

MULTI-BAND QMC: FORTRAN CODES EXPLAINED.

We start review of the code with parameter file which is used in most subroutines of the program.
 “param.dat” looks as :

```

    IMPLICIT REAL*8 (A-H,O-Z)
    parameter(L=16,Iwmax=2**10,Nom=Iwmax+1)
    parameter(Nlm=2,Nd=2*Nlm,Nf=Nlm*(Nd-1),Ns=1)
    complex*16 ci
    parameter (ci=(0.d0,1.d0) )
    parameter (pi=3.1415926535898d0)
    common/global/Beta,Zero,One,Two
    
```

Beta is $\beta = 1/T$, T is temperature.

Zero, One, Two are 0, 1, 2 correspondingly.

Where L is number of time slices, $Iwmax$ is number of ω points.

Nlm is number of orbitals,

Nd is maximum degeneracy (spin degeneracy “2” is taken into account). One can also say that is is maximum number of GFs treated in the system.

$Nf = 0.5 * Nd * (Nd - 1)$ is number of Ising spins.

Ns is real spins number (for AF case $Ns = 2$ and paramagnetic situation is treated with $Ns = 1$).

Meaning of ci, pi is clear.

```

    SUBROUTINE QMC(g0,g,dtau,u,iscf,idum,acc,nrat,nswpIN,nn)
C -----
C   IMPLICIT REAL*8 (A-H,O-Z)
    include 'param.dat'
    include 'mpif.h'
    DOUBLE PRECISION
    &   g0(L,Nd),   g(L,Nd)
    & , ga(L,L,Nd), gx(L,L,Nd), gst(L,L,Nd)
    & , v(L),      vn(L,Nf)
    & , del(L,L)
    & , xg(-L:L),  xgf(-L:L), G00(-L:L), xs(0:L)
    & , xf(Nd,Nf)
    & , u(Nf),     xlam(Nf)
    & , docc(Nf),  nn(Nf)
C
    DIMENSION s(2*L,1,Nf) ! it is to save memory space
    INTEGER ipoint(Nf,2)
    INTEGER myid,ierr,numprocs,master
C
    common/ising/ vn
    common/ifs/   if1,iff
    common/varmpi/myid,ierr,numprocs,master
C -----
    
```

Now we describe input data.

Weiss Green function (GF) $\mathcal{G}_0(\tau)$ is input function into QMC subroutine and $G(\tau)$ is output. In the program they are $g0(L, Nd)$ and $g(L, Nd)$ correspondingly. (L, Nd) means that GSs are defined in τ space (L slices) and for each degeneracy (Nd).

$ga(L, L, Nd), gx(L, L, Nd), gst(L, L, Nd)$ are working GSs they depend on τ and τ' and this is why all of them have (L, L, Nd) structure (time, time, Number_(of)_degeneracy). ga corresponds to G' in formulae above, gx is G and gst is used for storage ($gst = \text{Green Function } STorage$) of temporary GFs on each sweep.

Next line contains two arrays $v(L)$ and $vn(L, Nf)$. The first one is an auxiliary array to form array of Ising fields

$vn(L, Nf)$.

Array $del(L, L)$ is $L \times L$ identity matrix .

$xg(-L : L), xgf(-L : L), G00(-L : L)$ are temporary arrays used in “wrap-around” operation only.

$xf(Nd, Nf)$ contains the sign of the interaction (ie. +=ferro and -=antiferro) given the channel ($1 \rightarrow Nd$) and the pseudo spin ($1 \rightarrow Nf$). It is used only in clean update routine.

Matrix of Coulomb interactions is contained in $U(Nf)$ array.

And $xlam(Nf)$ contains information about all exponents used in Hubbard-Stratonovich transformations.

$docc(Nlm)$ is used to deal with double occupancy and it is unimportant for QMC itself.

Finally, $ipoint(Nf, 2)$ is enumeration array. It is used in calculation of determinants ratio.

Next step will be initialization.

```
if(idum.eq.0) idum=123457
nsweep=nsweepIN/numprocs
polar=0.5d0
ndirty=100
ncor=2
if(iscf.eq.1) then
  nwarm=100
else
  nwarm=30
endif
```

$idum$ is an arbitrary number to initialize random generator.

$nsweep$ is number of sweeps done per processor.

$ndirty$ is number of runs before the clean update which is necessary to check that the precision is not degraded.

$polar$ defines ratio of spin up and down.

$ncor = ?$

Next we initialize del , gst and xf arrays .

```
do in=1,Nf
  docc(in)=0.d0
  nn(in)=0.d0
enddo
C
do j=1,L
  do i=1,L
    del(i,j)=0.d0
    del(i,i)=1.d0
  enddo
enddo
c
do kc=1,Nd
  do j=1,L
    do i=1,L
      gst(i,j,kc)=0.d0
    enddo
  enddo
enddo
c
c initialize: xf
c xf contains in the sign of the interaction (ie. +=ferro
c and -=antiferro) given the channel (1-Nd) and the pseudo spin (1-Nf)
do i=1,Nd
  do j=1,Nf
    xf(i,j)=0.d0
  enddo
enddo
```

```

C
c      Sigma(i,ij) = 1 if i<j
c                = -1 if i>j
      ij=0
      do i=1,Nd-1
        do j=i+1,Nd
          ij=ij+1
          xf(i,ij)=1.d0
          xf(j,ij)=-1.d0
        enddo
      enddo

c initialization: ipoint(:2)
c first entry (1 -> Nf) is which pseudo spin flips
c and the second entry (1 -> 2) corresponds to
c 1 to ferro-coupling (+), and 2 to antiferro (-)
      ij=0
      do i=1,Nd-1
        do j=i+1,Nd
          ij=ij+1
          ipoint(ij,1)=i
          ipoint(ij,2)=j
        enddo
      enddo
C Structure of ipoint:
C 1 1 1 1 2 2 2 3 3 4
C 2 3 4 5 3 4 5 4 5 5

```

Before main loop we store all values of $\lambda_{mm'}$ in exponents (9).

```

do i=1,Nf
  z=dtau*u(i)/2.d0
  z=exp(z)
  xlam(i)=dlog(z+dsqrt(z**2-1.d0))
enddo

```

Next line is random generator initialization and following loops transform physical GF depending on τ into GF depending on τ and τ' plus we change sign of GF (historical reason).

```

c      initialization: random numbers generator
      r=ranw(idum)
c// Minus - change for QMC G-notation //
do kc=1,Nd
  do i=0,L-1
    g00(i)=-g0(i+1,kc)
  enddo
  do i=1,L-1
    g00(-i)=-g00(L-i)
  enddo
c
do ir=1,L
  do is=1,L
    gx(is,ir,kc)=g00(is-ir)
  enddo
enddo
enddo
c

```

Here we generate (initialize) field of auxiliary Ising spins $vn(L, Nf)$ for each time slice.

```

    if(iscf.eq.1) then
do iv=1,Nf
  xlam0=xlam(iv)
  call initial(xlam0,v,polar,idum)
  do i=1,L
    vn(i,iv)=v(i)
  enddo
enddo
endif

```

We do first clean update (full matrix inversion) and produce $G' \equiv ga$.

```

do kc=1,Nd
  call gnewclean(ga,vn,gx,del,xf,kc)
enddo

```

Read comments to learn meaning of kx , irr , $nrat$.

Here is very important point: START of the **MAIN** loop of simulations!

```

C -----
c Main loop does nsweeps: a clean update comes after ndirty dirty updates
c
c kx= # of measurements
c irr= # of accepted flips (after the warm-up)
c nrat= # of negative det encountered
C -----
  kx=0
  irr=0
  nrat=0
  kxmax=0
C
  DO 2 k=1,nsweep
    kk=mod(k,ndirty)
    kcor=mod(k,ncor)
    DO 5 j=1,L
      DO 55 iv=1,Nf

```

For a given spin configuration we make a spin flip and calculate the determinant ratio. Formulae (20), (21), (22), (23) are implemented. If determinant ratio (20) is negative we count such cases via $nrat$.

```

    dv=2.d0*vn(j,iv)
c// calculates the determinant ratio //
    kup=ipoint(iv,1)
    kdw=ipoint(iv,2)
    ratup=1.d0+(1.d0-ga(j,j,kup))*(dexp(-dv)-1.d0)
    ratdw=1.d0+(1.d0-ga(j,j,kdw))*(dexp(+dv)-1.d0)
    rat=ratup*ratdw
C
    if(rat.lt.0.)then
      nrat=nrat+1
    end if

```

Now we calculate determinant rat ratio (20) and compare it with a random number r and if $rat < r$ we accept spin flip, count it, and inset change into Ising spin configuration matrix vn .

```

        rat=rat/(1.d0+rat)
c// for num rec
        r=ranw(idum)
        if(rat.gt.r)then
            if(k.gt.nwarm)then
                irr=irr+1
            end if

```

We store the successful spin flip.

```

c
        vn(j,iv)=-vn(j,iv)

```

Depending on kk value (if it is equal to *ndirty* ($kk = 0$) then clean update should be done).

```

        if(kk.eq.0)then
            call gnewclean(ga,vn,gx,del,xf,kup)
            call gnewclean(ga,vn,gx,del,xf,kdw)
        else
            call gnew(ga,vn,j,iv,del,1.d0,kup)
            call gnew(ga,vn,j,iv,del,-1.d0,kdw)
        endif
    endif
55     CONTINUE
5     CONTINUE

```

Two loops in τ and in Ising spin are finished and we can store our measurement for a sweep. We store values of GF *gst* and *docc* – double occupancies.

```

c// store the measurements //
    if(kcor.eq.0.and.k.gt.nwarm)then
        kx=kx+1
        do kc=1,Nd
c// store GF //
            do jx=1,L
                do ix=1,L
                    gst(ix,jx,kc)=gst(ix,jx,kc)+ga(ix,jx,kc)
                enddo
            enddo
        enddo
    enddo
c
C -----
        ij=0
        do ik=1,2*Nlm-1
            do jk=ik+1,2*Nlm
                ij=ij+1
            do ix=1,L
C
                gak =1.-ga(ix,ix,ik)
                gakN =1.-ga(ix,ix,jk)
C
                docc(ij)=docc(ij)+gak*gakN
C
            enddo

```

```

                enddo
            enddo
C -----
c
    endif
c
2   CONTINUE ! end of the main QMC loop

```

After all simulations finished we can compute final GF, double occupancy, as well as the acceptance rate. Below we sum up GF and double occupancy.

```

C -----
c// acceptance rate :
    acc=dfloat(irr)/dfloat((nsweep-nwarm)*L*Nf)
c// normalization of GF //
    kxmax=kx*numprocs
    do kc=1,Nd
    do jx=1,L
        do ix=1,L
            gst(ix,jx,kc)=gst(ix,jx,kc)/dfloat(kxmax)
        enddo
    enddo
    enddo
C
    do k=1,Nf
        nn(k)=docc(k)/dfloat(kxmax*L)
    enddo

```

Getting back physical GF from QMC GF: $(\tau, \tau') \rightarrow (\tau - \tau')$.

```

c// wrap-around //
    DO kc=1,Nd
c// negative times (-) //
        do j=0,L-1
            xg(-j)=0.
            do i=1,L-j
                xg(-j)=xg(-j)+gst(i,i+j,kc)
            enddo
        enddo
c// positive times (+) //
        do j=1,L-1
            xg(j)=0.
            do i=1,L-j
                xg(j)=xg(j)+gst(i+j,i,kc)
            enddo
        enddo
c// it is enforced the time "antiperiodicity" //
        do i=1,L-1
            xgf(i)=(xg(i)-xg(i-L))/dfloat(L)
            xgf(i-L)=-xgf(i)
        enddo
        xgf(0)=xg(0)/dfloat(L)
        xgf(-L)=-xgf(0)

```

Finally, we must restore the sign of GF!

```

c// change MINUS sign from QMC //
  do i=0,L-1
    g(i+1,kc)=-xgf(i)
  enddo
c
  ENDDO ! kc=1,Nd
C -----
  And we have the output:  $g(L, Nd)$  !
C -----
  RETURN
  END

```

One can easily see correspondence between formulae (24) and (25) and subroutine “gnew” below. The key line is line with label 1.

SUBROUTINE gnew(g,vn,j,iv,del,xflag,kc) →
 SUBROUTINE gnew($G', V^m, \tau, m', \hat{I}, \pm 1, m$).
 $\pm 1 \equiv \begin{cases} +1 \rightarrow \uparrow \\ -1 \rightarrow \downarrow \end{cases}$;

```

c *****
c// dirty update of GF //
c
  SUBROUTINE gnew(g,vn,j,iv,del,xflag,kc)
  include 'param.dat'
  double precision g(L,L,Nd),vn(L,Nf),del(L,L),d(L,L)
c
  dv=xflag*2.*vn(j,iv)
  ee=exp(dv)-1.d0
  a=ee/(1.d0+(1.d0-g(j,j,kc))*ee)
  do i2=1,L
    do i1=1,L
1      d(i1,i2)=g(i1,i2,kc)+(g(i1,j,kc)-del(i1,j))*(a*g(j,i2,kc))
    enddo
  enddo
  do i2=1,L
    do i1=1,L
      g(i1,i2,kc)=d(i1,i2)
    enddo
  enddo
  RETURN
  END

```

To understand what is going on in this subroutine one should take a look at formulae (15), (16), (17), (18).

SUBROUTINE gnewclean(g,vn,gx,del,xf,kc) →
 SUBROUTINE gnewclean($G', V^m, G, \hat{I}, Q_{mm'}, m$)

```

C *****
c// clean update of GF //
c
  SUBROUTINE gnewclean(g,vn,gx,del,xf,kc)
  include 'param.dat'
  double precision
  & , g(L,L,Nd), vn(L,Nf)
  & , b(L,L), binv(L,L), del(L,L)
  & , gx(L,L,Nd), ee(L)

```

```

      & , xf(Nd,Nf)
c
do i=1,L
vv=0.d0
do iv=1,Nf
vv=vv+xf(kc,iv)*vn(i,iv)
enddo
ee(i)=dexp(vv)-1.d0
enddo
c
do j=1,L
do i=1,L
b(i,j)=del(i,j)-ee(j)*(gx(i,j,kc)-del(i,j))
enddo
enddo
c
call inverse(b,binv)
c
do i1=1,L
do i2=1,L
xdum=0.d0
do i=1,L

$$G' = A^{-1}G$$

xdum=xdum+binv(i1,i)*gx(i,i2,kc)
enddo
g(i1,i2,kc)=xdum
enddo
enddo
c
RETURN
END

```

This subroutine is obvious. It creates a random sequence of L spins.

```

C ***** ***** ***** ***** ***** *****
c// initialize the vector v of Ising fields //
c
subroutine initial(xlam,v,polar,idum)
include 'param.dat'
double precision v(L)
c
do i=1,L
s=1.d0
r=ranw(idum)
if(r.gt.polar) s=-1.d0
v(i)=xlam*s
enddo
RETURN
END
C ***** ***** ***** ***** ***** *****

```