

ctrl file

ctrl contains basis inputs to invoke lmf, lmfa, and lmchk.

ctrl has `Category-Token` structures. It is easier to understand it by examining the ctrl files generated by ctrlgenM1.py.

ctrl

ctrl is the basic input file. It is better to look into template of ctrl file generated by ctrlgenM1.py. Generated ctrl file is self-documented. See [Tutorial](#).

When we run fortran programs (lmf, lmfa, and lmchk),
we run ecalj/SRC/exec/ctrl2ctrlp.py (at your BINDIR) from the fortran program in order
to convert ctrl.foobar to ctrlp.foobar. See ~/ecalj/SRC/main/lmf.f90 for example.
Then we read ctrlp.foobar by SRC/subroutines/m_lmfini.f90.

Here is the list of `Category-Token`. Note that those with `.` at top is only for developers.
To see full list, look into `m_lmfini.f90` which reads in ctrlp file.

```
Category-Token      Input  cast  (size,min) -----
IO_VERBOS           opt   i4      1, 1      default= 30
  Verbosity for printout. Set from the command-line with --pr=xxx

IO_TIM              opt   i4v     2, 1      default= 1 1
  Turns CPU timing log. Value sets tree depth.
  Optional 2nd arg prints CPU times as routines execute.
  Args may be set through command-line: --time=#1,#2 --time=5,3 for
example.

. OPTIONS_HF        opt   lg      1, 1      default= F   T for non-
self-consistent Harris
```

```
STRUC_PLAT          reqd   r8v     9, 9      Primitive lattice vectors

STRUC_ALAT          reqd   r8      1, 1      Units of length (atomic
unit, a.u.)
STRUC_DALAT         opt    r8      1, 1      default= 0   added to alat
after reading inputs (only affecting to SPEC_ATOM_R/A case)
```

```
HAM_NSPIN           opt   i4      1, 1      default= 1   Set to 2 for
spin polarized calculations

HAM_REL             opt   i4      1, 1      default= 1
relativistic switch
```

0 for nonrelativistic Schrodinger equation
 1 for scalar relativistic Schrodinger equation

For SO=1 or 2, HAM_NSPIN must be 2

HAM_SO opt i4 1, 1 default= 0

Spin-orbit coupling (for REL=1)

0 : no SO coupling

1 : Add L.S to hamiltonian

2 : Add Lz.Sz only to hamiltonian

HAM_SOCAXIS opt r8v 3, 3 default= 0 0 1

SOC axis! Currently 0,0,1 (default) or 1,1,0 are effective for

HAM_SO=1

I think this works but not fully tested.

. HAM_GMAX reqd r8 1, 1

Energy cutoff for plane-wave mesh

* If HAM_GMAX is not parsed, attempt to read HAM_FTMESH:

HAM_FTMESH reqd i4v 3, 1

No. divisions for plane-wave mesh along each of 3 lattice
 vectors.

Supply one number for all vectors or a separate number for each
 vector.

. HAM_TOL opt r8 1, 1 default= 0.100D-05

wave function tolerance for FT mesh

. HAM_FRZWF opt lg 1, 1 default= F

Set to freeze augmentation wave functions for all species

HAM_FORCES opt i4 1, 1 default= 0

Controls the ansatz for density shift in force calculation.

-1 no force: no shift

1 free-atom shift 12 screened core+nucleus

HAM_XCFUN opt i4 1, 1 default= 2

Specifies local exchange correlation functional:

1 for Ceperly-Alder (VWN)

2 for Barth-Hedin (ASW fit)

103 for PBE-GGA (use xcpbe.F in ABINIT)

HAM_ScaledSigma opt r8 1, 1 default= 1

=\alpha_Q for QSGW-LDA hybrid. \alpha \times (\Sigma-Vxc^LDA) is
 added to LDA/GGA Hamiltonian.

. HAM_EWALD opt lg 1, 1 default= F

Make strux by Ewald summation

HAM_OVEPS opt r8 1, 1 default= 0.100D-06

Diagonalize hamiltonian in reduced hilbert space,

discarding part with evals of overlap < OVEPS

HAM_PWMODE opt i4 1, 1 default= 0

Controls APW addition to LMT0 basis. Use 11 usually.

1s digit:

LMT0 basis only

Mixed LMT0+PW

PW basis only

10s digit:

PW basis G is given at $q=0$

PW basis q -dependent. $|q+G|$ cutoff

(for jobgw=1, lmf automatically set PWMODE=11)

```
HAM_PWEMAX      opt    r8      1, 1      default= 0
  Include APWs with energy  $E < PWEMAX$  (Ry) ctrlgenM1.py set pwemax=3
. HAM_READP      opt    lg      1, 1      default= F !but =T
in ctrlgenM1.py
  Read Pnu and PZ (b.c. of radial func) from atmpnu.*(by lmf) when we
have no rst file

. HAM_V0FIX      opt    lg      1, 1      default= F
  Fix potential of radial functions-->Fix radial func. if READP=T
together
. HAM_PNUFIX      opt    lg      1, 1      default= F
  Fix b.c. of radial functions
```

```
SYMGRP          opt    chr      1, 0
  Generators for symmetry group
SYMGRPAF        opt    chr      1, 0
  One (or multiple) Extra Generator for adding anti ferro symmetry
```

SPEC_ATOM

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The following tokens are input for each species. See examples.

```
SPEC_ATOM       reqd    chr      1, 0
  Species label
SPEC_ATOM_Z      reqd    r8      1, 1
  Atomic number
. SPEC_ATOM_R     reqd    r8      1, 1
  Augmentation sphere radius rmax
* If token is not parsed, attempt to read the following:
  SPEC_ATOM_R/W   reqd    r8      1, 1
  rmax relative to average WS radius
* If token is not parsed, attempt to read the following:
  SPEC_ATOM_R/A   reqd    r8      1, 1
  rmax ratio to alat
. SPEC_ATOM_A     opt    r8      1, 1      default depends on
other input
```

Radial mesh point spacing parameter

```
. SPEC_ATOM_NR      opt    i4      1, 1      default= 51
    Number of radial mesh points
```

`. SPEC_ATOM_RSMH` reqd r8v 10, 1
Smoothing radii for basis. Gives l-cut max for base. We set $=1/2 \cdot R$ in ctrlgenM1.py

```
. SPEC_ATOM_EH      reqd    r8v      10, 0
    Kinetic energies for basis          We set EH=-1 in ctrlgenM1.py
```

`. SPEC_ATOM_RSMH2` opt r8v 10, 1
Basis smoothing radii, second group

```
. SPEC_ATOM_EH2     opt    r8v      10, 0
    Basis kinetic energies, second group We set EH=-2 in ctrlgenM1.py
```

`. SPEC_ATOM_LMX` opt i4 1, 1 default= 10
optional l-cutoff for basis. Convenient for debugging or tests.

`SPEC_ATOM_LMXA` opt i4 1, 1 default depends on other input
l-cutoff for augmentation
We use lmx=4 mainly but lmax=6 for 4f,5f atoms.

`SPEC_ATOM_MMOM` opt r8v 1, 1 default= 0
Starting mag. moms for each l channel.
For a channel with PZ, this is enforced to be zero.
See explanation for SPEC_ATOM_Q.

`SPEC_ATOM_PZ` opt r8v 1, 1 default= 0
Starting semicore log der. parameters
Add 10 to attach Hankel tail

`. SPEC_ATOM_LMXL` opt i4 1, 1 default depends on other input
lmax for which to accumulate rho,V in sphere

`. SPEC_ATOM_P` opt r8v 1, 1 default= 0
Starting log der. parameters for each l

`. SPEC_ATOM_Q` opt r8v 1, 1 default= 0
Starting valence charges for each l channel.
Q do not include semicore(PZ) electrons.
Charge configuration is shown by lmfa
WARN: This version cannot treat two valence channels per l (Q for a l-channel is zero if the l is with PZ).
This causes a problem typically in Li; then we can not treat both of PZ=1.9 and P=2.2 as valence.
To avoid this, use Q=0,1 together. This trick supply an electron to 2p channel; this trick works fine.

`. SPEC_ATOM_NMCORE` opt i4 1, 1 default= 0
spin-averaged core: jun2012takao
0(default): spin-polarized core
1 : spin-averaged core density is from spin-averaged potential

`. SPEC_ATOM_LFOCA` opt i4 1, 1 default depends on other input
FOCA switch 0(within MT):=1(frozenCore). Default: 1 for $z > 8$; 0 for $z \leq 8$

```
. SPEC_ATOM_KMXA      opt    i4        1, 1          default= 3
    k-cutoff for projection of wave functions in sphere.
. SPEC_ATOM_RSMA      opt    r8        1, 1          default depends on
other input
    Smoothing for projection of wave functions in sphere.
    input<0 => choose default * -input
. SPEC_ATOM_IDMOD     opt    i4v       1, 1          default= 0
    idmod=0 floats P to band CG, 1 freezes P, 2 freezes enu
. SPEC_ATOM_CSTRMX    opt    lg        1, 1          default= F
    Set to exclude this species when automatically resizing sphere radii
(SCLWSR>0)
. SPEC_ATOM_FRZWF     opt    lg        1, 1          default= F
    Set to freeze augmentation wave functions for this species
```

- The next three tokens are for LDA+U

```
SPEC_ATOM_IDU        opt    i4v       4, 1          default= 0 0 0 0
    LDA+U mode:  0 nothing, 1 AMF, 2 FLL, 3 mixed; +10: no LDA+U if
sigm.* exist
SPEC_ATOM_UH         opt    r8v       4, 1          default= 0 0 0 0
    Hubbard U for LDA+U
SPEC_ATOM_JH         opt    r8v       4, 1          default= 0 0 0 0
    Exchange parameter J for LDA+U
```

Core hole options

```
SPEC_ATOM_C-HOLE    opt    chr        1, 0
    Channel for core hole
SPEC_ATOM_C-HQ      opt    r8v       2, 2          default= -1 0
    Charge in core hole. Optional 2nd entry is moment of core hole:
    Q(spin1) = full + C-HQ(1)/2 + C-HQ(2)/2
    Q(spin2) = full + C-HQ(1)/2 - C-HQ(2)/2

. SPEC_ATOM_EREf     opt    r8        1, 1          default= 0
    Reference energy subtracted from total energy (we use little)
```

SITE

The following tokens are input for each site. See examples.

```
SITE_ATOM           reqd    chr        1, 0
    Species label
SITE_ATOM_POS       reqd    r8v       3, 1
    Atom coordinates, cartesian in alat
    * If token is not parsed, attempt to read the following:
SITE_ATOM_XPOS      reqd    r8v       3, 1
```

```

Atom POS. fractional(POSCAR direct) coordinates
SITE_ATOM_AF      opt    i4      1, 1      default= 0
    antiferro ID:=i and -i should be af-pair, we look for space-group
operation with spin-flip
SITE_ATOM_RELAX   opt    i4v     3, 1      default= 1 1 1
    relax site positions (lattice dynamics) or Euler angles (spin
dynamics)

```

Parameters for Brillouin zone integration ---

```

BZ_NKABC          reqd    i4v     3, 1
    No. qp along each of 3 lattice vectors.
    Supply one number for all vectors or a separate number for each
vector.

. BZ_BZJOB        opt    i4v     3, 1      default= 0
    0 centers BZ mesh at origin, 1 centers off origin
    Supply one number for all vectors or a separate number for each
vector.
    QSGW requires the default setting.

. BZ_METAL        opt    i4      1, 1      default= 3
    0 insulator only; 3 for metal (2 is for maintenance)

. BZ_TETRA        opt    lg      1, 1      default= T
    Tetrahedron integration

. BZ_N            opt    i4      1, 1      default= 0
    N>0: Polynomial order for Methfessel-Paxton sampling
    N=0: Conventional Gaussian sampling
    N<0: Broadening by Fermi-Dirac distribution
    To be used in conjunction with W= ; see next

. BZ_W            opt    r8      1, 1      default= 0.500D-02
    If BZ_N>=0, Line broadening for sampling integratio
    If BZ_N<0, Temperature for Fermi distribution (Ry)

BZ_ZBAK          opt    r8      1, 1      default= 0
    Homogeneous background charge

. BZ_SAVDOS        opt    i4      1, 1      default= 0
    Choose 0(F) or 1(T): Write dos.tot.* file (settings are NPTS and
DOS)

BZ_NPTS          opt    i4      1, 1      default= 2001
    No. DOS points (sampling integration)

BZ_DOSMAX        opt    r8      1, 1      default= 2.940
    Maximum energy to which DOS accumulated, relative to Efermi

. BZ_EFMAX        opt    r8      1, 1      default= 5
    Find evecs up to efmax

. BZ_NEVMX        opt    i4      1, 1      default= 0
    Find at most nevmx eigenvectors
    If NEVMX=0, program uses internal default
    If NEVMX<0, no eigenvectors are generated

BZ_FSMOM         opt    r8      1, 1      default depends on
other input

```

```

Fixed-spin moment (fixed-spin moment method)
BZ_FSMOMMETHOD    opt    i4        1, 1        default= 0
Method of Fixed-spin moment 0:original 1:discrete

```

Parameters for Ewald sums ---

```

. EWALD_TOL        opt    r8        1, 1        default= 0.100D-07
Ewald tolerance

```

Parameters for iterations ---

```

ITER_NIT           opt    i4        1, 1        default= 30
maximum number of iterations in self-consistency cycle
. ITER_NRMIX        opt    i4        1, 1        default= 80
lmfa rseq max iter
ITER_MIX           opt    chr       1, 0
Mixing rules for charge mixing. Syntax:
A[nmix][,b=beta][,bv=betv][,n=nit][,w=w1,w2][,nam=fn][,k=nkill][;...]
or
B[nmix][,b=beta][,bv=betv][,wc=wc][,n=#][,w=w1,w2][,nam=fn][,k=nkill]
ITER_CONV          opt    r8        1, 1        default= 0.100D-03
Tolerance in energy change from prior iteration for self-consistency
ITER_CONVC         opt    r8        1, 1        default= 0.100D-03
Tolerance in output-input charge for self-consistency
ITER_UMIX          opt    r8        1, 1        default= 0.500
Mixing parameter for densmat in LDA+U
ITER_TOLU          opt    r8        1, 1        default= 0
Tolerance for densmat in LDA+U
mmmixing parameters: A/B nmix wt: 0 -1 1.000000 1.000000 -9.000000
beta elin wc killj= 1.000000 -1.000000 0

```

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Parameters for dynamics and statics ---

```

DYN_MODE           opt    i4        1, 1        default= 0
0: no relaxation
4: relaxation: conjugate gradients
5: relaxation: Fletcher-Powell
6: relaxation: Broyden
DYN_NIT            opt    i4        1, 1        default= 1
maximum number of relaxation steps (statics) or time steps
(dynamics)
. DYN_HESS          opt    lg        1, 1        default= T
Read hessian matrix
. DYN_XTOL          opt    r8        1, 1        default= 0.100D-02
Convergence criterion in displacements

```

```

    XTOL>0: use length; <0: use max val; =0: do not use
. DYN_GTOL          opt    r8          1, 1          default= 0
    Convergence criterion in gradients
    GTOL>0: use length; <0: use max val; =0: do not use
. DYN_STEP          opt    r8          1, 1          default= 0.015
    Initial (and maximum) step length
. DYN_NKILL         opt    i4          1, 1          default= 0
    Remove hessian after NKILL iter

```

lmf console input

ecalj made from band structure part (lmf), and GW part (plus some additional functionalities such as Wannier).

lmfa (spherical atom for initial condition) and lmchk (crystal structure check) are by single core.

usage: lmf [--OPTION] [-var-assign] [extension]

```

usage:  lmfgwd [--OPTION] [-var-assign] [extension]

--help          List categories, tokens, and data program expects, and
quit
--show          Print control file after parsing by preprocessor,
                and echo input data as read from the control file
--pr=#1         Set the verbosity (stack) to values #1
--time=#1[,#2]  Print timing info to # levels (#1=summary; #2=on-the-
fly)

-vnam=expr      Define numerical variable "nam"; set to result of 'expr'
--jobgw=1 or 2  lmf-MPIK works as the GW driver (previous lmfgw-
MPIK)
--quit=band,    Quit after band
                Quit after initial setup. Convenient for check.
--quit=dmatu    Quit after dmatu
--quit=band     Quit after band calculation.
--quit=mkpot or --quit=dmat: Stop points. Surpress writing rst

NOTE: Read rst.* prior to atm.* file (No --rs options: 2022-6-20)
NOTE: Other command-line-options => Search "call cmdopt" in
SRC/*/*.f90

```

Search --quit option in [SRC/*/*.f90](#)