A comparison of semi-Lagrangian discontinuous Galerkin and spline based Vlasov solvers in four dimensions

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Abstract

The purpose of the present paper is to compare two semi-Lagrangian methods in the context of the four-dimensional Vlasov–Poisson equation. More specifically, our goal is to compare the performance of the more recently developed semi-Lagrangian discontinuous Galerkin scheme with the de facto standard in Eulerian Vlasov simulation (i.e. using cubic spline interpolation). To that end, we perform simulations for nonlinear Landau damping and a two-stream instability and provide benchmarks for the SeLaLib and sldg codes (both on a workstation and using MPI on a cluster).

We find that the semi-Lagrangian discontinuous Galerkin scheme shows a moderate improvement in run time for nonlinear Landau damping and a substantial improvement for the two-stream instability. It should be emphasized that these results are markedly different from results obtained in the asymptotic regime (which favor spline interpolation). Thus, we conclude that the traditional approach of evaluating numerical methods is misleading, even for short time simulations. In addition, the absence of any All-to-All communication in the semi-Lagrangian discontinuous Galerkin method gives it a decisive advantage for scaling to more than 256 cores.

Keywords: semi-Lagrangian discontinuous Galerkin method, Vlasov–Poisson equation, semi-Lagrangian methods, performance comparison, parallelization

1. Introduction

Many problems in computational plasma physics require the solution of kinetic equations. Due to the high dimensionality of these models, particle methods have been, and still are, heavily used to conduct these simulations. However, due to the rapid increase in computational performance, numerical methods based on the discretization of the entire phase space (the so-called Eulerian...
approach) have gained increasing attention in recent years (at least for four-
dimensional and certain five-dimensional problems). These Eulerian methods
offer superior accuracy and the possibility to resolve regions of the phase space
with low density, but are extremely demanding from a computational point of
view. Consequently, it is important to develop efficient numerical methods that
can be parallelized to state of the art supercomputers.

While standard techniques from fluid dynamics can be applied to kinetic
equations, these usually suffer from a severe restriction in the time step size due
to a Courant–Friedrichs–Lewy (CFL) condition. On the other hand, the use
of implicit methods would significantly increase the memory footprint, which
is already a scarce resource for high-dimensional simulations. This is an even
more pronounced issue for accelerators such as graphic processing units (GPUs)
and the Intel Xeon Phi (which in all likelihood will become more pervasive in
the future). Semi-Lagrangian methods offer a viable alternative. These methods
follow the characteristics backward in time. In many kinetic models (such as the
Vlasov–Poisson and Vlasov–Maxwell equations) applying a splitting procedure
results in sub-steps for which the characteristics can be determined analyti-
cally (if this is not the case an explicit ordinary differential equation solver is
usually used). For the Vlasov–Poisson equation this approach has been first
suggested in the seminal paper of Cheng & Knorr [5]. Later it was extended
in various ways to the Vlasov–Maxwell equation [7, 32, 29]. However, since the
endpoint of a characteristic curve does not necessarily coincide with the grid
used, an interpolation procedure has to be employed. An attractive choice is
to reconstruct the desired function by spline interpolation (see, for example,
[33, 23]), which according to [13] is still considered the de facto standard in
Vlasov simulations. The main downside of this approach is that a tridiagonal
linear system of equations has to be solved to construct the spline. While this
can be done very efficiently in a sequential implementation, for modern super-
computers (which require scaling to hundreds of thousands of cores) such global
dependencies seriously limit the utility of this algorithm.

On the other hand, the semi-Lagrangian discontinuous Galerkin method em-
ployes a piecewise polynomial approximation in each cell of the computational
domain (see, for example, [31, 12, 18, 30]). In the case of an advection equation
the discretized function is translated and then projected back to the appropriate
subspace of piecewise polynomial functions. This method, per construction, is
mass conservative and only accesses two adjacent cells in order to compute the
necessary projection (this is true independent of the order of the approxima-
tion). The local nature of this method also makes it easy to incorporate block
structured mesh refinement and employing limiters to maintain positivity. Pos-
itivity limiters were also investigated for spline interpolation (see [35, 36, 11]),
but these schemes are much more involved. For the semi-Lagrangian discontin-
uous Galerkin scheme mathematical rigorous convergence results are available
in [19, 18]. The method is fully explicit (i.e. no linear system has to be solved
to advance the solution in time) and thus it is easier to implement (especially
on parallel architectures) and shows a more favorable communication pattern.
We should note that some measures have been taken to improve the parallel
scalability of cubic spline interpolation \cite{10}. However, even for this approach a relatively large communication overhead is incurred. This is due to the fact that the boundary condition for the local spline reconstruction requires a large stencil if the desirable properties of the global cubic spline interpolation are to be preserved (the method derived in \cite{10} requires a centered stencil of size $21$). In addition, a very efficient mixed precision implementation is available for the semi-Lagrangian discontinuous Galerkin method \cite{14}. This is particular useful for modern accelerators (such as GPUs) and helps to alleviate the problem of memory scarcity on such systems. Furthermore, it is worth mentioning that a filamentation filtration strategy has been developed \cite{17} (for problems where numerical recurrence is an issue) and that the error propagation is superior to other semi-Lagrangian schemes \cite{34,20} (although to our knowledge no proof of this fact is available for the Vlasov equation).

Before proceeding, let us note that a range of other methods have been used to solve the Vlasov equation as well. A method that is related to the discontinuous Galerkin approach and which has been widely employed in plasma simulations is the van Leer scheme (see, for example, \cite{22,29,24,4}). What distinguishes the discontinuous Galerkin method from the van Leer scheme is that the coefficients in the corresponding basis expansion are stored directly in computer memory. In contrast, the van Leer scheme replaces these coefficients by performing an approximation using suitable differences on an equidistant grid. Its implementation and performance characteristics are thus more closely related to other finite difference schemes.

Despite the many advantages of the semi-Lagrangian discontinuous Galerkin method outlined above, there is certainly no universal agreement that this method will be a viable alternative for large scale high-dimensional kinetic simulation. Specifically, it is often argued, both by reviewers and in informal conversation, that these methods require more degrees of freedom to achieve the same precision (compared to say spline interpolation) and that this is not maintainable in a higher-dimensional setting. This argument is certainly not easily dismissed. Let us assume that instead of $n$ degrees of freedom per direction our numerical approach requires $\alpha n$, with $\alpha > 1$. Then the memory footprint and run time are increased by a factor $\alpha^d$, where $d$ is the dimensionality of the problem. For example, even a moderate increase, for example $\alpha = 1.2$, results in a substantial increase of memory footprint/run time (a factor of 2 for $d = 4$ and factor of 3 for $d = 6$).

Of course, the important question then becomes if the semi-Lagrangian discontinuous Galerkin method actually requires more degrees of freedom and if so by how much. Although this is a common problem for classic (i.e. non semi-Lagrangian) discontinuous Galerkin methods and has, for example, lead to so-called hybrid schemes (see, for example, \cite{27}), the situation considered here is quite different. Specifically, the problem is reduced to a sequence of one-dimensional advection equations which are then solved using the method of characteristics. In the present paper we will investigate if the semi-Lagrangian discontinuous Galerkin method requires more degrees of freedom compared to spline interpolation. This will be done in the context of the Vlasov–Poisson
equation
\begin{equation}
\partial_t f(t, x, v) + v \cdot \nabla_x f(t, x, v) + E(f)(x) \cdot \nabla_v f(t, x, v) = 0
\end{equation}
\begin{equation}
\nabla \cdot E(f)(x) = \int f(t, x, v) \, dv - 1, \quad \nabla \times E(f)(x) = 0.
\end{equation}

However, before we proceed and discuss the comparative efficiency of the semi-Lagrangian discontinuous Galerkin method (section 3) and the four-dimensional simulations conducted (sections 4 and 5), let us dwell a bit more on the argument (against the discontinuous Galerkin method) outlined above. To do that we will consider nonlinear Landau damping as a test problem and, for the time being, still restrict ourselves to the two-dimensional case (for reasons that will be apparent shortly; the four-dimensional variant of this problem is discussed in section 4). That is, we consider the initial value
\begin{equation}
f(0, x, v) = \frac{1}{\sqrt{2\pi}} (1 + \epsilon \cos(kx)) e^{-\frac{1}{2}v^2},
\end{equation}
where \(\epsilon = \frac{1}{2}\) and \(k = \frac{1}{2}\), on the domain \([0, 4\pi] \times [-6, 6]\). A splitting procedure is employed to obtain a sequence of advection equations which are discretized using either a semi-Lagrangian approach based on spline interpolation or the semi-Lagrangian discontinuous Galerkin approach (both numerical methods will be discussed in some detail in section 2). The error committed by these two numerical schemes is computed by comparing it to a reference solution (with a sufficiently fine resolution).

Many publications (see, for example, [16, 12, 6]), including work by the authors of this paper, have reported such results. To facilitate the present discussion, we have computed the error in the particle-density \(f\) at time \(t = 10\) as a function of the degrees of freedom (per direction). These results are shown on the left in Figure 1. We clearly observe that, at best, the discontinuous Galerkin method struggles in a direct comparison with spline interpolation. The sixth-order discontinuous Galerkin method (dG6) requires approximately 50% more degrees of freedom to obtain the same accuracy. Thus, it seems that the discussion has come to a close and the criticism outlined above seems to be completely justified. However, the major caveat here is that in practical applications no one is interested in solving nonlinear Landau damping only up to \(t = 10\). Now, if we look at the error at \(t = 50\) (right-hand side of Figure 1), the error does not appreciably decrease at all (up to 2048 degrees of freedom per direction) for any of the numerical methods considered. To run a simulation with 2048 degrees of freedom in four dimensions would require at least 280 TB of memory, which would stress even the world’s most powerful supercomputers (the number one system on the present TOP500 supercomputing list has 1310 TB of memory; thus such a simulation would be possible in principle, but certainly not very practical). Thus, while the argument provided is valid for very small times, it is all but pointless for realistic simulations. The question that we should ask instead is how well does a numerical method resolve the physics of the problem under consideration (for reasonable problem sizes).
Figure 1: The error, as a function of the number of degrees of freedom (per direction), for nonlinear Landau damping is shown at $t = 10$ (left) and $t = 50$ (right). The error is computed in the infinity norm and is normalized by the norm of the reference solution. The reference solution is computed using spline interpolation with $8192$ grid points (per direction). For all simulations a Strang splitting procedure with time step size $\tau = 0.1$ has been used.

The example considered here is two-dimensional and thus should not be considered the gold standard for higher-dimensional simulations. In fact, the reason why we consider this in two dimensions is that it is prohibitively expensive to run simulations with $2048^4$ or even more degrees of freedom. This paper will show that for the four-dimensional case the performance of a numerical method in the asymptotic regime is markedly different from the performance of interest in realistic simulations. More specifically, we will show that even though the semi-Lagrangian discontinuous Galerkin method is inferior to spline interpolation for small times, depending on the problem, similar or even superior performance can be achieved for more realistic simulations.

Of course, the importance of designing numerical methods that faithfully resolve the physics (as opposed to primarily decreasing the error in some norm) has been considered extensively in the context of long time integration (see, for example, [26]). The difference here is that we certainly would not consider $t = 50$ for nonlinear Landau damping a long-time result. In fact, to determine the long-time behavior of nonlinear Landau damping, numerical simulations, mostly in two dimensions, for at least up to $t = 1300$ have been considered (see, for example, [3] and [28]). In the literature much effort has been dedicated to numerical methods which conserve certain invariants (such as mass or energy) exactly and to methods which have good long time properties with respect to these quantities. This is also an important aspect for the Vlasov equation and many works exist that consider conservation of invariants (see, for example, [11, 23] for spline interpolation and [16, 12] for the discontinuous Galerkin approach). In addition, since the Vlasov equation is often used to simulate plasma instabilities, the time evolution of certain averaged quantities (most notably the total electric energy) are of significant interest. Both of these aspects will be considered in sections 4 and 5.
2. Numerical methods

To advance the numerical solution in time we use the splitting approach introduced in [5]. That is, we first consider the free-streaming sub-flow

\[ \partial_t f(t, x, v) + v \cdot \nabla_x f(t, x, v) = 0, \quad f(0, x, v) = g(x, v). \]  

(2)

The corresponding solution at time \( \tau \) is then denoted by

\[ e^{\tau A} g(x, v) = f(\tau, x, v). \]

For the second part of the splitting algorithm we consider the acceleration sub-flow

\[ \partial_t f(t, x, v) + E(x) \cdot \nabla_v f(t, x, v) = 0, \quad f(0, x, v) = g(x, v), \]

(3)

where the electric field \( E \) is determined from the (charge) density \( \rho(x) = \int g(x, v) \, dv \) and is thus taken constant during that step. This property actually follows from equation (3) as the translation is only in the \( v \)-direction and thus leaves the density invariant. We denote the corresponding solution by

\[ e^{\tau B} g(x, v) = f(\tau, x, v). \]

Note that, contrary to what the notation suggests, \( B \) is a nonlinear operator as the electric field depends on \( f \). Using this notation we can easily formulate a time step of the splitting algorithm. In all our numerical results we will use the second-order Strang splitting given by

\[ f_{m+1} = e^{\tau^2 A} e^{\tau B} e^{\tau^2 A} f_m, \]

where \( f_m(x, v) \) is an approximation of \( f(t_m, x, v) \) and \( \tau = t_{m+1} - t_m \) is the time step size. Let us note, however, that higher order splitting methods have been constructed as well (see, for example, [8]).

The main computational advantage of the splitting scheme is that the resulting sub-flows, given by equations (2) and (3), are in the form of an advection equation, where the advection speed is independent of the variables being advected. Thus, we can immediately solve equation (2) to obtain

\[ e^{\tau A} g(x, v) = g(x - \tau v, v) \]

and solve equation (3) to obtain

\[ e^{\tau B} g(x, v) = g(x, v - \tau E(x)). \]

Thus, by using the splitting approach outlined above we have reduced the task of computing a numerical approximation to the four-dimensional Vlasov–Poisson equation to computing two-dimensional translations in phase space. However, this can be further simplified by splitting equation (2) into an advection in the \( x_1 \) direction

\[ \partial_t f(t, x, v) + v_1 \partial_{x_1} f(t, x, v) = 0, \quad f(0, x, v) = g(x, v) \]
and an advection in the $x_2$ direction
\[ \partial_t f(t, x, v) + v_2 \partial_{x_2} f(t, x, v) = 0, \quad f(0, x, v) = g(x, v). \]

Let us denote the corresponding flows at time $\tau$ by $e^{\tau A_1} g(x, v)$ and $e^{\tau A_2} g$, respectively. Then since $[v_1 \partial_{x_1}, v_2 \partial_{x_2}] = 0$ (i.e. the two operators commute) we have
\[ e^{\tau A_2} e^{\tau A_1} g = e^{\tau A} g \]
and no further splitting error is incurred. A very similar procedure can be carried out for the acceleration sub-flow. Thus, the four-dimensional Vlasov–Poisson equation has been reduced to a sequence of one-dimensional advectons. This has significant advantages from a computational point of view and, in particular, implies that we only have to formulate our semi-Lagrangian scheme in a single dimension. We should remark that this approach can be extended to more complicated models, such as the Vlasov–Maxwell equations [7].

We will now describe the two competing approaches considered in this paper. First, for the semi-Lagrangian method based on spline interpolation an equidistant grid is used. However, for any grid point $(x_{1i}, x_{2j}, v_{1k}, v_{2l})$ the translations $x_{1i} - \tau v_{1k}$, $v_{1k} - \tau E_1(x_{1i}, x_{2j})$, etc., in general, do not coincide with a grid point (here $E_1$ denotes the first component of the electric field). Thus, we have to use an interpolation scheme in order to evaluate $g(x_{1i} - \tau v_{1k}, x_{2j}, v_{1k}, v_{2l})$, $g(x_{1i}, x_{2j}, v_{1k} - \tau E_1(x_{1i}, x_{2j}), v_{2l})$. This is done by cubic spline interpolation. Note that only splines in one space dimension have to be constructed. This is essential for the computational efficiency of the scheme. Further computational ramifications are discussed in more detail in section 3.

Second, the semi-Lagrangian discontinuous Galerkin scheme divides the domain into cells $C_{ijkl} = I_1^{(1)} \times I_2^{(2)} \times I_3^{(3)} \times I_4^{(4)}$, where the $I_1^{(1)}$, $I_2^{(2)}$, $I_3^{(3)}$, $I_4^{(4)}$ are one-dimensional intervals of length $h$. We further assume that a function $g$ is given such that $g|_{C_{ijkl}}$, i.e. the restriction of $g$ to the $i$th cell, is a polynomial of degree $\ell$. Then the function $g$ lies in the approximation space (note that we do not enforce a continuity constraint across cell interfaces). However, in general, this is not true for the translated function (for the other three translations we proceed in exactly the same way)
\[ (T_\tau g)(x_1, x_2, v_1, v_2) = g(x_1 - \tau v_1, x_2, v_1, v_2). \]

Thus, we perform an approximation by applying a projection operator $P$ and obtain $PT_\tau g$. In order to turn this into a sensible numerical scheme, it must hold that $Pf|_{C_{ijkl}}$ is a polynomial of degree $\ell$ for any function $f$. Then $PT_\tau g$ constitutes the sought-after approximation of $g(x_1 - \tau v_1, x_2, v_1, v_2)$. In the present case the operator $P$ is the $L^2$ projection on the (finite dimensional) subspace of cell-wise polynomials of degree $\ell$. Note that this projection is the only approximation made. There is some freedom in selecting appropriate degrees of freedom for this method. Two obvious choices are to use the coefficients in a Legendre expansion or the value of function evaluations at the Gauss–Legendre points. In our implementation we use the latter. Since the value at the beginning of each time step is (per assumption) a cell-wise polynomial of degree
ℓ, we can use Gauss–Legendre quadrature in order to analytically compute the coefficients required to implement the projection. The performance implications will be discussed in section 3.

3. Computational consideration & performance results

The goal of this section is to discuss the performance of the cubic spline and discontinuous Galerkin based semi-Lagrangian methods on modern computer architectures. We will do this both from a theoretical and a practical (i.e. measuring specific implementations of these algorithms) point of view.

As has been discussed in the previous section, only a set of one-dimensional advection equations have to be solved for both algorithms. Thus, to simplify the notation, we only consider the one-dimensional advection

\[ u^{m+1}(x) = u^m(x - v\tau) \]

in this section. This is, in fact, exactly how the numerical algorithm is implemented on a computer. A function is provided that solves the one-dimensional advection given by equation (4). This function is then executed multiple times with different parameters.

For spline interpolation, the function values \( u^m_i = u^m(x_i) \) are given at the grid points \( x_i \). The index \( i \) runs from 0 to \( n - 1 \), where \( n \) is the number of degrees of freedom. From this a spline that covers the entire domain has to be constructed. Since it would be wasteful to represent a spline as a collection of piecewise polynomials (as most of the coefficients of these polynomials are fixed by the continuity constraints), a B-spline basis is used instead. That is, the spline interpolation \( \tilde{u}^m \) of \( u^m_i \) is represented as

\[ \tilde{u}^m(x) = \sum_k \omega_k S(x - x_k), \]

where \( S \) is given by

\[
6S(x) = \begin{cases}
4 - 6(x/h)^2 + 3|x/h|^3 & 0 \leq |x| \leq h \\
(2 - |x/h|)^3 & h \leq |x| \leq 2h \\
0 & \text{otherwise}
\end{cases}
\]

and \( h \) is the grid spacing. The coefficients \( \omega_k \) are uniquely determined by the function values and can be obtained by solving the following linear system of equations (assuming periodic boundary conditions)

\[
\frac{1}{6} \begin{bmatrix}
4 & 1 & 0 & \cdots & 0 & 1 \\
1 & 4 & 1 & 0 & \cdots & 0 \\
0 & 1 & 4 & \ddots & \ddots & \ddots \\
0 & 0 & \ddots & \ddots & 0 \\
\vdots & \vdots & \ddots & 1 & 4 & 1 \\
1 & 0 & \cdots & 0 & 1 & 4
\end{bmatrix} \begin{bmatrix}
\omega_0 \\
\omega_1 \\
\vdots \\
\vdots \\
\omega_{n-1}
\end{bmatrix} = \begin{bmatrix}
u_0 \\
u_1 \\
\vdots \\
\vdots \\
u_{n-1}
\end{bmatrix}.
\]
Note that the matrix is tridiagonal and symmetric and thus efficient methods exist to solve this system. Once the spline is constructed it is used to solve equation \(4\) as follows

\[
 u_{i}^{m+1} = \tilde{u}^m(x_i - v\tau).
\]

Thus, the numerical algorithm proceeds in two steps. First, the spline is constructed by solving a linear system and this intermediate representation is then used to evaluate the numerical solution at the grid points. Both of these steps require \(O(n)\) memory and arithmetic operations, where \(n\) is the number of degrees of freedom. In the construction step this is due to the fact that linear complexity solvers are available for tridiagonal systems (such as the Thomas algorithm) and for the evaluation step this is true since the support of \(S\) is compact and independent of \(n\).

On modern computer hardware both of these steps are memory bound. That is, performance is dictated by how fast data can be read and written to memory (as opposed to how many arithmetic operations the computer can perform). In the present case we have to at least read all \(u_i^m\) and write all \(u_i^{m+1}\). In addition, we have to write and read all \(\omega_k\) at least once. The latter is more subtle than one might expect. Since in a sequential implementation only one one-dimensional array of \(\omega_k\) needs to exist at any given time, this array could conceivably be kept in cache (except for extremely large problem sizes) and then reads and writes to it would not count as a memory operation (in the sense that those do not negatively impact performance).

For the semi-Lagrangian discontinuous Galerkin scheme the degrees of freedom are given by function values \(u_{ij}^m \approx u^m(x_{i-1/2} + \xi_j)\), where \(i\) is the cell-index (which runs from 0 to \(n_C\)), \(x_{i-1/2}\) the left cell interface, and \(\xi_j\) is the \(j\)th Gauss–Legendre node scaled to the interval \([0,h]\). The indices \(j\) run from 0 to \(\ell\) (the degree of the polynomial approximant). The resulting scheme has order \(\ell + 1\) and we will henceforth denote it as dG(\(\ell + 1\)). It is clear that to perform the \(L^2\) projection on the \(i\)th cell only data from at most two adjacent cells is required. This is illustrated in Figure 2. We denote the index of these two cells by \(i^*\) and \(i^* + 1\), respectively. Since the projection is a linear operator we can write the resulting numerical scheme as follows

\[
 u_{ij}^{m+1} = \sum_l A_{jl} u_{i^* l}^m + \sum_j B_{jl} u_{i^*+1, j}^m,
\]

where \(A \in \mathbb{R}^{(\ell+1) \times (\ell+1)}\) and \(B \in \mathbb{R}^{(\ell+1) \times (\ell+1)}\) are matrices that only depend on \(v\). The matrices can be precomputed at the beginning of each time step and thus their cost will not be a major concern except for extremely coarse space discretizations. The derivation of their exact form is somewhat lengthy and we thus refer the interested reader to [15].

For the implementation equation \(5\) is employed. The size of the matrices is negligible in comparison to the cache size and we therefore do not have to count reads and writes to them as memory accesses. The present algorithm has to read all entries in \(u_{ij}^m\) and write to all entries in \(u_{ij}^{m+1}\); it thus requires \(O(n)\) memory operations, where \(n = n_C(\ell + 1)\) is the number of degrees of freedom.
Figure 2: Illustration of the semi-Lagrangian discontinuous Galerkin scheme. The piecewise polynomial function of degree $\ell$ (left picture) is translated by $v\tau$ (right picture). The translated function is then projected back to the approximation subspace (which requires data from at most two adjacent cells).

Technically, the algorithm requires $O(\ell n)$ arithmetic operations and thus does not scale linearly in the degrees of freedom. However, since, for reasonable values of $\ell$, this is still an extremely memory bound problem this is of no consequence (i.e. performance is dictated by the memory accesses and not by amount of arithmetic operations that need to be performed).

In an idealized setting both algorithms could achieve very similar levels of performance (both require us to read $u^n$ once and write to $u^{n+1}$ once). Of course, achieving this in an actual implementation is not a trivial task. However, it is our believe that the local nature of the discontinuous Galerkin approach simplifies this task, at least, to some extent. In the following we will compare the performance of two Vlasov solvers. SeLaLib [1] is a library that facilitate kinetic simulations based on the semi-Lagrangian approach. It has been used extensively (see, for example, [9,25,2]). The library is written in Fortran and provides an efficient implementation of the cubic spline based semi-Lagrangian discontinuous Galerkin method. SeLaLib is compared with our own implementation (called sldg) of the semi-Lagrangian discontinuous Galerkin approach. The code is written in C++ and due to the use of templates facilitates implementations that work in arbitrary dimensions, while still maintaining good performance. This architecture is described in [15]. In addition to solving the Vlasov equation, it has been used in non-traditional applications of semi-Lagrangian methods (such as for the Kadomtsev–Petviashvili equation that requires the solution of a nonlinear advection [21]).

Another major difference in the implementation of these two codes is how data is presented to the routines that perform the one-dimensional advectons. For SeLaLib the corresponding arrays are always contiguous in memory. That is, for the advection in the $x_1$-direction the elements in the $x_1$ direction are laid out sequentially in memory, for the advection in the $x_2$-direction the elements in the $x_2$ direction are laid out sequentially in memory, etc. This requires that the four dimensional array is converted (i.e. transposed) to the appropriate form for each advection. This approach has the advantage that the same code can be used for all advection steps and cache locality is ensured almost automatically. It also
Table 1: Comparison of performance and memory usage between SeLaLib (cubic spline interpolation) and sldg (fourth order semi-Lagrangian discontinuous Galerkin method). For both codes the nonlinear Landau damping initial value of section 4 is used and 64 and 128 degrees of freedom (dof) per direction are considered. The Workstation consists of two Intel Xeon E5-2630 v3 CPUs and 32 GB of DDR4 memory. One node of the VSC3 consists of two Intel Xeon E5-2650v2 and 64GB of DDR3 memory. The memory consumption is measured using the /usr/bin/time utility.

| Workstation | Time per step [s] | Memory used [GB] |
|-------------|------------------|-----------------|
|             | dof=64           | dof=128         |
| SeLaLib     | 0.194            | 3.25            |
|             | 0.85             | 13.0            |
| sldg (dG4)  | 0.105            | 1.62            |
|             | 0.28             | 4.2             |

| VSC3        | Time per step [s] |
|-------------|-------------------|
|             | dof=64            | dof=128         |
| SeLaLib     | 0.219             | 4.02            |
| sldg (dG4)  | 0.151             | 2.31            |

fits very well with spline interpolation, as the global algorithm for constructing the spline is pointed at one contiguous array of memory. The disadvantage of this approach is that the transpositions do not have a very favorable memory access pattern and thus incur a significant computational cost.

On the other hand, the semi-Lagrangian discontinuous Galerkin implementation uses the ability of abstractions in C++ to hide the actual memory layout from the function that performs the advection. Thus, it is still possible to write the one-dimensional advection code as if the arrays are contiguous, even though the actual memory layout is more complicated. As a consequence, no transpositions have to be carried out in this implementation. The main concern here is that cache locality has to be taken into account. However, the local nature of the discontinuous Galerkin approach makes this less problematic. To further alleviate this issue we use cache blocking for the dimension with the largest stride.

We have measured both the performance and the amount of memory used for these two codes. The results are shown in Table 1. For both methods one dual socket node of the described computer systems is used. Both codes are run using 16 parallel threads. We observe that the discontinuous Galerkin implementation is both faster (by approximately a factor of 2 on the newer Haswell CPUs and approximately a factor of 1.7 on the Ivy Bridge CPUs) and uses significantly less memory (by approximately a factor of 3), if the same number of degrees of freedom are used.

Both codes support parallelization using MPI. This is essential even to obtain good performance on a single node (i.e. for the results presented in Table 1). However, both codes can also be run on distributed memory clusters (supercomputers). In this case the issue of how to distribute the computational domain to the different MPI processes becomes important. Although this choice
of data distribution is identical for both codes it is instructive to discuss it. The main issue here is that most FFT libraries (in particular, FFTW, which is used in both codes) only allow a so-called pencil decomposition. Thus it is not straightforward to parallelize the Fourier transforms (without incurring a penalty associated with additional data transfer). Both codes avoid this problem by only parallelizing in the velocity directions. Then, the Fourier transforms can be conducted independently on each of the processes. For four/six dimensional simulations there are still two/three dimensions available for parallelization. The main difference in the two codes is that while sldg only performs halo communication (due to the local nature of the semi-Lagrangian discontinuous Galerkin scheme), SeLaLib requires two calls to MPI_Alltoall in each time step (due to the global data dependency inherent in constructing the spline).

We now investigate the scaling of both codes on the Vienna Scientific Cluster 3 (VSC3). For more information and details on the hardware of that system we refer the reader to \url{http://vsc.ac.at/systems/vsc-3/}. On the left hand side of Figure 3 we present strong scaling results (i.e. the problem size is kept constant as the number of cores is increased) for the problem considered in section 4. We observe that the discontinuous Galerkin scheme maintains its advantage independently of the number of cores the program is run on (i.e. both codes show a similar behavior with respect to scaling). On the right hand side of Figure 3 we present weak scaling results (i.e. the problem size increases proportionally to the number of cores). In this case we observe that sldg significantly outperforms SeLaLib (in particular, as we consider more than 256 cores). It is interesting to investigate the reason for this discrepancy in performance. To that end we have subtracted the time required for the calls to MPI_Alltoall, which are used to map the velocity grid into contiguous memory on each process, in the SeLaLib code from the total run time (note that even on a single node this operation requires a significant amount of time, as computing the transpose is a memory intensive operation). The resulting line is very similar to the run time of the sldg code. That is, almost all of the difference in performance between the two codes is explained by the global communication required for the spline interpolation.

4. Nonlinear Landau damping

In this section we consider nonlinear Landau damping in four dimensions. That is, we impose the initial value

\[
f(0, x_1, x_2, v_1, v_2) = \frac{1}{2\pi} \left(1 + \epsilon (\cos(kx_1) + \cos(kx_2))\right) e^{-\frac{(v_1^2 + v_2^2)}{2}}
\]

on the domain \([0, 4\pi]^2 \times [-6, 6]^2\). We set \(\epsilon = \frac{1}{2}\) and \(k = 0.5\). This is precisely the setup used in [23]. Strang splitting with step size \(\tau = 0.1\) is used for the time integration. We are interested in resolving the dynamics of nonlinear Landau damping up to \(t = 50\). More precisely, the characteristic growth and decay of the electric energy should be indistinguishable (in a plot) from the exact solution.
Figure 3: Strong and weak scaling for SeLaLib (cubic spline interpolation) and sldg (fourth order semi-Lagrangian discontinuous Galerkin method). For strong scaling the nonlinear Landau damping initial value of section 4 with 128 degrees of freedom per direction is used. For weak scaling $n^2 \times (\sqrt{n} \cdot 128)^2$ degrees of freedom, where $n$ is the number of nodes employed, are used. The simulations are conducted on the VSC3, each node of which consists of two Intel Xeon E5-2650v2 and 64GB of DDR3 memory.

until $t = 50$. To accomplish this for spline interpolation we need 128 grid points per direction (see Figure 1). Note that despite the fact that, as the results in Figure 1 show, the error in the density function is on the order of unity, the physically important quantity is resolved very well. Therefore, there is simply no advantage of running the integrator in the asymptotic regime (which in any case would not be feasible in four dimensions).

In the introduction second, fourth, and sixth order discontinuous Galerkin schemes were considered. It was already clear at that point that the second order scheme is not competitive. However, the order plots (for $t = 10$) presented there, seem to indicate that the sixth order discontinuous Galerkin approach performs best. However, this is misleading as numerical simulations have shown that for $t = 50$ (where the plots in the introduction give no sensible indication) the fourth order method should actually be favored. Therefore, in this section we will exclusively use dG4. Before proceeding, let us note, however, that for long time integration this is not necessarily the case. For example [16] found by performing numerical simulations in two dimensions that for long times (both for nonlinear Landau damping and for a bump on tail instabilities) the sixth order method can be significantly more efficient.

If we run the same simulation for the discontinuous Galerkin approach we see that the solution is somewhat worse. In particular, between $t = 40$ and $t = 50$ the growth rate of the electric energy is slightly increased compared to the reference solution. In order to remedy this we can increase the number of degrees of freedom per direction to 144 (corresponding to $\alpha = 1.125$). This means that in four dimensions 60% more degrees of freedom are required by the discontinuous Galerkin approach to achieve the same accuracy. However, as we have seen in section 3 the implementation of the discontinuous Galerkin scheme is also more efficient (by a factor of 2 on Haswell CPUs) and uses less
memory (by a factor of 3). Thus, the discontinuous Galerkin scheme retains an advantage in performance (approximately 25%). In addition, the amount of memory required is reduced by a factor of 2.

Figure 4: The electric energy (top left), the energy error (top right), the $L^2$ norm (bottom left), and the $L^1$ norm (bottom right) are shown as a function of time for the nonlinear Landau damping problem given in equation (6). The reference solution is computed using spline interpolation and 512 grid points (per direction).

In addition, Figure 4 shows the important physical invariants of energy, $L^2$ norm, and $L^1$ norm. Although the main concern for the nonlinear Landau damping is the time evolution of the electric energy, these invariants give additional diagnostics of how well physical properties of the exact solution are respected by a numerical method.

The error in energy is almost identical for both methods and is on the order of $10^{-3}$ (after a transient phase). This is due to the fact that the energy error for the present problem is dominated by the time integration error.

For collisionless plasma simulations the $L^2$ error is an important diagnostic. A decrease in the $L^2$ norm implies that our numerical scheme adds numerical diffusion to the simulation. Although most numerical methods introduce a certain amount of diffusion, it is clearly preferable to limit this as much as possible. For nonlinear Landau damping the discontinuous Galerkin method outperforms spline interpolation in this metric (in particular, between $t = 10$ and $t = 40$).
Finally, we look at the $L^1$ norm. For the exact solution mass and $L^1$ norm are equal in value and the $L^1$ norm is thus conserved. However, if a mass conservative numerical scheme commits a numerical error that introduces negative values, the result is a change in the $L^1$ norm. This is, in particular, a problem for schemes that are susceptible to oscillations (such as spectral methods). However, most numerical methods for the Vlasov equation experience this behavior to some extent. Let us emphasize that (contrary to say the Euler equations) this neither leads to numerical instabilities nor necessarily diminishes averaged quantities (such as the electric energy). However, since negative values make the physical interpretation of the density function difficult, it is still desirable to limit this behavior as much as possible. For the nonlinear Landau damping the discontinuous Galerkin scheme shows worse performance with respect to $L^1$ conservation compared to spline interpolation.

Overall we conclude that the discontinuous Galerkin implementation has a moderate advantage with respect to performance and $L^2$ conservation as well as a significant advantage with respect to memory consumption. On the other hand, spline interpolation has a moderate advantage with respect to $L^1$ conservation. If we assume that the present results generalize to the six-dimensional case (i.e. $\alpha = 1.125$ as above), then the performance of both methods would be evenly matched. This is due to the fact that $1.125^6 \approx 2$ more degrees of freedom would be required for the discontinuous Galerkin scheme, which exactly cancels the factor of 2 advantage the corresponding implementation provides.

5. Two-stream instability

In this section we consider the two-stream instability in four dimensions. That is, we impose the initial value

$$f(0, x_1, x_2, v_1, v_2) = (1 + \epsilon \cos(k x_1) \cos(k x_2)) f_{eq}(v_1, v_2)$$

(7)

with

$$f_{eq} = \frac{1}{8\pi} \left( e^{-(v_1-v_0)^2/2} + e^{-(v_1+v_0)^2/2} \right) \left( e^{-(v_2-v_0)^2/2} + e^{-(v_2+v_0)^2/2} \right),$$

where $\epsilon = 10^{-3}$, $k = 0.2$, and $v_0 = 2.4$. The computational domain is $[0, 10\pi]^2 \times [-6, 6]^2$. Strang splitting with step size $\tau = 0.1$ is used for the time integration.

This is an interesting problem as the electric energy is small (even decreases by three orders of magnitude compared to the initial value) during a quiescent phase (from $t = 0$ to approximately $t = 175$). However, eventually a physical instability develops that results in an exponential increase of the electric energy (approximately $t = 175$ to $t = 225$). This is followed by nonlinear saturation. In this nonlinear phase the electric energy oscillates around a constant value with an amplitude that decreases in time.

It turns out that to obtain agreement between a numerical and the exact solution for the quiescent phase is challenging. In fact, simulations conducted for a similar two-dimensional problem indicate that for spline interpolation this is
only achieved for 1024 grid points per direction (which is extremely expensive). Nevertheless, simulations with fewer degrees of freedom are still able to give pertinent information on the plasma instability. In particular, at which time the instability occurs, the growth rate of the instability, and the behavior of the electric energy in the nonlinear regime (i.e. after saturation).

Figure 5 shows a comparison of the fourth order discontinuous Galerkin method and spline interpolation for 32, 64, and 128 degrees of freedom (per direction). For the dG4 method already 32 degrees of freedom are sufficient in order to accurately predict the time and growth of the instability. On the other hand, this is obviously not true for spline interpolation (which shows an instability that is too late and does not reach the peak for the electric energy). While increasing the number of degrees of freedom for spline interpolation results in significant improvements with respect to these metrics, the discontinuous Galerkin approach always retains an advantage in the nonlinear phase. More specifically, spline interpolation exhibits a decay of the electric energy that is not present in the exact solution.

This behavior can be explained by looking at the time evolution of the $L^2$ norm (see Figure 5). We observe that the $L^2$ norm for dG4 is significantly better conserved compared to spline interpolation once we enter the nonlinear phase. Thus, more diffusion is introduced by spline interpolation which results in the corresponding (unphysical) decrease in electric energy observed (and the absence of the same artefact for the discontinuous Galerkin approach).

For the present example, the difference between performance in the asymptotic regime (valid only for small times) and performance for time scales of physical interest, can be highlighted very clearly. In fact, if we look at the plots in Figure 5 we see that in the quiescent phase, the numerical solution obtained with spline interpolation follows the exact solution somewhat longer. However, even for 128 degrees of freedom agreement is only achieved up to approximately $t = 30$ (for spline interpolation). Nevertheless, the dG4 method is clearly superior in terms of both predicting the onset of the instability and in providing qualitatively correct predictions for the nonlinear phase (i.e. in obtaining the physics of interest).

Let us further remark that the $L^1$ norm is significantly better conserved by the discontinuous Galerkin approach compared to spline interpolation. Both methods conserve energy quite well (at least up to $10^{-3}$). Although for the present simulation there is some advantage for spline interpolation.

We conclude that for the two-stream instability considered here, the discontinuous Galerkin approach is clearly the superior choice compared to spline interpolation.

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Figure 5: The time evolution of the electric energy for the two-stream instability given by equation (7) is shown for 32 (top left), 64 (top right), and 128 (center left) degrees of freedom. In addition, the energy error (center right), the $L^2$ norm (bottom left), and the $L^1$ norm (bottom right) are shown as a function of time. The reference solution is computed using spline interpolation and 512 grid points (per direction).
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