wannier90: A Tool for Obtaining
Maximally-Localised Wannier Functions

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Abstract

We present wannier90, a program for calculating maximally-localised Wannier functions (MLWF) from a set of Bloch energy bands that may or may not be attached to or mixed with other bands. The formalism works by minimising the total spread of the MLWF in real space. This done in the space of unitary matrices that describe rotations of the Bloch bands at each k-point. As a result, wannier90 is independent of the basis set used in the underlying calculation to obtain the Bloch states. Therefore, it may be interfaced straightforwardly to any electronic structure code. The locality of MLWF can be exploited to compute band-structure, density of states and Fermi surfaces at modest computational cost. Furthermore, wannier90 is able to output MLWF for visualisation and other post-processing purposes. Wannier functions are already used in a wide variety of applications. These include analysis of chemical bonding in real space; calculation of dielectric properties via the modern theory of polarisation; and as an accurate and minimal basis set in the construction of model Hamiltonians for large-scale systems, in linear-scaling quantum Monte Carlo calculations, and for efficient computation of material properties, such as the anomalous Hall coefficient. wannier90 is freely available under the GNU General Public License from http://www.wannier.org/.

Key words: Electronic structure, density-functional theory, Wannier function

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PROGRAM SUMMARY

Title of program: wannier90

Catalogue identifier:

Program Summary URL:

Program obtainable from: CPC Program Library, Queen’s University of Belfast, N. Ireland, or from the web-page [http://www.wannier.org/]

Licensing provisions: This program is distributed under then GNU General Public License v2.0 (see [http://www.gnu.org/] for details)

Computers for which the program has been designed and others on which it has been operable: any architecture with a Fortran 90 compiler

Operating systems under which the program has been tested: Linux, Windows, Solaris, AIX, Tru64 Unix, OSX

Programming Languages used: Fortran 90, perl

Libraries required:

- BLAS ([http://www.netlib.org/blas])
- LAPACK ([http://www.netlib.org/lapack])

Both available under open-source licenses.

Memory used to execute with typical data: 10 Mb

CPU time required to execute test cases: 1 min

No.of bits in a word: 32 or 64

Has the code been vectorised or parallelised?: No

Number of bytes in distributed program including test data, etc.: 4 600 000

Distribution format: gzipped tar

Keywords: Electronic structure, density-functional theory, Wannier functions

Nature of physical problem: Obtaining maximally-localised Wannier functions from a set of Bloch energy bands that may or may not be entangled.

Method of solution: In the case of entangled bands, the optimally-connected
subspace of interest is determined by minimising a functional which measures the subspace dispersion across the Brillouin zone. The maximally-localised Wannier functions within this subspace are obtained by subsequent minimisation of a functional that represents the total spread of the Wannier functions in real space. For the case of isolated energy bands only the second step of the procedure is required.

Unusual features of the program: Simple and user-friendly input system. Wannier functions and interpolated band structure output in a variety of file formats for visualisation.

References:


LONG WRITE-UP

1 Introduction

Within the independent-particle approximation, the electronic ground state of a periodic system may be solved in terms of a set of extended Bloch states $\psi_{nk}(r)$. These states are characterised by the good quantum numbers $n$ and $k$, which refer to the band index and crystal momentum, respectively. Although this choice is widely used in electronic structure calculations, alternative representations are available. For example, the Wannier representation constitutes a description in terms of localised functions labeled by $R$, the lattice vector of the cell in which the function is localised, and a band-like index $n$.

Wannier functions give a real-space picture of the electronic structure of a system. They provide insight into the nature of the chemical bonding and can be a powerful tool in the study of dielectric properties via the modern theory of polarisation.

The phase indeterminacy $e^{i\phi_n(k)}$ of an isolated Bloch state $\psi_{nk}(r)$ at each wave-vector $k$ results in the Wannier functions being non-unique. For a group of isolated bands, such as the valence states of an insulator, this indeterminacy is more general as they may undergo arbitrary unitary transformations $U_{mn}^{(k)}$ amongst themselves at each $k$. Two of the authors (NM and DV) developed a procedure that iteratively refines these degrees of freedom such that they lead
to Wannier functions that are well-defined and localised (in the sense that they minimise the second moment around their centres) [1]. We refer to this procedure as the Marzari-Vanderbilt (MV) scheme and the resulting Wannier functions as maximally-localised Wannier functions (MLWF).

The MV method was designed to be applicable to isolated sub-groups of bands, i.e., a group of bands that, though they may have degeneracies with each other at certain points, are separated from all other bands by finite gaps throughout the Brillouin zone. An important example is the set of valence bands of an insulator. In many applications, however, the bands of interest are not isolated and one is interested, for instance, in the partially filled bands of a metal close to the Fermi level. In this case the bands of interest lie within a limited energy range but are attached to, or cross with, other bands that extend further out in energy. Such bands are referred to as entangled bands.

The complication associated with treating entangled bands is that the states of interest hybridise with unwanted bands. As a result, the number of bands at each k-point in the relevant energy range may be greater than the number of Wannier functions $N$ that are required. Before the MV localisation procedure may be applied, a prescription for extracting $N$ bands at each k-point from the entangled manifold of states is required. Three of the authors developed just such a disentanglement procedure, allowing MLWF to be determined from a set of entangled bands [2]. This Souza-Marzari-Vanderbilt (SMV) strategy breaks down the procedure into two steps: first, the correct $N$-dimensional subspace of bands at every k is selected; and second, $N$ MLWF are localised from this subspace, exactly in the same way as for an isolated set of bands.

wannier90 is a tool for obtaining MLWF from a set of (possibly entangled) energy bands using the methods of MV and SMV. The principal ingredient that is required from an electronic structure calculation is the overlap matrix between the periodic part $|u_{nk}\rangle$ of Bloch states at neighbouring k-points (described in more detail in Section 2). This matrix is small and independent of the basis used in the underlying calculation to obtain the Bloch states. As a result wannier90 may be interfaced to any electronic structure code. At the time of writing, wannier90 is able to:

1. Disentangle an optimally-connected $N$-dimensional subspace of bands at each k-point and obtain the unitary transformations that generate MLWF from a given set of bands;
2. Output MLWF in a number of formats suitable for visualisation;
3. Generate interpolated band structures, densities of states and Fermi surfaces and output them in formats suitable for visualisation;
4. Output matrix elements of the Hamiltonian operator in the MLWF basis.

It is worth recording here a brief historical timeline of wannier90:
• **1997:** Two of the authors (NM and DV) develop and implement localisation algorithms in Fortran77; interface to the all-bands ensemble-DFT CASTEP code [3];

• **1998:** Algorithms implemented in the CPMD code [4] by Silvestrelli [5];

• **2001:** Three of the authors (IS, NM and DV) develop the SMV disentanglement extension; interface to a full-potential-linearized-augmented-plane-wave (FLAPW) code [6] and to any generic electronic-structure code quickly followed;

• **2006:** Two of the authors (AAM and JRY) completely rewrite the code in Fortran90, employing modern programming techniques and adding new functionality.

One significant benefit of the 2006 rewrite has been that in most cases wannier90 is faster than the original Fortran77 version by an order of magnitude, and in some by more than two. Another is that now the source code is easy to follow, which makes interfacing it to electronic structure codes more straightforward. Indeed, the first such example is already in place within the QUANTUM-ESPRESSO package [7].

The formalism has seen many and diverse applications: linear-scaling quantum Monte Carlo [8], photonic crystals [9,10], metal-insulator interfaces [11], as an efficient interpolator for the anomalous Hall effect [12] and electron-phonon couplings [13], and a powerful tool for the study of large-scale systems [14,15], to cite only a few. In addition, MLWF are playing an increasing role in bridging density-functional approaches and strongly-correlated ones, to derive model Hamiltonians or as a starting point for LDA+U or LDA+DMFT [16,17,18]. They are also closely related to muffin-tin orbitals [19,20].

The remainder of this paper is organised as follows. In Section 2 we review briefly the theoretical background to the problem of obtaining MLWF and a recent theoretical development in using them to interpolate band structures on arbitrarily fine k-point meshes. In Section 3 we give an overview of some of the more technical aspects of the wannier90 code. In Section 4 we describe the overall structure of the code. We demonstrate how to install and run wannier90 in Sections 5 and 6, respectively. Finally, in Section 7 we provide a number of example calculations and applications of the code.

## 2 Theoretical Background

For an isolated set of \( N \) Bloch bands \( \psi_{nk}(\mathbf{r}) \), a set of \( N \) Wannier functions \( w_{nR}(\mathbf{r}) = w_n(\mathbf{r} - \mathbf{R}) \), \( n \in [1,N] \), labelled by Bravais lattice vectors \( \mathbf{R} \), may
be constructed as

\[ |w_{nR}⟩ = \frac{V}{(2\pi)^3} \int_{\text{BZ}} \left[ \sum_{m=1}^{N} U_{mn}^{(k)} |ψ_{mk}⟩ \right] e^{-i\mathbf{k} \cdot \mathbf{R}} d\mathbf{k}, \tag{1} \]

where \( U^{(k)} \) is a unitary matrix that mixes the bands at wave-vector \( \mathbf{k} \), and the Brillouin zone (BZ) integral may also be interpreted as a unitary transformation. Different choices for \( U^{(k)} \) give rise to different Wannier functions, with different spatial spreads, that always describe the same manifold. The MV strategy consists of choosing the \( U^{(k)} \) that minimise the sum of the quadratic spreads of the Wannier functions about their centres:

\[
Ω = \sum_{n} \left( ⟨(\mathbf{r} - \mathbf{r}_n)^2⟩_n \right) = \sum_{n} \left( r^2 - 2\mathbf{r} \cdot \mathbf{r}_n + |\mathbf{r}_n|^2⟩_n \right) = \sum_{n} \left( ⟨r^2⟩_n - |\mathbf{r}_n|^2 \right) \tag{2} \]

where \( ⟨\hat{O}⟩_n \equiv \overline{\mathbf{r}}_n \equiv ⟨w_{n0}|\hat{O}|w_{n0}⟩ \).

It is worth noting that the spread functional may be decomposed into two contributions

\[
Ω = Ω_1 + Ω, \tag{3} \]

where

\[
Ω_1 = \sum_{n} \left[ ⟨w_{n0}|r^2|w_{n0}⟩ - \sum_{mR} |⟨w_{mR}|r|w_{n0}⟩|^2 \right] \tag{4} \]

and

\[
Ω = \sum_{n} \sum_{mR \neq n0} |⟨w_{mR}|r|w_{n0}⟩|^2. \tag{5} \]

It can be shown that each of these quantities is non-negative and that \( Ω_1 \) is gauge invariant, i.e., insensitive to changes in the matrices \( U_{mn}^{(k)} \) [1]. Therefore, the minimisation of the spread functional for an isolated set of bands just corresponds to minimising \( Ω \).
2.1 Reciprocal-Space Formulation

As shown by Blount [21], matrix elements of the position operator between Wannier functions may be expressed in reciprocal-space as

\[ \langle w_n | r | w_m \rangle = i \frac{V}{(2\pi)^3} \int e^{ik \cdot R} \langle u_n | \nabla_k | u_m \rangle \, dk \tag{6} \]

and

\[ \langle w_n | r^2 | w_m \rangle = -i \frac{V}{(2\pi)^3} \int e^{ik \cdot R} \langle u_n | \nabla^2_k | u_m \rangle \, dk, \tag{7} \]

where, as usual, the periodic part of the Bloch function is defined as \( u_{nk}(r) = e^{-i k \cdot r} \psi_{nk}(r) \). These expressions enable us to express the spread functional in terms of matrix elements of \( \nabla_k \) and \( \nabla^2_k \).

It is assumed that the Brillouin zone is discretised on a uniform Monkhorst-Pack mesh [22] on which the Bloch orbitals are determined. Thus, the gradient and Laplacian operators may be approximated by finite-difference formulas on the \( k \)-space mesh. If \( f(k) \) is a smooth function of \( k \), then

\[ \nabla_k f(k) = \sum_b \omega_b b [f(k + b) - f(k)] \tag{8} \]

and

\[ \langle f(k) | \nabla^2_k | f(k) \rangle = \langle \nabla_k f(k) \rangle^2 = \sum_b w_b [f(k + b) - f(k)]^2, \tag{9} \]

where \( \{b\} \) are vectors connecting mesh-point \( k \) to its nearest neighbours and \( \omega_b \) is a weight factor associated with each shell of neighbours \( b = |b| \). The choice of \( b \)-vectors and weights \( \omega_b \) is discussed in further detail in Section 3.2.

It is worth noting that a finite \( k \)-point mesh implies an approximation to both the self-consistent ground state that is obtained using that mesh and to the accuracy of the finite-difference representation of the operators \( \nabla_k \) and \( \nabla^2_k \). In principle, for a given mesh of \( k \)-points, the latter may be improved by using higher-order finite-difference expressions.

Having discretised Eqns. 6 and 7 in reciprocal space, the only information required for computing the spread functional is the overlap matrix

\[ M^{(k,b)}_{mn} = \langle u_{nk} | u_{m,k+b} \rangle. \tag{10} \]
After some algebra, the two components of the spread functional may be expressed as

$$\Omega_I = \frac{1}{N_{kp}} \sum_{k,b} \omega_b \sum_{m=1}^{N} \left[ 1 - \sum_{n=1}^{N} |M^{(k,b)}_{mn}|^2 \right]$$

and

$$\tilde{\Omega} = \frac{1}{N_{kp}} \sum_{k,b} \omega_b \left[ \sum_{n=1}^{N} (-\text{Im} \ln M^{(k,b)}_{nn} - \mathbf{b} \cdot \mathbf{r}_n)^2 + \sum_{m \neq n} |M^{(k,b)}_{mn}|^2 \right],$$

where $N_{kp}$ is the number of k-points in the Monkhorst-Pack grid and $\mathbf{r}_n$ is the centre of the $n^{th}$ Wannier function, given by

$$\mathbf{r}_n = -\frac{1}{N_{kp}} \sum_{k,b} \omega_b \mathbf{b} \text{Im} \ln M^{(k,b)}_{nn}.$$  

Using these expressions, the gradient of the spread functional with respect to infinitesimal unitary rotations of the $\psi_{n,k}(\mathbf{r})$ may be calculated as a function of $M^{(k,b)}_{mn}$. It is then straightforward to evolve $U^{(k)}_{mn}$ (and consequently $M^{(k,b)}_{mn}$) towards the solution of maximum localisation using a steepest-descents or conjugate-gradient minimisation algorithm.

It is worth noting that $\tilde{\Omega}$ may be further decomposed into band-diagonal and band-off-diagonal parts

$$\tilde{\Omega} = \Omega_D + \Omega_{OD},$$

where

$$\Omega_D = \sum_n \sum_{\mathbf{R} \neq 0} |\langle w_{nR} | \mathbf{r} | w_{n0} \rangle|^2$$

$$= \frac{1}{N_{kp}} \sum_{k,b} \omega_b \sum_{n=1}^{N} (-\text{Im} \ln M^{(k,b)}_{nn} - \mathbf{b} \cdot \mathbf{r}_n)^2,$$

$$\Omega_{OD} = \sum_{m \neq n} \sum_{\mathbf{R}} |\langle w_{mR} | \mathbf{r} | w_{n0} \rangle|^2$$

$$= \frac{1}{N_{kp}} \sum_{k,b} \omega_b \sum_{n=1}^{N} \sum_{m \neq n} |M^{(k,b)}_{mn}|^2.$$
2.2 The Case of Entangled Bands

The above description is sufficient to obtain ML WF for an isolated set of bands, such as the valence states in an insulator. In order to obtain ML WF for entangled energy bands, e.g., for metallic systems or the conduction bands of an insulator, we use the “disentanglement” procedure introduced by SMV [2].

The strategy proceeds as follows. An energy window (the “outer window”) is defined by the user such that at each k-point there are \( N_{\text{win}}^{(k)} \geq N \) states within the window. An orthonormal set of \( N \) Bloch states is obtained at each k-point, defining an \( N \)-dimensional subspace \( S(k) \), by performing a unitary transformation amongst the \( N_{\text{win}}^{(k)} \) states that fall within the energy window:

\[
|u_{nk}^{\text{opt}}\rangle = \sum_{m \in N_{\text{win}}^{(k)}} U_{mn}^{\text{dis}(k)} |u_{mk}\rangle,
\]

where \( U^{\text{dis}(k)} \) is a rectangular \( N_{\text{win}}^{(k)} \times N \) matrix.\(^1\)

Recall that \( \Omega_{I} \) is invariant under gauge transformations within a given subspace. Thus \( \Omega_{I} \) may be considered as a functional of \( S(k) \). The subspace selection proceeds by minimising \( \Omega_{I} \) with respect to the matrices \( U^{\text{dis}(k)} \) [2]. For a physical interpretation of this procedure, Eqn. 11 for \( \Omega_{I} \) may be rewritten as

\[
\Omega_{I} = \frac{1}{N_{kp}} \sum_{k,b} \omega_{\beta} \text{tr} \left[ \hat{P}_{k} \hat{Q}_{k+b} \right],
\]

where \( \hat{P}_{k} = \sum_{n=1}^{N} |u_{nk}\rangle \langle u_{nk}| \) is the projector onto \( S(k) \), and \( \hat{Q}_{k} = 1 - \hat{P}_{k} \). Now it can be seen that \( \Omega_{I} \) is a measure of the degree of mismatch between the subspaces \( S(k) \) and \( S(k+b) \). Minimising \( \Omega_{I} \) corresponds to choosing self-consistently at every \( k \) the subspace that has maximum overlap as \( k \) is varied.

Once the projected \( N \)-dimensional subspace at each k-point has been found, the MV localisation procedure described above may be used to minimise \( \Omega \) within that subspace and hence obtain MLWF. As an alternative to this two step minimisation, a procedure to minimise the total spread function \( \Omega \) has been proposed [23] and was found to give very similar MLWF to the present scheme.

\(^1\) As \( U^{\text{dis}(k)} \) is rectangular, this is a unitary operation in the sense that \( (U^{\text{dis}(k)})^{\dagger} U^{\text{dis}(k)} = I \).
It may be the case that the energy bands of the projected subspace do not correspond to any of the original energy bands due to mixing between states. In order to preserve exactly the properties of a system within a given energy range (e.g., around the Fermi level) a second inner energy window is introduced. States lying within this inner, or frozen, energy window are included unchanged in the projected subspace. The reader is referred to Ref. [2] for further details of the algorithm.

It is worth noting that the MLWF themselves are never explicitly constructed unless required for visualization or other post-processing purposes. Minimisation of the spread functional results in finding the converged \( U^{\text{dis}(k)} \) (if disentanglement was used) and \( U^{(k)} \). Along with \( M^{(k, b)} \), this is sufficient to define the centres and spreads of the MLWF. If the periodic parts of the Bloch wavefunctions are available, then the MLWF may be calculated on a real-space grid. For systems with time-reversal symmetry, we always find the MLWF corresponding to the minimum spread to be real, apart from a global complex phase factor, in empirical agreement with a recent claim of a mathematical proof [24].

### 2.3 Wannier Interpolation

Having found \( U^{\text{dis}(k)} \) and \( U^{(k)} \) for the target system, it is straightforward to express the Hamiltonian in the basis of MLWF. We first obtain the Hamiltonian in the basis of rotated Bloch states

\[
H^{(W)}(k) = (U^{(k)})^\dagger (U^{\text{dis}(k)})^\dagger H(k) U^{\text{dis}(k)} U^{(k)}
\]  

(21)

where \( H_{nm}(k) = \varepsilon_{nk} \delta_{nm} \). Next we find its Fourier sum

\[
H_{nm}(R) = \frac{1}{N_0} \sum_k e^{-i k \cdot R} H^{(W)}_{nm}(k).
\]  

(22)

This operation is done once and for all for each of the \( N_0 \) lattice vectors \( R \) lying in a supercell conjugate to the \( k \)-mesh (in practice we choose a Wigner-Seitz supercell centred on the origin [25]). \( H_{nm}(R) \) can be recognised as the matrix elements of the Hamiltonian between MLWF. Due to the strong localisation of the MLWF, the matrix elements \( H_{nm}(R) \) decay rapidly with \( R \). In the spirit of a Slater-Koster interpolation scheme [26] this allows us to compute the Hamiltonian on a much finer mesh of \( k \)-points in the original Bloch space. We can perform the inverse Fourier transform

\[
H_{nm}(k') = \sum_R e^{i k' \cdot R} H_{nm}(R),
\]  

(23)
to yield the interpolation of Eqn. 21 onto an arbitrary k-point $k'$. The final step is to diagonalise $H_{nm}(k')$ to obtain the interpolated band energies.

By construction, the interpolated band energies coincide with the true band-structure at the original k-points of the Monkhorst-Pack mesh (if the SMV scheme is used, this is only guaranteed to occur within the inner energy window). At intermediate k-points, the accuracy of the interpolation is dependent on the density of the original mesh [14,25].

Within wannier90 this interpolation procedure can be used to obtain plots of band structure, density of states and the Fermi surface at modest computational cost. The Wannier interpolation formalism is rather general and can also be used to interpolate arbitrary periodic operators [25].

3 Some Technical Aspects

3.1 Initial guess for iterative minimisation

The iterative minimisation of $\Omega_i$ begins with an initial guess for the subspaces $S(k)$. One possible choice is to start from the initial, random phases of the Bloch states provided by the ab initio code. Alternatively, we may define a set of $N$ trial functions $g_n(r), n \in [1, N]$, as an initial approximation to the $N$ MLWF. These are projected onto the cell-periodic parts of the $N^{(k)}_{\text{win}}$ Bloch eigenstates inside the energy window:

$$|\phi_{nk}\rangle = \sum_{m=1}^{N^{(k)}_{\text{win}}} A_{mn}^{(k)} |u_{mk}\rangle,$$  \hspace{1cm} (24)

where $A_{mn}^{(k)} = \langle u_{mk}| g_n \rangle$ is an $N^{(k)}_{\text{win}} \times N$ matrix. Orthonormalising the resulting $N$ functions $\{|\phi_{nk}\rangle\}$ via a Löwdin transformation, we find

$$|u_{nk}^{\text{opt}}\rangle = \sum_{m=1}^{N} (S^{-1/2})_{mn} |\phi_{mk}\rangle = \sum_{m=1}^{N^{(k)}_{\text{win}}} (AS^{-1/2})_{mn} |u_{mk}\rangle,$$ \hspace{1cm} (25)

where $S_{mn} = S_{mn}^{(k)} = \langle \phi_{mk}| \phi_{nk} \rangle = (A^\dagger A)_{mn}$, and $AS^{-1/2}$ is used as the initial guess for $U_{\text{dis}}^{(k)}$.

The same trial orbitals $\{|g_n\rangle\}$ can also be used to initialize the minimization of $\tilde{\Omega}$. Using a similar projection procedure to the one described above, an initial guess for the $N \times N$ unitary matrices $U^{(k)}$ is obtained at each k-point.
We now come to the choice of trial orbitals. As the minimisation scheme is quite robust, one option is to choose a set of spherically-symmetric Gaussian functions whose centres are chosen randomly; \texttt{wannier90} supports this option. On the other hand, a user may wish to apply chemical and physical insight in order to select a better starting point. An \textit{ab initio} code that is interfaced to \texttt{wannier90} may use any localised functions desired to construct the projections $A_{nm}^{(k)}$. For convenience, we have defined a standard set of projection functions that should suffice for most situations. This is the set of atomic orbitals of the hydrogen atom, which is a convenient choice for two reasons: first, analytical mathematical forms exist in terms of the good quantum numbers $n$, $l$ and $m$; and second, hybrid orbitals (sp, sp$^2$, sp$^3$, sp$^3$d, etc.) may be constructed by simple linear combination $|\phi\rangle = \sum_{nlm} C_{nlm} |nlm\rangle$, for some coefficients $C_{nlm}$. The angular parts of our projection functions are not the canonical spherical harmonics $Y_{lm}$ of the hydrogenic Schrödinger equation, but rather the \textit{real} (in the sense of non-imaginary) states $\Theta_{lm}$, $m_r \in [1, 2l + 1]$, obtained by unitary transformation of the $Y_{lm}$.

Each localised function is associated with a site and an angular momentum state. Optionally, one may define the spatial orientation, the radial form and the diffusivity for each function. \texttt{wannier90} is able to project onto functions with s, p, d and f symmetry, plus the hybrids sp, sp$^2$, sp$^3$, sp$^3$d, sp$^3$d$^2$. The user is referred to the documentation in the \texttt{wannier90} distribution for mathematical definitions and details on how to specify projection functions in the input file.

In general, the spread functional $\Omega$ has local minima and, occasionally, the minimisation becomes trapped in one. In other words, the final solution may depend on the initial choice for $S(k)$ and hence $\{g_n(r)\}$. In most cases, we find that these local minima give rise to MLWF that are complex, i.e., they have significant imaginary parts. In some cases, MLWF associated with local minima of $\Omega$ are found to be real and the reader is referred to Ref. [27] for more details.

### 3.2 Choosing the $b$ vectors

As discussed in Section 2.1, in order to compute the spread functional we require a finite-difference formula for $\nabla_k$. We now describe an automatic procedure to choose, for cells of arbitrary symmetry, the simplest such formula.

It is assumed that the Brillouin zone is discretised on a uniform Monkhorst-Pack mesh [22]. The vectors $\{b\}$ connect each mesh-point $k$ to its nearest neighbours. $N_{sh}$ shells of neighbours are included in the finite-difference formula, with $M_s$ vectors in the $s^{th}$ shell. For $\nabla_k$ to be correct to linear order we
require that (see Eqn. B1 of Ref. [1])

\[
\sum_{s} N_{sh} w_{s} \sum_{i} b_{i,s}^{\alpha} b_{i,s}^{\beta} = \delta_{\alpha \beta}
\]  

(26)

where \( b_{i,s}^{\alpha}, i \in [1, M_s], \) is the \( i \)th vector belonging to the \( s \)th shell with associated weight \( w_{s}, \) and \( \alpha \) and \( \beta \) run over the three Cartesian indices. If \( f(k) \) is a smooth function of \( k, \) then its gradient may be expressed according to Eqn. 8. For the case of a linear function \( f(k) = f_0 + g \cdot k, \) this finite-difference formula gives the exact result \( \nabla_{k,\alpha} = \sum b_{i,s} w_{i,s} b_{i,s}^{\beta} = \delta_{\alpha \beta} g_{\beta} = g_{\alpha}, \) where summation convention over repeated Greek indices is assumed.

In order to find the weights \( w_{s}, \) we notice that Eqn. 26 is symmetric in the Cartesian indices, therefore, there are six independent elements that may be expressed through a matrix equation

\[
Aw = q.
\]  

(27)

where \( q \) is a vector of length six containing the six elements of the lower triangular part of \( \delta_{\alpha \beta}. \) \( w \) is the vector of weights with length \( N_{sh}, \) and \( A \) is a \( 6 \times N_{sh} \) matrix given by \( A_{j,s} = \sum_{i} b_{i,s}^{\alpha} b_{i,s}^{\beta}, \) where there is a one-to-one correspondence between \( j \in [1, 6] \) and the six independent pairings of \( \alpha \) and \( \beta. \) \( A \) may be factorised using a singular value decomposition [28],

\[
A = UDV^{T},
\]  

(28)

which permits inversion of Eqn. 27 and solution for the shell weights,

\[
w = VD^{-1}U^{T}q.
\]  

(29)

Our automatic procedure for choosing the shells is as follows: we add shells in order of increasing \( |b|; \) with each additional shell we obtain the shell weights, and check to see if Eqn. 26 is satisfied; if not, then we add a new shell and repeat the procedure. A new shell may be linearly dependent on the existing shells, in which case one or more of the singular values will be close to zero and we reject the new shell. When the Bravais lattice point group is cubic, the first shell of nearest neighbours is sufficient. The maximum number of shells required is six (for triclinic symmetry). For the case of a very elongated unit cell it may be necessary for the routine to search through a large number of shells in order to find the correct combination.

With the above finite-difference formalism, the quadratic spread converges only polynomially with the sampling of the Brillouin zone. This slow convergence is a property of the spread operator, whereas the underlying MLWF
converge rapidly with the k-point density [29]. It may be possible to achieve improved accuracy and k-point convergence by using higher-order finite-difference formulas for $\nabla_k$, but this has not been explored.

In some cases, the automatic procedure outlined above might not find the shells and weights that give the most spherically symmetric representation of the position operator. When this happens, special care may be required to ensure that the most symmetric choice of shells is made [6], and this can be done explicitly (i.e., by hand) in the input file.

3.3 Specific Algorithms for Γ-Point Sampling

For isolated molecules or extended systems of large unit cell, single Γ-point sampling in the reciprocal space can provide an accurate description of the physical quantities of interest. When the Γ-point is sampled exclusively, the computational cost can be reduced by exploiting the symmetries of the overlap matrices $M^{(\Gamma,b)}$.

First, at Γ, the $b$-vectors are linear combinations of the primitive vectors of the reciprocal lattice, thus imposing the following conditions:

$$\psi_{n,\Gamma+b}(r) = \psi_{n,\Gamma}(r), \quad u_{n,\Gamma+b}(r) = e^{-ib \cdot r} u_{n,\Gamma}(r).$$

Then, it follows that $M^{(\Gamma,-b)}$ is the Hermitian conjugate of $M^{(\Gamma,b)}$,

$$M_{nm}^{(\Gamma,b)} = \langle u_m \Gamma | u_n,\Gamma+b \rangle = \langle u_n,\Gamma+b | u_m \Gamma \rangle^* = \langle u_n \Gamma | u_m,\Gamma-b \rangle^* = (M_{nm}^{(\Gamma,-b)})^*.$$  \hspace{1cm} (31)

Second, at Γ, the Bloch eigenfunctions may be chosen to be real. Then, the overlap matrices become symmetric, $M_{nn}^{(\Gamma,b)} = M_{nn}^{(\Gamma,-b)}$, and only the upper or the lower half of $M^{(\Gamma,b)}$ is independent.

wannier90 includes a Γ-only branch of algorithms that is able to exploit these symmetries. These algorithms may be activated by means of a logical keyword in the input file and they rely on the above relations being satisfied, i.e., the Bloch wavefunctions must be real.

There are several other advantages beside the significant decrease in the number of operations. In the disentanglement procedure [2], the diagonalisation of a complex Hermitian matrix at each iteration step is replaced by that of a real symmetric matrix, which ensures that $U_{\text{dis}(\Gamma)}$ is real. The localisation procedure in the Γ-only branch adopts an efficient algorithm proposed by Gygi et
This method minimises $\Omega_{OD}$ (not $\tilde{\Omega}$) by simultaneously diagonalizing a set of real symmetric matrices, $\{\text{Re} M^{(\Gamma,b)}, \text{Im} M^{(\Gamma,b)}\}$. The $U^{(\Gamma)}$ from this method is inherently real, giving real MLWF as a final product.

It is worth mentioning that these MLWF are not necessarily identical to those obtained from the MV localisation procedure unless $\Omega_{D}$ is strictly zero due to symmetry of the system.

### 3.4 Representative Timings and Convergence

In this section, timings and convergence characteristics for representative calculations are presented. The initial band-structure calculations are performed using the 	extsc{pwscf} code [7]. The self-consistent ground state is obtained, then the required overlap matrices and projections are calculated using the post-processing routine 	extsc{pw2wannier90}, supplied with the 	extsc{pwscf} distribution. 	extsc{wannier90} is then used to obtain the MLWF. The 	extsc{pwscf} and 	extsc{pw2wannier90} calculations were performed on a four-processor (dual dual-core) Intel Woodcrest 5130 2.0GHz desktop computer and the 	extsc{wannier90} calculations on a single processor of the same machine. The gauge-invariant and gauge-dependent spreads were converged to a tolerance of $10^{-10}$ Å$^2$.

#### 3.4.1 Crystalline Silicon

First, we consider a two-atom unit cell of crystalline silicon. A kinetic-energy cutoff of 25 Ry is used for the plane-wave expansion of the valence wavefunctions. The core-valence interaction is described by means of norm-conserving pseudopotentials in separable Kleinman-Bylander form [31]. Table 1 gives timings for obtaining the valence MLWF for various densities of Monkhorst-Pack k-point grid. Bond-centred s-type functions were used for the initial projection. It can be seen that the most time consuming part of the procedure is the computation of the band structure, followed by construction of $M^{(k,b)}_{mn}$ and $A^{(k)}_{mn}$, with 	extsc{wannier90} taking only 5-7% of the total time to perform the localisation of the MLWF and to write files for their visualisation.

In Figure 1 the effect of using different initial projections for the MLWF is demonstrated. It is worth noting that even when choosing randomly-centred s-type functions the correct MLWF are found in less than 10% of the total time.
<table>
<thead>
<tr>
<th>$N_{kp}$</th>
<th>$T_{SCF}$</th>
<th>$T_{M,A}$</th>
<th>$T_{W90}$</th>
<th>$N_{iter}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>64</td>
<td>7</td>
<td>5</td>
<td>&lt; 1</td>
<td>5</td>
</tr>
<tr>
<td>512</td>
<td>52</td>
<td>44</td>
<td>4</td>
<td>14</td>
</tr>
<tr>
<td>1728</td>
<td>190</td>
<td>142</td>
<td>13</td>
<td>30</td>
</tr>
<tr>
<td>4096</td>
<td>462</td>
<td>329</td>
<td>56</td>
<td>96</td>
</tr>
</tbody>
</table>

Table 1
Timings (in seconds) for the valence states of crystalline silicon with (i) different densities of Monkhorst-Pack mesh. $T_{SCF}$ is the time taken by pwscf to obtain the ground state wavefunctions at all $N_{kp}$ k-points of the Monkhorst-Pack mesh, $T_{M,A}$ is the time taken by pw2wannier90 to calculate $M_{mn}^{(k,b)}$ and $A_{mn}^{(k)}$, and $T_{W90}$ is the time taken by wannier90 to localise and plot the MLWF. $N_{iter}$ is the number of iterations required to minimise the total spread.

Fig. 1. Convergence of total spread $\Omega$ for crystalline silicon with an $8 \times 8 \times 8$ Monkhorst-Pack k-point grid. Solid curve: bond-centred s projection. Dashed curve: atom-centred sp$^2$-like projection. Dotted curve: atom-centred s and p projection. Dot-dashed curve: randomly centred spherical s-type projection. $\Omega_0$ is the converged total spread and is independent of the initial projection used.

3.4.2 Fullerene

We consider an isolated fullerene molecule ($C_{60}$) as a test case for the $\Gamma$-only algorithms. The molecule was placed in a cubic supercell of side-length 40 a$_0$ (21.2 Å), and the calculations were performed with $\Gamma$-point sampling, ultrasoft pseudopotentials [32], and a plane-wave cut-off of 30 Ry.

Fullerene has 120 valence states in total. For the localisation of the valence MLWF, two choices of initial projections are compared, one being 120 randomly-centred s-orbitals and the other being 120 s-orbitals placed on the MLWF centres generated by the former run. The disentanglement procedure is tested on the extended 150 states comprising 120 valence states and 30 states that cover the full $\pi^*$-manifold. Those 30 states are disentangled from the 100 unoccupied eigenfunctions spanning up to 7.5 eV above the HOMO level. The
final set of MLWF are found to consist of 60 atom-centred \( p_z \) orbitals and 90 bond-centred \( \sigma \)-bonding orbitals, similar to the MLWF shown in Fig. 4.

<table>
<thead>
<tr>
<th>Projection</th>
<th>( \Gamma )</th>
<th>( N )</th>
<th>( T_{M,A} )</th>
<th>( T_{W90} )</th>
<th>( T_{\text{dis}} )</th>
<th>( N_{\text{iter}}^{\text{dis}} )</th>
<th>( T_{\text{loc}} )</th>
<th>( N_{\text{iter}}^{\text{loc}} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>randomly-centred s</td>
<td>F</td>
<td>120</td>
<td>901</td>
<td>33</td>
<td>n/a</td>
<td>n/a</td>
<td>33</td>
<td>226</td>
</tr>
<tr>
<td></td>
<td>T</td>
<td>120</td>
<td>686</td>
<td>1</td>
<td>n/a</td>
<td>n/a</td>
<td>1</td>
<td>16</td>
</tr>
<tr>
<td>bond-centred s</td>
<td>F</td>
<td>120</td>
<td>893</td>
<td>18</td>
<td>n/a</td>
<td>n/a</td>
<td>18</td>
<td>117</td>
</tr>
<tr>
<td></td>
<td>T</td>
<td>120</td>
<td>694</td>
<td>&lt; 1</td>
<td>n/a</td>
<td>n/a</td>
<td>&lt; 1</td>
<td>11</td>
</tr>
<tr>
<td>bond-centred s,</td>
<td>F</td>
<td>150</td>
<td>1595</td>
<td>46</td>
<td>22</td>
<td>75</td>
<td>24</td>
<td>83</td>
</tr>
<tr>
<td>atom-centred p_z</td>
<td>T</td>
<td>150</td>
<td>1017</td>
<td>12</td>
<td>11</td>
<td>75</td>
<td>1</td>
<td>9</td>
</tr>
</tbody>
</table>

Table 2
Timings (in seconds) for MLWF in fullerene from general k-point and \( \Gamma \)-specific schemes. Different initial projections are tested. “T” in the second column (\( \Gamma \)) indicates that the \( \Gamma \)-only branch of algorithms is used. \( N \) is the number of MLWF obtained. \( T_{M,A} \) is the time taken to calculate \( M_{mn}^{(k,b)} \) and \( A_{mn}^{(k)} \), and \( T_{W90} \) is the total time taken by \textit{wannier90} to disentangle and localise MLWF. The individual time and the number of iterations for each of these operations are given as \( T_{\text{oper}} \) and \( N_{\text{iter}}^{\text{oper}} \) (where \text{oper}=\text{dis} or \text{loc}), respectively.

Timings for these calculations are summarized in Table 2. \( T_{M,A} \) decreases by more than 20% when the \( \Gamma \)-only branch is used, because only half the \( b \)-vectors are needed. The post-processing routine \textit{pw2wannier90} is not yet optimized to take full advantage of real wavefunctions, and we expect at least 50% of time reduction in the fully optimized case.

The minimisation method used in the \( \Gamma \)-only branch is apparently more efficient than the conjugate-gradient algorithm, especially when the initial projections are far from the final, converged MLWF. \( \Omega_D \) vanishes in the case of a simple cubic lattice, and \( \Omega \) from the two different minimization methods converges to an identical value up to the given tolerance. \( T_{\text{dis}} \) decreases by 50% for the reason discussed in Section 3.3, but \( N_{\text{iter}}^{\text{dis}} \) is the same, as it should be, and \( \Omega_I \) is identical within machine precision.

4 Structure of the Program

The schematic structure of the program is outlined in Fig. 2. Each box represents a Fortran90 module and the lines represent module dependencies. Modules only use data and subroutines from lower modules in the diagram. A description of the purpose of each module is given below.

- **constants**: definition of constants (e.g., \( \pi \))
• io: error handling, timing, and input and output units
• utility: commonly used operations such as conversion of Cartesian to fractional co-ordinates, string functions, matrix multiplication wrapper etc.
• parameters: all physical parameters relevant to the calculation and subroutines for reading them from the input file at the start of a calculation. Subroutines for writing checkpoint files for restarting previous runs.
• kmesh: set up of the framework for reciprocal-space derivatives and, if in post-processing mode, writing a file which communicates to an ab initio code information on how to calculate $M_{mn}^{(k,b)}$ and $A_{mn}^{(k)}$
• overlap: reading of the overlap matrices $M_{mn}^{(k,b)}$ and $A_{mn}^{(k)}$
• disentangle: if using entangled energy bands, finding the optimal subspace within a specified energy window by minimising the gauge-invariant spread
• wannierise: finding the unitary transformations $U_{mn}^{(k)}$ amongst the energy bands which give rise to MLWF
• plot: routines for output of Wannier functions, Fermi surface and band structure in file formats suitable for visualisation
• wannier_prog: the main program
• wannier_lib: library routines. wannier90 may be invoked directly from within an ab initio code as a set of library calls. The reader is referred to Section 6.1 for a brief description and to the documentation in the wannier90 distribution for full details.

5 Installation

wannier90 is distributed as a gzipped tar file (http://www.wannier.org/). On Linux platforms, for example, it may be unpacked by typing

> tar zxvf wannier90.tar.gz

which creates a directory containing the source files, documentation, examples etc.

Compilation is straightforward. From the config directory of the distribution the user should select the make.sys.plat file which corresponds most closely to the platform being used and copy it to the root directory of the distribution, renaming it make.sys in the process. The values of system-dependent parameters (e.g., the location of the BLAS and LAPACK libraries, the name of the Fortran compiler, the Fortran optimisation flags etc.) that are defined therein should be modified to correspond to the user’s particular system. Once this has been done, typing make from the root directory of the distribution will create an executable wannier90.x. Further details and make options may be found in the file INSTALL.readme in the root directory of the distribution.
6 Running wannier90

Before running wannier90 the user must perform a self-consistent first-principles calculation on the system of interest in order to obtain a set of Bloch energy bands from which MLWF may then be constructed. Once the Bloch bands have been computed wannier90 may be operated in a post-processing mode as described below.

The master input file for wannier90 is called seedname.win, where seedname is the prefix of all of the input and output files. The input system is designed to be simple and user-friendly and is described comprehensively in the documentation that is contained within the wannier90 distribution. An example input file is provided in Appendix A.

wannier90 must be run twice. On the first pass the command line option -pp must be used, as follows:

> wannier90.x -pp seedname

This causes the code to read the master input file seedname.win and generate
the file `seedname.nnkp` that contains all the information necessary to construct the overlap matrices $M_{mn}^{(k,b)}$ and $A_{mn}^{(k)}$ from the Bloch bands already obtained from a first-principles calculation. In order to interface an ab initio code to `wannier90` one needs to write subroutines that read `seedname.nnkp`, compute the overlap matrix $M_{mn}^{(k,b)}$ and, optionally, $A_{mn}^{(k)}$ from the Bloch bands and write these matrices in the appropriate format to files called `seedname.mmn` and `seedname.amn`, respectively. If using the disentanglement procedure or plotting a band structure, density of states or Fermi surface, `wannier90` also requires the eigenvalues $\varepsilon_{nk}$ corresponding to the Bloch states $\psi_{nk}(r)$, which should be written to a file called `seedname.eig`. The reader is referred to the documentation that is contained within the `wannier90` distribution for complete details of these files.

Once the necessary files have been written, `wannier90` must be run again, this time without any command-line options:

```
> wannier90.x seedname
```

On this pass, the code reads the overlap matrices and eigenvalues (if required) from file and performs the maximal-localisation procedure as outlined in Section 2, writing the output to a file `seedname.wout`.

At the time of writing, the PWSCF code (a part of the QUANTUM-ESPRESSO package [7]), which is available under GNU General Public License, has a `wannier90` interface in the form of a post-processing program called `pw2wannier90`. It is the authors’ hope that `wannier90` is a sufficiently useful tool for investigators to be motivated to write interfaces to other electronic structure codes.

### 6.1 Library mode

`wannier90` may also be compiled as a library and invoked with subroutine calls from within an ab initio code. The command

```
> make lib
```

creates a library `libwannier.a` in the root directory of the distribution. The library mode of `wannier90` works along exactly the same lines as the post-processing mode, described above. The formal difference is that in the latter, information is passed between the ab initio code and `wannier90` via files, such as `seedname.nnkp` and `seedname.mmn` etc., whereas, in the former, it is done via direct calls to library subroutines. As before, there are two stages: first, a call to the library subroutine `wannier_setup`, which returns the information necessary to construct the overlap matrices $M_{mn}^{(k,b)}$ and $A_{mn}^{(k)}$ from the Bloch bands; second, a call to the subroutine `wannier_run`, which takes as input the
$M_{mn}^{(k,b)}$ matrix (and $A_{mn}^{(k)}$ and the eigenvalues $\varepsilon_{nk}$, if required) and performs the maximal-localisation procedure as outlined in Section 2. The reader is referred to the documentation in the wannier90 distribution for further details.

7 Examples

7.1 Graphite

As an example we consider the generation of MLWF to describe the states around and below the Fermi level in Bernal (A-B-A) graphite. From the band-structure (Fig. 3) we expect that the minimum number of Wannier functions needed to describe these states is 10 per unit cell (2.5 per atom). We shall see that this choice corresponds to an intuitive chemical description of the system.

We perform the initial band-structure calculations using the pwscf code [7]. A kinetic-energy cutoff of 30 Ry is used for the plane-wave expansion of the valence wavefunctions. The core-valence interaction is described by means of norm-conserving pseudopotentials in separable Kleinman-Bylander form [31]. We obtain the self-consistent ground state using a 16×16×16 Monkhorst-Pack mesh of k-points [22] and a fictitious Fermi smearing [3] of 0.02 Ry for the Brillouin-zone integration. Then, we freeze the self-consistent potential and perform a non-self-consistent calculation on a uniform 6×6×6 grid of k-points. At each k-point we calculate the first 20 bands. The required overlap matrices and projections are calculated using the post-processing routine pw2wannier90, supplied with the pwscf distribution. Projections onto atom centred sp$^2$ and p$_z$ functions are used to construct the initial guess, and wannier90 is used to obtain the MLWF. The gauge-dependent and gauge-independent spreads converge to machine precision in 300 and 70 steps, respectively. The resulting MLWF are a set of six symmetry-related bond-centred $\sigma$-orbitals and four atom-centred p$_z$-orbitals, shown in Fig. 4 (as plotted with the XCrySDen package [33]). Similar MLWF were obtained for carbon nanotubes [14]. In Fig. 3 we show the band structure of graphite obtained using Wannier interpolation and compare it to the band structure obtained from a full first-principles calculation. Within the inner energy window the agreement is essentially perfect.

7.2 Lead

Let us consider the generation of MLWF to describe the states around and below the Fermi level in lead. As relativistic effects can be significant in heavy
Fig. 3. Band structure of graphite. Solid lines: original band structure from a conventional first-principles calculation. Dotted lines: Wannier-interpolated band structure. The zero of energy is the Fermi level.

Fig. 4. Isosurface contours of MLWF in graphite (red for positive value and blue for negative). [left] $\sigma$-type MLWF. [right] $p_z$-type MLWF.

elements, we include spin-orbit coupling in our electronic structure calculation. In lead, the 6s and 6p states form an isolated set of bands around the Fermi level. The ground-state structure of lead has both time-reversal and inversion symmetries, so that each band is two-fold degenerate. We will obtain a set of eight MLWF to describe these states.

Once again we perform the initial band-structure calculations using the pwscf code. A kinetic-energy cutoff of 45 Ry is used for the plane-wave expansion of the valence wavefunctions. The core-valence interaction is described by means of spin-orbit coupled norm-conserving pseudopotentials [34] in separable Kleinman-Bylander form. We obtain the self-consistent ground state using a $16 \times 16 \times 16$ Monkhorst-Pack mesh of k-points and a fictitious Fermi smearing [3] of 0.02 Ry for the Brillouin-zone integration. The self-consistent potential is frozen and we perform a non-self-consistent calculation on a uniform $12 \times 12 \times 12$ grid of k-points. Then we calculate the required overlap matrices and projections using pw2wannier90. For the initial projection we use orbitals with sp$^3$ symmetry, four projections onto spin-up states and four onto spin-down states.

wannier90 is used to obtain the MLWF. The gauge-dependent spread is converged in 200 steps. In Fig. 5 we show both the band structure and Fermi sur-
face obtained using Wannier interpolation. Although the spin-orbit induced splitting is large at certain points in the band structure (e.g., 3 eV at Γ), the Fermi surface is not significantly different from a scalar-relativistic calculation. The reader is referred to Ref. [12] for a detailed discussion of how to construct spinor Wannier functions for ferromagnetic systems with broken time-reversal symmetry.

Fig. 5. [left] Band structure of lead. Solid lines: original band structure from a conventional first-principles calculation. Dotted lines: Wannier-interpolated band structure. The zero of energy is the Fermi level. [right] Fermi surface of lead.

8 Conclusions

In conclusion, we present a new code called wannier90 for computing maximally-localised Wannier functions. wannier90 is freely available under the GNU General Public Licence [35]. It is very user-friendly and is written in Fortran90, employing modern programming techniques that enable the addition of further functionality, such as transport properties or interpolation of electron-phonon couplings, in an easy and modular fashion. wannier90 has been seamlessly interfaced to the PWSCF plane-wave DFT code [7] and, at the time of writing, interfaces to other codes, e.g., ABINIT [36], CASTEP [37] and FLEUR [38], are in progress. We hope that the availability of wannier90 will encourage the wider use of maximally-localised Wannier functions in the electronic structure community.

Acknowledgements

We would like to thank Stefano de Gironcoli, Malgorzata Wierzbowska and Maria Peressi for the interface to the PWSCF first-principles code [7]. This work was funded by DARPA-PROM and by the Laboratory Directed Research and
Development Program of Lawrence Berkeley National Laboratory under the Department of Energy Contract No. DE-AC02-05CH11231.

A Sample files

Input file:

```plaintext
num_bands = 20
num_wann = 10
dis_win_max = 19.2
dis_froz_max = 9.8
dis_num_iter = 400
num_iter = 100

Begin Atoms_Frac
  C1  0.0000000000  0.0000000000  0.7500000000
  C1  0.3333333333  0.6666666667  0.2500000000
  C2  0.0000000000  0.0000000000  0.2500000000
  C2  -0.3333333333 -0.6666666667  0.7500000000
End Atoms_Frac

Begin Unit_Cell_Cart
  2.1304215583 -1.2299994602  0.0000000000
  0.0000000000  2.4599989204  0.0000000000
  0.0000000000  0.0000000000  6.8000000000
End Unit_Cell_Cart

Begin Projections
  C1:sp2;pz
  C2:pz
End Projections

bands_plot = true

begin kpoint_path
  G  0.0000  0.0000  0.0000  M  0.5000 -0.5000  0.0000
  M  0.5000 -0.5000  0.0000  K  0.6667 -0.3333  0.0000
  K  0.6667 -0.3333  0.0000  G  0.0000  0.0000  0.0000
  G  0.0000  0.0000  0.0000  A  0.0000  0.0000  0.5000
end kpoint_path

mp_grid = 6 6 6
```
begin kpoints
0.00000000 0.00000000 0.00000000
0.00000000 0.00000000 0.16666667
0.00000000 0.00000000 0.33333333
0.00000000 0.00000000 0.50000000
0.00000000 0.00000000 0.66666667
0.00000000 0.00000000 0.83333333
0.00000000 0.16666667 0.00000000

.
.
.

0.83333333 0.83333333 0.66666667
0.83333333 0.83333333 0.83333333
End Kpoints

Output file (truncated):

Final State

<table>
<thead>
<tr>
<th>WF centre and spread</th>
<th>(x, y, z)</th>
<th>0.59341185</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>(-0.354954, 0.614798, 5.100000)</td>
<td>0.59341185</td>
</tr>
<tr>
<td>2</td>
<td>(-0.354954, -0.614798, 5.100000)</td>
<td>0.59341185</td>
</tr>
<tr>
<td>3</td>
<td>(0.709907, 0.000000, 5.100000)</td>
<td>0.59341190</td>
</tr>
<tr>
<td>4</td>
<td>(0.000000, 0.000000, 5.100000)</td>
<td>1.04236015</td>
</tr>
<tr>
<td>5</td>
<td>(0.354954, 1.845201, 1.700000)</td>
<td>0.59341120</td>
</tr>
<tr>
<td>6</td>
<td>(0.354954, 0.614798, 1.700000)</td>
<td>0.59341085</td>
</tr>
<tr>
<td>7</td>
<td>(1.420514, 1.230000, 1.700000)</td>
<td>0.59341100</td>
</tr>
<tr>
<td>8</td>
<td>(0.710140, 1.229999, 1.700000)</td>
<td>1.04371374</td>
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<tr>
<td>9</td>
<td>(0.000000, 0.000000, 1.700000)</td>
<td>1.04233128</td>
</tr>
<tr>
<td>10</td>
<td>(-0.710140, -1.229999, 5.100000)</td>
<td>1.04368702</td>
</tr>
<tr>
<td>Sum of centres and spreads</td>
<td>(2.130421, 3.689998, 33.999999)</td>
<td>7.73256084</td>
</tr>
</tbody>
</table>

Spreads (Ang^-2)  Omega I = 6.121019675

===================
Omega D = 0.032181361
Omega OD = 1.579353494

Final Spread (Ang^-2)  Omega Total = 7.732554530

References


[38] See http://www.flapw.de/