- In MD simulations (and actually many other applications) one of the central operations is the calculation of distances between atoms.
  - In MD this is needed in the energy and force calculation.
- Trivial calculation of distances between atoms:

```
do i=1,N
    do j=1,N
        if (i==j) cycle
        dx=x(j)-x(i);
        dy=y(j)-y(i);
        dz=z(j)-z(i);
        rsq=dx*dx+dy*dy+dz*dz
        r=sqrt(rsq)
        enddo
enddo
```

- This algorithm is  $O(N^2)$ , i.e. very slow when  $N \rightarrow \infty$ .
- But in practice we know the atoms move < 0.2 Å/time step. So a large fraction of the neighbours remain the same during one time step, and it seems wasteful to recalculate which they are every single time.

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# Constructing a neighbour list

- Solution: Verlet<sup>1</sup> neighbour list:
  - Make a list which contains for each atom *i* the indices of all atoms *j* which are closer to *i* than a given distance  $r_{\rm m}$ .  $r_{\rm m} > r_{cut}$ , the cutoff distance of the potential
  - The list is updated only every  $N_{\rm m}$  time steps.
  - $r_{\rm m}$  and  $N_{\rm m}$  are chosen such that

$$r_{\rm m} - r_{\rm cut} > N_{\rm m} \bar{v} \Delta t$$

where  $\bar{v}$  is a typical atom velocity and  $\Delta t$  the time step



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<sup>1.</sup> Loup Verlet, Phys. Rev. 159 (1967) 98.

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- An even better way to choose when to update the interval: after the neighbour list has been updated, keep a list of the maximum displacement of all atoms:
  - Make a separate table dxnei(i)
  - When you move atoms, also calculate dxnei(i)=dxnei(i)+dx
  - Calculate the two maximal displacements of all atoms:

```
drneimax=0.0; drneimax2=0.0
do i=1,N
    drnei=sqrt(dxnei(i)*dxnei(i)+dynei(i)*dynei(i)+dznei(i)*dznei(i))
    if (drnei > drneimax) then
        drneimax2=drneimax
        drneimax=drnei
    else
        if (drnei > drneimax2) then
            drneimax2=drnei
        endif
    endif
endif
enddo
```

- Now, when  $(drneimax+drneimax2) > r_m r_{cut}$  the neighbour list has to be updated.
- When the update is done, do dxnei(i)=0.0
- This alternative has two major advantages: the simulation does not screw up if one atom suddenly starts to move much faster than the average, and if the system cools down, the neighbour list update interval keeps increasing.

```
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```

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# **Constructing a neighbour list**

• In practice the neighbour list can look e.g. like the following:



- Here **NNei** is the number of neighbours of atom i.
- $j_1$ ,  $j_2$ , ... are the indices of neighbouring atoms (different for different atoms).
- So, if we would have a 64 atom system, where every atom has 4 neighbours, the neighbour list could look like this:

	neiał	hours	s of at	om 1		neiał	hours	ofat	om 2		neiał	hours	ofat	om 64
4	2	3	63	64	4	1	3	4	5	4	1	61	62	63

· A practical implementation of creating the list:

```
nlistbeg=1
do i=1.N
   nnei=0
                                                           Periodic boundaries
    do j=1,N
                                                           omitted for brevity.
        if (i==j) cycle
        dx=x(j)-x(i)
        dy=y(j)-y(i)
        dz=z(j)-z(i)
        rsq=dx*dx+dy*dy+dz*dz
        if (rsq <= rskincutsq) then</pre>
           nnei=nnei+1
            nlist(nlistbeg+nnei)=j
        endif
    enddo
   nlist(nlistbeg)=nnei
                                             ! Write in number of i's neighbours into list
   nlistbeg=nlistbeg+nnei+1
                                             ! Set starting position for next atom
enddo
```

- With the neighbour list, we can achieve a savings of a factor *N*<sub>m</sub> in calculating the distances to neighbours.
- But even using the neighbour list, our algorithm is still  $O(N^2)$ .

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# **Constructing a neighbour list**

- · Remedy: linked list / cell method
- Using a linked list and cellular division of the simulation cell, we can make the algorithm truly *O*(*N*):
  - Let's divide the MD cell into smaller subcells:  $M \times M \times M$  cells
  - The size of one subcell / is chosen so that

$$l = \frac{L}{M} > r_{\rm m},$$

where L = the size of the MD cell, and  $r_{\rm m}$  is as above.

• Now when we look for neighbours of atom *i* we only have to look through the subcell where *i* is, and its neighbouring subcells, but not the whole simulation cell. For instance if atom *i* is in cell 13:

The average number of atoms in a subcell is  $N_c = N/M^3$ .

 $\Rightarrow$  We have to go through 27*NN*<sub>c</sub> atom pairs instead of *N*(*N*-1).

• For some interaction potentials (symmetric ij pairs) it is actually enough to calculate every second neighbour pair (e.g. i > j) whence the number of pairs is further reduced by a factor of 2.

21	22	23	24	25
16	17	18	19	20
11	12	13	14	15
6	7	8	9	10
1	2	3	4	5

- · A practical implementation:
  - array HEAD:
    - size =  $M^3$
    - contains pointers to the table LIST
    - tells where the neighbours in subcell *m* start
  - array LIST:
    - size = *N*
    - element *j* tells where the next atom index of atoms in this cell is
- So the example below means that subcell 2 contains atoms 10, 9, 6, 4, and 3
- This representation is indeed enough to give all the atoms in all cells.
- A two dimensional array would of course also work, but would require much more memory, or dynamic allocation, both of which are less efficient.

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# Constructing a neighbour list

#### • Building the list:

assume a cubic case:				1							
• MD cell size = size(3)	HEAD	8	3								
• size of subcell =size()/M		L		' \							
MD cell centered on origin				. 🚽							
5			2	3	4	5	6	7	8	9	10
	LIST	0	1	0	3	2	4	5	7	6	9
do i=1.N											
head(i) = 0											
enddo											
do i=1,N											
icell = 1 + int((x(i)+size(1)/2)/size(1)*N	4) &										
int((y(i)+size(2)/2)/size(2)*1	4) * M 8	2									
int((z(i)+size(2)/2)/size(3)*1	4) * M '	* M									
list(i) = head(icell)											
head(icell) = i											
enddo											

- So the list LIST is filled in reverse order to the picture above.
- The above algorithm requires periodic boundaries. If the boundaries are open, an atom may get outside the cell borders, and the *icell* may point to the wrong cell.



• To account for possibly open boundaries properly things get a bit trickier:

```
• MD Cell size size (3)
   · MD cell centered on origin
   • Number of cells in different dimensions {\tt Mx}\,,~{\tt My}\,,~{\tt Mz}
   • Cell range 0 - Mx-1 and same in y and z
do i=1,N
    dx=x(i)+size(1)/2
     ! Check that we are really inside boundaries
    if (periodic(1) == 1 .and. dx < 0.0) dx=dx+size(1)
if (periodic(1) == 1 .and. dx > size(1)) dx=dx-size(1)
    ix=int((dx/size(1))*Mx)
    ! If not periodic, let border cells continue to infinity
    if (periodic(1) == 0) then
         if (ix < 0) ix=0
         if (ix \ge Mx) ix=Mx-1
    endif
     (and same thing for y and z)
    icell=(iz*My+iy)*Mx+ix
    list(i)=head(icell)
    head(icell)=i
enddo
```

21 22 23 24 25 16 17 18 19 20 11 12 13 14 15 6 7 8 9 10 1 2 3 4 5

· So the subcells at open boundaries continue out to infinity:

```
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```

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## **Constructing a neighbour list**

- Usually the linked list (LIST, HEAD) is used to generate a Verlet list
  - Decoding a linked list into a Verlet-list, as pseudocode:
  - Cell size size (3)
  - Number of cells Mx, My, Mz

```
do i=1,N
    do (Loop over 27 neighbouring cells: inx iny inz)
        icell=(inz*My+iny)*Mx+inx
        ! Get first atom in cell
        j=head(icell)
        do
            if (j==0) exit ! exit from innermost loop
            (get distance r between atoms i and j)
            if (r <= rneicut) then
               (accept neighbour)
            endif
            j=list(j)
        enddo
enddo
enddo</pre>
```

#### MD code mdmorse

- A simplified MD code mdmorse has been written for this course:
  - mdmorse simulates atom motion in a variety of metals (but only one metal at a time) with a simple Morse pair potential model.

$$V(r) = D[e^{-2\alpha(r-r_0)} - 2e^{-\alpha(r-r_0)}]$$

- The code has a Verlet neighbour list (but not a linked list) and the equations of motion are solved with the velocity Verlet method.
- The code is given in Fortran90 and C.
- The code can be downloaded from the course web page. (I'll email the exact location later.)
  - The code has the input parameter and output routines included.
  - Physically interesting subroutines have been removed from the code, so it does not work.
  - During the next few exercise sessions, you get the task of writing the missing subroutines.
  - Solutions will be provided and explained during the exercise sessions.
  - · You may either use your own or the provided solutions afterwards.

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## Structure of the mdmorse code

• Program files:

main.f90	Main program
inout.f90	Miscellaneous input and output stuff
modules.f90	Global variables
physical.f90*	Calculating T and E, and random number generators
neighbourlist.f90*	Getting the neighbour list
solve.f90 <sup>*</sup>	Solving the equations of motion
forces.f90*	Calculating the forces
Makefile	Makefile

- Files marked with \* contain the subroutines which will be filled up during the exercises
- C version: \*.c  $\rightarrow$  \*.f90 modules.f90  $\rightarrow$  global.h
- · Compiling the code:

#### make

- This has been tested to work at least on Linux systems with a GNU compilers (gfortran and gcc).
- You may have to change the compiler command in Makefile.

## Structure of the mdmorse code

• Input files (file names are hardcoded):

mdmorse.in	Miscellaneous parameters
atoms.in	Atom coordinates in XYZ format

• Running the program:

./mdmorse (or if you don't want to disturb other users nice ./mdmorse)

- Should be done in the same directory where the input files are.
- Output files:

• Note also that during the program running, the code writes out a large number of atom coordinates to a file <u>atoms.out</u>, which may grow very large.

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## Structure of the mdmorse code

#### • Input file mdmorse.in

Sample input file for mdmorse md program File format: \$identifier, then value. Rest is arbitrary comments Lines which do not begin with "\$" are all ignored

Identifier	Value	Comment
SinitialT	600 0	Initial temperature
yiniciali	000.0	initial temperature
\$desiredT	0.0	Variables for temperature control
Şbtctau	300.0	If btctau=0 no effect
\$bpctau	0.0	Variables for pressure control
\$bpcbeta	7.0e-4	If bpctau=0 no effect
\$desiredP	0.0	
\$mass	63.546	For Cu
\$xsize	18.126900793	Box size in each dimension
\$ysize	18.126900793	
\$zsize	18.126900793	
\$periodicx	1	1=periodic, 0=open
\$periodicy	1	
\$periodicz	1	
\$morseDe	0.3429	Morse potential parameters
\$morsealpha	1.3588	These values are for Cu
\$morseRe	2.866	
\$rpotcut	5.0	Potential cutoff
\$rskincut	6.0	Neighbour list cutoff
\$nupdate	5	Number of steps between n-list updates
\$nmovieoutput	100	Interval between atom movie output
\$deltat	2.0	Time step in simulation in fs
\$tmax	10000.0	Total simulation time
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#### Structure of the mdmorse code

• Input file atoms.in

• The file is a normal XYZ atom coordinate file:

500											
FCC	cell	made	by	makeFCC	with	a=	3.615	n=	5	5	5
Cu	- 8	8.133	75	-8.13	3375		-8.13	375			
Cu	- 6	6.3262	25	-6.32	2625		-8.13	375			

... and so forth the remaining 498 atom coordinates....

Cu	6.32625	8.13375	8.13375
Cu	8.13375	6.32625	8.13375

• Note that the cell is centered on the origin.

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## Structure of the mdmorse code

• Standard output (for the working code; F90 version):

```
----- mdmorse06 1.0F ---
Read in parameter $initialT
                                       value 300.000
Read in parameter $desiredT
                                       value 300.000
                                       value 300.000
Read in parameter $btctau
Read in parameter $bpctau
                                       value 500.000
Read in parameter $bpcbeta
                                       value 0.700000E-03
Read in parameter $desiredP
                                      value 0.00000
                                      value 63.5460
Read in parameter $mass
Read in parameter $xsize
                                      value 18.1269
Read in parameter $ysize
                                       value 18.1269
Read in parameter $zsize
                                      value 18.1269
Read in parameter Speriodicx
                                      value 1.00000
Read in parameter $periodicy
                                      value 1.00000
Read in parameter $periodicz
                                       value 1.00000
Read in parameter $morseD
                                      value 0.342900
                                      value 1.35880
value 2.86600
Read in parameter $morsealpha
Read in parameter $morser0
Read in parameter $rpotcut
                                      value 5.00000
Read in parameter $rskincut
                                      value 6.00000
                                      value 5.00000
Read in parameter $nupdate
Read in parameter $nmovieoutput
                                      value 100.000
Read in parameter $deltat
                                       value 2.00000
Read in parameter $tmax
                                       value 10000 0
                                       value 0.873440E+07
Read in parameter $seed
Using periodics (1=on, 0=off) 1 1 1
Morse potential parameters: D alpha r0 0.342900 1.358800 2.866000
Movie output selected every100 stepsDoing Berendsen temperature control with tau T300.000Doing Berendsen pressure control with tau beta500.000
Reading in 500 atoms described as FCC Cu; boxsize 18.1000 18
Initial atom temperature is 605.224802743929
Neighbour list update found 78.000 neighbours per atom
ec 2.000 605.225 0.07738 -3.03989 -2.96251 164.34551
                                                                               18 1000
Outputting atom movie at t = 2.000
ec 4.000 594.069 0.07538 -3.03868 -2.96330 163.82195
       4.000
          4.000 18.132452 18.132452 18.132452 5961.69346 163.82195
                                                                                  1.00015
bpc
                   574.307
                                0.07233
                                            -3.03638
                                                          -2.96405
                                                                      163.49694
         6.000
ec
. . .
```

#### Structure of the mdmorse code

· And so on. Here most things are self-explanatory.

• The "ec" and "bpc" lines contain the physically most interesting stuff in the following format:

	time(fs)	Т (К)	$E_{\mathrm{kin}}$ /at.	$E_{\mathrm{pot}}$ /at.	$E_{\rm tot}$ /at.	P(kbar)	(energies in eV)
ec	4.000	594.06	9 0.0753	8 -3.038	68 -2.96330	163.82195	
	time(fs)	$b_x(\text{\AA})$	$b_y(\text{\AA})$	$b_z(\text{\AA})$	$V(Å^3)$	P(kbar)	$\mu_{Berendsen}$
bpc	4.000	18.132452	18.132452	18.132452	5961.69346	163.82195	1.00015

• Output file atoms.out

• This file is in the XYZ format, but with the exception that column 5 contains the atom potential energy:

500							
mdmo	rse atom out	put at time	2.000	fs boxsize	18.1269	18.1269	18.1269
Cu	-9.053407	-9.061041	-9.048299	-3.085270			
Cu	-7.236810	-7.239921	-9.048988	-3.033905			
Cu	-7.241191	-9.049845	-7.246436	-3.035222			
Cu	-9.038484	-7.238137	-7.241429	-3.031141			
•							
•							
•							

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#### Structure of the mdmorse code

- Testing the incomplete code:
  - Even though the code is not complete, it should compile and run in the intermediate stages.
  - The output should look something like:

500 atoms described as FCC Cu; boxsize 18.1000 Reading in 18.1000 Neighbour list update found 0.26928E+06 neighbours per atom ec 2.000 0.000 0.00000 0.00000 0.00000 0.00000 Outputting atom movie at t = 2.000 4.000 0.000 0.00000 0.00000 0.00000 0.00000 ec

- I.e. the number of neighbours is nonsense, and the temperature is 0.
- When you start doing the exercises, this should change and interesting things will start to happen.
- Note: Old versions of *mdmorse* are not compatible with the new one.

## • Structure of the program



#### Introduction to atomistic simulations 2008

3. Neighbor lists and code  ${\tt mdmorse}$