## **Constructing a neighbour list**

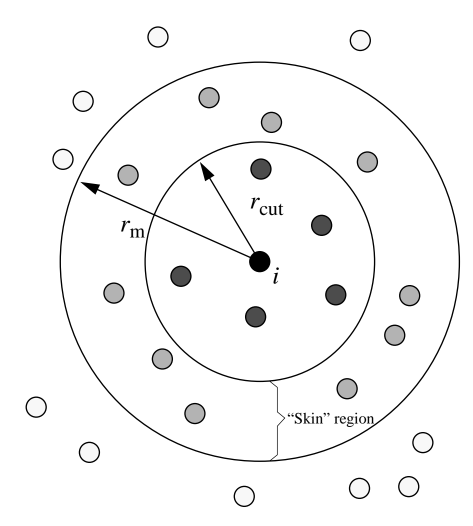
- In MD simulations (and actually many other applications) one of the central operations is the distances between atoms. In MD this is needed in the force calculation
- Trivial calculation of distances between atoms:

```
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  $N \to \infty$ .
- 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.

- Solution: Verlet 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  $\boldsymbol{N}_{\mathrm{m}}$  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

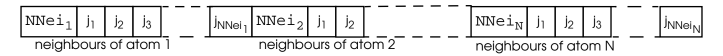


- An even better way to choose when to update the interval: after the neighbour list has been updated, keep a list of the maximum displacement of all atoms:
  - Make a separate table dxnei(i)
  - When you move atoms, also calculate dxnei(i)=dxnei(i)+dx
  - Calculate the **two** maximal displacements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

- Now when drneimax+drneimax2 >  $r_m$ - $r_c$  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.

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



- Here NNei<sub>1</sub> is the number of neighbours of atom i
- j<sub>1</sub>, j<sub>2</sub>, ... are the indices of neighbouring atoms
- So, if we would have a 64 atom system, where every atom has 4 neighbours, the neighbour list could look like this

- :



- A practical implementation of creating the list:

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

- ullet With the neighbour list, we can achieve a savings of a factor  $N_{
  m m}$  in calculating the distances to neighbours.
- But even using the neighbour list, our algorithm is still  $O(N^2)$ .

- 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  $27NN_c$  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:

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- array HEAD:

- **HEAD**:
- 10

- size =  $M^3$ 

- contains pointers to the table LIST - tells where the neighbours in subcell  $\boldsymbol{m}$  start
- LIST:

- 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.

- Building the list:

```
- assume a cubic case:

- MD cell size = size(3)
- size of subcell = size()/M
- MD cell centered on origin

HEAD:

8 3

LIST: 0 1 0 3 2 4 5 7 6 9
```

- 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 Mx, My, 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 >= 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
```

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

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

- 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>
```