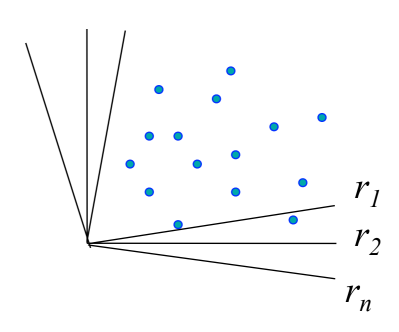


#### Molecular Simulations

Molecular dynamics: solve equations of motion  $r_1$   $r_2$   $r_n$ 

→ Monte Carlo: importance sampling



## Molecular Dynamics

- 4. Molecular Dynamics
  - 4.1.Introduction
  - 4.2.Basics
  - 4.3. Liouville formulation
  - 4.4. Multiple time steps

## 4. Molecular dynamics

4.2 Basics

#### "Fundamentals"

Theory:

$$F = m \frac{d^2r}{dt^2}$$

- Compute the forces on the particles
- Solve the equations of motion
- Sample after some # of timesteps

## 4. Molecular dynamics

4.3 Some practical details

# Molecular Dynamics

#### Initialization

- Total momentum should be zero (no external forces)
- Temperature rescaling to desired temperature
- Particles start on a lattice

#### **Force calculations**

- Periodic boundary conditions
- Order NxN algorithm,
- Order N: neighbor lists, linked cell
- Truncation and shift of the potential

#### Integrating the equations of motion

- Velocity Verlet
- Kinetic energy

## Molecular Dynamics

#### Algorithm 3 (A Simple Molecular Dynamics Program)

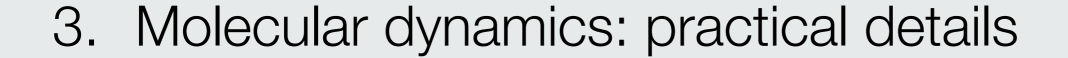
```
simple MD program
program md
                             initialization
call init
t = 0
                             MD loop
do while (t.lt.tmax)
                             determine the forces
   call force(f,en)
                             integrate equations of motion
   call integrate(f,en)
   t=t+delt
                             sample averages
   call sample
enddo
stop
end
```



3.3.1 Initialization

#### Algorithm 4 (Initialization of a Molecular Dynamics Program)

```
initialization of MD program
subroutine init
sumv=0
sumv2=0
do i=1, npart
  x(i)=lattice_pos(i)
                              place the particles on a lattice
                              give random velocities
  v(i) = (ranf() - 0.5)
                              velocity center of mass
  sumv=sumv+v(i)
                              kinetic energy
  sumv2=sumv2+v(i)**2
enddo
                              velocity center of mass
sumv=sumv/npart
sumv2=sumv2/npart
                              mean-squared velocity
                              scale factor of the velocities
fs=sqrt(3*temp/sumv2)
                              set desired kinetic energy and set
do i=1, npart
  v(i) = (v(i) - sumv) *fs
                              velocity center of mass to zero
                              position previous time step
  xm(i) = x(i) - v(i) *dt
enddo
return
end
```

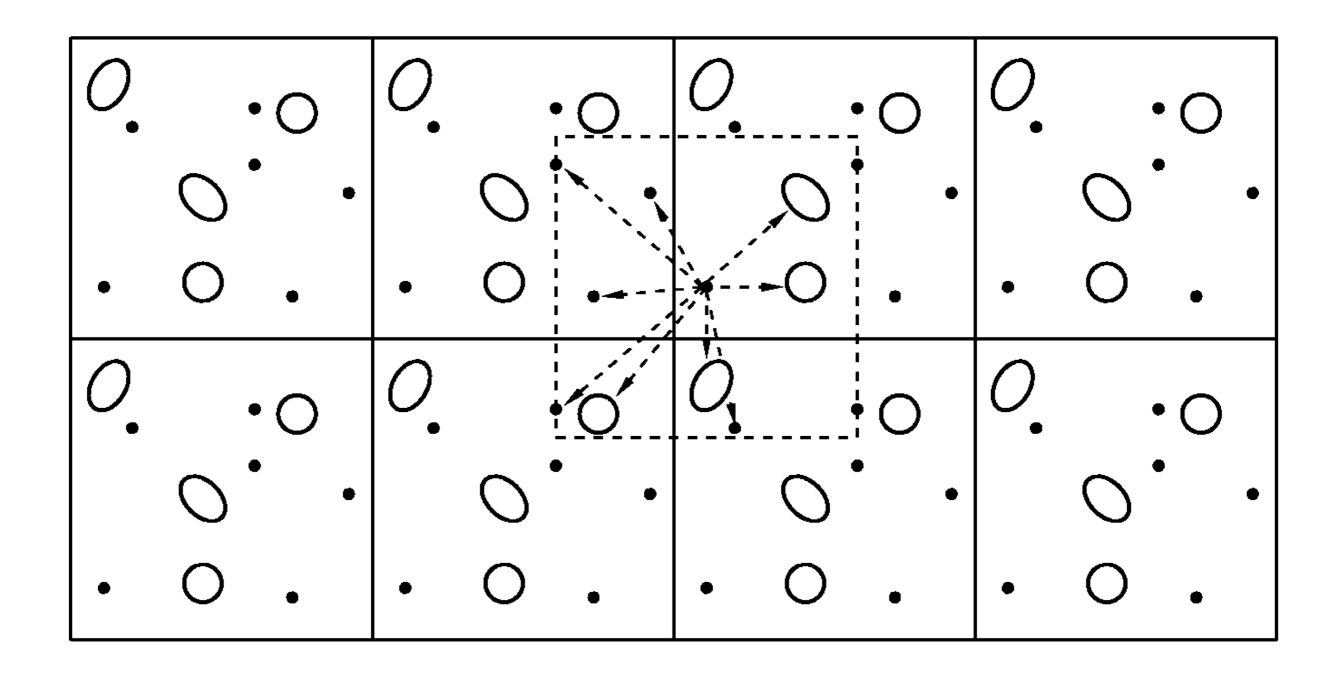


3.3.2 Force calculation

#### Algorithm 5 (Calculation of the Forces)

```
determine the force
subroutine force (f, en)
                                         and energy
en=0
do i=1, npart
   f(i) = 0
                                         set forces to zero
enddo
do i=1, npart-1
                                         loop over all pairs
  do j=i+1, npart
     xr=x(i)-x(j)
                                         periodic boundary conditions
      xr=xr-box*nint(xr/box)
      r2=xr**2
                                         test cutoff
      if (r2.lt.rc2) then
        r2i=1/r2
        r6i=r2i**3
        ff=48*r2i*r6i*(r6i-0.5)
                                         Lennard-Jones potential
                                         update force
        f(i) = f(i) + ff *xr
        f(j) = f(j) - ff *xr
        en=en+4*r6i*(r6i-1)-ecut update energy
      endif
  enddo
enddo
return
end
               Understanding Molecular Simulation
```

## Periodic boundary conditions



#### The Lennard-Jones potentials

The Lennard-Jones potential

$$U^{\sqcup}(r) = 4\varepsilon \left[ \left( \frac{\sigma}{r} \right)^{12} - \left( \frac{\sigma}{r} \right)^{6} \right]$$

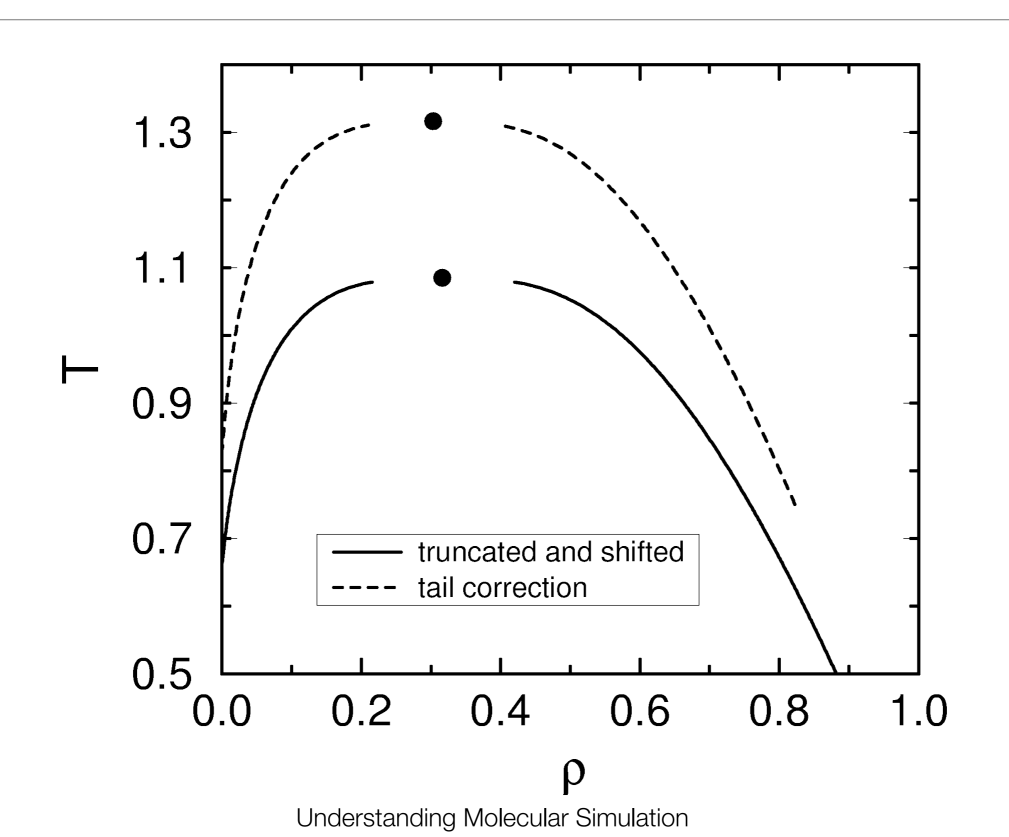
The truncated Lennard-Jones potential

$$U_{TR}^{\square}(r) = \begin{cases} U^{\square}(r) & r \leq r_{c} \\ 0 & r > r_{c} \end{cases}$$

The truncated and shifted Lennard-Jones potential

$$U_{TR-SH}^{LJ}(r) = \begin{cases} U^{LJ}(r) - U^{LJ}(r_c) & r \leq r_c \\ 0 & r > r_c \end{cases}$$

## The Lennard-Jones potentials

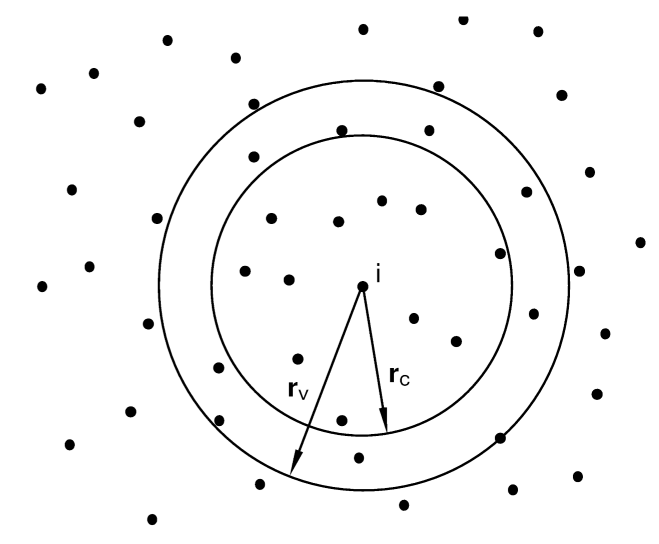


## Saving CPU-time

#### Cell list

# $r_{c}$

#### Verlet-list





3.3.3 Equations of motion

#### Algorithm 6 (Integrating the Equations of Motion)

```
integrate equations of motion
subroutine integrate(f,en)
sumv=0
sumv2=0
                                         MD loop
do i=1,npart
   xx=2*x(i)-xm(i)+delt**2*f(i)
                                         Verlet algorithm (4.2.3)
   vi=(xx-xm(i))/(2*delt)
                                         velocity (4.2.4)
                                         velocity center of mass
   sumv=sumv+vi
                                         total kinetic energy
   sumv2=sumv2+vi**2
                                         update positions previous time
   xm(i)=x(i)
                                         update positions current time
   x(i)=xx
enddo
                                         instantaneous temperature
temp=sumv2/(3*npart)
                                         total energy per particle
etot=(en+0.5*sumv2)/npart
return
end
```

#### Equations of motion

We can make a Taylor expansion for the positions:

$$r(t + \Delta t) = r(t) + \frac{dr(t)}{dt} \Delta t + \frac{d^2r(t)}{dt^2} \frac{\Delta t^2}{2!} + O(\Delta t^3)$$

The simplest form (Euler):

$$r(t + \Delta t) = r(t) + v(t)\Delta t + O(\Delta t^{2})$$
$$v(t + \Delta t) = v(t) + m\frac{df(t)}{dt}\Delta t$$

We can do better!

We can make a Taylor expansion for the positions:

$$r(t+\Delta t) = r(t) + \frac{dr(t)}{dt}\Delta t + \frac{d^2r(t)}{dt^2}\frac{\Delta t^2}{2!} + \frac{d^2r(t)}{dt^2}\frac{\Delta t^3}{3!} + O(\Delta t^4)$$

$$r(t - \Delta t) = r(t) - \frac{dr(t)}{dt} \Delta t + \frac{d^2r(t)}{dt^2} \frac{\Delta t^2}{2!} - \frac{d^2r(t)}{dt^2} \frac{\Delta t^3}{3!} + O(\Delta t^4)$$

When we add the two:

$$r(t + \Delta t) + r(t - \Delta t) = 2r(t) + \frac{d^2r(t)}{dt^2} \Delta t^2 + O(\Delta t^4)$$

Verlet algorithm

$$r(t + \Delta t) = 2r(t) - r(t - \Delta t) + f(t) \frac{\Delta t^2}{m} + O(\Delta t^4)$$

numerically not ideal

no need for velocities

Verlet algorithm:

$$r(t + \Delta t) = 2r(t) - r(t - \Delta t) + f(t) \frac{\Delta t^{2}}{m} + O(\Delta t^{4})$$

Velocity Verlet algorithm

$$r(t + \Delta t) = r(t) + v(t)\Delta t + f(t)\frac{\Delta t^{2}}{2m} + O(\Delta t^{4})$$

$$v(t+\Delta t) = v(t) + \frac{\Delta t}{2m} \left[ f(t+\Delta t) + f(t) \right]$$

to see the equivalence:

$$r(t+2\Delta t) = r(t+\Delta t) + v(t+\Delta t)\Delta t + f(t+\Delta t)\frac{\Delta t^{2}}{2m}$$
$$r(t) = r(t+\Delta t) - v(t)\Delta t - f(t)\frac{\Delta t^{2}}{2m}$$

adding the two

$$r(t+2\Delta t) = 2r(t+\Delta t) - r(t) + \left[v(t+\Delta t) - v(t)\right] \Delta t + \left[f(t+\Delta t) - f(t)\right] \frac{\Delta t^2}{2m}$$
with 
$$v(t+\Delta t) = v(t) + \frac{\Delta t}{2m} \left[f(t+\Delta t) + f(t)\right]$$

$$r(t+2\Delta t) = 2r(t+\Delta t) - r(t) + f(t+\Delta t) \frac{\Delta t^2}{m}$$

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## Lyaponov instability

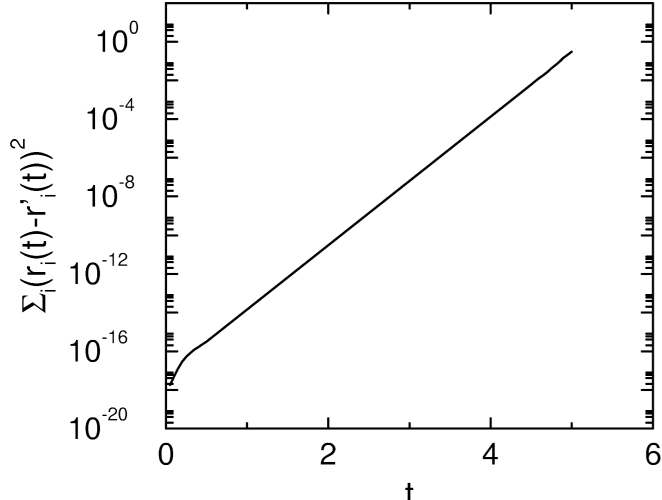
MD: reference trajectory

with initial condition:

$$(r_1(O), \dots, r_N(O), p_1(O), \dots, p_N(O))$$

 $\varepsilon = 10^{-10}$ 

MD: compare:  $(r_1(0), \dots, r_N(0), p_1(0), \dots, p_i(0) + \varepsilon, p_i(0) - \varepsilon, \dots, p_N(0))$ 



t
Understanding Molecular Simulation

## 4. Molecular dynamics:

4.4 Liouville Formulation

#### Liouville formulation

# the dot above, f, implies time derivative

Let us consider a function that *f* which depends on the positions and momenta of the particles:

$$f(p^N,r^N)$$

We can "solve" how f depends on time:

$$\dot{f} = \left(\frac{\partial f}{\partial r}\right)\dot{r} + \left(\frac{\partial f}{\partial \rho}\right)\dot{p}$$

Define the Liouville operator:

$$iL \equiv \dot{r} \left( \frac{\partial}{\partial r} \right) + \dot{p} \left( \frac{\partial}{\partial p} \right)$$

the time dependence follows from:

$$\frac{df}{dt} = iLf$$

with solution:

**beware:** the solution is equally useless as the differential equation

$$f = e^{iLt} f(0)$$

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In an ideal world it would be less useless:

$$iL \equiv \dot{r} \left( \frac{\partial}{\partial r} \right) + \dot{p} \left( \frac{\partial}{\partial p} \right)$$

Let us look at half the equation which has as solution:

$$iL_r \equiv \left(\frac{\partial}{\partial r}\right)\dot{r}$$

$$f = e^{iL_r t} f(0)$$

Taylor expansion:

$$e^{x} = 1 + x + \frac{x^{2}}{2!} + \frac{x^{3}}{3!} + \cdots$$

$$e^{iL_{r}t}f(0) = \left[1 + iL_{r}t + \frac{1}{2}(iL_{r}t)^{2} + \frac{1}{3!}(iL_{r}t)^{3} + \dots\right]f(0)$$

$$e^{iL_{r}t}f(0) = \left[1 + \dot{r}(0)t\left(\frac{\partial}{\partial r}\right) + \frac{1}{2}(\dot{r}(0)t)^{2}\left(\frac{\partial}{\partial r}\right)^{2} + \dots\right]f$$
 the operator iL<sub>r</sub> gives a shift of the positions

$$f(0+\dot{r}(0)t) = f(0)+\dot{r}(0)t\left(\frac{\partial f(0)}{\partial r}\right) + \frac{1}{2}(\dot{r}(0)t)^2\left(\frac{\partial f(0)}{\partial r}\right)^2 + \cdots$$

Hence:

$$e^{iL_rt}f(O)=f(O+\dot{r}(O)t)$$

The operation  $iL_r$  gives a shift of the positions

$$iL \equiv \dot{r} \left( \frac{\partial}{\partial r} \right) + \dot{p} \left( \frac{\partial}{\partial p} \right)$$

Similarly for the operator  $iL_p$  which has as solution:

$$iL_{p} \equiv \left(\frac{\partial}{\partial p}\right) \dot{p}$$

$$f = e^{iL_p t} f(0)$$

Taylor expansion:

$$e^{iL_{p}t}f(0) = \left[1 + iL_{p}t + \frac{1}{2}(iL_{p}t)^{2} + \frac{1}{3!}(iL_{p}t)^{3} + \dots\right]f(0)$$

$$e^{iL_{p}t}f(0) = \left[1 + \dot{p}(0)t\left(\frac{\partial}{\partial p}\right) + \frac{1}{2}(\dot{p}(0)t)^{2}\left(\frac{\partial}{\partial p}\right)^{2} + \dots\right]f(0)$$
the operator iL<sub>p</sub> gives a shift of the momenta

$$f(O+\dot{p}(O)t) = f(O)+\dot{p}(O)t\left(\frac{\partial f(O)}{\partial p}\right) + \frac{1}{2}(\dot{p}(O)t)^2\left(\frac{\partial f(O)}{\partial p}\right)^2 + \cdots$$

Hence

$$e^{iL_{p}t}f(O)=f(O+\dot{p}(O)t)$$

The operation  $iL_r$  gives a shift of the positions:

$$e^{iL_{r}t}f(O,O)=f(O,O+\dot{r}(O)t)$$

... and the operator  $iL_p$  a shift of the momenta:

$$e^{iL_{p}t}f(O,O)=f(O+\dot{p}(O)t,O)$$

This would have been useful if the operators would commute

$$e^{iLt}f(0,0) = e^{(iL_r + iL_p)t}f(0,0) \neq e^{iL_r t}e^{iL_p t}f(0,0)$$

Trotter expansion:

we have the non-commuting operators A and B:

then the following expansion holds:

$$e^{A+B} \neq e^A e^B$$

$$e^{A+B} = \lim_{P \to \infty} \left( e^{\frac{A}{2P}} e^{\frac{B}{P}} e^{\frac{A}{2P}} \right)^{P}$$

$$e^{iL_{r}t}f(0,0) = f(0,0+\dot{r}(0)t)$$
$$e^{iL_{p}t}f(0,0) = f(0+\dot{p}(0)t,0)$$

We can apply the Trotter expansion:

$$e^{A+B} = \lim_{P \to \infty} \left( e^{\frac{A}{2P}} e^{\frac{B}{P}} e^{\frac{A}{2P}} \right)^{P}$$

$$\Delta t = \frac{t}{P} \qquad \frac{iL_r t}{P} = iL_r \Delta t \qquad \frac{iL_p t}{2P} = iL_p \frac{\Delta t}{2}$$

These give as operations:

$$e^{iL_r\Delta t}f(p(t),r(t))=f(p(t),r(t)+\dot{r}(t)\Delta t)$$

gives us a shift of the position:

$$r(t + \Delta t) \rightarrow r(t) + \dot{r}(t) \Delta t$$

$$e^{iL_{p}\Delta t/2}f(p(t),r(t)) = f\left(p(t)+\dot{p}(t)\frac{\Delta t}{2},r(t)\right)$$

gives us a shift of the momenta:

$$p(t + \Delta t) \rightarrow p(t) + \dot{p}(t) \frac{\Delta t}{2}$$

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$$iL_r\Delta t$$
  $r(t+\Delta t) \rightarrow r(t) + \dot{r}(t)\Delta t$ 

$$iL_{\rho}\frac{\Delta t}{2}$$
  $p\left(t+\frac{\Delta t}{2}\right) \rightarrow p(t)+\dot{p}(t)\frac{\Delta t}{2}$ 

We can apply the Trotter expansion to integrate M time steps:  $t=M \times \Delta t$ 

$$f(t) = e^{iLt}f(0) = \left(e^{iL_{\rho}\frac{\Delta t}{2}}e^{iL_{r}\Delta t}e^{iL_{\rho}\frac{\Delta t}{2}}\right)^{M}f(0)$$

These give as operations:

$$e^{iL_{p}\frac{\Delta t}{2}} \qquad \rho\left(\frac{\Delta t}{2}\right) \to \rho(0) + \dot{\rho}(0)\frac{\Delta t}{2}$$

$$e^{iL_{p}\Delta t} \qquad r(\Delta t) \to r(0) + \dot{r}\left(\frac{\Delta t}{2}\right)\Delta t$$

$$e^{iL_{p}\frac{\Delta t}{2}} \qquad \rho(\Delta t) \to \rho\left(\frac{\Delta t}{2}\right) + \dot{\rho}(\Delta t)\frac{\Delta t}{2}$$
The gives after one step

which gives after one step

$$p(0) \to p(0) + \left[ f(0) + f(\Delta t) \right] \frac{\Delta t}{2}$$

$$r(0) \to r(0) + \dot{r} \left( \frac{\Delta t}{2} \right) \Delta t = r(0) + v(0) \Delta t + f(0) \frac{\Delta t^2}{2m}$$

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which gives after one step

$$r(0) \rightarrow r(0) + \dot{r}\left(\frac{\Delta t}{2}\right) \Delta t = r(0) + v(0) \Delta t + f(0) \frac{\Delta t^2}{2m}$$

$$p(O) \rightarrow p(O) + [f(O) + f(\Delta t)] \frac{\Delta t}{2}$$

Velocity Verlet algorithm

$$r(t + \Delta t) = r(t) + v(t)\Delta t + f(t)\frac{\Delta t^{2}}{2m}$$

$$v(t + \Delta t) = v(t) + \frac{\Delta t}{2m} \left[ f(t + \Delta t) + f(t) \right]$$

Velocity Verlet algorithm:

$$e^{iL_{\rho}\frac{\Delta t}{2}}e^{iL_{r}\Delta t}e^{iL_{\rho}\frac{\Delta t}{2}}$$

$$iL_r \Delta t : r(t + \Delta t) \rightarrow r(t) + v(t) \Delta t$$
  
 $iL_p \frac{\Delta t}{2} : v\left(t + \frac{\Delta t}{2}\right) \rightarrow v(t) + f(t) \frac{\Delta t}{2}$ 

Call force (fx)

Do while (t<tmax)

$$iL_{p}\frac{\Delta t}{2}: v\left(t+\frac{\Delta t}{2}\right) \rightarrow v(t)+f(t)\frac{\Delta t}{2}$$

vx=vx+delt\*fx/2

$$iL_r\Delta t: r(t+\Delta t) \rightarrow r(t)+v(t)\Delta t$$

x=x+delt\*vx

Call force (fx)

$$iL_{p} \frac{\Delta t}{2}$$
:  $v(t + \Delta t) \rightarrow v\left(t + \frac{\Delta t}{2}\right) + f(t + \Delta t) \frac{\Delta t}{2}$ 

vx=vx+delt\*fx/2

enddo

#### Liouville formulation

Velocity Verlet algorithm

$$r(t + \Delta t) = r(t) + v(t)\Delta t + f(t)\frac{\Delta t^{2}}{2m}$$
$$v(t + \Delta t) = v(t) + \frac{\Delta t}{2m} \left[ f(t + \Delta t) + f(t) \right]$$

**Transformations:** 

$$iL_{p} \Delta t/2: \quad r(t) \rightarrow r(t) \qquad \qquad iL_{r} \Delta t: \quad r(t + \Delta t) \rightarrow r(t) + v(t) \Delta t$$

$$v(t) \rightarrow v(t) + f(t) \Delta t/2m \qquad \qquad v(t) \rightarrow v(t)$$

$$J_{p} = Det \begin{vmatrix} 1 & 0 \\ \left(\frac{\partial f}{\partial r}\right) \frac{\Delta t}{2m} & 1 \end{vmatrix} = 1 \qquad \qquad J_{r} = Det \begin{vmatrix} 1 & \Delta t \\ 0 & 1 \end{vmatrix} = 1$$

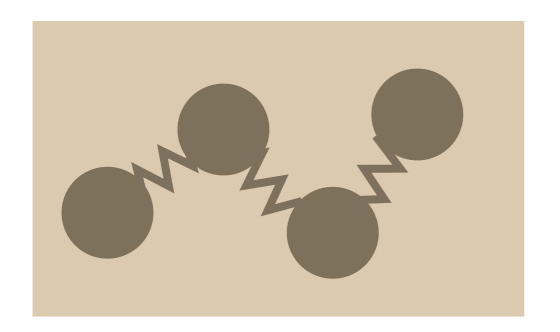
Three subsequent coordinate transformations in either r or r of which the Jacobian is one: Area preserving

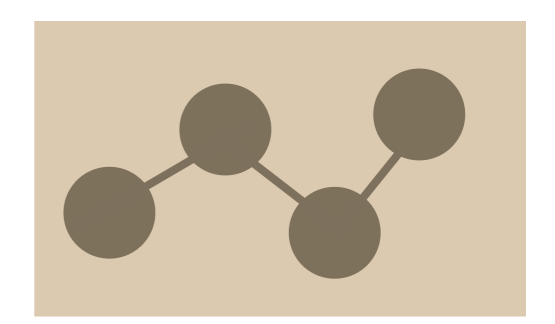
## 4. Molecular dynamics:

4.5 Multiple time steps

## Multiple time steps

What to do with "stiff" potentials?





- Fixed bond-length: constraints (Shake)
- Very small time step

$$iL_r \Delta t$$
:  $r(t + \Delta t) \rightarrow r(t) + v(t) \Delta t$   
 $iL_p \frac{\Delta t}{2}$ :  $v(t + \frac{\Delta t}{2}) \rightarrow v(t) + f(t) \frac{\Delta t}{2}$ 

We can split the force is the stiff part and the more slowly changing rest of the forces:  $f(t) = f_{Short}(t) + f_{Long}(t)$ 

This allows us to split the Liouville operator:

$$iLt = iL_rt + iL_{pShort}t + iL_{pLong}$$

The conventional Trotter expansion:

$$iLt = \left[iL_{pLong} \Delta t/2 \left[iL_{r} + iL_{pShort}\right] \Delta t \ iL_{pLong} \Delta t/2\right]^{M}$$

Now we can make another Trotter expansion:  $\delta t = \Delta t/m$ 

$$\left[iL_r + iL_{pShort}\right] \Delta t = \left[iL_{pShort} \delta t/2 iL_r \delta t iL_{pShort} \delta t/2\right]^m$$

The algorithm to solve the equations of motion

$$f(t) = f_{Short}(t) + f_{Long}(t)$$

$$iLt = \left[iL_{pLong} \Delta t/2 \left[iL_{r} + iL_{pShort}\right] \Delta t \ iL_{pLong} \Delta t/2\right]^{M}$$

$$\left[iL_{r} + iL_{pShort}\right] \Delta t = \left[iL_{pShort} \delta t/2 \ iL_{pShort} \delta t/2\right]^{m}$$

We now have 3 transformations:

$$iL_{pLong} \frac{\Delta t}{2}$$
:  $v\left(t + \frac{\Delta t}{2}\right) \rightarrow v(t) + f_{Long}(t) \frac{\Delta t}{2}$   
 $iL_{pShort} \frac{\delta t}{2}$ :  $v\left(t + \frac{\delta t}{2}\right) \rightarrow v(t) + f_{Short}(t) \frac{\delta t}{2}$   
 $iL_{r}\delta t$ :  $r(t + \delta t) \rightarrow r(t) + v(t)\delta t$ 

The steps are first iL<sub>pLong</sub> then m times iL<sub>pShort</sub>/iL<sub>r</sub> followed by iL<sub>pLong</sub> again

$$iL_{plong} \frac{\Delta t}{2} : v\left(t + \frac{\Delta t}{2}\right) \rightarrow v(t) + f_{long}(t) \frac{\Delta t}{2}$$

$$Call \ \, \text{force}(\mathbf{fx\_long}, \mathbf{f\_short})$$

$$v\mathbf{x} = v\mathbf{x} + \text{delt*} \mathbf{fx\_long}/2$$

$$Do \ \, \text{ddt=1,n}$$

$$iL_{pShort} \frac{\delta t}{2} : v\left(t + \frac{\delta t}{2}\right) \rightarrow v(t) + f_{Short}(t) \frac{\delta t}{2}$$

$$v\mathbf{x} = v\mathbf{x} + \text{ddelt*} \mathbf{fx\_short}/2$$

$$iL_{r} \delta t : r(t + \delta t) \rightarrow r(t) + v(t) \delta t$$

$$\mathbf{x} = \mathbf{x} + \text{ddelt*} \mathbf{vx}$$

$$Call \ \, \text{force\_short}(\mathbf{fx\_short})$$

$$iL_{pShort} \frac{\delta t}{2} : v\left(t + \frac{\delta t}{2}\right) \rightarrow v(t) + f_{Short}(t) \frac{\delta t}{2}$$

$$v\mathbf{x} = v\mathbf{x} + \text{ddelt*} \mathbf{fx\_short}/2$$

enddo

#### Algorithm 29 (Multiple Time Step)

```
Multiple time step, f_long is
subroutine
                                       the long-range part and f_short
    multi(f_long, f_short)
                                       the short-range part of the force
                                       velocity Verlet with time step \Delta t
vx=vx+0.5*delt*flong
                                       loop for the small time step
do it=1, n
                                       velocity Verlet with timestep \Delta t/n
   vx=vx+0.5*(delt/n)*f_short
   x=x+(delt/n) 2*vx
                                       short-range forces
   call force_short(f_short)
   vx=vx+0.5*(delt/n)*f_short
enddo
                                       all forces
call force_all(f_long,f_short)
vx=vx+0.5*delt*f_long
return
end
```