

Ordinary differential equations

The set of ordinary differential equations (ODE) can always be reduced to a set of coupled first order differential equations.

For example, Newton's law is usually written by a second order differential equation $m\ddot{\vec{r}} = F[\vec{r}, \dot{\vec{r}}, t]$. In Hamiltonian dynamics, the same problem leads to the set of first order equations $\dot{\vec{p}} = -\frac{\partial H}{\partial \vec{q}}$ and $\dot{\vec{q}} = \frac{\partial H}{\partial \vec{p}}$.

We will therefore concentrate on the system of equations

$$\frac{dy_i}{dx} = f_i(x, y_0, y_1, \dots, y_{N-1}) \quad (1)$$

To solve the problem, the **boundary conditions** need to be specified. They can be arbitrary complicated - for example a set of nonlinear equations relating values $y_i(x_l)$ and there derivatives $\dot{y}_i(x_l)$ at certain points x_l .

Boundary conditions can be divided into categories

- Initial value problems (all necessary conditions specified at starting point)
- Two point boundary problems (part of the conditions specified at one point x_0 and the rest at x_1).
- More complicated problems

In this chapter, we will concentrate on the Initial value problems and we will show in the *Density functional theory* chapter how to solve The two points boundary problem - by so called shooting method.

We will implement these methods

- Runge-Kutta method : general purpose routine
- Numerov's algorithm: $\ddot{y} = f(t)y(t)$ (for Schroedinger equation)
- Verlet algorithm: $\ddot{y} = F[y(t), t]$ (for molecular dynamics because it is more stable and preserves total energy)

In this chapter, we will concentrate on "most often" general purpose routine. In subsequent chapters we will implement the two other methods.

The simplest method for solving differential equations is Euler's method

$$y_{i+1} = y_i + hf(x_i, y_i) + O(h^2) \quad (2)$$

$$x_{i+1} = x_i + h \quad (3)$$

but it is **not advisable** to use it in practice.

When solving differential equation, we **usually** look for a very smooth function $y(x)$ and in order that the step size can be finite and precision loss is not very dramatic, it is recommended to use higher order routine. In addition, Euler's routine is very unstable because it is not "symmetric". However, keep in mind that sometimes (when the solution is not smooth function) high order is not necessary better.

The derivative $f(x_i, y_i)$ is taken at the beginning of the interval $[x_i, x_i + h]$. The stability and precision would increase if one could estimate derivative in the middle of the interval, i.e., $f(x_i + h/2, y(x_i + h/2))$.

The second order Runge-Kutta method implements the above idea

$$k_1 = hf(x_i, y_i) \quad (4)$$

$$y_{i+1} = y_i + hf\left(x_i + \frac{1}{2}h, y_i + \frac{1}{2}k_1\right) + O(h^3) \quad (5)$$

~~It is called second order, because error is of the order of h^3 . In general, the error of the~~

n -th order routine is $O(h^{n+1})$.

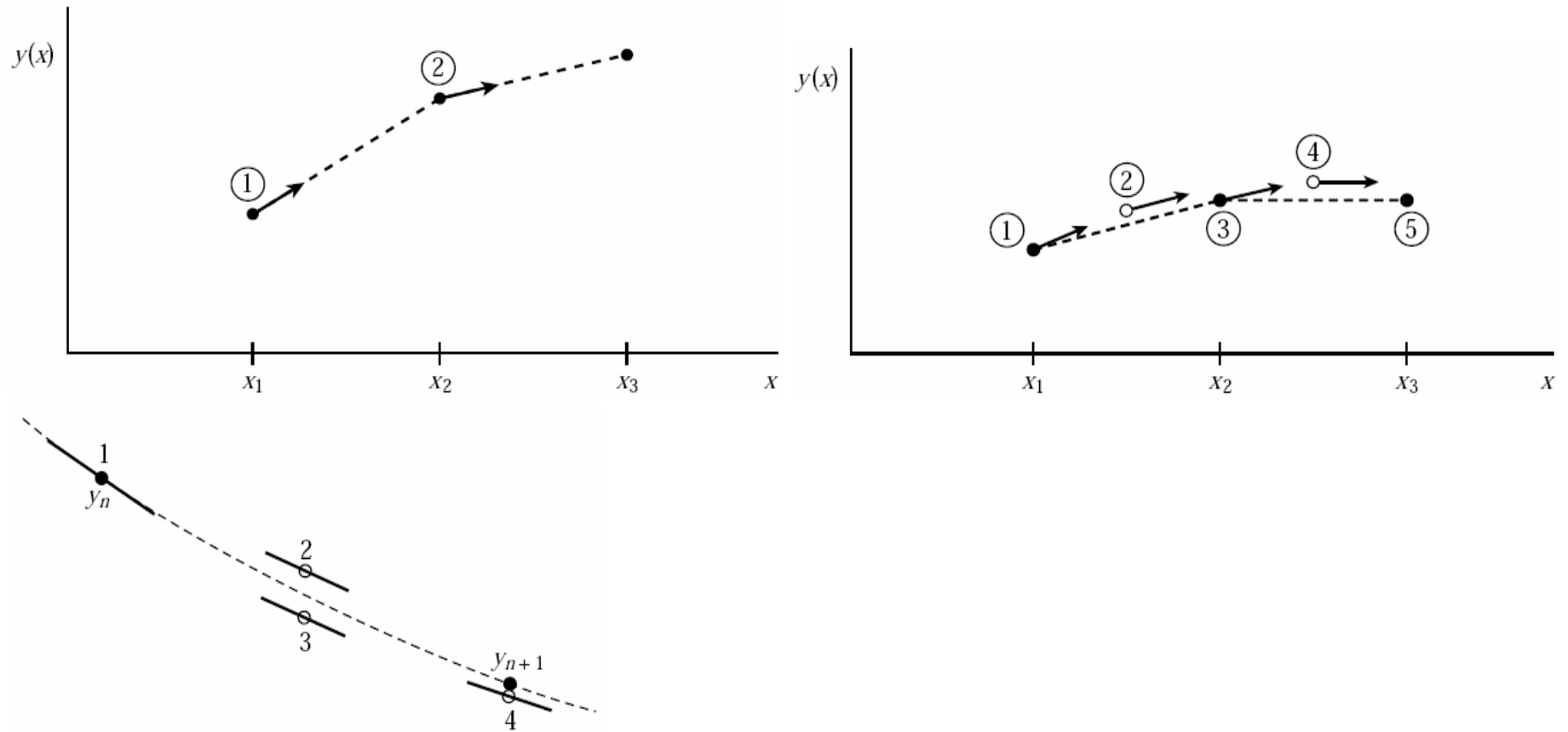


Figure 1: top left: Euler's algorithm, top right: Midpoint or Second order Runge-Kutta method
bottom: Forth order Runge-Kutta

Most popular is the **forth-order Runge Kutta** (RK4) method:

$$k_1 = hf(x_i, y_i) \quad (6)$$

$$k_2 = hf\left(x_i + \frac{1}{2}h, y_i + \frac{1}{2}k_1\right) \quad (7)$$

$$k_3 = hf\left(x_i + \frac{1}{2}h, y_i + \frac{1}{2}k_2\right) \quad (8)$$

$$k_4 = hf(x_i + h, y_i + k_3) \quad (9)$$

$$y_{i+1} = y_i + \frac{1}{6}k_1 + \frac{1}{3}k_2 + \frac{1}{3}k_3 + \frac{1}{6}k_4 + O(h^5) \quad (10)$$

How to understand the method? Looking at the above figure, we see:

- k_1 is the slope at the beginning of the interval;
- k_2 is the slope at the midpoint of the interval, using slope k_1 to determine the value of y at the point $x_i + h/2$ using Euler's method;
- k_3 is again the slope at the midpoint, but now using the slope k_2 to determine the y -value;
- k_4 is the slope at the end of the interval, with its y -value determined using k_3 ;

- in averaging the four slopes, greater weight is given to the slopes at the midpoint.

The RK4 method is a fourth-order method, meaning that the error per step is on the order of h^5 , while the total accumulated error has order h^4 . With only four function evaluations, for fourth order accuracy is extremely good.

The code for RK4 is given below:

```

////////////////////////////////////
////////// METHOD OF RUNGKE-KUTTA 4-th ORDER, FIXED STEP //////////
////////////////////////////////////
template <class functor, class container>
void rungek4(double x, double h, container& y, functor& derivs)
{
  // Runke-Kutta of fourth order for fixed time-step h.
  // derivs is a function which evaluates RHS of the ODE system, y contains values of dependent variables
  static container yt(y.size()), fk1(y.size()), fk2(y.size()), fk3(y.size()), fk4(y.size()); // temporary arrays
  double h2=h*0.5;
  double xh = x + h2;
  int N = y.size();
  derivs(x, y, fk1); // First step : evaluating k1
  for (int i=0; i<N; i++) yt[i] = y[i] + h2*fk1[i]; // Preparing second step by ty <- y + k1/2
  derivs(xh, yt, fk2); // Second step : evaluating k2
  for (int i=0; i<N; i++) yt[i] = y[i] + h2*fk2[i]; // Preparing third step by yt <- y + k2/2
  derivs(xh, yt, fk3); // Third step : evaluating k3
  for (int i=0; i<N; i++) yt[i] = y[i] + h*fk3[i]; // Preparing fourth step yt <- y + k3
  derivs(x+h, yt, fk4); // Final step : evaluating k4
  double h6=h/6.0; // Accumulate increments with proper weights
  for (int i=0; i<N; i++) y[i] += h6*(fk1[i]+2.0*(fk2[i]+fk3[i])+fk4[i]);
}

void simple_pendulum(double x, const function1D<double>& y, function1D<double>& dydx)
{
  dydx[0] = y[1];
  dydx[1] = -y[0];
}
...
for (int i=0; i<M; i++, t+=dh) rungek4(t, dh, y, simple_pendulum);

```

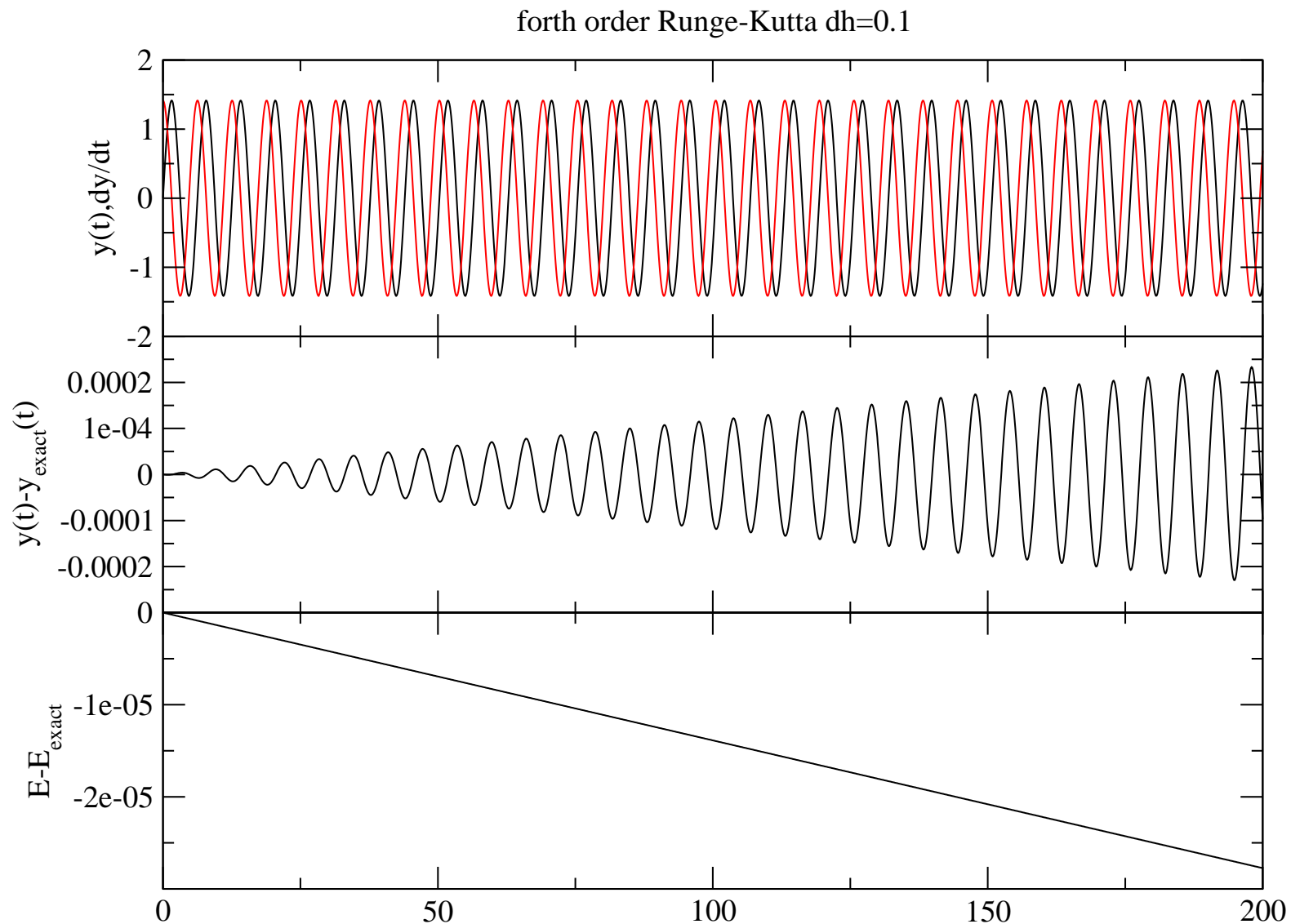


Figure 2: Fixed step Runge Kutta for simple pendulum. The error is increasing linearly with time and energy is being lost linearly with time.

In practice, it is usually better to use **adaptive step**. In this case, one needs some way of estimating error of each step and decrease the step size if necessary. Two types of algorithms are very popular

- **Step doubling**
- **Embedded methods**

Step doubling builds on the fact that when performing half of the step, the error is smaller and by comparing the error of full step and half step, we have good estimation of the error:

$$y(t + 2h) = y_1 + (2h)^5 C \quad \text{one big } 2h \text{ step} \quad (11)$$

$$y(t + 2h) = y_2 + 2h^5 C \quad \text{two small } h \text{ steps} \quad (12)$$

The difference of the two solutions is $\Delta = y_2 - y_1 - 30h^5 C$ which can serve as an error estimate Δ

$$\Delta \equiv y_2 - y_1 \quad (13)$$

The solution might even be "improved" to fifth order by evaluating

$$y(x + 2h) = y_2 + \frac{\Delta}{15} + O(h^6). \quad (14)$$

Embedded Runge-Kutta methods

For 4-th order RK method one needs 4 function evaluations, for higher order accuracy (of order M) one typically needs more function evaluations, namely, $M+2$.

The method due to Fehlberg needs 6 function evaluations for 5-th order accuracy. In addition, one can use the same 6 function values to get 4-th order accuracy. The difference can therefore be used as an error estimate.

The embedded fifth-order RK formulas are

$$k_0 = hf(x_i, y_i) \quad (15)$$

$$k_1 = hf(x_i + a_1h, y_i + b_{10}k_0) \quad (16)$$

$$\dots \quad (17)$$

$$k_5 = hf(x_i + a_5h, y_i + b_{50}k_0 + \dots + b_{54}k_4) \quad (18)$$

$$y_{i+1} = y_i + c_0k_0 + c_1k_1 + \dots + c_5k_5 + O(h^6) \quad (19)$$

$$y'_{i+1} = y_i + c'_0k_0 + c'_1k_1 + \dots + c'_5k_5 + O(h^5) \quad (20)$$

$$\Delta = y_{i+1} - y'_{i+1} = \sum_{i=0}^5 (c_i - c'_i)k_i \quad (21)$$

where coefficients are

i	a_i	c_i	c'_i	b_{i0}	b_{i1}	b_{i2}	b_{i3}	b_{i4}	b_{i5}
0		$\frac{37}{378}$	$\frac{2825}{27648}$						
1	$\frac{1}{5}$	0	0	$\frac{1}{5}$					
2	$\frac{3}{10}$	$\frac{250}{621}$	$\frac{18575}{48384}$	$\frac{3}{40}$	$\frac{9}{40}$				
3	$\frac{3}{5}$	$\frac{125}{594}$	$\frac{13525}{55296}$	$\frac{3}{10}$	$-\frac{9}{10}$	$\frac{6}{5}$			
4	1	0	$\frac{277}{14336}$	$-\frac{11}{54}$	$\frac{5}{2}$	$-\frac{70}{27}$	$\frac{35}{27}$		
5	$\frac{7}{8}$	$\frac{512}{1771}$	$\frac{1}{4}$	$\frac{1631}{55296}$	$\frac{175}{512}$	$\frac{575}{13824}$	$\frac{44275}{110592}$	$\frac{253}{4096}$	

The error estimate we have is for the forth-order value y'_{i+1} and is proportional to h^5 or $\Delta = Ch^5$. If we made a step h_1 and got the error Δ_1 , we can figure out what the step needs to be, to get the error of the order of Δ_0

$$h_0 = h_1 \left| \frac{\Delta_0}{\Delta_1} \right|^{1/5} \quad (23)$$

Although the error estimate is for the forth-order value, we obviously accept fifth order estimate y_{i+1} .

The above formula can be used in two ways

- If obtained error Δ_1 (by taking step h_1) is smaller than desired accuracy Δ_0 , we can increase step to $h_0 = h_1 \left| \frac{\Delta_0}{\Delta_1} \right|^{1/5}$ when taking the next step.
- If the obtained error is too big, we have to backtrack and take the same step again by choosing smaller step $h_0 = h_1 \left| \frac{\Delta_0}{\Delta_1} \right|^{1/5}$.

Backtracking is "expensive" because we throw away 5-6 function evaluations! To play it safe, we rather increase the step a little less than we should and decrease slightly more than we could

$$h_0 = \begin{cases} Sh_1 \left| \frac{\Delta_0}{\Delta_1} \right|^{0.2} & \Delta_0 > \Delta_1 \\ Sh_1 \left| \frac{\Delta_0}{\Delta_1} \right|^{0.25} & \Delta_0 < \Delta_1 \end{cases} \quad (24)$$

where $S \approx 0.9$ is safety factor.

We need routine to solve a system of coupled equations and therefore y_i is a vector of values, however, we treated Δ as a number. In the code, we take a number Δ to be the largest component of vector Δ_m since error of all components needs to be kept below desired accuracy.

Many times, the components (corresponding to different equations) differ dramatically in value. In many cases, we want to multiply different components with different factors when evaluating error

$$\Delta = \max(\Delta_m / \text{yscal}_m) \quad (25)$$

A good choice for scaling factors is

$$\text{yscal}_m = |y_m| + |f_m h| + 10^{-3} \quad (26)$$

where y_m is m -th component of the vector at each step i and f_m is the derivative at the same step. This ensures that the relative error is bounded rather than absolute.

The object implementating the adaptive Runge-Kutta method can be a `template` of `container` and the member function `Step` can be a `template` of a function which evaluates derivatives:

```

template <class container>
class VRungeKutta
{
    static const double SAFETY = 0.9, PGROW = -0.2, PSHRNK = -0.25, ERRCON = 1.89e-4;
    container yscal; // scaling factors
    double ht; // current time-step
    double eps; // Accuracy we check at each step
    double xt, t_stop; // start and stop time
    int nok, nbad; // number of bad and good steps
public:
    VRungeKutta(double t_start_, double t_stop_, int size, double accuracy=1e-6, double hmin=1e-4, double h1=0.1) :
        yscal(size), ht(h1), eps(accuracy), xt(t_start_), t_stop(t_stop_), nok(0), nbad(0){};
public:
    template <class functor>
    bool Step(double& x, container& y, functor& derivs){
        static container dydx(y.size()); // // current derivatives are local to this functions
        // Calculates derivatives when new step is taken
        derivs(xt,y,dydx);
        // good way of determining desired accuracy
        for (int i=0; i<y.size(); i++) yscal[i] = fabs(y[i]) + fabs(dydx[i]*ht) + 1e-3;
        double hnext, hdid;
        // Cals the Runge-Kutta routine
        RKStep(y, dydx, xt, ht, hdid, hnext, derivs); // Make one RK step
        if (hdid == ht) ++nok; else ++nbad; // good or bad step
        ht = hnext;
        x = xt;
        return x<t_stop;
    }
    double h(){return ht;}
private:
    template <class functor>
    void RKTry(container& y, container& dydx, double& x, double h, container& yout, container& yerr, functor& derivs);
    template <class functor>
    void RKStep(container& y, container& dydx, double& x, double htry, double& hdid, double& hnext, functor& derivs);
};

```

```
template <class container>
template <class functor>
void VRungeKutta<container>::RKStep(container& y, container& dydx, double& x, double htry,
    double& hdid, double& hnext, functor& derivs)
//Fifth-order Runge-Kutta step with monitoring of local truncation error to ensure accuracy and
//adjust stepsize. Input are the dependent variable vector y[...] and its derivative dydx[...]
//at the starting value of the independent variable x. Also input are the stepsize to be attempted
//htry, the required accuracy eps, and the vector yscal[...] against which the error is
//scaled. On output, y and x are replaced by their new values, hdid is the stepsize that was
//actually accomplished, and hnext is the estimated next stepsize. derivs is the user-supplied
//routine that computes the right-hand side derivatives.
{
    container yerr(y.size()), ytemp(y.size());
    double errmax;
    double h=htry; // Set stepsize to the initial trial value.
    for (;;) {
        RKTry(y, dydx, x, h, ytemp, yerr, derivs); // Take a step.
        errmax=0.0; // Evaluate accuracy.
        for (int i=0; i<y.size(); i++) errmax=std::max(errmax,fabs(yerr[i]/yscal[i]));
        errmax /= eps; // Scale relative to required tolerance.
        if (errmax <= 1.0) break; // Step succeeded. Compute size of next step.
        double htemp=SAFETY*h*pow(errmax,PSHRNK); // Truncation error too large, reduce stepsize.
        h=(h >= 0.0 ? std::max(htemp,0.1*h) : std::min(htemp,0.1*h)); // No more than a factor of 10.
        if ((x+h) == x) std::cerr<<"stepsize underflow in rkqs"<<std::endl;
    }
    if (errmax > ERRCON) hnext=SAFETY*h*pow(errmax,PGROW); // Step is too small, increase it next time
    else hnext=5.0*h; // No more than a factor of 5 increase.
    x += (hdid=h);
    for (int i=0; i<y.size();i++) y[i] = ytemp[i];
}
```

```

template <class container>
template <class functor>
void VRungeKutta<container>::RKTry(container& y, container& dydx, double& x, double h, container& yout, container& yerr,
    //Given values for variables y[...] and their derivatives dydx[...] known at x, use
    //the fifth-order Cash-Karp Runge-Kutta method to advance the solution over an interval h
    //and return the incremented variables as yout[...]. Also return an estimate of the local
    //truncation error in yout using the embedded fourth-order method. The user supplies the routine
    //derivs(x,y,dydx), which returns derivatives dydx at x.
{
    static double as[] = {0, 0.2, 0.3, 0.6, 1.0, 0.875};
    static double b10 = 0.2, b20=3.0/40.0, b30=0.3, b40 = -11.0/54.0, b50=1631.0/55296.0;
    static double b21 = 9.0/40.0, b31 = -0.9, b41=2.5, b51=175.0/512.0;
    static double b32 = 1.2, b42 = -70.0/27.0, b52=575.0/13824.0;
    static double b43 = 35.0/27.0, b53=44275.0/110592.0;
    static double b54 = 253.0/4096.0;
    static double c0=37.0/378.0,c2=250.0/621.0,c3=125.0/594.0,c5=512.0/1771.0;
    static double dc4 = -277.00/14336.0;
    static double dc0=c0-2825.0/27648.0,dc2=c2-18575.0/48384.0,dc3=c3-13525.0/55296.0,dc5=c5-0.25;
    int N=y.size();
    static container ak2(N), ak3(N), ak4(N), ak5(N), ak6(N), yt(N);

    for (int i=0; i<N; i++) yt[i] = y[i] + b10*h*dydx[i]; // First step.
    derivs(x+as[1]*h, yt, ak2); // Second step.
    for (int i=0; i<N; i++) yt[i] = y[i] + h*(b20*dydx[i]+b21*ak2[i]);
    derivs(x+as[2]*h,yt,ak3); // Third step.
    for (int i=0; i<N; i++) yt[i] = y[i] + h*(b30*dydx[i]+b31*ak2[i]+b32*ak3[i]);
    derivs(x+as[3]*h,yt,ak4); // Fourth step.
    for (int i=0; i<N; i++) yt[i] = y[i] + h*(b40*dydx[i]+b41*ak2[i]+b42*ak3[i]+b43*ak4[i]);
    derivs(x+as[4]*h,yt,ak5); // Fifth step.
    for (int i=0;i<N;i++) yt[i] = y[i] + h*(b50*dydx[i]+b51*ak2[i]+b52*ak3[i]+b53*ak4[i]+b54*ak5[i]);
    derivs(x+as[5]*h,yt,ak6); // Sixth step.
    // Accumulate increments with proper weights.
    for (int i=0;i<N;i++) yout[i] = y[i] + h*(c0*dydx[i]+c2*ak3[i]+c3*ak4[i]+c5*ak6[i]);
    // Estimate error as difference between fourth and fifth order methods.
    for (int i=0; i<N; i++) yerr[i] = h*(dc0*dydx[i]+dc2*ak3[i]+dc3*ak4[i]+dc4*ak5[i]+dc5*ak6[i]);
}

```

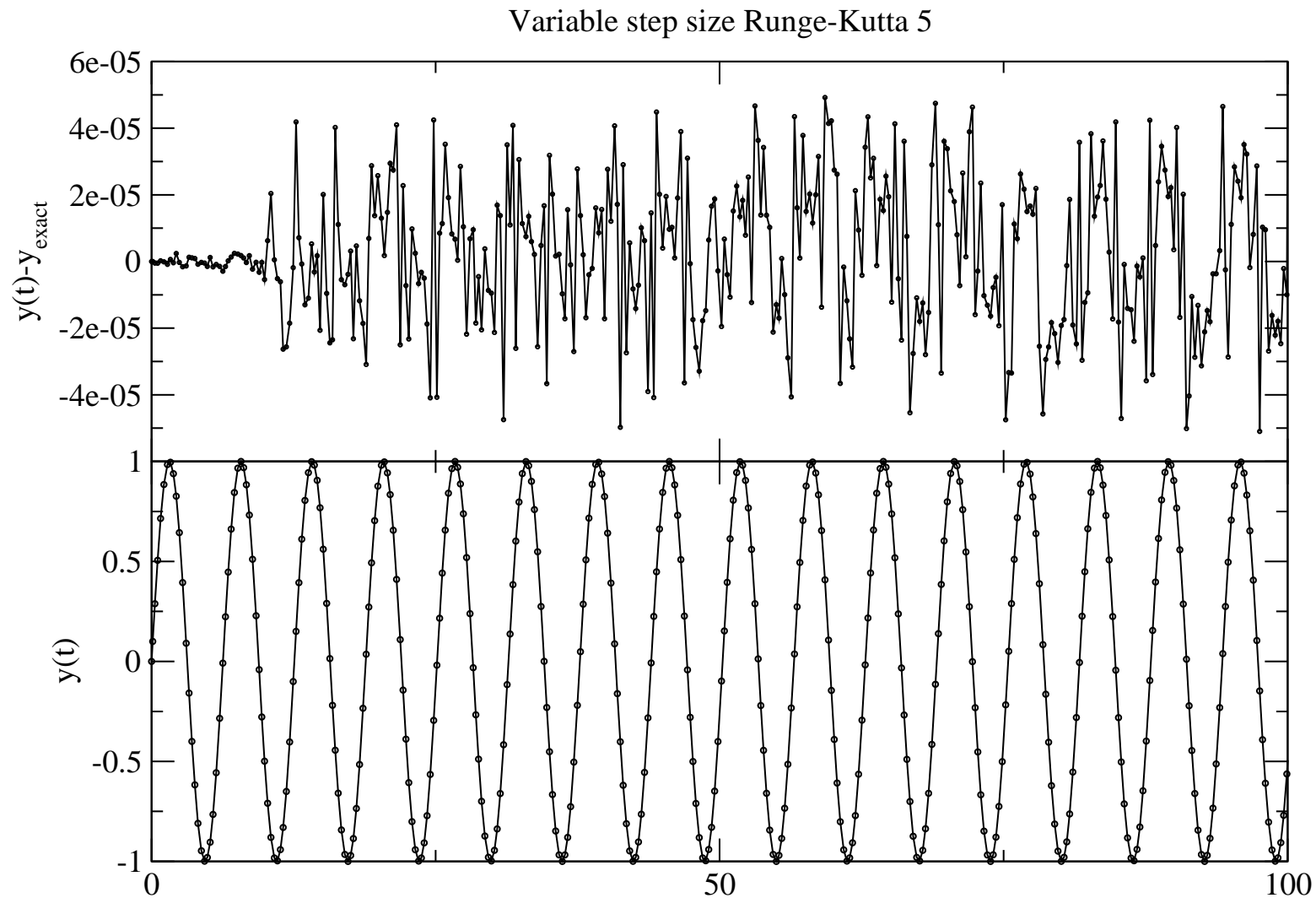
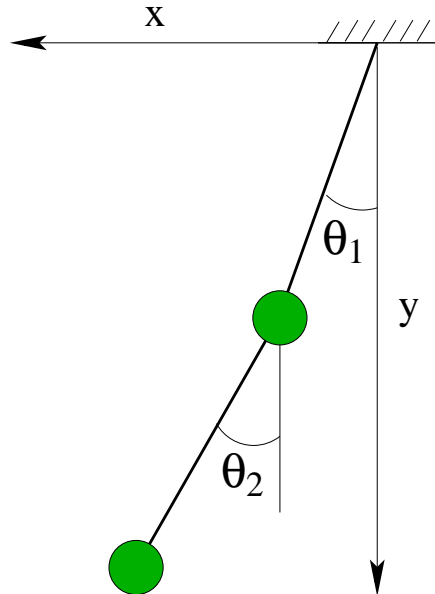


Figure 3: Variable Runge Kutta step for simple pendulum. The error is not increasing very fast with time.

A somewhat more interesting example is double pendulum. It is well known that double pendulum is chaotic for not too small energy and the Poincare plots in chaotic regime show nice patterns.



$$\vec{r}_1 = (l \sin \theta_1, l \cos \theta_1) \quad (27)$$

$$\vec{r}_2 = (l(\sin \theta_1 + \sin \theta_2), l(\cos \theta_1 + \cos \theta_2)) \quad (28)$$

$$T = \frac{1}{2} m \dot{\vec{r}}_1^2 + \frac{1}{2} m \dot{\vec{r}}_2^2 \quad (29)$$

$$V = 3mgl - m\vec{g}\vec{r}_1 - m\vec{g}\vec{r}_2 \quad (30)$$

$$L = \frac{1}{2} ml^2 [2\dot{\theta}_1^2 + \dot{\theta}_2^2 + 2 \cos(\theta_1 - \theta_2) \dot{\theta}_1 \dot{\theta}_2] - mgl [3 - 2 \cos \theta_1 - \cos \theta_2] \quad (31)$$

$$p_1 = \frac{\partial L}{\partial \dot{\theta}_1} = ml^2 [2\dot{\theta}_1 + \cos(\theta_1 - \theta_2) \dot{\theta}_2] \quad (32)$$

$$p_2 = \frac{\partial L}{\partial \dot{\theta}_2} = ml^2 [\dot{\theta}_2 + \cos(\theta_1 - \theta_2) \dot{\theta}_1] \quad (33)$$

$$H = \frac{1}{2ml^2} \frac{[p_1^2 + 2p_2^2 - 2p_1p_2 \cos(\theta_1 - \theta_2)]}{1 + \sin^2(\theta_1 - \theta_2)} + mgl[3 - 2 \cos \theta_1 - \cos \theta_2] \quad (34)$$

$$(35)$$

$$\dot{\theta}_1 = \frac{\partial H}{\partial p_1} = \frac{p_1 - p_2 \cos(\theta_1 - \theta_2)}{ml^2[1 + \sin^2(\theta_1 - \theta_2)]} \quad (36)$$

$$\dot{\theta}_2 = \frac{\partial H}{\partial p_2} = \frac{2p_2 - p_1 \cos(\theta_1 - \theta_2)}{ml^2[1 + \sin^2(\theta_1 - \theta_2)]} \quad (37)$$

$$\dot{p}_1 = -\frac{\partial H}{\partial \theta_1} = -2mgl \sin \theta_1 - C_1 + C_2 \quad (38)$$

$$\dot{p}_2 = -\frac{\partial H}{\partial \theta_2} = -mgl \sin \theta_2 + C_1 - C_2 \quad (39)$$

$$C_1 = \frac{p_1 p_2 \sin(\theta_1 - \theta_2)}{ml^2[1 + \sin^2(\theta_1 - \theta_2)]} \quad (40)$$

$$C_2 = \frac{[p_1^2 + 2p_2^2 - 2p_1p_2 \cos(\theta_1 - \theta_2)] \sin(2(\theta_1 - \theta_2))}{2ml^2[1 + \sin^2(\theta_1 - \theta_2)]^2} \quad (41)$$

$$\tilde{p} = \frac{p}{ml^2\omega_0} \quad (42)$$

$$\tilde{t} = t\omega_0 \quad (43)$$

$$\omega_0^2 = \frac{g}{l} \quad (44)$$

$$\tilde{p} \rightarrow p; \tilde{t} \rightarrow t \quad (45)$$

$$\dot{\theta}_1 = \frac{p_1 - p_2 \cos(\theta_1 - \theta_2)}{1 + \sin^2(\theta_1 - \theta_2)} \quad (46)$$

$$\dot{\theta}_2 = \frac{2p_2 - p_1 \cos(\theta_1 - \theta_2)}{1 + \sin^2(\theta_1 - \theta_2)} \quad (47)$$

$$\dot{p}_1 = -2 \sin \theta_1 - C_1 + C_2 \quad (48)$$

$$\dot{p}_2 = -\sin \theta_2 + C_1 - C_2 \quad (49)$$

$$C_1 = \frac{p_1 p_2 \sin(\theta_1 - \theta_2)}{1 + \sin^2(\theta_1 - \theta_2)} \quad (50)$$

$$C_2 = \frac{[p_1^2 + 2p_2^2 - 2p_1 p_2 \cos(\theta_1 - \theta_2)] \sin(2(\theta_1 - \theta_2))}{2[1 + \sin^2(\theta_1 - \theta_2)]^2} \quad (51)$$

The class which can be given to integration routine is

```
class DoublePendulum{
    double om02;
public:
    DoublePendulum(double om02_) : om02(om02_){}
    void operator()(double x, function1D<double>& y, function1D<double>& dydx)
    {
        double cs = cos(y[0]-y[1]), ss = sin(y[0]-y[1]);
        double t = 1/(1+ss*ss);
        dydx[0] = (y[2]-y[3]*cs)*t;
        dydx[1] = (2*y[3]-y[2]*cs)*t;
        double c1 = y[2]*y[3]*ss*t;
        double c2 = (y[2]*y[2]+2*y[3]*y[3]-2*y[2]*y[3]*cs)*cs*ss*t*t;
        dydx[2] = -2*om02*sin(y[0])-c1+c2;
        dydx[3] = -om02*sin(y[1])+c1-c2;
    }
    double Energy(const function1D<double>& y)
    {
        double cs = cos(y[0]-y[1]), ss = sin(y[0]-y[1]);
        return 0.5*(y[2]*y[2]+2*y[3]*y[3]-2*y[2]*y[3]*cs)/(1+ss*ss)+om02*(3-2*cos(y[0])-cos(y[1]));
    }
    double p2(double E, double y0, double y1)
    {
        if (E<=0) {std::cerr<<"Error in starting conditions!"<<std::endl; return 0;}
        double E2 = (E-om02*(3-2*cos(y0)-cos(y1)))*(1+sqr(sin(y0-y1)));
        if (E2<0) {std::cerr<<"Error in starting conditions!"<<std::endl; return 0;}
        else return sqrt(E2);
    }
};
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Poincare map From Wikipedia

In mathematics of dynamical systems, a Poincare map or Poincare section, named after Henri Poincare, is the intersection of a trajectory which moves periodically (or quasi-periodically, or chaotically), in a space of at least three dimensions, with a transversal hypersurface of one fewer dimension. More precisely, one considers a trajectory with initial conditions on the hyperplane and observes the point at which this trajectory returns to the hyperplane. The Poincare section refers to the hyperplane, and the Poincare map refers to the map of points in the hyperplane induced by the intersections.

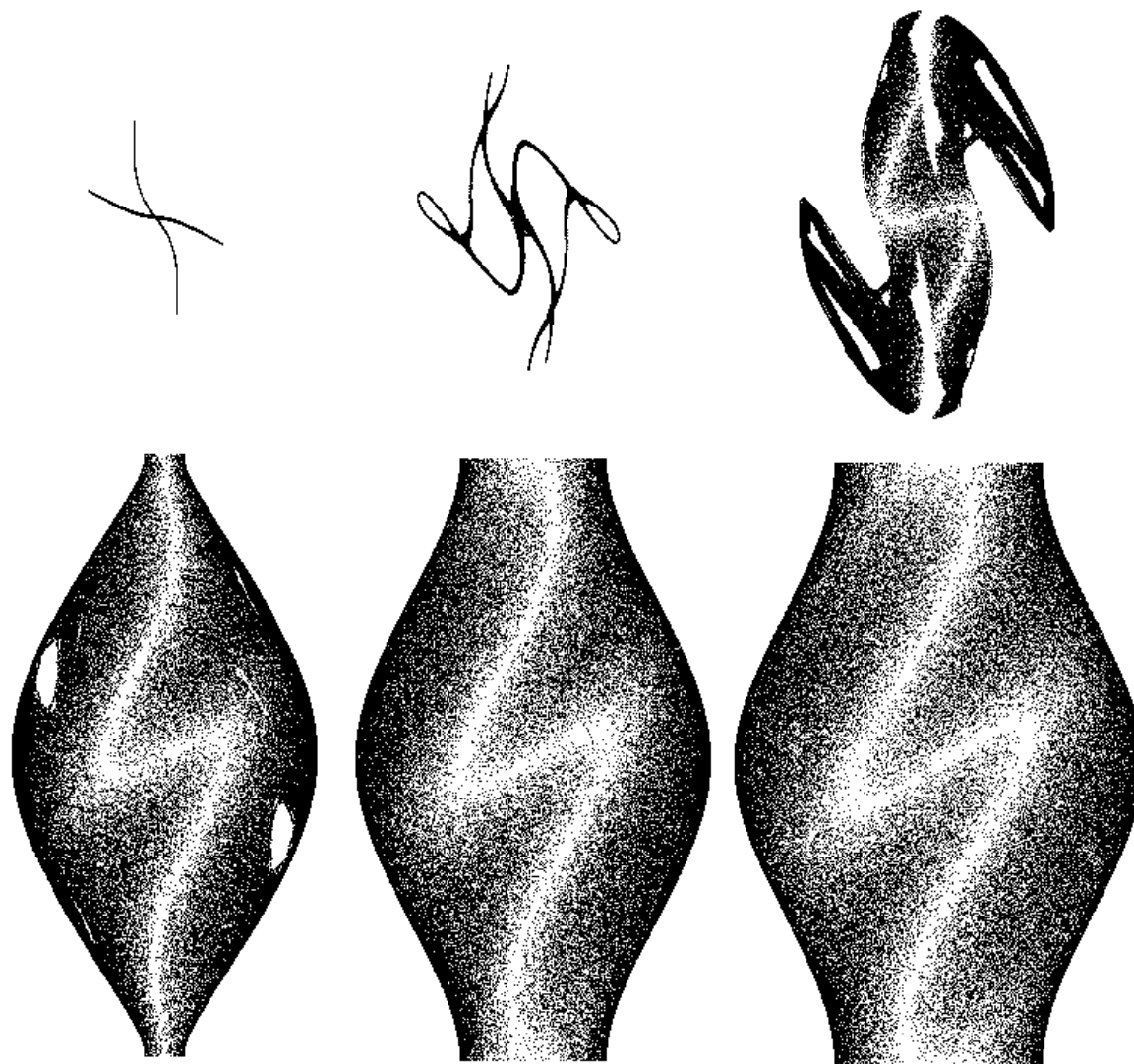


Figure 4: Poincare plots using θ_1 and θ_2 as variables. The point is plotted when $p_{\theta_1} = 0$.
Energies used to get the above plots are 5,10,15,20,25,30.

Homeworks

- 1 Simulate the motion of Earth in the solar system as a two body problem (taking into account only the Sun and Earth). Write the equation

$$m\ddot{\vec{r}} = -GmM \frac{\vec{r}}{r^3}$$

in dimensionless units or astronomical units (AU).

- 2 Simulate the three body problem and check how strong is the influence of Jupiter on motion of Earth.
- 3 Plot trajectories of Earth in case Jupiter's mass is 1000 times bigger than its actual mass.
- 4 Verify the existence of Kirkwood gaps. Simulate motion of Jupiter together with asteroids close to 2/1 Kirkwood gap with the following initial conditions

<i>Object</i>	<i>Radius(AU)</i>	<i>Velocity(AU/yr)</i>
<i>Jupiter</i>	5.2	2.755
<i>Asteroid1</i>	3.0	3.628
<i>Asteroid2</i>	3.276	3.471
<i>Asteroid3</i>	3.7	3.267

(52)

astronomical units (AU) are:

- length is measured in units of distance between Earth and Sun $\approx 1.5 \cdot 10^{11}$
- time can be measured in years