Exercises Lecture VIII Macrostates and microstates: equilibrium and entropy. Metropolis algorithm in the canonical ensemble: verifying Boltzmann's distribution

1. MC simulation of a simple N-particles model

Consider an ideal gas of N non interacting, distinguishable particles, **confined** in a box (fixed **V**) and **isolated** (fixed **E**), divided into left/right with the possibility for one particle at a time to pass through the separation wall, with equal probability from the left to the right or viceversa.

A macrostate is specified for instance by the number of particles on the left side, say n, that can correspond to different microstates depending on the list of the specific particles there. A Monte Carlo approach consists in generating a certain number of movements, randomly, and consider them as representative of all the possible movements. The program box.f90 is a possible implementation of the algorithm describing the time evolution of the system in terms of macrostates, i.e. –given an initial number of particles on the left, n– the approach to equilibrium and what the equilibrium macrostate is.

- (a) Choose N=4, 10, 20, 40, 80, and n=N initially. Make a plot of n (or, better, of n/N) with respect to time. What is the equilibration time τ_{eq} (=how many MC steps)?
- (b) Modify the program so that at each time step t it calculates the number of particles $\langle n(t) \rangle$ averaged over different runs (e.g. 5 runs). Make a plot to compare n(t) over the individual runs and averaged $\langle n(t) \rangle$.
- (c) (Optional; do it at home!) Compare the numerical value of < n(t) > with the exact analytic results for a simple case, for instance N=4.
- (d) (Optional) Consider only one run. Modify the program to calculate numerically the probability P_n of having at equilibrium a macrostate with n particles on the left, by simply counting the number of occurring microstates that correspond to the macrostate n and dividing for the total number of microstates generated in the time evolution. Plot the histogram P_n for N=20, 40, 80 and a "sufficiently" long run. Comment.
- (e) Modify the program to measure the statistical fluctuations at the equilibrium, by calculating the variance $\sigma^2 = \langle n^2 \rangle \langle n \rangle^2$, where the average is done over a time interval *after* reaching the equilibrium.
- (f) Determine $\langle n \rangle$ and $\sigma / \langle n \rangle$ at equilibrium for N=20, 40, 80. Which is the dependence of these quantities on N?
- (g) An alternative method to find the equilibrium macrostate is the calculation of the entropy S_n of the different possible macrostates, by looking at the one with maximum entropy. An efficient numerical implementation is feasible by evaluating the ratio $\mathcal{R}_n =$ sum of possible coincidences for each microstate/maximum number of possible coincidences for each microstate, then calculating $S_n \propto -\log \mathcal{R}_n$. The code entropy.f90 calculates \mathcal{R}_n and \mathcal{S}_n . Use it with N=10. Compare the numerically calculated \mathcal{S}_n with the analytical value.

2. Simulated annealing

Simulated annealing is a stochastic method for global energy minimization, considering the system starting from a sufficiently high temperature; at each temperature it goes towards equilibrium according to the Boltzmann factor (see the application of the Metropolis algorithm in the canonical ensemble); then the temperature is slightly reduced and the equilibration procedure is repeated, and so on, until a global equilibrium state is reached at T=0. The method can be efficiently used for function minimization, even if the function is not representing an energy. In program simulated_annealing.f90 it is implemented for the minimization of f(x) = (x + 0.2) * x + cos(14.5 * x - 0.3). Initial temperature, initial position and scaling factor for the temperature are input quantities. Test the program by choosing different initial parameters and scaling factor for the temperature.

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! box.f90
T
! simulation of the evolution of a physical system towards equilibrium:
! non interacting particles in a box divided into two parts;
! at each time step, one and only one particle (randomly choosen)
! goes from one side to the other one
module moduli_box
 implicit none
 public :: initial, move
 integer, public :: N,tmax
contains
 subroutine initial()
   print*," total number of particles N >"
   read*.N
   tmax = 10*N ! we choose the evolution time proportional to N
 subroutine move()
   integer :: nl,itime
   real :: r, prob
   nl = N ! we start with all the particles on the left side
   open(unit=2,file="box.out",action="write",status="replace")
   do itime = 1,tmax
     prob = real(nl)/N
                       ! fraction of particles on the left
     call random_number(r)
     if (r <= prob) then
        nl = nl - 1
     else
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nl = nl + 1
end if
write(unit=2,fmt=*)nl
end do
close(unit=2)
end subroutine move
end module moduli_box
program box
use moduli_box
! compare a random number with the fraction of particles on the left, nl/N:
! if r.le.nl/N we move one particle from the left to the right;
! elsewhere from the right to the left
call initial()
call move()
end program box
```

```
! initialize parameters
integer :: il,ir
print*, " total number of particles N (<=10)>"
read*, N
print*, " number of particles O<nl<N initially on the left (MACROstate)>"
read*,nl
if(nl<=0.or.nl>=N)then
    print*,' not acceptable, wrong nl'
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stop
      end if
   nr = N - nl ! number of particles on the right
   print*, " number of exchanges >" ! no. of evolution steps of the macrostate
   read*, nexch
   allocate(micro(0:nexch))
   micro(0) = 0
   write(*,fmt=*)'nleft =',nl
   write(*,fmt=*)'nright=',nr
   do il = 1,nl
            list left particles
      1
      mleft(il) = il
            quantity characterizing the initial macrostate
      !
      micro(0) = micro(0)*2 + 2
    end do
    do ir = 1,nr
      !
           list right particles
      mright(ir) = ir + nl
   end do
1
    print*, 'microstate(0)=',micro(0)
I.
    write(*,fmt="(a,i2,a,10(1x,i2))")'nleft =',nl,' labels=',mleft
    write(*,fmt="(a,i2,a,10(1x,i2))")'nright=',nr,' labels=',mright
I.
 end subroutine start
 subroutine exch()
    !
         exchange one particle on the left (ileft)
    !
         with one particle on the right (iright)
   real, dimension(2) :: r
   integer :: iexch,ileft,jleft,iright,jright
   do iexch = 1,nexch
      1
                choose randomly the labels of the two particles
      call random_number(r)
      ileft = int (r(1)*nl + 1)
                                  ! 1 =< ileft =< nl
      iright = int (r(2)*nr + 1)  ! 1 =< iright =< nr</pre>
      jleft = mleft (ileft)
      jright = mright(iright)
      mleft (ileft) = jright ! new particle on the left
      mright(iright) = jleft ! new particle on the right
             characterizing the microstate:
      1
      micro(iexch) = micro(iexch-1) + 2**jright - 2**jleft
    print*, 'microstate(',iexch,')=',micro(iexch)
    write(*,fmt="(a,i2,a,10(1x,i2))")'nleft =',nl,' labels=',mleft
    write(*,fmt="(a,i2,a,10(1x,i2))")'nright=',nr,' labels=',mright
!
    end do
 end subroutine exch
```

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!

```
subroutine output()
          calculate the ratio of coincidences with respect to the total number
    !
          of possible pairs, and consequently entropy
    !
    L
    integer :: ncoin, ncomp, iexch, jexch
    real :: rate,S
    ncoin = 0
    ncomp = nexch*(nexch-1)/2
          compare microstates: if coincident, count + 1;
    1
    1
          upgrade counter
    do iexch = 1,nexch-1
       do jexch = iexch+1, nexch
          if (micro(iexch) == micro(jexch)) ncoin = ncoin + 1
       end do
    end do
    !
          coincidence ratio
    rate = real(ncoin)/real(ncomp)
    if (rate > 0) then
       S = log(1.0/rate)
       print*, " numerically estimated entropy: S=",S
       else
       print*, " no coincidences! estimated entroty infinite! "
  end subroutine output
end module ma
program entropy
 use ma
  1
               total number of particles
       N:
              number of left particles (i.e. the MACROstate)
  1
       nl:
  1
       mleft(),mright(): labels of left and right particles
        micro: a "global" label for a microstate, here defined through
  I.
               mleft() : micro=sum_{il=1,nl} 2**(mleft(il))
  L
        nexch: total number of exchanges (evolution steps of the macrostate)
  1
  !
               microst.)
  call start()
  1
       the macrostates evolves (exchanging particles, the microstate changes)
  call exch()
       calculate the fraction of coincidence of microstates over all
  1
        the possible coincidences with the microstates and the entropy
  !
  call output()
  deallocate(micro)
end program entropy
```

```
! simulated_annealing.f90
! for function minimization; adapted from U. Schmitt, 2003-01-15
PROGRAM anneal
IMPLICIT NONE
INTEGER :: istep, nsteps
REAL, PARAMETER :: scale=0.5 ! should be chosen for specific function
REAL :: func, fx, fx_min, fx_new, temp, tfactor, x, x_min, x_new
REAL, DIMENSION(2) :: rand ! random numbers
x = 1.0; fx = func(x); fx_min = fx ! starting point for search
PRINT *, 'Starting from x = ', x, ', f(x) = ', fx
PRINT *, 'initial (high) temperature (e.g., 10)?' ! annealing schedule
READ *, temp
PRINT *, 'annealing temperature reduction factor (e.g., 0.9)?'
READ *, tfactor
PRINT *, 'number of steps per block (equilibration, e.g., 1000)?'
READ *, nsteps
Do WHILE (temp > 1E-5) ! anneal cycle
 DO istep = 1, nsteps
   CALL RANDOM_NUMBER(rand) ! 2 random numbers
   x_new = x + scale*SQRT(temp)*(rand(1) - 0.5) ! stochastic move
   fx_new = func(x_new) ! new object function value
   IF (EXP(-(fx_new - fx)/temp) > rand(2)) THEN ! success, save
     fx = fx_{new}
     x = x_{new}
   END IF
     write(1,fmt=*)temp,x,fx
   IF (fx < fx_min) THEN
     fx_min = fx
     x_min = x
     PRINT '(3ES13.5)', temp, x_min, fx_min
   END IF
 END DO
 temp = temp * tfactor ! decrease temperature
END DO
End PROGRAM anneal
REAL FUNCTION func(x) ! Function to minimize
Implicit NONE
REAL :: x
func = (x + 0.2) * x + \cos(14.5 * x - 0.3)
END FUNCTION
```