer boundary conditions. The goal of these

operators is to transform to characteristic variables and apply to the evolution equation of each characteristic mode suitable penalty terms, as in equations (3), (4) and (12).

Taking a time derivative of the energies defined in equations (16) and (17), using the evolution equations (18) and (19), and employing the SBP property along each direction, one gets

$$\dot{E}^l = h_y \sum_{j \leq 0} \sigma_{(y)j}^l [(u_{0j}^l, (A^x - 2S^l)u_{0j}^l) + 2(u_{0j}^l, S^l u_{0j}^r)] + h_x^l \sum_{i \leq 0} \sigma_{(x)i}^l (u_{i0}^l, (A^y - 2T^l)u_{i0}^l), \quad (20)$$

$$\dot{E}^r = h_y \sum_{j \leq 0} \sigma_{(y)j}^r [(u_{0j}^r, (-A^x - 2S^r)u_{0j}^r) + 2(u_{0j}^r, S^r u_{0j}^l)] + h_x^r \sum_{i \geq 0} \sigma_{(x)i}^r (u_{i0}^r, (A^y - 2T^r)u_{i0}^r), \quad (21)$$

where we have assumed that S and T are Hermitian matrices.

In order to control the interface terms in $\dot{E}^l + \dot{E}^r$, we can take

$$S^l = \frac{1}{\sigma_{(y)j}^l} [(\Lambda_a^+ + \delta_a^+) P_+^a + \delta_a^- P_-^a + \delta^0 P_0],$$

$$S^r = \frac{1}{\sigma_{(y)j}^r} [(-\Lambda_a^- + \delta_a^-) P_-^a + \delta_a^+ P_+^a + \delta^0 P_0],$$

where a sum over the index a is assumed and $\{P_+^a, P_-^a, P_0\}$ are projectors to the sub-spaces of eigenvectors of A^x with eigenvalues $\{\Lambda_a^+, \Lambda_a^-, \Lambda^0\}$, respectively. With this choice, $\dot{E}^l + \dot{E}^r$ becomes

$$\begin{aligned} \dot{E}^l + \dot{E}^r = h_y \sum_{j \leq 0} [(\Lambda_a^- - 2\delta_a^-) \|u_-^{a,l} - u_-^{a,r}\|^2 - (\Lambda_a^+ + 2\delta_a^+) \|u_+^{a,l} - u_+^{a,r}\|^2 - 2\delta^0 \|u_0^l - u_0^r\|^2] \\ + h_x^l \sum_{i \leq 0} \sigma_{(x)i}^l (u_{i0}^l, (A^y - 2P^l)u_{i0}^l) + h_x^r \sum_{i \geq 0} \sigma_{(x)i}^r (u_{i0}^r, (A^y - 2P^r)u_{i0}^r). \end{aligned} \quad (22)$$

Clearly, in order to obtain an estimate, the following conditions must be satisfied:

$$\Lambda_a^- - 2\delta_a^- \leq 0, \quad \Lambda_a^+ + 2\delta_a^+ \geq 0, \quad \delta^0 \geq 0,$$

which is analogous to the one-dimensional case, equations (9)–(11). Similarly, the outer boundary terms in $\dot{E}^l + \dot{E}^r$ (i.e. the sums over i) can be controlled on each domain separately. We need

$$u^r (A^y - 2T^r) u^r \leq 0, \quad u^l (A^y - 2T^l) u^l \leq 0.$$

We can therefore take, as in the one-dimensional case,

$$P^r = P^l = (\Lambda_a^+ + \delta_a^+) P_a^+,$$

where now P_a^+ are projectors to the spaces of eigenvectors of A^y of eigenvalues Λ_a^+ . With these choices, the final expression for the time derivative of the energy is

$$\begin{aligned} \dot{E}^l + \dot{E}^r = h_y \sum_{j \leq 0} [(\Lambda_a^- - 2\delta_a^-) \|u_-^{a,l} - u_-^{a,r}\|^2 - (\Lambda_a^+ + 2\delta_a^+) \|u_+^{a,l} - u_+^{a,r}\|^2 - 2\delta^0 \|u_0^l - u_0^r\|^2] \\ + h_x^l \sum_{i \geq 0} \sigma_{(x)i}^l (-\Lambda_a^- - 2\delta_a^-) \|u_-^{a,l}\|^2 + h_x^r \sum_{i \geq 0} \sigma_{(x)i}^r (-\Lambda_a^- - 2\delta_a^-) \|u_-^{a,r}\|^2 \end{aligned} \quad (23)$$

and, again, the possible ranges for the different δ s are, as in the 1D case, equations (9)–(11).

Note that nothing special has to be done at a corner, as each direction is treated and controlled independently.

2.3. The general case

The general case follows the same rules. Namely, we must add penalty terms on the characteristic modes corresponding to each of the boundary matrices separately and accordingly.

For example, what should be done at the vertices of three patches meeting in the cubed-sphere case discussed later in section 4.1.3? (The treatment at the edges follows along the same lines discussed above.) As we will see, we have three meshes with coordinates (at a constant radius) (a^1, b^1) , (a^2, b^2) and (a^3, b^3) arranged in a clockwise distribution according to the indices and intersecting at a point (see figure 18). In that case, taking into account the orientation of the coordinates on each patch, the contribution to the energy (without the penalty terms added to the evolution equations) is proportional to

$$(u_{0N}^1, (A^{a^1} + A^{b^1})u_{0N}^1) + (u_{00}^2, (A^{a^2} + A^{b^2})u_{00}^2) + (u_{0N}^3, (A^{a^3} + A^{b^3})u_{0N}^3).$$

Since the interfaces are aligned to the grids, we know that the normals coincide on both sides; therefore, we have

$$A^{a^1} = A^{a^3}, \quad A^{b^1} = A^{b^2}, \quad A^{b^3} = A^{a^2}.$$

So we include penalty terms on each side, including the endpoints of the grids in each direction (which constitute vertices and edges). Note that the characteristic modes at these points are computed with the normal with respect to the side that contains this direction. Consequently, points at edges/vertices of a (topologically) cubical grid will have two/three penalty terms.

3. High-order difference operators with diagonal norms

In this section, we analyse some aspects of Strand's 1D difference operators satisfying SBP with respect to diagonal metrics, when used in conjunction with the penalty technique to construct high-order schemes for handling domains with interfaces.

In particular, we discuss operators with accuracy of orders 2, 4, 6 and 8 at interior points. The requirement that these operators satisfy the SBP property with respect to diagonal norms implies that their accuracy orders at and close to boundaries are 1, 2, 3 and 4, respectively. We will therefore refer to these operators as D_{2-1} , D_{4-2} , D_{6-3} and D_{8-4} . Some of these operators are not unique, as the accuracy order and SBP requirements still leave in some cases additional freedom in their construction. Indeed, while the first two operators (D_{2-1} , D_{4-2}) are unique, D_{6-3} comprises a mono-parametric family and D_{8-4} a three-parametric one. This freedom can be exploited for several purposes, for instance, to minimize the operator's bandwidth or its spectral radius. While the former produces operators which are more compact, the latter can have a significant impact on the CFL limit when dealing with evolution equations. Indeed, for the D_{8-4} case, minimizing its bandwidth leads to a considerably larger spectral radius (though this does not happen in the D_{6-3} case) which, in turns, requires one to employ a rather small CFL factor for the fully discrete scheme to be stable if an explicit time integration is used.

To analyse this in each case, we numerically solve and discuss the eigenvalues of the amplification matrix multiplied by h of the advection equation with speed 1, $\dot{u} = \partial_x u$, under periodic boundary conditions. The periodicity is imposed through an interface with penalty terms and hence the scheme does depend on the penalty parameter δ and so will the discrete eigenvalues obtained. As discussed in section 2, in the case in which $\delta = -1/2$, SBP holds across the interface, and the energy for this model is strictly conserved. In other words, the amplification matrix is anti-symmetric and the eigenvalues are purely imaginary (see section 2). On the other hand, if $\delta > -1/2$, there is a negative definite interface term left after SBP, and a

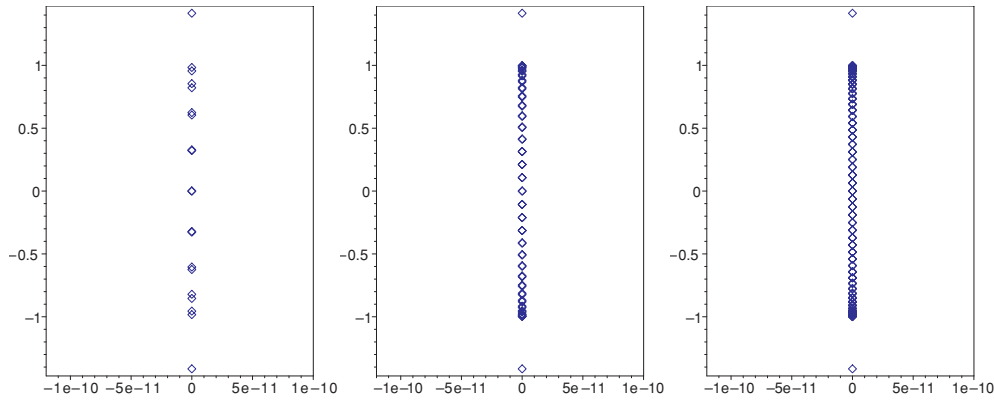


Figure 2. Numerically obtained eigenvalues (in the complex plane) corresponding to the D_{2-1} operator for $\delta = -1/2$ (purely imaginary case). From left to right the plots illustrate the results obtained with a grid containing 20, 60, 100 points, respectively. It is clear from the figures that these correspond, indeed, to a purely imaginary case.

negative real component in the eigenvalues must appear in the spectrum of the amplification matrix (see section 2).

We additionally discuss the global convergence factor for these operators and recall a feature associated with the mode with highest possible group speed at a given number of grid points, namely, that it travels in the ‘wrong’ direction and that the absolute value of its speed increases considerably with the order of the operator.

Appendix A lists, for completeness, some typos in [9] in some of the coefficients for these high-order operators.

3.1. Spectrum

In the following, we discuss the range of discrete eigenvalues obtained for the different derivative operators and their dependence on δ . We pay particular attention on the impact different values of δ and the chosen derivative operator have on the CFL limit.

3.1.1. Second order in the interior, first order at boundaries (D_{2-1} scheme). Figure 2 shows the eigenvalues obtained using 20, 60, 100 grid points and penalty term $\delta = -1/2$ (that is, the purely imaginary spectrum case, see section 2) for the D_{2-1} case. The maximum and minimum values are, approximately, ± 1.414 , and they seem to be related to the operator near the boundary, as their absolute value does not seem to increase with the number of points (instead, the region between the maximum and the minimum is filled out). As discussed below, in the higher order cases the maximum eigenvalues also seem to be related to the operator near the boundary.

Figure 3, in turn, shows the eigenvalues computed with 100 points and $\delta = 0, 1/10, 1/2$. A negative real part appears, as it should (based on the energy calculation), and the maximum in the imaginary axis slightly decreases (to approximately 0.999, not varying much among these three values). However, the maximum absolute value in the negative real axis grows quite fast with δ . For example, for $\delta = 1/10$, such a maximum already dominates over the maximum in the imaginary axis. The higher order operators analysed below behave similarly.

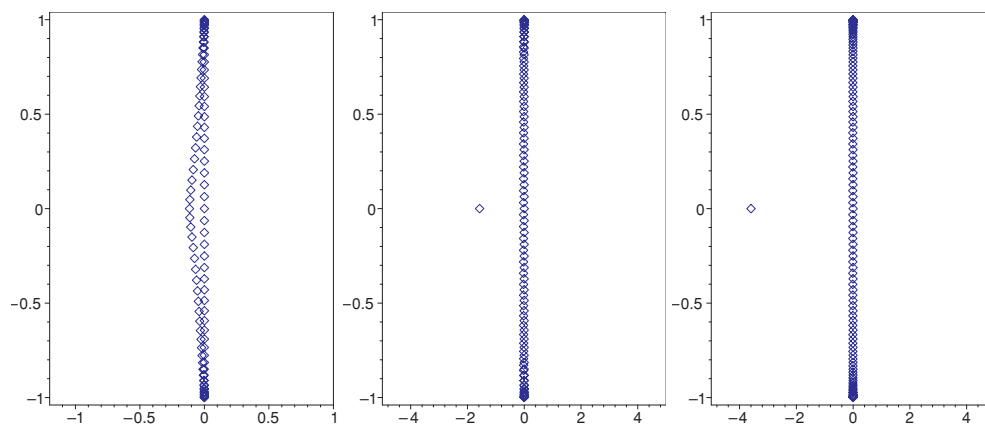


Figure 3. Eigenvalues corresponding to the D_{2-1} operator, obtained with a grid containing 100 points. From left to right the plots illustrate the behaviour for $\delta = 0, 1/10, 1/2$, respectively. As δ becomes larger, a larger (in magnitude) negative eigenvalue on the real axis is observed (note the left-most diamond at $y \simeq -1.6, -3.5$ in the middle and right-hand plots, respectively).

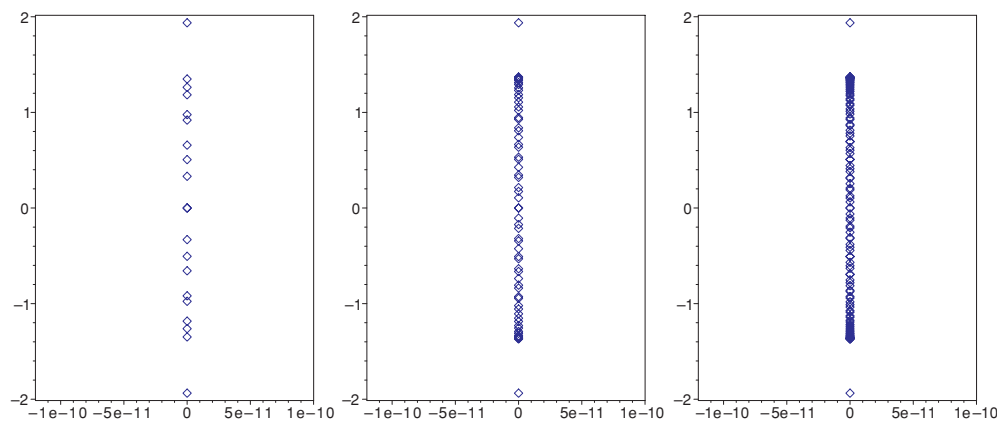


Figure 4. Numerically obtained eigenvalues corresponding to the D_{4-2} operator for $\delta = -1/2$ (purely imaginary case). From left to right the plots illustrate the results obtained with a grid containing 20, 60, 100 points, respectively. It is clear from the figures that these correspond, as in figure 2, to a purely imaginary case.

3.1.2. Fourth order in the interior, second order at and close to boundaries (D_{4-2} scheme). Figure 4 shows the equivalent of figure 2, but now for the D_{4-2} case. The maximum is slightly larger than the corresponding one for the D_{2-1} case: approximately 1.936.

Figure 5, in turn, shows the equivalent of figure 3 for the current case. As before, a negative real part appears and the maximum in the imaginary axis slightly decreases (in this case to roughly 1.371, not changing much among these three values of δ).

3.1.3. Sixth order in the interior, third order at and close to boundaries, minimum bandwidth case (D_{6-3} scheme). Figure 6 illustrates the equivalent of figures 2 and 4 for the D_{6-3} case. The maximum is roughly 2.129, slightly larger than those of the previous two cases. The behaviour for larger values of δ is similar to that found in the previous two cases, as

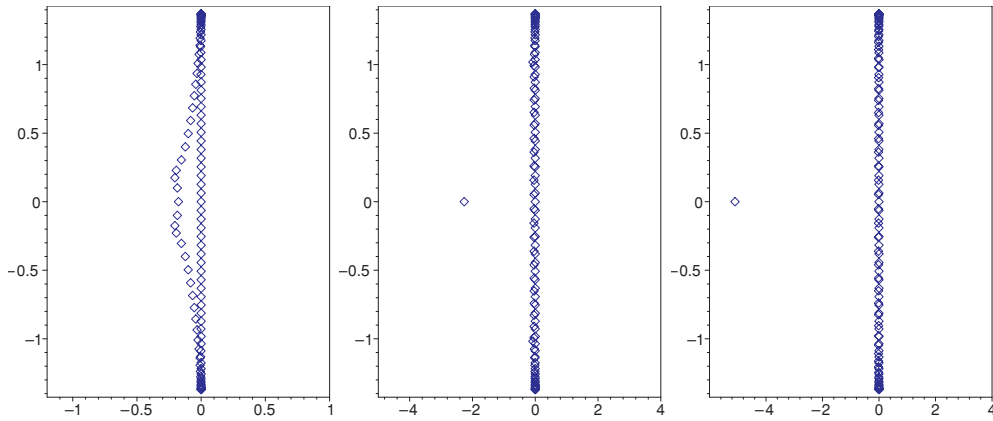


Figure 5. Eigenvalues corresponding to the D_{4-2} operator, obtained with a grid containing 100 points. From left to right the plots illustrate the behaviour for $\delta = 0, 1/10, 1/2$, respectively. As δ becomes larger, a larger (in magnitude) negative eigenvalue on the real axis is observed (note the left-most diamond at $y \simeq -2.5, -5$ in the middle and right-hand plots, respectively).

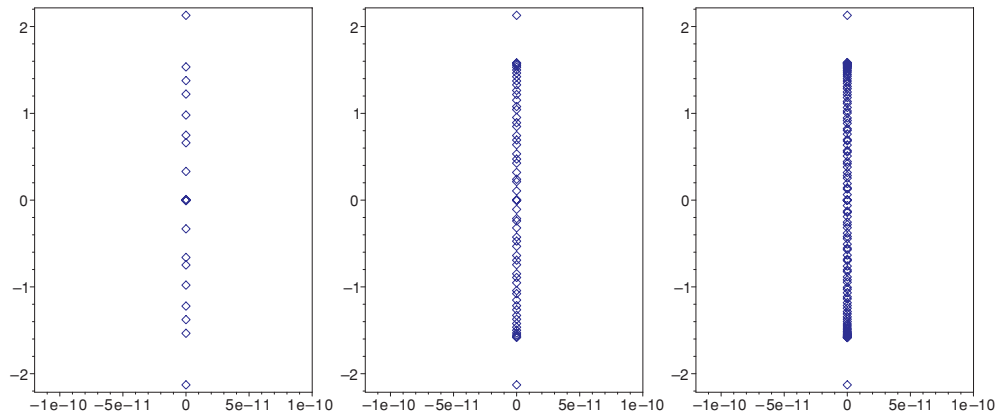


Figure 6. Numerically obtained eigenvalues corresponding to the minimum bandwidth D_{6-3} operator for $\delta = -1/2$ (purely imaginary case). From left to right the plots illustrate the results obtained with a grid containing 20, 60, 100, points respectively. As in figures 2 and 4, these correspond to a purely imaginary case.

seen in figure 7. The maximum in the imaginary axis again decreases slightly compared to the $\delta = -1/2$ case (to roughly 1.585) and does not change much among these three values of δ .

3.1.4. Eight order in the interior, fourth order at and close to boundaries (D_{8-4} scheme).

The D_{8-4} operator has three free parameters, denoted as x_1, x_2, x_3 both in [9] and here. As mentioned, these parameters can be freely chosen to satisfy a given criterion. For instance, they can be fixed so as to minimize the width of the derivative operator or yield as small a spectral radius as possible. As we discuss next, these options can yield operators with significantly different stability requirements as dictated by the CFL condition.

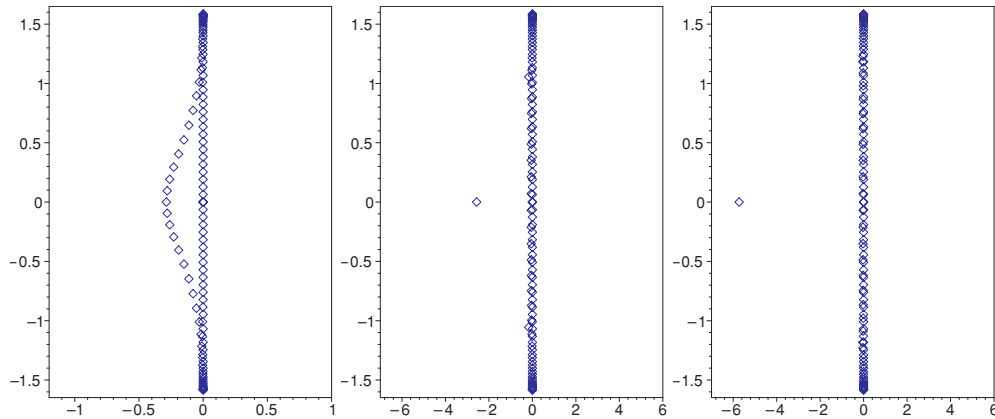


Figure 7. Eigenvalues corresponding to the D_{6-3} operator, obtained with a grid containing 100 points. From left to right the plots illustrate the behaviour for $\delta = 0, 1/10, 1/2$, respectively. As δ becomes larger, a larger (in magnitude) negative eigenvalue on the real axis is observed (note the left-most diamond at $y \simeq -2.5, -6$ in the middle and right-hand plots, respectively).

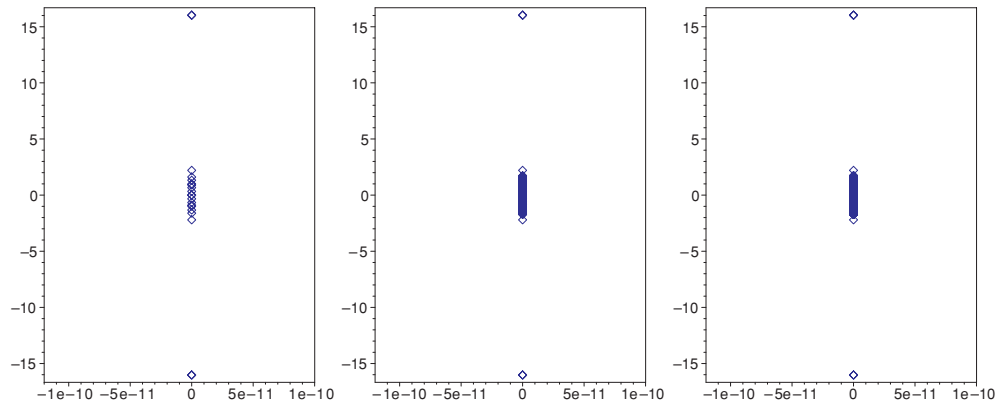


Figure 8. Numerically obtained eigenvalues corresponding to the minimum bandwidth D_{8-4} operator for $\delta = -1/2$ (purely imaginary case). From left to right the plots illustrate the results obtained with a grid containing 20, 60, 100 points, respectively. Although purely imaginary, the maximum (absolute) value in the vertical axis is approximately 16.

Minimum bandwidth operator. The minimum bandwidth case corresponds to the choice (see [9])

$$x_1 = \frac{1714\,837}{4354\,560}, \quad x_2 = -\frac{1022\,551}{30\,481\,920}, \quad x_3 = \frac{6445\,687}{8709\,120}. \quad (24)$$

Figure 6 shows for this minimum D_{8-4} bandwidth case the eigenvalues for 20, 60, 100 points, for $\delta = -1/2$ (purely imaginary case). While for the previous operators we have seen that the maximum eigenvalue increases slightly with the order of the operator, in this case the increase is quite large: the maximum is roughly 16.04. This translates into a CFL limit for this operator being almost an order of magnitude smaller than the limits for the previous operators. Additionally, variation of δ does not significantly affect this behaviour, as shown in figure 9. That figure shows the eigenvalues computed with 100 points and $\delta = 0, 1/10, 1/2$.

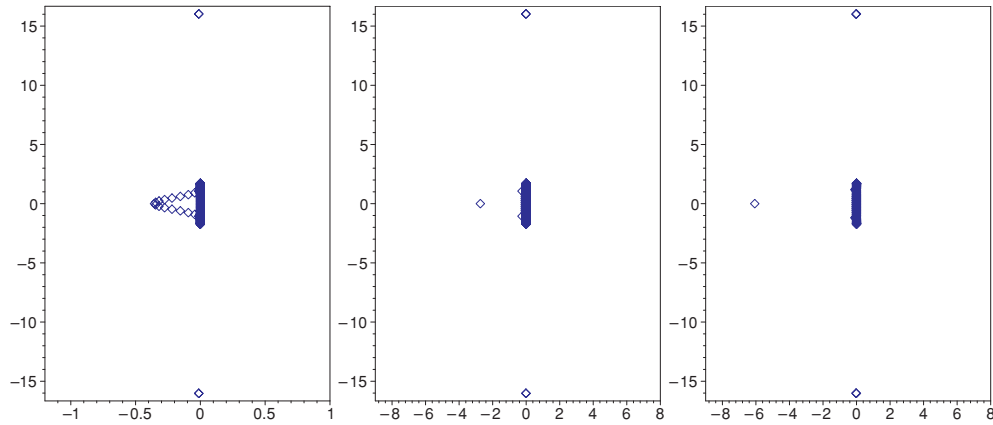


Figure 9. Eigenvalues for the minimum bandwidth D_{8-4} operator with $\delta = 0, 1/10, 1/2$ (from left to right) and 100 points. Values of δ larger than $-1/2$ introduce a negative real part in the spectrum but they have little effect on the maximum absolute value, which remains at approximately 16.

The qualitative behaviour when increasing δ is similar to that of the previous cases. A negative real part appears in the spectrum, and the maximum in the imaginary axis slightly decreases, to roughly 16.02, not varying much among these three illustrative values of δ . Such a large spectral radius for this operator motivates the search for another one, with a more convenient radius at the expense of not having the minimum possible bandwidth.

Optimized operator. We construct here an ‘optimized D_{8-4} operator’ (which we shall use from here on in the D_{8-4} case) in the sense that it has a spectral radius considerably smaller than that defined by equation (24). More precisely, through a numerical search in the three-parameter space, we have found that the following values

$$x_1 = 0.541, \quad x_2 = -0.0675, \quad x_3 = 0.748 \quad (25)$$

yield an operator whose maximum absolute eigenvalue in the purely imaginary case ($\delta = -1/2$) is

$$\lambda_{\max} = 2.242. \quad (26)$$

This maximum eigenvalue appears to be quite sensitive to these parameters. For example, truncating the above values to two significant digits,

$$x_1 = 0.54, \quad x_2 = -0.067, \quad x_3 = 0.75,$$

gives $\lambda_{\max} = 2.698$ and truncating even more, to just one digit,

$$x_1 = 0.5, \quad x_2 = -0.07, \quad x_3 = 0.7,$$

gives the large value $\lambda_{\max} = 71.76$. On the other hand, refining in the parameter search the values in equation (25) in one more digit did not change the maximum of equation (26) in its four digits shown here. The eigenvalues for $\delta = -1/2$ for this optimized D_{8-4} operator, given by the parameters of equation (25), are shown in figure 10, while figure 11 shows them for $\delta = 0, 1/10, 1/2$ and 100 points. As before, a negative real component appears and the maximum in the imaginary axis decreases (to around 1.754).

While completing this work we became aware of similar work by Svard, Mattson and Nordström [13], who constructed an optimized operator with different parameters by minimizing the spectral radius of the derivative itself (rather than that of the amplification

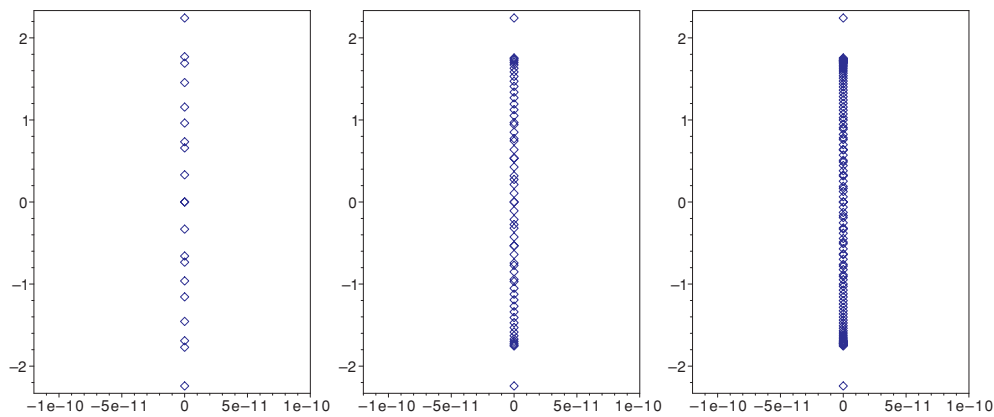


Figure 10. Eigenvalues for the optimized D_{8-4} operator with $\delta = -1/2$ (purely imaginary case) and 20, 60, 100 points (from left to right). Clearly, this modified operator has a much smaller spectral radius, compared to the minimum bandwidth one.

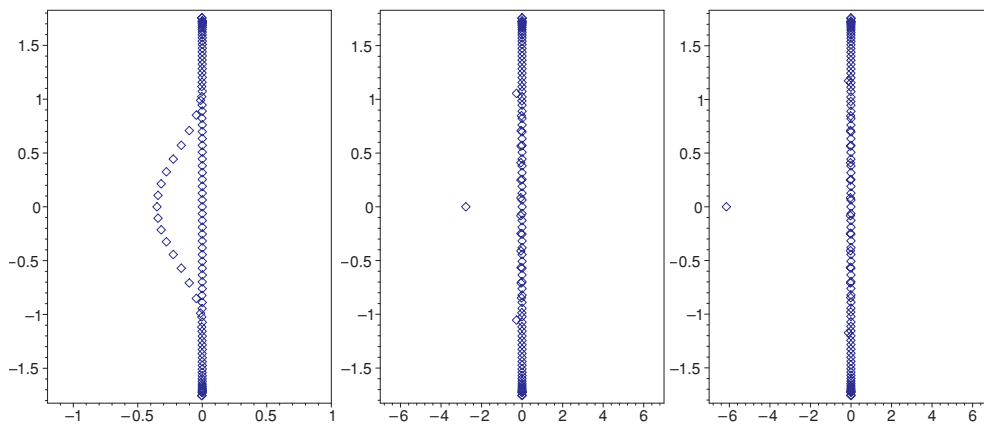


Figure 11. Eigenvalues for the optimized D_{8-4} operator, with 100 points and $\delta = 0, 1/10, 1/2$ (from left to right).

matrix of a toy problem with an interface, as in our case), obtaining $x_1 = 0.649$, $x_2 = -0.104$, $x_3 = 0.755$. When using these parameters in our toy problem with an interface, the resulting spectral radius (for 20 grid points) is $\lambda_{\max} = 2.241\,259$, while for the parameters we chose (cf equation (25)) $\lambda_{\max} = 2.241\,612$ [14].

3.2. Global convergence rate

In general, the global (say, in an L_2 norm) convergence factor for these operators will be dominated by the lower order at and close to boundaries. However, it is sometimes found that round-off values for the error in such a global norm are reached before this happens, and the convergence factor is different from the one expected from the boundary terms. The precise value is found to actually depend on the function being differentiated and whether one reaches round-off level. To illustrate the expected behaviour in a generic case, we consider

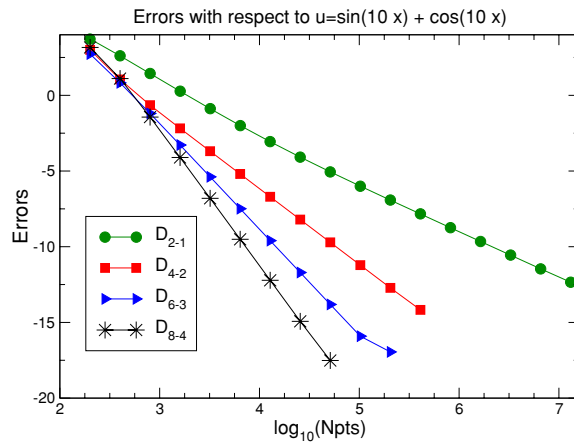


Figure 12. Logarithm (base 10) of the L_2 norms of the errors obtained when taking the discrete derivative of $\sin(10x) + \cos(10x)$ and comparing it with the analytical answer, using D_{2-1} , D_{4-2} , D_{6-3} , D_{8-4} operators, versus the number of grid points.

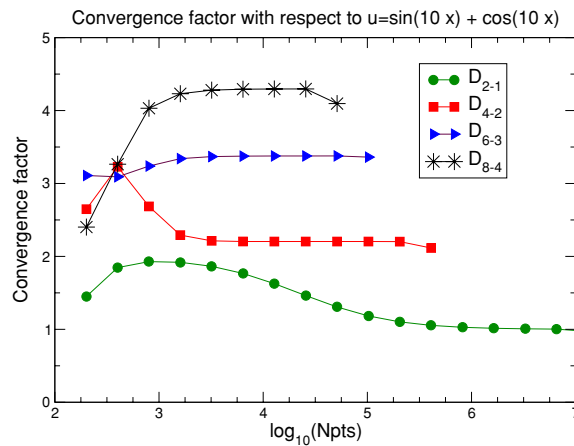


Figure 13. Convergence factors for the curves of figure 12. As in this case, the convergence in general will be dictated by the lower order the derivative operators have at and close to the boundaries. Note that the lines corresponding to the different operators terminate at sequentially fewer points. This is due to the corresponding errors reaching round-off levels, after which the convergence factor calculation ceases to have a sensible meaning.

the function $\sin(10x) + \cos(10x)$ in the domain $x \in [0, 2\pi]$.⁹ Figure 12 shows the error (with respect to the exact solution) when computing the discrete derivative versus the number of grid points, for the difference operators D_{2-1} , D_{4-2} , D_{6-3} and D_{8-4} . The errors in the L_2 norm are shown until round-off values are reached (further increasing the number of grid points causes the error to grow with the number of points involved). Figure 13, in turn, shows the convergence factors obtained.

⁹ This function is chosen (as opposed to, for example, $\cos(x)$ or $\sin(x)$) so as to ensure that relatively high frequency modes are present and no derivatives evaluate to zero at boundaries. The latter is important to examine the influence that the operator's lower order convergence at (and near) boundary points has.

3.3. Group speed

We now turn our attention to the group speed that different discrete modes have when the above considered operators are used. To simplify the discussion, we actually restrict ourselves to the periodic case, which lends itself to a clean analytical calculation. In this case, the operators of orders 2, 4, 6 and 8 satisfying SBP are the standard centred ones (D_0 , D_+ , D_- denote the standard centred second-order, and forward and backward first-order operators, respectively):

$$D^{(2)} = D_0, \quad (27)$$

$$D^{(4)} = D_0(I - h^2/6D_+D_-), \quad (28)$$

$$D^{(6)} = D_0(I - h^2/6D_+D_- + h^4/30D_+^2D_-^2), \quad (29)$$

$$D^{(8)} = D_0(I - h^2/6D_+D_- + h^4/30D_+^2D_-^2 - h^6/140D_+^3D_-^3). \quad (30)$$

In discrete Fourier space, the eigenvalues for these operators are, respectively,

$$\lambda_2 = i \sin(\zeta)/\zeta, \quad (31)$$

$$\lambda_4 = i \sin(\zeta)/\zeta(1 + 2 \sin(\zeta/2)/3), \quad (32)$$

$$\lambda_6 = i \sin(\zeta)/\zeta(1 + 2 \sin(\zeta/2)/3 + 8 \sin^4(\zeta/2)/15), \quad (33)$$

$$\lambda_8 = i \sin(\zeta)/\zeta(1 + 2 \sin(\zeta/2)/3 + 8 \sin^4(\zeta/2)/15 + 64 \sin^6(\zeta/2)/140), \quad (34)$$

where $\zeta = \omega h$ and ω is the associated wave number.

The highest possible frequency is $\omega = N/2$ (with N the number of grid points). For that frequency, $\zeta = \pi$ and the above eigenvalues are all zero. Therefore, the mode with the highest possible frequency for a given number of points does not propagate. Furthermore, if one examines the *group speed*,

$$v_g = \frac{d(\lambda\omega)}{d\omega}, \quad (35)$$

one finds that at $\zeta = \pi$ this speed is

$$v_g^2 = -1, \quad (36)$$

$$v_g^4 = -5/3 \approx -1.6, \quad (37)$$

$$v_g^6 = -33/15 \approx -2.2, \quad (38)$$

$$v_g^8 = -341/135 \approx -2.6. \quad (39)$$

Thus, for higher (than two) order operators, the velocity of this mode is higher than the continuum one (which is 1). But, more importantly, in all cases the speed has the opposite sign. Of course, this effect goes away with resolution, since the highest possible frequency moves to larger values as resolution is increased, but, still, is an effect to be taken into account. For instance, if noise is produced at an interface, it propagates backwards and with higher speed. Even though this effect is typically very small, it might be noticeable in highly accurate simulations or in simulations in which the solution itself decays to very small values (see section 4.3.2). This could also be a source of difficulties in the presence of black holes—or for this matter any system where some propagation speed changes sign—since the event horizon traps these high-frequency modes in a very narrow region and then releases them as low-frequency ones. We have observed this in some highly resolved one-dimensional simulations and explained an observed convergence drop which goes away when numerical dissipation is turned on.

4. Tests

In this section, we illustrate the behaviour of the aforementioned penalty technique, together with the choice of different derivative operators. We present tests in one, two and three dimensions. In particular, we implement the linearized Einstein equations (off a ‘gauge-wave’ spacetime [15, 16]) and propagation of scalar fields in black hole backgrounds. The former is cast in a way which yields a one-dimensional symmetric hyperbolic system with coefficients depending both in space *and* in time, while the latter provides a hyperbolic system of equations with space-varying coefficients and sets a conforming grid for spherical black hole excision.

Throughout this paper, we employ a fourth-order accurate Runge–Kutta time integrator. In a number of tests aimed at examining the behaviour of high-order operators, we adopt a sufficiently small time step Δt so that the time integrator does not play a role. Thus, we either choose a suitably small CFL factor or scale the time step quadratically with the grid spacing h .

4.1. One-dimensional simulations: linearizations around a gauge wave

As a first test we evolve Einstein’s equations in one dimension, linearized around a background given by

$$ds^2 = e^{A \sin(\pi(x-t))} (-dt^2 + dx^2) + dy^2 + dz^2. \quad (40)$$

This background describes flat spacetime with a sinusoidal coordinate dependence, of amplitude A , along the x direction. One of the interesting features of this testbed is that while a linear problem, the coefficients in the equations to solve are not only space but also time dependent.

The non-trivial variables for this metric are

$$\hat{g}_{xx} = e^{A \sin(\pi(x-t))}, \quad (41)$$

$$\hat{K}_{xx} = \frac{A}{2} \pi \cos(\pi(x-t)) e^{A/2 \sin(\pi(x-t))}, \quad (42)$$

$$\hat{\alpha} = e^{A/2 \sin(\pi(x-t))}, \quad (43)$$

$$\hat{\beta}^i = 0. \quad (44)$$

We evolve the linearized Einstein equations using the symmetric hyperbolic formulation presented in [17] with a dynamical lapse given by the homogeneous time-harmonic condition (defined by requiring $\square t = 0$). The formulation is cast in first-order form by introducing the variables $\mathcal{A}_x := \partial_x \alpha / \alpha$ and $d_{xxx} := \partial_x g_{xx}$. The equations determining the dynamics of the (first-order) perturbations, which we assume to depend solely on (t, x) , are obtained by considering linear deviations of a background metric given by equation (40). That is, we consider

$$\begin{aligned} g_{xx} &= \hat{g}_{xx} + \delta g_{xx}, & K_{xx} &= \hat{K}_{xx} + \delta K_{xx}, & d_{xxx} &= \hat{d}_{xxx} + \delta d_{xxx}, \\ \alpha &= \hat{\alpha} + \delta \alpha, & \mathcal{A}_x &= \hat{\mathcal{A}}_x + \delta \mathcal{A}_x, \end{aligned}$$

replace these expressions in Einstein’s equations and retain only first-order terms. The resulting equations are (henceforth dropping the δ notation)

$$\dot{\alpha} = -A\pi \cos(\phi)\alpha - K_{xx} + \frac{A\pi}{2\hat{\alpha}} \cos(\phi)g_{xx}, \quad (45)$$

$$\begin{aligned}\dot{\mathcal{A}}_x = & -\frac{1}{\hat{\alpha}}\partial_x K_{xx} + \frac{A\pi}{2\hat{\alpha}}\cos(\phi)K_{xx} - \frac{A\pi^2}{2\hat{\alpha}^2}(A\cos(\phi)^2 + \sin(\phi))g_{xx} \\ & + \frac{A\pi}{2\hat{\alpha}^2}\cos(\phi)d_{xxx} - \frac{A\pi}{2}\cos(\phi)\mathcal{A}_x + \frac{A\pi^2}{2\hat{\alpha}}\sin(\phi)\alpha,\end{aligned}\quad (46)$$

$$\dot{g}_{xx} = -A\hat{\alpha}\pi\cos(\phi)\alpha - 2\hat{\alpha}K_{xx}, \quad (47)$$

$$\begin{aligned}\dot{K}_{xx} = & -\hat{\alpha}\partial_x \mathcal{A}_x - \frac{A\hat{\alpha}\pi}{2}\cos(\phi)\mathcal{A}_x - \frac{A\pi^2}{4}(-2\sin(\phi) + A\cos(\phi)^2)\alpha \\ & - A\pi\cos(\phi)K_{xx} + \frac{A\pi}{4\hat{\alpha}}\cos(\phi)d_{xxx},\end{aligned}\quad (48)$$

$$\begin{aligned}\dot{d}_{xxx} = & A\hat{\alpha}\pi^2(\sin(\phi) - A\cos(\phi)^2)\alpha \\ & - A\hat{\alpha}\pi\cos(\phi)K_{xx} - A\hat{\alpha}^2\pi\cos(\phi)\mathcal{A}_x - 2\hat{\alpha}\partial_x K_{xx},\end{aligned}\quad (49)$$

where we have defined $\phi := \pi(x - t)$. This system is symmetric hyperbolic and the symmetrizer used to define the energy can be chosen so that the characteristic speeds which play a role in the energy estimate are 0 and ± 1 .

We consider here a periodic initial boundary-value problem on the domain $x \in [-1/2, 3/2]$, where periodic boundary conditions at $x = -1/2, 3/2$ are implemented through an interface with penalty terms, as described in section 2.

The system must satisfy two non-trivial constraint equations, corresponding to the definition of the variables d_{xxx} and \mathcal{A}_x (the linearized physical constraints are automatically satisfied by the considered ansatz). When linearized, these constraints are

$$0 = C_x = -\partial_x g_{xx} + d_{xxx}, \quad (50)$$

$$0 = C_{\mathcal{A}} = \mathcal{A}_x - \frac{1}{\hat{\alpha}}\left(\partial_x \alpha - \frac{A\pi}{2}\cos(\phi)\alpha\right). \quad (51)$$

In the first series of simulations, we adopt a CFL factor $\lambda = 10^{-3}$ ¹⁰ and consider relatively short evolutions corresponding to four crossing times. The D_{8-4} derivative is used, and dissipation is added through the dissipative operator constructed from $-\sigma h^7 D_+^4 D_-^4$, suitably modified at boundaries as explained in appendix B so as to make it non-positive definite with respect to the appropriate scalar product. Thus, the use of this dissipative operator does *decrease* the order of the spatial discretization by 1. The dissipation parameter used is $\sigma = 5 \times 10^{-4}$. Figure 14 exemplifies the behaviour observed in the convergence of the field K_{xx} (the other fields behave similarly). As time progresses, the convergence order obtained oscillates in a way that is consistent with the accuracy obtained at interior and boundary points.

Next, we adopt as a starting value for the CFL factor defined at the coarsest grid to be $\lambda = 0.2$ but in subsequent grids (refined by a factor of 2) we adopt $\lambda = 0.2/2^n$ with $n = 1, \dots, 3$. Figure 15 illustrates the behaviour observed; again, as time progresses the convergence order obtained oscillates inbetween the order of accuracy of interior and boundary points, with the additional effect of accuracy loss due to the accumulation of error as time progresses.

Finally, we compare the above results with those obtained in the ‘truly periodic case’, i.e. when periodicity is used explicitly to employ the same derivative operator at all points. We again consider cases where a sufficiently small CFL factor ($=10^{-3}$) is used or the time step

¹⁰ We do not claim that this is the largest possible CFL factor for which the errors due to time integration are negligible compared to the spatial ones, though. Actually, as discussed later, in some cases one can use larger CFL factors.

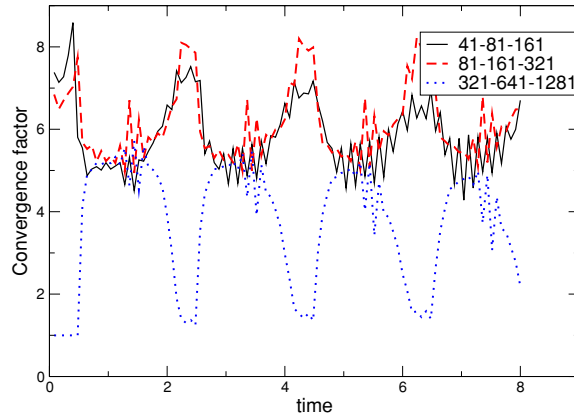


Figure 14. Evolutions of 1D linearized Einstein equations in a periodic domain, with periodicity enforced through an interface with penalty terms. Shown is the convergence factor for K_{xx} when using the D_{8-4} derivative, CFL factor $\lambda = 10^{-3}$ and dissipation $\sigma = 5 \times 10^{-4}$. While the convergence factors obtained with 41–321 points oscillate between the expected order at the boundary and that at the interior, the ones calculated with 321–1281 points are not meaningful when the pulse is located at interior points as the round-off level is reached.

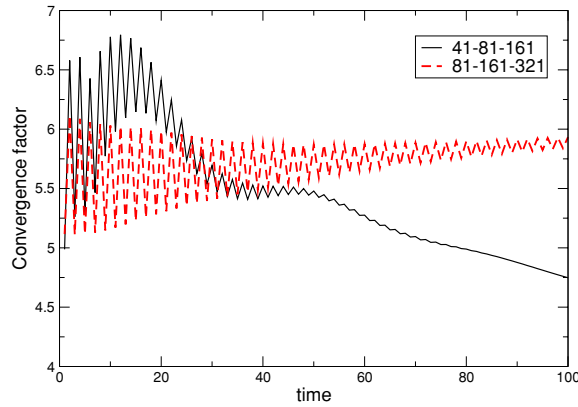


Figure 15. Same as previous figure, but with a decreasing CFL factor given by $\lambda = 0.2/2^n$ with $n = 1, \dots, 3$.

is scaled quadratically. Figures 16 illustrates the observed convergence rate for the field g_{xx} . As above, dissipation is added through a seventh-order dissipative operator (but now with no modification at boundaries needed) with same dissipative parameter: $\sigma = 5 \times 10^{-4}$. While the errors remain above round-off level, the observed convergence rate is consistent with the expected one of 7, as the orders of the derivative and dissipative operators employed are 8 and 7, respectively. Certainly, dissipation of higher order could have been introduced by simply employing the KO style operator $h^9(D_-D_+)^5$, but we have adopted this one to more directly compare with the case with interface boundaries. For the highest resolutions, the errors reach round-off level and the obtained convergence factors yield non-sensible values.

As an illustration of what is observed with other derivatives, we briefly discuss some simulations using the D_{6-3} operator and a fixed CFL factor (given, as before, by $\lambda = 10^{-3}$). Analogously to what was done above, dissipation is added here by extending—as discussed in appendix B—the operator $\sigma h^7 D_-^4 D_+^4$ at and near boundaries in order to make it non-positive

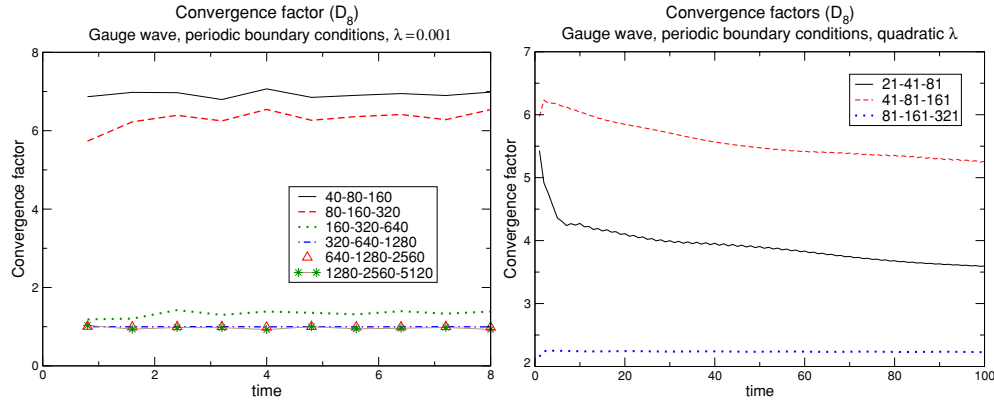


Figure 16. This figure shows evolutions similar to those of figures 14 and 15, the only difference being that periodicity is enforced here explicitly. The convergence factors for the metric component g_{xx} are shown.

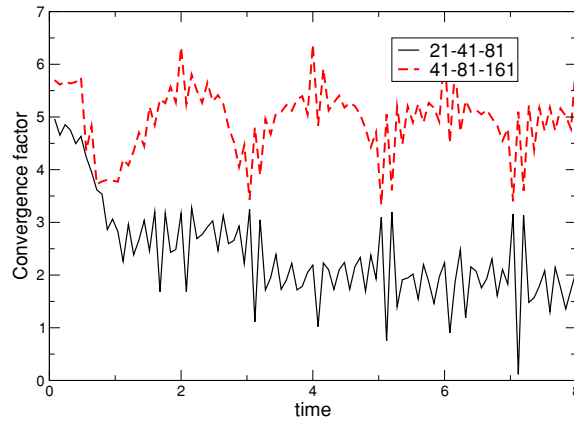


Figure 17. Similar to figure 14, but using the D_{6-3} derivative. The convergence factor for K_{xx} is shown.

definite with respect to the appropriate scalar product; a dissipation parameter $\sigma = 10^{-3}$ is used. The observed results are illustrated in figure 17, which shows the self-convergence factor for K_{xx} . As before, it oscillates between the order of the scheme in the interior and that at boundary points.

Summarizing, the presented results indicate that, in the case where boundaries are present, the worst-case scenario—as far as the expected convergence rate relates—is determined by the accuracy order at and close to boundary points.

4.1.1. Two- and three-dimensional simulations: problem set-up. In this section, we solve the wave equation for a scalar field ϕ propagating on a fixed background,

$$\nabla^a \nabla_a \phi = 0,$$

where ∇ is the covariant derivative associated with the metric of the background. We will consider two backgrounds: a 2D one consisting of the unit sphere with its standard metric and a 3D one consisting of a rotating Kerr black hole background.

We start by describing in more detail the equations solved and the multiple coordinate system used, and then present the actual results of the simulations.

4.1.2. A strictly stable scheme for the wave equation in a time-independent, curved spacetime. The wave equation in a time-independent background can be written, using any coordinates in which the metric is manifestly time independent, as

$$\dot{\phi} = \alpha \Pi, \quad (52)$$

$$\dot{\Pi} = \beta^i \alpha^{-1} D_i(\alpha \Pi) + h^{-1/2} D_i(h^{1/2} \beta^i \Pi + \alpha h^{1/2} H^{ij} d_j), \quad (53)$$

$$\dot{d}_i = D_i(\alpha \pi), \quad (54)$$

where $H^{ij} := h^{ij} - \alpha^{-2} \beta^i \beta^j$, h^{ij} is the inverse of the three-metric, $h = \det(h_{ij})$, α is the lapse and β^i the shift vector. The advantage of writing the equations in this way is that one can show that if D is any difference operator satisfying SBP, this form of the equations guarantees that the semi-discrete version of the physical energy is a non-increasing function of time. When the Killing field is timelike this means that there is a norm in which the solution is bounded for all times, thus suppressing artificial fast growing modes without the need of artificial dissipation (see [18] for details).

We now look at the characteristic variables and characteristic speeds with respect to a ‘coordinate’ observer. That is, the eigenfields and eigenvalues of the symbol $A^i n_i$, where A^i denotes the principal part of the evolution equations and n_i the normal to the boundary¹¹. The characteristic variables with non-zero speeds

$$\lambda^\pm = (\pm\alpha + \beta^k \hat{n}_k)(h^{ij} n_i n_j)^{1/2}$$

(where $\hat{n}_k = n_k(h^{ij} n_i n_j)^{-1/2}$) are

$$v^\pm = \lambda^\pm \Pi + \alpha H^{ij} \hat{n}_i d_j,$$

while the zero-speed modes are

$$v_i^0 = d_i - \hat{n}_i d_j \hat{n}^j.$$

4.1.3. Cubed-sphere coordinates. The topology of the computational domain in our 2D simulations is S^2 , the unit sphere, while in our 3D ones it is $S^2 \times R^+$. Since it is not possible to cover the sphere with a single system of coordinates which is regular everywhere, we employ multiple patches to cover it. A convenient set of patches is defined by the *cubed-sphere coordinates*, defined as follows (for a related definition, see for instance [19]).

Each patch uses coordinates a, b, c , where $c = \sqrt{x^2 + y^2 + z^2}$, the standard radial coordinate, is the same for the six patches (x, y, z are standard Cartesian coordinates). The other two coordinates, a, b , are defined as

- patch 0 (neighbourhood of $x = 1$): $a = z/x, b = y/x$,
- patch 1 (neighbourhood of $y = 1$): $a = z/y, b = -x/y$,
- patch 2 (neighbourhood of $x = -1$): $a = -z/x, b = y/x$,
- patch 3 (neighbourhood of $y = -1$): $a = -z/y, b = -x/y$,
- patch 4 (neighbourhood of $z = 1$): $a = -x/z, b = y/z$,
- patch 5 (neighbourhood of $z = -1$): $a = -x/z, b = -y/z$.

¹¹ If the boundary is aligned with the coordinate lines, as is indeed the case here, n_i would be either $(1, 0, 0)$, $(0, 1, 0)$ or $(0, 0, 1)$.

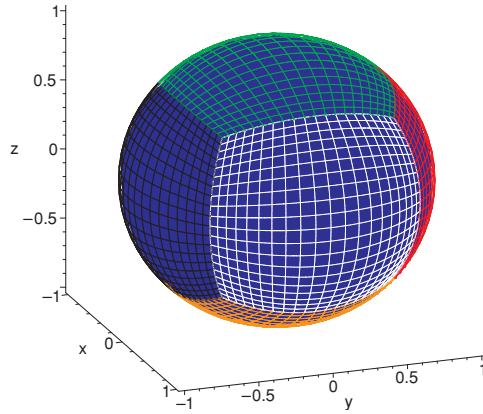


Figure 18. Cubed-sphere coordinates for S^2 .

Similarly, the inverse transformation is

- patch 0: $x = c/D$, $y = cb/D$, $z = ac/D$,
- patch 1: $x = -bc/D$, $y = c/D$, $z = ac/D$,
- patch 2: $x = -c/D$, $y = -cb/D$, $z = ac/D$,
- patch 3: $x = bc/D$, $y = -c/D$, $z = ac/D$,
- patch 4: $x = -ac/D$, $y = cb/D$, $z = c/D$,
- patch 5: $x = ac/D$, $y = cb/D$, $z = -c/D$,

with $D := \sqrt{1 + a^2 + b^2}$. This provides a relatively simple multi-block structure for S^2 which can be exploited to implement the penalty technique in a straightforward manner. Each patch is discretized with a uniform grid in the coordinates a and b , and the requirement of boundary points coinciding in neighbouring grids is indeed satisfied. Figure 18 shows this grid structure for 20×20 points on each patch.

4.2. 2D simulations

We now discuss simulations of the wave equation on the unit sphere in cubed-sphere coordinates, written in strictly stable first-order form (equations (52)–(54)). The metric used, therefore, is flat spacetime projected to the $r = 1$ slice, which in local coordinates is

$$ds^2 = -dt^2 + D^{-4}[(1 + b^2) da^2 + (1 + a^2) db^2 - 2ab da db],$$

where $D := \sqrt{1 + a^2 + b^2}$.

Figure 19 shows simulations using the D_{4-2} derivative and its associated dissipative operator constructed in appendix B, which we call KO6, using $n \times n$ points on each of the six patches, where $n = 41, 81, 161, 321$. The initial data for Π correspond to a pure $l = 2, m = 1$ spherical harmonic. The CFL factor used is $\lambda = 0.125$ and for each set of runs two values of dissipation are used: $\sigma = 10^{-2}$ and $\sigma = 10^{-3}$. As can be seen from the figure, the self-convergence factor obtained with these resolutions is above the lower value (2) expected from the order at the interfaces. The reason for the lower order at the interfaces not dominating is likely due to the fact that the initial data are an eigenmode of the Laplacian operator, and the solution at the continuum is just an oscillation in time of these initial data, without propagation across the interfaces. Indeed, the oscillations in the convergence factors in figure 19 appear when the numerical solution goes through zero, and the frequency at which this happens coincides approximately with the expected frequency at the continuum for this mode.

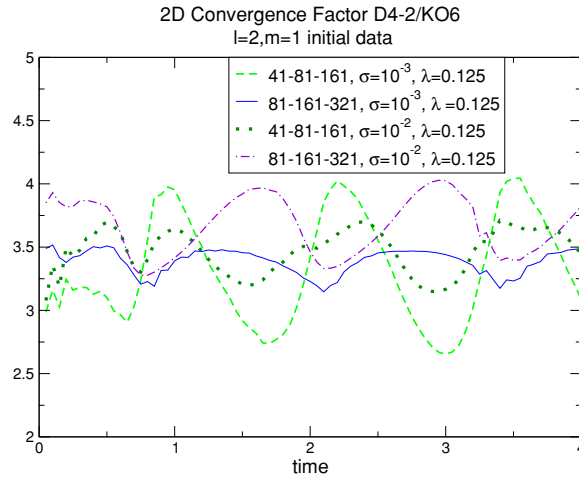


Figure 19. Evolutions of the wave equation on the unit sphere, using cubed-sphere coordinates and a pure $l = 2, m = 1$ spherical harmonic as initial data. Shown is the convergence factor when the D_{4-2} derivative and the KO6 dissipation operators are used.

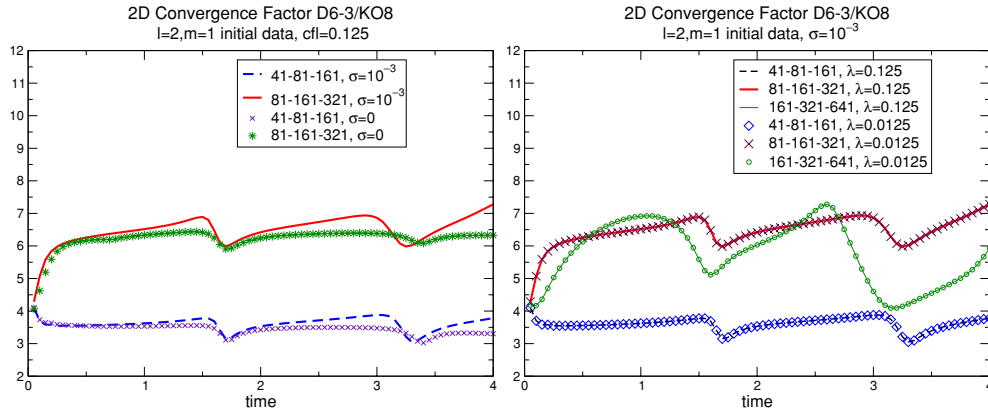


Figure 20. Same evolution equation and initial data as those used in figure 19, except that now the D_{6-3} derivative and KO8 dissipation are used.

The same initial data are now evolved with the D_{6-3} derivative and KO8 dissipation (again, see appendix B) and the results are shown in figure 20. As before, $\lambda = 0.125$ and $n = 41, 81, 161, 321$ points are used, but the values of dissipation shown are now $\sigma = 0$ (i.e., no dissipation) and $\sigma = 10^{-3}$. At the same resolutions, there is a small difference in the obtained convergence factors, depending on the value of σ , but with both values of this parameter the order of convergence is higher than the lower one expected from the time integrator if this one dominated. Figure 20 also presents a comparison made with a smaller CFL factor: $\lambda = 0.0125$, keeping the dissipation at $\sigma = 10^{-3}$. One more resolution is used (641 points) to look for differences between the solutions obtained with the two CFL factors, but they do not appear. This seems to suggest that at least in this case, and for these resolutions, it is not necessary to use too small a CFL factor in order to avoid the time integrator's lowest order to dominate over the higher spatial discretization (see figure 23 for another instance

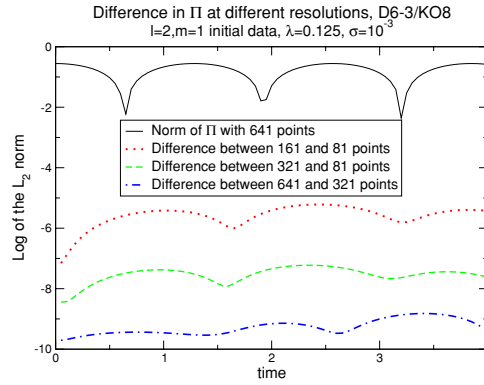


Figure 21. L_2 norms of the differences between the solutions at different resolutions, for the simulations of figure 20 with $\lambda = 0.0125$ and $\sigma = 10^{-3}$.

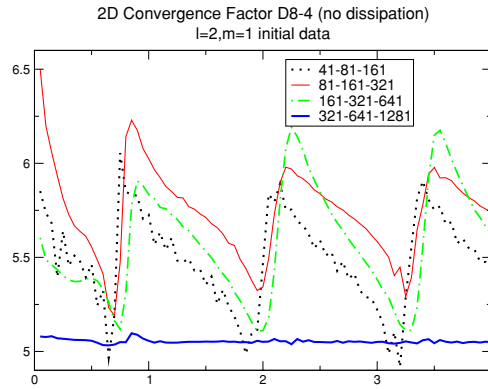


Figure 22. Simulations of the same equation and initial data as those of the previous figures, except that now the higher order, D_{8-4} derivative is used and no dissipation has been added.

where this happens). It is also worth pointing out that the difference between the two highest resolutions is not quite at round-off level, but it is rather small (of the order of 10^{-9} if scaled by the amplitude of the initial data), as shown in figure 21. That figure shows the L_2 norm of the differences between the solutions at different resolutions, for the simulations of figure 20 with $\lambda = 0.0125$ and $\sigma = 10^{-3}$.

Finally, figure 22 shows evolutions of the same initial data, with the D_{8-4} derivative and no dissipation. The CFL factor is decreased when resolution is increased, much as in section 4.1, so that the order of the time integration does not dominate over the higher one of the spatial discretization. That is, for the resolutions shown we used $\lambda = 0.25, 0.125, 0.0625, 0.03125, 0.015625$. The convergence factor obtained is also higher than that expected from the lower order at the interfaces (presumably for the same reason as before, the initial data adopted) and higher than that of the time integration.

4.3. Three-dimensional simulations

In this application, we consider fields propagating on a Kerr black hole background, as governed by equations (52)–(54), with the background metric written in Kerr–Schild form

and cubed-sphere coordinates used for the angular directions. Homogeneous maximally dissipative boundary conditions are used at the outer boundary, while no condition is needed at the inner one if it is *appropriately* placed inside the black hole so that it constitutes a purely outflow surface.

The Kerr–Schild metric in cubed-sphere coordinates. The Kerr metric in Kerr–Schild form is

$$ds^2 = \eta_{\mu\nu} + 2Hl_\mu l_\nu dx^\mu dx^\nu,$$

where $\eta_{\mu\nu}$ is the flat metric (with signature $(-, +, +, +)$),

$$H = \frac{mr}{r^2 + A^2 \cos^2 \theta}, \quad (55)$$

$$r^2 = \frac{1}{2}(\rho^2 - A^2) + \sqrt{\frac{1}{4}(\rho^2 - A^2)^2 + A^2 z^2}, \quad (56)$$

$$\rho^2 = x^2 + y^2 + z^2 \quad (57)$$

and l^μ is a null vector (with respect to both the flat metric and the whole metric).

Therefore, in order to write the above metric in cubed-sphere coordinates one needs to write $\eta_{\mu\nu}$ and l_μ in these coordinates. A straightforward change of coordinates of the first one gives

$$\eta_{\mu\nu} = -dt^2 + dc^2 + c^2 D^{-4}[(1 + b^2) da^2 + (1 + a^2) db^2 - 2ab da db],$$

with $D := \sqrt{(1 + a^2 + b^2)}$.

In Cartesian coordinates, the l_μ co-vector is, in turn,

$$l \equiv l_\mu dx^\mu = dt + \frac{rx + Ay}{r^2 + A^2} dx + \frac{ry - Ax}{r^2 + A^2} dy + \frac{z}{r} dz,$$

with $x^\mu = (t, x, y, z)$, which when changed to cubed-sphere coordinates (t, a, b, c) gives

$$l = dt + \frac{-c^2 a A^2 (a^2 - D^2)}{D^4 r (r^2 + A^2)} da + \frac{-c^2 A (D^2 r + A b a^2)}{D^4 r (r^2 + A^2)} db + \frac{c (D^2 r^2 + a^2 A^2)}{D^2 r (r^2 + A^2)} dc \quad \text{for patches 0–3}, \quad (58)$$

$$l = dt + \frac{-c^2 A (D^2 r b + a A)}{D^4 r (r^2 + A^2)} da + \frac{c^2 A (r a D^2 - A b)}{D^4 r (r^2 + A^2)} db + \frac{c (D^2 r^2 + A^2)}{D^2 r (r^2 + A^2)} dc \quad \text{for patch 4}, \quad (59)$$

$$l = dt + \frac{-c^2 A (-D^2 r b + a A)}{D^4 r (r^2 + A^2)} da + \frac{-c^2 A (r a D^2 + A b)}{D^4 r (r^2 + A^2)} db + \frac{c (D^2 r^2 + A^2)}{D^2 r (r^2 + A^2)} dc \quad \text{for patch 5}. \quad (60)$$

To write the wave equation, one also needs the inverse metric, which is

$$g^{\mu\nu} = \eta^{\mu\nu} - 2Hl^\mu l^\nu,$$

where all indices are raised with $\eta^{\mu\nu}$ (the inverse of the flat metric). The non-zero components of the latter are

$$\eta^{aa} = \frac{D^2(1 + a^2)}{c^2}, \quad (61)$$

$$\eta^{bb} = \frac{D^2(1+b^2)}{c^2}, \quad (62)$$

$$\eta^{cc} = 1, \quad (63)$$

$$\eta^{tt} = -1, \quad (64)$$

$$\eta^{ab} = \frac{abD^2}{c^2} \quad (65)$$

and the vector l^μ in the cubed-sphere coordinates is

$$l^\mu = \left[-1, \frac{-aA(rb - A)}{r(r^2 + A^2)}, \frac{A(a^2 - D^2)}{r^2 + A^2}, \frac{c(D^2r^2 + a^2A^2)}{D^2r(r^2 + A^2)} \right]^\mu \quad \text{for patches 0–3,} \quad (66)$$

$$l^\mu = \left[-1, -\frac{A(rb + aA)}{r(r^2 + A^2)}, \frac{A(ar - bA)}{r(r^2 + A^2)}, \frac{c(D^2r^2 + A^2)}{D^2r(r^2 + A^2)} \right]^\mu \quad \text{for patch 4,} \quad (67)$$

$$l^\mu = \left[-1, \frac{A(rb - aA)}{r(r^2 + A^2)}, -\frac{A(ar + bA)}{r(r^2 + A^2)}, \frac{c(D^2r^2 + A^2)}{D^2r(r^2 + A^2)} \right]^\mu \quad \text{for patch 5.} \quad (68)$$

4.3.1. Convergence tests. Figure 23 shows the differences, in the L_2 norm, between the numerical solutions at consecutive resolutions, using the D_{8-4} scheme, with no dissipation. The number of points in the angular directions is kept fixed to 16×16 points on each of the six patches, and the number of radial points ranges from 101 to 6401. The background is defined by a non-spinning black hole, and the inner and outer boundaries are at $1.9M$ and $11.9M$, respectively. Non-trivial initial data are given only to Π , in the form of a spherically symmetric Gaussian multipole:

$$\Pi(0, \vec{x}) = A \exp(r - r_0)^2 / \sigma_0^2, \quad (69)$$

with $r_0 = 5M$, $\sigma = M$, $A = 1$. The resulting convergence factors, the normalized differences $\|u_N - u_{2N}\| / \|u_N\|$ and the non-normalized ones, $\|u_N - u_{2N}\|$, are shown. The use of multiple patches not only allows for non-trivial geometries, but additionally one is able to define coordinates in a way such that resolution is adapted to the problem of interest. For example, in the geometry being considered, $S^2 \times R$, one employs a number of points in the angular direction limited by the expected multipoles of interest and concentrates resources to increase the number of points in the radial direction. As an example, the relative differences between the solutions at different resolutions shown in figure 23 reach values close to round-off, with modest computational resources. Even though the solution evolved here is spherically symmetric at the continuum, as discussed below the number of points used on each of the six patches that cover the sphere can reasonably resolve an $l = 2$ multipole. Next, figure 24 shows similar plots, but keeping the number of radial points fixed (to 101), using $N_a \times N_a$ points on each of the six patches in the sphere, with $N_a = 21, 41, 81$.

4.3.2. Tail runs. To illustrate the behaviour of the described techniques in 3D simulations, we examine the propagation of scalar fields on a Kerr black hole background. The numerical undertaking of such a problem has been previously treated using pseudo-spectral methods [20], which for smooth solutions allows the construction of very efficient schemes. As explained next, the combination of multi-block evolutions with high-order schemes also allows one to treat the problem quite efficiently. A detailed study of this problem will be presented elsewhere [21]; here we concentrate on two representative examples of what is achievable.

In the first case, we examine the behaviour of the scalar field propagating on a background defined by a black hole with mass $M = 1$ and spin parameter $a = 0.5$. Non-trivial initial

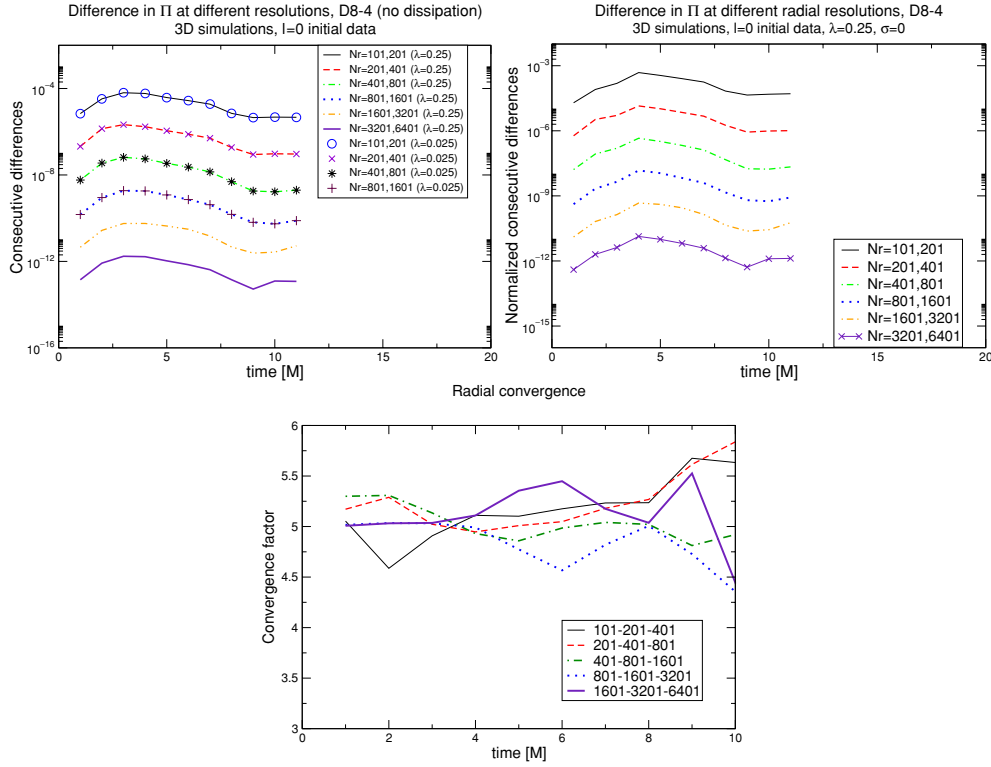


Figure 23. Convergence test in the radial direction, with CFL factors $\lambda = 0.25, 0.025$. Note that no appreciable difference is found between $\lambda = 0.25$ and a smaller value, and even with $\lambda = 0.25$ the convergence factor is not dominated by the time integrator.

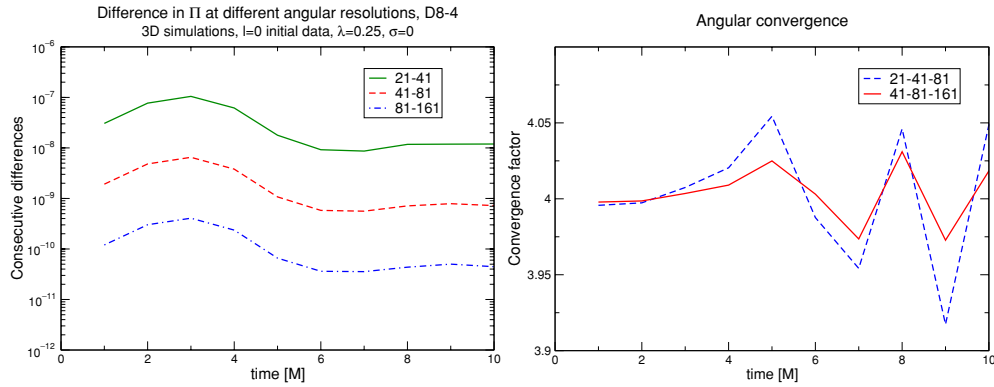


Figure 24. Convergence test in the angular direction, with a CFL factor $\lambda = 1/4$.

data are given only to Π , with a radial profile given by a Gaussian pulse as in equation (69), but with $r_0 = 20M$, and angular dependence given by a pure $l = 2$ multipole. The inner and outer boundaries are placed at $r = 1.8M$ and $r = 1001.8M$, respectively. We adopt a grid composed by six cubed-sphere patches, each of which is discretized with 20×20 points

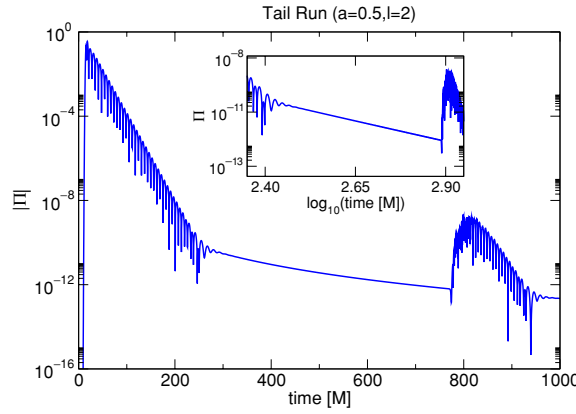


Figure 25. Behaviour of the time derivative of the scalar field for a pure multipole $l = 2$ initial data, on a Kerr background, with $a = 0.5$. Inner and outer boundaries are at $1.8M$ and $1001.8M$, respectively, and the six-patch grid is covered by $20 \times 20 \times 10001$ points on each patch. The D_{8-4} derivative is used, with no artificial dissipation. The noise at $t \approx 800M$ is due to the mode with group speed -2.6 discussed in section 3 hitting the outer boundary and reaching the observer at the black hole horizon. This noise goes away by either pushing the outer boundary, increasing resolution and/or adding dissipation. The average slope for Π in the interval $t \in [350M, 750M]$ gives a decay for Π of $t^{-4.07}$, in good agreement with the expected decay of t^{-4} . The inset shows a zoom in at the tail behaviour.

in the angular directions and 10001 points in the radial one. This translates into a relatively inexpensive calculation.

We adopt the D_{8-4} derivative operator, add no artificial dissipation and choose a CFL factor $\lambda = 0.25$. The salient features of the solution's behaviour observed are summarized in figure 25 which shows the time derivative of the scalar field as a function of time at a point in the equatorial plane on the event horizon. At earlier stages, the familiar quasi-normal ringing is observed. Next the late-time behaviour of the field reveals the expected tail behaviour as a fit in the interval $t \in [350M, 750M]$ gives a decay for Π of $t^{-4.07}$, which agrees quite well the expected decay of t^{-4} . This can be understood in terms of the generation of an $l = 0$ mode in the solution due to the spin of the black hole [20, 22]. Finally, noise can be observed appearing at $t \approx 800M$ due to the outer boundary. This noise, however, is not related to physical information propagating to the outer boundary and coming back (for this one would have to wait till $t \approx 2000M$) but, rather, is related to spurious modes with high group velocities travelling in the wrong direction, as described in section 3. As discussed there, for an eighth-order centred derivative the speed of this spurious mode is around -2.6 , which roughly matches with this noise appearing at $t \approx 800M$. We have checked that this boundary effect does go away with resolution, by introducing some amount of dissipation or by pushing the outer boundary farther out. Note that while the first two options allow one to observe the tail behaviour for much longer, eventually physical information would travel back from the outer boundary and ‘cavity’ effects which affect the decay would take place. Indeed, the behaviour would no longer be determined by a power law tail but by an exponential decay [23].

Figure 26 shows a similar run, in this case however the black hole is not spinning. A fit to the solution in the tail regime gives a decay for ϕ of $t^{-6.96}$, which again matches quite well the expected decay of t^{-7} [24].

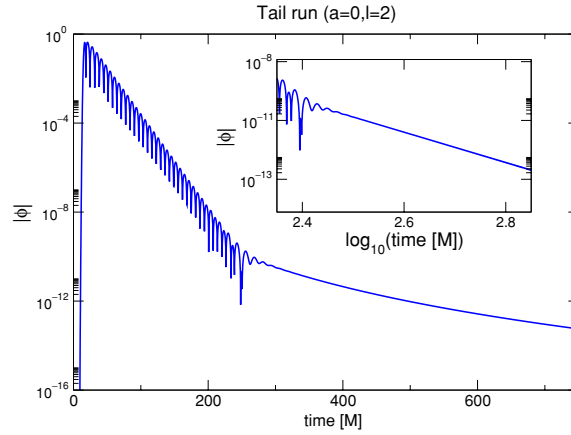


Figure 26. A simulation with the same parameters as those of figure 25, but with a non-spinning black hole. The average slope for Φ in tail regime gives a decay of $t^{-6.96}$, again in good agreement with the expected decay of t^{-7} .

5. Final comments

As illustrated in this work, the combination of the penalty technique together with those guaranteeing a stable single grid implementation for hyperbolic systems provides a way to achieve stable implementations of multi-block schemes of *arbitrary high order*. A similar penalty technique for multi-block evolutions is also being pursued in conjunction with pseudo-spectral methods [25].

The flexibility provided by multiple grids can be exploited to address a number of issues currently faced in simulations of Einstein's equations:

- The desire for a conforming inner boundary. This plays a central role in ensuring a consistent implementation of the excision technique together with a saving in the computational cost of the implementation.
- The need for a smooth outer boundary. This removes the presence of corners and edges which have proved difficult to deal with even at the analytical level [26, 27]. Furthermore, a smooth S^2 outer boundary simplifies tremendously the search for an efficient matching strategy to an outside formulation aimed to cover a much larger region of the spacetime with a formalism better suited to the asymptotic region (see, for example, [28–31]).
- The use of a grid that is better adapted to the description of wave phenomena as they propagate in the region far from the sources.

Acknowledgments

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Appendix A. Coefficients for high-order operators with diagonal metrics

For completeness, we point out here some misprints in [9] in some of the expressions for the diagonal metric cases.

- D_{2-1} : no typos.
- D_{4-2} : it says $\alpha_2 = -1/2$, but it should be $\alpha_2 = -1/12$. The expressions for q_{2i} are also missing; they should be the following:

$$q_{20} = \frac{8}{86} = q_{21} = -\frac{59}{86} = q_{23} = -q_{21}, \quad q_{24} = -q_{20}.$$

- D_{6-3} : no typos in the scalar product or coefficients for the derivative, neither in the general case nor in the minimum bandwidth one.
- D_{8-4} : the operator that has the minimum bandwidth is correct, but the three-parametric one has a typo in one of the coefficients (the scalar product is correct). It says

$$q_{06} = 49(-1244\,160x_1 + 18\,661\,400x_3 - 13\,322\,233)/17\,977\,668$$

when it should say

$$q_{06} = 49(-1244\,160x_1 + 18\,662\,400x_3 - 13\,322\,233)/17\,977\,668.$$

Appendix B. Dissipation for high-order difference operators with diagonal norms

The addition of artificial dissipation typically involves considering a dissipative operator Q_d which is non-positive with respect to the scalar product with respect to which SBP holds, i.e.,

$$\langle u, Q_d u \rangle \leq 0, \quad \forall u. \quad (\text{B.1})$$

We start the construction of such operators satisfying this property by defining them as

$$Q_d = (-1)^{m-1} \sigma h^n (D_+ D_-)^m \quad (\text{B.2})$$

on grid points lying within the range $[r, s]$ contained in the interval in which the weight used in the scalar product is 1. That is, the interval in which the difference operator is one of the centred ones of equations (27)–(30); for example, if the grid points range from 0 to N , then (r, s) must satisfy

$$r \geq 1, \quad s \leq N - 1 \quad \text{for } D_{2-1}, \quad (\text{B.3})$$

$$r \geq 4, \quad s \leq N - 4 \quad \text{for } D_{4-2}, \quad (\text{B.4})$$

$$r \geq 6, \quad s \leq N - 6 \quad \text{for } D_{6-3}, \quad (\text{B.5})$$

$$r \geq 8, \quad s \leq N - 8 \quad \text{for } D_{8-4}. \quad (\text{B.6})$$

As we will see later, in some cases our construction of dissipative operators imposes stricter constraints on the range of allowed values for r, s for each derivative.

If $n = 2m - 1$, the operator (B.2) is the standard Kreiss–Oliger dissipation (KO) [32], which is negative definite in the absence of boundaries (when the weight in the scalar product is identically 1). The choice $n = 2m - 1$ ensures that the added dissipation has the same ‘scale’ as the principal part (that is, length^{-1}) and that the resulting amplification factor is independent of resolution.

The addition of dissipation is commonly used to control unstable high-frequency modes which have associated amplification factors > 1 . In those cases, dissipation must be added such that the amount of dissipation on the high-frequency modes does not vary with respect to the resolution employed. This only occurs when $n = 2m - 1$. However, there are cases where the implementation is stable without the need of dissipation; here, one could opt to add dissipation to (partially) filter out higher frequency modes. In these cases, general values of n can be considered but the amount of dissipation will vary as the grid spacing is varied.

For each of the derivatives $D_{2-1}, D_{4-2}, D_{6-3}, D_{8-4}$, we seek to extend Q_d in equation (B.2) so that the resulting operator is negative definite with respect to the corresponding SBP scalar product. As mentioned above, we denote the points in which Q_d is given by equation (B.2) as $i = r, \dots, s$.

The identities of appendix C are used in the calculations needed for the construction of the operators below. These identities let one express the norm of the dissipation in the interior, which is proportional to $(-1)^{n-1}(u, (D_+ D_-)^n u)^{[r,s]}$, as

$$(-1)^{n-1}(u, (D_+ D_-)^n u)^{[r,s]} = \text{non-positive definite terms} + \sum_{i=0}^{r-1} u_i(\dots) + \sum_{s+1}^N u_i(\dots). \quad (\text{B.7})$$

Once this is done, the norm of the whole dissipative operator can be written as

$$\begin{aligned} (u, Q_d u)^{[0,N]} = & \text{non-positive definite terms} + \sum_{i=0}^{r-1} u_i[(\dots)_i + h\sigma_i Q_d u_i] \\ & + \sum_{s+1}^N u_i[(\dots)_i + h\sigma_i Q_d u_i] \end{aligned} \quad (\text{B.8})$$

and it is straightforward to make it non-positive definite. For example, by choosing

$$Q_d u_i = -\frac{(\dots)_i}{\sigma_i h}, \quad (\text{B.9})$$

which is how we proceed below. Note however this is not the only way to proceed. For instance, one could try to make the two sums of equation (B.8) cancel (as opposed to requiring the terms cancel at each grid point).

A more general approach for constructing dissipative operators that are negative definite with respect to the appropriate (SBP) scalar product has been recently presented by Mattson, Svard and Nordström [33].

B.1. Fourth derivative dissipation and KO type for D_{2-1}

In this case, the interior operator is

$$Q_d = -\sigma h^n (D_+ D_-)^2.$$

One can then split the norm of the dissipative operator over the whole grid into terms involving the left and right ranges $[0, r]$, $[s, N]$ (of the yet undefined operator) and the interior terms:

$$(u, Q_d u)_{\Sigma}^{[0,N]} = h \sum_{i=0}^{r-1} \sigma_i u_i Q_d u_i + h \sum_{s+1}^N \sigma_i u_i Q_d u_i - \sigma h^n (u, (D_+ D_-)^2 u)^{[r,s]},$$

where σ_i are the scalar product weights. Using the identities of appendix C, one can see that

$$\begin{aligned} (u, Q_d u)_{\Sigma}^{[0,N]} &= -\sigma h^n \|D_+ D_- u\|_{[r-1, s+1]}^2 + h \sum_0^{r-3} \sigma_i u_i Q_d u_i + h \sum_{s+3}^N \sigma_i u_i Q_d u_i \\ &\quad + h u_{r-2} (\sigma_{r-2} Q_d + \sigma h^{n-2} D_+^2) u_{r-2} + h u_{r-1} [\sigma_{r-1} Q_d - \sigma h^{n-2} \\ &\quad \times (2D_+ D_- - D_+^2)] u_{r-1} + h u_{s+2} (\sigma_{s+2} Q_d + \sigma h^{n-2} D_-^2) u_{s+2} \\ &\quad + h u_{s+1} [\sigma_{s+1} Q_d - \sigma h^{n-2} (2D_+ D_- - D_-^2)] u_{s+1}. \end{aligned}$$

There are several options now to control the contribution of these terms, the simplest one is

$$\begin{aligned} Q_d u_{r-2} &= -\frac{\sigma h^{n-2}}{\sigma_{r-2}} D_+^2 u_{r-2}, & Q_d u_{r-1} &= \frac{\sigma h^{n-2}}{\sigma_{r-1}} (2D_+ D_- - D_+^2) u_{r-1}, \\ Q_d u_{s+1} &= \frac{\sigma h^{n-2}}{\sigma_{s+1}} (2D_+ D_- - D_-^2) u_{s+1}, & Q_d u_{s+2} &= -\frac{2\sigma}{\sigma_{s+2}} h^{n-2} D_-^2 u_{s+2} \end{aligned}$$

and $Q_d = 0$ everywhere else, which implies $r \geq 2, s \leq N-2$. Note that, as anticipated, this is a further constraint on the possible range of r, s (cf equation (B.3)). The operator constructed not only satisfies the non-positivity requirement, but also transforms according to $D_- \rightarrow -D_+$ under the symmetry $i \rightarrow N-i$ (as it should). This dissipative operator is a trivial generalization of that constructed in [34].

Preferred choice. A KO-type dissipation for D_{2-1} , which we call KO4 (because it involves fourth derivatives in the interior) corresponds to

$$n = 3.$$

With this choice, the order of the scheme is 1 at at least two points (while in the absence of dissipation would be of order 1 at just one point). In general, the best option for r, s is

$$r = 2, \quad s = N-2,$$

as in that case the dissipation operator is non-trivial everywhere (while choosing $r > 2, s < N-2$ would imply that the dissipation is zero at—and even possibly close to—the boundaries). With this choice for r, s , the order of the scheme is 1 at the last two boundary points.

B.2. Sixth derivative dissipation and KO type for D_{4-2}

We now proceed to construct a higher derivative dissipative operator. As before, we adopt at the interior points the standard dissipation operator,

$$Q_d = \sigma h^n D_+^3 D_-^3.$$

Then, defining $v = D_+ D_- u$ and employing the identities of appendix C, one obtains

$$\begin{aligned} (u, Q_d u)_{\Sigma}^{[0,N]} &= -\sigma h^n \|D_- v\|_{[r-1, s+2]}^2 + h \sum_0^{r-1} \sigma_i u_i Q_d u_i + h \sum_{s+1}^N \sigma_i u_i Q_d u_i \\ &\quad + u_{r-3} h (-\sigma h^{n-3} D_+ v_{r-2} + \sigma_{r-3} Q_d u_{r-3}) \\ &\quad + u_{r-2} h (\sigma h^{n-3} (-h D_+^2 + 2D_+) v_{r-2} + \sigma_{r-2} Q_d u_{r-2}) \\ &\quad + u_{r-1} h [\sigma h^{n-3} (h(2D_+ D_- - D_+^2) v_{r-1} - D_+ v_{r-2}) + \sigma_{r-1} Q_d u_{r-1}] \\ &\quad + u_{s+1} h [\sigma h^{n-3} (h(2D_+ D_- - D_-^2) v_{s+1} + D_- v_{s+2}) + \sigma_{s+1} Q_d u_{s+1}] \\ &\quad + u_{s+2} h (\sigma h^{n-3} (-h D_-^2 - 2D_-) v_{s+2} + \sigma_{s+2} Q_d u_{s+2}) \\ &\quad + u_{s+3} h (\sigma h^{n-3} D_- v_{s+2} + \sigma_{s+3} Q_d u_{s+3}). \end{aligned}$$

Here again one has several options, the simplest one is

$$\begin{aligned}
Q_d u_{r-3} &= \frac{\sigma h^{n-3}}{\sigma_{r-3}} D_+ v_{r-2}, \\
Q_d u_{r-2} &= \frac{\sigma h^{n-2}}{\sigma_{r-2}} (h D_+^2 - 2 D_+) v_{r-2}, \\
Q_d u_{r-1} &= -\frac{\sigma h^{n-2}}{\sigma_{r-1}} [h(2 D_+ D_- - D_+^2) v_{r-1} - D_+ v_{r-2}], \\
Q_d u_{s+1} &= -\frac{\sigma h^{n-2}}{\sigma_{s+1}} [h(2 D_+ D_- - D_-^2) v_{s+1} + D_- v_{s+2}], \\
Q_d u_{s+2} &= \frac{\sigma h^{n-2}}{\sigma_{s+2}} (h D_-^2 + 2 D_-) v_{s+2}, \\
Q_d u_{s+3} &= -\frac{\sigma h^{n-3}}{\sigma_{s+3}} D_- v_{s+3}
\end{aligned}$$

and $Q_d = 0$ everywhere else. Note that, as in the D_{2-1} case, Q_d transforms under the symmetry $i \rightarrow N-i$ as it should. Note, however, that since we are modifying the dissipative operator at three points near the boundary, no further constraint occurs in the allowed range for r, s (cf equation (B.4)), unlike the D_{2-1} case.

Preferred choice. A KO-type dissipation for D_{4-2} , which we call KO6, corresponds to

$$n = 5.$$

In general, the best choice for r, s is

$$r = 4, \quad s = N - 4,$$

as in that case the dissipation does not reduce the order of the overall scheme¹². However, a drawback of our simplified method for constructing dissipative operators (which cannot be remedied by adopting different values for r, s) is that in this case the dissipation is zero at the last grid point.

B.3. Eighth derivative dissipation and KO type for D_{6-3}

As usual, we start with

$$Q_d = -\sigma h^n D_+^4 D_-^4, \quad i = r, \dots, s. \quad (\text{B.10})$$

Then, using the identities of appendix C, the norm of the dissipative operator results

$$\begin{aligned}
(u, Q_d u)_{\Sigma}^{[0,N]} &= h \sum_0^{r-1} \sigma_i u_i Q_d u_i + h \sum_{s+1}^N \sigma_i u_i Q_d u_i - \sigma h^n (u, (D_+ D_-)^4 u)_{[r,s]} \\
&= -\sigma h^n \|D_-^3 D_+ u\|_{[r,s+2]}^2 + u_{r-1} [-\sigma h^{n-3} (h^2 (2 D_+ D_- - D_+^2) w_{r-1} \\
&\quad - h D_+ w_{r-2} + D_+ \alpha_{r-2}) + \sigma_{r-1} h Q_d u_{r-1}] \\
&\quad + u_{r-2} [-\sigma h^{n-3} ((-h^2 D_+^2 + 2 h D_+) w_{r-2} - 3 D_+ \alpha_{r-2}) + \sigma_{r-2} h Q_d u_{r-2}] \\
&\quad + u_{r-3} [-\sigma h^{n-3} (-h D_+ w_{r-2} + 3 D_+ \alpha_{r-2}) + \sigma_{r-3} h Q_d u_{r-3}]
\end{aligned}$$

¹² Since the D_{4-2} operator already has four points close to each boundary where its order is 2, while the above dissipative operator has only three points where this happens.

$$\begin{aligned}
& + u_{r-4}(\sigma h^{n-3} D_+ \alpha_{r-2} + \sigma_{r-4} h Q_d u_{r-4}) + u_{s+4}(\sigma h^{n-3} D_+ \alpha_{s+2} + \sigma_{s+4} h Q_d u_{s+4}) \\
& + u_{s+3}[-\sigma h^{n-3}(h D_- w_{s+2} + 3 D_+ \alpha_{s+2}) + \sigma_{s+3} h Q_d u_{s+3}] \\
& + u_{s+2}[-\sigma h^{n-3}(h^2 D_-^2 + 2h D_-)w_{s+2} - 3 D_+ \alpha_{s+2}) + \sigma_{s+2} h Q_d u_{s+2}] \\
& + u_{s+1}[-\sigma h^{n-3}(h^2(2D_+ D_- - D_-^2)w_{s+1} + h D_- w_{s+2} + D_+ \alpha_{s+2}) + \sigma_{s+1} h Q_d u_{s+1}],
\end{aligned}$$

where $w = (D_+ D_-)^4 u$, $\alpha = D_- D_+ D_- u$. The simplest choice is given by setting to zero all these coefficients, which when expanding w and α gives

$$Q_d u_{r-1} = \frac{\sigma h^{n-4}}{\sigma_{r-1}} [h^2(2D_+ D_- - D_+^2)(D_+ D_-)^4 u_{r-1} - h D_+(D_+ D_-)^4 u_{r-2} + (D_+ D_-)^2 u_{r-2}], \quad (\text{B.11})$$

$$Q_d u_{r-2} = \frac{\sigma h^{n-4}}{\sigma_{r-2}} [(-h^2 D_+^2 + 2h D_+)(D_+ D_-)^4 - 3(D_+ D_-)^2] u_{r-2}, \quad (\text{B.12})$$

$$Q_d u_{r-3} = \frac{\sigma h^{n-4}}{\sigma_{r-3}} [-h D_+(D_+ D_-)^4 + 3(D_+ D_-)^2] u_{r-2}, \quad (\text{B.13})$$

$$Q_d u_{r-4} = -\frac{\sigma h^{n-4}}{\sigma_{r-4}} (D_+ D_-)^2 u_{r-2}, \quad (\text{B.14})$$

$$Q_d u_{s+4} = -\frac{\sigma h^{n-4}}{\sigma_{s+4}} (D_+ D_-)^2 u_{s+2}, \quad (\text{B.15})$$

$$Q_d u_{s+3} = \frac{\sigma h^{n-4}}{\sigma_{s+3}} [h D_-(D_+ D_-)^4 + 3(D_+ D_-)^2] u_{s+2}, \quad (\text{B.16})$$

$$Q_d u_{s+2} = \frac{\sigma h^{n-4}}{\sigma_{s+2}} [-(h^2 D_-^2 + 2h D_-)(D_+ D_-)^4 - 3(D_+ D_-)^2] u_{s+2}, \quad (\text{B.17})$$

$$a Q_d u_{s+1} = \frac{\sigma h^{n-4}}{\sigma_{s+1}} [h^2(2D_+ D_- - D_-^2)(D_+ D_-)^4 u_{s+1} + h D_-(D_+ D_-)^4 u_{s+2} + (D_+ D_-)^2 u_{s+2}] \quad (\text{B.18})$$

and $Q_d = 0$ everywhere else.

Preferred choice. If we want a KO-type dissipation for the D_{6-3} derivative, which we call KO8, we have to choose

$$n = 7.$$

There are six points near each boundary where the difference operator has order 3, while the above dissipation has four points where this happens. Therefore, the order of the whole scheme is not spoiled if

$$r = 6, \quad s = N - 6.$$

As in the D_{4-2} case, the drawback of our construction is that the dissipation is zero near boundaries; in this case, at the last two points.

B.4. Dissipation for D_{8-4}

In this case, we have not been able to write an interior KO dissipation which does not spoil the eighth-order accuracy of the derivative (that is, the one as in equation (B.2), with $m = 5$)

in the form of equation (B.8). Therefore, our simplified approach does not work in this case. However, since the expressions of the previous subsections are valid for a general scalar product, we can use, for example, expressions (B.10) and (B.11)–(B.18) with the weights corresponding to the D_{8-4} derivative and

$$n = 7, \quad r = 8, \quad s = N - 8.$$

This results in a KO dissipation of the KO8 type which is zero at the last four points, with the drawback that the order of the scheme is reduced to 7 at the interior points and 3 near boundaries (as opposed to 8 and 4, respectively, in the derivative itself).

Appendix C. Useful properties in the construction of dissipative operators

It is straightforward to show that with respect to the scalar product and norm

$$(u, v)^{[r,s]} \equiv \sum_{j=r}^s u_j v_j h_x, \quad (\|u\|^{[r,s]})^2 = (u, u)^{[r,s]},$$

the following properties hold:

$$(u, D_+ v)^{[r,s]} = -(D_- u, v)^{[r+1,s+1]} + u_j v_j|_r^{s+1}, \quad (C.1)$$

$$(u, D_- v)^{[r,s]} = -(D_+ u, v)_{[r-1,s-1]} + u_j v_j|_{r-1}^s, \quad (C.2)$$

$$(u, D_0 v)^{[r,s]} = -(D_0 u, v)_{[r,s]} + \frac{1}{2}(u_j v_{j+1} + u_{j+1} v_j)|_{r-1}^s. \quad (C.3)$$

The proofs for the first and third identities can be found in [35], and the second one is trivially obtained from the first.

The following equalities are also straightforward to check, though obtaining them becomes increasingly more cumbersome as more derivatives are involved:

$$\begin{aligned} (u, D_+ D_- v)^{[r,s]} &= -(D_- u, D_- v)^{[r+1,s+1]} + (u_j D_- v_j)_r^{s+1} \\ &= -(D_- u, D_- v)^{[r,s+1]} - u_{r-1} D_+ u_{r-1} + u_{s+1} D_- u_{s+1}, \\ (u, (D_+ D_-)^2 v)^{[r,s]} &= (D_+ D_- u, D_+ D_- v)^{[r-1,s+1]} + \frac{u_{s+1}}{h} (2D_+ D_- - D_-^2) v_{s+1} \\ &\quad - \frac{u_{s+2}}{h} D_-^2 v_{s+2} + \frac{u_{r-1}}{h} (2D_+ D_- - D_+^2) v_{r-1} - \frac{u_{r-2}}{h} D_+^2 v_{r-2}, \\ (u, (D_+ D_-)^3 v)^{[r,s]} &= -(D_- p, D_- w)_{[r-1,s+2]} \\ &\quad + \frac{u_{r-1}}{h} \left((2D_+ D_- - D_+^2) w_{r-1} - \frac{1}{h} D_+ w_{r-2} \right) \\ &\quad + \frac{u_{r-2}}{h} \left(-D_+^2 + \frac{2}{h} D_+ \right) w_{r-2} - \frac{u_{r-3}}{h^2} D_+ w_{r-2} \\ &\quad + \frac{u_{s+1}}{h} \left((2D_+ D_- - D_-^2) w_{s+1} + \frac{1}{h} D_- w_{s+2} \right) \\ &\quad + \frac{u_{s+2}}{h} \left(-D_-^2 - \frac{2}{h} D_- \right) w_{s+2} + \frac{u_{s+3}}{h^2} D_- w_{s+2}, \\ (u, (D_+ D_-)^4 v)^{[r,s]} &= (D_- q, D_- \alpha)_{[r,s+2]} \\ &\quad + \frac{u_{r-1}}{h} \left((2D_+ D_- - D_+^2) w_{r-1} - \frac{1}{h} D_+ w_{r-2} + \frac{1}{h^2} D_+ \alpha_{r-2} \right) \end{aligned}$$

$$\begin{aligned}
& + \frac{u_{r-2}}{h} \left(\left(-D_+^2 + \frac{2}{h} D_+ \right) w_{r-2} - \frac{3}{h^2} D_+ \alpha_{r-2} \right) \\
& + \frac{u_{r-3}}{h^2} \left(-D_+ w_{r-2} + \frac{3}{h} D_+ \alpha_{r-2} \right) - \frac{u_{r-4}}{h^3} D_+ \alpha_{r-2} \\
& - \frac{u_{s+4}}{h^3} D_+ \alpha_{s+2} + \frac{u_{s+3}}{h^2} \left(D_- w_{s+2} + \frac{3}{h} D_+ \alpha_{s+2} \right) \\
& + \frac{u_{s+2}}{h} \left(- \left(D_-^2 + \frac{2}{h} D_- \right) w_{s+2} - \frac{3}{h^2} D_+ \alpha_{s+2} \right) \\
& + \frac{u_{s+1}}{h} \left((2D_+ D_- - D_-^2) w_{s+1} + \frac{1}{h} D_- w_{s+2} + \frac{1}{h^2} D_+ \alpha_{s+2} \right),
\end{aligned}$$

where $w = D_+ D_- v$, $p = D_+ D_- u$, $q = D_-^2 D_+ u$, $\alpha = D_-^2 D_+ v$ and σ_i denotes the value of the scalar product at grid point i .

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