

ecalj manual

<https://github.com/tkotani/ecalj>

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Abstract

ecalj is an all-electron full-potential electronic-structure calculation package, especially for GW/QSGW calculations, in addition to its unique one-body problem solver which uses both augmented plane wave (APW) and the Muffin-tin orbital (MTO) simultaneously. Available freely from github.

This is based on the formulation in Refs. [1] and [2] (See Kotani2114QSGWinPMT.pdf and KotaniKinoAkai2015FormulationPMT.pdf at [ecalj/Document/Manual/](#)) on top of developments [3]. In **ecalj**, we can do QSGW and related calculations based on the $\text{PMT}=\text{LAPW}+\text{LMTO}$ method.

To get minimum usage for **ecalj**, read up to the Section.5 or Sec.8. (We can generate model Hamiltonian through Wannier functions([ecalj/MATERIALS/CuMLWF](#)), and so on, but not documented well yet...)

Be careful. This document still contain many bugs... Let me know bugs in this manual (try to fix things step by step). We need your help on this manual.

Requirement

To support of our **ecalj** activity, we need your acknowledgment to **ecalj** in your publications See [ecalj/README.org](#) at <https://github.com/tkotani/ecalj/>.

Contributions

There are many contributors to this package. T.Kotani currently maintain the **ecalj** package as a results of them.

Relation with Questaal at <http://www.questaal.org/>:

Questaal is one-another package to perform QSGW distributed by Mark van Schilfgaarde mainly in FP-LMTO, not in the PMT. In the package, there is the driver to use a modified version of the GW code in **ecalj** (GW part in **Questaal** is originated in **ecalj**). The FP-LMTO in **questaal** and **ecalj** have some similarities (for input files and outputs).

History

T.Kotani learned the LMTO-ASA-GW code by F.Aryasetiawan, which is on top of the Stuttgart's LMTO-ASA, mainly by van Schilfgaarde under O. Jepsen and O. K. Andersen <http://www2.fkf.mpg.de/andersen/LMTODOC/LMTODOC.html>. A FP-LMTO package is originally given by M.Methfessel, Mark van Schilfgaarde and their collaborators. T.Kotani developed an all-electron full-potential GW method on top of the FP-LMTO during his subbatical at 1999-2000 at the lab of M.Schilfgaarde. Then the QSGW method has developed with the help of S.Faleev and M.Schilfgaarde, first published in 2004. Then T.Kotani was adding new functionality such as spin fluctuation, impact ionization. In year 2008, T.Kotani started the $\text{PMT}=\text{LMTO}+\text{LAPW}$ method to remove problem of the FP-LMTO. After T.Kotani moved to Tottori-u at 2009, he spends time for the improvement of PMT method. T.Kotani and H.Kino showed that the PMT method can describe even the diatomic molecules efficiently and accurately. It turns out very limited number of APWs can remove such problems. Then T.Kotani have developed PMT-QSGW method. It is now stable, easy, and reliable rather than the previous version of QSGW on top of FP-LMTO.

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Change LOG:

- Jan2017 Add minimum description for syml generator at Sec.4.7.1
- Aug2019 slight modification for CMD35

1 Introduction

The `ecalj` is an first-principles electronic-structure calculations package with some unique features. With `ecalj`, we can do not only standard calculations (LDA/GGA/LDA+U, relaxation of atomic positions), but also the quasiparticle self-consistent GW calculations(QSGW), linear responses (charge and spin), Wannier functions (and U of them).

This is base on an unique one-body problem solver, the PMT method (=the Linearized APW+MTO method) [1]. Thus we identify the QSGW method implemented in `ecalj` as the PMT-QSGW method. Introduction to the PMT-QSGW is given in Sec.1.1. Today “QSGW” is accepted as a standard procedure in the electronic structure calculations [4].

First, see README.org shown at <https://github.com/tkotani/ecalj#ecalj-> (or `ecalj/README.org` in the package). Free to download `ecalj` package from it, and use it. The QSGW code is version controlled by git.

The `ecalj` is related to a FP-LMTO package `lmv7` seen at `ecalj/Document/Manual/CategoryAndToken.org` (or `html`).

The `lmv7` and `ecalj` are branched at year 2009. After branched, we added new features: simple install and test; all codes are in `f90` (no C compiler); new methods, especially PMT-QSGW; MPI parallelization for QSGW; simple usage with automatic setting of default files by `python ver2`; a small tool to convert VASP POSCAR to `ecalj`, and so on. PMT-QSGW shows more stable convergence than the previous version, FP-LMTO-GW ¹.

`ecalj` package mainly consists of two parts. One is one-body part(PMT part) (in `ecalj/lm7K/`), the other is many-body part(GW part) (in `ecalj/fpgw/`).

One-body part (PMT part), based on [1]

We can perform standard calculations such as LDA,GGA,LDA+U, atomic position relaxation, and so on. In addition, the PMT part has an interface to perform GW (and QSGW) calculation: the one-body part can include a given non-local exchange-correlation potential stored in a file `sigm.*`. ² The QSGW calculation is performed by a script `gwsc`, which has an iteration loop calling the one-body program (`lmf`) and many-body part (GW part) alternatively. The many-body part generate the file `sigm.*`. See Fig.6 and around.

Many-body part (GW part), based on [2]

As the inputs for the GW calculation, we have to supply the eigenfunctions and the eigenvalues from the one-body part to the GW part. The eigenfunctions re-expanded by the two types of basis functions, the atomic-like argumentation functions in the muffin-tin(MT) spheres, and the plane-waves in the interstitial region, say, the interstitial plane-wave (IPW) hereafter. IPW is defined as the usual plane waves in the interstitial region, but zero within MTs'. The IPWs + "atomic like functions within MTs" make "a basis set to expand eigenfuncitons". See Eq.(17) of [2].

We need another basis set to expand "product of eigenfuncions". That is, the mixed product bases (MPB)

[5, 6], which consists of the two kinds of bases (caution: do not mixed up with the basis set for eigenfuncitons); (i)the local atom-centered functions confined to MT spheres, so-called the product basis; (ii) IPW. The product-basis are calculated from products of solutions to the Schrödinger equation within the MT sphere. The Coulomb matrix v , the dynamically screened Coulomb interaction W , and so on, are expanded in the MPB. It can virtually span all the space made of product of eigenfunctions (but, in practical calculations, we need to use a small size of the bases to reduce computational time). We include full energy-dependence of $W(\omega)$. See Sec.3 of [2].

¹In cases to treat magnetic systems which have intrinsic magnetic fluctuations, we may need to be careful about initial condition or mixing procedure to get convergence. In cases, we need to start from LDA+U results as initial condition from which we start QSGW. Let me know about such trouble.

²In the case of using `sigm.*`, total energy shown in the console output (also in `save.*`) are just the indicator, not the meaningful total energy

Recently T.Kotani includes the Wannier function generator, which was originally developed by T.Miyake and H.Kino on top of previous version of GW part. Thus the Wannier functions (including effective interactions) can be generated in the PMT-QSGW. (U parameters in the full-screening and cRPA).

1.1 Uniqueness of the `ecalj` package.

We will explain two unique points of `ecalj`.

PMT:

Central part in an electronic structure packages is one-body problem solver. It means how to calculate eigenvalues/eigenfunctions for a given one-body potential. Inversely, we have to generate new one-body potential for given eigenfunctions/eigenvalues based on the density functional theory (DFT) in the LDA or GGA (In the followings, LDA means both of LDA and GGA). Then we can make the electron density self-consistent by iterations until converged, and obtain total energy of ground states. Then we can calculate atomic forces by perturbation. Based on such an one-body problem solver, we can implement kinds of methods; e.g, dielectric function, magnetic susceptibility, transport and so on. Furthermore, we can implement higher-level approximations such as the QSGW method explained below. An one-body problem solver (in linear methods) are characterized by

- (i) linear combinations of what basis set to represent eigenfunctions;
- (ii) how to represent electron density and one-body potential.

In `ecalj`, we use the PMT method [1, 7] as the one-body problem solver. The PMT method is a new all-electron full potential method. It uses not only the augmented plane waves (APW) but also the muffin-tin orbitals (MTO) together, in addition to the local orbital (lo's), to represent the eigenfunctions (no other methods use two kinds of augmented waves together). Thus eigenfunctions are expanded in the linear combinations of the APWs, MTOs, and the lo's. The formulation is clarified in Ref.[1]. Then the electron density and the one-body potential are given in the "3-components representation". That is, the electron density (one-body potential) is divided into three components,

"smooth part + onsite muffin-tin (MT) part - counter part".

Here the counter part is in order to remove smooth part within MTs. This formalism (Soler-Williams formalism [8, 9]) is also used in the projected augmented wave (PAW) method such as VASP.

We now usually use highly localized MTOs together with APWs of low energy cutoff ($3 \sim 4$ Ry).³ I think this is promising not only for efficient DFT/QSGW scheme, but also for kinds of applications in future.

QSGW:

In `ecalj`, we can perform the GW calculation. The usual GW approximation is so-called "one-shot GW" starting from LDA. It usually only calculates differences between the quasi-particle energies (QPEs) and the LDA eigenvalues by a perturbation (only diagonal part of self energy for the LDA eigenfunctions). Its ability is limited; it may fail when its starting point (eigenfunctions and eigenvalues supplied by LDA) is problematic. This is the reason why we originally develop the QSGW method. The QSGW now becomes popular and taken as a possible candidate to go beyond current limitation of such GW and LDA/GGA [4]. In principle, results given by QSGW do not depend on LDA anymore; the LDA are only used to prepare initial condition for self-consistency iteration cycle of the QSGW calculation⁴.

Usually the QPEs obtained by QSGW reproduce experiments better than LDA. For example, the band gap by GGA for GaAs is about 0.5 eV in contrast to the experimental value of 1.69 eV⁵. On the other hand, the QSGW predicts about 1.8~1.9eV, a few tenth of eV larger than experiment (for practical use, we sometimes use "hybrid functional between QSGW and

³current implementation have not yet efficiently use this locality; this must allow us to speed up one-body problem solver.

⁴Exactly speaking, we use LDA idea for efficient implementation of QSGW; thus obtained results are slightly dependent on the choice of LDA or GGA

⁵We undo electron-phonon effect (0.06eV) and spin-orbit effect (0.11eV) from the true the experimental value 1.52 eV.

LDA” so as to obtain smaller band gap). Even in the case of NiO and so on, the QSGW gives reasonable results (there is a tendency to give a little larger band gaps than experiments). This is in contrast to the case of the one-shot GW applied to NiO, where we can not have good agreement with experiments because the starting points in LDA is problematic.

The `ecalj` have other functions. LDA+U, atomic forces and relaxation (in GGA/LDA), core level spectroscopy and so on. In addition, we can calculate dielectric functions and magnetic responses from QPEs and the quasiparticle eigenfunctions given by LDA/QSGW. But total energy in QSGW is still in research (shown total energies in QSGW calculations are dummy now).

The QSGW calculations are very time-consuming; roughly speaking, it takes 10 or more times expensive than usual one-shot GW (although we can reduce computational time by choosing computational conditions). Thus the size of systems which we can treat is limited to ten atom in a cell or something, say, with a node of 16 cores; computation may require a week or so to have reasonable convergence. (heavy atoms require longer computational efforts, light atoms faster; non-magnetic systems are easier. We still have much room to accelerate the method, but not have done yet so much. Minimum MPI parallelization is implemented). The computational effort is $\propto N^4$ in the most time-consuming part of QSGW.

1.1.1 What do we expect for QSGW?

Let us recall hybrid functionals such as B3LYP, and LDA+U. In hybrid functional methods, we use $V_{xc} = (1-\alpha)*LDA + \alpha*(\text{Fock exchange like term})$, where α is taken to be ~ 0.25 usually⁶. The α can be dependent on materials; for metals α should be almost zero. For larger band gap insulator, α becomes larger.⁷ Despite of success of the functional, its ability is limited. For example, it is known that a hybrid functionals fail to describe metals such as bcc Fe. On the other hand, we have LDA+U method which succeeded to describe materials including localized electrons. However, it contains kinds of ambiguity and U is chosen by hand. The important part of the hybrid functional methods and LDA+U is the non-local potential. It is missing in the DFT. As we discussed above, they give some success but not satisfactory. We somehow need to have a method to determine high-quality non-local potential (a substitution of the exchange-correlation potential in LDA). It is the QSGW method.

Note two important aspects of non-local potential (missing character in the local potential used in DFT). One is the onsite non-locality; it is also taken into account by LDA+U model. However, note that relative shift of O(2p) band with respect to the center of 3d band is not in LDA+U. The other is the off-site non-locality (mainly between nearest neighbors), which may relate to LUMO-HOMO gap. A non-local potential can behave a projector which push down only the HOMO states (valence band) to lower energy. This can be in the hybrid functional but not in LDA+U.

In the QSGW, we determine such a non-local potential with the calculation of the GW method, in a self-consistent manner (we repeat GW calculations until converged). We can expect QSGW much more than hybrid methods/LDA+U. Roughly speaking, because the QSGW automatically determine U of LDA+U, or alpha of the hybrid functionals. More accurately speaking, we determine not only G_0 but also W (the screened Coulomb interaction) self-consistently. Here W corresponds to U and alpha. Thus QSGW gives reasonable results even if it is applied to metals such as Fe. For systems with metallic screening, it gives small non-locality (results are close to those of LDA). For systems with large band gap, QSGW gives large enough non-locality (like $0.25*(\text{Fock exchange})$).

Since we now need to treat complex systems, e.g, metal on insulator, it is very essential to treat kinds of materials on a same footing.

The main purpose of QSGW is to determine an one-particle effective Hamiltonian H^0 ⁸, which describes the quasiparticle picture (or independent-particle picture) for the system we calculate. In other words, QSGW divides the full many-body Hamiltonian H into $H = H^0 + (H - H^0)$. The screened Coulomb interaction W is determined self-consistently in

⁶exactly speaking, we have range cutoff for Fock exchange term in the HSE functional in addition

⁷If you use $\alpha = 1$ (Hartree-Fock limit), the band gap of Si becomes 20eV or something.

⁸people often pronounce this H-naught

the QSGW iteration cycle.

In comparison with LDA, we see differences;

- Band gap. QSGW tends to give slightly larger than experiments. It looks systematic.
 - Band width. Usually, sp bands are enlarged (except very low density case such as Na). This is the case for homogeneous electron gas. As for localized bands like 3d electrons, they can be narrowed.
 - Relative position of bands. e.g. O(2p) v.s. Ni(3d). More localized bands tends to get more deeper. Exchange splitting between up and down (like LDA+U) get larger. In cases such as NiO, magnetic moment become larger; closer to experimental values.
 - Hybridization of 3d bands with others. QSGW tends to make eigenfunctions localized.
- However, reality is complexed, and not so simple in cases.

1.2 Rule in this manual

- **This font** is for executable file(program) or shell scripts.
- `echo 3|hbasfp0` means doing `hbasfp0` with the argument '3' supplied as the standard input (`read(*,*)` in fortran).
- **This font** is for files, directories, contents of files, or variables used in codes.
- `ctrl.si,rst.si` and so on mean the case of Si. You may need to replace the extension `si` for your case. (this extension is given by user. Lower case, number, and underscore [a-z0-9_] are allowed.) In the followings, `ctrl.*` means a file with such an extension.
- There are files named `foobarU` and `foobarD`, which are for up spin (`isp=1`) and for down spin (`isp=2`), respectively; for example, `SEXU` and `SEXD`. We sometimes use `foobarU` to denote `foobarU` and `foobarD` together.
- **k** vector in the Brillouin zone is called as **q** or **k**.

1.3 What can we do with the ecalj package?

At Feb.2015, what we can do is as follow. We have limited parallelization. (e.g. k point parallel).

- LDA/GGA LDA+U, calculations, atomic forces and relaxation. Spin-orbit is included only for co-linear spin-density cases.
- Quasi-particle(QP) energy in the 1st-iteration from LDA. (one-shot *GW*)
Make band plot for LDA and the QP energies.
- Spectrum function of the self-energy Σ . Life time (imaginary part) of QPs.
- Dielectric function, and its inverse. (including local-field effect or not).
- QP self-consistent *GW*(QSGW)
- magnetic susceptibility.
- Wannier function. (not only one-body part, but also effective interaction *W* and cRPA)

2 Install

Install and minimum tests are easy; even in a note PC, e.g., we can use gfortran in Ubuntu. In Thinkpad T480s (2018) install may take five minutes, and all tests may take another fifteen minutes.

For productive runs, we may need multi-cores. Current implementation for parallelization by MPI is limited (not so much especially for the dielectric function part yet). Thus, probably, it may be not so efficient to use too many cores.

Follow the instruction of [ecalj/README.org](https://github.com/tkotani/ecalj/README.org).
or we can see the same one at <https://github.com/tkotani/ecalj/README.org>. We have a command `ecalj/InstallAll.ifort` and so on. This command installs `ecalj` and run a series of install tests automatically.

2.1 Binaries and Scripts

Main binaries and Scripts contained in `ecalj` are

- `ctrlgenM1.py`
Generate default input file `ctrl.*` from the structure file `ctrls.*`. The latter file only contains information of crystal structure.
- `lmfa`
Spherical atom calculation as initial condition, and core charge.
- `lmchk`
Check atomic positions, crystal symmetry, and computational conditions. Check ID of MuffinTins (MTs of atoms specified by `ctrl` file is re-ordered by computer.
- `lmf` and `lmf-MPIK` LDA/GGA, LDA+U calculations. (or we can use Vxc in QSGW instead). We mainly use `lmf-MPI` (k-parallel version) instead of `lmf`.
- PROCAR mode of `lmf`: Fat band mode.
`mpirun -np 4 lmf-MPIK --mkprocar --band:fn=sym1 mgo` gives PROCAR
(Try an example `/ecalj/MATERIALS/MgO-PROCAR/`. Run `./job` at the directory).
- `gwsc`
QSGW calculation
- `job_band, job_tdos, job_pdos`
band, fermi surface, tdos, pdos plot.
- `job_fermisurface`
Run `job_fermisurface cu -np 4 10 10 10`. This write down all band energies (around E_f) for 10x10x10 in BZ. Then we can view it with `xcrysden` as `xcrysden --bxsf fermiup.bxsf` (usage of `xcrysden` need to be written here). With `-allband` added for `job_fermisurface`, we can write all the bands. So, we need this option if you like to write iso-energy surface at high (or low) energy.
- `epsPP_lmfh`
Dielectric function without local field correction (LFC).
- `eps_lmfh`
Dielectric function with LFC
- `epsPP_lmfh_chipm`
Non-interacting transverse spin polarization.
- `gw_lmfh`
One-shot GW calculation. This also show life-time of QPs (`QPU_lmf`). (we need make it parallelized...)
- `genMLWF`
Wannier functions and matrix elements of W on it. A implementation of cRPA included.
- `dqpu`
A small python script to compare `QPU.*` files (eigenvalues are compared) numerically.

2.2 tests

Install. ifort run tests at ecalj/TestInstall.

In the following, si:gw_lmfh means '>make si_gw_lmfh' at ecalj/TestInstall/; this test is performed with the Makefile at the directory.

si:gw_lmfh/	Results: QPU
si:gWSC/	: QPU,log.si
gas:gWSC/	: QPU,log.gas
nio:gWSC/	: QPU,log.nio
fe_epsPP_lmfh_chipm:	: ChiPM*
gas:eps_lmfh/	: EPS*
gas:epsPP_lmfh/	: EPS*

(These are just samples; not for practical calculations)

2.3 Directory structure

```
ecalj
  InstallAll.ifort, Install.gfortran ! install and test
  Document/
  fpgw/      ! full potential GW code
  fpgw/Wannier ! MaxLoc Wannier and cRPA code.
  lm7K/      ! PMT method
  MATERIALS/ !job_materials.py contains examples.
  StructureTool/ ! POSCAR converter, and structure tool
  GetSymb1/ ! Brillouin zone and symmetry lines generator and visualizer.
  TestInstall/ ! Install test; this is invoked from Install.*
  TOOLS/      !Tools for developers
```

3 Theory (note)

aaa Except the technical details, we need to know minimum for these theories.

- DFT in LDA/GGA.
- GW
- QSGW

There are literatures as for GW. A recent one is Ref.[4]. In addition, it is better to know the basics of the PMT method (LAPW+LMTO method) [1]. Here is a small note for *GW*.

Green function

The Green's function $G(\mathbf{r}, \mathbf{r}', \omega)$ is the central quantity in the GW calculation. In the one-particle theory (mean-field theory/non-interacting case), the Schrödinger eq. is

$$(i\frac{\partial}{\partial t} - H_0)\psi(\mathbf{r}, t) = 0 \quad (1)$$

Here H_0 is the one-particle Hamiltonian which contains electrostatic potential plus exchange-correlation potential V_{xc} (here we don't care how it is given. This should be static but can be non-local as Fock term). When we have (unknown) source term $J(\mathbf{r}, t)$ instead of 0 in the right hand side of Eq. (1), we can calculate $\psi(\mathbf{r}, t)$ by multiplying inversion of the operator $(i\frac{\partial}{\partial t} - H_0)$. This is the (non-interacting) Green function $G(\mathbf{r}, \mathbf{r}', \omega)$ defined as

$$(i\frac{\partial}{\partial t} - H_0)G = \delta(t - t')\delta(\mathbf{r} - \mathbf{r}'). \quad (2)$$

As this shows, G is the inverse matrix of the operator $(i\frac{\partial}{\partial t} - H_0)$, (we pay attention to the boundary condition, especially for time-direction: retarded, advanced or time-ordered). Thus we can write $G = 1/(i\frac{\partial}{\partial t} - H_0) = 1/(\omega - H_0)$. In other words, the Green function G is the integration kernel in order to convert a differential equation Eq. (2) to an integral equation. This is the same as the case of Poisson equation (Laplacian) for the electrostatic problem; then $1/|\mathbf{r} - \mathbf{r}'|$ is the Green function for the conversion (then we calculate electro static field $\phi(\mathbf{r})$ for give source term $\rho(\mathbf{r})$).

In the case of electro static problem, we have two ways to solve the problem. One is the direct method to solve differential equation, the other is using such integration kernel. They are equivalent, but the latter is easy to handle and suitable for numerical calculations (under the assumption of "superposition law", that is, linear response). Although $G(\mathbf{r}, \mathbf{r}', t - t')$ contains time variables (to describe wave propagation), essentials are the same.

Green function: many body case

$G(\mathbf{r}, \mathbf{r}', \omega)$ is the quantity which is well defined even in the many-body perturbation theory. It is defined as the expectation value as $\langle 0|\hat{\psi}(\mathbf{r}t)\hat{\psi}^\dagger(\mathbf{r}'t')|0\rangle$ (this is not accurate. See literatures..). Here we use the second-quantized field-operator $\hat{\psi}$. This definition of G is a natural extension of the one-particle case.

However, there are differences in the G of one-particle case and many-body case. In the one-particle case, we have no interaction between electrons. Thus an electron added to the ground state (which is by filling electrons up the Fermi energy) is the eigenstate. An electron going on can not be an eigenstate, because of the correlation effect and exchange effect.

Let me explain the correlation effect. In contrast to this, one-particle state (=the quasi-particle(QP)) in the many-body theory can be not the eigenstate; it moves in the sea of other electrons and holes, which can be excited to other states, or an moving electron can move with polarized cloud of other electrons and holes. Its motion is affected by other electrons. It will lose energy gradually; the QP can have life time. This may be identified as the correlation effect.

Let me explain the exchange effect. This is due to the Fermi statistics. The electron, which you focus on, can not be distinguished from other electrons. (Thus mu-on can not feel this effect. G for mu-on in solids only contains pure correlation effect). From a point of view, this exchange effect is interpreted as a hopping effect, sudden jump from an electron to another electron (actually the Hartree-Fock theory gives zero effective mass at the Fermi energy for metallic systems).

We can include both of the effects in the GW calculation at the lowest order. Exactly speaking, these two effects are really mixed up. However, in GW (since GW is at the lowest order), these two are clearly separated.

The rigorous equation for G is

$$\left(i\frac{\partial}{\partial t} - H_0 + (\Sigma(\mathbf{r}, \mathbf{r}', t - t') - V_{xc})\right) G = \delta(t - t')\delta(\mathbf{r} - \mathbf{r}'). \quad (3)$$

Only the difference is adding $\Sigma(\mathbf{r}, \mathbf{r}', t - t') - V_{xc}$. In other words, all unknown effects are pushed into this term. Many body effect are pushed into (downfolded or projected into) the self-energy (=dynamical one-particle potential) $\Sigma(\mathbf{r}, \mathbf{r}', t - t')$. We don't treat many-body quantities such as $X(\mathbf{r}_1 t_1, \mathbf{r}_2 t_2, \mathbf{r}_3 t_3)$ directly in the GW calculations.

The GW method is how to give the $\Sigma(\mathbf{r}, \mathbf{r}', t - t')$ in the lowest order. Because of the long-range property of the Coulomb interaction, we need to have a special technique (not a simple perturbation). $\Sigma(\mathbf{r}, \mathbf{r}', t - t')$ is calculated as the product of $G \times W$, where G is (usually) just the non-interacting Green function given in Eq. (2). W is the dynamically screened Coulomb interaction calculated in RPA.

QSGW

Note that Eq. (3) completely cancels the effect of $V_{xc}(\mathbf{r}, \mathbf{r}')$ since it is included in H_0 . Thus this division is not meaningful if we can solve a problem completely. However, what we can do is only the perturbation. Quantities for H_0 is taken as the basic quantities. On it, we apply the perturbation. Equivalently we can have a division $H = H_0 + (H - H_0)$, where H is the many-body Hamiltonian.

In principle, we can start from any V_{xc} . However, as long as we use perturbation theory, we have to minimize the effect of $\Sigma(\mathbf{r}, \mathbf{r}', t - t') - V_{xc}$. The degree of freedom of the choice of V_{xc} is used to minimize it. This is the self-consistent perturbation theory idea for QSGW.

If $\Sigma(\mathbf{r}, \mathbf{r}', t - t') - V_{xc}$ give small effect, we can use the concept of the QP (independent particle picture) as the basis to evaluate physical quantities.

We have to determine best (or optimum) V_{xc} . How to do it? An idea is that we should determine it so that the size of $\Sigma(\mathbf{r}, \mathbf{r}', t - t') - V_{xc}$ should be smallest (how to measure the size?). Or, based on the Landau-Silin's theory, the QPs contained in G should be reproduced by H_0 . In anyway, this ends up with the self-consistent perturbation theory. For trial V_{xc} (and electron density), we have H_0 ; from H_0 we can calculate $\Sigma(\mathbf{r}, \mathbf{r}', t - t')$. Then we have to determine new V_{xc} in a method. This is repeated until converged. No unique choice for the method. Thus we test some possible ways and now we usually use a standard procedure as described in Ref. [2].

4 LDA/GGA calculations and Plots

Calculations are performed by following steps. These steps are detailed in the following sub-sections.

To identify a set of files used for a material we calculate, we use an extension to files. For example, files explained below are with extensions (only lower case allowed) of materials. For example, `ctrls.cu` and `ctrl.cu`. In this case `cu` is the extension. Any extension works. Other possible examples are `ctrls.lagao3`, `ctrl.wgantest1`, and so on.

1. Write crystal structure file `ctrls.*`, which contains crystal structure. It can be by hand, or convert it from POSCAR (in VASP). There is a tool to convert between POSCAR and `ctrls.*` (See `ecalj/StructureTool/README.txt` for the tool. we have `vasp2ctrl` and `ctrl2vasp`. Type them without arguments to see help.)
2. Generate `ctrl.*` from `ctrls.*` by a script `ctrlgenM1.py`,⁹
Here `ctrl.*` is the main control file which contains all required information to perform calculations. `ctrl.*` contains not only the content of `ctrls.*`, but also other information needed for calculations. If necessary, we edit the generated `ctrl.*` file before next step.
3. Check crystal structure. `lmchk` is to confirm the crystal structure (space-group symmetry and so on). `lmchk` is applied not to `ctrls.*` but to `ctrl.*`. `ctrls.*` never used in the following steps.

CAUTION for a known bug! : If a crystal structure is only slightly different from a structure with higher symmetry, the `lmchk` may give a wrong crystal symmetry. In such a case, you have to “standardize structure” by VESTA or some other tools. This occurs e.g., when we use a structure numerically relaxed by VASP.

4. Run `lmfa` (calculations of spherical atoms (MT sites) in the cell). It also calculates core eigenfunctions and valence electron charge to set up initial condition. Then we run main calculation of LDA by `lmf`. It repeats iterations, and end up with converged results in LDA. Main result (electron density satisfying self-consistency) is stored in restart file `rst.*` (binary file). It finished within a second.
5. Run LDA/GGA calculations. We can run the LDA/GGA calculation by `lmf` or `lmf-MPIK` (-MPIK means kpoint parallel version).
6. Post processing.
Plot energy band, DOS, PDOS, by running scripts. We can use scripts `job.band` for band plot (need `sym1.si` file (symmetry line for band plot)). We also have `job.pdos`, `job.tdos`, `job.fermi` and so on for DOS, PDOS, fermi surfaces. Since we use gnuplot to plot them, meanings of obtained data is apparently clear.

4.1 Write crystal structure file, `ctrls`

Let me show some samples of crystal structure files `ctrls.*`.

```
Cu: ~/ecalj/lm7K/TESTsamples/Cu/ctrls.cu
-----from here -----
% const da=0 alat=6.798
STRUC  ALAT={alat} DALAT={da}
        PLAT=  0.0 0.5 0.5    0.5 0.0 0.5    0.5 0.5 0.0
SITE    ATOM=Cu POS=0 0 0
-----to here -----
```

```
GaAs: ecalj/lm7K/TESTsamples/GaAs/ctrl.gaas
-----from here -----
#id = GaAs
%const bohr=0.529177 a=5.65325/bohr
STRUC
```

⁹`ctrlgenM1.py` exists originally at `ecalj/lm7K/` (`ctrlgenM1.py` was already copied to your `BINDIR=` defined in `ecalj/Install.ifort` in the installation).

```

        ALAT={a}
        PLAT=0 0.5 0.5 0.5 0 0.5 0.5 0.5 0
SITE
        ATOM=Ga POS=0.0 0.0 0.0
        ATOM=As POS=0.25 0.25 0.25
-----to here -----
SrTiO3: ecalj/lm7K/TESTsamples/SrTiO3/ctrls.srtio3
-----from here -----
%const da=0 au=0.529177
%const d0=1.95/au a0=2*d0 v=a0^3 a1=v^(1/3)
HEADER  SrTiO3 cubic
STRUC   ALAT={a1} DALAT={da}
        PLAT=1 0 0 0 1 0 0 0 1
SITE
        ATOM=Sr POS=1/2 1/2 1/2
        ATOM=Ti POS= 0 0 0
        ATOM=O POS=1/2 0 0
        ATOM=O POS= 0 1/2 0
        ATOM=O POS= 0 0 1/2
-----to here -----

```

Lines starting from '#' are neglected as comment lines. Lines starting from '% const' define variables and set values (in these cases, **da**, **alat**, and **bohr**, and so on). Then the variable **alat** is referred to as **{alat}**; in the cu case, **{alat}** means 6.798. Lines not start from '#' nor '%' are main content in **ctrls.***.¹⁰

Note that we have two tags of "categories" "STRUC" and "SITE". ("HEADER" tag is also; but it is just for user's memo shown in console output). These tags should start from the first column. Thus **ctrls.*** is divided into multiple "categories". In a category, we have "tokens" such as ALAT, DLAT, PLAT. These under STRUC category. ALAT+DALAT specify unit of length in this ctrl file. These are in a.u. (= bohr radius=0.529177Å).

The unit cell is given by PLAT (as noted, ALAT+DALAT as unit). In the above example of GaAs, three primitive cell vectors specified by nine numbers after PLAT=; they give three primitive vectors; PLAT1=(0,0,0.5), PLAT2=(0.5, 0.0, 0.5), and PLAT3=(0.5, 0.5, 0). DALAT is convenient to change lattice constant; but it is fixed to be zero here; thus no effect in this example.

Note that SITE category can have multiple ATOM tokens. The number of ATOM token under SITE should be the same as number of atoms in the primitive cell. In the case of GaAs; SITE contain multiple ATOM tokens. POS= just next to ATOM is taken as subtokens under ATOM token.¹¹ In cases, we specify such subtokens as SITE_ATOM_POS.

In the SITE category, we place atoms (MT names) in the primitive cell. In these cases we use defaults atomic symbol (MT names) for ATOM. POS is in the Cartesian coordinate (in the unit of ALAT+DALAT).

To test ecalj, you may make a test directory and copy a **ctrls.*** to your directory. If you have VESTA and ecalj/StructureTool/ installed, you can see its structure by

```
$ viewvesta ctrls.cu
```

(here \$ means command prompt).

NOTE: As written in ecalj/README, you have to install VESTA and **viewvesta**. Then set VESTA= at the top of ecalj/Structure/viewvesta, and make softlink to it. The command **viewvesta**(~/ecalj/StructureTool/viewvesta.py) generate **POSCAR_cu.vasp** first, then send it to VESTA. **viewvesta** also accept **POSCAR_cu.vasp** directly. Except names starting from **ctrl** and **ctrls**, **viewvesta** sends the name to VESTA directly. We need extension '.vasp' to recognize it is written in VASP

¹⁰For these variables, we can overlay values when we start programs. For example, 'lmf -vdalat=0.1 si' means that **alat** is recorded in save.si file.

¹¹This may looks slightly uncomfortable since the end of range of ATOM is not clearly shown; it end just at the next ATOM token or new category.

format. We have samples in `~/ecalj/StructureTool/sample`.

A tool `vasp2ctrl` converts POSCAR..vasp to ctrls.. “-help” show a small help.

- `ecalj/StructureTool/` is not tested well. Not believe it so much... We will fix it on your request. Another possible way is using `cif2cell`.

If you have a cif file, run

```
cif2cell foobar.cif -p vasp --vasp-cartesian --vasp-format=5
```

And convert POSCAR to ctrls. `cif2cell` is available from github.

In `ctrls.srtio3`, we use an expression `1/2` to give POS. We can use mathematical expression instead of values. Mathematical expressions such as “+ - */ sqrt(...)” are recognized. (instead of `3**2`, use `3^2`. Use parenthesis, and no space for an expression). We can use default atomic symbols (to check default atom name (MT name) type `ctrlgenM1.py --showatomlist`). Instead of such default symbols, we can use your own symbol as

```
SITE
ATOM=M1 POS=1/2 1/2 1/2
ATOM=M2 POS= 0 0 0
ATOM=O POS=1/2 0 0
ATOM=O POS= 0 1/2 0
ATOM=O POS= 0 0 1/2
SPEC
ATOM=M1 Z=38
ATOM=M2 Z=22
ATOM=O Z=8
```

. Then we have to add extra category SPEC where we set Z number. (You can use `Z=37.5` for virtual crystal approximation, however, you can not do it in ctrls now. Edit it in ctrl file. Such a procedure will be explained in other place.xxx)

This is an example for Antiferro NiO:

```
#id = NiO
%const bohr=0.529177 a=7.88
STRUC ALAT={a} PLAT= 0.5 0.5 1.0 0.5 1.0 0.5 1.0 0.5 0.5
SITE ATOM=Niup POS= .0 .0 .0
ATOM=Nidn POS= 1.0 1.0 1.0
ATOM=O POS= .5 .5 .5
ATOM=O POS= 1.5 1.5 1.5
SPEC
ATOM=Niup Z=28 MMOM=0 0 1.2 0
ATOM=Nidn Z=28 MMOM=0 0 -1.2 0
ATOM=O Z=8 MMOM=0 0 0 0
```

In this case, we define Niup and Nidn sites. These are recognized as Ni atom because of given Z number in SPEC. The subtoken `MMOM=Ms,Mp,Md,Mf...` re to specify number of magnetic moments (μ_B) for s,p,d,f channels (difference of up - down electrons within MT sites) as initial condition. In this case, we set `n(up)-n(down)=1.2` for Niup site for d channel. Even just one ATOM name is given by yourself, all ATOM in SPEC should be given (in this case SPEC for O should be given).

We can see other samples in `~/ecalj/1m7K/TESTsamples/*/ctrls.*`. (we also have a sample generator. See later section.) Note that ctrls file is jut in order to generate default ctrl file in the followings. Not from ctrls but from ctrl, we can start calculations. (thus ctrls is not needed if we prepare ctrl file directory).

It is possible to add `RELAX= 0 0 1` after `SITE_ATOM_POS`; this means structure relaxation along z-axis (also need to set DYN category given in Document/Manual/CategoryAndToken.org). but its defaults are given (but commented out) automatically in the ctrl file generated by the procedure described in the following section). We detail it in other place xxx.

After `ctrl.*` is generated as shown below, we can run a command `lmchk` to check whether crystal structure is correctly given or not. It finishes in a second. It shows symmetry information, and so on used in the calculation.

CAUTION!: Positions of atoms are not necessarily fixed by `ctrl.*` when you restart calculation with `rst.*` file, because atomic positions are read from `rst.*`. We need to pay attention when we use DYN option because `lmf` run may save relaxed atomic positions into the `rst.*`. As “`lmf -help`” shows, `lmf si --rs=1,1,1,0,0` can read atomic position from `ctrl` file.

4.2 Generate default ctrl from ctrls by ctrlgenM1.py

To run programs of `lm7K` (`lmfa`, `lmf`, `lmchk`) in `ecalj`, we need an input file `ctrl.*`, which contains not only structures but also other settings. To generate `ctrl.*` from `ctrls.*`, we have a command “`ctrlgenM1.py`” (written in python 2.x and call fortran programs(`lmfa`,`lmchk`) internally). Two steps required to complete `ctrl` file: (i) we give reasonable options when we run `ctrlgenM1.py`. Then (ii) we may edit the `ctrl` file afterward. In anyway, `ctrl` file is the starting point of calculations; `ctrls` is required just in order to generate `ctrl`.

At first, try `ctrlgenM1.py` without arguments. It shows help. To generate `ctrl` from `ctrls`, type

```
$ ctrlgenM1.py cu --nk1=8
```

Here `cu` specifies `ctrls.cu`. The option `--nk1=8` means the number of division of the Brillouin zone for integration. It means $8 \times 8 \times 8$ division. If we like to use $8 \times 8 \times 4$, we have to supply three arguments `--nk1=8 --nk2=8 --nk3=4`. The above command gives following console output.

```
$ ctrlgenM1.py cu --nk1=8
=== INPUT arguments (--help gives default values) ===
--help      Not exist
--showatomlist  Not exist
--nspin=1
--nk=8
--xcfun=vw   !(bh,vwn,pbe)
--systype=bulk !(bulk,molecule)
--insulator  Not exist !(do not set for --systype=molecule)

...
```

OK! A template of `ctrl` file, `ctrlgen2.ctrl.cu`, is generated.

As we see above, options which you specified are shown at the beginning of the console output (in this case `--nk1=8`). Others such as `--nspin=1` are default settings. If we like to perform spin-polarized calculations, we add other option ‘`--nspin=2`’ as

```
ctrlgenM1.py nio --nspin=2 --nk1=6
```

(NOTE: In the spin-polarized case, we need to set initial condition of size of magnetic moment at each atoms. Set it in `ctrls.*` as in the previous section, or edit `MMOM` of `ctrl` file (`MMOM=s p d f ...`) to be like `MMOM=0 0 1.2`). The `ctrlgenM1.py` generates `ctrl` file named as `ctrlgenM1.ctrl.cu`. To do calculations, copy it to `ctrl.cu` so that `lmf` can recognize it.

```
cp ctrlgenM1.ctrl.cu ctrl.cu
```

4.3 crystal structure checker: lmchk

Do `lmchk` to confirm that we can let `lmf` know correct crystal structure. It also shows crystal structure information, equivalent sites, site index and so on.

```
lmchk --pr60 cu (--pr# gives more informations if # is number)
```

Then it reads `ctrl.cu`. `--pr60` is an option of verbose. Bigger number gives more information.

- Lattice info, Space group symmetry operations (in `lmf` format), and their generators (these operations can be generated from a few of them.)

See `SYMGRP` in `Document/Manual/CategoryAndToken.org`. about how to represent the operations.

- Show atomic positions in ctrl file.
- Tabulate MT radius and distance between atomic sites.

(lmchk -help shows help, but difficult to see. Not need to read it first.)

lmchk is also shows atom (MT site) id (position and class(equivalent positions)). This is needed to interpret PDOS.

CAUTION for a known bug! See 3 in Sec.4.:

4.4 ctrl file

It is not necessary to look into ctrl file first, although some details are explained in the generated ctrl file. Please compare obtained results by lmf with those by other packages or literatures; let me know if you find something strange or your questions.

It is necessary to edit ctrl file to use full ability of lmf. For example, LDA+U, atomic position relaxation, core level spectroscopy, Change setting of default MTO and lo, better mixing procedure for stable convergence; higher accuracy, and so on.

But a few of ctrl file is easy to modify. Search these words and read explanations embedded in ctrl file.

(1)XCFUN

(choice of XC—it is not need to repeat ctrlgenM1.py). It is also possible to change number of k points for sampling, to modify crystal structure slightly, and so on; all things needed are in ctrl. It is not needed to repeat ctrlgenM1.py again.

(2)SO

To obtain correct dispersion around top of valence at Γ point for GaAs, we need to set SO=1 and NSPIN=2. QSGW calculation (by gwsc) do not allow this option now; Thus we run lmf (or lmf-MPIK) with such settings changed in ctrl file, after QSGW is converged.

lmf --input shows what can we write in ctrl file. But more than half are not for users, but for developers (or irrelevant now).

4.5 Run LDA/GGA calculations, and get convergence

Here we show how to get converged results from a ctrl file.

At first, we need initial guess of charge density. It can be given by a super position of atomic charge density. To obtain the charge density, we solve atoms first. It is by

```
$ lmfa gaas | tee llmfa
```

It takes just a few seconds. Here tee is a command of Linux. It keeps console output (standard output) to a file (llmfa in this case).

Then try

```
$ grep conf llmfa
```

. Then you see a key point that

```
conf:SPEC_ATOM= Ga : --- Table for atomic configuration ---
conf:  isp 1  int(P) int(P)z  Qval  Qcore  CoreConf
conf:   1 0      4  0      2.000  6.000 => 1,2,3,
conf:   1 1      4  0      1.000 12.000 => 2,3,
conf:   1 2      4  3     10.000  0.000 =>
conf:   1 3      4  0      0.000  0.000 =>
conf:   1 4      5  0      0.000  0.000 =>
conf:-----
conf:SPEC_ATOM= As : --- Table for atomic configuration ---
conf:  isp 1  int(P) int(P)z  Qval  Qcore  CoreConf
conf:   1 0      4  0      2.000  6.000 => 1,2,3,
conf:   1 1      4  0      3.000 12.000 => 2,3,
conf:   1 2      4  3     10.000  0.000 =>
```

```

conf:      1  3      4  0      0.000  0.000 =>
conf:      1  4      5  0      0.000  0.000 =>
conf:-----

```

This is an initial electron distribution, and how we divide core and valence. In this case core charge Q_{core} are “6 electron for s channel=1s,2s,3s and 12 electron for 2s and 3p”. Q_{core} is treated by frozen core approximation. See Sec.2.5 in Ref.???. Q_{val} means electrons for valence s,p,d channels. The valence channels are 4s,4p,4d,4f (if we set $E_{\text{H}}=s,p,d,f$) in this case The $\text{int}(P)z$ column is for local orbital. Thus we have 3d treated as local orbital. (ecalj allow add one local orbital per l .)

The isp index means spin (1 or 2), since $-\text{nspin}=1$ (when we invoke `ctrlgenM1.py`) for GaAs, no $\text{isp}=2$ exist. In summary we have 4s,4p,4d,3d,4f as valence. This means we use corresponding number of MTOs and local orbitals.

After `lmfa`, let us start main calculation.

```
$ lmf cu
```

In unix, we can save console output to `llmf` by `$ lmf cu | tee llmf`. As it starts iteration calculations, it shows similar output again and again (it is a little too noisy now). Then you end up with self-consistent result as

```

.....
it 8 of 30    ehf=  -3304.895853    ehk=  -3304.895853
From last iter    ehf=  -3304.895856    ehk=  -3304.895855
diffe(q)=  0.000003 (0.000007)    tol=  0.000010 (0.000010)    more=F
c ehf=-3304.8958531 ehk=-3304.8958529
Exit 0 LMF
CPU time:    7.024s    Mon Aug 19 02:03:19 2013    on

```

`it 8 of 30` means it stop at 8th iteration, although we set maximum number of iteration 30. Note that this number is given by `ITER_NIT=30` in `ctrl.cu`. `ehf` and `ehk` are the ground state energy in Ry. They are calculated in a little different procedure. Although they are different during iterations, it finally get to be the almost the same number. (But they can be slightly different even converged for large systems. But you don't need to care it so much).

NOTE: `ehk`:Hohenberg-Kohn energy, `ehf`: Harris-Faulkner energy.

“`grep diffe lllmf`” shows how the changed of total energy (and charges) during iteration. `diffe` mean changes of energy with previous iteration, `(q)` is for electron density difference as well. See also `save.*` file, which only show `ehk` and `ehf` obtained by each iteration.

“`grep gap lllmf`” shows how the band gap changes (in the usual setting), two same numbers per iteration are shown now.

Thus we do have ground state energy. Although output of `lmf` is long, most of all are to monitor convergence (we will shrink it). As long as it converged well, you don't need to look into it in detail. Eigenvalues are shown as

```

bndfp: kpt 1 of 4, k=  0.00000  0.00000  0.00000    ndimh = 122
-1.2755 -1.2008 -1.2008 -0.2052 -0.2052 -0.2052 -0.0766 -0.0766 -0.0766
-0.0174 -0.0174 -0.0174  0.1094  0.1095  0.1095  0.2864  0.2864  0.4170
 0.4170  0.4736  0.6445  0.6445  0.6445

```

This is at $k=0.00000\ 0.00000\ 0.00000$. (because of historical reason, two same `bndfp`: are shown in each iteration; two band path method). “`lmf cu | grep -A6 BZWTS`” shows the Fermi energy (for insulator, we see band gap). Deep levels which gives little dependence on k are core like levels. These are in Ry; zero level is not so meaningful (for convenience, it is simply determined from the potential at MT boundaries).

`rst.*` contains is the main output which contains electron density. `mix.*` is a mixing file (which keeps iteration history). When you restart `lmf` again, it read `rst.cu` and `mix.cu`. If you start from `lmfa` result, please remove them. We can do parallel calculation with `lmf-MPIK`, we can invoke it with `mpirun -np 8 lmf-MPIK cu`. It should give the same answer.

4.6 DOS, PDOS plot

We already have script to plot dos, band, and pdos from the result of lmf self-consistent calculations. We have scripts

```
job_tdos, job_band, job_pdos
```

. Look into these scripts, and then you see how to plot them.

For total DOS plot, it is better to check ctrl file; **BZ_TETRA=1**(this is default; thus make sure that **BZ_TETRA** do not exist or **BZ_TETRA=1**). In addition, it might be better to enlarge number of k point **NKABC** in ctrl file to have smooth curve. Then we do

```
job_tdos cu
```

This shows total DOS as The range of DOS and division of total DOS is given by DOSMAX

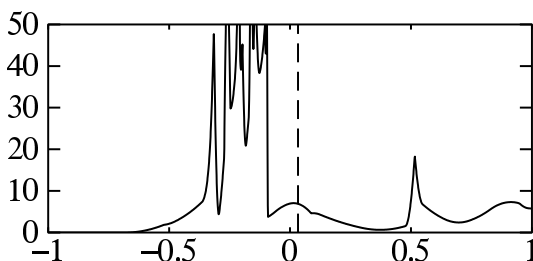


Figure 1: DOS(Cu)

and NPTS in ctrl. Edit tdos.cu.glt for gnuplot for your presentation. Please look into **job_tdos** file in your bin directory. It is a small script.

For PDOS plot,

```
job_pdos cu
```

It shows figures (number of figures are number of atoms in the cell) in gnuplot (they are written in the same position on X-window; move top one a little). The command **job_pdos** is a little time-consuming because we use no symmetry to distinguish all lm channels. (PDOS is not yet implemented for SO=1 case; spin-orbit coupling $L\hat{S}$ is added.) We can edit script of gnuplot (pdos.site*.glt) for your purpose. To plot again, run

```
gnuplot -persist pdos.site001.cu.glt
```

In principle, meanings of all data files are shown (see at the bottom of console output about lm ordering in a line), thus not so difficult to rewrite *.glt. For example, to plot eg and t2g separately. (NOTE: site id is shown by lmchk).

WARNING: Usually lmf and so on recognize options such as -v option. For example, 'lmf gaas -vnspin=2' or 'lmf gaas -vso=1'. This option changes values of variables defined in % **const** section. This is recorded in save.* file, and also shown at the top of console output. However, job_tdos and so on, do not yet accept these options. Thus we need to modify ctrl file without using -v option. Or you need to write these option to these command by hand (we will fix this problem in future.)

4.7 Band plot

For band plot, we have to set symmetry lines along which we plot eigenvalues. Collections **syml.*** are in ecalj/MATERIALS/. (On Jan2017, we prepared automatic generator of syml.* See Sec.4.7.1). Choose and modify one of them and rename it. I will gather other samples soon. 'BZ wikipedia' or something else will help you to interpret it.

To do band plot, we need **syml.cu** in your directory.

```
$ cp ~/ecalj/MATERIALS/Cu/syml.cu .
```

Then check **syml.cu**; it is

```

21 .5 .5 .5      0 0 0   L Gamma
21  0 0 0      1 0 0   Gamma X
0   (this is the terminator line)

```

We supply ten data for each lines (integer, three real, three real, two words). First line means, we calculate eigenvalues for \mathbf{k} points from $\mathbf{k}=(0.5,0.5,0.5)$ to $\mathbf{k}=(0,0,0)$. "L Gamma" are names of two end points (.5 .5 .5) and (0 0 0) in this case. These names are used in a gnuplot script for band plot (bndplot.isp*.glt). Second line means, we calculate eigenvalues for \mathbf{k} points from $\mathbf{k}=(0,0,0)$ to $\mathbf{k}=(1,0,0)$. 3rd line means calculation just stop here. Units of \mathbf{k} are in $2\pi/\text{ALAT}$ (or $2\pi/(\text{ALAT}+\text{DALAT})$ if DALAT exist.). A line starting from '#' is neglected (comment line).

To do band plot, run

```
$ job_band cu
```

. This is for both nspin=1 and nspin=2 (These scripts try to determine the Fermi energy first. You may skip it in cases (but need to change the script)).

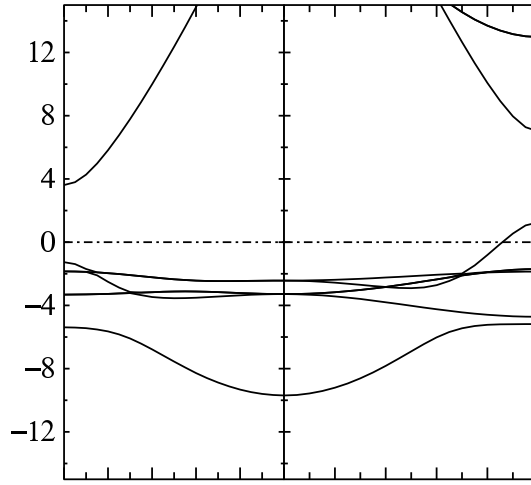


Figure 2: band plot(Cu)

4.7.1 syml generator

Jan2017, we added directory `ecalj/GetSyml`, which contains automatic generator of symmetry-line file `syml.*` from `ctrl.*`). Simultaneously, we plot a picture of BZ showing the symmetry lines by matplotlib. See `ecalj/GetSyml/README`. We use open softwares `spglib` and `seekpath` (these should be installed separately). We have to add acknowledgement to them (see README))when we use this generator.

4.8 Useful samples: `ecalj/MATERIAL/`

Not only `ecalj/lm7K/TestSamples` (some of them are by older version), We have a material database in `ecalj/MATERIALS/`. Move to the directory, and type

```
$ ./job_materials.py
```

Then it shows a help. You see

```

...
=== Materials in Materials.ctrls.database are:===
2hSiC 3cSiC 4hSiC AlAs AlN AlNzb AlP AlSb Bi2Te3 C
CdO CdS CdSe CdTe Ce Cu Fe GaAs GaAs_so GaN
GaNzb GaP GaSb Ge HfO2 HgO HgS HgSe HgTe InAs
InN InNzb InP InSb LaGaO3 Li MgO MgS MgSe MgTe
Ni NiO PbS PbTe Si SiO2c Sn SrTiO3 SrVO3 YMn2
ZnO ZnS ZnSe ZnTe ZrO2 wCdS wZnS
...

```

. For these simple materials (now 57 materials), input files can be generated, and run them automatically by a command `./job_materials.py` below. The ctrls are stored in `ecalj/MATERIALS/Materials.ctrls.database` in a compact manner (in addition, options passed to `ctrlgenM1.py` and options to `lmf-MPIK` are included). See `ecalj/MATERIALS/README` about how to add new material to it; it is not difficult. The command `./job_materials.py` gives `ctrls.*` for these materials from descriptions in the `Materials.ctrls.database`. And then it generates ctrl file by calling `ctrlgenM1.py` internally, and run `lmfa lmf-MPIK` successively (when no `--noexec`).

Try `./job_materials.py Fe --noexec`. (not `fe` but `Fe` as it shown above) at `ecalj/MATERIALS/`. Then it makes a directory `Fe/` and set `ctrl.fe` (also `ctrls.fe`) in the directory. Without '`--noexec`', it does calculation for `Fe` successively. As for `NiO` and `Fe`, we see that `./job_materials.py` gives `SPEC_ATOM_MMOM` in generated ctrls and ctrl files. (Look into `ctrls.fe`; we need `SPEC` section when we add `MMOM`.)

Try `job_materials.py GaAs Si`.

Then directories `GaAs/` and `Si/` are generated. See `save.*` files containing total energies iteration by iteration. Starting from `ctrl.*` in these directory, the command perform DFT calculations (Console output is stored in `llmf`, `save.*` gives total energies. `rst.*` contains self-consistent density, from which we can calculate energy bands and so on).

`./job_materials --all --noexec` generates ctrls and ctrl files of these materials. `./job_materials --all` do self-consistent LDA calculations for materials (it takes an hour or more. To change the number of cores for `lmf-MPIK`, set option `-np` (number of core). See help of `./job_materials` (type this without arguments).

To make band plot and so on for `Fe`, follow instructions already explained.

```
$ ./job_materials.py Fe (and need to type return)
(If you like start over, remove Fe/ under it first).
$ cd Fe
$ ./job_materials.py fe
  (but it might be better to do --noexec, and observe Fe/ctrls.fe and
Fe/ctrl.fe first. grep conf llmf shows the initial electron distribution).
$ cat save.fe (this shows total energies of each iteration. 'c ' at
the first column gives converged result. 'h ' is from atm file.)
  If it does not ends with 'c ...' line, something strange
occurs. see llmf (console out put of lmf is saved to llmf).
$ cp ../syml.fe .
$ job_band fe -np 4
  (As I said, this shell script do not yet accept
options to lmf. Look into the script).
  (This calculate fermi energy first for safe; it takes
some time)
$ job_tdos fe
$ job_pdos fe (as I said, this supress space-group symmetry, thus time consuming).
```

At the end of `job_pdos`, we show a help which `pdos` data is where (In `pdos` file, we have 26 numbers a line; first is energy, 2-26 are `pdos` for `s,p,d,f,g`; which is which are shown in the help). See `jobllmf` file also (it contains options to invoke `lmf`. This is shown in `save.*`. In principle, options in `jobllmf` should be passed to band plot and so on. But not yet implemented (it is not so difficult; I have to do it).

After doing `./job_materials foobar`, you may like move it back to original... In such a case, `git` works. At `ecalj/`, do

```
$ mv MATERIALS MATERIALS.bk
$ git checkout MATERIALS
```

Then you can see `MATERIALS/` is moved back to just downloaded one.

4.9 How to add spin-orbit coupling?

In the LDA level calculations, we can use

SO=0 (no SO), SO=1(LdotS), or SO=2(LzSz) schemes.
 However, SO=1 is not the non-collinear method
 (z-direction is assumed to be spin direction; in cases
 we may need to primitive cell so that the z-direction
 is the spin direction).

Within LDA, we have two possible ways.
 (A) Do LDA and/or QSGW with SO=0 first.
 Then apply the spin-orbit coupling by perturbation.
 or (B) do self-consistent LDA calculations with SO=1 (or =2).
 For semiconductors, we think you observe little differences.

Let me explain the case (A).
 After converged with nspin=1, create new directory and copy
 ctrl.gas, rst.gas, sigm.gas, QGpsi,
 to it. Then we set
 nspin=2
 METAL=3 (usually this is default)
 SO=1 (this is ldots calculation off-diagonal elements included).
 Q=band (we do not change potential.)
 in ctrl.gas.
 Then run
 >lmf gas >& llmf_SO
 You can see "band gap with SO" by
 > grep gap llmf_SO.
 Then you can see two same lines.
 VBmax = 0.101949 CBmin = 0.236351 gap = 0.134402 Ry = 1.82786 eV
 VBmax = 0.101949 CBmin = 0.236351 gap = 0.134402 Ry = 1.82786 eV
 (two lines per iteration is shown in metal mode).
 This is the band gap with SO as a first-order perturbation
 on top of the "QSGW without SO". When you use ctrl file generated by
 ctrlgenM1.py. You can do the above procedure with
 >lmf --rs=1,0 gas -vnit=1 -vso=1 -vnspin=2 -vmetal=3 --quit=band
 (--rs=1,0 read rst.gas but not write rst.gas. Run lmf --help.
 The switch -v (-vso=1 in this case) replaces so=0 with so=1.
 This is recorded in save.gas file).

For band plot, you can use the same procedure
 for the case without SO. (Look into the shell script job_band.
 You have to modify it so that
 '--rs=1,0 gas -vnit=1 -vso=1 -vnspin=2 -vmetal=3 --quit=band'
 is added to arguments for >lmf --band:sym1 ...).
 (--quit=band is not necessary if we like to renew eigenfunctions
 self-consistent. Anyway we expect little differences.)

\noindent QSGW with SO:\\
 For given sigm file, it is possible to attain self-consistency with SO=1(or=2) with keeping sigm (the
 However, this imply that Vxc is fixed in QSGW,
 it is not necessary better than the above procedure.

4.10 PROCASR (VASP format) generator

PROCASR mode for lmf (not yet in lmf-MPIK) Band weight decomposition. (Size of circles
 show the size of components. Superposed on band plot). See ecalj/MATERIALS/MgO_PROCAR/README,

May19.2014

4.11 efermi.lmf

This is generated by lmf(lmf-MPI), which is used in the job_band.

4.12 Effective mass calculation

New band plot and effective mass calculation (curvature):
(see sample at ecalj/MATERIALS/mass_fit_test/ and its README).

We now read sym1.* in lm7K/fp/bndfp.F. Thus job_band is changed
(nspin=1 or nspin=2 is automatically choosed by job_band command).
We do not use plbnd anymore. If necessary, you can modify
"writeband" subrouitne in lm7K/fp/bndfp.F by yourself.

Follow steps to get effective mass...

1.

New sym1 read label of k point. It is shown by the gnuscript
file "bandplot.glt". Type "gnuplot -p bandplot.isp1.glt" and so on,
when you like to remake band plot.

2.

New sym1 allow a special input suitable to determine effective mass
for semiconductor. An example of new sym1 is (this is a case of GaAs)

----- sym1 example start -----
#NOTE: ndiv2, ninit2 nend2 etolv etolc are for mass mode.
#ndiv qlleft(1:3) qright(1:3) llabel rlabel ndiv2 ninit2 nend2 etolv(Ry) etolc(Ry)
5 0 0 0 .5 .5 .5 Gamma L 257 1 32 0.1 0.01
5 0 0 0 1. 0 0 Gamma X
5 0 0 0 .75 .75 0 Gamma K 257 1 32 0.1 0.01
----- end -----

New feature is start from the next columns of symmetry points labels.

In this example, 257 means a line connecting 0 0 0 (Gamma) and .5 .5 .5 (L) is divided
into 256. Then we only calculate from the 1st point to the 32th point
among 257 points.

Then we see not only band*.spin*, but also
we have files such as "Band001Sym1001Spin1.dat"
files which contains eigenvalues for selected bands, that is,
we make a table of detailed plot of energy bands
whose energy E at Gamma (exactly speaking, at
left-end point), is evaltop-etolv (Ry) < E < econtop + etolc(Ry).

(As its head line shows, Band*Sym1*.dat file contains data
isym1,iq, ib,isp, QPE-EF, QPE-QPE(start), |q|, mass=2*2*(QPE-QPE(start))/|q|**2).
here QPE(start) is QPE at the left-end of sym line.)

This sample is only for Gamma-L and Gamma-K.

Let us use this example for GaA and run job_band.

(e.g., job_band gaas -np 2 -vnspin=2 -vso=1;

note that -vfoobar=xxx replace value of foobar with xxx in ctrl.gaas)

Then you can see not only bandplot.isp1.glt.

Then you can see usual energy bands plot in addition to the dense energy bands
superposed on it. The dense energy bands are controlled by
the part "257 1 32 0.1 0.01". In the current version, set all lines

of etolv and etolc are the same.

There are useful data, not only the eigenvalues in "Band001Sym1*Spin*.mass" files. But, anyway, we have to determine effective mass from the eigenvalues on the dense symmetry mesh.

3.

Least square fit by gnuplot.

We have an example is at ~/ecalj/MATERIALS/mass_fit_test

Look into ./job and run it.

See README in it.

For your convenience, we have dE/dk in the bnd*.spin* files.

This is useful to determine the Fermi surface.

See the efermi.lmf to read the Fermi energy.

4.13 density and $|\text{eigenfunction}|^2$ plot

We can generate density and $\sum_n |\Psi_{kn}(\mathbf{r})|^2$ by --density mode (after 18mar2016) Usages are

```
>mpirun -np 24 lmf-MPIK si --density
```

This is for density. We have smrho.xsf, and rho1mt*(1st components) and rho2mt*(2nd components).

```
>mpirun -np 24 lmf-MPIK si --density,iq=1,ib=5
```

iq=1 means Gamma point. See index shown by the console output of lmf. The index ib is the band index.

```
>mpirun -np 24 lmf-MPIK si --density,iq=1,ib=5,6,7
```

In this case we have sum for ib=5,6,7.

Use xcrysden as

```
>xcrysden --xsf smrho.xsf
```

Then open menu 'Tools→DataGrid'. Then plot isosurface and/or planes. For example, you need to set isovalue, and/or turn on 'display color plane' buttons.

4.14 LDA+U

There is a document of file://../BACKUP/MarksOriginalDoc/fp.html#ldaplusu. But I include all required things here. Usage is slightly changed. We have examples at ~/ecalj/MATERIALS/erasldau,GdNldau. I am going to add some samples.

To run LDA+U calculaiton, we need two extra steps in addition to usual LDA calculaitons:

Step.(1) Set three tokens in ctrl file. LDA+U type, Parameters U and J.

Step.(2) Prepare occnum.ext (initial condition of diagonal part of the density matrix).

Let us explain one by one.

Step.(1)

For the point (1), we have a new line in SPEC_ATOM sections. In ecalj/MATERIALS/erasldau, it is

```
IDU= 0 0 2 2 UH= 0 0 0.1 0.632 JH= 0 0 0 0.055
```

The SPEC_ATOM.IDU token tells lmf that no U is to be added to the s or p channels, but that a U is to be added to the d and f channels. IDU=2 specifies LDA+U functional style 2; this is the "Fully Localized Limit" described in Liechtenstein, PRB 52, R5467 (1995)). IDU=1 specifies the "Around Mean Field" functional (Petukhov, PRB 67, 153106 (2003)). U=0.1 Ry

is included on the d orbital, and $U=0.632$ is included on the f orbital. Additionally $J=0.055$ is put on the f orbital.

Density matrix:

It is stored in `dmats.ext`. In principle, the density-matrix is read from and written to the file `dmats.ext`. Thus, in principle, we need a `dmats.ext` as an initial condition. However, we usually use diagonal-only density matrix `occnum.ext` instead. If no `dmats.ext`, we read `occnum.eras` instead. We explain it afterwards. Two density-matrices (for each spin) are written to this file in a $(2l+1)$ by $(2l+1)$ block for each l block for which a U is defined. `dmats.ext` is an ASCII file which you can read, and it's quite useful to interpret what's going on. The diagonal parts are the occupation numbers and are the most important. It is in the spherical harmonics (not the real harmonics). The harmonics is defined in the `ecaljdetail.tex`.

Note that in the LDA case, complete information is contained in the density, stored in file `rst.ext`. In the LDA+U, the Density-matrix is stored in file `dmats.ext`. Thus complete information is contained in the combination of `rst.ext` and `dmats.ext`.

The density matrix is printed out in the standard output. It can be represented on the real harmonics. To check it, run

```
>lmf eras --quit=dmats
```

. This stops promptly without side effect. Then the symmetrized density matrix represented on the spherical harmonics and real harmonics are shown (we make symmetrization for the given SYMOPS in `ctrl` file).

Step. (2)

Usually it is too tedious to supply `dmats.ext` itself. Instead, you can supply an "occupation numbers" file `occnum.ext`, which is a starting guess for the density-matrix (its diagonal part). `occnum.ext` has one line of $(2l+1)$ numbers for the occupation numbers of the first spin, following by a line with the occupation numbers for the second spin. In the case of `ecalj/MATERIALS/erasldau`, we have `occnum.eras`.

Er has 11 f electrons, 7 of which go into the majority channel and 4 into the minority channel. There is some choice in which m states to fill and which to keep empty. A key point is that the self-consistent solution you end up with will depend on this choice. The ErAs test uses the following input file for `occnum.eras` :

```
0 0 0 0 0
0 0 0 0 0
1 1 1 1 1 1 1
0 1 1 1 1 0 0
```

The first and second lines are occupation numbers for the majority and minority d channel; the third corresponds to the majority f channel where all states are taken to be filled. The last line corresponds to the minority f channel. In this case, $m=-2,-1,0,1$ are filled and $m=-3,2,3$ are empty. As the script notes, different choices of starting occupation numbers lead to different self-consistent solutions. The one with the lowest energy is that which satisfies Hund's rule ($m=0,1,2,3$ filled and $m=-3,-2,-1$ empty).

It is possible to use 'real harmonics (see `ecalj/fpgw/exec/jobpdos` for definition)' instead of 'spherical harmonics'. As for `occnum.ext`, you can use

```
% real
0 0 0 0 0
0 0 0 0 0
1 1 1 1 1 1 1
0 1 1 1 1 0 0
```

. Here '`% real`' means on the basis of real harmonics. Check whether this is correctly supplied or not by `>lmf --quit=dmats eras`.

To run with spin-orbit coupling, set `HAM.SO=1`, then run `!grep IORBTM -A20 lmf`, here `lmf` is the standard output of `lmf`. Then you see orbital moments at 'IORBTM'.

NOTE for gwsc:

When you just like to generate initial condition for gwsc by LDA+U, you have to exit right after `sigm` generated (add `exit` right after `$echo_run echo == $ix 'iteration over=='` in

gwsc). Then gwsc stops right after `sigm` generated. Then you have to remove or comment out `IDU` (or set all zero) in `ctrl` file. Then run `lmf` together with obtained `sigm` file. This is because `sigm` contains LDA+U kind of effect. After this, check energy band (`jobband`) for check. And run usual `gwsc`— then you can perform QSGW with the LDA+U’s initial condition.

4.15 QSGW after LDA+U

We modified `lm7K/subs/m_rdctrl.F(feb2019)`, so that An option which makes “QSGW after LDA+U” easy to run. A line for atom section

```
IDU=0 0 0 12  UH=0 0 0 .7  JH=0 0 0 0
```

works in `ctrl` file. Here `12=10+2` instead of `2` means that we set `UH=JH=0` when `sigm.*` exits. Thus we can perform QSGW(without U term) calculaiton, after LDA+U calculaiton. This can be convenient for cases when we control the initial condition of QSGW.

4.16 partially occupied core-hole

See `file://../BACKUP/MarksOriginalDoc/fp.html#ldaplusu` around. Search the keyword `ldau` in `fp/lmfp.F` → `sudmtu.F` which read `occnum.gdn`.

Core-hole procedure is one of the important procedure. An example setting for Nd case is

```
-----
ATOM=Nd Z=60 R=3.00
C-HOLE=4f C-HQ=-11 P=6.5,6.5,5.5,5.2<-- this line is added
-----
```

1. C-HOLE=4f means that 4f core is to set “core hole”.
2. P=6.5,6.5,5.5,5.2 is to set principle (fractal) quantum number explicitly for valence electrons. To set this, try `lmfa nd|grep conf` first. The you can see the electronic configuration of an atom. Key point here is to set “5.2” for f channel. Thus 4f is treated as core and 5f as valence. Fractional parts (0.2 of 5.2) means the radial functon contains nodes as 5f and closer to the energy right on the change of 4f to 5f. If 5.9, it is closer to the enrgy almost have noded of 6f.
3. C-HQ means that corehole is . Thus we have only $14-11=3$ electrons for 4f (spherically occupied).
4. Run `lmfa`. If it does not show that `freeat (warning) atom not neutral`, it is the neutral atom. You can also chage the message tagged by `Add core hole`: Simultaneously you can control initial total valence charge by `Q=...`

Very recently (aug2019), I fixed a bug of `EFERMI`(fermi energy for GW calculaiton, given by `fpgw/main/heftet.m.F`) for the case C-HOLE is supplied.

5 How to run QSGW calculation?

In the QSGW, we calculate a *non-local exchange-correlation potential* $V^{xc}(\mathbf{r}, \mathbf{r}')$, by a procedure of GW calculation (very time-consuming part). Then difference $V^{xc}(\mathbf{r}, \mathbf{r}')V - V_{xc}^{LDA}(\mathbf{r})\delta(\mathbf{r} - \mathbf{r}')$ is stored into `sigm.*` file. The potential file `sigm` is a key to perform QSGW calculations as seen in Fig.6. The `sigm` contains static non-local potential $\Sigma_{QSGW} - V_{xc}^{LDA}$.

Then, we again do one-body calculation by `lmf` (or `lmf-MPIK`) where we add this `sigm` to one-body potential ; when we run `lmf` or `lmf-MPIK`(k-parallel mpi version), `sigm` is read and added to the one-body potential if we have `HAM_RDSIG=12` in `ctrl.*`. Thus this means that we replace $V_{xc}^{LDA}(\mathbf{r})\delta(\mathbf{r} - \mathbf{r}')$ with $V^{xc}(\mathbf{r}, \mathbf{r}')$.

This iteration cycle is performed by a script “gwsc” as we explain later on. (In the default setting of `ctrl1.*` file, `lmf` try to read `sigm.*` file as long as it exists. If not, do `lmf` or `lmf-MPIK` calculation.

To start QSGW calculation by `gwsc`, we need not only `ctrl.si`, but also another input file `GWinput`. Its template `GWinput.tmp` can be generated by the Step.2 as follows.

As a summary, you have to follow steps below in order to perform QSGW calculation.

- Step 1. Perform from the Step.1 thru the Step.4 (up to `lmfa`) in Sec.4.5. (as same as the case of LDA/GGA).

You don't need to perform LDA calculation in advance, since `gwsc` perform LDA/GGA calculation at its beginning. (It means that we start from the one-body Hamiltonian H_0 in LDA/GGA as initial condition. In cases, LDA/GGA give poor initial conditions for QSGW; in such a case, we may need another trick to prepare starting point.).

[Caution: We have to use the same `LMXA` (l in the expansion of eigenfunctions in each MT) for all the MT spheres. (This is due to historical reason; we may need to fix this.)] `xxx` need to explain `LMA xxx`

- Step 2. Run the script `mkGWIN_lmf2`.

The purpose of this script is to get `GWinput.tmp`. Other files generated are not used in the following stage.

- Step 3. Edit `GWinput.tmp` and save it as `GWinput`.

`GWinput` is the input file describing the computational conditions for GW calculation. Usually, the default setting gives reasonable results. To reduce computational time, we may use `lcutmx(atom)=2` for oxygen sites (this may be also other small atoms.). `grep lcut GWinput -A1` shows `lcutmx` for each atomic sites) These step 2. and step 3. are just only to get `GWinput`.

- Step 4. Run the script `gwsc`.

5.1 GWinput

In order to perform QSGW, one another input file `GWinput` (no extension) is necessary in addition to `ctrl.*`. Thus all input files for QSGW is just two files, `ctrl.*` and `GWinput`. A template `GWinput` can be generated by a script `mkGWIN_lmf2`. You may have to modify it in cases for your purpose.

Let us start from `ctrls.si`;

```
#id = Si
%const bohr=0.529177 a= 5.43095/bohr
STRUC
  ALAT={a}
  PLAT=0 0.5 0.5 0.5 0 0.5 0.5 0.5 0
SITE
  ATOM=Si POS=0.0 0.0 0.0
  ATOM=Si POS=0.25 0.25 0.25
```

. Do `ctrlgenM1.py si --tratio==1.0 --nk1=6` and copy `ctrlgenM1.ctrl.si` to `ctrl.si`. NOTE: the option `--tratio=1.0` means we use touching MT; this can be checked by `lmchk si`; since defaults is almost unity (`--tratio=0.97`), this is irrelevant, just to explain options.

We have to write `GWinput`. The default is given automatically by a command `mkGWIN_lmf2`;

```
$ lmfa si (lmfa is needed to do in advance).
$ mkGWIN_lmf2 si
.....
== Type three integers n1 n2 n3 for Brillowin Zone meshing for GW! ==
n1=
```

Then it pause and ask numbers. You have to type three numbers as 2+ return + 2+return+2 return.

```
== Type three integers n1 n2 n3 for Brillowin Zone meshing for GW! ==
n1= 2
n2= 2
n3= 2
2 2 2
```

...(skip)...

OK! GWinput.tmp is generated!

Generated file is GWinput.tmp; you have to copy it to GWinput.

```
$ cp GWinput.tmp GWinput
```

These '2 2 2' you typed is reflected in a section 'nln2n3 2 2 2' in GWinput. This means 2x2x2 (8 points in 1st BZ). You can edit it, and change it to e.g. 'nln2n3 4 4 4' if you like to calculate self-energy on dense BZ mesh 8x8x8.

The template of GWinput is usually not so bad. But it may give a little expensive setting (or not very good enough in cases).

NOTE: GWinput allows some options. Some newer points are

1. For the atoms without 3d involved, you can use lcutmx=2. Typically we use lcutmx=2 for oxygen.
2. GaussianFilterX0 0.0001 given in GWinput is effective to stabilize the convergence for metals.

5.2 Run gwsc script

Let us perform QSGW calculation. For this purpose, we use a script gwsc. We need to do lmfa in advance. Then do (not need to do lmf);

```
gwsc (number of iteration+1) -np (number of nodes) (id of ctrl)
```

If (number of iteration+1)=0, it gives one-shot calculation from LDA. But it is different from the usual one-shot; since it calculates off-diagonal elements of self-energy also, we can plot energy band dispersion. In cases (for usual semiconductors), it can give rather reasonable results in comparison with experiments from practical point of view.

This is an example of one iteration of QSGW cycle. (now a little different but essentially similar)

```
takao@TT4:~/ecalj/test1$ gwsc 0 -np 2 si
gwsc 0 -np 2 si
### START gwsc: ITER= 0, MPI size= 2, TARGET= si
--- No sigm nor sigm.$TARGET files for starting ---
---- goto sc calculation with given sigma-vxc --- ix=,0
No sigm ---> LDA caculation for eigenfunctions
      Start mpirun -np 2 /home/takao/ecalj/TestInstall/bin/lmf-MPIK si > llmf_lda
OK! --> Start echo 0| /home/takao/ecalj/TestInstall/bin/lmf gw si > llmf gw00
OK! --> Start echo 1| /home/takao/ecalj/TestInstall/bin/qg4 gw > lqg4 gw
OK! --> Start echo 1| mpirun -np 2 /home/takao/ecalj/TestInstall/bin/lmf gw-MPIK si > llmf gw01
OK! --> Start /home/takao/ecalj/TestInstall/bin/lmf2 gw > llmf2 gw
OK! --> Start echo 0| /home/takao/ecalj/TestInstall/bin/rdata4 gw_v2 > lrd data4 gw_v2
OK! --> Start echo 1| /home/takao/ecalj/TestInstall/bin/heft et > left et
OK! --> Start echo 1| /home/takao/ecalj/TestInstall/bin/hch knw > lch knw
OK! --> Start echo 3| /home/takao/ecalj/TestInstall/bin/hbas fp0 > lbas C
OK! --> Start echo 3| mpirun -np 2 /home/takao/ecalj/TestInstall/bin/hvcc fp0 > lvcc C
OK! --> Start echo 3| mpirun -np 2 /home/takao/ecalj/TestInstall/bin/hsfp0_sc > lsxC
OK! --> Start echo 0| /home/takao/ecalj/TestInstall/bin/hbas fp0 > lbas
OK! --> Start echo 0| mpirun -np 2 /home/takao/ecalj/TestInstall/bin/hvcc fp0 > lvcc
OK! --> Start echo 1| mpirun -np 2 /home/takao/ecalj/TestInstall/bin/hsfp0_sc > lsx
OK! --> Start echo 11| mpirun -np 2 /home/takao/ecalj/TestInstall/bin/hx0 fp0_sc > lx0
OK! --> Start echo 2| mpirun -np 2 /home/takao/ecalj/TestInstall/bin/hsfp0_sc > lsc
OK! --> Start echo 0| /home/takao/ecalj/TestInstall/bin/hqpe_sc > lqpe
OK! --> == 0 iteration over ==
OK! --> Start mpirun -np 2 /home/takao/ecalj/TestInstall/bin/lmf-MPIK si > llmf_gwscend.0
OK! ==== All calculation finished for gwsc 0 -np 2 si ====
```

Here `echo (integer)` is read in at the beginning of the code. To see it, please look into `gwscript` (`gwsc` is at `ecalj/fpgw/exec/` and copied to your `bin/` by `make install2`). In anyway, this console output shows calculations finished normally.

Now we get `rst.si` and `sigm.si` file which contains (static version of) self-energy minimis V_{xc}^{LDA} . What we did is the one-shot GW from LDA result; but note that we calculate not only diagonal elements but also off-diagonal elements.

We can write energy dispersion (band plot) in the same manner in LDA. To do it, we need `rst.si`, `sigm.si`, `ctrl.si`, `QGpsi`. (but `QGpsi` is quickly reproduced). After you have `syml.si` (e.g. in `ecalj/MATERIALS/`), Do

```
$ job_band si
```

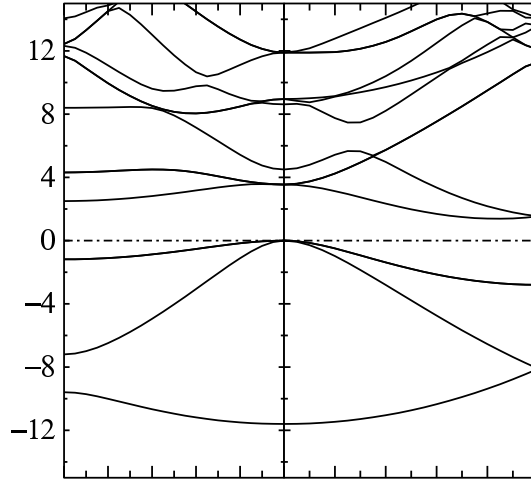


Figure 3: Si, one-shot GW with off-diagonal elements

You can observe large band gap as shown in the Fig.5.2. (To see it again, `gnuplot bnds.gnu.si -p`. All plots are in `gnuplot`, thus it is easy to replot it as you like).

We have QPU file (and also QPD for `spin=2`), which contains content of the diagonal part of self-energy. It will be explained elsewhere.

You can make total DOS and PDOS plot by

```
$ job_tdos si
$ job_pdos si
```

CAUTION:pdos plot is not allowed for `so=1`. (even `tdos-l` ask to t.kotani.)

To get final QSGW results, we have to repeat iteration until eigenvalues are converged. Note that total energy shown by console output `llmf` (and also shown in save file) is not so meaningful in the QSGW; we just take it as an indicator to check convergence. Let us repeat 5 iteration more. "`-np 2`" means one core to use.

```
$ gwsc 5 -np 2 si
### START gwsc: ITER= 5, MPI size= 2, TARGET= si
--- sigm is used. sigm.$TARGET is softlink to it ---
---- goto sc calculation with given sigma-vxc --- ix=,0
we have sigm already, skip iter=0
---- goto sc calculation with given sigma-vxc --- ix=,1
...(skeip here) ...

OK! --> == 5 iteration over ==
OK! --> Start mpirun -np 2 /home/takao/ecalj/TestInstall/bin/lmf-MPIK si > llmf_gwscend.0
OK! ==== All calclation finished for gwsc 0 -np 2 si ====
```

Note that we do skip 0th iteration (it is for one-shot from LDA) since we start from `rst.si` and `sigm.si` given by one-shot LDA. Thus we do just five iterations. Information of eigenvalues are in `QPU.(number)run` files. (for magnetic systems with `nspin=2`), we have `QPD.(number)run` also). Check it by `ls`;

```
$ ls QPU.*run
QPU0.run  QPU.1run  QPU.2run  QPU.3run  QPU.4run  QPU.5run
```

(These are overwritten when we again repeat `gwsc`; be careful.) Note that `QPU0.run` was old one when you did 1-shot GW from LDA at the beginning. In anyway `*.0run` are confusing files; remove them).

In order to check convergence calculations going well during iteration, do

```
$ grep gap llmf*
```

This shows how band gap changes in `llmf.*run` files. In metal cases, we need to compare QPU file, magnetic moment or `grep '[xc] save.*`; this shows end of lmf iteration. Energy is not so meaningful but can be indicator to convergence.

Let us check convergence of the QSGW calculations. For this purpose, it is convenient to take a difference of QPU(QPD) files by a script `dqpu`. These files are human readable. To compare `QPU4.run` and `QPU5.run`, do

```
$ dqpu QPU.3run QPU.4run
```

Then we see a list of numbers (these are the differences of values in QPU files). Then it shows at the bottom as

```
Error! Difference>2e-2 between:  QPU.4run  and  QPU.5run
:  sum(abs(QPU-QPD))= 0.05736
```

but you don't need to care it so much. You rather need to check the difference of values. I can say most of all difference (especially around the Fermi energy are) are almost 0.00eV or 0.01eV, we can judge QPEs are converged. If not converged well, you may need to repeat `gwsc` again. (when the size of two QPU files are different, `dqpu` stops.)

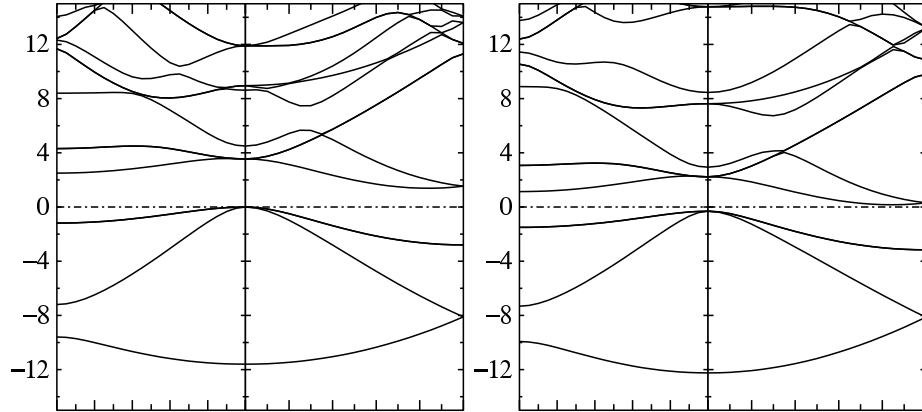


Figure 4: band plot(Si, QSGW one-shot test) and band(Si) (GGA)

5.3 Spectrum function: How to calculate $\langle qn | \Sigma(\omega) | qn \rangle$

We have an example at `ecalj/MATERIALS/SiSigma/`, where you can just type `job`. It calls a shell script `gwsigma`, which is just a modification of `gwsc` for spectrum function plotting. If you have `sigm.*`, it will automatically read it as in the case of `gwsc`.

By the script `gwsigma`, we calculate the diagonal elements $\langle \psi(\mathbf{q}, n) | \sigma_c(\omega) | \psi(\mathbf{q}, n) \rangle$. Thus we need to set \mathbf{q} and band index n for which we calculate. In addition, we need to set resolution of ω .

1. Set `<QPNT>` section. This section is to set the \mathbf{q} point, and band index for which we calculate the self energy. In addition, energy mesh for plotting is set.

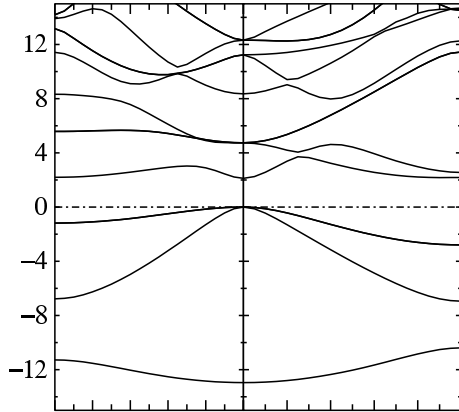


Figure 5: band(GaAs), QSGW (test case)

(A) Set q point set

If you set

```
*** all q -->1, otherwise 0; up only -->1, otherwise 0
      1          0
-----
```

You will have self-energy for all irreducible k points. This may be needed for $A(\omega)$.
or

You have to set all q points as

```
*** q-points, which should be in qbz., See KPNTin1BZ.
```

```
      3          <--- number of readin q point
1      0.0000000000000000      0.0000000000000000      0.0000000000000000 <--1st number is irrel
2     -0.5000000000000000      0.5000000000000000      0.5000000000000000
3      0.0000000000000000      0.0000000000000000      1.0000000000000000
-----
```

To know allowed q points on regular mesh point, run the command "mkGWIN_lmf2", then supply n_1, n_2, n_3 . The template of GWinput.tmp contains all possible q points. Edit it.

NOTE: Anyq option can allow you to specify any q points by shifted mesh technique.
(if necessary, but only for some special purpose).

(B) Band index set

It is specified by the section

```
*** no. states and band index for calculation.
```

```
2
```

```
4 5
```

means the self-energy for band index 4 and 5. Just two bands.

If you like to plot self energy from 1 through 8, use

```
*** no. states and band index for calculation.
```

```
8
```

```
1 2 3 4 5 6 7 8
```

If you need 17 bands for example, it should be 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 in addition to 17 (number of bands at the first line).

(C) energy mesh set

At the bottom of <QPNT> section, we have

```
-----  
*****  
0.01 2.0  
-----
```

Two real number should be supplied.

These are dwplot and omegamaxin, read in hsf0.m.F by a line

```
read (ifqpnt,*,err=2038,end=2038) dwplot,omegamaxin
```

dwplot (=0.01 Ry) is mesh for self energy.

omegamaxin=(2.0 Ry) means the range "-2 Ry to 2 Ry" for self-energy plot.

Note that imaginary part of Sigma is given as the convolution of ImW(omega) and the pole of (esmr in GWinpout gives energy smearing of the pole). Resolution for Im W (near omega=0) is b

I think that the resolution of self-energy is ~ 0.05 eV in the default setting.

This is because $\Delta E \sim \Delta E_{\text{esmr}} \sim 0.05 \text{ eV}$.

2 Run gwsigma. This will run

```
echo 4| mpirun -np 24 hsf0,
```

after dielectric function is calculated.

Then we have SEComg.UP (DN) files, Look for file handle, ifoutsec,

for the file in fpgw/main/hsf0.m.F to see format for the file.

(not hsf0.sc.m.F but hsf0.m.F). Search a line

```
open(ifoutsec,file='SEComg'//sss) (around hsf0.m.F L1052)
```

You can find that we use following lines to plot SEComg.*.

```
-----
```

```
write(ifoutsec,"(4i5,3f10.6,3x,f10.6,2x,f16.8,x,3f16.8)")
```

```
& iw,itq(i),ip,is, q(1:3,ip), eqx(i,ip,is),
```

```
& (omega(i,iw)-ef)*rydberg(), hartree*zsec(iw,i,ip) !,sumimg
```

This means we use energy in eV.

iw: omega index

itq(iq): band index specified by <QPNT>

ip: k point index specified by <QPNT>

is: spin index

q: q vector (cartesian in 2pi/alat)

eqx: eigenvalue in eV. (I think relative to the Fermi energy)

(omega(i,iw)-ef)*rydberg(): omega relative to the Fermi energy

hartree*zsec(iw,i,ip): Self energy. real and imaginary part.(complex, two values)

You can only repeat echo 4| mpirun -np 24 hsf0

when you change setting in <QPNT> section.

3 Example. There is an example MATERIALS/SiSigma/

```
plot 'SEComg.UP' u ($9):($10) w l,' u ($9):($11) w l
```

can give a plot for Re (Sigma_c(omega)) and Im(Sigma_c).

```
9th: energy in eV (omega(i,iw)-ef)*rydberg()
```

```
10th: real part Re hartree*zsec(iw,i,ip)
```

```
11th: imag part Im hartree*zsec(iw,i,ip)
```

4 To get integrated spectrum function (DOS), we need to superpose all the spectrum function (All q points and all band index). Be careful about the degeneracy (multiplicity) for each q points. You have to build it from SEComg file. To know the multiplicity, search following lines of keyword Multiplicity in the console output of qg4gw (lqg4gw).

Anyway, consider about "is it worth to do?" To confirm your result, use sum rule (sum of spectrum weight). And pay attention to the relation between real and imag parts (Hilbert transformation).

6 gwsc script to perform QSGW

6.1 outputs of gwsc

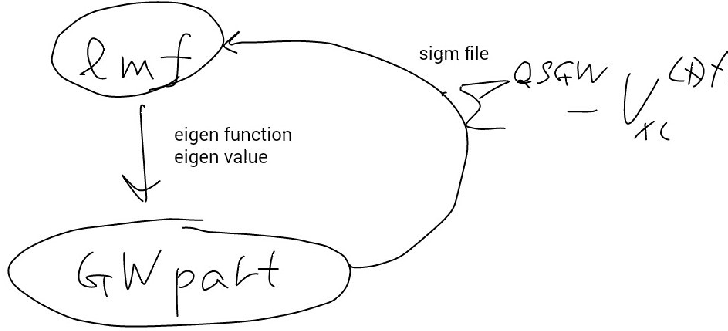


Figure 6: Shell script `gwsc` to perform QSGW contains an iteration loop to make `sigm` (and eigenvalues, eigenfunctions) self-consistent. The iteration loop is written in `ecalj/fpgw/exec/gwsc` (a bash script). Exactly speaking, we have to pass all the required data (not only eigenvalues and eigenfunctions, but also crystal structure, $\mathbf{q}+\mathbf{G}$ vectors, symmetry information and so on) to `GW part`. The purpose of the `GW part` is to calculate $\Sigma_{\text{QSGW}} - V_{\text{xc}}^{\text{LDA}}$.

When `gwsc` runs normally, it gives console output as follows. This is a case for `ctrl.gaas` for

```
>gwsc 10 -np 24 gaas
. Without arguments, typing gwsc shows usage as
>An example of usage: gwsc 5 -np 4 si, where 5 means 5+1 iterations
. We recommend you do look into the script gwsc. It uses run_arg, which is a special subroutine
of bash; but not so difficult to understand it. (In the followings, we assume /home/binx/ is
your bin directory at which we have all binaries for ecalj.)

### START gwsc: ITER= 10, MPI size= 24, TARGET= gaas
--- No sigm nor sigm.$TARGET files for starting ---
---- goto sc calculation for given sigma-vxc --- ix=,0
No sigm ---> LDA caculation for eigenfunctions
OK! --> Start echo --- | mpirun -np 24 /home/binx/lmf-MPIK gaas > llmf_lda
OK! --> Start echo 0 | /home/binx/lmfgw gaas > llmfgw00
OK! --> Start echo 1 | /home/binx/qg4gw > lqg4gw
OK! --> Start echo 1 | mpirun -np 24 /home/binx/lmfgw-MPIK gaas > llmfgw01
OK! --> Start echo --- | /home/binx/lmf2gw > llmf2gw
... (preparation stage ends here; start main stage) ...
OK! --> Start echo 0 | /home/binx/rdata4gw_v2 > lrdata4gw_v2
OK! --> Start echo 1 | /home/binx/heftet > leftet
OK! --> Start echo 1 | /home/binx/hchknw > lchknw
OK! --> Start echo 3 | /home/binx/hbasfp0 > lbasC
OK! --> Start echo 3 | mpirun -np 24 /home/binx/hvccfp0 > lvccC
OK! --> Start echo 3 | mpirun -np 24 /home/binx/hsfp0_sc > lsxC
OK! --> Start echo 0 | /home/binx/hbasfp0 > lbas
OK! --> Start echo 0 | mpirun -np 24 /home/binx/hvccfp0 > lvcc
OK! --> Start echo 1 | mpirun -np 24 /home/binx/hsfp0_sc > lsx
OK! --> Start echo 11 | mpirun -np 24 /home/binx/hx0fp0_sc > lx0
OK! --> Start echo 2 | mpirun -np 24 /home/binx/hsfp0_sc > lsc
OK! --> Start echo 0 | /home/binx/hqpe_sc > lqpe
OK! --> Start echo --- | mpirun -np 24 /home/binx/lmf-MPIK gaas > llmf_oneshot
... (this is the end of main stage) ...
== 0 iteration over ==
---- goto sc calculation for given sigma-vxc --- ix=,1
OK! --> Start echo --- | mpirun -np 24 /home/binx/lmf-MPIK gaas > llmf
OK! --> Start echo 0 | /home/binx/lmfgw gaas > llmfgw00
... (lines here omitted) ...
OK! --> Start echo 0 | /home/binx/hqpe_sc > lqpe
== 1 iteration over ==
... (lines here omitted) ...
== 2 iteration over ==
... (this repeat until ITER= 10(the first argument to gwsc) ) ...
```

This shows that `gwsc` invoke `lmf-MPIK`, `lmfgw`, `qg4gw`, ... successively. `echo 3|hbasfp0` means running a fortran program `hbasfp0` with the argument '3' from the standard input

(`read(*,*)` in fortran code). We can divide these successive calls to “preparation stage” and “main stage”. Preparation stage is just to prepare eigenfunctions and so on which are required for the “main stage” of GW calculation. At the end of “main stage”, we have the potential file `sigm`.

As it shows, console output are going to `l*` files.

6.2 Preparation stage of `gwsc`

At the end of this stage, we get required eigenfunctions, BZmesh data, and so on, which are required for the main stage. Note that `echo 0 | lmfgw` means supply an integer to the fortran program `lmfgw` from standard input by `read(*,*)`.

- **lmf-MPIK** (k-parallel version of `lmf`)
This is the one-body calculation for given `sigm.gaas`. At the beginning, we do not have `sigm.gaas`. In this case `lmf-MPIK` just perform LDA/GGA calculation.
- **echo 0 | lmf_{gw}**:
Get some small information files to start `qg4gw`. If you type just `lmfgw`, and observe what occurs. It shows a menu and pauses (asking you to supply an integer); then we supply 0 in this case. (If we do `echo 0 | lmfgw`, no pause occurs.)
- **echo 1 | qg4gw** : Get **k** points used in the *GW* calculations and the corresponding **G** vectors. And what is the irreducible k point (See console output of `qg4gw`. `gwsc` keeps it in `lqg4gw`).
Since we use the offset-Gamma method for BZ integration for $G \times W$, we need shifted mesh points to calculate *W* at offset-Gamma points. The **q** vectors of offset-Gamma method is in `Q0P` file. (If you have two points in `Q0P`, we see two shifted mesh points in addition to regular mesh points.) Remember that cutoff of **G** is given by `QpGcut_psi` and `QpGcut_cou` in `GWinput`. (Based on the experiences, we use smaller `QpGcut_cou` to reduce computational time. Explained in other section xxx).
- **echo 1 | lmf_{gw}-MPIK** : Calculate the LDA eigenfunctions, eigenvalues, and $\langle \psi | V_{xc}^{LDA} | \psi \rangle$ at the irreducible **k** points (shown at the bottom of output `lqg4gw` of `qg4gw`.)
- **lmf2gw**: store these data into `DATA4GW_V2` and `CphiGeig`, whose I/O is controlled by a key subroutine `gwinput_v2.f`.

6.3 Main stage of `gwsc`

We can start the main stage of *GW* calculation from these files;

`GWinput DATA4GW_V2 CphiGeig QGpsi QGcou Q0P QIBZ SYMOPS BZDATA HAMindex CLASS`;
these files contains eigenfunctions and so on in the manner of Eq.(17) of [2], eigenvalues and other required information. This is the starting point of the *GW* calculation.

- `GWinput` : computational conditions.
- `DATA4GW_V2` : Crystal structures and so.
- `CphiGeig` : Eigenvalues and Eigenfunctions
- `QGpsi` : **q** and **G** vector for the eigenfunctions(**q** means **k** in the previous section),
- `QGcou` : **q** and **G** vector for the Coulomb matrix
- `Q0P` : **q** points near **q**=0 instead of **q**=0 (offset Gamma points)
- `QIBZ` : irreducible **q** points (This is also contained in `BZDATA`).
- `SYMOPS` : point group operation
- `BZDATA` : **q** points date (and tetrahedron weights if necessary) for BZ integrals.
- `HAMindex` : Hamiltonian index, all required complex index for Hamiltonian of PMT method. (See the top of subroutine `write_hamindex` in `lm7K/subs/m_hamindex.F`. This is also in `fpgw/gwsrc/m_hamindex.F`. Identical files are in two different directory— it should be avoided in future.)

- **CLASS** : class information or atomic sites (equivalent sites).

With these files, we do the main stage as

- **rdata4gw_V2**: Read DATA4GW_V2 and so on, and decompose it into files required in the followings. And calculate PPOVL* files (overlap matrix of interstitial plane waves. Because of technical reasons some different types of PPOVL* with **q**-point index).
- **heftet** : Get the Fermi energy **EFERMI** by tetrahedron method. It is used in **hx0fp0**.
- **hchknw** : stores the number of required ω points along real-axis into **NW**. (NW is not essentially used, but is supposed to exist in the followings.)
- **echo 3|hbasfp0**: gives the product basis for Core exchange.
- **echo 0|hvccfp0**: gives the Coulomb matrix for the Core exchange.
- **echo 3|hfp0_sc** : gives the Core exchange part of the self-energy. ¹²
- **echo 0|hbasfp0**: gives the product basis.
- **echo 0|hvccfp0**: gives the Coulomb matrix v .
- **echo 1|hfp0_sc** : gives the exchange part of the self-energy.
- **echo 11|hx0fp0_sc** : gives the correlated part of the screened Coulomb interaction $W - v$.
- **echo 2|hfp0_sc** : gives the correlated part of the self-energy.
- **echo 0|hqpe_sc**: gather data and write down final results into **sigm**, **QPU**, **TOTE.UP** files.

6.4 Other functions (or scripts)

In addition to **gw_lmfh**, there are some other additional scripts and functions.

- **gw_lmfh** : The one-shot *GW* calculation. Lifetime(impact ionization rate) of QPs.
- **gwsc** : QSGW calculation explained here
- **epsPP_lmfh**, **eps_lmfh** : Dielectric function without or with local-field effects.
- run-mode 4 of **hfp0**: to plot the spectrum function $\Sigma(\omega)$. (need to be fixed again probably).
- **epsPP_lmfh_chipm** : non-interacting spin susceptibility. One-degree of freedom like Rigid moment approx. After it ends, you need to do **calj_nlfc_metal** and/or **calj_summary_mat** to get the full spin susceptibility.
- **genMLWF** : Wannier function and its matrix elements of the Screened Coulomb interaction.

7 Usage problems, QandA error messages.

1. Bandplot for FSMOMMETHOD/=0

Even when you use FSMOMMETHOD/=0 in GWinput for gwsc,

you need to set FSMOMMETHOD=0 (or comment it out) when you run job_band_nspin2.

¹²Correlation part due to cores is neglected. In future, we will switch to a version without PB for core part to reduce computational time and for numerical accuracy.

[If you run job_band_nspin2 with FSMOMMETHOD/=0, it make a shift (adding bias magnetic field).]

2. Note that ctrlgenM1.py automatically set this for --systype=molecule.

Then we have

TETRA=0

N=-1 #Negative is the Fermi distribution function W= gives temperature.

W=0.001 #W=0.001 corresponds to T=157K as shown in console

In addiiton, FSMOM (n_up-n_down) is needed (FSMOMMETHOD=1)if we have magnetic moment.

3. core>evalence message.

Ecore is grater than Evalence.

For save, we do not allow this.

Complare ECORE file and valence levels, shown in log file or console output.

4. If you see a error message from lmf (e.g., internally called in the gwsc script).;

Exit -1 rdsigm: Bloch sum deviates more than allowed tolerance (tol=5e-6)

You have to enlarge RSRNG so that lmf finsh normally.

5. Back ground charge and fractional Z.

You can use fractional numbers for ATOM_Z, and also can set valence charge by BZ_ZBAK (I removed BZ_VAL).

You see console out put, e.g,

```
"Charges:  valence    19.80000    cores      8.00000    nucleii   -28.00000
             hom background    .20000    deviation from neutrality:    0.00000
```

. This is a case with BZ_ZBAK=.2.

NOTE: at the first iteration, Charges: shows such as

```
Charges:  valence      8.00000    cores    20.00000    nucleii   -28.00000
             hom background    0.12300    deviation from neutrality: 0.12300
because of the initial condition by superposition of atoms. It show
deviation seems nonzero. But charge should be conserved from the
next iteration.
```

6. Not converged in metal. --->mixing may help

For example, if you try metal such as Bi2Sr2CuO6, it may fail at LDA/GGA level.

Then use ITER MIX=A2,b=.2. or something (.2 means it only mix 20% of output to give new input for next iteration). Then I see convergence. (b is the mixing parameter.

7. Use PZ or not.

If spillout of core is not so small (more then 0.05 or something.), it is better to use PZ(1o). Treat the core as valecne.

Bi4d is such a case. Maybe use PZ=0,0,4.9

8. Core treatment

See 10.1103/PhysRevB.76.165106 (Eq.35 and after).

Now I usually not use CORE2 (CORE1 only).

9. ERROR EXIT! rgwina: 2nd wrong 1 valence

This may be because you use wrong GWinput.

Back it up. And run mkGWIN_lmf2 (any n1 n2 n3 is fine).

10. Known bug.

Error occurs when system is anisotropic such as CuAlTe2.

Temporary fix is "Add token NPWPAD=100 in HAM category".

(guess of used APW fails (more than expected)).

CuAlTe2, CuGaTe2 cases.

11. Known bug

a little unstable when metal GGA, especially when we have large empty regions.

8 Cautions for usage

1. == not meaningful total energy in QSGW==

Total energy shown in QSGW mode in current version is not meaningful. (just treat as an indicator to convergence).

2. == Do we use VWN or GGA for QSGW? ==

In principle, QSGW results should not depends on VWN or GGA (XCFUN=1 or 103 in ctrl). But there is minor dependence, because

1. frozen core density.
2. core eigenfunctions.
3. radial basis functions
4. Slight numerical reason

(This is probably because Sigma-interpolation procedure But not exactly figured out yet → affect about 0.02eV as for band gap for GaAs.). In anyway, use VWN (HAM_XCFUN=1) as standard. And such technical things affects, 0.05 eV level of error for band gap.

3. EH and EH2 : For si, if EH and EH2 are the same, the following error occurred.

```
fexit,fexit2,fexit3 error retval=          -1
Exit -1 zhev_tk2: nev /=nevx something wrong.
```

The large EH, EH2 get to be meaningless. We usually use up to ~ 2 . (If you use very large EH such as $E \sim 10$, I am not so sure weather it is)

4. The options about the product basis within MT. (SeungWoo's memo)

```
<PRODUCT_BASIS>
tolerance to remove products due to poor linear-independency
0.100000D-02 ! =tolopt
```

When the product basis are made, we may have poorly linear independent basis. For example, one in the set $\{f_1, f_2, \dots, f_n\}$ would be almost give by a linear-combination of others. We need to make the linear-independent set. Therefore, after calculating the overlap matrix $\langle f_i | f_j \rangle$. We do diagonalization, then we remove eigenvectors corresponding to small eigenvalues than `tolopt`. See the **hbasfp0** command in **gwsc**

`lcutmx(atom)` = maximum l-cutoff for the product basis.

4 4 4 2 2 4 4

For $\phi_1 \times \phi_2$ case, $|l_1 - l_2| \leq l_{tot} \leq |l_1 + l_2|$. So 'lcutmx' changes the maximum cutoff for the l_{tot} . The order is the same as the order of atoms in the **ctrl** file.

atom	l	nnvv	nnc	!
1	0	3	3	
1	1	3	2	
1	2	2	1	

'atom' means the atom number identified in the **ctrl** file.

'l' is the angular momentum quantum number.

'nnvv' is the number of radial functions (valence) on the augmentation-waves.

'nnc' is the number of radial functions for core.

The latter two ones, 'nnvv' and 'nnc', will be understood more clearly if you see the following ones.

atom	l	n	occ	unocc	!	Valence(1=yes,0=no)
1	0	1	1	1	!	5S_p -----
1	0	2	0	0	!	5S_d
1	0	3	1	1	!	4S_l
1	1	1	1	1	!	5p_p
1	1	2	0	0	!	5p_d
1	1	3	1	1	!	4p_l

Above options are about the product basis set within MT (Valence).

'atom' and 'l' are explained above. 'nnvv' for 'atom = 1 and l = 0' was '3' so this case we have 3 basis ('n = 1, 2, 3')

'n' is the degree of freedom of the radial function, ϕ . 'n = 1' means ϕ , 'n = 2' means $\dot{\phi}$, and 'n = 3' means kind of $\ddot{\phi}$, which the dot above the letter represents the differentiation with respect to the energy. So 'n = 1 and 2' is related to the linearization of the radial function and 'n = 3' is the local orbital which is restricted in MT. The local orbital can be modified changing 'PZ' in the **ctrl** file. Finally, the number of the basis set which is needed for expanding eigenfunctions is $(l+1)^2 \times n$.

'occ' and 'unocc' mean that we use only ones that checked as '1', in other words we neglects '0' cases for making product basis. Be careful for confusion with name 'occ' and 'unocc'. These don't mean that occupied or unocc. When making product basis, $M = \phi_1 \times \phi_2$, 'occ' corresponds to ϕ_1 and 'unocc' to ϕ_2 . For example,

atom	l	n	occ	unocc	!	Valence(1=yes,0=no)
1	0	1	1	1	!	5S_p -----
2	3	1	0	1	!	4f_p

If the options are like the above, the product basis will be consists of $(\phi_1 = \phi_{atom=1,l=0}) \times (\phi_2 = \phi_{atom=1,l=0})$, $(\phi_{atom=1,l=0} \times \phi_{atom=2,l=3})$. As you can see, $(\phi_1 = \phi_{atom=2,l=3})$ is skipped.

In the **ctrl** file, 'EH' controls the l part. As for 'EH', (s, p, d, f) are used but **GWinput** file uses (s, p, d, f, g). 'EH' : HEAD part. 'GWinput' : contains TAIL part... need more explanation.

atom	l	n	occ	unocc	ForX0	ForSxc	!	Core (1=yes, 0=no)
------	---	---	-----	-------	-------	--------	---	--------------------

1	0	1	0	0	0	0	! 1S -----
1	0	2	0	0	0	0	! 2S
1	0	3	0	0	0	0	! 3S

Above options are about the product basis set within MT (Core).

'nnc' for 'atom = 1 and l = 0' was '3' so this case we have 3 basis ('n = 1, 2, 3')

Finally, for the convergence check, we can modify the following three things, (i) tolerance, (ii) lcutmx, and (iii) occ and unoccu.

== one show QSGW (not one-shot GW) ==

one-shot QSGW can be useful in cases.

As it contains off-diagonal part, we can resolve band tanglement problem in Ge (no band gap).

== Restart calculation in lda ==

lmf(lmf-MPIK) read rst.* in defaluts.

rst contains electron density.

If rst is already converged, it stops after two iteration.

rst contains atomic positions.

So, in order to read atomic positions change in ctrl,

Use options shown in lmf --help.

== Restart calculation in qsgw ==

To remove mixsigm* (mixing for sigm), maybe required.

== iteration check ==

First, watch console output of gwsc (do redirect to output file)

Need to check OK! signs arrayed on 1st columns.

gwsc iteration is time cosuming,

So we need to check calculations are normally going on or not.

Memory inefficiency.

Set 'KeepEigen off' an 'KeepPPOVL' off.

In fact, out code is still inefficient for memory usage.

grep gap llmf ---> minimum gap at mesh point.

see save.* ,or grep '[xc]' save.*

the end of iteration of lmf is shown as x or c.

(if failed, QPU file.

dqpu QPU.4run QPU.3run

As for usual semi-conductor, accuracy abou t0.1 eV is limit of current implementation.

Set vwn (xcfun=1) looks better (stable) for GW.

\$grep rms lqpe*

shows

... rmsdel=2.44D-04

```
... rmsdel=4.91D-03
... rmsdel=2.44D-04
... rmsdel=3.37D-04
```

If rmsdel is getting to be smaller, it is on convergence path.
 (but in magnetic cases, it may give be too good even not yet going to
 be converged..., because magnetic energy is so small)

grep diffe llmf ---> difference of energies of each iteration.

ehf (harris energy)
 ehk (Hohenberg kohn energy)

== emax cutoff for APWs. ==

We can not use so many APWs in current version,
 because of overcompleteness (this is because null vector within MTs),
 In anyway, use pwemax=3 as standard (test it with 4 or 5).
 To avoid failure of calculation, we may use smaller MT radius for
 alkali, and alkali-earth elements.
 In feature, I think we can introduce pseudopotentials for these atoms only.

== Check Used MTO

Near beginig of console output, what MTO you use is shown as: (GaAs case).
 sugcut: make orbital-dependent reciprocal vector cutoffs for tol= 1.00E-06

spec	l	rsm	eh	gmax	last term	cutoff
Ga	0*	1.13	-1.00	6.579	1.19E-06	1459
Ga	1*	1.13	-1.00	7.028	1.26E-06	1807
Ga	2*	1.13	-1.00	7.475	1.09E-06	2109
Ga	3	1.13	-1.00	7.920	1.06E-06	2637
Ga	0*	1.13	-2.00	6.579	1.19E-06	1459
Ga	1*	1.13	-2.00	7.028	1.26E-06	1807
Ga	2	1.13	-2.00	7.475	1.09E-06	2109
As	0*	1.18	-1.00	6.300	2.13E-06	1243
As	1*	1.18	-1.00	6.720	1.26E-06	1471
As	2*	1.18	-1.00	7.140	1.37E-06	1837
As	3	1.18	-1.00	7.558	1.05E-06	2229
As	0*	1.18	-2.00	6.300	2.13E-06	1243
As	1*	1.18	-2.00	6.720	1.26E-06	1471
As	2	1.18	-2.00	7.140	1.37E-06	1837

== gwsc cause error stop.

Have you ever changed MTO setting? Consistent with GWinput?

== QSGW for Fe.

It is better to use 3p as core. Furthermore, 3d+4d as valence is better.
 Thus we need to set PZ=0,3.9,4.5
 I also got aware that emax_sigm should be large enough ($4\sim 5$ Ry)
 to have smooth band dispersion. n1n2n3 can be 10x10x10.

== RSRNGE: enlarge RSRNGE ==

Use RSRNGE=10 or so (in cases, RARNGE=20 or more is required),
for large number of k points. Try and enlarge it if it fails with a
message "Exit -1 rdsigm: Bloch sum deviates more than allowed tolerance (tol=5e-6)".
We will have to make it automatic in future.

Detailed memo (for developers) is at the bottoem of ecalj/Document/BACKUP/MarksOriginalDoc/gw.htm

== QOP check

In cases, it is better to use QOPchoice=2 instead of default QOPchoice=1.
(For slabs, QOPchoice=2 may be better; need check more. In anyway,
it is problematic to use unbalanced k points for anisotropic cell).
See Copmuter Physics Comm. 176(2007)1-13).

=== When calculation in LDA level fails ===

when calculation fails in LDA level.

- (1) smaller MT
- (2) fewer PW. smaller pwemax.
- (3) core as semicore.

=== LDA+U ===

not yet written...

=== MAE by rotating crystal ===

(we have a sample at lm7K/TESTsmamples/MAEtest/, but only in GGA/LDA).

=== spin wave ===

J calculation.

=====

If not stable convergence in gwsc, try to set
mixbeta 0.5
(and/or mixpriorit 3 or something)
at the begining of sigma.

=====

cleargw (directory):

This command clean up up intermediate files under (directory).

This recursively into deeper level. Be careful, or edit it.

I use it as '>cleargw .''

Magnetic moment within MTs are shown as

charges:		old	new	
smooth		17.240314	17.240740	...
mmom		0.000024	-0.000010	
site	1	6.207135	6.206590	
mmom		1.062276	1.062991	<--- here
site	2	6.207115	6.206834	
mmom		-1.062323	-1.062958	<--- here
site	3	1.172718	1.172918	
mmom		0.000011	-0.000011	
site	4	1.172718	1.172918	
mmom		0.000011	-0.000011	

In this case, MTsite1 has 1.062991 and MTsite2 has -1.062958.

```
>grep 'lin mix' -A30 llmf
```

can take out this message (if console output is in llmf).

```
-----
ORBITAL MOMENT in pertubation:
-----
```

Try

```
>lmf nio --rs=1,0 -vso=1 --quit=band >llmf
```

After converged, try

```
>grep IORBTM -A20 llmf
```

Then llmf shows orbital moment in first order perturbation.

(Here --rs=1,0 read rst.* file but not change it. See >lmf --help.

--quit=band means quit just after band calculation.)

== EPS mode,

Check Im part of chi0 is smoothly damping at high energy (typically 1Ry or larger energy range). If there is some large Im part remains, something strange (usually due to orthogonality problem of eigenfunctions when you set low q).

Related source codes are in ecalj/lm7K/ .

A command ecalj/lm7K/ctrlgenM1.py can generate 'standard input file (ctrl file)' just from a given crystal structure file called as ctrls file.

Binaries are lmf and lmf-MPIK (MPI k-parallel version).

8.1 lmf -help

lmf -help show option of --rs=(five numbers); this let lmf know how to read atm.* file which is the initial atom file by lmfa.

9 Wannier function

We can generate Wannier functions (maximally localized Wannier Functions or similar) by a script **genMLWF**. It automatically perform cRPA calculation successively. (If it is not necessary, insert 'exit' in **genMLWF**, after it performs **lmaxloc2**).

Try to run examples in `ecalj/MATERIALS/Sample_MLWF/`. Read `README` in it. To run the script `genMLWF`, we need to get `GWinput` by editing `GWinput.tmp`. (`mkGWin_lmf2` contains default Wannier section). In addition, we have some settings (energy windows and so on). This is the example of the initial conditions for Cu case. 5 is the number of Wannier function. The most left one means ϕ index and the right one of it is $\dot{\phi}$ index. They are written in the `@MNLA_CPHI` file.

Then we can run `genMLWF`. After it finished, we can analyze its results. (if you don't need Wannier function plot, You can skip a line of `wanplot` in `genMLWF`. Then we don't need to set `vis_wan_*` options.)

9.1 lwmatK1 and lwmatK2

If you input the following command

```
>grep Wan lwmatK*
```

You will get the following results. (This case : Cu cases)

lwmatK1:	Wannier	1	1	24.644475	0.000000 eV	
lwmatK1:	Wannier	1	2	24.644576	0.000000 eV	
lwmatK1:	Wannier	1	3	25.471361	0.000000 eV	
lwmatK1:	Wannier	1	4	24.644575	0.000000 eV	
lwmatK1:	Wannier	1	5	25.470946	0.000000 eV	
lwmatK2:	Wannier	1	1	0.000000 eV	-21.263759	-0.000000 eV
lwmatK2:	Wannier	1	2	0.000000 eV	-21.263839	0.000000 eV
lwmatK2:	Wannier	1	3	0.000000 eV	-21.931033	-0.000000 eV
lwmatK2:	Wannier	1	4	0.000000 eV	-21.263839	-0.000000 eV
lwmatK2:	Wannier	1	5	0.000000 eV	-21.930702	-0.000000 eV

Wannier Branch now under developing (imported from T.Miyake's Wannier and H.Kino's).

A. make at `ecalj/fpgw/Wannier/` directory, and do make, and make install.

(need to check Makefile first). You first have to install `fpgw/exec/` in advance.

B. Samples are at these directories.

`MATERIALS/CuMLWFs` (small samples),

`MATERIALS/CuMLWF/`

`MATERIALS/CuMLWFs/`

`MATERIALS/FeMLWF/`

`MATERIALS/NiOMLWF/`

`MATERIALS/SrVO3MLWF/`

C. With `GWinput` and `ctrl.*`, run

```
>genMLWF
```

at these directories.

In `GWinput`, we supply settings to generate Wannier functions. (Sorry, not documented yet..)

D. After `genMLWF`, do

```
>grep Wan lwmatK*
```

then compare these with `Result.grepWanlwmatK`

These are onsite effective interactions (diagonal part only shown).

`*.xsf` are for plotting the Maximally localized Wannier functions.

Anyway, documentation on Wannier is on the way half.

Time consuming part (and also the advantage) is for effective interaction in RPA.

Look into the shell script `genMLWF`; you can skip last part if you don't need the effective interaction

10 ctrl file details

A ctrl file is usually generated from a ctrls file by the `ctrlgenM1.py` (a crystal structure file is not “ctrl” but “ctrls”). It contains self explanation. Here we give complementary explanations to it. Let us Look into a ctrl file. This is a head part of `ctrl.cu` generated by `ctrlgenM1.py`:

```
### This is generated by ctrlgenM1.py from ctrls
### For tokens, See Document/Manual/CategoryAndToken.org.
### Do lmf --input to see all effective category and token ###
### It will be not so difficult to edit ctrlge.py for your purpose ###
VERS    LM=7 FP=7          # version check. Fixed.
IO       SHOW=T VERBOS=35 TIM=2,2
        # SHOW=T shows readin data (and default setting at the begining of
        console output)
        # It is useful to check ctrl is read in correctly or not
        (equivalent with --show option).
        # larger VERBOSE gives more detailed console output.
SYMGRP find # 'find' evaluate space-group symmetry automatically.
        # Usually 'find is OK', but lmf may use lower symmetry
...

```

Note that # means comment lines. We can also use lines starting from `% const ...` to define variables and set constant.

We see “categories” such as `VERS`, `IO`, and so on. The beginning of categories are starting from the first column. Under categories, we have “tokens” such as `VERBOSE`. Thus we specify full name of token `VERBOSE` under category `IO` as `IO_VERBOSE`.

- `IO_TIM` is for debugging. It shows which subroutines are called and so on. Bigger number shows deeper subroutines.
- `SYMGRP` is a category without token under it; we set generators of space group (See explanation in previous paragraph). When we set `find`, it automatically calculate symmetry of crystal lattice. If we like to enforce symmetry, set some of generators which are shown by `lmchk`.
- We see `ctrls` is embedded in the ctrl by `ctrlgenM1.py`.

```
... (skip) ...
% const  da=0 alat=6.798
STRUC    ALAT={alat} DALAT={da}
        PLAT=  0.0 0.5 0.5  0.5 0.0 0.5   0.5 0.5 0.0
        NL=4 NBAS= 1 NSPEC=1
SITE     ATOM=Cu POS=0 0 0
... (skip) ...

```

`NL`, `NBAS`(number of `SITE`) and `NSPEC`(number of `SPEC`) are automatically added by `ctrlgenM1.py`. It is possible to deform unit cell by adding some optional tokens under `STRUC` category. Search `STRUC` in an old document (which may be still effective) `BACKUP/MarksOriginalDoc/lmto.html`. However, it is a little complicated. For new calculations, it is better to find some examples first.

- **SITE** category: As for MT sites, we have two categories. (1)SPEC(species) and (2)SITE(specify centers of atoms(species) in primitive cell). As for SPEC, we specify MTs(radius, Z, MTOs on it) appeared in the cell. These are defined subtokens under SPEC_ATOM=foobar (we have multiple SPEC_ATOM=foobar).

Then we place these MTs at SITE sections in the cell. At SITE, we specify atomic sites (What SPEC_ATOM is placed to positions by POS) in a primitive cell. We set POS= by direct form (Cartesian) but with the unit of ALAT+DLAT. Total number of SITE (number of tokens SITE_ATOM) is the number of atoms in the primitive cell. Setting POS= under SITE_ATOM=foobar means that we place MT named as foobar defined in SPEC_ATOM=foobar. In addition, we can set SITE_ATOM.RELAX, if you like to find relaxed structure (we simultaneously set DYN category) in LDA. As for relaxation, see `LaGa0_relax/ctrl1.lagao` example, and read DYN in Document/Manual/CategoryAndToken.org. The SITE_ATOM=foobar (with same foobar with different POS) are not necessarily equivalent with respect to the space group operation of a system. Thus SITE_ATOM=foobar are divided into “classes” which are connected by the operation. The lmf automatically judge “classes” (see also info by lmchk). Thus not need to specify it, but it may be better to check it. A sample is `lmchk lagao` at `~/ecalj/lm7K/TESTsamples/LaGa0_relax`

- **SPEC** category: In ctrls, we have not yet specified contents of SPEC; we have just given default symbols or only Z= when we use non-default names (shown by `ctrlgenM1.py -showatomlist`). The command `ctrlgenM1.py` adds default SPEC sections.

We have some SPEC_ATOM, under which we give subtokens such as SPEC_ATOM_R(MT radius), SPEC_ATOM_Z(nucleus charge), cutoff parameters of angular momentum, and so on. These SPEC_ATOM is refereed to in SITE.

An example of SPEC category is

SPEC

```

ATOM=Fe Z=26 R=1.70
      KMXA={kmtx} LMX=3 LMXA=4 NMCORE=1
      PZ=0,3.9,4.5
      EH=-1 -1 -1 -1 RSMH=0.85 0.85 0.85 0.85
      EH2=-2 -2 -2 RSMH2=0.85 0.85 0.85
      MMOM=0 0 2 0

      ATOM=... (then the similar block of ATOM= are repeated.)
      ...

```

Under the token ATOM=Fe, we have subtokens SPEC_ATOM_Z, SPEC_ATOM_R, and so on. Subtokens Z= is the nucleus charge and R= MT radius. Note that Fe is just a name to distinguish MT sphere in the cell. If you set SPEC_ATOM_Z=27, it is recognized as Co (since Z=27). LMX=3 is the maximum l of MTOs. Thus maximum l of MTO is l=3. The maximum of l to expand electron density and potential within MT is LMXA (in contrast to usual LAPW), we can use quite small LMXA such as LMXA=4. NMCORE=1 means we calculate core density without non magnetically-polarization. This can reduce computational confusion.

PZ is to set local orbital (if not, no local orbitals). EH and RSMH are to specify first set of MTOs.(We can check how local orbitals are set by lmfa explained in the next section). EH2 and RSMH2 are to specify second set of MTOs.

After PZ=, we have three numbers. These are numbers for s,p,d,f,g,... channels. Zero means not exist. You can use space or comma(,) as delimiter. Here not only the integer part of principle quantum number, but also the fractional part should be supplied (If PZ=0,3,4, it does not work.) Now PZ=3.9 for p and PZ=4.5 for d. This means we use local orbital for 3p, and local orbital for 4d (fractional parts (continuous principle quantum number) are large ~ 0.9 for core like orbital, and smaller for extended orbital ~ 0.3 or something. See Logarithmic Derivative Parameters in Document/BACKUP/MarksOriginalDoc/lmto.html. This is a little confusing, thus we will explain this in appendix. See Sec.10.1.

EH(damping factor), and RSMH (where the smooth Hankel function bent) determines MTOs (or its envelope function as a smooth Hankel function). We now set four numbers for them. Thus we set MTOs s,p,d,f with EH=-1 and RSMH=0.85. Our current test shows that RSMH is one half of R (that is, $0.85=1.70/2$, but minimum RSMH is 0.5) and not need to be dependent on s,p,d,f. (If LMX=2, s,p,d are allowed and no f MTOs.) EH is -1; not need to change except test purpose. In a similar manner, EH2 and RSMH2 for second set of MTOs are given. Just three numbers means these for s,p,d.

MMOM=s,p,d,f... gives initial condition of magnetic moment in μ_B (number of up-down electron).

In cases such as As, the local orbital given by default ctrl is responsible of rather deep core, and it is not need to be treated as valence electrons. In such a case, we don't need local orbital.

In the case of AntiFerro-II NiO, it contains two NiO in a primitive cell. Thus it is reasonable to have two SPEC_ATOM as Ni1 and Ni2, although subtokens under ATOM=Ni1 and ATOM=Ni2 (e.g. SPEC_ATOM_EH for them) are the same except initial condition of magnetic moment of MMOM=s,p,d,f... See example of NiO.

The minimum help of call Category.token_subtoken are listed with minimum explanation with

```
$ lmf --input
```

It gives a long output. But many of them are experimental and not need to manage them. A part of it is

```
Token          Input  cast  (size,min)
-----
... ..

STRUC_ALAT      reqd   r8      1, 1
  Scaling of lattice vectors, in a.u.
... ..
```

This is an minimum explanation of it. "reqd" means "required" (no default). r8 means it read with real number, 1,1 means that ALAT=xxx should contain one number minimum (max is also one) (See also STRUC_PLAT, and so on).

There are kinds of examples in ecalj packages. Please look into their ctrls.* and ctrl.* These are in lm7K/TESTsample/* and ecalj/CMDsamples. In addition, ecalj/MATERIALS contain many samples (need a command); see a later subsection.

As for what is shown in `$ lmf --input`, most of important tokens are already described in the ctrl file generated by `ctrlgenM1.py`. So, we don't need to care many options shown by it.

For QSGW calculation:

We need a setting in ctrl file to read `sigm` file (`HAM_SIGP`). It is simplified now, and not need to care it so much. As we set `RDSIG=12` in defaults, `lmf` read `sigm` file and add it to one-body potential as long as `sigm.*` exist.

NOTE for old users: We now set `SIGP[MODE=3 EMAX=9999.]` in ctrl file to read self-energy in `lmf` (or `lmf-MPIK`). This is because we use very localized MTOs (similar with the Maxloc Wannier). Our test shows reasonable results and this simplify algorithms. In my previous version, we asked you to use `SIGP[MODE=3 EMAX=2.0]` where `EMAX` is a little (0.5Ry) less than `emax_sigm`. If something strange occurs, try this setting).

- In principle, QSGW result should not depended on the choice of `XCFUN`. However, it can affect slightly. In our tests, it seems slightly better to use `VWN` (`XCFUN=1`) for QSGW calculations. (BUT need to check more...)

10.1 How to set local orbitals

As we stated, do `"lmfa |grep conf"` to check used MTO basis.

We have to set `SPEC_ATOM_PZ=?,?,?,?`

(they ordered as `PZ=s,p,d,f,...`) to set local orbitals.

`lmv7` (originally due to ASA in Stuttgart) uses a special terminology "continious principle quantum number for each `l`", which is just relatated to the logalismic derivative of radial funcitons at MT boundary. It is defined as

`P= principleQuantumNumber + 0.5-1/pi*atan(r* 1/phi dphi/dr),`

where `phi` is the radial function for each `l`. For example,

`P= n.5` for `l=0` of free electron (flat potential) because `phi=r^0`,

`P= n.25` for `l=1` because `phi=r^1`;

`P= n.147584` for `l=2` because `phi=r^2`; `P=, n.102416 n.077979` for `l=3,4`.

(Integer part can be changed). See Logarithmic Derivative Parameters in <http://titus.phy.qub.ac.uk/packages/LMT0/lmto.html#section2>

Its fractioanl part `0.5-atan(1/phi dphi/dr)` is closer to unity for core like orbital, but closer to zero for extended orbitals.

Examples of choice:

Ga p: in this case, choice 0 or choice 2 is recommended.

We usually use `lo` for semi-core, or virtually unoccupied level.

(0)no `lo` (4p as valence is default treatment without `lo`.)

3p core, 4p valence, no `lo`: default.

Then we have choice that `lo` is set to be for 3p,4p,5p.

(1)3p `lo` ---> 4p val (when 3p is treated as valence)

```

3d semi core, 4d valence
Set PZ=0,3.9
(P is not required to set. *.9 for core like state. It is just an initial condition.)
(2)5p lo ---> 4p val (PZ>P)
Set PZ=0,5.5
5.5 is just simply given by a guess (no method have yet
implemented for
    If 5.2 or something, it may fails
    because of poorness in linear-dependency. We may need to observe
    results should not change so much on the value of PZ.

(3xxx)4p lo ---> 5p val (we don't use this usually. this is for test purpose)
4p lo, 5p valence
Set PZ=0,4.5 P=0,5.5 (In this case, set P= simultaneously).
(NOTE: zero for s channel is to use defalut numbers for s)

Ga d: (in this case, choice 0 or choice 1 is recommended).
(0)no lo (3d core, 4d valecne, no lo: default.)
    Then we have choice that lo is set to be for 3d,4d,5d.
(1) 3d lo ---> 4d val (when 3d is treated as valence)
    Set PZ=0,0,3.9 (P is not required to set)
(2) 5d lo ---> 4d val (PZ>P)
    Set PZ=0,0,5.5
(3xxx) 4d lo ---> 5d val (this is for test purpose)
    Set PZ=0,0,5.5 P=0,0,4.5
    (NOTE: zero for s,p are to use defalut numbers )

If you like to read from atm.ga file instead of rst file(if exist).
You have to do lmf --rs=1,1,0,0,1, for example. See lmf --help
Becase rst file keeps the setting of MT0, thus change in ctrl is not
reflected without the above option to lmf.
=====

```


11 GWinput details

11.1 generate a template of GWinput

As in the previous section, we need two input files `ctrl.si` and `GWinput`. In principle, these two determines final results uniquely. A template `GWinput.tmp` is generated by `mkGWIN_lmf2`. Required files are

Input files

- `ctrl.si` : The control file for PMT method.

(Recently modified `mkGWIN_lmf2` runs `lmfa` internally. If you use older version, do `lmfa` in advance).

Output files

- `GWinput.tmp` : A file including computational conditions for the *GW* calculation. In addition, it specifies the **k** points for which you calculate the QP energy.

When you run `mkGWIN_lmf`, it asks you to supply three numbers for BZ integration as
== Type three integers n1 n2 n3 for Brillouin Zone meshing for GW! ==
n1=

Then you need to type a number e.g. as "2" for n1. Then you need to repeat it for n2 and n3 as

n1= 2

n2= 2

n3= 2

. These numbers specifies what **k** points in BZ is used for BZ integration (In this case, $2 \times 2 \times 2 = 8$ **k** point in the 1st BZ is used. Roughly speaking, we need $4 \times 4 \times 4$ (or $6 \times 6 \times 6$) to get band gap for Si and so on, with ≈ 0.1 eV accuracy.)

Then you have to edit `GWinput.tmp` and copy it to `GWinput`. We details the `GWinput` in later chapter.

We need to repeat `mkGWIN_lmf2` when you change MTO sections in `ctrl` file (adding PZ case, and so on).

11.2 overview of GWinput

(Because of historical reason, input file is different from `ctrl.*`).

The main input files is `GWinput`. This controls the setting of *GW* calculation. The file `GWinput` consists of structures as

keyword1 data1

keyword2 data2

...

In each lines, it consists of keyword and data. Data can be single or plural. As for keywords, upper case or Lowercase is not distinguished. All keywords should start from 1st column (no space at head). Order of lines are irrelevant. As for logical variable, you can use anything "true, yes, on, 1, T" for .true., and anything "false, no, off, 0, F" for .false.

Or we have "tag sections" in `GWinput` specified by `<PRODUCT_BASIS>`, `<QPNT>`, `<PBASMAX>`, `<QforEPS>`, and `<QforEPSL>`. (`<PRODUCT_BASIS>` is requires for all kinds of calculations. `<PBASMAX>` is optional. `<QforEPS>` and/or `<QforEPSL>` are required for epsilon mode). It is like

`<PRODUCT_BASIS>`

tolerance to remove products

```

0.100000D-02 ! =tolopt
lcutmx(atom)
3 3
atom 1
...
</PRODUCT_BASIS>

```

. In these tag sections, you have to keep format for its own (usually numbers are read by free format `read(*,*)`).

The fundamental readin routine for `GWinput` is a subroutine `getkeyvalue` defined in `gwsrc/keyvalue.F`. `getkeyvalue` is a general and convenient readin routine in full use of the f90 features. Read a head part of the file and try to do "grep getkeyvalue *.F" in `gwsrc/` or `main/` so as to see how to use it (test routine is `main/kino_input_test.F`.)

So the `GWinput` consists of three sections

1. General section
2. `<PRODUCT_BASIS>` section
3. `<QforEPS>`, `<QforEPSL>` section (only effective for dielectric function mode)
4. `<QPNT>` section (only effective for one-shot mode)
5. `<PBASMAX>` section (optional)

We will explain each by each in the followings.

11.3 General section

In general section, it looks like

```
! ##### From GWINO #####
n1n2n3      1      1      1 ! for BZ meshing in GW
QpGcut_psi   4.000 !(See unit_2pioa for unit) |q+G| cutoff for eigenfunction.
QpGcut_cou   3.000 !(See unit_2pioa for unit) |q+G| cutoff for Coulomb and W.
unit_2pioa off ! off --> a.u.; on--> unit of QpGcut_* are in 2*pi/alat
alpha_OffG   1.000 !(a.u.) Used in auxially function in the offset-Gamma method.
!emax_chi0   99999.000 !(Ry) emax cutoff for chi0 (Optional)
emax_sigm    3.000 !(Ry) emax cutoff for Sigma

! ##### FREQUENCIES from GWIN_V2 #####
dw           0.005000 !(a.u.) energy-mesh (bin width size) along real axis.
omg_c        0.040 !(a.u.) energy-mesh is twiced at omg_c
! coarser mesh for higher energy. Width get to be doubled at omg_c.
iSigMode     3 ! QSGW mode switch for gwsc. use =3.
niw           10 ! Number of frequencies along Im axis. Used for integration to get Sigma_c
! E.g. try niw=6 and niw=12
delta        -0.10D-05 !(a.u.) Broadening of x0. negative means tetrahedron method.
! used by hx0fp0. You get smeard x0 with abs(delta).
deltaw       0.020000 !(a.u.) Mesh for numerical derivative to get the Z factor
esmr         0.003000 !(Ry) used by hsfp0. Keep esmr smaller than band gap for insulators
! Poles of G^LDA are treated as if they have width esmr in hsfp0.
! Change esmr for metals. See DOSACC*---especailly around Ef.
GaussSmear on ! Gaussian or Rectangular smearing for Pole of G^LDA with esmr for hsfp0.
```

1. BZ integration.

n1n2n3 3 integers as N_1, N_2, N_3 (no default); They are ≥ 0 .

Brillouin Zone mesh for integration is determined by keywords **BZmesh** and **n1n2n3**. Current version only allow regular mesh point including Gamma point for $G \times W$. But not that **Chi_RegQbz** below allow you to use off-Gamma mesh for $W(\omega)$ (and dielectric function mode).

We usually take '4 4 4', '6 6 6' or '8 8 8' for GaAs. For metal such as Fe, '10 10 10' or more is better.

Chi_RegQbz (on or off)

Chi_RegQbz = on (default): Use regular mesh (including gamma) for eps calculation.

Chi_RegQbz = off : Use off-Gamma mesh (Not including gamma) for eps calculation.

(In cases, **Chi_RegQbz** off gives faster convergence as for **n1n2n3**; not only for GW, but also for dielectric functions **eps_lmfh**.)

2. Plane wave ($\mathbf{q} + \mathbf{G}$) cutoff

QpGcut_psi 1 real (no default)

QpGcut_Cou 1 real (no default)

unit_2pioa 1 logical (no default)

We have two cutoff for $\mathbf{q} + \mathbf{G}$. **QpGcut_psi** is the cutoff of $|q + G|$ for the IPW in the expansion of the eigenfunctions. **QpGcut_Cou** is for the IPW of the interactions v, D, W .

Its unit is specified by `unit_2pioa` ; "off" means unit in a.u. and "on" means in unit of $\frac{2\pi}{\text{alat}}$. (alat is length scale unit in ctrl.*).

Rule of thumb: `QGcut_psi` is a little (usually 0.5 or so) larger than `QpGcut_cou`. It becomes accurate if we use large `QpGcut_cou`. But it enlarge size of IPW(interstitial plane wave) part of Mixed product basis. For test, try 2.7, 3.2, 3.7 for `QGcut_cou` (and add 0.5 or 1 for `QGcut_psi`). Larger one is expensive.

We expand eigenfunctions in the Muffin-tin division of the space. See Eq.[?] in Ref.[?]; in the current *GW* implementation, we use very simple form of eigenfuncitons (not by the 3-component formalism in the [?]).

Thus the form of expansion is just related to the division of space; not directly related to the difference among LAPW, LMTO, and PMT.

3. Eigenfunctions within MTs (no parameters setting for them).

The radial functions (phi and phidot for each l), corresponding to the true parts, (= 2nd component in the 3-component formalism [?]), are automatically determined already in the one-body part of program `lmf-MPIK`.

4. Cutoff for used bands.

`emax_chi0`: 1 real (optional,default= ∞), in Ry

`emax_sigm`: 1 real (optional,default= ∞ ; We usually use 3 Ry).

`emax_sigm` is the maximum limit of the self-energy (measured from the Fermi energy). See the paper [?] which shows how the results are affected by `emax_sigm`. But in cases, small `emax_sigm` can give poor dispersion curve (slightly unnatural behavior) because of sudden cutoff by `emax_sigm`. However, we like to use smaller value to reduce computational time.

That is, larger is better, but expensive (And note that we simultaneously need to use empty spheres when we use large `emax_sigm`, as shown in [?],).

Generally speaking, accuracy less than $\sim 0.1\text{eV}$ (for bandgap) is allowance of current technique. Probably, it may be possible to have better accuracy, but it may ask us to repeat many calculations with changing conditions to confirm stability.

`nband_chi0`: 1 integer (optional,default= ∞)

`nband_sigm`: 1 integer (optional,default= ∞)

These specify how many bands you use in `hx0fp0` (for chi0) and in `hsfp0` (for sigma). Higher bands above them are neglected.

5. Energy mesh related parameters.

— new after march2016 —

`HistBin_dw`: 1 real (a.u.). Mesh width along real axis for $W(\omega)$.

`HistBin_ratio`: 1 real (a.u.).

`HistBin_dw` and `HistBin_ratio` specify real space bins which we accumulate imaginary part weight of polarization functions. The bins are (see `frhis` in `ecalj/fpgw/gwsc/m_freq.F`)

$$\omega_i = b * (\exp(a * (i - 1)) - 1), \quad (4)$$

where $[\omega_i, \omega_{i+1}]$ ($i = 1, 2, \dots, \text{nwhis}+1$) is the i -th bin. `HistBin_dw` is bin width at $\omega = 0$. The ratio ω_{i+1}/ω_i for large ω is $\exp(a)=\text{HistBin_ratio}$. This choice of getting coarser at high energy is because we think $W(\omega)$ around $\omega \sim 0$ gives most important

contribution to the GW approximation. If histogram bins are too wide, dielectric function can be less accurate, but results may be not so much affected.

In GW calculation, the Plasmon pole is important. It is determined not by the Drude weight; it usually gives small contribution to the Plasmon pole (for example, Si is described well by a Plasmon pole model, but Si has no Fermi surface). We expect that the GW results are not so sensitive to the choice of `HistBin_dw`, `HistBin_ratio` usually. We may use fine mesh when we plot quantities such as $W(\omega)$ near $\omega = 0$.

The `ecalj` gives $\bar{W}(\omega = 0) \sim 0$ for metal; where $\bar{W}(\omega)$ is the effective interaction averaged in the Γ -cell [10]. And $\bar{W}(\omega)$ get closer to v for larger ω .

Default value of them are given in `fpgw/gwsrc/m_freq.F`. They are

```
call getkeyvalue("GWinput","HistBin_ratio",oratio, default=1.03d0)
call getkeyvalue("GWinput","HistBin_dw",dw, default=1d-5) !a.u.
```

These are safer settings. Thus new `mkGWIN_lmf2` (`GWinput.tmp` generator) gives default lines in `GWinput` as

```
HistBin_dw 2d-3 ! 1d-5 for metal !(a.u.) bin width along real axis at omega=0.
HistBin_ratio 1.08 ! 1.03 for check. frhis(iw)= b*(exp(a*(iw-1))-1),
  where a=ratio-1.0 and dw=b*a
```

. This is given in the `conv2gwinput.F`.

— old version. before 14march2016 —

`dw` : 1 real (a.u.). Mesh width along real axis for $W(\omega)$.

`omg_c` : 1 real (a.u.).

`dw` and `omg_c` determines ω mesh along real axis to calculate $W(\omega)$...

(WARNING! Some of my examples may show as if they are in "(Ry)". But they are Wrong!)

`delta` : 1 real (a.u.). We usually use very small number as -1d-8 for `gw_lmfh`, `eps` mode and so.

xxx does this really make the stabilization? xxx This is the size of δ in denominator of Π (EQ.xxx). But (I think) we can (or can not) use it so as to make broadening for theoretical test (maybe not exactly corresponding to δ). or when you make calculation stabilized (Takao need to check this point again xxx!)

[Old note. Need check xxx: In `gw_lmf`, it is used for broadening of x_0 when it call `hx0fp0`. Then `delta` is δ is EQ.32. The sign of `delta` is just used as a flag whether you use the tetrahedron method of dielectric constant [11] or not; minus sign means "Use the tetrahedron method for D "; plus sign means you do it by simple sum. You can usually use this default setting. But it might be possible to use a larger value to smear the fine structures on the energy-dependence of W in cases. This might be necessary if W is so energy-dependent and `dw` is not so small to resolve the structure—but I don't know.]

`niw` : 1 integer.

Number of integration points along the imaginary axis(FIG.1) to get Σ_c . See routines `wint*` called from `sxcf*.F`, which is called from the main routine `hsfp0.m.F` (or `hsfp0.sc.m.F` in the QSGW case). The integration points are $i\omega'(n) = i(1/x(n) - 1)$, where $x(n)$ is the usual Gaussian-integration points for the interval $[0,1]$. In addition,

we give the special analytical treatment for the peaky part at $\omega' = 0$. Our tests show `n1w=6` for Si is good enough for 0.01 eV accuracy. The convergence as for `n1w` is quite good. This integration scheme has been developed by Ferdi Aryasetiawan. The number of points should be the one of 6,10,12,16,20,24,32,40, or 48. It is because we use a subroutine `gauss` in `/gwsr/mate.F` prepared by Ferdi. We will replace better one in future. See II-F in Ref.I.

`GaussSmear` : 1 logical

`esmr` : 1 real (Ry). Used by `hsfp0` (and `hsfp0.sc` for QSGW).

Poles of the Green function G^{LDA} are treated as if they have width `esmr` in `hsfp0`. If `GaussSmear` is on, each pole of G^{LDA} is smeared by a Gaussian function with $\sigma = \text{esmr}$ in the calculation of `hsfp0`. If `GaussSmear` is off, we assume rectangular smearing for the poles. Usually it is necessary to take rather smaller value than band gap for insulators. Try to use 0.003 or so in the case of Si and `GaussSmear=on`.

For metal, this `esmr` is somehow related to how we capture the Fermi surface; In principle, we have to take the limit $n1n2n3 \rightarrow \infty$ and `esmr` $\rightarrow 0$). However, we may inevitably use some finite `esmr` to make calculations converged.)

`deltaw` : 1 real (a.u.) only for one-shot case.

`deltaw` is the interval for the numerical derivative $\frac{\partial \Sigma(\omega)}{\partial \omega}$ in EQ.8. We calculate $\langle \psi^{\mathbf{k}n} | \Sigma(\epsilon^{\mathbf{k}n} + \text{deltaw}) | \psi^{\mathbf{k}n} \rangle$ and $\langle \psi^{\mathbf{k}n} | \Sigma(\psi^{\mathbf{k}n} - \text{deltaw}) | \psi^{\mathbf{k}n} \rangle$ in addition to $\langle \psi^{\mathbf{k}n} | \Sigma(\epsilon^{\mathbf{k}n}) | \psi^{\mathbf{k}n} \rangle$. From these values, we can calculate two Z (or second-derivative of $\Sigma(\omega)$), as shown in SECU. It will help to see whether the used `deltaw` is O.K. or not.

6. Offset-gamma point.

`QOPChoice 0` : 1 integer

`QOP_Choice` gives how to determine the offset gamma points. Initially we take them as

1: —q— is ten times smaller than regular mesh.(default)

2: —q— is average in the Gamma cell (cell of BZ including Gamma point).

Then we choose only inequivalent \mathbf{q} points based on the point group symmetry. Obtained offset gamma points is given in `QOP` file.

`alpha_offG` : 1 real (a.u.)

`alpha_offG` corresponds to α in EQ.48. `alpha_offG=1d0` is usually good in the sense that it seems to be almost a limit at $\alpha \rightarrow 0$. So you can usually fix it as `alpha_offG=1d0`, and check the convergence as for `n1n2n3`.

7. core orthogonalization (default=off)

`CoreOrth` 1 logical — recently, this option is not maintained — Better to use local orbital instead, so that core charge do not spill out.

If this is on, we enforce cores orthogonalized to valence ϕ and $\dot{\phi}$ (these appear in II-C in Ref.I). This procedure enforces the correct orthogonal condition, thus we have correct behavior for the dielectric function at $\mathbf{q} \rightarrow 0$. However, it may deform core functions too much, especially in the case of shallow 3d (or maybe 4d) cores. So we don't recommend use this option, even though then the orthogonality condition is somehow broken. Anyway you can check whether it affects to results or not by this switch.

8. QP self-consistent GW.

`iSigMode` 1 integer (no default).

This is required for QSGW calculation by the script `gwsc`. We have some possible ways to make GW self-consistent (how we determine V_{xc} from calculated $\Sigma(\omega)$). We now mostly use `iSigMode=3`.

- 3: Use $\text{Re} \frac{\Sigma_{nn'}(\epsilon_n) + \Sigma_{nn'}(\epsilon_{n'})}{2}$ (mode-A in [?]).
 1: Use $\Sigma_{nn'}(E_F) + \delta_{nn'}(\Sigma_{nn'}(\epsilon_n) - \Sigma_{nn'}(E_F))$ (mode-B in [?]).
 5: Use $\delta_{nn'}\Sigma_{nn}(\epsilon_n)$ (Eigenvalue-only self-consistency, keeping the eigenfunctions as given)).
 See /gwsrsc/sxsf_fal2.sc.F, which is called from the main routine hsfp0_sc (this is the routine to calculate self-energy)).

9. Others

KeepEigen 1 logical (default=on)

These are for memory usage. When KeepEigen is on, eigenfunctions (Eigen) are kept in memory during calculation. If you have not enough memory in your machine, use them off. Then you can save memory usage. However, then we may have too frequent access to files. So %CPU might get lower. Be careful to use these options.

Verbose 1 integer (default=0) If 0, it gives minimum standard output. If 40 or higher, it shows too much output. (these verbosity control is not well-organized yet).

10. **LFC@Gamma**, **EIBZmode**, **multitet** are for test purpose.xxx

11.4 <QPNT> section

(only for one-shot GW. Not suitable to make band plot in BZ.) This section is to specify the q points and bands index for which you calculate the QP energies (QPE). An example is

```
<QPNT>
--- Specify the q and band indeces, for which we evaluate the self-energy ---
*** all q -->1, otherwise 0; up only -->1, otherwise 0
          0          0
*** no. states and band index for calculation.
          3
    15 16 17
*** q-points, which should be in qbz. See KPNTin1BZ.
          2
    1 0.0000000000000000 0.0000000000000000 0.0000000000000000
    2 0.2886751358562925 -0.5000000000000000 0.0000000000000000
    3 0.0000000000000000 0.0000000000000000 0.3073140749846343
</QPNT>
```

Numbers are read by free format read(5,*), thus the numbers should be separated by space. At the next line to the first ***, you have to give two numbers used as flags. Both of them takes 0 or 1. 1st one is whether you calculate QPE for all q points (in IBZ) or not. If it is 1, you calculate QPE for all q. If it is 0, you calculate them only for q points specified within this file. In the case of metal where you want to calculate the Fermi energy for QPE, you need to calculate all the eigenvalues somehow above the Fermi energy (If you put 1, it is safer but too time-consuming). The second number is whether you calculate QPE for both spins or not. It is usually 0. In the case of antiferro material, it should be 1.

From the next line to the second ***, you have to specify the states for which you calculate the QPE. In this example, you calculate the 3 bands of QPE for 15th, 16th, and 17th eigenfunctions (they are ordered from the bottom).

From the next line to the third ***, you have to specify the q points. The first numbers of each line are dummy. In this case, you calculate QPE for two q points. The third q point is neglected because 2 is given at first.

When you generate GWinput.tmp, you see all the possible q points are listed (these q points should be a part of the regular mesh points).

In the QSGWmode (gwsc), this section is neglected (then we calculate all QPE on regular mesh points); so its hsfp0_sc part is quite expensive (usually it takes time more than hx0fp0).

Additional Note —————

`QPNT_nbandrange` num1 num2 (two integers).

This override setting in `<QPNT>`. (I think this switch may still work, but not checked recently).

`AnyQ` on (default is off)

If this is on, you can specify any Q point which is not on the mesh point. For the purpose, we need to prepare eigenfunctions at extra **k** points. But it is automatic. In order to make the computation efficient. Even in this case, from the computational view, it is better to choose **q** on the two times finer divided mesh (or three times finer divided **k** mesh). This is used for Fig.6 in Phys. Rev. B 74, 245125 (2006).

11.5 set QPNT for eps mode (QforEPS section)

For eps modes (scripts `eps_*`, which are for linear responses. See Sec.16), you have to specify q point in the following ways.

1. `QforEPSIBZ` on

Then all Q point in IBZ are used.

2. Use section as

```
<QforEPS>
0d0 0d0 0.01d0
0d0 0d0 0.02d0
0d0 0d0 0.04d0
0d0 0d0 0.08d0
</QforEPS>
```

In addition, you can specify Q points as

```
<QforEPSL>
0d0 0d0 0d0 1d0 0d0 0d0 8
0d0 0d0 0d0 .5d0 .5d0 0d0 8
</QforEPSL>
```

This is along the line— 8 point along the line (not left-end q; so omitting 0 0 0). The first line means line (0d0 0d0 0d0)—(1d0 0d0 0d0) is divided to 8. So we have 7 points, (0.125 0 0), (0.25 0 0),... (1 0 0).

11.6 <PRODUCT_BASIS> section

This section is to define product basis to expand W and so. Numbers are read by free format read(5,*), thus the numbers should be separated by space. The line number in this section is meaningful (you can not add comment lines).

```

<PRODUCT_BASIS>
tolerance to remove products due to poor linear-independency
0.100000D-04 !=tolopt; larger gives smaller num. of product basis. See lbas and lbasC, which are
lcutmx(atom) = maximum l-cutoff for the product basis. =4 is required for atoms with valence d, li
4 3
atom 1 nnvv nnc ! nnvv: num. of radial functions (valence) on the augmentation-waves, nnc: num
1 0 2 3
1 1 2 2
1 2 3 0
1 3 2 0
1 4 2 0
2 0 2 1
2 1 2 0
2 2 2 0
2 3 2 0
2 4 2 0
atom 1 n occ unocc ! Valence(1=yes,0=no)
1 0 1 1 ! 4S_p -----
1 0 2 1 0 ! 4S_d
1 1 1 1 1 ! 4P_p
1 1 2 0 0 ! 4P_d
1 2 1 1 1 ! 4D_p
1 2 2 0 0 ! 4D_d
1 2 3 1 1 ! 3D_l
1 3 1 0 1 ! 4f_p
1 3 2 0 0 ! 4f_d
1 4 1 0 0 ! 5g_p
1 4 2 0 0 ! 5g_d
2 0 1 1 1 ! 2S_p -----
2 0 2 0 0 ! 2S_d
2 1 1 1 1 ! 2P_p
2 1 2 0 0 ! 2P_d
2 2 1 1 1 ! 3d_p
2 2 2 0 0 ! 3d_d
2 3 1 0 1 ! 4f_p
2 3 2 0 0 ! 4f_d
2 4 1 0 0 ! 5g_p
2 4 2 0 0 ! 5g_d
atom 1 n occ unocc ForX0 ForSxc ! Core (1=yes, 0=no)
1 0 1 0 0 0 0 ! 1S -----
1 0 2 0 0 0 0 ! 2S
1 0 3 0 0 0 0 ! 3S
1 1 1 0 0 0 0 ! 2P
1 1 2 0 0 0 0 ! 3P
2 0 1 0 0 0 0 ! 1S -----
</PRODUCT_BASIS>

```

This section is read in the free format in fortran. So, e.g., 0.01 works as same as 0.10000D-01. The line order is important (you have to keep the order given by `GWinput.tmp`). Be careful atom atom id—`lmf` may re-order it and pass it to `gw` code. Look into `LMTO` file (generated by `mkGWIN_lmf2`); which contains crystal structure information after such re-ordering by `lmf`. I used `!` to make clear that things after `!` are comments. But `!` is not meaningful – just the expected numbers of data separated by blank(s) are read for each line from the beginning of lines.

- `0.100000D-02 ! =tolopt` controls a number of Product basis to expand the Coulomb interaction within MTs. `tolopt` is a criterion to remove the poorly linear-independent product basis. Note that the product basis, which is to expand the Coulomb interaction, is different from the basis to expand eigenfunctions. In our experience, `0.100000D-02` (=0.001) is not so bad. If you like to reduce computational time use 0.01 or so, but a little dangerous in cases. With 0.0001, we can check stability on it. (note: By supplying multiple numbers, we can specify `tolopt` atom by atom. Remember `lmchk` gives atom ID.)

- `lcutmx(atom)` is the l cutoff of product basis for atoms in the primitive cell (do `lmchk` for atom id). In the case of Oxygen, we can usually use `lcutmx=2` (need check by the difference when you use `lcutmx=2` or `lcutmx=4`). Then the computational time is reduced well.
- (dec2014:<PBASMAX> is not checked recently; see `fpgw/main/hbasfp0.m.F` and `fpgw/gwsrsrc/basfnf.F`.) You can use <PBASMAX> section to override this setting. It is given as

```
<PBASMAX>
1  5 5 5 3 3
2  5 5 3 2 3
3  3 3 2 2 2
</PBASMAX>
```

The first number is for atom index (fixed), and other are product basis for each l channel.

- The integer numbers in 4th line `lcutmx` gives the maximum angular momentum l for the product basis for each atomic site. In our experience, `lcutmx=4` is required when the semi-core (or valence) $3d$ electrons exist and we want to calculate the QP energies of them.
- Keep a block starting from "atom l nnvv nnc ..." as it originally generated in `GWinput.tmp`. It just shows that how many kinds of radial functions for cores and valence electrons for each atom and l . `nnvv=2` in the case of ϕ and $\dot{\phi}$; `nnvv=3` in the case to add the local orbital in addition.
- There are two blocks after the line "atom l n occ unocc :Valence(1=yes, 0=no)" and after "atom l n occ unocc ForX0 ForSxc ! Core (1=yes, 0=no)". These are used to choose atomic basis to construct the product basis. The product basis are generated from the products of two atomic basis.

`GWinput.tmp` generated by `mkGWIN_lmf2` contains labels on each orbitals as `4S_p`, `4S_d`, `4P_p`... Here `4S_p` is for ϕ_{4s} ; `4S_d` for $\dot{\phi}_{4s}$; `3D_l` for ϕ_{3d}^{local} . Capital letter just after the principle-quantum number means the orbital is used as 'Head of MTO'; lowercase means just used only as the 'tail of MTO'.

The switches for columns labeled as `occ` and `unocc`. take 0 (not included) or 1 (included). With the switch, we can construct two groups of orbitals, `occ` and `unocc`. In this sample `GWIN_V2` as for atom 1, $\{\phi_{4s}, \dot{\phi}_{4s}, \phi_{4p}, \phi_{4d}, \phi_{3d}^{\text{local}}, \phi_{3s}^{\text{core}}, \phi_{3p}^{\text{core}}\}$ consist the group `occ`, and $\{\phi_{4s}, \phi_{4p}, \phi_{4d}, \phi_{3d}^{\text{local}}, \phi_{4f}\}$ consists the group `unocc`. So the any product of combinations $\{\phi_{4s}, \dot{\phi}_{4s}, \phi_{4p}, \phi_{4d}, \phi_{3d}^{\text{local}}, \phi_{3s}^{\text{core}}, \phi_{3p}^{\text{core}}\} \times \{\phi_{4s}, \phi_{4p}, \phi_{4d}, \phi_{3d}^{\text{local}}, \phi_{4f}\}$ are included as for the basis of the product basis. As for atom 2, $\{\phi_{2s}, \phi_{2p}, \phi_{3d}\} \times \{\phi_{2s}, \phi_{2p}, \phi_{3d}, \phi_{4f}\}$ are included.

- Core section: (not worth to read, since we currently use no `CORE2`, `A=B=C=0`.)

Each line of the last section of **Product BASIS** forms

```
atom  1      n  occ unocc  ForX0 ForSxc :CoreState(1=yes, 0=no)
   1    2    1    A    x      B      C
```

At first you have to understand the concept of `CORE1` and `CORE2` in EQ.35 Ref.I. However, in our recent calculations, we do not use "CORE2" generally. So, in such a case, set `A=B=C=0`. And treat shallow cores (above Efermi-2Ry or so) as valence electron by "local orbital method" in `lmf`.

- Be careful. Current version is inconvenient... Need to repeat `mkGWIN_lmf2` to generate `GWinput` template when you add PZ (local orbital).

[(Note: you can skip here if you don't use CORE2.)

Each of **A**,**x**,**B**,**C** takes 0 or 1. There are some possible combination of these switches;

1. If you take (**A x B C**)= (1 0 1 1), then the core is included in core2. In other words, this core is treated in the same manner of the valence electron.
2. If you take (**A x B C**)= (0 0 0 0), then the core is included in core1. The (exchange only) self-energy related to this core is included in **SEXcore**. **C** is the key switch which determine whether it is included in core1 or core2. There could be another option.
3. If you take (**A x B C**)= (1 0 0 1). This core is in core2. But it is not included in the calculation of *D* and *W*. This core is only included for SEX and SEC calculations.

These three kinds of choices are reasonable ones but we can consider some another choice. In the following, we show how these switches (**A**,**B**,**C**) affect executions called from **gw_lmfh** (essentially as same as **gw_lmf**).

- **hbasfp0**(mode 3) :Product basis for exchange due to core.
We include the **C**=0 cores as a part of the product basis as if **A**=1 **x**=0.
 - **hsfp0**(mode 3): exchange mode for core.
 Σ_x only due to the **C**=0 cores are calculated.
 - **hbasfp0** (mode1): Product basis.
Only see the switch **A** and **x**. The product basis is generated from (occupied \times unoccupied), where **A**=1 core is included as one of the occupied basis.
 - **hsfp0** (mode 1): exchange mode.
Only see the switch **C**. Σ_x due to valence and due to **C**=1 cores are calculated.
 - **hx0fp** (mode 1): *W* – *v* calculation.
Only see the switch **B**. *W* is calculates using all the valence and **B**=1 cores.
 - **hsfp0** (mode 2): correlation mode.
Only see the switch **C**. Σ_c due to valence and due to **C**=1 are calculated.
- After you perform **gw_lmfh** or anything, you find output files **lbas** by **hbasfp0** (mode1), and/or **lbasc** by **hbasfp0** (mode3) for core. These contains important information about how many and how product basis are chosen. E.g. '**grep nbloch lbas**' shows how many product basis are used in the calculations.

11.7 ANFcond (we can skip here since we do not check this option now. Need fix this if necessary.

This file is used in **hx0fp0** in the calculation of *W* – *v* (or rather Π in the program) to specify the antiferro condition.

Note : Now only for the case that (a translation vector + spin flip) is a symmetry operation.

This should be given by hand. For the cases of not antiferro, this file should not exist. Even if **ANFcond** does not exists for antiferro case, **hx0fp0** works but it requires about two time computational efforts.

The existence of this file means the Antiferro condition is used for x0k
Product basis $B(\{\mathbf{r}\}-\{\mathbf{a}\})$ is translated to $B(\{\mathbf{r}\}-\{\mathbf{a}\}-\mathbf{Af})= B(\{\mathbf{r}\}-\{\mathbf{a}\}'-\mathbf{T}_0)$
1d0 1d0 1d0 ! Af=Antiferro translation vector in Cartesian.
1 2
2 1
3 4
4 3

The first line specifies the Antiferro translation vector. From the second line, we specify that atom i in the primitive cell is mapped to what atom $j(i)$ in the cell with the opposite spin by the translation. In this case, $j(1) = 2, j(2) = 1, j(3) = 4, j(4) = 3$. You have to be careful as for the true atomic position used in the GW calculations can be different from the given atomic positions in `ctrl.MnO`. The true atomic positions is written in `LMTO`.

In the case of one-shot GW (`gw_lmf` and `gw_lmfh`), it may be better to set "up only" QPE, so that you only calculate QPE of up spins at the same time.

In the case of `gwsc`, we just calculate QPE for up spins automatically (QPNT section is neglected).

12 Main Output Files of GW part

12.1 QPU

This is the main output¹³ in human readable format.

An example of one-shot GW by `gw_lmfh` is (In the case of QSGW, Z ($Z = 1$) is not shown):

```
=====
quasiparticle energies MAJORITY
=====
E_shift=  0.4263273221017709D+00  0.6075150850568627D+00  0.7046628446164018D+00 eV

      q      state SEx  SExcore SEc   vxc   dSE  dSEnoZ  eLDA   eQP  eQPnoZ  eHF  Z   2Z*Simg ReS(
0.0 0.0 0.0 1  -29.56  -1.97  10.40 -20.22 -0.52  -0.90 -19.08 -19.42 -19.71 -30.81 0.58 0.95  -21.
0.0 0.0 0.0 2  -30.52  -2.24  10.09 -21.53 -0.70  -1.14 -18.06 -18.58 -18.93 -29.72 0.61 0.96  -22.
0.0 0.0 0.0 3  -20.67  -1.87   5.97 -16.85  0.19   0.28  -7.20  -6.83  -6.65 -13.32 0.67 0.66  -16.
...
```

From the 6h line, we have the eigenvalue data. All of the unit of energy is in eV. We should note that the zero-level of these values `eLDA` `eQP` `eQPnoZ` can be changed by `hqpe`. This `eLDA` - `E_shift` are the eigenvalues relative to a Fermi energy determined by the smearing method. Detailed value of `eLDA` is in `TOTE2.UP`. Detailed value of `eLDA` - `E_shift` is in `TOTE.UP`.

`q` : `k` vector

`state`: Band index n , which is from the lowest eigenvalue (not include cores).

`SEx`: $= \langle \Psi_{\mathbf{k}n} | \Sigma_{\mathbf{x}}^{\text{core2+valence}}(\mathbf{r}, \mathbf{r}') | \Psi_{\mathbf{k}n} \rangle$

`SExcore`: $= \langle \Psi_{\mathbf{k}n} | \Sigma_{\mathbf{x}}^{\text{core1}}(\mathbf{r}, \mathbf{r}') | \Psi_{\mathbf{k}n} \rangle$

`SEc`: $= \langle \Psi_{\mathbf{k}n} | \Sigma_{\mathbf{c}}^{\text{core2+valence}}(\mathbf{r}, \mathbf{r}', \epsilon_n(\mathbf{k})) | \Psi_{\mathbf{k}n} \rangle$

`vxc`: LDA exchange correlation energy. $\langle \Psi_{\mathbf{k}n} | V_{\mathbf{x}\mathbf{c}}^{\text{LDA}}([n_{\text{total}}], \mathbf{r}) | \Psi_{\mathbf{k}n} \rangle$

`dSE`: $Z_{n\mathbf{k}} \times \text{dSEnoZ}$

`dSEnoZ`: $\langle \Psi_{\mathbf{k}n} | \Sigma_{\mathbf{x}}^{\text{core1}}(\mathbf{r}, \mathbf{r}') + \Sigma_{\mathbf{x}\mathbf{c}}^{\text{core2+valence}}(\mathbf{r}, \mathbf{r}', \epsilon_n(\mathbf{k})) | \Psi_{\mathbf{k}n} \rangle - \langle \Psi_{\mathbf{k}n} | V_{\mathbf{x}\mathbf{c}}^{\text{LDA}}([n_{\text{total}}], \mathbf{r}) | \Psi_{\mathbf{k}n} \rangle$
 $= \text{SEx} + \text{SExcore} + \text{SEc} - \text{vxc}$

`eLDA`: LDA eigenvalues. $\epsilon_n(\mathbf{k})$

`eQP`: QP energy. $\epsilon_n(\mathbf{k}) + \text{dSE}$

`eQPnoZ`: QP energy without Z . $\epsilon_n(\mathbf{k}) + \text{dSEnoZ}$

`eHF`: HF energy of 1st iteration. $\epsilon_n(\mathbf{k}) + \text{SEx} + \text{SExcore} - \text{vxc}$

`Z`: Z factor. $Z_{n\mathbf{k}}$

`2Z*Simg`: Quasi-particle life time. $2Z_{n\mathbf{k}} \times \text{Im} \langle \Psi_{\mathbf{k}n} | \Sigma_{\mathbf{c}}^{\text{core2+valence}}(\mathbf{r}, \mathbf{r}', \epsilon_n(\mathbf{k})) | \Psi_{\mathbf{k}n} \rangle$

(Is this really the usual definition of the life time?—don't believe me)

`ReS(eLda)`: $\text{Re} \langle \Psi_{\mathbf{k}n} | \Sigma_{\mathbf{x}}^{\text{core1}}(\mathbf{r}, \mathbf{r}') + \Sigma_{\mathbf{x}\mathbf{c}}^{\text{core2+valence}}(\mathbf{r}, \mathbf{r}', \epsilon_n(\mathbf{k})) | \Psi_{\mathbf{k}n} \rangle$

12.2 XCU

LDA exchange-correlation. Detailed data of above `vxc`.

12.3 SEXU

Exchange part of the self-energy due to valence electrons. Detailed data of above `SEx`.

12.4 SEXcoreU

Exchange part of the self-energy due to core. Detailed data of above `SExcore`.

¹³Note that QPU also implies QPD and so on. U is for up D is for down spins.

12.5 SECU

Correlation part of the self-energy. Detailed data of above SEc.

12.6 TOTE.UP (TOTE.DN)

This is a central output. It contains LDA and QP energies. These values are relative to a Fermi energy determined by the smearing method. It contains two kind of QP energies QP QPnoZ. The first line contains the Fermi energy in Ry determined by the smearing method. It is also shown in the end of DOSACC.Ida.

12.7 TOTE2.UP (TOTE2.DN)

This is a central output. It contains zero-level shifts from TOTE.UP. The first line contains the Fermi energy in eV (= the Fermi energy in TOTE.UP but it is in Ry) and three energy shifts E_shift, which are the same values in the 4th line of QPU.

Note that all *.chk files are just to check calculations (not read in by successive executions).

12.8 DOSACC.Ida

This lists all the eigenvalues in ascendant order. States with almost the same eigenvalues are degenerated states. The 4th column contains number of electrons up to the eigenvalue.

12.9 DOSACC2.Ida

This is similar with DOSACC.Ida. But we remove the degeneracy.

12.10 Core_ibas*.*.chk

Used core eigenfunctions.

12.11 VXCFP.chk

This contains eigenvalues and $\langle \psi_{\mathbf{k}n} | V_{\text{xc}} | \psi_{\mathbf{k}n} \rangle$ in both units, Ry and eV. See below.

12.12 The Fermi energies in this GWcode.

We mainly have two kinds of Fermi energy $E_{\text{FEERMI}}^{\text{smear}}$ $E_{\text{FEERMI}}^{\text{tetra}}$.

1. At first eigenvalues given by lmf gw is in VXCFP.chk. You can see

```
%head VXCFP.chk
### LDA exchange correlation ###
#   qvec                ikp iband    eigen      VXC(ntotal)      VXC(nvalence)    eigen(eV)
  0.0000  0.0000  0.0000  1  1      -0.96932423      -1.00727912      0.00000000      -13.18843159
...
```

These are raw values. TOTE contains the eigenvalues but relative to a Fermi energy $E_{\text{FEERMI}}^{\text{smear}}$ which is determined by the smearing method. It is also shown at the top part of output files lxx_sf and lsc_sf. And you also see the value at the end of DOSACC.Ida.

This is the head of TOTE.UP;

%head TOTE.UP

```

      43      8  8.520283353474250E-003
      0.0000000  0.0000000  0.0000000  1  1  -0.1330435686590073D+02 -0.1322984339282777D+02
      0.0000000  0.0000000  0.0000000  2  1  -0.7555264915356062D+00 -0.6267595395613325D+00
      ...

```

Here $E_{\text{FEERMI}}^{\text{smear}} = 8.520283353474250\text{E}-003$. From the second lines, they are LDA eigenvalues and QP energies (Z included and Z=1); they are relative to the $E_{\text{FEERMI}}^{\text{smear}}$.

-13.18843159 eV - $E_{\text{FEERMI}}^{\text{smear}}$ (which should be translated into in eV) = -0.1330435686590073D+02 eV. Here -13.18843159 is the value in VXCFF.chk shown above.

2. There is the another Fermi energy $E_{\text{FEERMI}}^{\text{tetra}}$, which is used by mode 11 (or mode 1) of `hx0fp0` in `gw_lmfh`. It is determined by `heftet` and stored in `EFERMI`.
3. `hqpe` gives `TOTE2.UP` and `QPU`. They contains the same values. You can see `eLDA` `eQP` `eQPnoZ` `Z` not only in `QPU` but also in `TOTE2.UP`. At top lines of `TOTE2.UP`, you see

%head TOTE2.UP

```

      43      8  0.1159252712507000D+00  0.7555207081466229D+00  0.6267572296579150D+00  0.6
      0.0000000  0.0000000  0.0000000  1  1  -0.1254883615775411D+02 -0.1260308616316985D+02
      0.0000000  0.0000000  0.0000000  2  1  -0.5783388983382487D-05 -0.2309903417430093D-05
      0.0000000  0.0000000  0.0000000  3  1  -0.1369098933889923D-05 -0.6195200397129952D-07
      0.0000000  0.0000000  0.0000000  4  1  0.0000000000000000D+00  0.0000000000000000D+00
      ..

```

, where a number in first line $E_{\text{FEERMI}}^{\text{smear}} = 0.1159252712507000\text{D}+00$ eV = 8.520283353474250E-003 Ry, the same as the previous one. This is a case when you did `hqpe` with augment 4 (it means we set the 4th-band eigenvalue zero). Another 3 values in the first line are shifts from `TOTE`. Shown `eshift(eLDA)` = 0.7555207081466229D+00 eV. E.g., the second line shows -0.1254883615775411D+02 eV = -0.1330435686590073D+02 (in `TOTE`) + `eshift(eLDA)` eV.

When you do `hqpe metal`, three shifts at the first line in `TOTE2.UP` is determined so as to give the eigenvalues relative to the Fermi energies shown in `EFERMI`, `EFERMI.QP1`, and `EFERMI.QPz=1`. These are Fermi energies by tetrahedron method.

As for `gwband_lmf`, it recalculates eigenvalues for all **q** along SYML. Then the default "zerolevel" = $E_{\text{FEERMI}}^{\text{smear}} - \text{eshift(lda)}$. Because the eigenvalues given by this band-mode are presumably the same, we have

Shown LDA eigenvalue

```

= -13.18843159 (raw data by band mode—same as that in VXCFF.chk) - zerolevel
= (-13.18843159 - EFERMIsmear) + eshift(lda).
= -0.1330435686590073D+02 ( this is in TOTE.UP) + eshift(lda)
= -0.1254883615775411D+02 ( this is in TOTE2.UP).

```

It means that values in `TOTE2.UP` recovers. But if raw data by band mode is different from it, these is a trouble. It does not recover the values in `TOTE2.UP`(=QPU).

As for the QPE, we calculate the difference from LDA values in `TOTE2.UP` at first, and add the difference to the Shown LDA eigenvalue.

13 mkGWIN_lmf2 and its I/O Files

(QPNT.chk contains irreducible k point for given n1 n2 n3; KPTin1BZ.gwinit.chk contain all k points in Brillouin Zone).

The purpose of the script **mkGWIN_lmf2** is to give a template **GWinput.tmp**. The script is complicated because of historical reasons. However, its essential is simple; we calls three executions in this script as

```
echo 0 | lmf gw si
echo 1 | gwinit
echo -100 | qg4gw
```

. We explain each by each.

13.1 echo 0|lmfgw

— makes SYMOPS LATTC CLASS NLAindx.

Input files

- **GWIN0** : This is a file, which contains your supplied n1 n2 n3 when you invoke the script. This file is given within the script of mkGWIN_lmf2 (as "here document").

```
cat <<EOF >GWIN0
n1 n2 n3
  $n1 $n2 $n3
cut
  4.0 3.0
alpha
  1
Number of bands
  999 99999.0
  999 3.0
EOF
```

- **ctrl.si** : Master input file of lmf calculation.

Output files

- **LATTC** : contains the information of primitive translation vectors, lmxa and konf. See ??
- **SYMOPS** : The point group operations. See ??
- **CLASS** : Equivalent atomic positions are called as 'class'. This small file contains a map between atomic site and 'class'.
- **NLAindx** : This file contains indexes (p_{valence}, l, a) for orbitals in the MT. (p_{valence} is radial function index, a is atomic site index). Eigenfunctions are expanded in this order.
- **ldima** : Number of MTOs for each atomic site. (this is used only from **hqpe_sc**—QSGW mode).
- **ves*** : not meaningful at this stage
- **rhoMT*** : not meaningful at this stage

13.2 gwinit

— Get GWIN_V2.tmp and QPNT.tmp

Input files

- **GWIN0** :
- **LATTC** :
- **SYMOPS** :
- **NLAindx** :

Output files

- **GWIN_V2.tmp** : A part of GWinput.tmp
- **QPNT.tmp** : A part of GWinput.tmp
- (**KPNTin1BZ.gwinit.chk**) : check KPNT in the 1BZ.

If SYML exist, **gwinit** gives also a template **QPNTforSYML.tmp** suitable for such SYML. Here SYML specify how to plot the energy band. See explanation for **bandplot** script.

Note that LATTC SYMOPS CLASS NLAindx are overwritten when you execute **gw_lmfh** because we repeat **echo 0|lmf** at the head of **gw_lmfh**.

13.3 echo -100|qg4gw

— Generate GWinput.tmp

Input files

- GWIN0 : (copy of GWIN0.tmp by **gwinit**)
- GWIN_V2 : (copy of GWIN_V2.tmp by **gwinit**)
- QPNT : (copy of QPNT.tmp by **gwinit**)

Output files

- GWinput : (this is copied to GWinput.tmp)

This command "echo -100|qg4gw" is a file converter from these two files into GWinput. And it is copied to GWinput.tmp. (**mkGIN_lmf** keeps GWinput if it exist before you invoke it.).

14 gwsc script and its I/O Files

In `gwsc`, we have a loop of QSGW self-consistency. Look into the `gwsc` script. In each iteration, we perform these fortran programs;

```
NO_MPI=0 #this is used for non-mpi versions of fortran program.

### self-consistent calculation for given Sigma(self-energy) ###

    run_arg '---' $MPI_SIZE $nfpwgw /lmf-MPIK    llmf $TARGET

### Preparation stage #####

argin=0; run_arg $argin $NO_MPI $nfpwgw /lmfgw      llmf gw00 $TARGET
argin=1; run_arg $argin $NO_MPI $nfpwgw /qg4gw      lqg4gw #Generate required q+G v
argin=1; run_arg $argin $MPI_SIZE $nfpwgw /lmfgw-MPIK llmf gw01 $TARGET
        run_arg '---' $NO_MPI $nfpwgw /lmf2gw      llmf2gw #reform data for gw

### Main stage of gw #####

argin=0; run_arg $argin $NO_MPI $nfpwgw /rdata4gw_v2 lrd4gw_v2 #prepare files
argin=1; run_arg $argin $NO_MPI $nfpwgw /hefttet     leftet # A file EFERMI for hx0fp0
argin=1; run_arg $argin $NO_MPI $nfpwgw /hchknw      lchknw # A file NW, containing nw

## Core part of the self-energy (exchange only) ##

argin=3; run_arg $argin $NO_MPI $nfpwgw /hbasfp0     lbasC # Product basis generation
argin=3; run_arg $argin $MPI_SIZE $nfpwgw /hvccfp0   lvccC # Coulomb matrix for lbasC
argin=3; run_arg $argin $MPI_SIZE $nfpwgw /hsfp0_sc   lsxC # Sigma from core1

## Valence part of the self-energy Sigma ##

argin=0; run_arg $argin $NO_MPI $nfpwgw /hbasfp0     lbas # Product basis generation
argin=0; run_arg $argin $MPI_SIZE $nfpwgw /hvccfp0   lvcc # Coulomb matrix for lbas
argin=1; run_arg $argin $MPI_SIZE $nfpwgw /hsfp0_sc   lsx # Exchange Sigma
argin=11; run_arg $argin $MPI_SIZE $nfpwgw /hx0fp0_sc lx0 $lx0_para_option #x0 part
argin=2; run_arg $argin $MPI_SIZE $nfpwgw /hsfp0_sc   lsc #correlation Sigma
argin=0; run_arg $argin $NO_MPI $nfpwgw /hqpe_sc      lqpe #all Sigma are combined.

run_arg:
```

Here a subroutine of bash `run_arg` was used, which is given in `ecalj/lm7K`; it just invoke a command with the argument `$argin` (this is read by `read(*,*)` in fortran). In cases with `MPI_SIZE/=0`, `mpirun` is invoked. Console out put go to `l*` files. For example,

```
    argin=2; run_arg $argin $MPI_SIZE $nfpwgw /hsfp0_sc lsc #correlation Sigma
```

invokes `hsfp0_sc` with argument '2' by `mpirun` with the `-np $MPI_SIZE`. Console outputs are written into logfiles such as `lqpe`. `$nfpwgw` contains path to the execution binaries.

In the following, We explain input/output files for each fortran program. Note that “`echo 0|lmfgw`” means invoking `lmfgw` with `argin=0`.

14.1 echo 0| lmfgw si

See Sec.13.1.

14.2 echo 1| qg4gw

This makes **q** points, and **G** vectors for these **q**. (**q** was **k** in previous sections.) Main routine of qg4gw is fpgw/main/qg4gw.m.F and calls fpgw/gwsrsc/mkqg.F

Input files

- GWinput :
- LATTC :
- SYMOPS :

Output files

- QGpsi : (bin) q and G vector for the eigenfunctions.
- QGcou : (bin) q and G vector for the Coulomb matrix
- Q0P : offset- Γ points which are the replacement of the $q=0$ points. See section??.
- QIBZ : q points in the Irreducible BZ.
- BZDATA : (bin) BZ data for integration (include tetrahedrons if necessary). See e.g. main/hxOfp0.sc.F and search "call read_BZDATA", which is a readin routine of this file defined in rwbzdata.F.
- KPTin1BZ.mkqg.chk : list of q in the 1st BZ for check.
- QBZ : q point in the 1st BZ.
- EPSwklm : Required information for the BZ integration (mainly in order to evaluate the weight in the Γ cell). See Eq.xxx in [?].

14.3 echo 1|lmfgw si

Calculate eigenfunctions, eigenvalues and $\langle\psi|H_{KS}|\psi\rangle$

Input files

- ctrl.si :
- rst.si : (bin) Restart file of the lmf calculation. It contains all information
- sigm.si : (bin) If this exist and
- QGpsi, QGcou, Q0P : :
- NLAindx : :

Output files

- gwa.si : (bin) atomic data
- gwb.si : (bin) band data
- gw1.si : (bin) $\langle\psi|H_{KS}|\psi\rangle$
- gw2.si : (bin) $\langle\psi|H_{KS} - V_{xc}(n_{total})|\psi\rangle$.
- vxc.si, evvec.si : (bin) used in hqpe.sc.m.f as "v_xc" and "evvec").

vxc.si contains $\langle\psi|V_{xc}(n_{total})|\psi\rangle$ including off-diagonal part. evvec.si contains eigenfunctions.

- normchk.si : norm check (only for check) This is like this

```
> head -20 normchk.si
#      IPW      IPW(diag)      Onsite(tot)      Onsite(phi)      Total
      0.436015      0.805123      0.563972      0.562573      0.999988
      0.339134      0.620353      0.660515      0.656881      0.999649
      0.339133      0.620353      0.660516      0.656882      0.999649
      0.339133      0.620353      0.660516      0.656882      0.999649
      0.507738      0.648515      0.492040      0.487673      0.999778
...
```

This check is sometimes important for debugging and to determine the cutoff parameter **QGcut_psi**. The first line (corresponding to 1st band of 1st q point) means that total normalization almost unity = 0.999988 = 0.436015 + 0.563972. Because we expand the MTO by IPW, the normalization is a bit different from unity, especially for higher bands. You can see that it get closer to unity for larger QGcut_psi, though it does not reach to unity because of some contribution of the higher angular momentum contribution within MT. [Values of Onsite(phi) are not correctly shown in the case when you use local orbital.]

Due to historical reason, data in vxc.si and exec.si and others contains duplicated data.

14.4 lmf2gw

All the required information are stored into DATA4GW_V2 and CphiGeig.

Input files

- gwa.si :
- gwb.si :
- gw1.si :
- gw2.si :
- Q0P :
- CLASS :
- NLAindx :

Output files

- DATA4GW_V2 : (bin) Main data for GW calculations.
I/O of DATA4GW_V2 is controlled by `gwinput.f`, which contains detailed information.
- CphiGeig : (bin) Eigenfunctions for GW calculations.
- VXCFP.chk : Eigenvalue and Vxc check (only used for check)
It is like this;

```
### LDA exchange correlation ###
#   qvec      ikp iband   eigen   VXC(ntotal)   VXC(nvalence)   eigen(eV)   VXC(ntotal)(eV)   VXC(nvalence)
0.0000 0.0000 0.0000 1 1   -0.68505346   -0.91850436   0.00000000   -9.32070032   -12.49698668   0.00000000
0.0000 0.0000 0.0000 1 2    0.19292662   -0.99853478   0.00000000   2.62492096   -13.58586453   0.00000000
0.0000 0.0000 0.0000 1 3    0.19292763   -0.99853469   0.00000000   2.62493477   -13.58586334   0.00000000
0.0000 0.0000 0.0000 1 4    0.19292777   -0.99853461   0.00000000   2.62493664   -13.58586222   0.00000000
...
```

Here `VXC(nvalence)` is not used now. The eigenvalue in `eigen` is in Ry.

— This is the end of the preparation stage. —

From here, the main stage.

14.5 rdata4gw_v2

— Read DATA4GW_V2 and some files, and decompose it into files required in the following GWsteps. (checked! dec2014)

Input files

- GWinput :
- DATA4GW_V2 :
- CphiGeig :
- QGpsi :
- QGcou :
- Q0P :
- QIBZ :
- SYMOPS : points group operations.

Output files

- hbe.d : data size
- Coreibas*.*.chk : core eigenfunctions just for check.
- LMTO : basic data for the crystal.
- EValue : (bin) valence eigen value
- ECORE : core data and core eigenvalues
- CPHI : (bin) Coefficients of eigenfunctions as for the atomic-like argumentation waves in MTs'.
- GEIG : (bin) Coefficients of eigenfunctions as for IPW.
- PHIVC : (bin) All the radial functions.
- @MNLA.CPHI : index set for CPHI. This is not refereed just a check write.
- @MNLA.core : index set for core. This is not refereed just a check write.
- VXCFP : (bin) this is for diagonal elements of $V_{xc}^{LDA}(n_{total})$.
- PPOVLI.* : (bin) Overlap matrix of IPW. xxxxxxxxxxxx

- PPOVLG.* : (bin) PPOVLG Overlap matrix xxxxxxxxxxxxf IPW. not exactly the the overlap matrix. see around line 500 in rdata4gw.m.f
- PPOVL0 : (bin) xxxxxxxxxxxxxx
- HVCCIN : (bin) Required inputs for hvccfp0. Information in this files.
- NQIBZ : q point info. Only used for parallel test mode.
- normchk.dia : Norm check. These numbers should be almost the same as those in normchk.si

These files are input for the following steps. The name of file *fooU* means that it relates to up-spin. We have *fooD* files in the case of spin-polarized calculation with **nspin=2**.

14.6 echo 1|heftet

— Get the Fermi energy EFERMI by tetrahedron method. It is used in **hx0fp0**.

Input files

- EVU :
- BZDATA :
- GWinput :
- ECORE : (dummy)
- SYMOPS : (dummy)
- LMTO :
- hbe.d :

Output files

- EFERMI : contains Fermi energy given by the tetrahedron method. It is used in **hx0fp0** but not in **hsfp0**.
- DOSACC.la,DOSACC2.la : They are lists of the all the eigenvalues from the bottom. DOSACC2.la is a list to show only the un-degenerated eigenvalues. They are just check write. But it is an indicator for you to determine **esmr** in **GWinput**.

14.7 hchknw

— Calculate the required number of ω points along real axis.

This NW is not essentially used in **gw_lmfh** (but required as a dummy file). Only used in **gw_lmf**.

Input files

- BZDATA :
- GWinput :
- ECORE : (dummy)
- SYMOPS : (dummy)

Output files

- NW : contains number of ω points.

14.8 echo 3|hbasfp0

— Make product basis.

Mode 3 is for the core states. It generate a product basis on each MT suitable to expand to calculate the exchange part due to core. See explanations for the input file of **GWinput**.

Input files

- LMTO :
- PHIVC :
- GWinput :

Output files

- BASFP* : (bin) Product basis functions
- PPBRD_V2.* : (bin) Radial integrals on each MT, symbolically written as $\int \phi(r)\phi(r)B(r)dr$
- PHIV.chk : Valence radial functions (for check).

14.9 echo 0| hvccfp0

— Calculate the Coulomb matrix in the Mixed basis

Input files

- HVCCIN :
- Q0P :
- BASFP* :

Output files

- VCCFP : The Coulomb matrix expanded in the mixed basis
- Mix0vec : This is used only for dielectric-constant calculation (mode 2 or 3 of **hx0fp0**). This contains the expansion of the plane wave $\exp(i\mathbf{q}\mathbf{r})$ in the mixed basis. See Usuda's note.

14.10 echo 3|hsfp0

— Exchange part of the self-energy for the core

Input files

- GWIN_V2,LMTO,ECORE :
- CLASS :
- hbe.d :
- Q0P :
- PPBRD_V2_* :
- CPHI :
- GEIG :
- VCCFP :
- PPOVL :

Output files

- SEXcoreU : The core part of the exchange self-energy for \mathbf{q} and band index specified in $\langle \text{QPNT} \rangle$. See ??.

14.11 echo 0|hsfp0

— Make product basis for the valence part.

14.12 echo 1|hsfp0

— Exchange part of the self-energy for the valence part.

Output files

- XCU : The LDA exchange self-energy for \mathbf{q} and band index specified in $\langle \text{QPNT} \rangle$. See ??.
- SEXU : The valence part of the exchange self-energy for \mathbf{q} and band index specified in $\langle \text{QPNT} \rangle$. See ??.

14.13 echo 11|hx0fp0

— Screened Coulomb interaction W (Sergey mode)

Input files

- GWinput, LMTO, Ecore, EVU :
- NW : dummy
- hbe.d :
- Q0P :
- PPBRD_V2_* :
- CPHI,GEIG :
- PPOVL :
- VCCFP :
- ANFcond : (optional) This file is to specify antiferro condition. This should not exist for other cases. This file should be given by hand.

Output files

- WV.d : size of the dielectric function
- WVR : (bin) $W - v$ in the expansion of mixed basis along the real axis

- WVI : (bin) $W - v$ in the expansion of mixed basis along the imaginary axis

14.14 echo 12|hsfp0

— Correlation part of the self-energy(Sergey mode)

Input files

- GWinput, LMTO, Ecore, SYMOPS : These are readin by **genallcf_v3**.
- CLASS, hbe.d, EVU, Q0P :
- PPBRD_V2.* :

Radial integrals on each MT, symbolically written as $\int \phi(r)\phi(r)B(r)dr$. These are generated by **hbasfp0**.

- CPHI,GEIG :
- PPOVL :
- WV.d, WVR, WVI :

Output files

- SECU : The correlation part of the self-energy for **q** and band index specified in <QPNT>. See ??.

14.15 echo 0|hqpe

— Summarize the output

Input files

- SEXcoreU,XCU,SEXU,SECU : See ??.

Output files

- QPU : The QP energies and related value summary in human interface. See ??.
- TOTE : The detailed values of the QP energies. See ??.
- TOTE2 : The detailed values of the QP energy. See ??.

NOTE: For example, if you do **{echo 4|\$hqpe}{hqpe}**, it just shift zero level of QPE, so that 4th line (counted from top) eigenvalue (in QPU) is to be zero.

15 Check list for convergence on GW calculations

Results could be dependent on cutoff parameters in `GWinput`. In my opinion, generally speaking, it is so easy to have convergence more than $\sim 0.1\text{eV}$ for band gap...

- Number of k points `n1n2n3`.

Probably $4 \times 4 \times 4$ (or $6 \times 6 \times 6$) are reasonable choice for insulator in the case of two atoms such as GaAs. In other words, “used periodic cell volume” = $4 \times 4 \times 4 \times (\# \text{ of atoms}) \times (\text{Volume per atom})$. For example, for $2 \times 2^3 = 16$ atoms per cell, we can use $2 \times 2 \times 2$ instead of $4 \times 4 \times 4$ (this is the same as in the case of LDA).

For metals, $12 \times 12 \times 12$ or more per atom may be required. But it depends on case by case.

- MTO’s and APWs. Basis for eigenfunctions.

If we like to get “best converged results, we may need to use large enough MTO’s. (I think the default setting is reasonable; but there is a room to change MTO settings in `ctrl.*`). But, in cases, we can not use large APW cutoff (`pwemax`) more than $3.0 \sim 4.0$ Ry because of poor linear dependency of basis set (calculation in LDA level fails). In such a case, we need to use “smaller MT radius `R=`”. Then we may need semi-core as local orbital when “its spillout in the outside of MTs is too large”.

In `GWinput`, we can set the number of unoccupied states which you take into account by `emax_chi0`, `emax_sig`, `nband_chi0`, and `nband_sig`. But we usually unset them except `emax_sig` for `gWSC`.

(We may need a kind of completeness of the basis set; the completeness could be important from the view of ‘Coulomb hole’ picture.)

NOTE: Current PMT-QSGW method [?] expand the static version of self-energy of QSGW just in the basis of MTO’s (no APWs). Thus the expansion can be unsatisfactory (it depends on case by case, and required convergence). In such a case, we inevitably have to use empty spheres (MTO’s) which is for empty region.

- Cores.

We usually use all cores as `core1` (exchange only core). If necessary, it is better to treat shallow cores by the local orbitals (such cores are treated as valence).

—followings are obsolete —

If we treat cores by `core2` (not only for exchange, but also in dielectric functions), we have to be careful about the core wave orthogonalization with respect to valence eigenfunctions; this is a little complicated; probably it is better not to use `core2`.

(This is related to `CoreOrth` (only for `core2`). If it affects so much, `D` function might be too poor due to the poor orthogonality condition between core and valence.) (NOTE: `CoreOrth` is not maintained recently)

- `QpGcut_psi` IPW cutoff to expand eigenfunctions in the interstitial region. We usually use `QpGcut_psi 4.0`. Usually not so bad. Larger is better but expensive. You may test calculations with `QpGcut_psi 3.0`, and how much difference of results.
- `QpGcut_cou` IPW cutoff to expand the Coulomb interaction in the interstitial region (Mixed product basis(MPB) consist of this IPW and PB). interstitial region. We usually use `QpGcut_cou 3.0`. Usually not so bad. Larger is better but expensive. You may test calculations with `QpGcut_cou 2.5`, and how much difference of results.
- Product basis section.
At least, `1cutmx=4` will be necessary for atoms with d electrons. (but `1cutmx=2` look reasonable for oxygen, `1cutmx=1` for hydrogen. Need check).
- `esmr`.
In our experience, `esmr=0.003000` (default) is reasonable. But there is a room to check stability on it for metals. In principle, for larger `n1n2n3`, we can use smaller `esmr`.

- `dw, omg_c` `niw`

It will be worth to try to check how much the results changed due to them. But usually `dw=0.005`, `omg_c=0.04` is not so bad a choice. As for `niw=10` seems to be not so bad usually, but it is safer to check the convergence on it (test cases with `niw=6,10,12,16`).

- `deltaw`. ~ 0.01 a.u. will be not so bad. See two Z values shown in SXCU. It is better to try to check how about the dependence on this.
- `chi_regqbz off`. (on is default). If off, we use off-Gamma mesh (Gamma point is between mesh points) for dielectric functions when we perform GW or QSGW. “chi_regqbz off” may accelerate convergence on number of k points.
- `PWMODE=1` in ctrl This used fixed number of G vector determined at $q=0$

16 Linear response calculations

With these scripts for linear response calculations, **eps***, we can calculate **q**-dependent dielectric function $\epsilon(\omega, \mathbf{q})$ (and v , W) (and χ for spin fluctuation). But (because of numerical reason), we can not use $\mathbf{q} = 0$ limit. (if $|\mathbf{q}|$ is too small, we have numerical problem, zero divided by zero, because we have not implemented the version to use $\mathbf{q} = 0$.)

- **eps_lmfh**
Dielectric function epsilon with local field correction. Expensive calculation (we may need to reduce number of wing parts in future...).
 - **epsPP_lmfh**
epsilon without local field correction. $1 - \langle e^{i\mathbf{q}\mathbf{r}} | v | e^{i\mathbf{q}\mathbf{r}} \rangle \langle e^{i\mathbf{q}\mathbf{r}} | (\chi^0) | e^{i\mathbf{q}\mathbf{r}} \rangle$
 - **epsPP_lmfh_chipm**
For spin susceptibility. This essentially calculate non-interacting spin susceptibility. Then it is used for the calculation of full spin susceptibility with **util/calj_*.F** programs (small quick programs). See spin wave paper. See spin susceptibility section Sec.??.
 - (not maintained now; we will recover this) **eps_lmfh_chipm**
This gives full non-interacting spin susceptibility. Testing. We have to determine U (Stoner I) for the determination of full spin susceptibility. TDLDA? or so?
 - (This is old mode --- not maintained) **epsPP_lmfh_chipm_q**
For spin susceptibility, spin susceptibility $\langle e^{i\mathbf{q}\mathbf{r}} | \chi(q, \omega) | e^{i\mathbf{q}\mathbf{r}} \rangle$ In this script, You have to assign that isp=1 is majority, isp=2 is minority. This is with long wave approximation.
-
- We use the histogram method (the Hilbert transformation method); we first calculate its imaginary parts with the tetrahedron technique for dielectric functions. Then we get its real part by the Hilbert transformation.
You need to choose **HisBin_dw, HisBin_ratio**. The width of histogram bins are getting larger when omega gets larger. dw is the size of histogram-bin width at omega=0. At omega=omg_c, its width gets twiced.
- To plot dielectric function with reasonable resolution, it might be better to set **dw** 0.001 and **omg_c** 0.1 for example. You may have to choose small enough omega for spin wave mode as 0.001 Ry (Or smaller). omg_c is given like 0.05 Ry or so. But sometimes it can be like 1Ry.
- **epsPP_lmfh** only calculation an matrix element of dielectric function for $\exp(i\mathbf{q}\mathbf{r})$. Thus very faster than **eps_lmfh** mode. It uses a a special product basis set for cases without inversion (problem is in how to expand $\exp(i\mathbf{q}\mathbf{r})$ in the MPB; the product basis is not from phi and phidot, but from spherical Bessel functions).

In ***_lmfh_*** modes(I now use little for ***_lmf_*** modes), you can use small enough delta. Use small enough delta ($\approx 10^{-8}$ a.u.) for spin wave modes (also you can use it for dielectric function and GW). This is necessary because pole is too smeared if you use larger delta.

16.1 eps_lmfh, epsPP_lmfh: the dielectric functions

You can invoke the script, e.g. as **"eps_lmfh si"**.

Specify **q** point in **<QforEPS>** or so. Mesh for ω is specified by **dw, omg_c**.

The obtained data are in **EPS*.dat** and **EPS*.nlfc.dat**. **EPS*.nlfc.dat** contains the result without local-field correction **EPS*.dat** contains the result with local-field correction (this is generated only for **eps_lmfh**. Both of them contains

q(1:3), ω , $\text{Re}(\epsilon)$ $\text{Im}(\epsilon)$, $\text{Re}(1/\epsilon)$, $\text{Im}(1/\epsilon)$

in each line.

- This code works OK only for **q** is near 0. Be careful for $\mathbf{q} \rightarrow 0$ limit. Too small **q** can give strange spectrum at high energy (real part is affected by it)

Because $\mathbf{q} \rightarrow 0$ gives too large cancellation effects (the denominator and numerator go to zero—it means we need very accurate orthogonalization between occupied and unoccupied states). This is a kind of disadvantage of our method (though there is an advantage— our code can calculate dielectric function even for metal as far as you use large enough number of \mathbf{k} point.)

- The calculate of dielectric functions usually requires so many k point. For example, for Si, $n1\ n2\ n3 = 4\ 4\ 4$ is too small. It gives too large dielectric constants ~ 19.4 though the converged value should be ~ 13 . (we need $10 \times 10 \times 10$ or more like $20 \times 20 \times 20$ for some reasonable results). For GaAs, we observed that reasonable $\epsilon(\omega)$ requires rather large number of \mathbf{q} points like $15 \times 15 \times 15$ or $20 \times 20 \times 20$ for `n1n2n3`. This is too time-consuming to get result (but you can use “very small product basis”(just sp polarization for this purpose; it makes speed up so much). Or, you can calculate “ $\epsilon(\omega)$ without LFC”. See section for `eps_PP_lmfh`.

- Core orthogonalization problem (only when `core2` is used)
—`CoreOrth` is not maintained recently — `CoreOrth` gives so serious effect for $\epsilon(\omega)$, if you include some cores as “`core2`” in the product basis setting. (This means that you includes transitions from “`core2` to valence” in the calculation of $\epsilon(\omega)$).

Then you have to use “`CoreOrth on`”. Without it, you will have rather large imaginary part at rather high energy. Such transitions from core to higher valence bands is artificial due to the incomplete orthogonality between core and the higher bands. However, shallow d semi-core might be deformed too much by this option. Try to plot `Core_*.chk` files, which contains core radial functions. Anyway, it is better to treat shallow core as valence by “local orbital”.

16.2 epsPP_lmfh: the dielectric function(No LFC— faster)

You can calculate ϵ without LFC by `epsPP_lmfh`. It is very faster than `eps_lmfh`.

To calculate $\epsilon(\mathbf{q}, \omega)$ without LFC accurately, the best basis set for the expansion of the Coulomb matrix within MT is apparently not the product basis, but the Bessel functions corresponding to the plane waves $\exp(i\mathbf{q}\mathbf{r})$. We use such a basis in this mode. However, our experience shows that the changes are little even with the usual product basis (we don’t describe this here).

16.3 How to calculate correct dielectric function?

(this subsection is essentially OK... but need to clean it up. dec2014)

There are problems to calculate correct epsilon.

At first, we talk about `epsPP_lmfh`, which is No LFC. Main problem are

1. Convergence for number of \mathbf{k} point(specified by `n1n2n3`).

Roughly speaking, $20 \times 20 \times 20$ is required for not-so-bad results for Fe and Ni.

It is better to do $30 \times 30 \times 30$ to see convergence check.

However, in the case of ZB-MnAs (maybe because of simple structure around Ef), it requires less \mathbf{q} points.

figs are for GaAs.

fig001: `n1n2n3` convergence for `Chi_RegQbz = on` case.

fig002: `n1n2n3` convergence for `Chi_RegQbz = off` case.

(`Chi_RegQbz` is explained in General section in this manual).

As you see, \mathbf{k} points convergence looks a little better in `Chi_RegQbz=off`

(mesh not including gamma). However a little problem is that its threshold around 0.5eV is too high and slowly changing.

fig003: Alouanis' (from Arnaud) vs. 'Chi_RegQbz = on' vs. 'Chi_RegQbz = off'
 As you see, the threshold of the Red line (20x20x20 Chi_RegQbz=on) and Alouani's
 are almost the same, but the red line is too oscillating at the low energy part.
 On the other hand, 'Chi_RegQbz = off' in Green broken line is not so satisfactory
 at the low energy part.

fig.gas_eps_kconf.pdf shows the convergence behavior of epsilon for

2. $q \rightarrow 0$ convergence (this is related to whether Chi_RegQbz=on or off).

If you use very small q like $q=0.001$ in GaAs, it can cause a problem.

Use $q=0.01$ or larger (maybe $q=0.02$ or more is safer).

Very small q can give numerical error for high-energy region.

In fig004, we show the high energy tail part of $\text{Im } \epsilon(\omega)$ for GaAs case.
 At $q=0.01$ (this means $q = 2\pi/\text{alat} * (0.001)$), the imaginary part
 is a little too large. Less than 80eV, $q=0.02$ gives good results when compared with
 other high q results, though it still has noise above 80eV.

In fig005, I showed the same results compared with Alouani's (his is up to 40eV).
 Both gives rather good agreements. As you see, $q=0.06$ or above might be necessary
 to get reasonable convergence for high energy part above 40eV.

We have to be careful for this poor result in high energy part--- it may effect
 low-energy $\text{Re } \epsilon$ through KK relation. However this can be very small
 enough.

In fig.gas_eps_qconv.jpg, we checked the convergence of $\epsilon(\omega=0, q)$ for $q \rightarrow 0$.
 As you see, it gives convergence, however, $q=0.01$ is a little out of
 curve---this should be because of the poor result in the high energy part.
 so $q=0.02$ or $q=0.03$ is safer, and you can get ϵ within 1 percent accuracy.

3. Including Core for dielectric constant is dangerous.

It can cause very poor results if you include core part in GWinput.

You need to include core just as valence (with local orbital).

In fig008, we showed core effects. It starts from $\approx 16\text{eV}$
 (this is core to conduction transition).

fig007 showed the check about the q point dependence---even with large q ,
 it would not change.

These shows that the core excitation can have larger energy range.

This is in contrast to the valence case

(then the most of excitation is limited to less than 10eV).

We have to be careful for such high-energy excitation... The LMT0 basis might
 be not so good for high energy.

4. basis set.

Use $Q_{\text{Gcut_psi}} \approx 3.0$ a.u. or so (as same as GW calculation).

In the case of epsPP* mode,

$Q_{\text{Gcut_cou}}$ can be very small--- In our codes now,

$\text{ngc} \geq 1$ should be for all q vector shown in lqg4gw02 (output of echo 2|qg4gw).

[In principle, it should be only for the q vector for which we calculate epsilon.

But there is a technical poor result in our code---

(maybe) a problem here; the plane-wave part of the eigenfunction generated
 in lmf2gw is not correctly passed to lmf2gw when $\text{ngc}=0$].

-- eps_lmfh: including LFC -----

To include eps with LFC, do eps_lmfh.

But lcutmx=2 seems to be good enough to get 0.5 percent error (maybe better than this).

Test it 10x10x10 or so. (I need to repeat if necessary).

Further you can use smaller QpGcut_cou like 2.2 or so,

with rather smaller product basis (up to p timed d, not including f).

Note: epsPP_lmfh is designed to use good basis to calculate eps

without LFC. This is usually in agreement with what you obtained by eps_lmfh;

however it can give slight difference when you use small product basis.

---Summary -----

So in conclusion, I think a best way to do is

1. set $q=0.02$ [$q=2\pi/\text{alat}(0\ 0\ 0.02)$] or so for GaAs case.

If you want to check, do $q=0.03$ and $q=0.06$ also.

‘‘Chi_RegQbz = off’’ is better for materials like GaAs with direct gap.

2. You can use small QpGcut_cou but all ngc should be one or more.

3. As for the Product basis setting in epsPP* scripts, only

lcutmx and tolerance (this can be like 0.001 or so) are relevant.to determine $\text{eps}(\omega=0, q=0)$.

E.g. set lcutmx=4 or so.

5. To get eps with LFC, set QpGcut_cut as xxx, and set lcutmx=2 where

4. Do nk=20 18 16 and take interpolation

(occupied sp) \timex (unoccupied spd) are included.

But correct EPS*.nolfc.d is rather from epsPP_lmfh script.

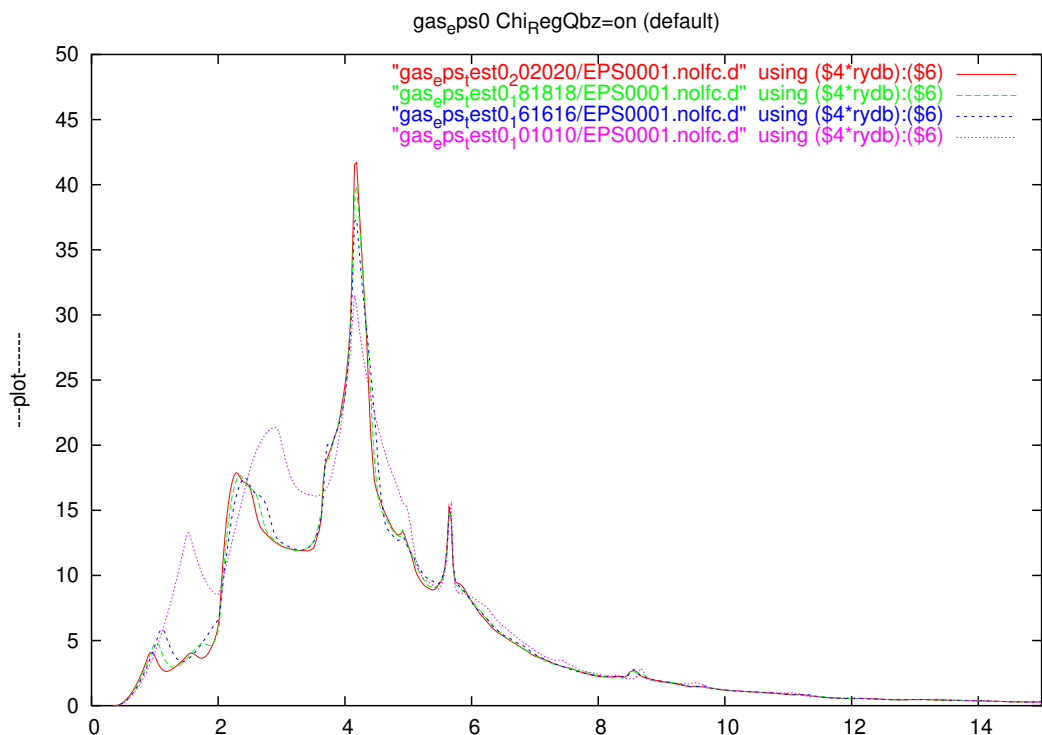


fig001

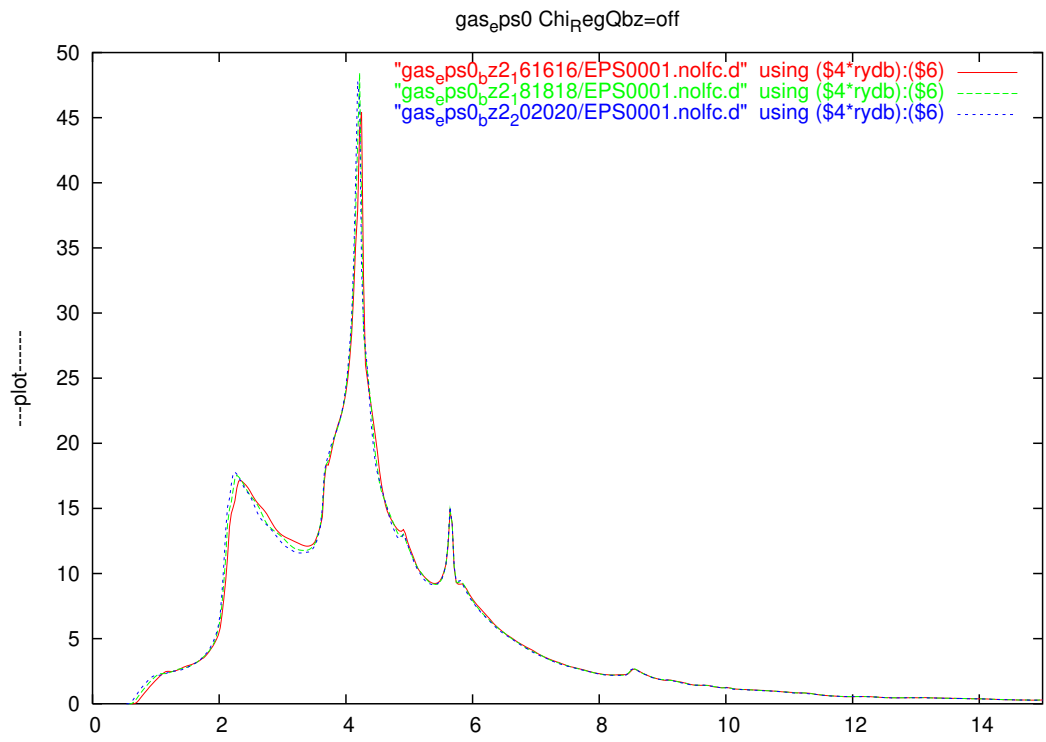


fig002

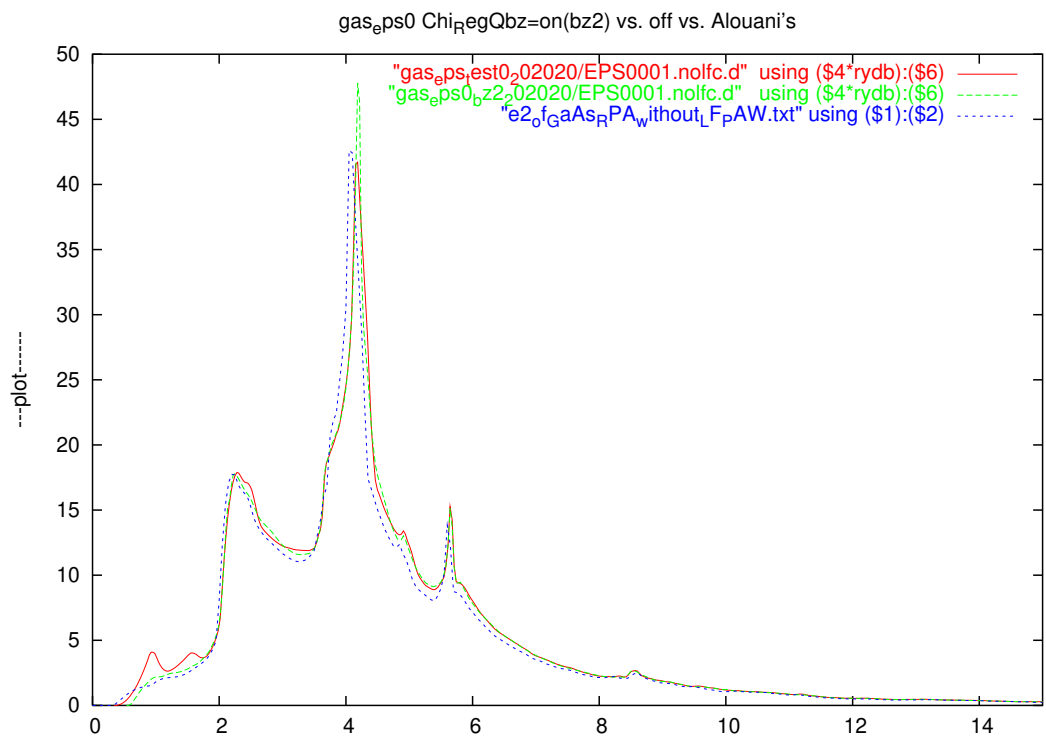


fig003

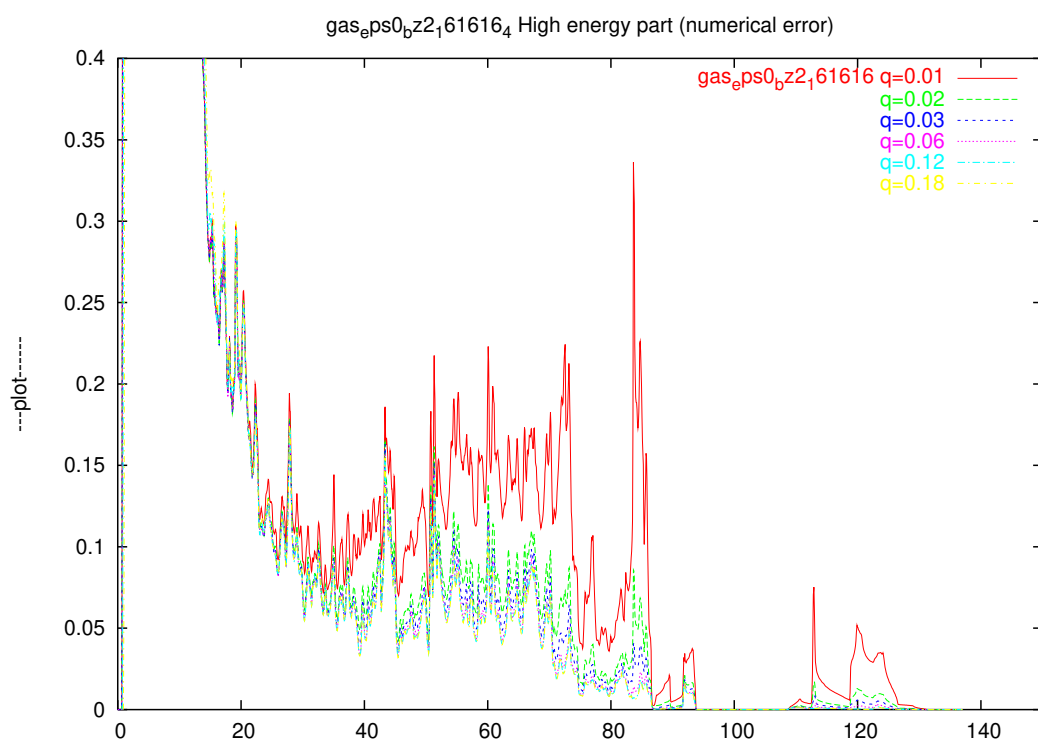


fig004

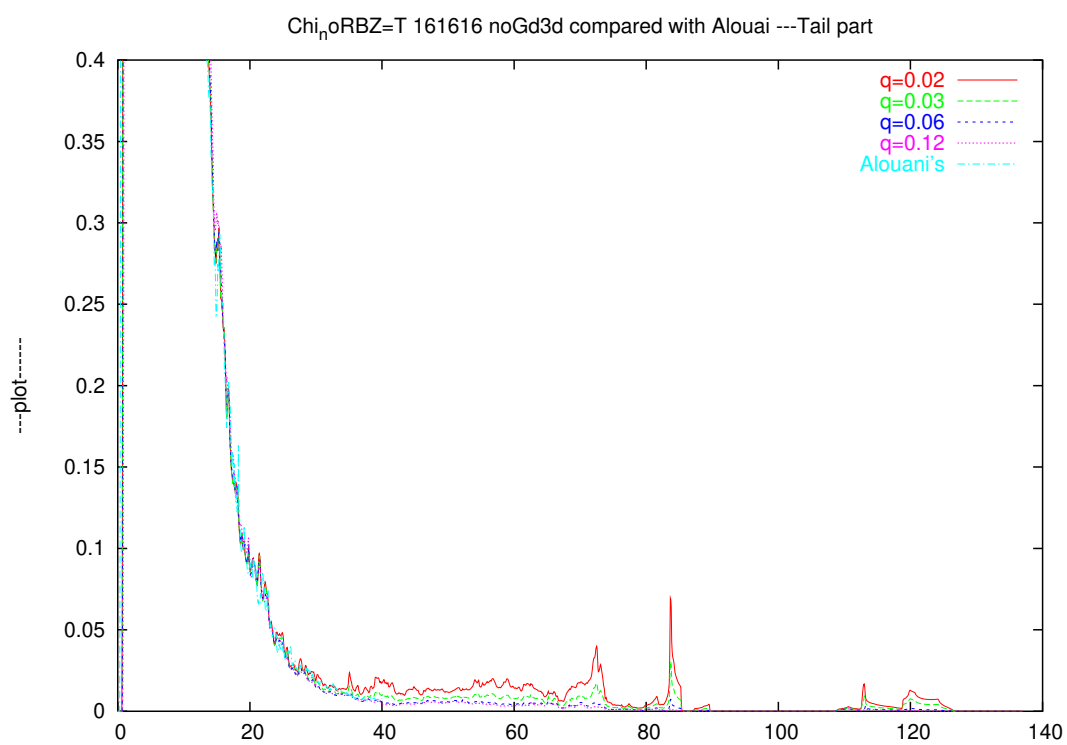


fig005

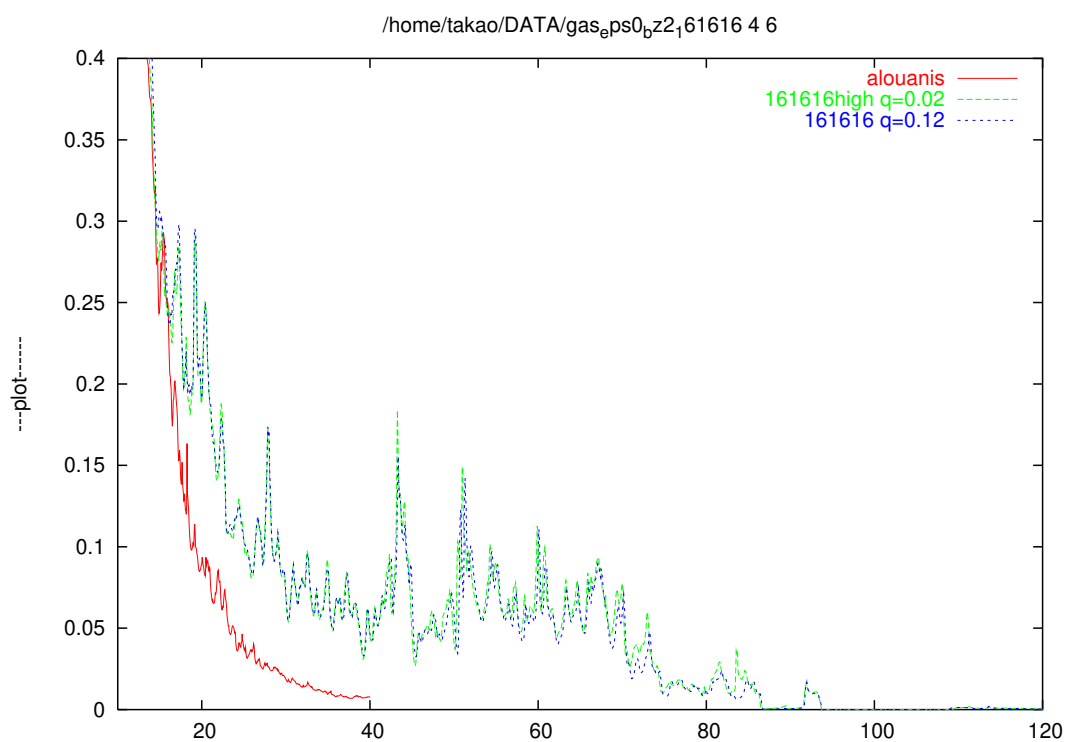


fig007

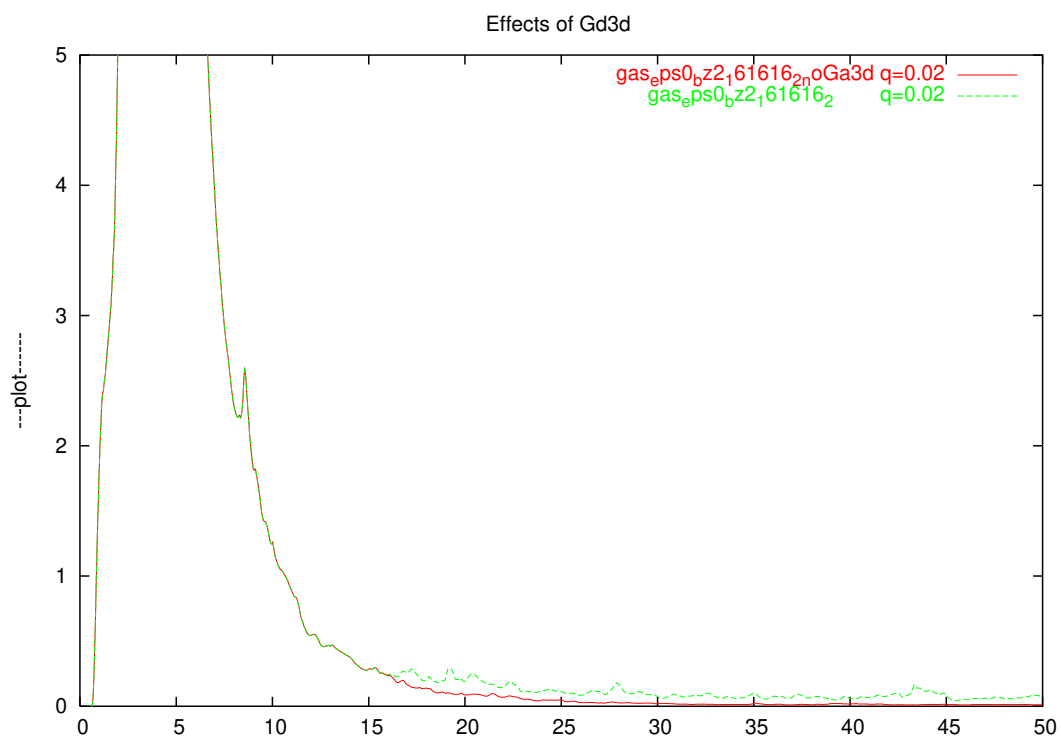


fig008

17 Utility

1. **structure tool** `viewvesta,ctrl2vasp,vasp2ctrl`
2. **Other calculation examples** A recent paper by Deguch et al [12] contains a link to a sample package.
3. **cif to PROCAR**
`./cif2cell --vasp-cartesian --vasp-format=5 cifs/BaTiO3_cubic.cif`

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