Six Lectures on the Geometric Integration of ODEs

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1 General tools

"A major task of mathematics is to harmonize the continuous and the discrete, to include them in one comprehensive mathematics, and to eliminate obscurity from both." (E. T. Bell, Men of Mathematics)

1.1 Introduction

Motion is described by differential equations, which are derived from the laws of physics. In the simplest case, they read $m\frac{d^2x}{dt^2} = F(t, x, \frac{dx}{dt})$ —Newton's second law. These equations contain within them not just a statement of the current acceleration experienced by the object(s), but all the physical laws relevant to the particular situation. Finding these laws and their consequences for the motion has been a major part of physics since the time of Newton. For example, the equations tell us the space in which the system evolves (its *phase space*, which may be ordinary Euclidean space or a curved space such as a sphere); any symmetries of the motion, such as the left–right or forwards–backwards symmetries of a pendulum; and any special quantities such as energy, which for a pendulum is either conserved (if there is no friction) or decreases (if there is friction). Finally and most importantly, the laws describe how all motions starting close to the actual one are constrained in relation to each other. These laws are known as *symplecticity* and *volume preservation*.

"A gyroscope is an emissary from a six-dimensional symplectic world to our three-dimensional one; in its home world its behavior looks simple and natural." (Yuri Manin)

Standard methods for simulating motion, called *numerical integrators*, take an initial condition and move the objects in the direction specified by the differential equations. They completely ignore all of the above hidden physical laws contained within the equations. Since about 1990, new methods have been developed, called *geometric integrators*, which obey these extra laws. Since this is physically natural, we can hope that the results will be extremely reliable, especially for long-time simulations.

Before we tell you all the advantages, three caveats:

- The hidden physical law usually has to be known if the integrator is going to obey it. For example, to preserve energy, the energy must be known.
- Because we're asking something more of our method, it may turn out to be computationally more expensive than a standard method. Amazingly (because the laws are so natural?) sometimes it's actually much cheaper.
- Many systems have multiple hidden laws, for which methods are currently known which preserve any one law but not all simultaneously.

Now the advantages:

- Simulations can be run for enormously long times, because there are no spurious non-physical effects, such as dissipation of energy in a conservative system;
- By studying the structure of the equations, very simple, fast, and reliable geometric integrators can often be found;

- In some situations, results can be guaranteed to be *qualitatively* correct, even when the motion is chaotic. This allows one to study systems in a "quick and dirty" mode and explore the system thoroughly, while retaining reliability;
- For some systems, even the actual quantitative errors are much smaller for short, medium, and long times than in standard methods.

Chapter 2 discusses a case where all of these nice features are realized: the solar system.

The first lecture is about general tools which will be useful later on, the second discusses the question "why bother?", and the third to sixth lectures are about how to preserve various specific properties.

These lectures were delivered at ANODE, the Auckland Numerical ODEs workshop, in July 1998. Naturally, they are tailored to our own research interests. They are intended to be suitable for a student's first exposure to the subject, and we have preserved their informality. We are very grateful to John Butcher for inviting us to speak, to all the organizers of ANODE, and especially to Nicolas Robidoux for transcribing the lectures. ANODE and the authors are supported by the Marsden Fund of the Royal Society of New Zealand, the Australian Research Council and the EPSRC. The written form was prepared at the MSRI, Berkeley, supported in part by NSF grant DMS-9701755.

1.2 The exact flow of an ODE, and general properties of integrators

We first define the exact flow (or solution) of an ordinary differential equation (ODE) and discuss what properties one would like an integrator to have. Let x(t) be the exact solution of the system of ordinary equations (ODEs)[†]

$$\frac{dx}{dt} = f(x), \quad x(0) = x_0, \quad x \in \mathbb{R}^m.$$
(1.1)

The exact flow φ_{τ} is defined by

$$x(t+\tau) = \varphi_{\tau}(x(t)) \quad \forall t, t \in \mathcal{F}$$

For each fixed time step τ, φ is a map from phase space to itself, i.e. $\varphi_{\tau} : \mathbb{R}^m \to \mathbb{R}^m$.

Three properties of exact flows

(i) (Self-adjointness) The flow has the continuous group property

$$\varphi_{\tau_1} \circ \varphi_{\tau_2} = \varphi_{\tau_1 + \tau_2} \quad \forall \tau_1, \tau_2 \in \mathbb{R}.$$

$$(1.2)$$

In particular,

$$\varphi_{\tau} \circ \varphi_{-\tau} = Id \tag{1.3}$$

Hence the exact flow is *self-adjoint*:

$$\varphi_{\tau} = \varphi_{-\tau}^{-1}. \tag{1.4}$$

(ii) (Taylor expansion)

$$x(\tau) = x(0) + \tau \frac{dx}{dt}(0) + \tau^2 \frac{1}{2} \frac{d^2x}{dt^2}(0) + \dots$$

Substitute

$$\frac{dx}{dt} = f(x)$$
$$\frac{d^2x}{dt^2} = (df)\frac{dx}{dt} = (df)f$$

Hence

$$\varphi_{\tau}(x_0) = x_0 + \tau f(x_0) + \frac{1}{2}\tau^2 (df(x_0))f(x_0) + \dots$$
(1.5)

[†] Nonautonomous ODEs dx/dt = f(x, t) can be formulated autonomously as $dx/dt = f(x, x_{m+1})$, $dx_{m+1}/dt = 1$. The geometric integrator is applied to this "extended" system (if possible), and then $t = x_{m+1}$ substituted.

General tools

(iii) (Formal exact solution)

$$\varphi_{\tau}(x) = e^{\tau \sum_{i=1}^{n} f_{i}(x) \frac{\partial}{\partial x_{i}}}(x)$$

$$:= \exp(\tau f)(x)$$
(1.6)

It's impossible to construct integrators with the continuous group property (1.2) for any reasonably general class of ODEs. The closest one can come is to preserve self-adjointness.

Properties of integrators

In general we don't know the flow φ_{τ} , so we seek maps ψ_{τ} that approximate φ_{τ} . We call such ψ_{τ} integrators. Some properties of integrators:

(i) (Self-adjointness) It is useful for ψ_{τ} to be self-adjoint, i.e.,

$$\psi_{\tau} = \psi_{-\tau}^{-1}$$

(ii) (Order of an integrator) The order of accuracy of ψ_{τ} is p, if the Taylor series of ψ_{τ} and the exact flow φ_{τ} agree to order p:

$$\psi_{\tau}(x) - \varphi_{\tau}(x) = \mathcal{O}(\tau^{p+1})$$

(iii) (Consistency) A necessary property of ψ_{τ} is that it be consistent, i.e., first order accurate, i.e.,

$$\psi_{\tau}(x) = x + \tau f(x) + \mathcal{O}(\tau^2)$$

Note: It is not difficult to show that every self-adjoint integrator is of even order. There are three types of integrators:

- (i) Integrators that form a group
- (ii) Integrators that form a symmetric space
- (iii) Integrators that form a semigroup

1.3 Integrators that form a group

Suppose we have a set G of integrators which may or may not be consistent. If, for all integrators ψ_{τ} and χ_{τ} in G, we have

$$\psi_{\tau} \circ \chi_{\tau} \in G$$

and

$$\psi_{\tau}^{-1} \in G,$$

we say the integrators form a group. That is, they are a group where the group operation is composition of maps.

Examples of integrators that can form a group are

- (i) symplectic integrators (Lecture 3)
- (ii) symmetry-preserving integrators (Lecture 4)
- (iii) volume-preserving integrators (Lecture 5)
- (iv) integral-preserving integrators (Lecture 6)

For example, for the group of integral-preserving integrators there is a real function I(x) (the *integral*) such that $I(x) = I(\varphi_{\tau}(x))$ for all x: the value of the integral I is preserved by the integrator. Therefore it is also preserved by $\varphi_{\tau} \circ \chi_{\tau}$ and by φ_{τ}^{-1} : the integrators form a group.[†]

These groups are infinite-dimensional groups of diffeomorphisms. They share many but not all of the properties of Lie groups; various extensions of the concept of Lie groups from finite to infinite dimensions have been proposed. One approach is the theory of "Lie pseudogroups" of diffeomorphisms. Cartan discovered in 1913 that in a sense there are just 6 fundamental Lie pseudogroups: the group of all diffeomorphisms; those preserving a symplectic, volume, or contact structure; and those preserving a symplectic or volume structure up to a constant. These correspond to different generic types of dynamics.

† If φ_{τ}^{-1} exists, which it does for the methods of Lecture 6, but not necessarily for projection methods.

How to construct integrators that form a group

The main way to construct integrators that form a group is through *splitting methods*. Splitting methods work for all cases (1)-(4) above, and are discussed further in Lecture 3.

We illustrate splitting for integral-preserving integrators. Assume we don't know an integral-preserving integrator for the vector field f, but f can be split into two vector fields f_1 and f_2 , each with the same integral as f:

$$f(x) = f_1(x) + f_2(x)$$

and assume that we do know integral preserving integrators ψ_1 (resp. ψ_2) for f_1 (resp. f_2) separately.

Then we obtain an integral-preserving integrator ψ for f by composition:

$$\psi_{\tau} = \psi_{2,\tau} \circ \psi_{1,\tau}$$

This is a consistent method for f, because it is the map $\psi_{\tau}: x \mapsto x''$ given by

$$\begin{aligned} x' &= x + \tau f_1(x) + \mathcal{O}(\tau^2) \\ x'' &= x' + \tau f_2(x') + \mathcal{O}(\tau^2) \\ &= x + \tau (f_1(x) + f_2(x)) + \mathcal{O}(\tau^2) \end{aligned}$$

Splitting methods are very easy to program—one merely calls routines for ψ_1 and ψ_2 in turn. Thus the problem becomes:

- (i) How to split vector fields while staying in the appropriate class;
- (ii) How to construct integrators in the appropriate group;
- (iii) How to compose those integrators so as to get an integrator of the original vector field of the desired order.

Each of these will be considered in these lectures.

1.4 Integrators that form a symmetric space

Suppose we have a set G of integrators with the property that, for all integrators ψ_{τ} and χ_{τ} in G, we have

$$\psi_{\tau} \circ \chi_{\tau}^{-1} \circ \psi_{\tau} \in G. \tag{1.7}$$

Then G is an example of the algebraic object known as a symmetric space, a set G together with a binary operation * obeying the axioms

$$\begin{aligned} x*x &= x\\ x*(x*y) &= y\\ x*(y*z) &= (x*y)*(x*z)\\ *y &= y \Rightarrow y = x \text{ for all } y \text{ sufficiently close to } x\end{aligned}$$

In our case the integrators G form a symmetric space by taking

x

$$\psi_{\tau} * \chi_{\tau} := \psi_{\tau} \circ \chi_{\tau}^{-1} \circ \psi_{\tau}.$$

Notice that every group also forms a symmetric space, but not vice versa: a group may have subsets which are closed under (1.7) but not under simple composition.

- The two most important examples of integrators that form a symmetric space are
- (i) Self-adjoint integrators.
- (ii) Integrators that possess time-reversal symmetry (Lecture 4).

Proof of (1): Let $\theta_{\tau} = \psi_{\tau} \circ \chi_{\tau}^{-1} \circ \psi_{\tau}$. Then

$$\begin{aligned} \theta_{-\tau}^{-1} &= \left(\psi_{-\tau} \circ \chi_{-\tau}^{-1} \circ \psi_{-\tau}\right)^{-1} \\ &= \psi_{-\tau}^{-1} \circ \chi_{-\tau} \circ \psi_{-\tau}^{-1} \\ &= \psi_{\tau} \circ \chi_{\tau}^{-1} \circ \psi_{\tau} \\ &= \theta_{\tau}. \end{aligned}$$

General tools

How to construct integrators that form a symmetric space

There are two main ways:

(i) (Projection methods) If ψ_{τ} is any integrator, then the "projection"

$$\chi_{\tau} := \psi_{\tau/2} \circ \psi_{-\tau/2}^{-1}$$

is self-adjoint.

(ii) (Splitting methods) If f can be split into two vector fields

$$f(x) = f_1(x) + f_2(x)$$

such that we have self-adjoint integrators ψ_1 and ψ_2 for f_1 and f_2 separately, then we obtain a self-adjoint integrator ψ for f from the symmetric composition

$$\psi_{\tau} := \psi_{1,\tau/2} \circ \psi_{2,\tau} \circ \psi_{1,\tau/2}.$$

These are generalized to other symmetric spaces in Lecture 4. The projection is almost miraculous, because it starts with *any* integrator. There is no analogous projection for groups.

1.5 Integrators that form a semigroup

A set G of integrators forms a semigroup if for all integrators ψ_{τ} and χ_{τ} in G, we have $\psi_{\tau} \circ \chi_{\tau} \in G$, but not necessarily $\psi_{\tau}^{-1} \in G$.

These arise from properties that only hold for forwards time:

- (i) systems with a Lyapunov function (Lecture 6)
- (ii) systems which contract phase space volume

For example, if the Lyapunov function is decreasing as t increases, it is *increasing* as t decreases, and even the flow $\varphi_{\tau}^{-1} = \varphi_{-\tau}$ does not have the Lyapunov property. This means one cannot use backwards time steps when composing these integrators, which can be proved to limit the order of composition methods to 2.

1.6 Creating higher order integrators: composition methods

Having obtained a geometric integrator ψ_{τ} , a higher order method can be obtained from the composition

$$\chi_{\tau} = \psi_{\alpha_{n}\tau} \psi_{-\alpha_{n-1}\tau}^{-1} \psi_{\alpha_{n-2}\tau} \psi_{-\alpha_{n-3}\tau}^{-1} \cdots \\ \cdots \psi_{\alpha_{n-3}\tau} \psi_{-\alpha_{n-2}\tau}^{-1} \psi_{\alpha_{n-1}\tau} \psi_{-\alpha_{n}\tau}^{-1}$$

which has been chosen to be self-adjoint, i.e. $\chi_{\tau} = \chi_{-\tau}^{-1}$. Here the number of integrators n and the coefficients α_n can be adjusted to obtain the desired order. High order methods can also be designed in the context of splitting methods, using the two flows $\varphi_{1,\tau}$ and $\varphi_{2,\tau}$ of f_1 and f_2 respectively. One uses the composition

$$\varphi_{1,\beta_1\tau}\varphi_{2,\gamma_1\tau}\cdots\varphi_{1,\beta_n\tau}$$

However, these two approaches turn out to be equivalent [7].

Example 1 If ψ is self-adjoint, then a fourth-order integrator is obtained as follows:

$$\chi_{\tau} = \psi_{\gamma\tau} \circ \psi_{(1-2\gamma)\tau} \circ \psi_{\gamma\tau}$$

where $\gamma := (2 - 2^{1/3})^{-1}$.

This example is generalized in Theorem 2 below. Note that $1 - 2\gamma < 0$.

1.6 Composition methods

An example of composition methods: the generalized Yoshida method

Theorem 2 (Yoshida, Qin, and Zhu) Let ψ be a self-adjoint integrator of order 2n. Then

$$\chi_{\tau} = \psi_{\gamma\tau} \circ \psi_{(1-2\gamma)\tau} \circ \psi_{\gamma\tau}, \quad \gamma = (2 - 2^{\frac{1}{2n+1}})^{-1}$$

is a self-adjoint integrator of order 2n + 2.

Proof Let

$$\psi_{\tau}(f) = \varphi_{\tau}(f) + \delta \tau^{2n+1} + \dots$$

Then, using the flow property of the exact flow φ_{τ} ,

$$\psi_{\gamma\tau} \circ \psi_{(1-2\gamma)\tau} \circ \psi_{\gamma\tau} = \varphi_{\tau}(f) + (\gamma^{2n+1} + (1-2\gamma)^{2n+1} + \gamma^{2n+1}) \,\delta\tau^{2n+1} + \dots$$

which has order 2n + 1 if γ is as given in the theorem. However, it is self-adjoint by construction, so it has even order, hence the order is 2n + 2.

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Composition methods

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2 Why preserve structure?

2.1 Introduction

Let's start with an example of a simulation of the outer solar system by Jack Wisdom and coworkers. Part of its appeal is the long history of modelling the solar system. The people who do this are not from the numerical analysis community, but they have their own history of methods which they have developed and tweaked.

In the 1980's, a special-purpose supercomputer, the "Digital Orrery", simulated the outer planets for 845 million years. With a lot of tweaking, an energy error of about 10^{-9} was achieved with a time step of 45 days (a six month calculation!). A calculation with a very high order symmetric multistep method achieved an energy error of about 10^{-10} in a 3 million year simulation, with a time step of 0.75 days. In a completely different approach, Laskar (1990) used classical perturbation theory (expanding in mass ratios and eccentricities about circular orbits) to eliminate the fast (annual) frequencies. This required 250,000 terms, but a time step of 500 years could be taken.

All of these attempts were roundly routed by the calculation of Jack Wisdom et al., using a very simple, elegant symplectic integrator. Their billion year simulation with a time step of 7.5 days gave an energy error of only 2×10^{-11} . Moreover, only one force evaluation was used per time step, making the method very fast.

Roughly speaking, they wrote the ODE as a sum of uncoupled Kepler 2-body problems and the potential which couples the planets: $f = f_1 + f_2 = f_{\text{Kepler}} + f_{\text{coupling}}$. Each f_i is a Hamiltonian system, and the flow $\varphi_{\tau,i}$ of each can be found exactly and quickly (the 2-body problems using an elegant method of Gauss).



Fig. 2.1. Energy error of leapfrog applied to the whole solar system over 10^8 years (Wisdom et al.)



Fig. 2.2. Energy error after application of corrector χ_{τ} .

The time stepping is simply the simplest composition $\psi_{\tau} = \varphi_{\tau,2} \circ \varphi_{\tau,1}$ —a form of the "leapfrog" method. Since the flow of Hamiltonian ODEs is symplectic, and symplectic maps form a group, ψ_{τ} is symplectic. Moreover, they found a "corrector" χ_{τ} such that

$$\chi_{\tau} \circ \psi_{\tau} \circ \chi_{\tau}^{-1} = \varphi_{\tau} + \mathcal{O}(m^2 \tau^3)$$

where $m \approx |f_2/f_1| \approx 10^{-3}$ is the mass ratio between Jupiter and the sun. (The result after *n* time steps is $\chi_\tau \circ \psi_\tau^n \circ \chi_\tau^{-1}$, so that χ_τ only needs to be evaluated once, no matter how long the simulation.) This method:

- is symplectic;
- is one-step;
- is explicit;
- is second order;
- uses one force evaluation per time step;
- exploits classical analysis, namely the exact solution of the 2-body problem;
- preserves total linear and angular momentum;
- is self-adjoint and reversible;
- has an extra factor of $m^2 = 10^{-6}$ in its local truncation error, compared to classical methods;
- for moderate times ($\approx 2 \times 10^7$ years), has linear growth of global errors, compared to quadratic growth for classical methods;
- has bounded energy errors for long times.

This is almost a dream situation, where geometric integration has lead to a simple method with vastly improved local (time τ), global (time T), and structural (time ∞) errors. This calculation discovered chaos in the outer solar system with a Lyapunov time, the time for the separation between nearby orbits to grow by a factor e, of 20 million years. Over the billion year calculation, they would separate by $e^{50} \approx 10^{22}$, and integration errors would be magnified by this amount also. Thus, the final angular positions of the planets are not expected to be accurate. However, we can be confident that the qualitative or statistical properties of the solution are correct.

2.2 Phase space and phase flow

"In phase space, no one can hear you scream."

(Caltech T-shirt)

The fundamental idea to keep in mind is to think in phase space. It's a simple idea but one which you

Why preserve structure?



Fig. 2.3. Inclination of Pluto over 10^9 years, showing chaos. Even after 10^9 years the inclination has reached a new maximum.

have to keep reminding yourself of: a simple definition in a dynamical systems class just isn't enough. Considering that differential equations were studied for 200 years before Poincaré adopted this point of view, this may not be too surprising.

"Consider the fluid molecules which initially form a certain figure F_0 ; when these molecules are displaced, their ensemble forms a new figure which will be deformed in a continuous manner, and at the instant t the envisaged ensemble of molecules will form a new figure F."

(Poincaré, Celestial Mechanics, 1899)

In a trajectory $\varphi_t(x_0)$, one thinks of the initial condition x_0 as fixed, and the time t increasing; in the flow map $\varphi_\tau(x)$, one thinks of all initial conditions x flowing forward for some fixed time τ . We'll only consider one-step methods, so that the numerical approximation for one time-step τ is a map

$$\psi_{\tau}: \mathbb{R}^m \to \mathbb{R}^m.$$

Now classical approximation theory, e.g. for Runge-Kutta methods, shows that chaos always wins: the best bound that can be obtained in general for a method of order p is

$$\left|\psi_{\tau}^{T/\tau}(x) - \varphi_{T}(x)\right| \leq (\Delta t)^{p} C \frac{e^{\Lambda T} - 1}{\Lambda}$$

The precise value of Λ depends on the Lipschitz constant of the vector field and on the method, but $\Lambda > 0$ and consequent exponential growth of error cannot be avoided in general. But dynamical systems theory teaches that ψ can be "close" to φ in other ways: their phase portraits may be qualitatively or even quantitatively similar; the stability of their orbits may be the same; for strange attractors, their Lyapunov exponents or fractal dimensions may be close.

The pendulum: theory

Systems can have many geometric or structural properties. Before we get into definitions, let's look at the planar pendulum. It is a two-dimensional system with phase space \mathbb{R}^2 , and dynamics

$$\dot{q} = p, \quad \dot{p} = -\sin q \tag{2.1}$$

where q is the angle of the pendulum, and p its angular momentum. (Here we are taking $q \in \mathbb{R}$, the covering space of the actual angle.) Here are some of the properties of the pendulum:

- It conserves the total energy $\dot{H} = \frac{1}{2}p^2 \cos q$. That is, its flow stays on the level sets of this function. Because this is a two-dimensional system, these level sets are curves in the plane.
- Being a Hamiltonian system, its flow is symplectic. For two-dimensional systems, this is equivalent to being area-preserving.



Fig. 2.4. Phase portrait and flow of the pendulum (from Hairer and Wanner). The area of each cat is preserved in time, the manifestation of symplecticity. Energy, whose levels sets are the curves shown, is preserved. Rotation by 180° $((q,p) \mapsto (-q,-p))$ is a symmetry, while flipping up-down $(p \mapsto -p)$ is a reversing symmetry.

It has one discrete symmetry and one discrete reversing symmetry (see Lecture 4). The symmetry, (q, p) → (-q, -p), maps the vector field into itself; the reversing symmetry, (q, p) → (q, -p), maps the vector field into minus itself. Imagining flowing along one of the solution curves, you can see that the motion of the reflected points is constrained.

Because this is such a simple system, preserving any of these three properties gives a geometric integrator with good long-time behavior for almost all initial conditions. A picture of its phase portrait will look very similar to the true phase portrait; we'll see examples of this in Section 2.7. By contrast, standard methods (e.g. Euler's method) destroy the qualitative phase portrait completely.

2.3 Philosophy of geometric integration

In any numerical study, one should

- examine any geometric or structural properties of the ODE or its flow;
- design numerical methods which also have these structural properties; and
- examine the consequences, hopefully over and above the immediate ones.

This encourages us to

- confront questions of phase space and degrees of freedom;
- think about the significance of local, global, and qualitative errors; and
- think about the kinds of tools and functions allowed in numerical analysis.

For example, multistep methods do not define a map on phase space, because more than one initial condition is required. They can have geometric properties, but in a different (product) phase space, which can alter the effects of the properties. (See Fig. 2.12.) This puts geometric integration firmly into the "single step" camp. If a system is defined on a sphere, one should stay on that sphere: anything else introduces spurious, non-physical degrees of freedom.

The *direct* consequences of geometric integration are that we are

• studying a dynamical system which is close to the true one, and in the right class; and



Fig. 2.5. Phase portrait of a symplectic integration, from Channell and Scovel. 10^5 time steps for 10 different initial conditions are shown. Smooth curves ("KAM tori") correspond to regular, quasiperiodic motion; clouds correspond to chaotic motion.

• this class may have restricted orbit types, stability, and long-time behavior.

In addition, because the structural properties are so natural, some *indirect* consequences have been observed. For example,

- symplectic integrators have good energy behavior;
- symplectic integrators can conserve angular momentum and other conserved quantities;
- geometric integrators can have smaller local truncation errors for special problems, and smaller global truncation errors for special problems/initial conditions (even though they're larger in the "generic" case);
- some problems (particle scattering, isospectral problems) can have errors tending to zero at long times.

Here's a pictorial survey showing what you can expect from geometric integration. Fig. 2.5 appears in Channell and Scovel [3], one of the first symplectic integration papers. Orbits starting on the smooth curves ("invariant circles") stay on them forever. Of course, the orbit may be going around the circle at the wrong speed, but the "orbital error" does not grow in time. Compare this to the traditional approach to numerical integration, with its overwhelming emphasis on the estimation and control of local errors. The idea that errors grow in time and, once committed, cannot be undone, was deeply ingrained. Pictures like Fig. 2.5 did a lot to revise this traditional point of view.

Other orbits in Fig. 2.5 are chaotic, and their position errors grow exponentially. But, they can never jump across the invariant circles, and because it's the *right kind* of chaos (namely, the solution of some nearby Hamiltonian system), statistical observations of this chaos will have small errors.

2.4 Types of geometric properties

Study the list in the Table. The left hand column gives properties of vector fields, and the right hand column gives the corresponding properties of their flow. It's the right hand property that must be preserved by the integrator. Usually the flow properties are named the same as the ODE property.

(The standard example of a *Lie group* G is the set of orthogonal 3×3 matrices, $A^T A = I$, which represent rotations. Its Lie algebra \mathfrak{g} is the set of antisymmetric 3×3 matrices. G is a manifold whereas \mathfrak{g} is a linear space, a much simpler object to work with.)

To bring some order to this table, consider the following features.

• Is the structure linear in some sense?

All of the ODE properties are linear in f, but all of the flow properties are nonlinear in φ , except

ODE $\dot{x} = f(x)$		flow φ_t , derivative $d\varphi_t$
Hamiltonian Poisson source-free symmetric reversible Lie group isospectral integral dissipative	$\begin{split} f &= J\nabla H(x), \ J = \begin{pmatrix} 0 & I \\ -I & 0 \end{pmatrix} \\ f &= J(x)\nabla H(x) \\ \nabla \cdot f &= 0 \\ dS.f &= f \circ S \\ -dR.f &= f \circ R \\ f &= a(x)x, \ x \in G, \ a \in \mathfrak{g} \\ f &= [b(x), x], \ x, \ b \in \mathfrak{g} \\ f \cdot \nabla I &= 0 \\ f \cdot \nabla V &\leq 0 \end{split}$	$d\varphi^{T} J d\varphi = J \text{ (symplectic)}$ $d\varphi^{T} J d\varphi = J \circ \varphi$ $\det d\varphi = 1 \text{ (volume preserving)}$ $S \circ \varphi = \varphi \circ S$ $R \circ \varphi^{-1} = \varphi \circ R$ $\varphi \in G$ $eigenvalues \lambda(x) \text{ constant}$ I(x(t)) = I(x(0)) $V(x(t)) \leq V(x(0))$



for linear symmetries. Symplecticity, Poisson, and reversibility are quadratic; volume preservation and isospectrality are degree m when $x \in \mathbb{R}^m$.

- Does the structure appear explicitly or implicitly in the ODE? Hamiltonian, Poisson, Lie group, and isospectral ODEs are explicit (e.g. $f = J\nabla H$ generates all Hamiltonian ODEs); the rest are implicit—there are side conditions which f has to satisfy.
- Does the flow property depend on φ or dφ?
 Symplecticity, Poisson, and volume preservation depend on the Jacobian dφ. This makes them harder to preserve.

These will be explored further in the other lectures. Briefly, it is easier to work on *linear* and *explicit* properties, so we concentrate on bringing all flow properties into this form. (See §3.1 on splitting.) This has been achieved for all the properties in the Table, but not for some of their nonlinear generalisations and combinations.

A major justification for geometric integration comes from *backward error analysis*. This theoretical tool writes the integrator ψ_{τ} as the time- τ flow of *some* vector field \tilde{f} , i.e. $\psi_{\tau}(f) = \varphi_{\tau}(\tilde{f})$. If the method is of order p, we have $\tilde{f} = f + \mathcal{O}(\tau^p)$. Then, in many cases one can argue that since ψ_{τ} is in some class (e.g. symplectic), the perturbed vector field must be in the appropriate class too (e.g. Hamiltonian). So we know that by studying the dynamics of the method, we are at least studying dynamics in the right class. The reliability of the results then depends on the "structural stability" of the original system: a difficult problem, but a standard one in dynamical systems.

In the Hamiltonian case, $\tilde{f} = J\nabla \tilde{H}$ for some Hamiltonian \tilde{H} , which is conserved by the method. Since we don't know \tilde{H} and can only measure the original energy H, it (H) will be seen to oscillate, but (if the levels sets of H and \tilde{H} are bounded) will not drift away from its original level.

Technically, one suspends the map ψ_{τ} to a time-dependent flow $\varphi_{\tau}(\tilde{g}(x,t))$, from which, when ψ_{τ} is analytic, nearly all the time dependence can be removed by a change of variables, giving $\tilde{f}(x) + \mathcal{O}(e^{-1/\tau}, t)$. This introduction of an exponentially small nonautonomous term is inevitable, because most maps, even those close to the identity, are not actually flows. If the time step is too large these exponentially small terms can actually pollute the calculation, and one observes, for example, the energy drifting.

2.5 Miscellaneous topics

Some other branches of geometric integration are

• ODEs on manifolds, such as homogeneous spaces. Although ultimately one can only compute in a linear space, it's best to formulate the method on the manifold and transfer to coordinates as late as possible. A special case is when the manifold is a Lie group [4]; Lie group methods are one of the major themes in geometric integration which we don't have space to discuss here.



Fig. 2.6. Flow on a family of invariant tori. From V.I. Arnol'd, Small denominators and problems of stability of motion in classical and celestial mechanics, *Uspehi Mat. Nauk (Russ. Math. Surv.)* **18** (1963) no. 6 (114) 91–192.

- Mapping methods approximate the equations in x as well as in t, for example, by Taylor series. Maps defined by series can then be manipulated analytically.
- When evaluating *Lyapunov exponents* one should try to preserve their structure, e.g., that the Jacobians used are symplectic or volume-preserving.
- For *partial differential equations* one can either discretize in space first, seeking a finite-dimensional version of, e.g., the Hamiltonian structure, or discretize space-time directly.
- One can discretize phase space itself and study *lattice maps*, a form of cellular automata. This has been used in studies of the effect of roundoff error.
- Instead of trying to construct special methods that preserve particular properties, one can see how well standard methods do. Usually the property has to be fairly robust, e.g., dissipation of the type $d|x|^2/dt < 0$ for |x| > R is studied, instead of $dV/dt \le 0$ for all x. This approach is thoroughly treated in Stuart and Humphries, Dynamical Systems and Numerical Analysis.

2.6 Growth of global errors

The global error is $\psi_{\tau}^{T/\tau}(x) - \varphi_{\tau}(x)$ where T is a large, but fixed, time. Geometric integrators are not expressly designed to control the global error. Nevertheless, sometimes it grows linearly in a symplectic integrator and quadratically in a standard integrator. This will make the symplectic integrator superior if T is large enough.

This property has been observed in many systems of different types. It is associated with preservation of *invariant tori* by the method. An invariant torus is a subset of initial conditions, topologically a torus, which orbits starting on stay on for all forwards and backwards time. A torus is *preserved* if the integrator has an invariant torus of its own, which tends to the torus of the ODE as $\tau \to 0$.

Invariant tori

Invariant tori are ubiquitous in dynamics. They're found in:

- Hamiltonian systems (tori have dimension n/2);
- reversible systems (when orbits intersect the symmetry plane; tori often have dimension n/2);
- volume-preserving systems (tori have any dimension < n).

They are important because they

- form positive-measure families of neutrally stable orbits, which
- mostly persist under small perturbations of the system;
- form "sticky sets," dominating behavior of nearby orbits on intermediate time scales

Nearby orbits diverge like

• $\mathcal{O}(1)$ on same torus



Fig. 2.7. Cross-section of the tori in Fig 2.6 after perturbation (Arnol'd). Some are destroyed and replaced by chaos, some persist.

- $\mathcal{O}(T)$ on a nearby or perturbed torus
- $\mathcal{O}(T^2)$ if $\mathcal{O}(T)$ drift across tori
- $\mathcal{O}(T, e^{\lambda T})$ on nearby chaotic orbits; λ depends on the order of resonance, but can be very small.

Therefore, in an integrator we should try to preserve tori of the correct dimension. In a standard method, they are not preserved, and orbits drift transversely, leading to $\mathcal{O}(T^2)$ growth of global errors. If the torus is preserved, orbits only move around the torus at a slightly wrong angle or speed, leading to $\mathcal{O}(T)$ errors.

It turns out to be an extraordinarily subtle question to determine when which tori persist under which perturbations. Finally, in the 1960's, conditions were found by Kolmogorov, Arnol'd, and Moser under which most tori do persist under appropriate perturbations, although some are destroyed. This forms the subject of KAM theory.

For Hamiltonian systems, an appropriate perturbation is Hamiltonian, so the results apply to symplectic integrators.

In between invariant tori, or if tori were destroyed by taking too large a time step, orbits can be chaotic. But, because of the nearby tori, exponential separation can be very slow, and the linear error growth can dominate for long times.

2.7 The pendulum: numerical experiments

We illustrate the above points on the simplest meaningful example, the pendulum (Eq. (2.1)). The simplest symmetric, reversible, self-adjoint symplectic method is leapfrog:

$$q' = q + \frac{1}{2}\tau p$$
$$p' = p - \tau \sin q$$
$$q'' = q' + \frac{1}{2}\tau p'$$

The results of this method are shown in Fig. 2.8 for a small time step ($\tau = 0.1$) and in Fig. 2.9 for a much larger time step ($\tau = 1$). Even for the larger time step, the left-right and up-down symmetries are preserved, as are most of the invariant circles, as promised by KAM theory for symplectic integrators. Chaos is significant only in a small neighbourhood of the homoclinic orbit connecting (π , 0) and ($-\pi$, 0).

A symplectic method which is not symmetric or self-adjoint is shown in Fig. 2.10; the lack of symmetry

Why preserve structure?



Fig. 2.8. 1000 times steps of symplectic leapfrog applied to the pendulum, time step $\tau=0.1.$



Fig. 2.9. As in Fig. 2.8, but $\tau = 1$.



Fig. 2.10. A nonsymmetric symplectic integration of the pendulum, τ = 0.1.



Fig. 2.11. 10^6 time steps of leapfrog at $\tau = 1$, showing a chaotic orbit.



Fig. 2.12. A symplectic multistep method: the torus has dimension 2 instead of 1 as in Figs. 2.8–2.10. A single orbit is shown, with the first 50 time steps marked by \times .

is plain to see. In this case, invariant circles are still preserved. In higher-dimensional systems, there is a more complicated interaction between symplecticity and reversibility.

What is the effect of the chaos created by the numerical integrator? Fig. 2.11 shows one chaotic orbit of leapfrog at the large time step $\tau = 1$, obtained with initial condition (q, p) = (0, 1.8). It was found to have a large Lyapunov exponent of 10^{-2} . By $T \sim 100$, the chaos would dominate the numerical errors. by contrast, with initial condition (q, p) = (0, 1.6), the Lyapunov exponent is already reduced to 10^{-7} , and phase errors (moving around the circle at the wrong speed) would dominate until $T \sim 10^7$. Thus, even when the numerical orbit does not lie on an invariant torus, the preservation of *some* invariant tori nearby helps a great deal.

In Section 2, we talked about the importance of staying in the right phase space. The multistep method $x_{n+1} = x_{n-1} + 2\tau f(x_n)$ is a map on the *product* phase space $\mathbb{R}^2 \times \mathbb{R}^2$. It can be shown to be symplectic in this larger space, but its KAM tori have dimension 2, instead of 1 as in the real system. When projected to the original phase space, they fill out a solid region, instead of a curve—a disaster for long-time simulations. This effect is illustrated in Fig. 2.12.

2.8 Summary

Systems may have many geometric or structural features. Integrators must balance costs, local, global, and long-time errors, stability, and structural preservation. You can't expect to do well at all of these

simultaneously! Also, numerical studies can have different goals. Demanding very small local errors for a large class of ODEs tilts the balance in favour of highly-developed standard methods; seeking reliability over long times with simple, fast methods tilts in favour of geometric integrators.

The remaining lectures look at preserving different properties. Here we sum up what is known about preserving several properties at once.

- (i) Symplecticity and energy: If, by "integrator", we mean that the method is defined for all Hamiltonian ODEs, then by a theorem of Ge, this is impossible. For exceptional ("completely integrable") problems, such as the free rigid body, this can be done.
- (ii) Symplecticity and integrals apart from energy: Not known, although doable in principle.
- (iii) Symplecticity and linear symmetries: Achieved by, e.g., the implicit midpoint rule.
- (iv) Poisson and linear symmetries: Not known.
- (v) Volume preservation and linear symmetries: Not known.
- (vi) Integrals and linear symmetries: Sometimes possible using the Harten, Lax and Van Leer discrete gradient (see Lecture 6).
- (vii) Volume and an integral: Can be done by splitting for all systems with some integrals and for some systems with any integrals. Not known in general.

Extending the concept of geometric integration to PDEs is much less developed. Work has been done, e.g., on integral preservation [7], symmetry preservation [8], and Lagrangian (variational) structure [9].

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Symplectic integrators: A case study of the molecular dynamics of water.

"Chemistry is a science, but not Science; for the criterion of true science lies in its relation to mathematics" (Kant)

"Chemistry will only reach the rank of science when it shall be found possible to explain chemical reactions in the light of their causal relations to the velocities, tensions and conditions of equilibrium of the constituent molecules; that the chemistry of the future must deal with molecular mechanics by the methods and in the strict language of mathematics, as the astronomy of Newton and Laplace deals with the stars in their courses" (Du Bois Reymond)

This quote (from D'Arcy Thompson's *On Growth and Form*) could not be more apt: symplectic integrators, developed to deal with the stars in their courses, are now applied to the velocities of molecules.

There are many fine surveys of symplectic integration, so here we'll discuss *Poisson systems*, or noncanonical Hamiltonian systems, and how they arose in a study of water. Water, the "king of polar fluids," has many strange phases and anomalous properties, which statistical mechanics has a hard time explaining. Therefore people turn to numerical simulations.

3.1 Splitting

Recall the problem of splitting—how can we write $f = f_1 + f_2$ so that the f_i retain some properties of f? The idea is to represent all f in the given class explicitly by a "generating function." Then we split the generating function. This can be done for Hamiltonian systems by splitting the Hamiltonian. Look at Table 2.1: Hamiltonian systems are expressed explicitly.

Example 3 Hamiltonian systems. The generating function is the Hamiltonian H.

$$f = J\nabla H = J\nabla \left(\sum_{i} H_{i}\right) = J\nabla H_{1} + \ldots + J\nabla H_{n}.$$

Properties due to J, which is not split, are retained—symplecticity. Properties due to H, which is split, are lost—conservation of H.

Example 4 Systems with an integral. The generating function is the skew-symmetric matrix function J.

$$f = J\nabla H = \Big(\sum_{i} J_i\Big)\nabla H = J_1\nabla H + \ldots + J_n\nabla H$$

Properties due to J, which is split, are lost—symplecticity. Properties due to H, which is not split, are retained—conservation of H.

We'll return to systems with an integral in Lecture 6, and see how to apply splitting to volume-preserving systems in Lecture 4.

3.2 Poisson systems

Consider a standard, canonical Hamiltonian system.

$$\dot{x} = J \nabla_x H(x), \quad J = \begin{pmatrix} 0 & I \\ -I & 0 \end{pmatrix}.$$

It only has this special form when written in special variables. If we apply an arbitrary change of variables, writing the system in terms of y = g(x), it becomes

$$\begin{split} \dot{y} &= dg \cdot \dot{x} \\ &= dg \cdot J \nabla_x H(x) \\ &= dg \cdot J \cdot dg^T \nabla_y H(x) \\ &= \widetilde{J}(y) \nabla_y \widetilde{H}(y), \end{split}$$

where

$$\widetilde{J} = dg \cdot J \cdot (dg)^T$$
$$\widetilde{H}(y) = H(x).$$

This is an example of a "Poisson system," the most obvious change being that the matrix J now depends on y. However, the class of Poisson systems is invariant under changes of variables. Since the history of mathematics and of physics is a history of requiring invariance under more operations, it seems we should study Poisson systems in their own right.

(There are many other motivations for the introduction of Poisson systems, from PDEs, systems on Lie groups and other manifolds, and symmetry reduction.)

An important special case are the "Lie"-Poisson systems. Let $x \in \mathbb{R}^m$ be an element of a Lie algebra. Let $[x_i, x_j] = \sum_{k=1}^m c_{ij}^k x_k$ be the Lie bracket. Let $J_{ij} = [x_i, x_j]$, so that the entries of J are linear functions of x. Then

$$\dot{x} = J(x)\nabla H(x)$$

or

$$\dot{x}_i = \sum_{j,k} c_{ij}^k x_k \frac{\partial H}{\partial x_j}$$

is called a Lie-Poisson system.

Example 5 The free rigid body in \mathbb{R}^3 . The variables are π_1 , π_2 , π_3 , the angular momenta of the body in body-fitted coordinates.

$$J = \begin{pmatrix} 0 & \pi_3 & -\pi_2 \\ -\pi_3 & 0 & \pi_1 \\ \pi_2 & -\pi_1 & 0 \end{pmatrix}$$
$$H = \frac{1}{2} \left(\frac{\pi_1^2}{I_1} + \frac{\pi_2^2}{I_2} + \frac{\pi_3^2}{I_3} \right)$$

Here the Lie algebra is so(3), the antisymmetric 3×3 matrices.

3.3 Splitting into solvable pieces

Earlier we showed how to split a vector field into appropriate pieces, and how to compose their flows. But, it is still important to be able to apply a geometric integrator to each piece. Here we achieve this by requiring the pieces to be (easily) integrable.

Observation I If $J_{ij} = 0$ for $1 \le i, j \le k < n$ and $H = H(x_1, \ldots, x_k)$, then the ODEs are

$$\dot{x} = \begin{pmatrix} 0 & * \\ * & * \end{pmatrix} \begin{pmatrix} * \\ 0 \end{pmatrix}$$

$$\dot{x}_{i} = \begin{cases} 0 & i = 1, 2, \dots, k \\ \sum_{j,l} c_{ij}^{l} x_{l} f_{j}(x_{1}, \dots, x_{k}) & i = k+1, \dots, m \end{cases}$$

which are linear with constant coefficients, hence easily solved (although the coefficients depend parametrically on the other variables x_1, \ldots, x_k).

Observation II Systems with $H = \sum_{i} H_i(x_i)$ can be split into easily solved parts. The rigid body has this form.

Observation III Quadratic Hamiltonians can be diagonalized, i.e., put in the form of Observation II, and hence split.

3.4 Molecular dynamics

The basic steps in a molecular dynamics simulation are the following.

- (i) Take a large sea of particles.
- (ii) Impose boundary conditions (e.g. 3D periodic) and impose constancy of any three out of the four quantities pressure, volume, temperature, and number of particles; the fourth is determined.
- (iii) Find a classical model of the interparticle forces.
- (iv) Move the particles for a long time.
- (v) Collect statistics of the motion.

Applications are to exploring states of matter (phase transitions, liquid, liquid crystal, colloidal), protein folding, the design of large organic molecules and drugs, nanotechnology. It's a big field.

Current limits are about 10^8 simple atoms on a supercomputer, 10^5 simple atoms on a workstation, and 10^2-10^3 water molecules on a workstation. For water, even $27 = 3 \times 3 \times 3$ molecules with periodic boundary conditions are enough to see solid, liquid, and gas phases.

What's the best way to move the particles? The method should

- obey the physical laws;
- exhibit the correct statistical equilibrium in the face of chaos; and
- be fast and cheap, since forces are expensive to evaluate.

In fact, the forces are *so* expensive that users don't want to evaluate them more than once per time step. For decades they've been using the Verlet method for point masses:

$$H = \text{kinetic} + \text{potential} = \frac{1}{2}p^2 + V(q),$$
$$q_{n+1} = q_n + \tau p_n$$
$$p_{n+1} = p_n - \tau \nabla V(q_{n+1})$$

We now know that it's so good because it's the simplest symplectic integrator, and comes from splitting the Hamiltonian.

How can we extend the Verlet algorithm to non-symmetric molecules like water? Many approaches have been considered.

- Move each atom separately. This involves modelling the interatomic forces, which means simulating the many modes of vibration within each molecule. Their time scale is very short and they are not believed to affect the macroscopic properties of water, which rules out this approach.
- Model as a rigid body. This is the preferred option. It can be done in various ways:
- Consider the molecule as a set of particles subjected to constraints on interatomic distances and angles. This is possible, but constraints lead to expensive, implicit methods. They are needed for problems involving flexible chains such as proteins.
- Model as a rigid body, with orientation represented by Euler angles or unit quaternions. The Hamiltonian in these variables is complicated and nonseparable; this makes symplectic methods expensive. In the most popular variant, unit quaternions are used and the ODEs are passed to a black-box solver.
- Represent orientation by a 3×3 orthogonal matrix, and update this by rotations only.

This last is what we now do. First, let's look at the free rigid body.



Fig. 3.1. The phase portrait of the free rigid body (from Bender & Orszag, Advanced Mathematical Methods for Scientists and Engineers.) Orbits on a sphere of constant angular momentum $|\pi|$ are shown. There are three pairs of fixed points, corresponding to rotation about each of the three principal axes; two are stable and one is unstable. To observe passage along the "homoclinic orbit" joining the unstable pair, hold a hammer with the head horizontal and toss so it rotates once about a left-right axis.

3.5 The free rigid body

Let the angular momentum vector be π , the orientation be $Q \in SO(3)$, i.e. $Q^T Q = Id$; the Hamiltonian for the free rigid body is

$$H = \frac{1}{2} \left(\frac{\pi_1^2}{I_1} + \frac{\pi_2^2}{I_2} + \frac{\pi_3^2}{I_3} \right).$$

The phase portrait for this system is shown in the famous Figure 3.1.

As noted above, splitting methods work excellently for Lie-Poisson systems with Hamiltonians of this type. The flow of $H = \frac{\pi_1^2}{2I_1}$ is

$$\pi(t) = R\pi(0)$$
$$Q(t) = Q(0)R^{T}$$

where

$$R = \begin{pmatrix} 1 & 0 & 0 \\ 0 & \cos\theta & -\sin\theta \\ 0 & \sin\theta & \cos\theta \end{pmatrix}, \quad \theta = t \frac{\pi_1}{I_1}.$$

This decomposes the motion into three elementary rotations. The method is fast, accurate, reversible and symplectic. Q is always orthogonal up to round-off error, because it is updated only by rotations.

How does this fit into a full simulation of water? For each molecule the variables are the q, the position of the centre of mass; p, the linear momentum; Q; and π . The total energy has the form

$$H = T^{\text{rotation}}(\pi) + T^{\text{translation}}(p) + V(q, Q)$$

together with the constraints $Q^T Q = Id$. We apply a Verlet-like splitting into kinetic and potential parts. For each molecule, we have

Potential part:

$$\begin{split} \dot{q} &= 0\\ \dot{Q} &= 0\\ \dot{p} &= -\frac{\partial V}{\partial q} =: f \quad \text{(force)}\\ \dot{\pi} &= (Q^T f) \times x \quad \text{(torque)} \end{split}$$

where x is the point at which the force acts. Since the positions and orientations are here held constant, these equations are easy to solve.

Kinetic part:

$$q = p$$

$$\dot{Q} = Q (skew(I^{-1}\pi))$$

$$\dot{p} = 0$$

$$\dot{\pi} = \pi \times I^{-1}\pi$$

where $I = \text{diag}(I_1, I_2, I_3)$ is the inertia tensor. The centres of mass undergo free, straight-line motion, while the orientations move as a free rigid body. The latter could be solved explicitly, although this has never been implemented in a production code; in practice, we approximate its flow by the previously-given splitting method.

Composing these pieces gives an analogue of the Verlet method for this non-canonical Hamiltonian system. The final method uses only one force evaluation per time step, but is still explicit, symplectic, reversible, and conserves total linear and angular momentum (because each piece does). As expected for such a method, energy errors are bounded in time. When implemented in the existing research code ORIENT using existing error criteria, this method was about ten times faster than the old method (2-level leapfrog with Bulirsch-Stoer extrapolation).

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4

Symmetries and reversing symmetries

4.1 Symmetries of ODEs

A symmetry is a map $h : \mathbb{R}^m \to \mathbb{R}^m$ from phase space to itself, such as $x \mapsto -x$. In a system with symmetries, the vector field at the two points x and h(x) are related to each other. This is shown for the pendulum in Fig. 2.4. Under the 180° rotation $(q, p) \mapsto (-q, -p)$, arrows (the vector field) map to arrows: a symmetry. Under the reflection $p \to -p$, arrows map to arrows if we also reverse their direction: a reversing symmetry. The analogous properties for flows can be seen by tracing along the flow lines.

Symmetries and reversing symmetries both reduce the possible complexity of the phase portrait, and should be preserved.

In reversible Hamiltonian systems, reversing symmetries are a bit easier to preserve than symplecticity (although one can have both, if desired). For example, for simple mechanical systems there are explicit, variable-step-size reversible methods.

Consider the ODE $\dot{x} = f(x)$ under the change of variables y = h(x). The new system is

$$\dot{y} = f(y) := ((dh \cdot f)h^{-1})(y).$$

Definition 6 The vector field f has symmetry h if $f = \tilde{f}$, i.e., if $dh \cdot f = fh$. The vector field f has reversing symmetry h if $f = -\tilde{f}$, i.e., $dh \cdot f = -fh$, and f is called reversible.

The notation fh indicates composition, i.e., (fh)(x) = f(h(x)).

Example 7 The pendulum. For the vector field

$$f: \dot{q} = p, \ \dot{p} = -\sin q$$

we have

$$h_1: \ \widetilde{q} = q, \ \widetilde{p} = -p \Rightarrow \widetilde{q} = -\widetilde{p}, \ \widetilde{p} = \sin \widetilde{q},$$

—a reversing symmetry;

$$h_2: \ \widetilde{q} = -q, \ \widetilde{p} = p \Rightarrow \widetilde{q} = -\widetilde{p}, \ \widetilde{p} = \sin \widetilde{q},$$

—a reversing symmetry; and so

$$h_1 \circ h_2: \ \widetilde{q} = -q, \ \widetilde{p} = -p \Rightarrow \dot{\widetilde{q}} = \widetilde{p}, \ \dot{\widetilde{p}} = -\sin \widetilde{q}$$

is a symmetry. So the pendulum has "reversing symmetry group" (the group of all symmetries and reversing symmetries)

$$\Gamma = \{ \mathrm{id}, h_1, h_2, h_1 h_2 \}.$$

In general, half of the elements of Γ are symmetries, and the composition of two reversing symmetries is a symmetry.

We will use S for a symmetry and R for a reversing symmetry.

Definition 8 The fixed set of S is

$$fix(S) := \{x : x = S(x)\}.$$

The fixed set is invariant under the flow of f. So preserving symmetries is one way of staying on a submanifold.

Example 9 A nonlinear symmetry. For the pendulum, the elements of Γ were all linear maps. Here is an example of a matrix ODE with a nonlinear symmetry. It is related to the famous Toda lattice. Let

$$X, L_0 \in \mathbb{R}^{n \times n},$$
$$\dot{X} = B(XL_0X^{-1})X,$$
$$B(L) = L_+ - L_-,$$

where L_+ (L_-) is the upper (lower) triangular part of L. This system has $h(X) = X^{-T}$ as a symmetry. The fixed set is $X = h(X) = X^{-T}$ or $XX^T = I$, i.e., $X \in O(n)$, the orthogonal group. A symmetrypreserving integrator for this system would also have O(n) as an invariant set.

4.2 Symmetries of maps

Definition 10 A map ψ has h as a symmetry if $h\psi = \psi h$, i.e.,

 $\psi = \mathcal{N}_h \psi := h^{-1} \psi h.$

A map ψ has h as a reversing symmetry if $h\psi = \psi^{-1}h$, i.e.

$$\psi = \mathcal{N}_h \mathcal{I} \psi := h^{-1} \psi^{-1} h$$

The important property of the operators \mathcal{N}_h , \mathcal{I} is how they act on compositions of maps. \mathcal{N}_h acts as an *automorphism*, i.e.

$$\mathcal{N}_h(\psi_1\psi_2) = (\mathcal{N}_h\psi_1)(\mathcal{N}_h\psi_2),$$

while \mathcal{I} acts as an *antiautomorphism*, i.e.

$$\mathcal{I}(\psi_1\psi_2) = (\psi_1\psi_2)^{-1} = \psi_2^{-1}\psi_1^{-1} = (\mathcal{I}\psi_2)(\mathcal{I}\psi_1)$$

For a map, having a (reversing) symmetry is equivalent to being in the fixed set of an (anti)automorphism. Therefore, we study how to construct maps in such fixed sets. We shall see that for antiautomorphisms this is relatively simple, while for automorphisms it is unsolved.

Thus, paradoxically, we know how to construct reversible integrators, (which is good, because reversibility brings good long-time behavior, e.g., through invariant tori), but not symmetric integrators, which looks at first sight simpler.

4.3 Covariance

Why are Runge-Kutta methods called linear methods? One explanation is that they are linearly *covariant*. Consider methods ψ which associate to each ODE f a map $\psi_{\tau}(f)$, where τ is the time step.

Definition 11 A method ψ is h-covariant if the following diagram commutes.

$$\begin{array}{cccc} \dot{x} = f(x) & \stackrel{x=h(y)}{\longrightarrow} & \dot{y} = \tilde{f}(y) \\ \downarrow & \downarrow & \downarrow \\ \widetilde{x} = \psi_{\tau}(f)(x) & \stackrel{x=h(y)}{\longrightarrow} & \widetilde{y} = \psi_{\tau}(\widetilde{f}) \end{array}$$

That is, if

$$\psi = \mathcal{K}_h \psi := h^{-1} \psi((dh \cdot f)h^{-1})h$$

where \mathcal{K}_h is an automorphism.

In words, we get the "same" integrator whether we take the ODE in variables x or y. Notice that if h is a symmetry of f, then $f = \tilde{f}$, and hence h is a symmetry of ψ . So an h-covariant method is automatically h-symmetric, even if we don't know what the symmetry is!

So, we should classify methods by their covariance group.

Example 12 Euler's method

$$\psi_{\tau}(f): x \mapsto x + \tau f(x)$$

(or, more generally, any Runge-Kutta method), is covariant under any affine map x = Ay + b.

Example 13 The exact solution $\varphi_{\tau}(f)$ is covariant under all maps x = h(y).

Example 14 The splitting method for $\dot{q} = f(p)$, $\dot{p} = g(q)$,

$$\psi_{\tau}(f): q' = q + \tau f(p), \ p' = p + \tau g(q')$$

is covariant under all maps $\tilde{q} = h_1(q)$, $\tilde{p} = h_2(p)$. It is not even covariant under linear maps which couple the q, p variables.

When composing symmetric or reversible methods, we can use the properties

- (i) The fixed sets of automorphisms form a group.
- (ii) The fixed sets of antiautomorphisms form a symmetric space.

Property (1) is immediate, while property (2) follows from the following

Lemma 15 (The Generalized Scovel Projection.) Let \mathcal{A}_{-} be an antiautomorphism with order 2, *i.e.* $\mathcal{A}_{-}^{2} = \text{id.}$ Let $\chi = \mathcal{A}_{-}\chi$, *i.e.* $\chi \in \text{fix } \mathcal{A}_{-}$. Then

$$\psi \chi \mathcal{A}_{-} \psi \in \operatorname{fix} \mathcal{A}_{-} \quad \forall \psi$$

Proof $\mathcal{A}_{-}(\psi\chi\mathcal{A}_{-}\psi) = \mathcal{A}_{-}^{2}\psi\mathcal{A}_{-}\chi\mathcal{A}_{-}\psi = \psi\chi\mathcal{A}_{-}\psi.$

Example 16 For any (anti)automorphism \mathcal{A} , we have $\mathcal{A}(\mathrm{id}) = \mathrm{id}$ and hence $\mathcal{A}(\chi^{-1}) = \mathcal{A}(\chi)^{-1}$. Therefore, $\chi \in \mathrm{fix} \mathcal{A}_{-} \Rightarrow \chi^{-1} \in \mathrm{fix} \mathcal{A}_{-}$. Taking $\psi = \mathcal{A}_{-} \psi$ gives the symmetric space property $\psi \chi^{-1} \psi \in \mathrm{fix} \mathcal{A}_{-}$.

Example 17 $\chi = id \Rightarrow \psi A_- \psi \in fix A_-$. This gives a way of constructing elements fixed under any antiautomorphism, starting from any element.

Example 18 With $\mathcal{A}_{-}\psi_{\tau} := \psi_{-\tau}^{-1}$ and $\chi = \mathrm{id}$, this builds self-adjoint methods of the form $\psi_{\tau}\psi_{-\tau}^{-1}$.

Example 19 With $\mathcal{A}_{-}\psi := h^{-1}\psi^{-1}h$, $h^{2} = \mathrm{id}$, and $\chi = \mathrm{id}$, this builds reversible methods of the form $\psi h^{-1}\psi^{-1}h$.

4.4 Building symmetric methods

This is unsolved except in two simple cases.

- (i) If the method is h-covariant and h is a symmetry, then the method is h-symmetric.
- (ii) If the symmetry group H is linear and the map ψ belongs to a linear space, then we can average over H:

$$\overline{\psi} := \frac{1}{|H|} \sum_{h \in H} \psi h$$

is *H*-symmetric.

Since preserving symmetries is difficult, we should try not to destroy symmetries in the first place, by doing non-symmetric splittings, for example.



Fig. 4.1. The discrepancy for $\psi_3 = 01101001$ applied to Example 9, where the base method ψ_0 is Euler. Notice how the symmetry error is drastically reduced every 2, then every 4, then every 8 time steps.

Hamiltonian systems

A symplectic integrator preserves an integral I if it preserves the symmetry associated with that integral, namely, the flow of $J\nabla I$.

Example 20 Angular momentum $I = q \times p$. Here $J \nabla I$ has the form $\dot{q} = a(q)$, $\dot{p} = b(p)$, whose flow has the form of Example 14. Therefore, splitting methods for $H = \frac{1}{2}p^2 + V(q)$ preserve angular momentum.

Example 21 *Quadratic integrals* are associated with linear symmetries, so are preserved by any linearly covariant symplectic integrator, such as the midpoint rule (by Noether's theorem).

4.5 Building reversible methods

Here the situation is much nicer.

Theorem 22 Let Γ be a group of automorphisms and antiautomorphisms. Let φ be fixed under the automorphisms. Then

$$\psi = \varphi \mathcal{A}_{-} \varphi$$

is fixed under \mathcal{A}_g for all $g \in \Gamma$, where \mathcal{A}_- is any antiautomorphism in Γ .

For example, if all symmetries are linear, then we can use this theorem to construct integrators having the full reversing symmetry group.

4.6 Approximately preserving symmetries

The composition used in Lemma 15 is so nice that it would be nice to use it for symmetries as well as reversing symmetries. Although it doesn't eliminate the symmetry error, it does reduce it by one power of the time step.

Theorem 23 Let \mathcal{A}_+ be an automorphism of order 2. Let ψ_{τ} be a method with $\psi = \mathcal{A}_+ \psi + \mathcal{O}(\delta)$. Let $\psi_1 := \psi \mathcal{A}_+ \psi$. Then $\psi_1 = \mathcal{A}_+ \psi_1 + \mathcal{O}(\tau \delta)$, where $\delta = \mathcal{O}(\tau)$.

Proof The proof is an illustration of backward error analysis and manipulation of flows considered as exponentials. We write the map ψ as the flow of a vector field consisting of a part S which has the

symmetry and a part M which does not:

$$\psi = \exp(\tau S + \delta M).$$

Therefore,

$$\mathcal{A}_+\psi = \exp(\tau S + \delta N)$$

for some vector field N. Now

$$\psi_1(\mathcal{A}_+\psi_1)^{-1} = (\psi \,\mathcal{A}_+\psi)(\mathcal{A}_+\psi \,\psi)^{-1}$$
$$= \psi \,\mathcal{A}_+\psi \,\psi^{-1} \,(\mathcal{A}_+\psi)^{-1}$$
$$= \exp([\tau S + \delta M, \tau S + \delta N] + \ldots)$$
$$= \exp(\tau \delta[S, N - M] + \ldots)$$

Usually, the initial symmetry error δ will be $\mathcal{O}(\tau^{p+1})$ for a method of order p, and this composition reduces it to $\mathcal{O}(\tau^{p+2})$. The idea can be applied iteratively: if

$$\psi_{n+1} = \psi_n \mathcal{A}_+ \psi_n,$$

then ψ_n has symmetry error $\mathcal{O}(\tau^n \delta)$. This gives methods of the form

$$\psi_2 = \psi \,\mathcal{A}_+ \psi \,\mathcal{A}_+ \psi \,\psi = `0110',$$

$$\psi_3 = 01101001$$

and so on, given by the initial elements of the famous 'Thue-Morse' sequence.

In the matrix example given previously, it is desired to leave the fixed set $XX^T = I$ invariant. This could be done by, e.g., the midpoint rule, but this is implicit and, given the form of the ODE, very expensive. Instead, one can use a simple explicit method for ψ , and reduce the symmetry error to any desired order using ψ_n . This leaves X orthogonal to any desired order.

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Background on symmetries

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Symmetric integration

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Volume-preserving integrators

Remember that the ODE

$$\frac{dx}{dt} = f(x)$$

is source-free (or divergence-free) if

$$\nabla \cdot f = \sum_{i=1}^{m} \frac{\partial f_i}{\partial x_i} = 0$$

for all x. Let $df = (\partial f_i / \partial x_j)$ be the derivative of f and $A = \partial \varphi_\tau / \partial x$ be the Jacobian of its flow. A evolves according to

$$\frac{dA}{dt} = dfA, \quad A(0) = Id$$

and one can show that

$$\frac{d}{dt}\det A = \operatorname{tr}(df)\det(A)$$

Consequently, if $\nabla \cdot f = \operatorname{tr}(df) = 0$, then det A = 1 for all time; the flow is volume preserving.

Volume preserving systems may be seen as one of the very few fundamental types of dynamics; their flows belong to one of the "Lie pseudogroups" of diffeomorphisms. They arise in tracking particles in incompressible fluid flow, in perturbations of Hamiltonian systems, and in discretizations of the wave equations of mathematical physics; volume preservation (and not symplecticity, for example) is the key conservation law underlying statistical mechanics. An example comparing volume- and non-volume-preserving integration is shown in Figs. 5.1-5.3.

The integrator ψ_{τ} is volume preserving (VP) if

$$\det\left(\frac{\partial\psi_{\tau,i}}{\partial x_j}\right) = 1$$

for all x. There are two general ways to construct VP integrators:

- (i) the splitting method, and
- (ii) the correction method.

5.1 Volume-preserving splitting method

Starting with the system of ODEs

$$\frac{dx_1}{dt} = f_1(x)$$

$$\vdots$$

$$\frac{dx_m}{dt} = f_m(x)$$



Fig. 5.1. Orbit of the 3D "ABC" flow computed with a second-order volume-preserving integrator. The system is $\dot{x} = A \sin z + C \sin y$, $\dot{y} = B \sin x + A \cos z$, $\dot{z} = C \sin y + B \cos x$, with parameters A = B = 1, C = 2 and initial conditions (2, 5, 0). The phase space \mathbb{T}^3 is here viewed along the z-axis, the long axis of the torus. The integration time is 750, equivalent to 240 circuits of the z-axis, and the time step is $\tau = 0.1$. The orbit lies on a torus and its regular, quasiperiodic behaviour is apparent.



Fig. 5.2. As in Fig. 5.1, but computed with the a non-volume preserving second order Runge-Kutta method with the same time step. The computed flow is not quasiperiodic and the amplitude of the motion gradually decreases (the last 3% of the orbit is shown in bold). However, this method does preserve the 16 linear symmetries of the ODE [4.2], which may explain why the results are better than in Fig. (5.3)

we substitute

$$f_m(x) = \int \frac{\partial f_m}{\partial x_m} dx_m$$
$$= -\int \sum_{i=1}^{m-1} \frac{\partial f_i}{\partial x_i} dx_m$$



Fig. 5.3. As in Fig. 5.1, but computed with MATLAB's ODE45 routine. The use of time-adaptivity has broken the spatial symmetries, with drastic consequences.

with appropriately chosen constants of integration, to get the equivalent form

$$\frac{dx_1}{dt} = f_1(x)$$

$$\vdots$$

$$\frac{dx_{m-1}}{dt} = f_{m-1}(x)$$

$$\frac{dx_m}{dt} = -\sum_{i=1}^{m-1} \int \frac{\partial f_i}{\partial x_i} dx_m$$

Now we split f, writing f as the sum of the m-1 vector fields

$$\frac{dx_i}{dt} = 0 \quad i \neq j, m$$
$$\frac{dx_j}{dt} = f_j(x)$$
$$\frac{dx_m}{dt} = -\int \frac{\partial f_j}{\partial x_j} dx_m$$

for $j = 1, \ldots, m - 1$. Note that:

- (i) Each of these m-1 vector fields is source-free.
- (ii) We have split one big problem into m-1 small problems. But we know the solution to each small problem! They each correspond to a two-dimensional Hamiltonian system

$$\frac{dx_j}{dt} = \frac{\partial H_j}{\partial x_m}$$
$$\frac{dx_m}{dt} = -\frac{\partial H_j}{\partial x_j}$$

with Hamiltonian $H_j := \int f_j(x) dx_m$, treating x_i for $i \neq j, m$ as fixed parameters. Each of these 2D problems can either be solved exactly (if possible), or approximated with any symplectic integrator ψ_j . Even though ψ_j is not symplectic in the whole space \mathbb{R}^m , it is volume-preserving.

A volume-preserving integrator for f is then given by

$$\psi = \psi_1 \circ \psi_2 \circ \ldots \psi_{m-1}.$$

Example 24 (for illustration only) The 3D Volterra system

$$\frac{dx_1}{dt} = x_1(x_2 - x_3)
\frac{dx_2}{dt} = x_2(x_3 - x_1)
\frac{dx_3}{dt} = x_3(x_1 - x_2)$$

is source-free, and splits as

$$\begin{split} & \frac{dx_1}{dt} = x_1 x_2 - x_1 x_3 \\ & \frac{dx_2}{dt} = 0 \\ & \frac{dx_3}{dt} = -x_2 x_3 + \frac{1}{2} x_3^2 \\ & \frac{dx_1}{dt} = 0 \\ & \frac{dx_2}{dt} = x_2 x_3 - x_2 x_1 \\ & \frac{dx_3}{dt} = x_1 x_3 - \frac{1}{2} x_3^2 \\ \end{split} \right\} \tilde{f}_2, \end{split}$$

where volume-preserving integrators for \tilde{f}_1 and \tilde{f}_2 are given by the implicit midpoint rule,

$$x' = x + \tau \tilde{f}_1\left(\frac{x+x'}{2}\right), \quad x'' = x' + \tau \tilde{f}_2\left(\frac{x'+x''}{2}\right).$$

Note that the x_3^2 terms were not in the original system, but on combining the two steps they cancel.

The splitting is an example of a generating function method: we construct source-free f's without any side conditions.

5.2 The volume-preserving correction method

The simplest case is the semi-implicit method

$$x'_{1} = x_{1} + \tau f_{1}(\tilde{x})$$

$$\vdots$$

$$x'_{m-1} = x_{m-1} + \tau f_{m-1}(\tilde{x})$$

$$x_{m} = \int^{x'_{m}} J(\tilde{x}) dx'_{m}$$

where

$$J := \det\left(\frac{\partial x'_i}{\partial x_j}\right)_{i,j=1,\dots,m-1}$$

and

 $\tilde{x} = (x_1, \dots, x_{m-1}, x'_m).$

For a proof of consistency and volume-preservation, see [3].

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Example 25 (for illustration only) For the 3D Volterra system

$$\frac{dx_1}{dt} = x_1(x_2 - x_3)
\frac{dx_2}{dt} = x_2(x_3 - x_1)
\frac{dx_3}{dt} = x_3(x_1 - x_2)$$

we get

$$\begin{aligned} x_1' &= x_1 + \tau x_1 (x_2 - x_3') \\ x_2' &= x_2 + \tau x_2 (x_3' - x_1) \end{aligned}$$

 and

$$J = \begin{vmatrix} \frac{\partial x'_1}{\partial x_1} & \frac{\partial x'_1}{\partial x_2} \\ \frac{\partial x'_2}{\partial x_1} & \frac{\partial x'_2}{\partial x_2} \end{vmatrix}$$
$$= \begin{vmatrix} 1 + \tau (x_2 - x'_3) & \tau x_1 \\ -\tau x_2 & 1 + \tau (x'_3 - x_1) \end{vmatrix}$$
$$= 1 + \tau (x_2 - x_1) + \tau^2 (x_2 x'_3 + x_1 x'_3 - x'^2_3)$$

and the last component of the method is $x_3 = \int^{x'_3} J dx'_3$ or

$$x_3 = x'_3 + au x'_3 (x_2 - x_1) + rac{ au^2}{2} \left(x_2 x'^2_3 + x_1 x'^2_3 - rac{2}{3} x'^3_3
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The splitting method

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6

Integrators that preserve integrals and/or Lyapunov functions

Definition 26 I(x) is a (first) integral or a conserved quantity of an ODE if

$$\frac{d}{dt}I(x(t)) = 0$$

for solutions x(t) of the ODE $\frac{dx}{dt} = f(x), x \in \mathbb{R}^m$.

By the chain rule, this requires $\sum \frac{dI}{dx_i} \frac{dx_i}{dt} = 0$ for all solutions x(t), or equivalently

$$\sum \frac{dI}{dx_i} f_i(x) = f \cdot \nabla I = 0$$

for all x.

Definition 27 V(x) is a Lyapunov function if

$$\frac{d}{dt}V(x) \le 0.$$

Equivalently,

$$f \cdot \nabla V \leq 0$$
 for all $x \in \mathbb{R}^m$.

ODEs with one or more first integrals occur frequently in physics. Many examples come from two main classes:

(i) Hamiltonian systems. For example, the pendulum

$$\frac{dx_1}{dt} = x_2$$
$$\frac{dx_2}{dt} = -\sin(x_1)$$

where x_1 is the angular position of the pendulum and x_2 its angular momentum, has the first integral

$$I(x_1, x_2) = \frac{1}{2}x_2^2 - \cos(x_1).$$

(ii) **Poisson systems.** For example, the free rigid body with moments of inertia I_1 , I_2 , and I_3 , and angular momentum π_1 , π_2 , π_3 in body-fixed coordinates,

$$\frac{d\pi_1}{dt} = \left(\frac{1}{I_2} - \frac{1}{I_3}\right) \pi_2 \pi_3$$
$$\frac{d\pi_2}{dt} = \left(\frac{1}{I_3} - \frac{1}{I_1}\right) \pi_3 \pi_1$$
$$\frac{d\pi_3}{dt} = \left(\frac{1}{I_1} - \frac{1}{I_2}\right) \pi_1 \pi_2$$

has the first integral

$$I(\pi_1, \pi_2, \pi_3) = \pi_1^2 + \pi_2^2 + \pi_3^2,$$



Fig. 6.1. An orbit of the Kepler 2-body problem with the eccentricity of the Hale–Bopp comet, computed with an integral-preserving method (left) and Runge-Kutta (right).

which is the body's total angular momentum.

6.1 Preserving a first integral

Before presenting the theory, here is a picture. Fig. 6.1 shows an orbit in the Kepler problem; it's an ellipse. This remains true if you use an integral preserving method. With a standard method such as Runge-Kutta, the orbit spirals down to the origin.

The general method we present is as follows:

- (i) For every ODE with a first integral I(x), we construct an equivalent "skew-gradient system";
- (ii) we discretize the skew-gradient system to a "skew discrete-gradient system";
- (iii) we show that the skew discrete-gradient system has the same integral I(x).

More specifically,

(i) Given the system $\frac{dx}{dt} = f(x)$ and first integral I(x) such that $\frac{d}{dt}I(x(t)) = 0$, we construct the equivalent skew gradient system

$$\frac{dx}{dt} = S\nabla I, \quad S^T = -S;$$

(ii) we discretize this to the skew discrete gradient system

$$\frac{x'-x}{\tau} = S\overline{\nabla}I(x,x')$$

where $\overline{\nabla}$ is a "discrete gradient;" (iii) we show that I(x') = I(x).

Constructing an equivalent skew gradient system

We want to solve $S\nabla I = f$ for the antisymmetric matrix S, where f and the integral I are given. Because I is an integral,

$$\frac{d}{dt}I = \frac{dx}{dt} \cdot \nabla I = f^T \nabla I = 0.$$

One solution for S is

$$S = \frac{f(\nabla I)^T - (\nabla I)f^T}{|\nabla I|^2}$$

but S is not unique. In particular, if the critical points of I (points where $\nabla I(x) = 0$) are nondegenerate, then there is an S which is as smooth as f and ∇I . Sometimes, as in Poisson systems, S is already known and does not need to be constructed.

Discretizing the skew-gradient system to a skew discrete-gradient system

A discrete gradient $\overline{\nabla}I$ is defined by the two axioms

$$I(x') - I(x) = (\overline{\nabla}I) \cdot (x' - x)$$

$$\overline{\nabla}I(x, x') = \nabla I(x) + \mathcal{O}(x' - x).$$

For any such discrete gradient we can construct the skew discrete-gradient system

$$\frac{x'-x}{\tau} = \widetilde{S}\overline{\nabla}I$$

where \widetilde{S} is any consistent antisymmetric matrix, such as $\widetilde{S}(x, x') = S((x + x')/2)$. This disarctization has the same integral L.

This discretization has the same integral I:

$$I(x') - I(x) = (\overline{\nabla}I) \cdot (x' - x)$$

= $\tau (\overline{\nabla}I)^T S(\overline{\nabla}T)$
= 0

Examples of discrete gradients

The problem is reduced to finding discrete gradients $\overline{\nabla}$ satisfying the axioms. The general solution is known. Here are two particular solutions:

(i) (Itoh and Abe)

$$\overline{\nabla}I(x,x')_i := \left(I(x^{(i)}) - I(x^{(i-1)})\right) / (x'_i - x_i),$$

where

$$x^{(i)} := (x'_1, \dots, x'_i, x_{i+1}, \dots, x_m)$$
 .

(ii) (Harten, Lax and Van Leer)

$$\overline{\nabla}I(x,x') := \int_0^1 \nabla I\left(x + \xi(x'-x)\right) \, d\xi$$

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Fig. 6.2. Evolution of an integral I(x) in a 3D system with three fourth-order methods. QT4: Integral preserving; RK4: Classical Runge-Kutta; LM4: linear multistep.

Numerical example preserving one integral

Consider the system

$$\frac{dx}{dt} = yz + x(1 + 0.1y^2 + y^3 + 0.1y^5)$$
$$\frac{dy}{dt} = -x^2 + z^2 - 0.1x^2y^2$$
$$\frac{dz}{dt} = -z - xy - y^3z$$

which has first integral

$$I = \frac{x^2}{2} + \frac{y^4}{4} + y + \frac{z^2}{2},$$

but is not Hamiltonian or Poisson. I(x) has compact level surfaces, so staying on them means that the numerical integration is stable. This system is written as a skew-gradient system as

$$\frac{d}{dt} \begin{pmatrix} x \\ y \\ z \end{pmatrix} = \begin{pmatrix} 0 & x(1+0.1y^2) & y \\ -x(1+0.1y^2) & 0 & z \\ -y & -z & 0 \end{pmatrix} \nabla I$$

and a comparison between a skew-discrete gradient integrator and two classical methods is shown in Fig. 6.2.

6.2 Preserving a Lyapunov function

An integral is a function that is preserved in time, dI/dt = 0. A Lyapunov function decreases in time, $dV/dt \leq 0$. These also arise frequently:

(i) **Gradient systems** (these arise in dynamical systems theory)

$$\frac{dx}{dt} = -\nabla V(x)$$

Here $\frac{dV}{dt} = -(\nabla V)^T \nabla V = -|\nabla V|^2 \leq 0$, so V is a Lyapunov function.

(ii) Systems with dissipation For example, the damped pendulum with friction $\alpha \ge 0$,

$$\frac{dx_1}{dt} = x_2$$
$$\frac{dx_2}{dt} = -\sin(x_1) - \alpha x_2$$

has Lyapunov function $V(x_1, x_2) = \frac{1}{2}x_2^2 - \cos(x_1)$, because

$$\frac{dV}{dt} = -\alpha x_2^2 \le 0$$

(iii) Systems with an asymptotically stable fixed point Here the construction of the Lyapunov function is a standard (although difficult) problem in dynamical systems.

These systems can be discretized very similarly to systems with an integral. Namely,

(i) Given the system $\frac{dx}{dt} = f(x)$ with Lyapunov function V, we construct the equivalent "lineargradient system"

$$\frac{dx}{dt} = L\nabla V$$

where L is negative semidefinite, i.e. $v^T L v \leq 0$ for all vectors v;

(ii) we take the linear-discrete gradient system

$$\frac{x'-x}{\tau} = L\overline{\nabla}V(x,x'),$$

(iii) we show that $V(x') \leq V(x)$.

Note that L is not necessarily symmetric. This is important, because as the dissipation tends to zero, we want L to smoothly tend to an antisymmetric matrix, to recover the integral-preserving case.

Constructing an equivalent linear-gradient system

We want to solve $L\nabla V = f$ where f and V are given, $\frac{dV}{dt} = f \cdot \nabla V \leq 0$, and L is a negative semidefinite matrix. One solution is

$$L = \frac{f(\nabla V)^T - (\nabla V)f^T + (f \cdot \nabla V)I}{|\nabla V|^2}.$$

One can check that $v^T L v = |v|^2 (f \cdot \nabla V) / |\nabla V|^2 \leq 0$, so that L is negative semi-definite, and that $L \nabla V = f$. As with skew-gradient systems, the matrix L is not unique, and special care is required near critical points of V.

$The \ linear-discrete \ gradient \ system$

By analogy with the integral-preserving case, we check that the linear-discrete gradient system has the same Lyapunov function as the original system:

$$V(x') - V(x) = (\overline{\nabla}V) \cdot (x' - x)$$
$$= \tau (\overline{\nabla}V)^T L(\overline{\nabla}V)$$
$$\leq 0.$$

Numerical example

The damped pendulum

$$\frac{dx_1}{dt} = x_2$$
$$\frac{dx_2}{dt} = -\sin(x_1) - \alpha x_2$$

with Lyapunov function $V(x_1, x_2) = \frac{1}{2}x_2^2 - \cos(x_1)$, can be written in the linear-gradient form

$$\frac{d}{dt}\begin{pmatrix}x_1\\x_2\end{pmatrix} = \begin{pmatrix}0 & 1\\-1 & -\alpha\end{pmatrix}\begin{pmatrix}\sin(x_1)\\x_2\end{pmatrix} = L\nabla V.$$

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Fig. 6.3. The damped pendulum, computed with a linear-discrete gradient method. Orbits spiral in correctly even if the dissipation rate tends to zero.

(Note that L is negative semi-definite, because the eigenvalues of its symmetric part are 0 and $-\alpha$.) Using the Itoh–Abe discrete gradient we get the discretization

$$\frac{1}{\tau} \begin{pmatrix} x_1' - x_1 \\ x_2' - x_2 \end{pmatrix} = L \overline{\nabla} V$$
$$= \begin{pmatrix} 0 & 1 \\ -1 & -\alpha \end{pmatrix} \begin{pmatrix} \frac{-\cos(x_1') + \cos(x_1)}{x_1' - x_1} \\ \frac{x_2' + x_2}{2} \end{pmatrix},$$

whose phase portrait is sketched in Fig. 6.3. The behaviour of the non-dissipative Euler's method is quite different. It increases energy near p = 0 for all time steps τ . Globally, for $\tau > 2\alpha$ orbits move out across the separatrix; and for $\alpha < \tau < 2\alpha$ there are spurious asymptotically stable periodic orbits inside the separatrix.

Extensions and generalizations

- (i) The above methods can be generalized to ODEs with any number of integrals and/or Lyapunov functions.
- (ii) There are discrete gradient methods of order 2, but higher order is desirable. For systems with an integral, this can be done by composition. For Lyapunov functions, the maps form only a semigroup, so the order cannot be increased beyond 2 by composition.
- (iii) Given a system in skew- or linear-gradient form, the matrix can be split, leading to 2D systems with the same integral or Lyapunov function.
- (iv) Satisfactory treatment of nonautonomous ODEs is still an open problem.

References

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