Author response to the comments of the Referee Dr. Eigil Kaas.

First of all, we would like to warmly thank Dr. Kaas for his very helpful comments and further advises regarding their implementation. These allowed us to improve the paper, in particular, adding several new directions in the model evaluation and comparison with other approaches. Below, we provide a point-by-point response to the comments.

The Word file with all changes tracked is given as a Supplementary zip archive.

General comments

A main issue is the numerical accuracy. Probably it is rather difficult to perform an analytic von Neuman analysis with calculation of amplification factors etc. However, this can easily be (and should be) done numerically for different harmonic wave numbers using different time steps (and fft’s back and forth). Such an analysis should include a comparison with a known scheme, i.e., a classical semi-Lagrangian scheme with cubic interpolation.

Thank you for this suggestion and advises on its implementation! We added a section with the numerical spectral analysis and comparison with two classic schemes: the semi-Lagrangian and the Bott algorithms. The new section is 4.5.

The scheme has some tendency to introduce anti-diffusion for certain shapes and a smoother is therefore introduced in a rather non rigorous way (equation 20). This smoothing seems important in order to obtain reasonable results. However, the nature of the smoother is rather unusual and a more detailed analysis of its performance is requested. In particular it must be clarified in a quantitative way why the value of k is set to approximately 0.08. What is the exact value that is used in the different runs (hopefully the same value in all runs)?

We certainly used the same value in all runs with the smoother, the tests can only be with or without it. The paper is now modified with explicit references to whether the smoother was used or not.

We also expanded its description is section 4.4, put explicit analysis of its impact and optimal strength in spectral analysis section and in all tests where it was used. Finally, we added a new section 7.3 in Discussion, where outlined the cases requiring its application.

The idea of the smoother came from the qualitative observation that the scheme with mass centres fixed in the middle of the grid cells is quite similar to a simple upwind algorithm, which is very diffusive and thus has no gradient-sharpening tendencies. The relaxation towards this scheme was
then suggested as a way to counter the anti-diffusivity. The idea is particularly tempting because the additional costs are completely negligible. Specific value of the relaxation factor has been found from numerical tests, which also confirmed its stability in a variety of cases. The actual value of 0.08 was found as the lowest distortion that damps the anti-diffusion in the tests leading to practically non-distorted gradients. In spectral analysis, this corresponds to lowering the amplification factor below 1 for (almost) all wave numbers at (almost) all times.

As far as I can judge, the scheme has similarities to the CIP and CIP-CLSR schemes by Xiao and others - see below under Specific comments.

We have added the scheme to the introduction, and drew attention to the generic idea behind the whole class of such schemes: to use ”something” that would allow for higher resolution than the formal grid cell size. Apart from this general idea, the similarities seem to be quite limited, less significant than with, e.g. the Egan & Makhoney scheme. The CSLR takes rational functions for describing the within-cell concentration profile. However, there are no slabs and centres of masses, which are the corner stone of the Galperin’s approach. Instead, CSLR used concentrations at the interface points to define the profiles. The brings about another principal difference: the interface points are affected by two neighbouring cells, while in Galperin’s approach each cell completely decides on its slab when it is formed. The information is passed only with the transported mass.

Specific comments

l. 146 Units are not just mol or kg but mol/kg and kg/kg since they represent mixing ratios.

In fact, the model operates with masses in the grid cells and, for chemistry, with concentrations, but mmr is, indeed, mol/kg and kg/kg. Clarification added.

l. 159 "The Eq. (1)" should be replaced by "Eq. (1)"

l. 173 "Eulerian dynamics of SILAM". It is proposed to change this to "Eulerian transport scheme of SILAM". Otherwise the reader may be confused to think that the scheme is used for the atmospheric dynamics (on-line coupling), which is not the case.

l. 186 "The Eq. (5)" should be replaced by "Eq. (5)"

l. 205 "in the Eq. (5)" should be replaced by "in Eq. (5)"
l. 291 "In application to Galperin scheme" should be "For application to the Galperin scheme" or "For application in the Galperin scheme"

Figure 2 It is disturbing that the legends overlap the figures. Please, use one common legend, e.g., between the two upper and two lower panels. Also, in the caption "Legend includes ..." should be replaced with "The Legend includes ...".

All the deficiencies have been corrected

l. 253 In my opinion the review of similar schemes is not complete without a discussion of the CIP-CLSR scheme by Xiao (2002). In its essence this scheme is quite similar to the new scheme presented in the manuscript. In CIP, the total field is represented by the cell averages as well as the interphase values between cells. In this way the interphase values play the same role as the center of mass locations in the Galperin scheme. .... Thus, a careful discussion of the essential difference between CIP / CIP-CLSR and the new scheme is requested.

We included the references to the introduction. Please see above remarks on the schemes similarity.

l. 320-334 Apparently the Galperin scheme is somehow antidiffusive - although not for all shapes. The distortions seem to be a rather serious issue. The smoothing effectively introduced via the expression in eq. (20) seems, as also noted by the authors, to be a non-rigorous fix. It is, however, obvious that the numerical diffusion is enhanced when this "filter" is introduced. An analysis of this should be included in the "numerical von Neuman analysis" mentioned under General comments.

The impact of this smoother is included in the sections with test analysis, plus two dedicated sections discuss it now. Please see above.

Sect. 6.2. It is really good that all these tests have been run. I don’t think, however, that the weaknesses of the scheme are commented/recognised in sufficient depth. You should discuss the rather widespread noise away from the advected entities. The most depressing result is probably the tendency for the scheme to unmix even when the smoother is on (Figure 11). A general discussion of the Figures 7 through 11 should be included, which directly compare the results with some of the schemes in LI4.
The discussion has been strongly expanded in section 7.2 and the issues of the noise far from the advected bodies and the “unmix” included. We added new chart showing the mixing diagnostic, in addition to previously available numerical information printed on the “bow” charts. The corresponding discussion is added. We also highlighted that origin of some of the issues, first of all, the non-monotonic background, is not the scheme per-se (the 1D algorithm) but rather the dimensional split used for its implementation in 3D (well, 2D in that specific case). Similarly, the tendency to unmix the initially homogeneous field is due to this very splitting: the 1D algorithm transports a flat plain perfectly.

We also added the discussion on the smoother, which improves the linearity but: (i) it has low power against the background fluctuations, largely responsible for the unmix, (ii) we limited its power ($\varepsilon=0.08$!) in order to avoid excessive diffusivity. Therefore, remnants of the unmix still remained visible. We made it clear in the revised paper.

l. 476 “Eq. (21)” should be "in Eq. (21)". However, I am a little confused since this equation relates to the vertical direction, while the tests described here are horizontal. I suspect that the equation number should be (20) and not (21). The correlated tracer runs are run both with and without the smoothing. It is unclear, however, if the smoother in (20) was applied to all other runs or if it has been included or not on an ad hoc basis. This must be clarified.

We checked the numbering (of course, eq. 20, not 21) and made it clear throughout the paper what pictures represent the runs with / without the smoother.

l. 506-514 It makes little sense to compare the performance of the Galperin scheme on one computer with that of other transport schemes, which are run on a different computer. One should instead compare directly with the performance of another transport scheme on the same computer. I suggest a simple (non-conserving) semi-Lagrangian scheme with bi-cubic interpolation. For a model like SILAM it is highly relevant to test the multi-tracer efficiency, as it was done for instance in Kaas et al. (2013), which you refer to. This should be done.

We have reviewed the efficiency section, moved it to Discussion (now it is section 7.4) and provided more extensive analysis. In particular, it now includes the comparison with Bott scheme in its standard implementation available from internet – and with a simple semi-Lagrangian scheme made by ourselves. We also made a new subsection with analysis of the model scalability with regard to the number of species, horizontal resolution, and time step.
Regarding the comparison with HEL and CSLAM, in our experience the difference between the modern computers is usually comparatively small (tens of %) and Linpack benchmarks give a good representation of it for the purposes of dispersion modelling. The impact of implementation quality, however, can be large – e.g. we got a factor of 2-3 by quite basic optimization of our code for the Galperin scheme. We found Linpack scores for the processor used by Kaas et al – 20GFlops (now included in the paper, sorry for the omission), – similar to 18GFlops of our notebook. We also found another comparison of these CPUs with a different series of tests, which confirmed the similarity within some 20% margin. Therefore, the similarity between the machines is confirmed, with probably weaker computer used in our tests: Linpack is marginally lower and the bus capacity is quite low (a usual feature of notebooks). Therefore, we think that our comparison is fair: tests are identical, computers are similar and, most importantly, the implementations of the schemes come from their authors, i.e. their quality is as high as one can get. Together with comparison with “reference” implementation of Bott algorithm and the semi-Lagrangian scheme, this should give sufficient credibility to the efficiency assessment of the scheme. These issues are now explained in the paper.
Construction of the Eulerian atmospheric dispersion model
SILAM based on the advection algorithm of M. Galperin

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1. Abstract

The paper presents the transport module of the System for Integrated Modelling of Atmospheric Composition SILAM v. 5 based on the advection algorithm of Michael Galperin. This advection routine, so far weakly presented in international literature, is positively defined, stable at any Courant number, and efficient computationally. We present the rigorous description of its original version, along with several updates that improve its monotonicity and shape preservation, allowing for applications to long-living species in conditions of complex atmospheric flows. The scheme is connected with other parts of the model in a way that preserves the sub-grid mass distribution information that is a corner-stone of the advection algorithm. The other parts include the previously developed vertical diffusion algorithm combined with dry deposition, a meteorological pre-processor, and chemical transformation modules.

Quality of the advection routine is evaluated using a large set of tests. The original approach has been previously compared with several classic algorithms widely used in operational
dispersion models. The basic tests were repeated for the updated scheme and extended with
real-wind simulations and demanding global 2-D tests recently suggested in literature, which
allowed positioning the scheme with regard to sophisticated state-of-the-art approaches. The
advection scheme performance was fully comparable with other algorithms, with a modest
computational cost.

This work was the last project of Dr. Sci. Michael Galperin who untimely passed away on 17
March 2008.

Keywords: advection schemes, numerical algorithms, dispersion modelling, Eulerian model.

2. Introduction

One of the key problems in atmospheric composition modelling is the accuracy and reliability
of numerical schemes. A less appreciated but important issue is the consistency of the
approaches applied in different modules of the modelling system. Usually, model construction
follows process-wise split (Yanenko 1971; Marchuk 1995; Seinfeld & Pandis 2006), thus
considering separately the advection and diffusion algorithms, physical and chemical
transformations, and dry and wet deposition. In practical model developments, features of the
transport algorithms, first of all, advection scheme, largely shape-up the model and determine
its area of application.

2.1. Advection schemes

There are numerous types of advection schemes currently employed in atmospheric dispersion
models. Two major categories refer to Lagrangian or Eulerian treatment of tracers: as small-
size masses (Lagrangian particles) or as the concentration fields discretised in a prescribed grid. The Eulerian schemes, the primary subject of this paper, can be divided to flux-form finite volume, semi-Lagrangian, or expansion-function schemes. The expansion-function schemes approximate the solution with a given set of basis functions and, in turn, can be divided to spectral, pseudospectral and finite-element approaches. Some classic schemes are also based on finite-difference approximations of the advective term of the transport equation. The basic principles of all these approaches were formulated several decades ago and, with certain modifications, are still in use. Many modern schemes combine several approaches. The large diversity of the advection algorithms is explained by a long list of requirements to such schemes. The most important ones are: positive definition, minimal numerical diffusion, limited non-monotonicity and non-linearity, stability with regard to high Courant number (the number of the model grid cells passed within one advection time step), small phase error, local and global mass conservation, and high numerical efficiency. Some of these requirements contradict to each other. For example, numerical diffusion “blurs” the resulting patterns but also makes them smoother, thus improving the monotonicity.

The finite-difference schemes involve direct discretization of the dispersion equation and various interpolation functions to evaluate derivatives of the concentration fields (see reviews of (Richtmyer 1962; Leith 1965; Roach 1980), as well as section 3.1 in (Rood 1987); specific examples are, for instance, (Russell & Lerner 1981; Van Leer 1974; Van Leer 1977; Van Leer 1979). These schemes, being once popular, usually suffer from large numerical diffusion and limited stability, which sets stringent limitations to the Courant number usually requiring it to be substantially less than one. Therefore, the interest has gradually shifted towards flux, finite-element, and semi-Lagrangian schemes. The flux schemes represent the transport via mass fluxes at the grid cell borders, which are calculated from concentrations in the neighbouring cells and wind characteristics. They are
inherently mass conservative and have become popular in atmospheric chemistry transport models (Kukkonen et al. 2012). Probably the most widely used flux-type scheme is the one made by A. Bott (Bott 1989; Bott 1992; Bott 1993), especially if one would count the numerous Bott-type schemes (see examples in (Syrakov 1996; Syrakov & Galperin 1997; Syrakov & Galperin 2000; Walcek & Aleksic 1998; Walcek 2000; Yamartino 1993), which are based on the same principle but involve different approximation functions, monotonicity and normalization procedures, etc.

The semi-Lagrangian schemes have been among the most widely used approaches for decades, with numerous algorithms using its basic concept [Crowley, 1968; Egan and Mahoney, 1972; Pedersen and Prahm, 1974; Pepper and Long, 1978; Prather, 1986; Smolarkiewicz, 1982; Staniforth and Cote, 1991 and references therein], [Lowe et al., 2003; Sofiev, 2000], etc. In the forward form, these schemes consider the transport of mass starting from the grid mesh points (departure points) and calculate the masses at the grid points closest to the arrival point. Backward algorithms start from arrival grid points and find the grid points near the departure point. The schemes can be based on tracking either grid points or grid cells along their trajectories. The gridpoint-based schemes are not inherently mass-conserving, whereas the volume-based schemes achieve mass conservation by integrating the mass over the departure volume. They can sometimes be described as a combination of finite-volume and semi-Lagrangian methods (Lin & Rood 1996, 1997). Stability of these schemes can be ensured for a wide range of Courant numbers (Leonard 2002). A general review can be found in (Lauritzen et al. 2011), whereas a comparison of 19 modern schemes is discussed in (Lauritzen et al. 2014), hereinafter referred to as L14.

Modelling in spectral space is another approach with long history (Ritchie 1988; Kreiss & Oliger 1972; Zlatev & Berkowicz 1988; Prahm & Christensen 1977) but not widely used today.
One of the main problems of the existing schemes is substantial numerical diffusion originating from the finite-step discretization along space and time. Seemingly inevitable in Eulerian context, this phenomenon, however, does not exist in Lagrangian advection schemes, which do not contain explicit discretization of particle movement. Lagrangian domain is a continuous space rather than a set of pre-defined grid meshes and the position of the particles can be calculated precisely. As a result, numerical diffusion of purely Lagrangian schemes is always zero – at a cost of strongly non-monotonous concentration fields due to limited spatial representativeness of a single Lagrangian particle and a limited number of particles.

One of ways to reduce the diffusivity of an Eulerian scheme is to store additional prognostic variables to describe the state of each grid cell with higher spatial resolution than the formal cell size: a sub-grid mass distribution. This can take a form of extra conservation equation(s) for, e.g., first- or higher-order moments (Egan & Mahoney 1972; Prather 1986). There are other approaches that use different kind of the extra information. For instance, the Conservative Semi-Lagrangian schemes (Yabe & Aoki 1991; Yabe et al. 2001) use a semi-Lagrangian step to evaluate the mixing ratio at cell interfaces, and then use the interface values along with the cell integrals to derive an interpolant representing the sub-grid distribution.

In a series of works, Michael Galperin developed a semi-Lagrangian scheme that used the sub-grid information on mass centre location inside the cell. The scheme was made fully non-diffusive for isolated plumes, positively defined, and very efficient computationally (Galperin et al. 1994; Galperin et al. 1995; Galperin et al. 1997; Galperin 1999; Galperin & Sofiev 1998; Galperin & Sofiev 1995; Galperin 2000). The early version of this scheme applied in the large-scale modelling by (Sofiev 2000) resembled the “moving-centre” approach widely used in aerosol dynamics models (Kokkola et al. 2008) and shared its characteristic weakness – high non-monotonicity. The later developments substantially reduced it without damaging.
other features (Galperin 1999; Galperin 2000). Further development of this scheme is the subject of the current paper.

### 2.2. Horizontal and vertical diffusion, dry deposition

Diffusion algorithms are less diverse than advection schemes. The physical ground for one of the common diffusion parameterizations is described in details by (Smagorinsky 1963), who suggested a formula for grid-scale scalar eddy-diffusivity based on the model resolution and wind speed derivatives, thus connecting the numerical features of the simulations and hydrodynamics. It is widely used in chemical transport models (Kukkonen et al. 2012).

The dry deposition is usually accounted for as a boundary condition to the vertical advection-diffusion equation. Computation of dry deposition for gases practically always follows the electrical analogy, for which one of the first comprehensive parameterizations was suggested by (Wesely 1989). Among the extensions of this approach, one was suggested by Sofiev, (2002), who combined it with vertical diffusion and connected with the Galperin advection scheme. The algorithm used effective mean diffusion coefficient over thick layers calculated from high-resolution meteorological vertical profiles, the direction also recommended by Venkatram & Pleim (1999). These thick layers were determined using the subgrid information available from the advection scheme, which increased the accuracy of both algorithms (Sofiev 2002).

For aerosol species, the electrical analogy is not correct (Venkatram & Pleim 1999). Compromising approaches suggested by (Slinn 1982; Zhang et al. 2001) and updated by Petroff and Zhang (2010) involve numerous empirical relations, sometimes on thin ground. More rigorous approach unifying the gas and aerosol deposition parameterizations into a single solution was developed by (Kouznetsov & Sofiev 2012).
2.3. Organization of the paper

The current paper describes the Eulerian transport algorithm of the System for Integrated modeling of Atmospheric composition SILAM v.5, which is based on the advection scheme of Michael Galperin with several updates. The paper is organised as follows. Section 3 describes the original algorithm of M. Galperin and positions the scheme among other approaches. Section 4 presents the improvements made during its implementation and testing in SILAM. The Section 5 outlines the scheme interconnections with other model parts. Standard and advanced model tests are shown in section 6. Finally, discussion in the section 7 includes analysis of the scheme performance in the tests, as well as comparison with other algorithms.

Following the SILAM standards, all units throughout the paper are the basic SI: [mole] for chemicals, [kg] for aerosols without chemical speciation, [m] for distance and size, [sec] for time, etc. The model operates with concentrations, [mol m\(^{-3}\)] or [kg m\(^{-3}\)]. Some of the below tests are formulated in mixing ratios [mol mol\(^{-1}\)] or [kg kg\(^{-1}\)].

3. Background

3.1. Basic equations

We consider the forward dispersion equation with the first-order K-theory-based closure for diffusion:

\[ L \varphi = E, \quad \text{where} \quad L = \frac{\partial}{\partial t} + \sum_{i=1}^{3} \left( u_i \frac{\partial}{\partial \xi_i} \right) - \sum_{i=1}^{3} \rho \mu_{ij} \frac{\partial}{\partial \xi_j} \frac{1}{\rho} + \zeta. \]

Here \( \varphi \) is concentration of the pollutant, \( t \) is time, \( E \) is emission term, \( \xi_i, i = 1..3 \) denote the three spatial axes, \( u_i \) are components of the transport velocity vector along these axes, \( \mu_{ij} \) are
components of the turbulent diffusivity tensor, $\rho$ is air density, and $\zeta$ represents transformation source and sink processes.

Equation (1) is considered on the time interval $t \in (t_n, t_{n+1})$ in the domain

$$\{\xi \in \Xi = [h_1, H] \times \Omega, \text{where the heights } h_1 \text{ and } H \text{ are the lower and upper boundaries of the }$$

computational domain and $\Omega$ is the horizontal computational area with border $\partial \Omega$. The initial conditions are:

$$\varphi|_{t_n} = \varphi_0(\xi)$$

Boundary conditions depend on type of the simulations. In a general form, they constrain concentration and/or its first derivative:

$$\alpha \frac{\partial \varphi}{\partial \xi} |_{i,j \in \Xi} + \beta \varphi|_{i,j \in \partial \Xi} = \gamma$$

Here the values of $\alpha$, $\beta$, and $\gamma$ depend on type of the boundary. In particular, dry deposition at the surface $\xi_3 = h_j$ is described via $\alpha = \mu_{j3}$, $\beta = -v_d$ (dry deposition velocity), $\gamma = 0$; prescribed concentration $\varphi_l$ at the lateral boundaries $\xi_{l,3} \in \partial \Omega$ implies $\alpha = 0$, $\beta = 1$, $\gamma = \varphi_l$, etc. A global domain would require periodic longitudinal conditions.

3.2. Advection scheme of Michael Galperin

The current section presents the advection algorithm suggested by Michael Galperin in 2000s as a contribution to Eulerian transport scheme of SILAM. The idea of the scheme can be found in a short methodological publication of (Galperin 2000) (in Russian) and conference materials (Galperin 1999; Sofiev et al. 2008). It is very briefly outlined by (Petrova et al. 2008) (hereinafter referred to as P08) but no systematic description exists so far.
For the 1-D case, let us define the simulation grid, \( \xi_i = x_i \), as a set of \( I \) grid cells \( i = 1...I \). Let the coordinate of the centre of the \( i \)-th grid cell be \( x_i \), and coordinates of the cell left- and right-hand borders be \( x_i - 0.5 \) and \( x_i + 0.5 \), respectively. The 1-D cell size is then \( V_i = x_{i+0.5} - x_{i-0.5} \). The advected field \( \varphi \) in each grid cell \( i \), is defined via the total mass in the cell \( M_i \) and position of the centre of mass \( X_i \):

\[
M_i = \int_{x_i-0.5}^{x_i+0.5} \varphi(x)dx
\]

\[
X_i = \frac{1}{M_i} \int_{x_i-0.5}^{x_i+0.5} x\varphi(x)dx
\]

Let us represent the mass distribution in each grid cell via the rectangular slab:

\[
\pi_i^n(x) = \begin{cases} 
\frac{1}{2\omega_i^n}, & |x - X_i^n| \leq \omega_i^n \\
0, & \text{otherwise}
\end{cases}
\]

where \( n \) is time step and \( \omega_i^n = \min \left( |X_i^n - x_i - 0.5|, |X_i^n - x_i + 0.5| \right) \) is distance from the centre of mass \( X_i^n \) to the nearest border of the cell \( i \). Eq. (5) defines the widest unit-volume slab that can be confined inside the cell (Figure 1) for the given centre of mass.

The advection scheme consists of a transport step and a reprojection step. At the transport step, each slab \( \pi_i \) is moved along the velocity field \( u(x) \). Advection of the slab does not change its shape within the time step \( \delta t = t_n + 1 - t_n \) but can move it anywhere over the domain or bring outside. In essence, the slab transport is replaced with advection of its mass centre, which during this time step becomes analogous to a Lagrangian particle:

\[
X_i^{n+1} = X_i^n + \int_{t_n}^{t_n+1} u(X_i^n, t) dt
\]
where $u(X^n_i, t)$ is wind speed at the mass centre location.

The original Galperin scheme employed wind at the cell mid-point $x_i$ and used explicit first-order time discretization: $u(x^n_i, t_n) = u^n_i$. Then the transported slab is given by:

$\tilde{\pi}_i^n(x) = \pi_i^n(x - u^n_i \Delta t)$

Following the transport step (7), the masses $M_k$ and centres of mass $X_k$ of the receiving set of cells $k \in K$ are updated using the transported slabs $\tilde{\pi}_i^n$:

$M_k^{n+1} = \sum_{i=1}^{N_k} \alpha_{i,k} M_i^n$

$X_k^{n+1} = \frac{1}{M_k^{n+1}} \sum_{i=1}^{N_k} \beta_{i,k} M_i^n.$

where $\alpha_{i,k} = \int_{x_i-0.5}^{x_i+0.5} \tilde{\pi}_i^n(x) dx$ and $\beta_{i,k} = \int_{x_i-0.5}^{x_i+0.5} x \tilde{\pi}_i^n(x) dx$ correspond to the mass and the first moment fractions arriving from the cell $i$ into cell $k$. The integrals are easy to evaluate due to the simple form of $\tilde{\pi}_i^n(x)$ in Eq. (5). In essence, Eq. (8) describes a mass-conservative projection of the advected slab to the computation grid.

The coefficients $\alpha_{i,0} = \int_{-\infty}^{0.5} \tilde{\pi}_i^n(x) dx$ and $\alpha_{i,1} = \int_{0.5}^{\infty} \tilde{\pi}_i^n(x) dx$ determine the transport outside the domain through the left and right borders, respectively, i.e. the scheme is fully accountable and mass-conservative since $\sum_k \alpha_{i,k} = \int_{-\infty}^{\infty} \tilde{\pi}_i^n(x) dx = 1$ for each $i$. Also, since the functions $\pi_i^n(x)$ are nonnegative, the coefficients $\alpha_{i,k}$ are nonnegative, and consequently $M_k^{n+1} \geq 0$ as long as $M_i^n \geq 0$ for all $i$. It means that the scheme is positively defined for any simulation setup: $u$, $\Delta t$, and discretization grid.
In multiple dimensions, the slabs are described by the total mass in multidimensional cell and centres of mass along each dimension. In two dimensions, an analogue of Eq. (5) will be:

$$\Pi^{n}_{i,j}(x, y) = \pi^{n}_{i,j}(x)\pi^{n}_{i,j}(y)$$

where the functions $\pi_{i,j}(x)$ and $\pi_{i,j}(y)$ depend on $X_{i,j}$ and $Y_{i,j}$, respectively. The advection step in form of (7) and the slab projection integrals (8) are then defined in 2D space.

However, a simpler procedure used in the original scheme is obtained with dimensional splitting, where the transport in each dimension is processed sequentially with the grid projection applied in-between. For an x-y split, the intermediate distribution for each row $j$ is obtained by setting:

$$\Pi^{n+1/2}_{i,j}(x, y) = \tilde{\pi}^{n}_{i,j}(x)\pi^{n}_{i,j}(y),$$

evaluating $\alpha_{i,k}$ and $\beta_{i,k}$ from $\tilde{\pi}^{n}_{i,j}(x)$ and updating $M_{i,j}$, $X_{i,j}$ and $Y_{i,j}$. Since

$$\int_{y_{i,j} - 0.5}^{y_{i,j} + 0.5} \pi_{i,j}(y)dy = 1 \text{ and } \int_{y_{i,j} - 0.5}^{y_{i,j} + 0.5} y\pi_{i,j}(y)dy = Y_{i,j}^{n},$$

the two-dimensional slab projection simplifies to:

$$M^{n+1/2}_{k,j} = \sum_{i=1}^{N} \alpha_{i,k}M^{n}_{i,j},$$

$$X^{n+1/2}_{k,j} = \frac{1}{M^{n+1/2}_{k}} \sum_{i=1}^{N} \beta_{i,k}M^{n}_{i,j},$$

$$Y^{n+1/2}_{k,j} = \frac{1}{M^{n+1/2}_{k}} \sum_{i=1}^{N} \alpha_{i,k}M^{n}_{i,j}Y^{n}_{i,j}.$$
3.3. Relations of Galperin scheme to other approaches

The Galperin scheme shares some features with other approaches (see reviews (Rood 1987) and (Lauritzen et al. 2011)). Arguably the closest existing scheme is the finite-volume approach of (Egan & Mahoney 1972), hereinafter referred to as EM72, and (Prather 1986), hereinafter P86. The main similarity between these schemes is the representation of the mass distribution via a set of slabs (rectangular in EM72 and continuous polynomial distributions in P86), one per grid cell, with the mass centre identified via the slab first moment, plus additional constraints. During the EM72 and P86 advection step, mass and the first moment are conserved, similarly to the reprojection step (8). However, this expires the similarity.

There are several principal differences between the EM72/P86 and Galperin algorithms. Firstly, in EM72 the slab width is computed via the second moment (variance) of the mass distribution in the grid cell. P86 uses the second moments to constrain the shape of the polynomials. As a result, this moment has to be computed and stored for the whole grid, and the corresponding conservation equation has to be added, on top of those for the mass and the first moment. The Galperin’s approach does not require the second moment, instead positioning the slab against the cell wall. In fact, EM72 pointed out that the second moment can be omitted but did not use the wall-based constrain in such “degenerated” scheme, which severely affected its accuracy.

Secondly, EM72 uses the movements of the slabs in adjacent grid cells to calculate the mass flows across the border. Such local consideration requires the Courant number to be less than 1: the so-called “portioning parameter” (a close analogy to the Courant number in the scheme) is limited between 0 and 1. The same limitation is valid for P86 approach. Galperin’s scheme can be applied at any Courant number and its reprojection step can rather be related to (Lin & Rood 1996).
4. Updates of the scheme in SILAM v.5

There are several features of the original scheme, which make it difficult to use in large-scale chemical transport simulations. These are listed here and the corresponding improvements are introduced in the following sub-sections.

- The scheme is formulated with zero inflow boundary conditions
- Real-wind tests have shown that the scheme has difficulties in complex wind and complex-terrain conditions, similar to EM72 (Ghods et al. 2000)
- The explicit forward-in-time advection (7) is inaccurate

In addition, the accuracy of the dimension split was increased via symmetrisation: the order of dimensions in SILAM routines is inverted each time step: x → y → z → z → y (Marchuk 1995).

4.1. Lateral and top boundary conditions

The open boundary for the outgoing masses is kept in SILAM regional simulations. The inflow into a limited-area domain is defined via prescribed concentration at the boundary (3), α = 0, β = 1, γ = ϕ. The mass coming into the domain during a single time step is equal to:

\[ M_{in}^{u} = \phi(x_{0.5}) \frac{\int M(x_{0.5})}{(u(x_{0.5})) \delta t} \]
\[ M_{in}^{l} = \phi(x_{l.0.5}) \frac{\int M(x_{l.0.5})}{(-u(x_{l.0.5})) \delta t} \]

Here \( N(u) \) is Heaviside function (= 1 if \( u > 0 \), = 0 if \( u \leq 0 \)). Assuming the locally-constant wind we obtain that \( M_{in}^{u} \) is distributed uniformly inside the slab similar to that of (5). For, e.g., the left-hand-border, the continuous form will read:

\[ \Pi_{in}^{-1}(x) = \begin{cases} \phi(x_{0.5}) \frac{\int N(u(x_{0.5}, t))}{(u(x_{0.5}, t)) \delta t}, & x \in \left[ x_{0.5}, x_{0.5} + u(x_{0.5}, t) \right] \frac{\int N(u(x_{0.5}, t))}{(u(x_{0.5}, t)) \delta t} \\ 0, & otherwise \end{cases} \]
It is then projected to the calculation grid following (8).

The top boundary follows the same rules as the lateral boundaries. At the surface, the vertical wind component is zero, which is equivalent to closure of the domain with regard to advection.

Global-domain calculations require certain care: SILAM operates in longitude-latitude grids, i.e. it has singularity points at the poles and a cut along the 180E line. For longitude, if a position of a slab part appears to be west of -180E or east of 180E, it is increased or decreased by 360 degrees, respectively. Resolving the pole singularity is done via reserving a cylindrical reservoir over each pole. The radius of the reservoirs depends on the calculation grid resolution but is kept close to 2 degrees. The calculation grid reaches the borders of the reservoirs, whose mean concentrations are used for the lateral boundary conditions:

\[
\varphi \big|_{y_1=y_{2,5.5}} = \varphi_{S\_pole} (t, z) \\
\varphi \big|_{y_1=y_{2,10.5}} = \varphi_{N\_pole} (t, z)
\]

Vertical motion in the pole cylinders is calculated separately using vertical wind component diagnosed from global non-divergence requirement.

**4.2. Extension of the scheme for complex wind pattern**

The original Galperin scheme tends to accumulate mass at stagnation points where one of the wind components is small. Similar problems were reported by (Ghods et al. 2000) for the EM72. Ghods et al. (2000) also suggested an explanation and a generic principle for solving the problem: increasing the number of points at which the wind is evaluated inside the grid cell. For application in the Galperin scheme, it can be achieved by separate advection of each slab edge instead of advecting the slab as a whole. This allows for shrinking and stretching the slab following the gradient of the velocity field. Formally, this can be written as follows.
Let’s again consider the 1-D slab that has been formed according to (5). Its edges are:

\[
X_{L,i} = X_i - \omega_i, \quad X_{R,i} = X_i + \omega_i
\]

The advection step takes the wind velocity at the left and right slab edges and transports them in a way similar to (6) with the corresponding wind velocities. The new slab is formed as a uniform distribution between the new positions of the edges:

\[
\tilde{X}^{i+1,i}(x) = \begin{cases} 
\frac{1}{X_{R,i} - X_{L,i}}, & X_{L,i} \leq x \leq X_{R,i} \\
0, & \text{otherwise}
\end{cases}
\]

Where \( \tilde{X}_{L,i}^i, \tilde{X}_{R,i}^i \) are the new positions of the slab edges at the end of the time step. This new slab is then projected following Eq. (8).

### 4.3. Changing wind along the mass-centre trajectory

The explicit advection step (7) is inaccurate and can be improved under assumption of linear change of wind inside each grid cell, with values at the borders coming from the meteo input:

\[
u(x) = u(x_{i-0.5}, t_n) \frac{(x - x_{i-0.5})}{(x_{i+0.5} - x_{i-0.5})} + u(x_{i+0.5}, t_n) \frac{(x - x_{i+0.5})}{(x_{i+0.5} - x_{i-0.5})}, \quad x_{i-0.5} \leq x \leq x_{i+0.5}
\]

Then, the trajectory equation (6) can be piece-wise integrated analytically for each slab edge.

Let’s denote \( \Delta u = u_{i+0.5} - u_{i-0.5}, \quad \Delta t = t_{n+1} - t_n, \quad \alpha = \Delta u / \Delta t \) and consider the transport starting at, e.g. \( x_{i-0.5} \). Then the time needed for passing through the entire cell, \( \Delta x = x_{i+0.5} - x_{i-0.5} \) is:

\[
T_{\text{eff}} = \log(1 + \alpha \Delta x u_i) / \alpha
\]

If \( \Delta t < T_{\text{eff}} \), the point will not pass through the whole cell and stop at:

\[
x_{\alpha} = x_{i-0.5} + u_i (\exp(\alpha \Delta t) - 1) / \alpha
\]
Applying sequentially (18) and (19) until completing the model time step \( \Delta t \), one obtains the analytical solution for the final position of the slab edges. This approach neglects the change of wind with time. However, the integration method is robust, since the linearly interpolated wind field is Lipschitz-continuous everywhere, which in turn guarantees the uniqueness of the trajectories of \( X_L \) and \( X_R \). Therefore, using the analytic solution (18) and (19), the borders of the slabs will never cross.

4.4. Reducing the shape distortions

The original scheme tends to artificially sharpen the plume edges and to gradually redistribute the background mass in the vicinity of the plume towards it (Figure 2, blue shapes). Similar “antidiffusive” distortions were also reported by P08 and by EM72 – for their scheme.

The reason for the feature can be seen from Eq. (8): if a large mass is concentrated at one of the grid cell sides, the centre of mass becomes insensitive to the low-mass part of the cell, i.e. a small mass that appears there from the neighbouring cell is just added to the big slab with little effect on its position and width.

A cheap, albeit not rigorous, way to confront the effect is to compensate it via additional small pull of the mass centre towards the cell midpoint before forming the slab for advection:

\[
\hat{X}_i'' = x_i + (X_i'' - x_i)(1-\varepsilon),
\]

where \( \varepsilon \) is smoothing factor. The adjusted mass centre \( \hat{X}_i'' \) is then used to form the slab (5).

The way this smoother works gets clearer if one notices that the Galperin scheme becomes similar to the upwind algorithm if the locations of the centres of masses are always forced to the middle of the grid cells at the beginning of time step. The upwind scheme is very diffusive, and relaxation towards it confronts the anti-diffusive features of the Galperin approach. Actual value of \( \varepsilon \) cannot be easily obtained from some optimization problem. Its
increase from 0 up to 1 gradually makes the scheme more and more diffusive, with the above
distortions becoming negligible by $\varepsilon \approx 0.08$ (Figure 2, red shapes). This behaviour and the
value were similar for various Courant numbers and tests. It is also seen from the spectral
features of the scheme in the next section – and further discussed in relation to scheme tests.

### 4.5. Analysis in frequency space

The non-linearity introduced by the coupling of cell mass and centre of mass in Eq. (8)
makes formal stability and convergence analysis after Charney et al., (1950) difficult. However, the features of the scheme can be investigated numerically following the approach of Kaas & Nielsen, (2010).

The scheme was run for 200 time steps in a 1D domain with 100 grid points. For each
wavenumber up to 25, the scheme was initialized with the corresponding sine function, and
run with Courant numbers ranging from 0.05 to 0.95 in steps of 0.05. This allowed evaluating
the spectrally resolved root mean squared error (RMSE) and, after a Fourier transform, the
spectral amplification factor (cumulative for the 200 steps) for each wavenumber. The
amplification factor quantifies the scheme’s ability to resolve the corresponding harmonics,
while the RMSE additionally includes the effect of phase errors and possible spurious modes.

Since the scheme is formulated for nonnegative concentrations, a constant background $B = 1$
is added to each waveform.

Figure 3 presents the amplification factor and RMSE for the Galperin scheme without the
smoother (panels a,d) and with it, $\varepsilon = 0.08$ (panels c,f). Furthermore, the impact of doubling
the background component to twice the wave amplitude is shown (panels b,e). In the case of
$B = 1$, the scheme without smoother shows only minor damping of all considered
wavenumbers ($k$ up to 25). The RMSE has a maximum for $k$ between 5 and 10 but stays
almost constant from $k = 10$ to $k = 25$. This shows the scheme’s ability to resolve sharp
gradients when there is no significant background. The cumulative amplifying factors for
some wavelengths exceed one but this does not imply instability since the single-step
amplifying factors fluctuate depending on the positions of centres of mass. If the integration is
continued over large number of timesteps (not shown), the solution converges to a
combination of rectangular pulses (a similar feature was mentioned in EM72 for that scheme).
Introducing the smoothing \( \epsilon = 0.08 \) resulted in strong attenuation of high-frequency
components and increased the RMSE for wavenumbers above \( \sim 10 \). Since the smoothing
factor effectively damps the fluctuations of the centres of mass, the amplification factors are
below one for all wavenumbers. Adding a background term also reduces the responsiveness
of the mass centres to newly coming amounts (see Eq (8)), which leads to a similar damping
of the higher frequencies in Figure 3c, f.
To further investigate the spectral response of the scheme, it was evaluated with a broadband
input:

\[
f(x) = \sin(2\pi x \cos(20\pi x)) + B \]

Figure 4, left panel, depicts the power spectral densities for the exact and numerical solutions
after a single revolution with CF=0.7 and 100 grid points. The corresponding solutions are
shown in the right panel. For the comparison, results are also shown for the 4th order Bott
(1989) scheme without shape preservation, and for a generic non-conservative upstream semi-
Lagrangian scheme with cubic spline interpolation.

With \( B = 1 \), all schemes capture the first spectral peaks around \( k = 10 \) and therefore resolve
most of the spectral content. Without smoother, the Galperin scheme follows the spectrum of
the true solution also resolves the spectral features around \( k = 30 \). Application of the
smoother leads to damping effect throughout the spectrum, including the spurious high-
frequency components, such as the peak at $k = 40$. This illustrates the use of the smoother for reducing over- and undershoots, as discussed in Section 4.4. Similarly to the single-harmonic tests, the situation changes in presence of a significant background ($B = 2$). Regardless of smoothing, the Galperin scheme damps the spectral peaks starting around $k = 10$, which corresponds to the reduction of amplitude visible in the numerical solution.

5. Connection of the advection scheme with other SILAM modules

Construction of the dispersion model using the Galperin advection scheme as its transport core is not trivial because all other modules should support the use of the sub-grid information on positions of the mass centres. In some cases it is straightforward but in others one can only make the module to return them undamaged back to advection.

5.1. Vertical axis: combined advection, diffusion, and dry deposition

For the vertical axis, SILAM combines the Galperin advection with the vertical diffusion algorithm following the extended resistance analogy (Sofiev 2002), which considers air column as a sequence of thick layers. Vertical slabs within these layers are controlled by the same 1-D advection, which is performed in absolute coordinates – either z- or p- depending on the vertical (height above the surface or hybrid). Settling of particles is included into advection for all layers except for the first one, where the exchange with the surface is treated by the dry deposition scheme.

The centres of masses are used but not modified by diffusion: the effective diffusion coefficient between the neighbouring thick layers is taken as an inverse of aerodynamic resistance between the centres of mass of these layers (Sofiev 2002):
The effective thickness $\Delta z_{i,j+1}$ is taken proportional to pressure drop between the centres of mases, which assures equilibration of mixing ratios due to diffusion. In the lowest layer, the dry deposition velocity is calculated at the height of centre of mass $Z_1$ following the approach of (Kouznetsov & Sofiev 2012).

The advantages of using the mass centres as the vertical diffusion meshes are discussed in details by (Sofiev 2002), where it is shown that the effect can be substantial if an inversion layer appears inside the thick dispersion layer. Then the location of the mass centre above / below the inversion can change the up / down diffusion fluxes by a factor of several times.

5.2. Emission-to-dispersion interface

Emission data is the only source of sub-grid information apart from the advection itself: location of the sources is transformed into the mass centre positions of their emission.

For point sources, the mass is added to the corresponding grid cell and centres of masses are updated:

\[
\hat{M}_{ijk} = M_{ijk} + M_{\text{em}} \\
\hat{X}_{ijk} = (X_{ijk} M_{ijk} + M_{\text{em}} X_{\text{em}}) / \hat{M}_{ijk} \\
\hat{Y}_{ijk} = (Y_{ijk} M_{ijk} + M_{\text{em}} Y_{\text{em}}) / \hat{M}_{ijk} \\
\hat{Z}_{ijk}^{\text{em}} = (Z_{ijk}^{\text{em}} M_{ijk} + M_{\text{em}} Z_{\text{em}}^{\text{em}}) / \hat{M}_{ijk}
\]

where $M_{\text{em}}$ is the mass emitted to the cell $(i,j,k)$ during the time step, $X_{\text{em}}, Y_{\text{em}}$ are the coordinates of the source in the grid and $Z_{\text{em}}^{\text{em}}$ is the effective injection height in the layer $k$, equal to middle of the layer if no particular information is available.
For area sources, the approach depends on the source grid. If it is the same as the computational one, the mass centre is put to the middle of the cell (no extra information can be obtained). If the grids are different, the source is reprojected. For each computational grid cell \((i,j)\), the centre of mass of emission is:

\[
X_{em,ij} = \frac{\int_{(x,y) \in (i,j)} x M(x,y) \, dx \, dy}{\int_{(x,y) \in (i,j)} M(x,y) \, dx \, dy}; \quad Y_{em,ij} = \frac{\int_{(x,y) \in (i,j)} y M(x,y) \, dx \, dy}{\int_{(x,y) \in (i,j)} M(x,y) \, dx \, dy}
\]

Where \(M(x,y)\) denotes the original source distribution. After that, the procedure is the same as in the case of point source (23).

5.3. Meteo-to-dispersion interface

Modifications described in section 4 require staggered wind fields, which have to be provided by the meteo pre-processor (unless they are directly available from the input data). Moreover, the pre-processor needs to ensure consistency between the flow and air density fields (Prather et al. 1987; Rotman et al. 2004; Robertson & Langner 1999). This is particularly important with the present advection scheme, since mixing ratio perturbations caused by the mass-flow inconsistency are not suppressed by numerical diffusion.

The wind pre-processing follows the idea of “pressure fixer”, which means adding a correction \(\delta V\) to the original horizontal wind field \(V_0\) such that for their sum, the vertical integral of mass flux divergence corresponds to the surface pressure tendency:

\[
\int_0^t \nabla \cdot (V_0 + \delta V) \, dp = -\frac{\partial p_s}{\partial t},
\]

where the surface pressure tendency \(\partial p_s / \partial t\) is evaluated from the meteorological input data. The correction \(\delta V\) is not uniquely determined, and SILAM adopts the algorithm of Heimann.
& Keeling (1989), where the correction term is given by the gradient of a two-dimensional potential function:

\[ \delta \nabla = \nabla \psi(x, y). \]

Substituting (26) into (25) yields a Poisson equation for \( \psi(x, y) \), which is solved to subsequently recover \( \delta \nabla \). The obtained correction flux is then distributed within the column proportionally to the air mass in each layer, ending up with the corrections to the horizontal winds. The vertical wind is then evaluated in each column to enforce the proper airmass change in each cell.

5.4. Chemical module interface

This interface is implemented in a very simple manner: the mass centres are not affected by the transformations. Chemical module deals exclusively with concentrations in the grid cells. The newly created mass is added to the existing one, thus accepting its centre position in the cell. If some species did not exist before the transformation the new mass centre is put to the middle point of the cell.

6. Testing the Galperin advection algorithm

6.1. Standard tests

A set of basic tests and comparison with some classical approaches has been presented by Galperin (1999) and P08 for the original scheme, along with Bott, Holmgren, and several other schemes. Their main conclusions were that the scheme is very good for sharp-edge patterns: in particular, it transports delta functions without any distortions. It had, however, issues with long slopes, smooth shapes, etc, where the tendency to gradually convert them to a collection of rectangles was noticeable.
Addressing these concerns, tests used during the scheme improvements and implementation in SILAM included puff-over-background, conical and sin-shaped peaks and dips, etc (some examples are shown in Figure 2); divergent 1-D high-Courant wind test in 1-D divergent wind field (Figure 5), constant-level background field in eight vortices with stagnation points (Figure 6), and rotation tests for various shapes (Figure 7).

The scheme stays stable at arbitrarily high Courant numbers and handles the convergence and divergence of the flows (Figure 5).

Transport and rotation tests of the improved scheme maintain low distortions of the shapes: the $L_2$ norm of the error varies from 0.1% up to 3.8% of the initial-shape norm – for the most challenging task in Figure 7. The effect of the improvements in comparison with the original scheme is demonstrated in Figure 2, where the blue contours show the results of the original scheme. In particular, application of the smoothing Eq. (20) reduced the distortions of smooth shapes (red curves), largely resolving the concerns of P08: Figure 2b presents the same test as one of the P08 exercises. However, the smoother also leads to a certain numerical viscosity of the scheme, so its use in problems requiring non-diffusive schemes (e.g., narrow plumes from accidental releases) should be avoided.

The test with eight vortices was difficult for the original scheme (Figure 6a) due to its insufficient sub-grid resolution but the improvements (15) - (16), section 0, resolved the problem (Figure 6b). This refinement is instrumental for complex-topography domains.

6.2. Global 2-D tests

Performance of Galperin’s advection scheme in global spherical domain was assessed with the collection of demanding tests of (Lauritzen et al. 2012). The cases are designed to evaluate the accuracy of transport schemes at a wide range of resolutions and Courant numbers. The tests used a prescribed non-divergent 2D velocity field defined on a sphere and consisting of...
deformation and rotation, so that the initial concentration pattern is reconstructed at the end of
the test, $t=T$, providing the exact solution $\phi(t=0) = \phi(t=T)$.

Four initial concentration distributions were used (Figure 8): “Gaussian hills” with unity
maximum value, “cosine bells” with background of 0.1 and maxima of 1, “slotted cylinders”
– rough pattern with 0.1 background and 1 maximum level, and “correlated cosine bells” –
distribution obtained from “cosine bells” with a function:

$\phi_{\text{ccb}}(t) = 0.9 - 0.8 \phi_{\text{cb}}^2$

The tests were run with SILAM on a global regular non-rotated lon-lat grid, with $R=6400$ km
and $T=12$ h. Spatial resolutions were: 6, 3, 1.5, 0.75, 0.375, and 0.1875 degrees, each run with
mean Courant numbers of $-5.12$, $-2.56$, $-0.85$ (for 6° grid they correspond to the model time
step of $T/12=1$ h, $T/24=30$ min, and $T/72=5$ min), total 18 runs for each initial pattern.

Examples of the most challenging runs with slotted cylinders at $t=T/2$ and at $t=T$ are shown in
Figure 9 and Figure 10, respectively. The corresponding error fields are collected in Figure 11
as decimal logarithms of the absolute difference between the corresponding field in Figure 10
and the slotted-cylinder initial shape of Figure 8. The main complexity of the test was in
reproducing the very tiny sharp-edge structures obtained from the cylinder cut at $t=T/2$ – and
then returning them back by $t=T$. The pictures, together with the error field at $t=T$ (Figure 11)
show that already 24 time steps allow the scheme to make the shape recognisable (3°, C=5.12
pattern), whereas 48 time steps allow for main details to show up. Expectedly, certain
deviations at the cylinder edge remain at any resolution – as visible from the error fields.

Deviation of the resulting field $\phi_L = \phi(t=T)$ from the initial shape $\phi_0 = \phi(t=0)$, was
considered in three spaces: $L_2$, $L_\infty$, $L_1$. The corresponding distance metrics are defined as
follows:
where $\mathcal{S}[]$ is an area-weighted sum over latitude and longitude. The values of these three
metrics for all model runs are presented in Figure 12. The main interest of these curves is that
they show the rate of the scheme convergence (straight grey lines correspond to the first- and
second-order convergence rates). Expectedly, the rates depend on the transported shape (the
smoother the shape the faster convergence) and on the norm used. Thus, the scheme
converges in $L_1$ faster than in $L_2$, whereas in $L_\infty$ no convergence in case of sharp edges is an
expected result. The rate in the $L_2$ norm is in-between the first- and the second order, whereas
in $L_4$ it is close to the latter one.

Advection should also keep the local ratio of the tracer’s concentrations. Such ratio between
“cosine bells” and “correlated cosine bells” was calculated at $t=T/2$ and $t=T$. Since these
initial patterns are related by eq. (27), the concentration fields during the tests should maintain
the same relation. The scatter plots of the concentrations in these two tests give an indication
on how the ratio is kept. Ideal advection would keep all points on a line given by Eq. (28).

The results of the tests for $t=T/2$ are shown in Figure 13, where the results with and without
the smoother in Eq. (20) are presented. The smoother improves the scheme linearity, i.e. it can
be recommended to chemical composition computations, which usually also tolerate some
numerical viscosity.

### 6.3. Global 3-D test with real wind

Testing the scheme with real-wind conditions has one major difficulty: there is no accurate
solution that can be used as a reference. An exception is simulations of constant-mixing-ratio
3D field, which, once initialised, must stay constant throughout the run. Deviation from this
constant is then the measure of the model quality. Such test verifies both the scheme and the
meteo-to-dispersion interface, which has to provide the consistent wind fields.
The constant-vmr test was set with winds taken from ERA-Interim archive of ECMWF, for an arbitrarily selected month of January 1991 (Figure 15). The model was initialised with vmr = 1 and run with 3° of lon-lat resolution and time step of 30 minutes (max Courant number exceeding 13 in the stratosphere and reaching up to 2-3 in the troposphere). The model top was closed at 10Pa, which corresponds to the top level of the ERA-Interim fields. The procedure described in the section 5.3 was used to diagnose the vertical wind component. The results of the test are shown in Figure 15, which depicts the model state after 240 hours of the run, panel a) showing the boundary-layer vmr, and panel b) presenting it in the stratosphere. The zonally-averaged vertical cross-section is shown in panel c. Green colours in the pictures correspond to less than 1% of the instant-field error.

An important message is that the limited distortions about 1-2% are visible in a few places but they are not related to topography, rather being associated with the frontal zones and cyclones. The comparatively coarse spatial and temporal resolution of the test makes the associated changes of the wind quite sharp, so that the dimension-split errors start manifesting themselves. Smoother flows in the stratosphere posed minor challenges for the scheme. The L2-error (not shown) is approximately proportional to the model time step.

### 7. Discussion

The presented SILAM v.5 transport module is based on semi-Lagrangian advection scheme of M. Galperin with subgrid information available through the positions of centres of masses. It poses certain challenges in implementation. Firstly, one has to organise the sub-grid information use and transmission between the advection and other model units. Secondly, the scheme requires storage of four full fields for each transported species (mass and moments) and care should be taken to maintain an efficient exchange between the processors and the

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**Deleted:** Figure 12

**Deleted:** Efficiency of Galperin advection scheme

Evaluation of the scheme efficiency is always very difficult as it depends on computer, parallelization, compiler options, etc. Nevertheless, some basic characteristics of the scheme have been deduced from comparison of the simple cases for classical schemes (Galperin 2000). It was shown to be 2.3 ∼ 15 times faster than, e.g., Bott scheme depending on implementation, specific test, etc.

For the L14 tests, the run with 0.75 degree resolution and 120 time steps (took 47 seconds) can be related to performance of HEL and CSLAM schemes, which were tested against the same collection by (Kaas et al. 2013). With all ambiguity of the runtime parameter, it took about 200 seconds for HEL and 300 seconds for CSLAM, i.e., about 4 and 6 times longer than for SILAM. Our tests were run on a simple notebook with dual-core hyperthreaded Intel Core i5-540M CPU and 4 GB of RAM (Intel Linpack benchmark). We used GNU compiler with -O3 optimization without parallelization, which corresponds to the settings of (Kaas et al. 2013).

In SILAM applications, advection is parallelised using the shared-memory OMP technology, whereas the MPI-based domain split is being developed. The OMP parallelization is readily applicable along each dimension, thus exploiting the dimensional split of the advection scheme. For MPI, care should be taken to allow for a sufficient width of the buffer areas to handle the Courant > 1 cases.

The original scheme was formulated for the bulk mass of all transported tracers, thus performing the advection step for all species at once: the tracer's mass in the slab definition (5) was the sum of masses of all species. This algorithm is faster than the species-wise advection and reduces the number of the moments per dimension down to one regardless the number of tracers. It can also be useful in case of strong chemical bonds between the species in coarse-grid and sub-optimal Courant number: as seen from Figure 11, such runs can have noticeable non-linearity between the tracer concentrations. The bulk advection does not have the non-linearity problem instead loses much of its quality if the species have substantially different lifetimes in the atmosphere, are emitted from substantially different sources or otherwise decorrelated in space.
computer memory. Thirdly, possibility to run with high Courant numbers and MPI parallelization via horizontal domain split can be utilised only if the MPI split allows for sufficient buffer zones. Finally, the better performance of the advection at Courant number greater than 1 challenges the implementation of other modules, first of all, chemistry and emission. Indeed, introduction of emitted mass once per long time step would result in a broken plume unless the mass is spread downwind over the corresponding distance. Similar problems show up in chemical transformation calculations. At present, the actual SILAM applications are performed with Courant close to but mostly smaller than one to avoid such problems.

The above challenges are mostly technical and their solution allows the scheme to demonstrate strong performance with low computational costs.

In particular, by attributing the release from point source to its actual location one can reduce the impact of the common problem of Eulerian models: point release is immediately diluted over the model grid cell. This substantially improves the transport though does not solve the problem completely: (i) the chemical module still receives the diluted plume concentration, (ii) the slab size in case of the source near the centre of the grid cell will still be as large as the grid cell itself. A more accurate solution would be the plume-in-grid or similar approaches, which is being built in SILAM. Another example of the sub-grid information usage is utilisation of full meteorological vertical resolution to calculate effective values of meteorological variables for thick dispersion layers (Sofiev 2002).

The model can operate at any Courant number (Figure 5). Its time step is limited not by grid cell size but by a spatial scale of the wind-shear field, i.e. has to satisfy much less restrictive Lipshitz criterion, which relates spatial and temporal truncation errors (Pudykiewicz et al. 1985). It follows from the advection step (6) and the reprojection step (8), which do not
restrict new positions of the slabs: they can find themselves anywhere in the grid or outside it after the time step is made.

SILAM heavily relies on such features of Galperin’s scheme as mass conservation and accountability: the scheme provides complete mass budget including transport across the domain boundaries. In particular, nesting of the calculations is straightforward and does not need the relaxation buffer at the edges of the inner domain: the inflow through the boundaries is described by the same slabs as the main advection. The scheme is also shape-preserving – in the sense this term is used by L14, – i.e. it does not result in unphysical solutions, such as negative mixing ratio. Some distortions are still possible (Figure 2), which can be reduced by the smoother described in section 4.4, eq. (20).

7.1. Standard advection tests

Evaluating the Galperin’s scheme with the simple tests (Figure 2 - Figure 7), one can point out the known issues of the classical schemes resolved in Galperin’s approach: high-order algorithms suffer from numerical diffusion, oscillations at sharp gradients (require special efforts for limiting their amplitude), high computational costs and stringent limits to Courant number. None of these affect the Galperin scheme.

The main issue noticed during the implementation of the original scheme was the unrealistically high concentrations near the wind stagnation points. Thus, the concentration pattern at the test Figure 6a resembles the situation of divergent wind field. However, it is not the case: the 2D wind pattern is strictly solenoidal. The actual reason is insufficient resolution of the advection grid: one centre of mass point is not enough if spatial scale of the wind variation is comparable with the grid cell size. Tracking the edges of the slab rather than its centre resolves the problem (Figure 6b).
The other challenging tasks for Galperin’s algorithm were those with smooth background and soft gradients, a frequent issue for semi-Lagrangian schemes, which is easily handled by more diffusive approaches. This feature was visible in the P08 tests where the scheme noticeably distorts the Gaussian and conical plumes. For the puff-over-background pattern, the scheme makes a single low-mass dip in the vicinity of the puff, which receives this mass (Figure 2).

From formal point of view, the scheme does not conserve the higher moments inside the grid cell, which becomes a problem when the pattern changes at a spatial scale shorter than the grid cell size. The smoothing step (20) may be advised despite it has no rigorous ground and, as in L14 evaluation of other schemes, may damage some formal quality scores (adding this step introduces numerical viscosity - Figure 2).

7.2. Global 2D and real-wind advection tests

The application of the scheme to highly challenging tests of (Lauritzen et al. 2012) allowed its evaluation in a global 2-D case and comparison with the state-of-the-art schemes evaluated by L14 and (Kaas et al. 2013).

Performing these tests with different spatial and temporal resolutions, as well as Courant numbers, suggested that the scheme has an “optimal” Courant number for each spatial resolution where the error metrics reach their minimum, so that the increase of temporal resolution is not beneficial. Indeed, in Figure 12, the low-Courant runs are by no means the most accurate. This is not surprising: for an ideal scheme, increasing the grid resolution and reducing the time step should both lead to gradual convergence of the algorithm, i.e. the error metrics should reduce. For real schemes, higher temporal resolution competes with accumulation of the scheme errors with increasing number of steps. Convergence in L14 tests was still solid for all fixed-Courant-number series (Figure 12) but excessive temporal resolution (specific for each particular grid cell size) was penalised by higher errors.

Similarly, the most-accurate representation of the correlated patterns is obtained from the runs...
with the intermediate Courant numbers (Figure 13). This seems to be a common feature: the same behaviour was noticed by L14 for several schemes.

High optimal Courant numbers, however, should be taken with care. For L14, the smooth wind fields reduced the dimension-split error and made the long time steps particularly beneficial.

It is also seen (Figure 11) that the best performance, in case of near-optimal Courant, is demonstrated by the high-spatial-resolution simulations, which have reproduced both the sharp edges of the slotted cylinders, the flat background and the cylinder’s top planes. The scheme demonstrated convergence rate higher than one for all metrics and all tests with smooth initial patterns. Even for the most stringent test with the slotted cylinders, the scheme showed the first-order convergence rate in the L1 norm (Figure 12).

Among the other features of the solution, one can notice a certain inhomogeneity of the background field away from the transported bodies. The error is very small ($< 10^{-4}$) for high-resolution cases (Figure 11) and $< 0.1\%$ for inexpensive setups, such as $\Delta \lambda = 0.75$, $C=2.56$. For coarser resolutions, it grows. The inhomogeneity also grows with Courant number, which is opposite to decreasing error of representation of the shapes themselves. The issue originates from the dimension-split error in polar areas, where the spatial scale of wind change becomes comparable with the distance passed by the slabs within one time step.

Similar non-monotonicity of background is visible for some schemes tested by L14.

Unfortunately, no error fields are given there but Figs. 7-10 there are comparable with our Figure 9 (results without smoother). With few exceptions (schemes TTS-I and LPM, notations of L14), all algorithms manifested such patterns unless filters are applied. For some schemes (SFF-CSLAM3, SFF-CSLAM4, UCISOM-CS, CLAW, and CAM), these inhomogeneities are visible also for the tests with shape-preserving filters. One should note however that the
0.1 level, which distinguishes between the two violet colours in Figure 9 and Fig. 7-10 of L14, corresponds to the background level in slotted-cylinder test. As a result, even a very small deviation leads to appearance of such shapes in the plots (note stripes in background of Figure 8).

Comparing the so-called “minimal resolution” threshold for \( L_2 \) norm of cosine bells to reach 0.033 (Figure 3 of L14) for SILAM was about 0.75°, which puts it in the middle of that multi-model chart (specific place depends on whether the shape preservation is considered or not).

Another criterion can be the optimal convergence of \( L_2 \) and \( L_{\infty} \) norms for Gaussian hills:

about 1.7-1.8 for SILAM – is again in the middle of the L14 histograms, in the second half if the unlimited schemes (without shape-preservation filters) are considered and in the first half if the unphysical negative concentrations are suppressed (since the Galperin advection is strictly positively defined, no extra efforts needed to satisfy this requirement).

Interestingly, the L14 tests were limited with 3° as the coarsest resolution, and it was pointed out that the schemes start converging only when a certain limit, specific for each scheme, is reached. The SILAM results show similar behaviour only for the lowest Courant number (red lines in Figure 12), which indeed required appropriate resolution to start working. Higher Courant setups were much less restrictive (the errors decrease with growing resolution also for coarse-grids) and, as already pointed out, often worked better than the low-Courant runs (similar to many L14 schemes).

The scheme demonstrated limited distortion of pre-existing functional dependence – see the cosine bells and correlated cosine bells tests, Eq (27) (Figure 13). Formal scores suggested by (Lauritzen et al. 2012a) calculated for the Galperin scheme are shown in Figure 14. Notations are: \( L_\infty \) “overshooting”, describes the values that fell outside the rectangular [0.1:1] (Figure 13), \( L_\infty \) “shape-preserving unmix”, describes the values inside that rectangular but outside the...
“lens” formed by its diagonal \((0.1, 1) – (1, 0.1)\) and the curve, and \(l_c\), “real mixing”, describes the values inside the “lens”. Comparison with L14 (Fig. 15, middle panel) shows that the Galperin scheme outperforms CLAW, SLFV-ML, SLFV-SL, and all setups of ICON schemes, being close to CAM-SE, MIPAS, and HOMME, and trailing behind the runs with CSLAM, HEL, SFF, and UCISCOM schemes.

A peculiarity of the mixing diagnostic scores is that they are significantly affected by the background areas far from the advected bells, which occupy only a small fraction of the domain (Figure 8). As a result, small background fluctuations discussed above in application to slotted cylinders (see error field in Figure 11) contribute significantly to the mixing diagnostic scores too. In particular, the high-Courant simulations, which accurately reproduce the bells themselves (the dots are close to the curve in the scatter-plots Figure 13), still show poor formal scores due to non-zero width of the cloud near the location \((0.1, 1)\), where all background dots should arrive. This issue contributes most-significantly into “overshooting” part of the error, but also to other two components.

Expectedly, the smoother improves the mixing diagnostic scores, mainly affecting the representation of the bells themselves (Figure 13). This is in contrast with the schemes tested in L14, where the shape-preservation filters mostly removed the penalty for overshooting the background but rarely improved the other two components, sometimes worsening them.

Following the conclusions of the section 4.4 and 1D tests, we used the smoothing factor of 0.08, which is a compromise between the scheme diffusivity and distortion reduction. As a result, some non-linearity exists also in the smoothed solution. The test showed that simple increase of temporal resolution leads to increase of the number of steps and related reprojections, which then worsen the representation of the bells – but improved the background field by reducing the dimension-split errors. A synchronous rise of the resolution
in time and space with the same Courant number (columns in Figure 13) showed better results for higher-resolving setups.

Further investigating the flat-field behaviour in complex wind patterns, the simulations with the constant-vmr initial conditions (Figure 15) were performed, showing that the model has no major problem in keeping the homogeneous distribution: deviations do not exceed few % with no relation to topography. The existing ups and downs of the vmr are related to cyclones and atmospheric fronts, which challenge the dimension-splitting algorithm rather than the core 1D advection (it transports the homogeneous field perfectly – no distortion was found after \(10^5\) steps regardless the Courant number). Increasing the resolution leads to lower “unmix” of the pattern (not shown). This experiment refines the “optimal-Courant” recommendation of the L14 test, which had smoother wind fields and, consequently, higher optimal Courant number. For real-life applications, especially with coarse grid, it may be necessary to choose time step short enough to ensure comparable levels of time- and space-wise truncation errors (Pudykiewicz et al. 1985). This case also argues for developing the 2D implementation of Galperin scheme, which would eliminate the horizontal dimension split.

7.3. Where to use the smoother

When deciding of whether to apply the smoother Eq (20), one has to keep in mind that Galperin scheme is always positively defined and does not need a shape-preserving filter to provide a “physically meaningful” solution, i.e. without negative values. It is free from this caveat. The purpose of the smoother is only to reduce the non-linear distortions of fields.

The smoother has both positive and negative impact on the scheme performance. Among the positive ones are: (i) it damps the distortions of smooth shapes and gradients (section 4.4), (ii) it reduces the amplification factor precluding it from exceeding one even for few time steps (section 4.5), (iii) it reduces the unmixing problem (Figure 14). Its negative features are: (i) the obtained solution is diffusive (section 4.4), (ii) moderate and high frequencies in the
solution spectrum are damped (section 4.5). (iii) formal scores and convergence rates are lower in some tests (sections 6.2 and 7.2). The smoother has little impact on background inhomogeneity.

Most of positive and negative features coincide with impact of shape-preserving filters (e.g., L14), despite the different idea and formulations.

Since the smoother computational cost is negligible, one can decide whether to apply it depending only on the problem at hands. Strict interconnections between the species, smooth patterns and tolerance to diffusion form a case for the smoother. Conversely, sharp plumes over zero background (e.g., accidental release case) argue against it.

The smoother impact grows monotonically with its parameter $\epsilon$. Numerous tests showed that the distortions and above-one amplification factor essentially disappear at $\epsilon \sim 0.08$, where the diffusivity also becomes significant. This value appeared stable with regard to Courant number and setup of the tests.

7.4. **Efficiency of Galperin advection scheme**

Evaluation of the scheme efficiency is always very difficult as it strongly depends on the algorithm implementation, but also on computer, parallelization, compiler options, etc. Nevertheless, basic characteristics of the scheme can be deduced from comparison of its original version with several classical schemes made by Galperin (2000). It included, in particular, EM72 and Bott, which appeared >5 and >3 times slower, respectively. Comparison with another implementation of Bott routine by (Petrova et al. 2008) showed 7-15 times difference depending on tests. The updated scheme version, however, is bound to be heavier.

It is also worth putting it in line with modern approaches.

In this section, the efficiency of the updated Galperin scheme is evaluated from several points of view: (i) the scalability with regard to the number of transported species, spatial and
temporal resolution, specifics of the problem at hands, (ii) comparison with “standard implementation” of the Bott algorithm and semi-Lagrangian scheme, (iii) comparison of the runtime in the L14 tests with HEL and CSLAM schemes.

7.4.1. SILAM run time vs number of species, temporal and spatial resolution

The scalability of the scheme and the whole SILAM model was tested in real-wind global simulations for arbitrarily taken three days (15-17.5.2012). The reference run was set with 0.5° resolution, 6 vertical layers, time step of 30 minutes, and one aerosol species. Two types of emission were considered: an artificial one-hour-long source filling-up the whole 3D domain, and the SILAM-own wind-blown dust emission model, which created dust plumes from sandy areas of Sahara. Vertical diffusion, which is coupled with vertical 1D advection, was turned off for artificial source test but turned on for dust source in order to allow the model to quickly populate the upper layers of the domain. Then, the number of aerosol species, spatial and temporal resolutions were repeatedly doubled (one change at a time).

The model was run in a single-processor mode but compiled with O3 optimization and OMP code pre-processing. Runs were made in a notebook with Intel Core i7 processor and repeated in a workstation with Intel Xeon E5. The scaling differed by 10-20%, which was considered to be negligible.

The results (Figure 16) highlight the scalability of the scheme and its implementation in SILAM. The species-unrelated time of horizontal 2D advection (Figure 16a, offset in regression line) is ~30% of a single-species computation time (represented via slope). This “overhead” is, in fact, the transport-step integrals Eq. (17) - (19), which are computed only once and used for all species. Higher overhead of the vertical advection is due to necessity to handle the uneven vertical layers, which makes it scaling just 20% better than 2D horizontal one. It also has larger species-independent overhead.
With chemical module turned off, advection constitutes ~85% of the total model run time.

Since the scheme operates with the source grid cells, it can check that $M_i^n > 0$ before going into computations, which gives a very substantial speed-up in case of limited-volume plumes (Figure 16b). In Saharan dust run, the horizontal advection time is about twice lower, whereas the vertical advection, even together with diffusion, becomes all but negligible, owing to efficient filtering of zero-columns in comparison with lon- or lat-stripes.

A faster-than-proportional growth of the horizontal advection time with increasing resolution (Figure 16c, normalised run time) is a result of growing Courant number: for 4-times smaller grid cell (0.25° lon-lat resolution) the time step of 30 minutes means $C \gg 1$ over large part of the domain. As a result, transport integrals Eq. (17) - (19) have to be analysed over longer paths. Still, the growth is much smaller than the cost of 4-fold reduction of time step, which makes the high-C computations attractive. Vertical advection is not affected and its time is proportional to the number of columns to analyse.

Time spent by advection is practically proportional to the temporal resolution (Figure 16c), i.e. it follows the number of times the advection is computed in the run.

**7.4.2. Comparison with efficiency of other schemes**

Comparison with other schemes is arguably the most uncertain part of the exercise: the scheme efficiency is strongly dependent on the quality of the implementation (note the different results for Bott scheme obtained by Galperin, (2000) and (Petrova et al. 2008)). To obtain reproducible results, we made this comparison against the “standard implementation” of the Bott code available from internet (http://www2.meteo.uni-bonn.de/forschung/gruppen/tgwww/people/abott/fortran/fortran_english.html, visited 28.09.2015). Since our code is also available (see section 8), this comparison is reproducible.
The test with $10^4$ time steps, 2000 gridpoints in 1D periodic grid, Courant number = 0.1, and one species took: 0.92 sec for Galperin scheme (~0.3 sec for cell border advection, ~0.6 sec for slab reprojection) and 0.85 sec for Bott scheme. This confirms the expectation that the updates of the Galperin scheme from its initial version about-tripled its run time, which is now similar to that of the Bott scheme. However, the Galperin scheme still scales better with the number of species: as shown in previous section, only reprojection is multiplied with the number of species, whereas Bott scheme does not have such saving possibility.

The above numbers should be considered as indicative only since the environment for the tests was completely artificial: the schemes were used as a stand-alone code applied in 1D space. Galperin scheme needed only one moment instead of three, which would be the case of 3D advection. Despite very limited extra computations, this would still raise the memory exchange. The Bott scheme was taken without shape-preservation filter, which would be needed for any real-life applications.

The tests were also made for our own implementation of the semi-Lagrangian scheme (took ~50% longer than the above timing) but its efficiency was not carefully verified.

The L14 tests allowed rough benchmarking of the SILAM implementation of the scheme in 2D tasks. In particular, the run with 0.75° resolution and 120 time steps can be related to performance of HEL and CSLAM schemes, which were tested against the same test collection by (Kaas et al. 2013). Extrapolating the charts of Fig.13 of (Kaas et al. 2013) to one species (the range given there is 2-20 species), the test takes about 190 seconds for HEL and 300 seconds for CSLAM but only 47 seconds for SILAM, i.e. the difference was about 4 and 6 times, respectively.

Formal benchmarks of the computers, the main uncertainty in this comparison, are essentially the same: Kaas et al used Intel Core2 Duo E6550 processor (Intel Linpack 20 GFlops).
http://www.techpowerup.com, visited 8.10.2015. Our tests were run on a simple notebook
with mobile Intel Core i5-540M Duo (Intel Linpack 18.5 GFlops). These CPUs were also
compared in http://www.cpubenchmark.net (visited 8.10.2015), which also put them within
20% from each other, albeit i5-540M was put forward. Memory bandwidth of our notebook,
as always for compact computers, was modest: 7.2 GB/s (STREAM test).
http://www.cs.virginia.edu/stream/ref.html accessed 5.10.2015). We used GNU compiler with
–O3 optimization without parallelization, similar to Kaas et al. (2013).

7.4.3. Further boosting the scheme efficiency: parallelization
In SILAM applications, advection is parallelised using the shared-memory OMP technology,
whereas the MPI-based domain split is being developed. The OMP parallelization is readily
applicable along each dimension, thus exploiting the dimensional split of the advection
scheme. For MPI, care should be taken to allow for a sufficient width of the buffer areas to
handle the Courant > 1 cases.
The original scheme was formulated for the bulk mass of all transported tracers, thus
performing the advection step for all species at once: the tracer’s mass in the slab definition (5)
was the sum of masses of all species. This is much faster than the species-wise advection
and reduces the number of the moments per dimension down to one regardless the number of
tracers. It is also useful in case of strong chemical binds between the species because the bulk
advection keeps all pre-existing relations between the species. However, transport accuracy
diminishes if the species have substantially different life times in the atmosphere, are emitted
from substantially different sources or otherwise decorrelated in space.
8. Code availability

SILAM is a publicly available model. Our experience shows however that its successful application critically depends on the user’s modelling skills and understanding of the model concepts. Therefore, SILAM is available on-request basis from the authors of this paper, who also provide support in the initial model installation and setup. The model description, operational and research products, as well as reference documentation, are presented at


9. Summary

Current paper presents the transport module of System for Integrated modeLling of Atmospheric coMposition SILAM v.5, which is based on the improved advection routine of Michael Galperin combined with separate developments for vertical diffusion and dry deposition. The corner stone of the advection scheme is the subgrid information on distribution of masses inside the grid cells, which is generated at the emission calculation stage and maintained in a consistent way throughout the whole model, including chemical transformation, deposition, and transport itself. This information, albeit requiring substantial storage for handling, allows for accurate representation of transport.
The scheme is shown to be particularly efficient for point sources and sharp gradients of the concentration fields, still showing solid performance for smooth patterns. The most challenging task was found to be the puff-over-plain test, where the scheme showed noticeable distortions of the concentration pattern. Application of a simple smoother efficiently reduces the problem at a cost of non-zero viscosity of the resulting scheme. Advanced tests and comparison with state-of-art algorithms confirmed the compromise between the efficiency and accuracy. SILAM performance was fully comparable with the other algorithms, outperforming some of them.

Among the future developments, implementation of the scheme in 2D space and replacement of the smoother with extensions of the core advection algorithm, are probably the most pressing ones.

10. Acknowledgments

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Deleted: introduction of physically grounded horizontal diffusion procedure


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Figure captions

Figure 1. Advection step of the scheme of M. Galperin

Figure 2. Shape preservation tests: a) step, b) triangle peak, c) sin-shaped dip, d) sin-shaped peak. Sequential positions are shown, 'r' denotes the scheme without smoother, 'r_diff' - with it. The legend includes the number of times steps made. Wind is from left to right, Courant = 0.4.

Figure 3. Spectral analysis for 1D. Panels a)-d) Amplification Factor (AF) and RMSE, respectively for the Galperin scheme without smoother; b)-c) AF and RMS for Galperin scheme with large background; c), f) AF and RMSE for Galperin scheme with smoother ϵ=0.08.

Figure 4. Example of input and output spectra for broadband input to the advection schemes with zero and nonzero background level. Left panels: exact and numerical solutions. Right panels: power spectrum densities initially and after one revolution. Top: B = 0, bottom: B = 1.0.

Figure 5. Linear-motion tests with a constant-release point source at X0 and varying wind speed along x-axis. Upper panel: Courant number, lower panel: concentration [arbitrary unit]. Wind blows from left to right.

Figure 6. Test with eight non-divergent 2-D vortices. Left panel: test of the original scheme (5)-(7), time step 8; right panel: improved scheme (15)-(16), time step 50. Both tasks were initialised with constant value 0.4, also used as boundary conditions. Without smoother.

Figure 7. Double-vortex rotation tests: a rectangular split between the vortices (upper panels); three single-cell peaks and two connected rectangles (middle panels); sin- and cone-shaped surfaces (lower panels). A series of time steps shown in the left panels, except for the low panel (shown t=361). Right panels: error field after 1 full revolution (obs 10-fold more sensitive scale and relative L2 norm given above each plot). Max Courant ~ 1.5. Grid dimensions = 400 x 200. Without smoother.

Figure 8. Initial shapes of the puffs for the 2-D global test on the sphere.

Figure 9. Half-period (t=T/2) shapes for the 2D global test with slotted cylinders for different spatial and temporal resolutions. Without smoother.

Figure 10. Final shapes (t=T) for the 2-D global tests with slotted cylinders for different spatial and temporal resolutions. Without smoother.

Figure 11. The error fields for the final shapes of Figure 10 as compared with slotted cylinder initial shape in Figure 8. Without smoother.

Figure 12. Dependence of the performance metrics L1, L2, and L∞ for the spherical 2D tests with initial shapes of Figure 8. Dashed straight lines mark the slope for the first and second order of convergence. Without smoother.

Figure 13. Mixing preservation test for cosine bells and correlated cosine bells (27) at t=T/2. Each two lines show the tests without (upper line) and with (lower line) smoother (28).
**Figure 14.** A histogram of mixing diagnostic (stacked) for the same resolutions, Courant number and smoother factor as in Figure 13. Metrics are (see text and (Lauritzen et al. 2012) for more details): $I_r$ is “real mixing”, $I_u$ is “range-preserving unmixing”, $I_o$ is “overshooting”. Values are relative to the reference CSLAM performance in L14 tests. Picture is comparable with panel b) of Fig. 15 in L14.

**Figure 15.** Constant-vmr test with real-wind conditions after 122 hrs. a) vmr within boundary layer, b) vmr above the tropopause, c) zone-average vertical cross-section of vmr. Without smoother.

**Figure 16.** Scalability of Galperin advection scheme and SILAM model. Panel a) Full-grid runtime for different number of species, b) sparse-plume run time for different number of species, c) full-grid run time for varying horizontal grid resolution, d) full-grid run time for varying time step.
Figure 1. Advection step of the scheme of M. Galperin

\[ u \]

Formation of slabs

\[ X_i^n = X_i^{n+1} + \omega \delta t \]

Move with wind

\[ \hat{X}_i^n = X_i^n + u \delta t \]

Reprojection

\[ X_i^{n+1} = X_i^{n+1} \]

\[ t_{n+1} = t_n + \delta t \]

\[ \hat{X}_i^* \rightarrow (X_i^{n+1}, X_{i+1}^{n+1}) \]
Figure 2. Shape preservation tests: a) step, b) triangle peak, c) sin-shaped dip, d) sin-shaped peak. Sequential positions are shown. ‘r’ denotes the scheme without smoother, ‘r_diff’—with it. The legend includes the number of times steps made. Wind is from left to right, Courant = 0.4.
Comment [MS1]: May be, scale for the application factor should contain 1 as a edge.
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Deleted: Figure 11. Linearity test for cosine bells and correlated cosine bells (26) at $t=T/2$. Each two lines show the tests without (upper line) and with (lower line) smoother (20).
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