Time Integration (with hands-on examples using SUNDIALS)

Presented to

ATPESC 2023 Participants

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Time integrators in the HPC "landscape"

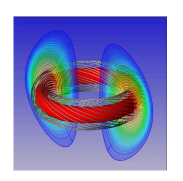
Most models of physical systems are formulated in terms of the rate of change of some variable, e.g.

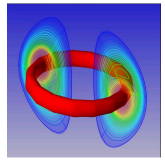
Newton's 2nd law:

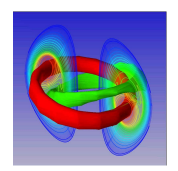
$$\mathbf{f} = m\mathbf{a} \quad \Rightarrow \quad \frac{\mathrm{d}\mathbf{v}}{\mathrm{d}t} = \frac{\mathbf{f}}{m}$$

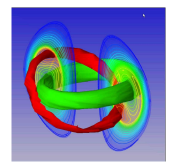
- Chemical rate equations:
$$A+B\to P \quad \Rightarrow \quad \frac{\mathrm{d}[P]}{\mathrm{d}t}=k(T)[A][B]$$

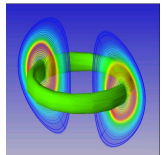
Time integrators are used to track changes in solutions as time proceeds, allowing studies of the 'evolution' of a model.

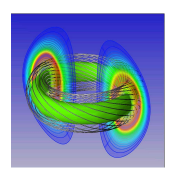












"Sawtooth" reconnection in a tokamak (NIMROD)

Time integrators in the HPC "landscape"

Unlike spatial discretization or visualization that live at the bottom/top of the software stack, respectively, time integrators typically live in the "middle." Consider some PDE systems,

$$\partial_t \rho + \nabla \cdot (\rho \mathbf{u}) = 0$$

$$\partial_t \mathbf{u} + \mathbf{u} \cdot \nabla \mathbf{u} + \frac{\nabla p}{\rho} = \mathbf{g}$$

$$\partial_t \mathbf{u} + \mathbf{u} \cdot \nabla \mathbf{u} - \nu \nabla^2 \mathbf{u} = -\nabla \left(\frac{p}{\rho_0}\right) + \mathbf{g}$$

$$\partial_t e + \mathbf{u} \cdot \nabla e + \frac{p}{\rho} \nabla \cdot \mathbf{u} = 0$$

$$\nabla \cdot \mathbf{u} = 0$$

Using a "method of lines" approach, after spatial discretization, one considers the resulting ODE/DAE system:

$$\dot{y} = f(t, y), \quad y(t_0) = y_0$$
 $F(t, y, \dot{y}) = 0, \quad y(t_0) = y_0, \quad \dot{y}(t_0) = \dot{y}_0$

- y contains all discretized solution components; f or F encodes the physics & spatial discretization

Time integrator overview

- Let $y_n \approx y(t_n)$, $t_{n+1} = t_n + \Delta t_n$. Then instead of requiring the solution at all time values, we only compute the solution at the finite set of times $\{t_n\}_{n=0...N}$.
- A "time marching" scheme computes these time-evolved solutions using a prescribed update formula:

$$y_{n+1} = \Phi(\Delta t_n, y_{n+1}, y_n, \cdots)$$

e.g., explicit Euler, $y_{n+1} = y_n + \Delta t_n f(t_n, y_n)$ and implicit Euler, $y_{n+1} = y_n + \Delta t_n f(t_{n+1}, y_{n+1})$

- Time integrator types (explicit, implicit, IMEX):
 - If Φ depends on y_{n+1} then the method is *implicit*, and requires a nonlinear solve of the form

$$F(y) \equiv y - \Phi(\Delta t_n, y, y_n, \dots) = 0$$

- If Φ does not depend on y_{n+1} then the method is *explicit*, in that the updated solution may be explicitly constructed using known data.
- Implicit-explicit (IMEX) methods arise when only some parts of Φ depend on y_{n+1} .
- Multirate methods use different time step sizes $\Delta t_n \gg \delta t_n$ to evolve separate problem components.

Time integrator overview (continued)

- Time integration methods have multiple mechanisms for achieving increased accuracy:
 - "One-step" methods use multiple internal stages per step [Runge-Kutta, Rosenbrock].
 - More work per-step; amenable to problems with spatial adaptivity & hyperbolic effects.
 - "Multistep" methods retain a longer history of previous solutions [Adams-Bashforth, BDF].
 - Less work per-step; amenable to problems with strong reaction and diffusion effects.
- Linear stability: a method is numerically stable if for a desired Δt_n , roundoff error stays "controlled" throughout the simulation (vs growing out of control). For a brief refresher, see here.
 - "A-stable": linearly stable no matter the Δt_n this is only possible with implicit methods¹.
 - Otherwise, the method has a maximum stable step size Δt_n for any given problem (in PDEs, this is frequently given by the *CFL condition*, wherein $\Delta t_n \propto \Delta x$ or $\Delta t_n \propto \Delta x^2$).
 - Stability ≠ accuracy just because a solution does not blow up, it is not necessarily accurate.

¹So-called "exponential" methods are explicit and may be A-stable, but require *significantly* more work per-step than traditional explicit methods. I know of no open-source HPC library that provides these.

Choosing between explicit and implicit methods

Explicit Methods	Implicit Methods
+ easy to conceptualize	+ less/nonexistent stability limits
+ easy to code	+ steps over fastest dynamics
+ no algebraic solvers required	 requires algebraic solvers
 stability limits on step sizes 	 solvers generally couple all solution unknowns
 tracks fastest dynamics 	 increased code complexity

- IMEX: a bit of both one chooses the splitting to balance 'cheaper' algebraic solvers and stability.
- "Stiffness" helps us choose: "The stepsize needed to maintain stability of the forward Euler method is much smaller than that required to represent the solution accurately." (Ascher and Petzold, 1998)
 - Depends on Jacobian eigenvalues, system dimension, accuracy requirements, length of simulation.
 - For stability, stiff problems generally require implicit or IMEX methods, with robust implicit solvers.
- DAEs nearly always require implicit methods to maintain stability due to the algebraic constraint.
- Multirate methods may be preferable if the 'slow' operator is much more costly than the 'fast'.

Adaptive time-step selection

- Stability alone should never dictate the time steps used in an application.
- Given a maximum stable step size, adaptive methods select Δt_n to obtain a desired solution accuracy:
 - At each internal step, computes both the solution and an estimate of the error introduced in that step.
 - If that *local truncation error* is small enough the step is accepted; otherwise a new step size is chosen that should provide sufficient accuracy, and the step is recomputed.
 - Advanced "error controllers" adapt these step sizes to meet a variety of objectives:
 - minimize failed steps
 - maximize step sizes
 - maintain smooth transitions in the step sizes as integration proceeds
- Temporal adaptivity can lead to much more efficient (and accurate) results.

"Solving" Initial-Value Problems with SUNDIALS

SUNDIALS' integrators consider initial-value problems of a variety of types:

- Standard IVP [CVODE]: $\dot{y}(t) = f(t, y(t)), \quad y(t_0) = y_0$

- Linearly-implicit, split [ARKODE]: $M \dot{y}(t) = f_1(t, y(t)) + f_2(t, y(t)), \quad y(t_0) = y_0$

- Multirate [ARKODE/MRIStep]: $\dot{y} = f^F(t,y) + f^S(t,y), \quad y(t_0) = y_0$

- Differential-algebraic form [IDA]: $F(t, y(t), \dot{y}(t)) = 0, \quad y(t_0) = y_0, \quad \dot{y}(t_0) = \dot{y}_0$

By "solve" we adapt time steps (and/or method order) to meet user-specified tolerances:

$$\left[\frac{1}{N}\sum_{k=1}^{N} \left(\frac{\operatorname{error}_{k}}{\operatorname{rtol}|y_{k}| + \operatorname{atol}_{k}}\right)^{2}\right]^{1/2} < 1$$

- $-\operatorname{error} \in \mathbb{R}^N$ is the estimated temporal error in the time step
- $-y \in \mathbb{R}^N$ is the previous time-step solution
- $-\operatorname{rtol}\in\mathbb{R}$ encodes the desired relative solution accuracy (number of significant digits)
- $-\operatorname{atol}\in\mathbb{R}^N$ is the 'noise' level for any solution component (protects against $y_k=0$)

Other DOE Time Integration Packages

PETSc's TS module provides a unified interface for implicit, explicit, & IMEX ODEs and DAEs:

$$F(t, y, \dot{y}) = G(t, y), \quad y(t_0) = y_0$$

- $-F(t,y,\dot{y})$ stiff portion; G(t,y) nonstiff portion
- <u>Trilinos</u> includes both an older Rythmos module for ODEs and DAEs:

$$F(t, y(t), \dot{y}(t)) = 0, \quad y(t_0) = y_0, \quad \dot{y}(t_0) = \dot{y}_0$$

As well as a newer *Tempus* module for ODEs:

$$M(t,y) \dot{y}(t) = G(t,y) + F(t,y),$$
 $y(t_0) = y_0$
 $M \ddot{y}(t) + C \dot{y}(t) + K y(t) + F(t) = 0,$ $y(t_0) = y_0, \ \dot{y}(t_0) = \dot{y}_0$

- Top: G(t, y) stiff, F(t, y) nonstiff; Bottom: Newmark integrators for second-order ODEs
- All perform temporal adaptivity, and provide a rich set of algebraic solvers for implicit time integration methods.

Implicit methods require a nonlinear solver for F(x) = 0

The PETSc team is presenting on nonlinear solvers in sessions parallel to this one, so I'll only give a high-level idea, leaving details for them.

Nonlinear solvers must be iterative, since few nonlinear equations admit analytical solutions.

The two largest classes of nonlinear solvers are fixed-point vs Newton-based.

- FP typically use only *F* and converge linearly, but may have a large domain of convergence.
- Newton uses both F and the Jacobian $J(x) \equiv \frac{\partial F(x)}{\partial x}$ (or an approx.):
 - Each iteration requires a linear solve with the matrix J (see linear solvers talks this AM).
 - Typically converge quadratically (or superlinearly, depending on how well J is solved).
 - For most problems, Newton is algorithmically scalable as the mesh is refined, the number of iterations remains fixed, so scalability hinges on the linear system solver.

Why use a solver library (instead of "rolling your own")

- Many applications (particularly early on) implement their own numerical methods to remove software dependencies. While functional, the methods may be overly simplistic (e.g., straight from "Numerical" Recipes") or buggy, may not leverage modern hardware, and/or omit advanced "expert" features.
- Solver libraries, on the other hand, are typically bug-free, heavily tested, and admit numerous benefits:
 - Time adaptivity provide approximate solutions with requested accuracy and minimal work.
 - Seamless integration with scalable algebraic solver libraries for implicit and IMEX problems.
 - Native support for cutting-edge GPU hardware via CUDA, HIP, oneAPI, Kokkos, RAJA, ...
 - Advanced options for later use: temporal root-finding, sensitivity analysis, solution constraints, ...
- For more information:
 - SUNDIALS: https://computing.llnl.gov/projects/sundials
 - PETSc: https://www.mcs.anl.gov/petsc/
 - Trilinos: https://trilinos.github.io/

Hands-on lessons

Switch over to web-based hands-on lesson instructions – webpage

Agenda:

- 1. Explicit time integration (HandsOn1.exe)
- 2. Implicit / IMEX time integration (HandsOn2.exe)
- 3. Preconditioning (HandsOn3.exe)

Take Away Messages

- SUNDIALS, PETSc, and Trilinos provide a wide variety of high quality, scalable ODE/DAE integrators and nonlinear solvers.
- PDEs can be converted to ODEs/DAEs via spatial semi-discretization, and then integrated using ODE/DAE libraries.
- Stiffness is an important characteristic of ODEs, and helps dictate which methods are appropriate for any given problem.
- Adaptive time-stepping provides an inexpensive means to combine algorithmic efficiency and solution quality.
- Scalability of implicit and IMEX methods hinges on selection of robust and scalable algebraic solvers; while Newton methods can handle nonlinearities, robustness and scalability of the inner linear solver is critical (and often problem-dependent).

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EXASCALE COMPUTING PROJECT

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Linear Stability – A brief refresher

A fundamental question for any time integration method is how well it handles errors due to floating-point roundoff. To this end, we consider the simple "Dahlquist" test problem:

$$y'(t) = \lambda y(t), \quad y(0) = 1$$

- Here, y corresponds to the normalized floating-point error, and λ to the largest eigenvalue of the Jacobian of a prototypical ODE right-hand side function (assumed to satisfy $\Re(\lambda) < 0$).
- The true solution to this problem is just $y(t) = e^{\lambda t}$, which decays to zero as $t \to \infty$, indicating that roundoff errors should decay as the simulation proceeds.
- The numerical method, on the other hand, computes approximate solutions

$$y_{n+1} = \Phi(\Delta t_n, y_{n+1}, y_n, \ldots)$$

that may (or may not) similarly satisfy the similar requirement that $y_n \to 0 \text{ as } n \to \infty$.

• Generally, this decay in numerical roundoff error will only occur for specific values of $\Delta t \lambda = z \in \mathbb{C}$. We therefore define the *stability region* for a method as $S = \{z \in \mathbb{C} : \Phi_z(y_n) \to 0 \text{ as } n \to \infty\}$

Linear Stability Example – Explicit Euler

Consider the explicit Euler method: $y_{n+1} = y_n + \Delta t f(t_n, y_n)$

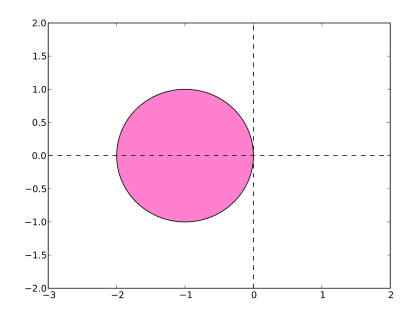
For the Dahlquist test problem, this becomes

$$y_{n+1} = y_n + \Delta t \lambda y_n = (1 + \Delta t \lambda) y_n = (1 + \Delta t \lambda)^2 y_{n-1} = \dots = (1 + \Delta t \lambda)^{n+1} y_0 = (1 + \Delta t \lambda)^{n+1}$$

which only decays to zero if $|1+\Delta t\lambda|<1$.

Hence the explicit Euler linear stability region is

$$S = \{ z \in \mathbb{C} : |1 + z| < 1 \}$$



From https://en.wikipedia.org/wiki/Euler_method

Linear Stability Example – Implicit Euler

Consider the implicit Euler method: $y_{n+1} = y_n + \Delta t f(t_{n+1}, y_{n+1})$

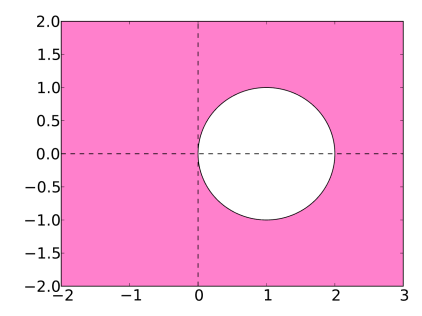
For the Dahlquist test problem, this becomes

$$y_{n+1} = y_n + \Delta t \lambda y_{n+1} \quad \Leftrightarrow \quad y_{n+1} = (1 - \Delta t \lambda)^{-1} y_n = \dots = (1 - \Delta t \lambda)^{-(n+1)}$$

which only decays to zero if $|1 - \Delta t \lambda| > 1$.

Hence the explicit Euler linear stability region is

$$S = \{ z \in \mathbb{C} : |1 - z| > 1 \}$$



From https://en.wikipedia.org/wiki/Backward Euler method