

# Scientific Computing: An Introductory Survey

## Chapter 9 – Initial Value Problems for Ordinary Differential Equations

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# Outline

- 1 Ordinary Differential Equations
- 2 Numerical Solution of ODEs
- 3 Additional Numerical Methods



# Differential Equations

- Differential equations involve derivatives of unknown solution function
- *Ordinary differential equation* (ODE): all derivatives are with respect to single independent variable, often representing time
- Solution of differential equation is *function* in infinite-dimensional space of functions
- Numerical solution of differential equations is based on finite-dimensional approximation
- Differential equation is replaced by algebraic equation whose solution approximates that of given differential equation



# Order of ODE

- *Order* of ODE is determined by highest-order derivative of solution function appearing in ODE
- ODE with higher-order derivatives can be transformed into equivalent first-order system
- We will discuss numerical solution methods only for first-order ODEs
- Most ODE software is designed to solve only first-order equations



# Higher-Order ODEs, continued

- For  $k$ -th order ODE

$$y^{(k)}(t) = f(t, y, y', \dots, y^{(k-1)})$$

define  $k$  new unknown functions

$$u_1(t) = y(t), u_2(t) = y'(t), \dots, u_k(t) = y^{(k-1)}(t)$$

- Then original ODE is equivalent to first-order system

$$\begin{bmatrix} u_1'(t) \\ u_2'(t) \\ \vdots \\ u_{k-1}'(t) \\ u_k'(t) \end{bmatrix} = \begin{bmatrix} u_2(t) \\ u_3(t) \\ \vdots \\ u_k(t) \\ f(t, u_1, u_2, \dots, u_k) \end{bmatrix}$$



## Example: Newton's Second Law

- Newton's Second Law of Motion,  $F = ma$ , is second-order ODE, since acceleration  $a$  is second derivative of position coordinate, which we denote by  $y$
- Thus, ODE has form

$$y'' = F/m$$

where  $F$  and  $m$  are force and mass, respectively

- Defining  $u_1 = y$  and  $u_2 = y'$  yields equivalent system of two first-order ODEs

$$\begin{bmatrix} u_1' \\ u_2' \end{bmatrix} = \begin{bmatrix} u_2 \\ F/m \end{bmatrix}$$



## Example, continued

- We can now use methods for first-order equations to solve this system
- First component of solution  $u_1$  is solution  $y$  of original second-order equation
- Second component of solution  $u_2$  is velocity  $y'$



# Ordinary Differential Equations

- General first-order system of ODEs has form

$$\mathbf{y}'(t) = \mathbf{f}(t, \mathbf{y})$$

where  $\mathbf{y}: \mathbb{R} \rightarrow \mathbb{R}^n$ ,  $\mathbf{f}: \mathbb{R}^{n+1} \rightarrow \mathbb{R}^n$ , and  $\mathbf{y}' = d\mathbf{y}/dt$  denotes derivative with respect to  $t$ ,

$$\begin{bmatrix} y_1'(t) \\ y_2'(t) \\ \vdots \\ y_n'(t) \end{bmatrix} = \begin{bmatrix} dy_1(t)/dt \\ dy_2(t)/dt \\ \vdots \\ dy_n(t)/dt \end{bmatrix}$$

- Function  $\mathbf{f}$  is given and we wish to determine unknown function  $\mathbf{y}$  satisfying ODE
- For simplicity, we will often consider special case of single scalar ODE,  $n = 1$





# Initial Value Problems

- By itself, ODE  $y' = f(t, y)$  does not determine unique solution function
- This is because ODE merely specifies *slope*  $y'(t)$  of solution function at each point, but not actual value  $y(t)$  at any point
- Infinite family of functions satisfies ODE, in general, provided  $f$  is sufficiently smooth
- To single out particular solution, value  $y_0$  of solution function must be specified at some point  $t_0$



## Initial Value Problems, continued

- Thus, part of given problem data is requirement that  $\mathbf{y}(t_0) = \mathbf{y}_0$ , which determines unique solution to ODE
- Because of interpretation of independent variable  $t$  as time, think of  $t_0$  as initial time and  $\mathbf{y}_0$  as initial value
- Hence, this is termed *initial value problem*, or *IVP*
- ODE governs evolution of system in time from its initial state  $\mathbf{y}_0$  at time  $t_0$  onward, and we seek function  $\mathbf{y}(t)$  that describes state of system as function of time



# Example: Initial Value Problem

- Consider scalar ODE

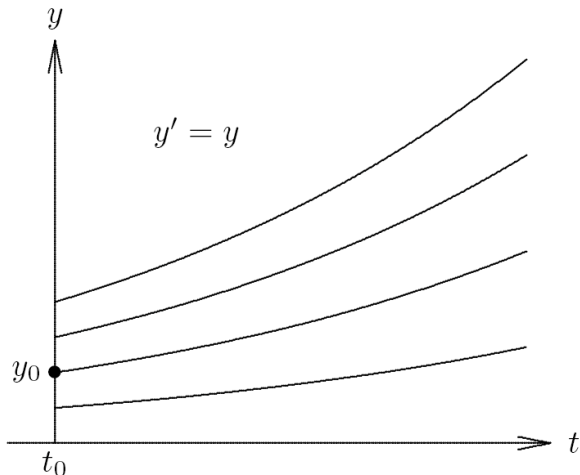
$$y' = y$$

- Family of solutions is given by  $y(t) = ce^t$ , where  $c$  is any real constant
- Imposing initial condition  $y(t_0) = y_0$  singles out unique particular solution
- For this example, if  $t_0 = 0$ , then  $c = y_0$ , which means that solution is  $y(t) = y_0e^t$



# Example: Initial Value Problem

Family of solutions for ODE  $y' = y$



# Stability of Solutions

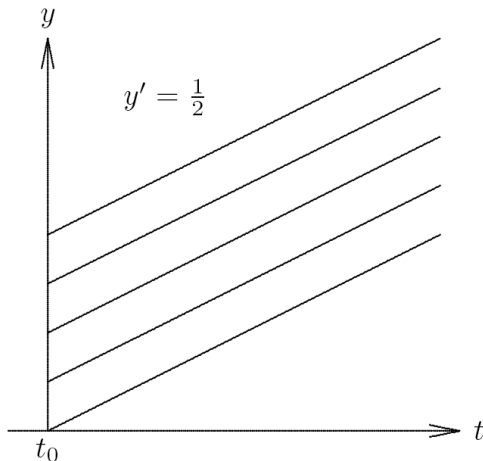
Solution of ODE is

- *Stable* if solutions resulting from perturbations of initial value remain close to original solution
- *Asymptotically stable* if solutions resulting from perturbations converge back to original solution
- *Unstable* if solutions resulting from perturbations diverge away from original solution without bound



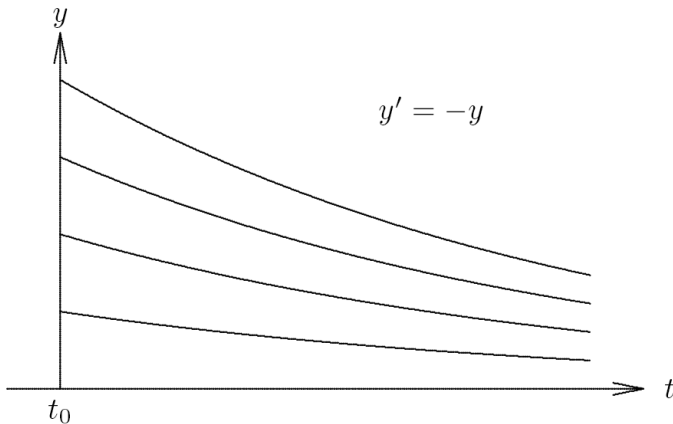
# Example: Stable Solutions

Family of solutions for ODE  $y' = \frac{1}{2}$



# Example: Asymptotically Stable Solutions

Family of solutions for ODE  $y' = -y$



## Example: Stability of Solutions

- Consider scalar ODE  $y' = \lambda y$ , where  $\lambda$  is constant.
- Solution is given by  $y(t) = y_0 e^{\lambda t}$ , where  $t_0 = 0$  is initial time and  $y(0) = y_0$  is initial value
- For real  $\lambda$ 
  - $\lambda > 0$ : all nonzero solutions grow exponentially, so every solution is unstable
  - $\lambda < 0$ : all nonzero solutions decay exponentially, so every solution is not only stable, but asymptotically stable
- For complex  $\lambda$ 
  - $\text{Re}(\lambda) > 0$ : unstable
  - $\text{Re}(\lambda) < 0$ : asymptotically stable
  - $\text{Re}(\lambda) = 0$ : stable but not asymptotically stable





# Example: Linear System of ODEs

- Linear, homogeneous system of ODEs with constant coefficients has form

$$\mathbf{y}' = \mathbf{A}\mathbf{y}$$

where  $\mathbf{A}$  is  $n \times n$  matrix, and initial condition is  $\mathbf{y}(0) = \mathbf{y}_0$

- Suppose  $\mathbf{A}$  is diagonalizable, with eigenvalues  $\lambda_i$  and corresponding eigenvectors  $\mathbf{v}_i$ ,  $i = 1, \dots, n$
- Express  $\mathbf{y}_0$  as linear combination  $\mathbf{y}_0 = \sum_{i=1}^n \alpha_i \mathbf{v}_i$
- Then

$$\mathbf{y}(t) = \sum_{i=1}^n \alpha_i \mathbf{v}_i e^{\lambda_i t}$$

is solution to ODE satisfying initial condition  $\mathbf{y}(0) = \mathbf{y}_0$



## Example, continued

- Eigenvalues of  $A$  with positive real parts yield exponentially growing solution components
- Eigenvalues with negative real parts yield exponentially decaying solution components
- Eigenvalues with zero real parts (i.e., pure imaginary) yield oscillatory solution components
- Solutions stable if  $\operatorname{Re}(\lambda_i) \leq 0$  for every eigenvalue, and asymptotically stable if  $\operatorname{Re}(\lambda_i) < 0$  for every eigenvalue, but unstable if  $\operatorname{Re}(\lambda_i) > 0$  for any eigenvalue



# Stability of Solutions, continued

- For general nonlinear system of ODEs  $\mathbf{y}' = \mathbf{f}(t, \mathbf{y})$ , determining stability of solutions is more complicated
- ODE can be linearized locally about solution  $\mathbf{y}(t)$  by truncated Taylor series, yielding linear ODE

$$\mathbf{z}' = \mathbf{J}_f(t, \mathbf{y}(t)) \mathbf{z}$$

where  $\mathbf{J}_f$  is Jacobian matrix of  $\mathbf{f}$  with respect to  $\mathbf{y}$

- Eigenvalues of  $\mathbf{J}_f$  determine stability locally, but conclusions drawn may not be valid globally



# Numerical Solution of ODEs

- Analytical solution of ODE is closed-form formula that can be evaluated at any point  $t$
- Numerical solution of ODE is table of approximate values of solution function at discrete set of points
- Numerical solution is generated by simulating behavior of system governed by ODE
- Starting at  $t_0$  with given initial value  $y_0$ , we track trajectory dictated by ODE
- Evaluating  $f(t_0, y_0)$  tells us slope of trajectory at that point
- We use this information to predict value  $y_1$  of solution at future time  $t_1 = t_0 + h$  for some suitably chosen time increment  $h$



# Numerical Solution of ODEs, continued

- Approximate solution values are generated step by step in increments moving across interval in which solution is sought
- In stepping from one discrete point to next, we incur some error, which means that next approximate solution value lies on *different* solution from one we started on
- Stability or instability of solutions determines, in part, whether such errors are magnified or diminished with time



# Euler's Method

- For general system of ODEs  $\mathbf{y}' = \mathbf{f}(t, \mathbf{y})$ , consider Taylor series

$$\begin{aligned} \mathbf{y}(t+h) &= \mathbf{y}(t) + h\mathbf{y}'(t) + \frac{h^2}{2}\mathbf{y}''(t) + \dots \\ &= \mathbf{y}(t) + h\mathbf{f}(t, \mathbf{y}(t)) + \frac{h^2}{2}\mathbf{y}''(t) + \dots \end{aligned}$$

- Euler's method* results from dropping terms of second and higher order to obtain approximate solution value

$$\mathbf{y}_{k+1} = \mathbf{y}_k + h_k \mathbf{f}(t_k, \mathbf{y}_k)$$

- Euler's method advances solution by extrapolating along straight line whose slope is given by  $\mathbf{f}(t_k, \mathbf{y}_k)$
- Euler's method is *single-step* method because it depends on information at only one point in time to advance to next point



# Example: Euler's Method

- Applying Euler's method to ODE  $y' = y$  with step size  $h$ , we advance solution from time  $t_0 = 0$  to time  $t_1 = t_0 + h$

$$y_1 = y_0 + hy'_0 = y_0 + hy_0 = (1 + h)y_0$$

- Value for solution we obtain at  $t_1$  is not exact,  $y_1 \neq y(t_1)$
- For example, if  $t_0 = 0$ ,  $y_0 = 1$ , and  $h = 0.5$ , then  $y_1 = 1.5$ , whereas exact solution for this initial value is  $y(0.5) = \exp(0.5) \approx 1.649$
- Thus,  $y_1$  lies on different solution from one we started on



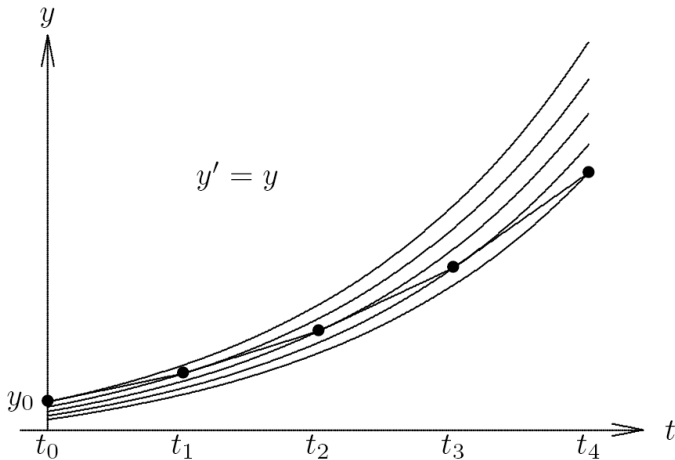
## Example, continued

- To continue numerical solution process, we take another step from  $t_1$  to  $t_2 = t_1 + h = 1.0$ , obtaining
$$y_2 = y_1 + hy_1 = 1.5 + (0.5)(1.5) = 2.25$$
- Now  $y_2$  differs not only from true solution of original problem at  $t = 1$ ,  $y(1) = \exp(1) \approx 2.718$ , but it also differs from solution through previous point  $(t_1, y_1)$ , which has approximate value 2.473 at  $t = 1$
- Thus, we have moved to still another solution for this ODE





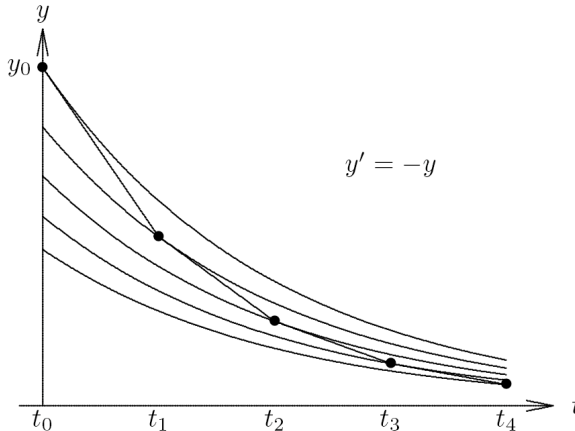
## Example, continued



For unstable solutions, errors in numerical solution grow with time



## Example, continued



For stable solutions, errors in numerical solution may diminish with time

< interactive example >



# Errors in Numerical Solution of ODEs

- Numerical methods for solving ODEs incur two distinct types of error
  - *Rounding error*, which is due to finite precision of floating-point arithmetic
  - *Truncation error* (*discretization error*), which is due to approximation method used and would remain even if all arithmetic were exact
- In practice, truncation error is dominant factor determining accuracy of numerical solutions of ODEs, so we will henceforth ignore rounding error



# Global Error and Local Error

Truncation error at any point  $t_k$  can be broken down into

- **Global error**: difference between computed solution and true solution  $\mathbf{y}(t)$  passing through initial point  $(t_0, \mathbf{y}_0)$

$$\mathbf{e}_k = \mathbf{y}_k - \mathbf{y}(t_k)$$

- **Local error**: error made in one step of numerical method

$$\mathbf{l}_k = \mathbf{y}_k - \mathbf{u}_{k-1}(t_k)$$

where  $\mathbf{u}_{k-1}(t)$  is true solution passing through previous point  $(t_{k-1}, \mathbf{y}_{k-1})$

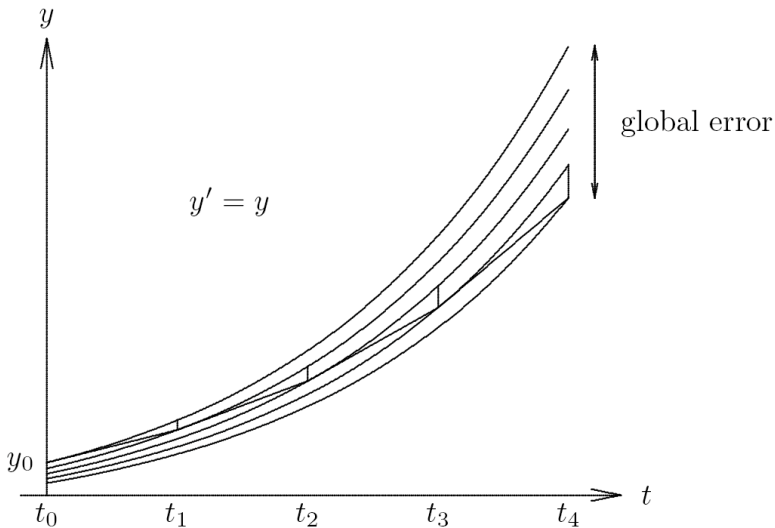


## Global Error and Local Error, continued

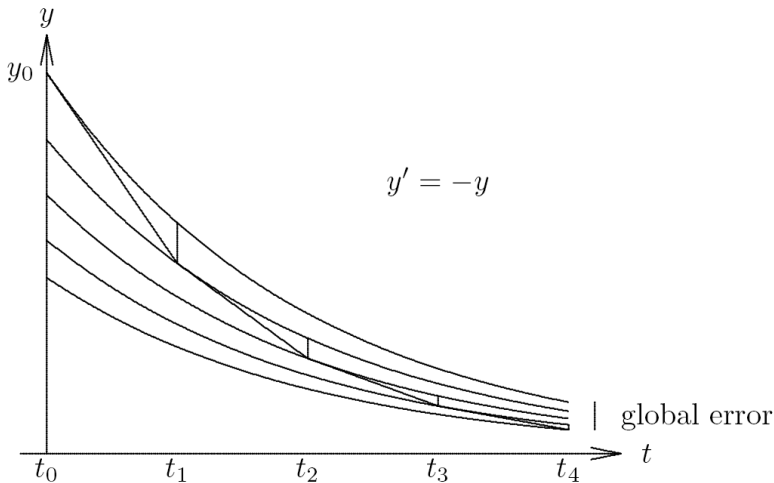
- Global error is not necessarily sum of local errors
- Global error is generally greater than sum of local errors if solutions are unstable, but may be less than sum if solutions are stable
- Having small global error is what we want, but we can control only local error directly



# Global Error and Local Error, continued



# Global and Local Error, continued



# Order of Accuracy

- *Order of accuracy* of numerical method is  $p$  if

$$\ell_k = \mathcal{O}(h_k^{p+1})$$

- Then local error per unit step,  $\ell_k/h_k = \mathcal{O}(h_k^p)$
- Under reasonable conditions,  $e_k = \mathcal{O}(h^p)$ , where  $h$  is average step size

< interactive example >





# Stability

- Numerical method is *stable* if small perturbations do not cause resulting numerical solutions to diverge from each other without bound
- Such divergence of numerical solutions could be caused by instability of solution to ODE, but can also be due to numerical method itself, even when solutions to ODE are stable



# Determining Stability and Accuracy

- Simple approach to determining stability and accuracy of numerical method is to apply it to scalar ODE  $y' = \lambda y$ , where  $\lambda$  is (possibly complex) constant
- Exact solution is given by  $y(t) = y_0 e^{\lambda t}$ , where  $y(0) = y_0$  is initial condition
- Determine stability of numerical method by characterizing growth of numerical solution
- Determine accuracy of numerical method by comparing exact and numerical solutions



## Example: Euler's Method

- Applying Euler's method to  $y' = \lambda y$  using fixed step size  $h$ ,

$$y_{k+1} = y_k + h\lambda y_k = (1 + h\lambda)y_k$$

which means that

$$y_k = (1 + h\lambda)^k y_0$$

- If  $\text{Re}(\lambda) < 0$ , exact solution decays to zero as  $t$  increases, as does computed solution if

$$|1 + h\lambda| < 1$$

which holds if  $h\lambda$  lies inside circle in complex plane of radius 1 centered at  $-1$

< interactive example >



# Euler's Method, continued

- If  $\lambda$  is real, then  $h\lambda$  must lie in interval  $(-2, 0)$ , so for  $\lambda < 0$ , we must have

$$h \leq -\frac{2}{\lambda}$$

for Euler's method to be stable

- *Growth factor*  $1 + h\lambda$  agrees with series expansion

$$e^{h\lambda} = 1 + h\lambda + \frac{(h\lambda)^2}{2} + \frac{(h\lambda)^3}{6} + \dots$$

through terms of first order in  $h$ , so Euler's method is first-order accurate



# Euler's Method, continued

- For general system of ODEs  $\mathbf{y}' = \mathbf{f}(t, \mathbf{y})$ , consider Taylor series

$$\begin{aligned}\mathbf{y}(t+h) &= \mathbf{y}(t) + h\mathbf{y}'(t) + \mathcal{O}(h^2) \\ &= \mathbf{y}(t) + h\mathbf{f}(t, \mathbf{y}(t)) + \mathcal{O}(h^2)\end{aligned}$$

- If we take  $t = t_k$  and  $h = h_k$ , we obtain

$$\mathbf{y}(t_{k+1}) = \mathbf{y}(t_k) + h_k \mathbf{f}(t_k, \mathbf{y}(t_k)) + \mathcal{O}(h_k^2)$$

- Subtracting this from Euler's method,

$$\begin{aligned}\mathbf{e}_{k+1} &= \mathbf{y}_{k+1} - \mathbf{y}(t_{k+1}) \\ &= [\mathbf{y}_k - \mathbf{y}(t_k)] + h_k [\mathbf{f}(t_k, \mathbf{y}_k) - \mathbf{f}(t_k, \mathbf{y}(t_k))] - \mathcal{O}(h_k^2)\end{aligned}$$



## Euler's Method, continued

- If there were no prior errors, then we would have  $\mathbf{y}_k = \mathbf{y}(t_k)$ , and differences in brackets on right side would be zero, leaving only  $\mathcal{O}(h_k^2)$  term, which is local error
- This means that Euler's method is first-order accurate



# Euler's Method, continued

- From previous derivation, global error is sum of local error and *propagated* error
- From Mean Value Theorem,

$$\mathbf{f}(t_k, \mathbf{y}_k) - \mathbf{f}(t_k, \mathbf{y}(t_k)) = \mathbf{J}_f(t_k, \boldsymbol{\xi})(\mathbf{y}_k - \mathbf{y}(t_k))$$

for some (unknown) value  $\boldsymbol{\xi}$ , where  $\mathbf{J}_f$  is Jacobian matrix of  $\mathbf{f}$  with respect to  $\mathbf{y}$

- So we can express global error at step  $k + 1$  as

$$\mathbf{e}_{k+1} = (\mathbf{I} + h_k \mathbf{J}_f) \mathbf{e}_k + \boldsymbol{\ell}_{k+1}$$

- Thus, global error is multiplied at each step by *growth factor*  $\mathbf{I} + h_k \mathbf{J}_f$
- Errors do not grow if spectral radius  $\rho(\mathbf{I} + h_k \mathbf{J}_f) \leq 1$ , which holds if all eigenvalues of  $h_k \mathbf{J}_f$  lie inside circle in complex plane of radius 1 centered at  $-1$



# Stability of Numerical Methods for ODEs

In general, growth factor depends on

- Numerical method, which determines form of growth factor
- Step size  $h$
- Jacobian  $\mathbf{J}_f$ , which is determined by particular ODE





# Step Size Selection

- In choosing step size for advancing numerical solution of ODE, we want to take large steps to reduce computational cost, but must also take into account both stability and accuracy
- To yield meaningful solution, step size must obey any stability restrictions
- In addition, local error estimate is needed to ensure that desired accuracy is achieved
- With Euler's method, for example, local error is approximately  $(h_k^2/2)\mathbf{y}''$ , so choose step size to satisfy

$$h_k \leq \sqrt{2 \text{ tol} / \|\mathbf{y}''\|}$$



# Step Size Selection, continued

- We do not know value of  $y''$ , but we can estimate it by difference quotient

$$y'' \approx \frac{y'_k - y'_{k-1}}{t_k - t_{k-1}}$$

- Other methods of obtaining error estimates are based on difference between results obtained using methods of different orders or different step sizes



# Implicit Methods

- Euler's method is *explicit* in that it uses only information at time  $t_k$  to advance solution to time  $t_{k+1}$
- This may seem desirable, but Euler's method has rather limited stability region
- Larger stability region can be obtained by using information at time  $t_{k+1}$ , which makes method *implicit*
- Simplest example is *backward Euler method*

$$\mathbf{y}_{k+1} = \mathbf{y}_k + h_k \mathbf{f}(t_{k+1}, \mathbf{y}_{k+1})$$

- Method is implicit because we must evaluate  $\mathbf{f}$  with argument  $\mathbf{y}_{k+1}$  before we know its value



## Implicit Methods, continued

- This means that we must solve algebraic equation to determine  $\mathbf{y}_{k+1}$
- Typically, we use iterative method such as Newton's method or fixed-point iteration to solve for  $\mathbf{y}_{k+1}$
- Good starting guess for iteration can be obtained from explicit method, such as Euler's method, or from solution at previous time step

< interactive example >



## Example: Backward Euler Method

- Consider nonlinear scalar ODE  $y' = -y^3$  with initial condition  $y(0) = 1$
- Using backward Euler method with step size  $h = 0.5$ , we obtain implicit equation

$$y_1 = y_0 + hf(t_1, y_1) = 1 - 0.5y_1^3$$

for solution value at next step

- This nonlinear equation for  $y_1$  could be solved by fixed-point iteration or Newton's method
- To obtain starting guess for  $y_1$ , we could use previous solution value,  $y_0 = 1$ , or we could use explicit method, such as Euler's method, which gives  $y_1 = y_0 - 0.5y_0^3 = 0.5$
- Iterations eventually converge to final value  $y_1 \approx 0.7709$



# Implicit Methods, continued

- Given extra trouble and computation in using implicit method, one might wonder why we bother
- Answer is that implicit methods generally have significantly larger stability region than comparable explicit methods



# Backward Euler Method

- To determine stability of backward Euler, we apply it to scalar ODE  $y' = \lambda y$ , obtaining

$$y_{k+1} = y_k + h\lambda y_{k+1}$$

$$(1 - h\lambda)y_{k+1} = y_k$$

$$y_k = \left( \frac{1}{1 - h\lambda} \right)^k y_0$$

- Thus, for backward Euler to be stable we must have

$$\left| \frac{1}{1 - h\lambda} \right| \leq 1$$

which holds for *any*  $h > 0$  when  $\text{Re}(\lambda) < 0$

- So stability region for backward Euler method includes entire left half of complex plane, or interval  $(-\infty, 0)$  if  $\lambda$  is real



# Backward Euler Method, continued

- Growth factor

$$\frac{1}{1 - h\lambda} = 1 + h\lambda + (h\lambda)^2 + \dots$$

agrees with expansion for  $e^{\lambda h}$  through terms of order  $h$ , so backward Euler method is first-order accurate

- Growth factor of backward Euler method for general system of ODEs  $\mathbf{y}' = \mathbf{f}(t, \mathbf{y})$  is  $(\mathbf{I} - h\mathbf{J}_f)^{-1}$ , whose spectral radius is less than 1 provided all eigenvalues of  $h\mathbf{J}_f$  lie outside circle in complex plane of radius 1 centered at 1
- Thus, stability region of backward Euler for general system of ODEs is entire left half of complex plane





# Unconditionally Stable Methods

- Thus, for computing stable solution backward Euler is stable for any positive step size, which means that it is *unconditionally* stable
- Great virtue of unconditionally stable method is that desired accuracy is only constraint on choice of step size
- Thus, we may be able to take much larger steps than for explicit method of comparable order and attain much higher overall efficiency despite requiring more computation per step
- Although backward Euler method is unconditionally stable, its accuracy is only of first order, which severely limits its usefulness



## Trapezoid Method

- Higher-order accuracy can be achieved by averaging Euler and backward Euler methods to obtain implicit *trapezoid method*

$$\mathbf{y}_{k+1} = \mathbf{y}_k + h_k (\mathbf{f}(t_k, \mathbf{y}_k) + \mathbf{f}(t_{k+1}, \mathbf{y}_{k+1})) / 2$$

- To determine its stability and accuracy, we apply it to scalar ODE  $y' = \lambda y$ , obtaining

$$y_{k+1} = y_k + h (\lambda y_k + \lambda y_{k+1}) / 2$$

$$y_k = \left( \frac{1 + h\lambda/2}{1 - h\lambda/2} \right)^k y_0$$

- Method is stable if

$$\left| \frac{1 + h\lambda/2}{1 - h\lambda/2} \right| < 1$$

which holds for any  $h > 0$  when  $\text{Re}(\lambda) < 0$



## Trapezoid Method, continued

- Thus, trapezoid method is unconditionally stable
- Its growth factor

$$\begin{aligned} \frac{1 + h\lambda/2}{1 - h\lambda/2} &= \left(1 + \frac{h\lambda}{2}\right) \left(1 + \frac{h\lambda}{2} + \left(\frac{h\lambda}{2}\right)^2 + \left(\frac{h\lambda}{2}\right)^3 + \dots\right) \\ &= 1 + h\lambda + \frac{(h\lambda)^2}{2} + \frac{(h\lambda)^3}{4} + \dots \end{aligned}$$

agrees with expansion of  $e^{h\lambda}$  through terms of order  $h^2$ , so trapezoid method is second-order accurate

- For general system of ODEs  $\mathbf{y}' = \mathbf{f}(t, \mathbf{y})$ , trapezoid method has growth factor  $(\mathbf{I} + \frac{1}{2}h\mathbf{J}_f)(\mathbf{I} - \frac{1}{2}h\mathbf{J}_f)^{-1}$ , whose spectral radius is less than 1 provided eigenvalues of  $h\mathbf{J}_f$  lie in left half of complex plane < interactive example >



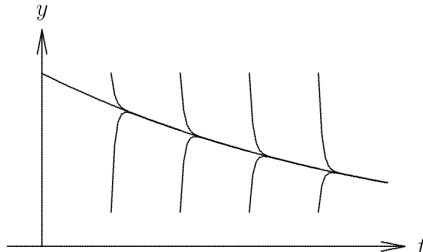
## Implicit Methods, continued

- We have now seen two examples of implicit methods that are unconditionally stable, but not all implicit methods have this property
- Implicit methods generally have larger stability regions than explicit methods, but allowable step size is not always unlimited
- Implicitness alone is not sufficient to guarantee stability



# Stiff Differential Equations

- Asymptotically stable solutions converge with time, and this has favorable property of damping errors in numerical solution
- But if convergence of solutions is too rapid, then difficulties of different type may arise
- Such ODE is said to be *stiff*



## Stiff ODEs, continued

- Stiff ODE corresponds to physical process whose components have disparate time scales or whose time scale is small compared to interval over which it is studied
- System of ODEs  $\mathbf{y}' = \mathbf{f}(t, \mathbf{y})$  is stiff if eigenvalues of its Jacobian matrix  $\mathbf{J}_f$  differ greatly in magnitude
- There may be eigenvalues with
  - large negative real parts, corresponding to strongly damped components of solution, or
  - large imaginary parts, corresponding to rapidly oscillating components of solution



## Stiff ODEs, continued

- Some numerical methods are inefficient for stiff ODEs because rapidly varying component of solution forces very small step sizes to maintain stability
- Stability restriction depends on rapidly varying component of solution, but accuracy restriction depends on slowly varying component, so step size may be more severely restricted by stability than by required accuracy
- For example, Euler's method is extremely inefficient for solving stiff ODEs because of severe stability limitation on step size
- Backward Euler method is suitable for stiff ODEs because of its unconditional stability
- Stiff ODEs need not be difficult to solve numerically, provided suitable method is chosen



## Example: Stiff ODE

- Consider scalar ODE

$$y' = -100y + 100t + 101$$

with initial condition  $y(0) = 1$

- General solution is  $y(t) = 1 + t + ce^{-100t}$ , and particular solution satisfying initial condition is  $y(t) = 1 + t$  (i.e.,  $c = 0$ )
- Since solution is linear, Euler's method is theoretically exact for this problem
- However, to illustrate effect of using finite precision arithmetic, let us perturb initial value slightly





## Example, continued

- With step size  $h = 0.1$ , first few steps for given initial values are

$t$	0.0	0.1	0.2	0.3	0.4
exact sol.	1.00	1.10	1.20	1.30	1.40
Euler sol.	0.99	1.19	0.39	8.59	-64.2
Euler sol.	1.01	1.01	2.01	-5.99	67.0

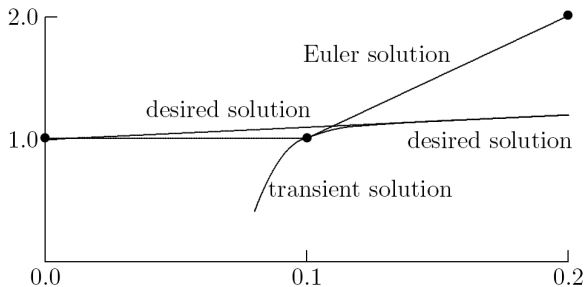
- Computed solution is incredibly sensitive to initial value, as each tiny perturbation results in wildly different solution
- Any point deviating from desired particular solution, even by only small amount, lies on different solution, for which  $c \neq 0$ , and therefore rapid transient of general solution is present

< interactive example >



## Example, continued

- Euler's method bases its projection on derivative at current point, and resulting large value causes numerical solution to diverge radically from desired solution
- Jacobian for this ODE is  $J_f = -100$ , so stability condition for Euler's method requires step size  $h < 0.02$ , which we are violating



## Example, continued

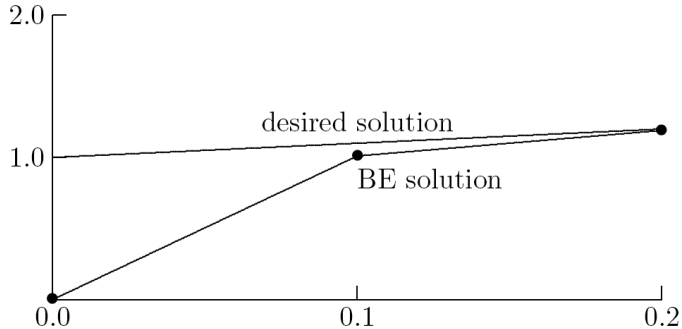
- Backward Euler method has no trouble solving this problem and is extremely *insensitive* to initial value

$t$	0.0	0.1	0.2	0.3	0.4
exact sol.	1.00	1.10	1.20	1.30	1.40
BE sol.	0.00	1.01	1.19	1.30	1.40
BE sol.	2.00	1.19	1.21	1.30	1.40

- Even with very large perturbation in initial value, by using derivative at next point rather than current point, transient is quickly damped out and backward Euler solution converges to desired solution after only few steps
- This behavior is consistent with unconditional stability of backward Euler method for stable solutions



## Example, continued



< interactive example >

# Numerical Methods for ODEs

There are many different methods for solving ODEs, most of which are of one of following types

- Taylor series
- Runge-Kutta
- Extrapolation
- Multistep
- Multivalued

We briefly consider each of these types of methods



# Taylor Series Methods

- Euler's method can be derived from Taylor series expansion
- By retaining more terms in Taylor series, we can generate higher-order single-step methods
- For example, retaining one additional term in Taylor series

$$\mathbf{y}(t+h) = \mathbf{y}(t) + h \mathbf{y}'(t) + \frac{h^2}{2} \mathbf{y}''(t) + \frac{h^3}{6} \mathbf{y}'''(t) + \dots$$

gives second-order method

$$\mathbf{y}_{k+1} = \mathbf{y}_k + h_k \mathbf{y}'_k + \frac{h_k^2}{2} \mathbf{y}''_k$$



# Taylor Series Methods, continued

- This approach requires computation of higher derivatives of  $\mathbf{y}$ , which can be obtained by differentiating  $\mathbf{y}' = \mathbf{f}(t, \mathbf{y})$  using chain rule, e.g.,

$$\begin{aligned}\mathbf{y}'' &= \mathbf{f}_t(t, \mathbf{y}) + \mathbf{f}_y(t, \mathbf{y}) \mathbf{y}' \\ &= \mathbf{f}_t(t, \mathbf{y}) + \mathbf{f}_y(t, \mathbf{y}) \mathbf{f}(t, \mathbf{y})\end{aligned}$$

where subscripts indicate partial derivatives with respect to given variable

- As order increases, expressions for derivatives rapidly become too complicated to be practical to compute, so Taylor series methods of higher order have not often been used in practice

< interactive example >



# Runge-Kutta Methods

- *Runge-Kutta methods* are single-step methods similar in motivation to Taylor series methods, but they do not require computation of higher derivatives
- Instead, Runge-Kutta methods simulate effect of higher derivatives by evaluating  $f$  several times between  $t_k$  and  $t_{k+1}$
- Simplest example is second-order *Heun's method*

$$\mathbf{y}_{k+1} = \mathbf{y}_k + \frac{h_k}{2} (\mathbf{k}_1 + \mathbf{k}_2)$$

where

$$\mathbf{k}_1 = \mathbf{f}(t_k, \mathbf{y}_k)$$

$$\mathbf{k}_2 = \mathbf{f}(t_k + h_k, \mathbf{y}_k + h_k \mathbf{k}_1)$$





# Runge-Kutta Methods, continued

- Heun's method is analogous to implicit trapezoid method, but remains explicit by using Euler prediction  $\mathbf{y}_k + h_k \mathbf{k}_1$  instead of  $\mathbf{y}(t_{k+1})$  in evaluating  $\mathbf{f}$  at  $t_{k+1}$
- Best known Runge-Kutta method is classical fourth-order scheme

$$\mathbf{y}_{k+1} = \mathbf{y}_k + \frac{h_k}{6} (\mathbf{k}_1 + 2\mathbf{k}_2 + 2\mathbf{k}_3 + \mathbf{k}_4)$$

where

$$\mathbf{k}_1 = \mathbf{f}(t_k, \mathbf{y}_k)$$

$$\mathbf{k}_2 = \mathbf{f}(t_k + h_k/2, \mathbf{y}_k + (h_k/2)\mathbf{k}_1)$$

$$\mathbf{k}_3 = \mathbf{f}(t_k + h_k/2, \mathbf{y}_k + (h_k/2)\mathbf{k}_2)$$

$$\mathbf{k}_4 = \mathbf{f}(t_k + h_k, \mathbf{y}_k + h_k \mathbf{k}_3)$$

< interactive example >



## Runge-Kutta Methods, continued

- To proceed to time  $t_{k+1}$ , Runge-Kutta methods require no history of solution prior to time  $t_k$ , which makes them *self-starting* at beginning of integration, and also makes it easy to change step size during integration
- These facts also make Runge-Kutta methods relatively easy to program, which accounts in part for their popularity
- Unfortunately, classical Runge-Kutta methods provide no error estimate on which to base choice of step size



# Runge-Kutta Methods, continued

- Fehlberg devised *embedded Runge-Kutta* method that uses six function evaluations per step to produce both fifth-order and fourth-order estimates of solution, whose difference provides estimate for local error
- Another embedded Runge-Kutta method is due to Dormand and Prince
- This approach has led to automatic Runge-Kutta solvers that are effective for many problems, but which are relatively inefficient for stiff problems or when very high accuracy is required
- It is possible, however, to define *implicit* Runge-Kutta methods with superior stability properties that are suitable for solving stiff ODEs



# Extrapolation Methods

- *Extrapolation methods* are based on use of single-step method to integrate ODE over given interval  $t_k \leq t \leq t_{k+1}$  using several different step sizes  $h_i$ , and yielding results denoted by  $Y(h_i)$
- This gives discrete approximation to function  $Y(h)$ , where  $Y(0) = \mathbf{y}(t_{k+1})$
- Interpolating polynomial or rational function  $\hat{Y}(h)$  is fit to these data, and  $\hat{Y}(0)$  is then taken as approximation to  $Y(0)$
- Extrapolation methods are capable of achieving very high accuracy, but they are much less efficient and less flexible than other methods for ODEs, so they are not often used unless extremely high accuracy is required and cost is not significant factor < interactive example >



# Multistep Methods

- *Multistep methods* use information at more than one previous point to estimate solution at next point
- *Linear multistep* methods have form

$$\mathbf{y}_{k+1} = \sum_{i=1}^m \alpha_i \mathbf{y}_{k+1-i} + h \sum_{i=0}^m \beta_i \mathbf{f}(t_{k+1-i}, \mathbf{y}_{k+1-i})$$

- Parameters  $\alpha_i$  and  $\beta_i$  are determined by polynomial interpolation
- If  $\beta_0 = 0$ , method is explicit, but if  $\beta_0 \neq 0$ , method is implicit
- Implicit methods are usually more accurate and stable than explicit methods, but require starting guess for  $\mathbf{y}_{k+1}$



## Multistep Methods, continued

- Starting guess is conveniently supplied by explicit method, so the two are used as *predictor-corrector* pair
- One could use corrector repeatedly (i.e., fixed-point iteration) until some convergence tolerance is met, but it may not be worth expense
- So fixed number of corrector steps, often only one, may be used instead, giving *PECE* scheme (predict, evaluate, correct, evaluate)
- Although it has no effect on value of  $\mathbf{y}_{k+1}$ , second evaluation of  $\mathbf{f}$  in PECE scheme yields improved value of  $\mathbf{y}'_{k+1}$  for future use



## Multistep Methods, continued

- Alternatively, nonlinear equation for  $\mathbf{y}_{k+1}$  given by implicit multistep method can be solved by Newton's method or similar method, again with starting guess supplied by solution at previous step or by explicit multistep method
- Newton's method or close variant of it is essential when using an implicit multistep method designed for stiff ODEs, as fixed-point iteration fails to converge for reasonable step sizes

< interactive example >



## Examples: Multistep Methods

- Simplest second-order accurate explicit two-step method is

$$\mathbf{y}_{k+1} = \mathbf{y}_k + \frac{h}{2}(3\mathbf{y}'_k - \mathbf{y}'_{k-1})$$

- Simplest second-order accurate implicit method is trapezoid method

$$\mathbf{y}_{k+1} = \mathbf{y}_k + \frac{h}{2}(\mathbf{y}'_{k+1} + \mathbf{y}'_k)$$

- One of most popular pairs of multistep methods is explicit fourth-order Adams-Bashforth predictor

$$\mathbf{y}_{k+1} = \mathbf{y}_k + \frac{h}{24}(55\mathbf{y}'_k - 59\mathbf{y}'_{k-1} + 37\mathbf{y}'_{k-2} - 9\mathbf{y}'_{k-3})$$

and implicit fourth-order Adams-Moulton corrector

$$\mathbf{y}_{k+1} = \mathbf{y}_k + \frac{h}{24}(9\mathbf{y}'_{k+1} + 19\mathbf{y}'_k - 5\mathbf{y}'_{k-1} + \mathbf{y}'_{k-2})$$





# Examples: Multistep Methods

- *Backward differentiation formulas* form another important family of implicit multistep methods
- BDF methods, typified by popular formula

$$\mathbf{y}_{k+1} = \frac{1}{11}(18\mathbf{y}_k - 9\mathbf{y}_{k-1} + 2\mathbf{y}_{k-2}) + \frac{6h}{11}\mathbf{y}'_{k+1}$$

are effective for solving stiff ODEs

< interactive example >



# Multistep Adams Methods

- Stability and accuracy of some Adams methods are summarized below
  - Stability threshold* indicates left endpoint of stability interval for scalar ODE
  - Error constant* indicates coefficient of  $h^{p+1}$  term in local truncation error, where  $p$  is order of method

Explicit Methods		
Order	Stability threshold	Error constant
1	-2	1/2
2	-1	5/12
3	-6/11	3/8
4	-3/10	251/720

Implicit Methods		
Order	Stability threshold	Error constant
1	$-\infty$	-1/2
2	$-\infty$	-1/12
3	-6	-1/24
4	-3	-19/720

- Implicit methods are both more stable and more accurate than corresponding explicit methods of same order



# Properties of Multistep Methods

- They are not self-starting, since several previous values of  $y_k$  are needed initially
- Changing step size is complicated, since interpolation formulas are most conveniently based on equally spaced intervals for several consecutive points
- Good local error estimate can be determined from difference between predictor and corrector
- They are relatively complicated to program
- Being based on interpolation, they can efficiently provide solution values at output points other than integration points



## Properties of Multistep, continued

- Implicit methods have much greater region of stability than explicit methods, but must be iterated to convergence to enjoy this benefit fully
  - PECE scheme is actually explicit, though in a somewhat complicated way
- Although implicit methods are more stable than explicit methods, they are still not necessarily unconditionally stable
  - No multistep method of greater than second order is unconditionally stable, even if it is implicit
- Properly designed implicit multistep method can be very effective for solving stiff ODEs



# Multivalued Methods

- With multistep methods it is difficult to change step size, since past history of solution is most easily maintained at equally spaced intervals
- Like multistep methods, *multivalued methods* are also based on polynomial interpolation, but avoid some implementation difficulties associated with multistep methods
- One key idea motivating multivalued methods is observation that interpolating polynomial itself can be evaluated at any point, not just at equally spaced intervals
- Equal spacing associated with multistep methods is artifact of representation as linear combination of successive solution and derivative values with fixed weights



## Multivalued Methods, continued

- Another key idea in implementing multivalued methods is representing interpolating polynomial so that parameters are values of solution and its derivatives at  $t_k$
- This approach is analogous to using Taylor, rather than Lagrange, representation of polynomial
- Solution is advanced in time by simple transformation of representation from one point to next, and it is easy to change step size
- Multivalued methods are mathematically equivalent to multistep methods but are more convenient and flexible to implement, so most modern software implementations are based on them



# Example: Multivalued Method

- Consider four-value method for solving scalar ODE

$$y' = f(t, y)$$

- Instead of representing interpolating polynomial by its value at four different points, we represent it by its value and first three derivatives at single point  $t_k$

$$\mathbf{y}_k = \begin{bmatrix} y_k \\ hy'_k \\ (h^2/2)y''_k \\ (h^3/6)y'''_k \end{bmatrix}$$



## Example: Multivalue Method

- By differentiating Taylor series

$$y(t_k + h) = y(t_k) + hy' + \frac{h^2}{2}y'' + \frac{h^3}{6}y''' + \dots$$

three times, we see that corresponding values at next point  $t_{k+1} = t_k + h$  are given approximately by transformation

$$\hat{y}_{k+1} = B y_k$$

where matrix  $B$  is given by

$$B = \begin{bmatrix} 1 & 1 & 1 & 1 \\ 0 & 1 & 2 & 3 \\ 0 & 0 & 1 & 3 \\ 0 & 0 & 0 & 1 \end{bmatrix}$$





## Example: Multivalue Method

- We have not yet used ODE, however, so we add correction term to foregoing prediction to obtain final value

$$\mathbf{y}_{k+1} = \hat{\mathbf{y}}_{k+1} + \alpha \mathbf{r}$$

where  $\mathbf{r}$  is fixed 4-vector and

$$\alpha = h(f(t_{k+1}, y_{k+1}) - \hat{y}'_{k+1})$$

- For consistency, i.e., for ODE to be satisfied, we must have  $r_2 = 1$
- Remaining three components of  $\mathbf{r}$  can be chosen in various ways, resulting in different methods, analogous to different choices of parameters in multistep methods



## Example, continued

- For example, four-value method with

$$\mathbf{r} = \left[ \frac{3}{8} \quad 1 \quad \frac{3}{4} \quad \frac{1}{6} \right]^T$$

is equivalent to implicit fourth-order Adams-Moulton method given earlier



# Multivalued Methods, continued

- It is easy to change step size with multivalued methods: we need merely rescale components of  $y_k$  to reflect new step size
- It is also easy to change order of method simply by changing components of  $r$
- These two capabilities, combined with sophisticated tests and strategies for deciding when to change order and step size, have led to development of powerful and efficient software packages for solving ODEs based on variable-order/variable-step methods



# Variable-Order/Variable-Step Solvers

- Such routines are analogous to adaptive quadrature routines: they automatically adapt to given problem, varying order and step size of integration method as necessary to meet user-supplied error tolerance efficiently
- Such routines often have options for solving either stiff or nonstiff problems, and some even detect stiffness automatically and select appropriate method accordingly
- Ability to change order easily also obviates need for special starting procedures: one can simply start with first-order method, which requires no additional starting values, and let automatic order/step size selection procedure increase order as needed for required accuracy

