

Density functional theory (DFT) and the concepts of the augmented-plane-wave plus local orbital (L)APW+lo method

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Walter Kelm

J.C.Slater

O.K.Andersen

Electronic structure of solids and surfaces



hexagonal boron nitride on Rh(111) 2x2 supercell (1108 atoms per cell)

Phys.Rev.Lett. 98, 106802 (2007)



K.Schwarz, P.Blaha, S.B.Trickey, Molecular physics, **108**, 3147 (2010)

Wien2k is used worldwide by about 2600 groups





1965

1975

- Walter Kohn: density functional theory (DFT),
- J.C.Slater: augmented plane wave (APW) method, 1937
- O.K.Andersen: Linearized APW (LAPW),
- Wien2k code: developed during the last 35 years
 - In the year 2000 (2k) the WIEN code (from Vienna) was called wien2k
 - One of the most accurate DFT codes for solids
 - All electron, relativistic, full-potential method
 - Widely used in academia and industry

Applications:

- <u>solids</u>: insulators , covalently bonded systems, metals
- Surfaces: catalysis
- Electronic, magnetic, elastic , optical ,...properties
- Many application in literature
- See <u>www.wien2k.at</u>



K.Schwarz, Computation of Materials Properties at the Atomic ScaleInTech, (2015) Chapter 10, 275-310,open access bookISBN 978-953-51-2126-8 (2015)dx.doi.org/10.5772/59108

K.Schwarz, P.Blaha, Electronic structure of solids and surfaces with WIEN2k in Practical Aspects of Computational Chemistry I: An Overview of the Last Two Decades and Current Trends,

J.Leszczyncski, M.K.Shukla (Eds),

Springer Science+Business Media B.V. (2012) Chapter 7, 191-207, ISBN 978-94-007-0918-8

S.Cottenier, Density Function Theory and the family of (L)APW methods : A step by step introduction 2002-2012 /(2nd edition); ISBN 978-90-807215-1-7 Freelv available at: http://www.wien2k.at/reg-user/textbooks I

K.Schwarz, P.Blaha, S.B.Trickey, *Electronic structure of solids with WIEN2k* Molecular physics, **108**, 3147 (2010)

K.Schwarz, *DFT calculations of solids with LAPW and WIEN2k* Solid State Chem.**176**, 319-328 (2003)

K.Schwarz, P.Blaha, G.K.H.Madsen, *Electronic structure calculations of solids using the WIEN2k package for material Sciences* Comp.Phys.Commun.**147**, 71-76 (2002)







Atomic structure

- Periodic boundary condition (approximation)
- Quantum mechanical treatment
 - DFT (functionals) and beyond (GW, DMFT, RPA, BSE, ...)
- How to solve the QM (basis set)
 - LAPW method and local orbitals as implemented in WIEN2k
- Applications
 - Structure, surfaces, core-level spectra, NMR, hyperfine, Wannier,...
- Software development
 - Accuracy, efficiency, system size, user-friendliness, commercial
- Insight and understanding
 - Analysis to find trends, computer experiments (artificial cases)
- Combination of expertise
 - Chemistry, physics, mathematics, computer science, application





Iarge scale applications to simulate "real" materials

- at the atomic scale
- A proper quantum mechanical treatment (accurate)
 - develop more accurate DFT functionals
 - beyond DFT (GW, DMFT, BSE, RPA...)
- Efficiency (make calculations faster):
 - improve numerics, parallelization, algorithms (iterative diagonalization)
- calculate "new" properties
 - for direct comparison with experiment





A crystal is represented by a unit cell

- We assume periodic boundary condition (approximation)
- The unit cell is repeated to infinity (makes calculations feasible)
- A real crystal is finite (with surfaces, impurities, defects ...)
- Nano materials differ from bulk
- Symmetry helps (space group, Bloch theorem, ...)

In theory

- The atomic structure is an input and thus well defined.
- Artificial structures can be studied too

In experiment

- The atomic structure is not perfectly known
- Single crystals, micro crystals, powder samples, nano
- e.g. by X-ray: averaged with uncertainties (defects, disorder)





Crystal structure

- Unit cell (defined by 3 lattice vectors) leading to 7 crystal systems
- Bravais lattice (14)
- Atomic basis (Wyckoff position)
- Symmetries (rotations, inversion, mirror planes, glide plane, screw axis)
- Space group (230)
- Wigner-Seitz cell
- Reciprocal lattice (Brillouin zone)
- Electronic structure
 - Periodic boundary conditions
 - Bloch theorem (k-vector), Bloch function
 - Schrödinger equation (HF, DFT)





Assuming an ideal infinite crystal we define a unit cell by

Unit cell: a volume in space that fills space entirely when translated by all lattice vectors.

The obvious choice:

a parallelepiped defined by **a**, **b**, **c**, three basis vectors with

the best **a**, **b**, **c** are as orthogonal as possible

the cell is as symmetric as possible (14 types)



A unit cell containing one lattice point is called primitive cell.





$$\mathbf{a} = \mathbf{b} = \mathbf{c}$$
$$\alpha = \beta = \gamma = 90^{\circ}$$

primitive





I (bcc)

body centered

face centered



F (fcc)









7 Crystal systems and 14 Bravais lattices

Triclinic	1	"no" symmetry
Monoclinic (P, C)	2	Two right angles
Orthorhombic (P, C, I, F)	4	Three right angles
Tetragonal (P, I)	2	Three right angles + 4 fold rotation
Cubic (P, I, F)	3	Three right angles + 4 fold + 3 fold
Trigonal (Rhombohedral)	1	Three equal angles (\neq 90°)+ 3 fold
Hexagonal	1	Two right and one 120° angle + 6 fold





Form connection to all neighbors and span a plane normal to the connecting line at half distance





Fraction of atoms on surface (black) or edge (red) as function of particle size n



- The electronic structure requires a QM treatment
- The main scheme is density functional theory (DFT)
 - It is a mean field approach and requires approximations
 - According to Hohenberg Kohn, it is sufficient to know the electron density of a system to determine its total energy. The many electron wave function (which depends on many variables) is not needed. In principle this is an enormous simplification, but in practice approximations must be made.
 - The direction of improving the QM treatment is summarized pictorially in Jabob's ladder:
- There are schemes which go beyond DFT:
 - GW method (for excitations or band gaps)
 - The Bethe Salpeter equation (BSE) for excitons (core hole electron)
 - Dynamical mean field theory (DMFT) based on DFT (wien2wannier)





$$\left[-\frac{1}{2}\nabla^2 + V(r)\right]\Psi(r) = E\Psi(r)$$

1-dimensioanl case:

V(x) has lattice periodicity ("translational invariance"): V(x)=V(x+a)The electron density $\rho(x)$ has also lattice periodicity, however, the wave function does NOT:

$$\rho(x) = \rho(x+a) = \Psi^*(x)\Psi(x) \qquad but:$$

$$\Psi(x+a) = \mu\Psi(x) \implies \mu^*\mu = 1$$

Application of the translation τ g-times:

$$\tau^g \Psi(x) = \Psi(x + ga) = \mu^g \Psi(x)$$



The wave function must be uniquely defined: after G translations it must be identical (G a: periodicity volume):

$$\tau^{G}\Psi(x) = \Psi(x + Ga) = \mu^{G}\Psi(x) = \Psi(x)$$

$$\Rightarrow \mu^{G} = 1$$

$$\mu = e^{2\pi i \frac{g}{G}}$$

$$g = 0, \pm 1 \pm 2, \dots$$

$$Def.: \qquad k = \frac{2\pi}{a} \frac{g}{G}$$

$$\mu = e^{ika}$$

Bloch condition: $\Psi(x+a) = e^{ika} \Psi(x) = \Psi_k$





Wave functions with Bloch form:



Replacing k by k+K, where K is a reciprocal lattice vector, fulfills again the Bloch-condition. $\rightarrow k$ can be restricted to the first Brillouin zone.

$$e^{i\frac{2\pi}{a}K} = 1 \qquad \qquad -\frac{\pi}{a} < k < \frac{\pi}{a}$$



Concepts when solving Schrödingers-equation in solids









Two communities in solid state theory

	LDA bandstructure	many body theory
+	 material-specific, "ab initio" often successful, quantitative 	 electronic correlations qualitative understanding
(1)))	 effective one-particle approach 	 model Hamiltonian





Ab-initio Hamiltonian

(non-relativistic/Born-Oppenheimer approximation)





LDA bandstructure corresponds to





Coulomb potential:

- nuclei
- all electrons
- including self-interaction

Quantum mechanics:

- exchange
- correlation
- (partly) cancel self-interaction

Jacob's ladder: The grand challenge:

- Find a functional which leads to "DFT heaven": predictive power with chemical accuracy
- J. Perdew's "Jacob's ladder" into DFT heaven:



- unoccupied orbitals (ACFDT-RPA)
- occupied orbitals (hybrid-DFT)
- meta-GGAs (kinetic energy density τ)
- GGA ($\nabla \rho$)

accuracy (computer time)

- LDA (ρ)
- Hartree













Hohenberg-Kohn theorem: (exact)

The total energy of an interacting inhomogeneous electron gas in the presence of an external potential $V_{ext}(r)$ is a functional of the density ρ

$$E = \int V_{ext}(\vec{r})\rho(\vec{r})d\vec{r} + F[\rho]$$

Kohn-Sham: (still exact!)

$$E = T_o[\rho] + \int V_{ext} \rho(\vec{r}) d\vec{r} + \frac{1}{2} \int \frac{\rho(\vec{r})\rho(\vec{r}')}{|\vec{r}' - \vec{r}|} d\vec{r} d\vec{r}' + E_{xc}[\rho]$$



In KS the many body problem of interacting electrons and nuclei is mapped to a one-electron reference system that leads to the same density as the real system.



- We divide the density of the N-1 electron system into the total density n(r) and an exchangecorrelation hole:
- Properties of the exchange-correlation hole:
- Locality
- Pauli principle
- the hole contains ONE electron
- The hole must be negative
- The exchange hole affects electrons with the same spin and accounts for the Pauli principle
- In contrast, the correlation-hole accounts for the Coulomb repulsion of electrons with the opposite spin. It is short range and leads to a small redistribution of charge. The correlation hole contains NO charge:



 $\overline{n}(r_0,r) = n(r) + h(r_0,r)$



$$dr h_c(r_0, r) = 0$$

Kohn-Sham equations

$$LDA, GGA$$

$$E = T_o[\rho] + \int V_{ext}\rho(\vec{r})d\vec{r} + \frac{1}{2}\int \frac{\rho(\vec{r})\rho(\vec{r}')}{|\vec{r}' - \vec{r}|}d\vec{r}d\vec{r}' + E_{xc}[\rho]$$
1-electron equations (Kohn Sham)

$$\{-\frac{1}{2}\nabla^2 + V_{ext}(\vec{r}) + V_c(\rho(\vec{r})) + V_{xc}(\rho(\vec{r}))\}\Phi_i(\vec{r}) = \varepsilon_i\Phi_i(\vec{r})$$

$$\frac{-1}{2}\nabla^2 + V_{ext}(\vec{r}) + V_c(\rho(\vec{r})) + V_{xc}(\rho(\vec{r}))\}\Phi_i(\vec{r}) = \varepsilon_i\Phi_i(\vec{r})$$

$$\frac{-1}{2}\nabla^2 + V_{ext}(\vec{r}) + V_c(\rho(\vec{r})) + V_{xc}(\rho(\vec{r}))\}\Phi_i(\vec{r}) = \varepsilon_i\Phi_i(\vec{r})$$

$$\frac{-1}{2}\nabla^2 + V_{ext}(\vec{r}) + V_c(\rho(\vec{r})) + V_{xc}(\rho(\vec{r}))\}\Phi_i(\vec{r}) = \varepsilon_i\Phi_i(\vec{r})$$

$$\frac{-1}{2}\nabla^2 + V_{ext}(\vec{r}) + V_c(\rho(\vec{r})) + V_{xc}(\rho(\vec{r}))\}\Phi_i(\vec{r}) = \varepsilon_i\Phi_i|\rho(\vec{r})|$$

$$\frac{-1}{2}\nabla^2 + V_{ext}(\vec{r}) + V_c(\rho(\vec{r})) + V_{xc}(\rho(\vec{r}))\}\Phi_i(\vec{r}) = \varepsilon_i\Phi_i|\rho(\vec{r})|$$

$$\frac{-1}{2}\nabla^2 + V_{ext}(\vec{r}) + V_c(\rho(\vec{r})) + V_{xc}(\rho(\vec{r})) + V_{xc}(\rho(\vec{r}))$$

$$\frac{-1}{2}\nabla^2 + V_{ext}(\vec{r}) + V_c(\rho(\vec{r})) + V_{xc}(\rho(\vec{r})) + V_{xc$$

New (better ?) functionals are still an active field of research





- Every observable quantity of a quantum system can be calculated from the density of the system ALONE (Hohenberg, Kohn, 1964).
- The density of particles interacting with each other can be calculated as the density of an auxiliary system of non-interacting particles (Kohn, Sham, 1965).







With 80 years



- 1923 born in Vienna
- 1938 had to leave Vienna
- 1946 Univ. Toronto (master, Math)
- 1948 Harvard (PhD, Schwinger)
- Carnegie Mellon, Pittsburgh (Luttinger), Bell Labs, Copenhagen, Washigton (Seattle), Paris, Imperial College (London), ETH Zürich, Unv. California, San Diego
- 1979 Founding director Santa Barbara, California
- 1964 Hohenberg Kohn
- 1965 Kohn Sham
- 1998 Nobel prize: Chemistry
- 2016, he died on April 19



Walter Kohn, Nobel Prize 1998 Chemistry







"Self-consistent Equations including Exchange and Correlation Effects" W. Kohn and L. J. Sham, Phys. Rev. 140, A1133 (1965) 50 years ago

Literal quote from Kohn and Sham's paper:"... We do not expect an accurate description of chemical binding."

DFT ground state of iron









GGA follows LDA







Single-electron picture

Koopmans theorem: $P = \Delta E \approx \varepsilon_i$



Slater's transion state: Cl atom



K.Schwarz Chem. Physics <u>7</u>, 100 (1975)



CoO AFM-II total energy, DOS





CoO

- in NaCl structure
- antiferromagnetic: AF II
- insulator
- *t_{2g} splits into a_{1g} and e_g*
- GGA almost spilts the bands









FeF₂: GGA works surprisingly well









Lattice parameters (Å)							
	Exp.	LDA	PBE	WC			
Со	2.51	2.42	2.49	2.45			
Ni	3.52	3.42	3.52	3.47			
Cu	3.61	3.52	3.63	3.57			
Ru	2.71	2.69	2.71	2.73			
Rh	3.80	3.76	3.83	3.80			
Pd	3.88	3.85	3.95	3.89			
Ag	4.07	4.01	4.15	4.07			
Ir	3.84	3.84	3.90	3.86			
Pt	3.92	3.92	4.00	3.96			
Au	4.08	4.07	4.18	4.11			

3d elements:

 PBE superior, LDA much too small

4d elements:

LDA too small, PBE too large *New functional Wu-Cohen (WC)*

Z.Wu, R.E.Cohen, PRB 73, 235116 (2006)

5d elements:

 LDA superior, PBE too large



accuracy: "DFT limit"



BN Sn Testing of DFT functionals: error of theoretical lattice Pb Aa **LDA** GaAs Ge parameters for a large GaAs LiCI InF AlAs GaAs LiC **AM05** Ŧ MaO Pd variety of solids (Li-Th) Au CeC Rb AIAs GaP AIA CeO Pt SiC CeO_{2 Ir} CeO CeO. CeO CeO AIP Gal GaP Si CaC Pd NbN WC С Aq PBEso HfN SIC Cu me mae mre mare (Å) (Å) (%) (%) TiC Та ZrC ZrC ZrC Та 1.32 LDA -0.058 0.058 -1.32 Nb Gab d d Aa NhC 7rN CaO PBE CaO TIC ZrN ZrN NaC TiN 0.68 SO-GGA -0.014 0.029 -0.37 NaE ZrC TIN HfC. TIN NiAl Та Ba TIN LiCI VN 0.67 **PBEsol** -0.005 0.029 -0.17 TiC Та CoA NiA VC Ra -0.03 0.68 WC 0.000 0.031 Cu +-- LDA Sr ----- SOGGA — PBEsol AM05 0.005 0.035 0.01 0.77 A── WC AM05 TPSS 0.055 1.05 1.18 PBE 0.051 - PBE $^{-1.5}_{100(a_0^{calc} - a_0^{expt})/a_0^{expt})}$ -3.5-3 -2.5 -2 0.5 1 1.5 2 2.5 З 3.5





• better GGAs and meta-GGAs (ρ , $\nabla \rho$, τ):

- usually improvement, but often too small.
- LDA+U: for correlated 3d/4f electrons, treat strong Coulomb repulsion via Hubbard U parameter (cheap, "empirical U" ?)
- Exact exchange: imbalance between exact X and approximate C
 - hybrid-DFT (mixing of HF + GGA; "mixing factor"?)
 - exact exchange + RPA correlation (extremely expensive)
- **GW:** gaps in semiconductors, expensive!
- Quantum Monte-Carlo: very expensive
- DMFT: for strongly correlated (metallic) d (f) -systems (expensive)


Approximations for
$$E_{\rm xc}$$

► LDA:
$$E_{\rm xc}^{\rm LDA} = \int f(\rho(\mathbf{r})) d^3 r$$

• GGA:
$$E_{\rm xc}^{\rm GGA} = \int f(\rho(\mathbf{r}), |\nabla \rho(\mathbf{r})|) d^3r$$



• MGGA: $E_{\rm xc}^{\rm MGGA} = \int f(\rho(\mathbf{r}), |\nabla \rho(\mathbf{r})|, \nabla^2 \rho(\mathbf{r}), t(\mathbf{r})) d^3 r$

$$\blacktriangleright \text{ LDA+}U: E_{\text{xc}}^{\text{LDA+}U} = E_{\text{xc}}^{\text{LDA}} + E_{\text{ee}} - E_{\text{dc}}$$

• GGA+U:
$$E_{\rm xc}^{\rm GGA+U} = E_{\rm xc}^{\rm GGA} + E_{\rm ee} - E_{\rm dc}$$

• hybrid:
$$E_{xc}^{hybrid} = E_{xc}^{DFT} + \alpha (E_x^{HF} - E_x^{DFT})$$

where

$$E_{\mathbf{x}}^{\mathrm{HF}} = -\frac{1}{2} \sum_{\sigma} \sum_{\substack{n,\mathbf{k}\\n',\mathbf{k'}}} w_{\mathbf{k}} w_{\mathbf{k}'} \int \int \frac{\psi_{n\mathbf{k}}^{\sigma*}(\mathbf{r})\psi_{n'\mathbf{k}'}^{\sigma*}(\mathbf{r}')\psi_{n\mathbf{k}'}^{\sigma}(\mathbf{r})\psi_{n\mathbf{k}}^{\sigma}(\mathbf{r}')}{|\mathbf{r}-\mathbf{r}'|} d^{3}r d^{3}r'$$







Only for certain atoms and electrons of a given angular momentum *l*

$$E_{\mathrm{xc}}^{\mathrm{hybrid}} = E_{\mathrm{xc}}^{\mathrm{DFT}}[\rho^{\sigma}] + \alpha \left(E_{\mathrm{x}}^{\mathrm{HF}}[n_{m_{i}m_{j}}^{\sigma}] - E_{\mathrm{x}}^{\mathrm{DFT}}[\rho_{\ell}^{\sigma}] \right)$$

$$E_{\mathrm{x}}^{\mathrm{HF}}[n_{m_{i}m_{j}}^{\sigma}] = -\frac{1}{2} \sum_{\sigma} \sum_{m_{1},m_{2},m_{3},m_{4}}^{\ell} n_{m_{1}m_{2}}^{\sigma} n_{m_{3}m_{4}}^{\sigma} \langle m_{1}m_{3} | v_{\mathrm{ee}} | m_{4}m_{2} \rangle$$

$$\langle m_{1}m_{2} | v_{\mathrm{ee}} | m_{3}m_{4} \rangle = \sum_{k=0}^{2\ell} a_{k} F_{k}$$

The Slater integrals F_k are calculated according to P.Novák et al., phys.stat.sol (b) **<u>245</u>**, 563 (2006)





 $\left| -\frac{1}{2} \nabla^2 + V(r) \right| \Psi_i^k = \varepsilon_i^k \Psi_i^k$

- Ψ cannot be found analytically
- complete "numerical" solution is possible but inefficient Ansatz: $\Psi_k = \sum c_{k_n} \Phi_{k_n}$
 - Inear combination of some "basis functions"
 - different methods use different basis sets !
 - finding the "best" wave function using the variational principle:

$$\langle E_k \rangle = \frac{\left\langle \Psi_k^* \middle| H \middle| \Psi_k \right\rangle}{\left\langle \Psi_k^* \middle| \Psi_k \right\rangle} \qquad \frac{\partial E_k}{\partial c_{k_n}} = 0$$

• this leads to the famous "Secular equations", i.e. a set of linear equations which in matrix representation is called "generalized" eigenvalue problem"

HC = ESC

H, S : hamilton and overlap matrix; C: eigenvectors, E: eigenvalues



plane waves

- pseudo potentials
- PAW (projector augmented wave) by P.E.Blöchl
- space partitioning (augmentation) methods
 - LMTO (linear muffin tin orbitals)
 - ASA approx., linearized numerical radial function
 - + Hankel- and Bessel function expansions
 - full-potential LMTO
 - ASW (augmented spherical wave)
 - similar to LMTO
 - KKR (Korringa, Kohn, Rostocker method)
 - solution of multiple scattering problem, Greens function formalism
 - equivalent to APW
 - (L)APW (linearized augmented plane waves)

LCAO methods

Gaussians, Slater, or numerical orbitals, often with PP option)









pseudopotential plane wave methods



• plane waves form a "complete" basis set, however, they "never" converge due to the rapid oscillations of the atomic wave functions χ close to the nuclei

 let's get rid of all core electrons and these oscillations by replacing the strong ion—electron potential by a much weaker (and physically dubious) pseudopotential

 Hellmann's 1935 combined approximation method







- "real" potentials contain the Coulomb singularity -Z/r
- the wave function has a cusp and many wiggles,
- chemical bonding depends mainly on the overlap of the wave functions between neighboring atoms (in the region between the nuclei) →







APW (J.C.Slater 1937)

- Non-linear eigenvalue problem
- Computationally very demanding
- LAPW (O.K.Anderssen 1975)
 - Generalized eigenvalue problem
 - Full-potential
- Local orbitals (D.J.Singh 1991)
 - treatment of semi-core states (avoids ghostbands)
- APW+lo (E.Sjöstedt, L.Nordstörm, D.J.Singh 2000)
 - Efficiency of APW + convenience of LAPW
 - Basis for



K.Schwarz, P.Blaha, G.K.H.Madsen, Comp.Phys.Commun.**147**, 71-76 (2002)

K.Schwarz, DFT calculations of solids with LAPW and WIEN2k Solid State Chem.**176**, 319-328 (2003)

K.Schwarz, P.Blaha, S.B.Trickey, Molecular physics, **108**, 3147 (2010)



APW Augmented Plane Wave method





 Dasis set.

 PW:
 $e^{i(\vec{k}+\vec{K}).\vec{r}}$

 Atomic partial waves
 join

 $\sum_{\ell m} A_{\ell m}^{\kappa} u_{\ell}(r',\varepsilon) Y_{\ell m}(\hat{r}')$

 $u_{l}(r,\varepsilon)$ are the numerical solutions of the radial Schrödinger equation in a given spherical potential for a particular energy ε A_{lm}^{K} coefficients for matching the PW



Numerical search for those energies, for which the det[H-ES] vanishes. Computationally very demanding. "Exact" solution for given MT potential!

Linearization of energy dependence



expand u_l at fixed energy E_l and add $\dot{u}_l = \partial u_l / \partial \mathcal{E}$

A_{Im}^k, *B_{Im}^k*: join PWs in value and slope

→ General eigenvalue problem (diagonalization)

→ additional constraint requires more PWs than APW





Atomic sphere approximation (ASA)

- overlapping spheres "fill" all volume
- potential spherically symmetric

- "muffin-tin" approximation (MTA)
 - non-overlapping spheres with spherically symmetric potential +
 - interstitial region with V=const.





"full"-potential

no shape approximations to V







SrTiO₃

Full potential

Muffin tin approximation



TiO₂ rutile

 The potential (and charge density) can be of general form (no shape approximation)

$$V(r) = \begin{cases} \sum_{LM} V_{LM}(r) Y_{LM}(\hat{r}) & r < R_{\alpha} \\ \sum_{K} V_{K} e^{i\vec{K}.\vec{r}} & r \in I \end{cases}$$

 Inside each atomic sphere a local coordinate system is used (defining LM)









For example: Ti



- Valences states
 - High in energy
 - Delocalized wavefunctions
- Semi-core states
 - Medium energy
 - Principal QN one less than valence (e.g. in Ti 3p and 4p)
 - not completely confined inside sphere (charge leakage)
- Core states
 - Low in energy
 - Reside inside sphere

1 Ry =13.605 eV



Local orbitals (LO)







 $\Phi_{IO} = [A_{\ell m} u_{\ell}^{E_{1}} + B_{\ell m} \dot{u}_{\ell}^{E_{1}} + C_{\ell m} u_{\ell}^{E_{2}}]Y_{\ell m}(\hat{r})$

LOs

- are confined to an atomic sphere
- have zero value and slope at R
- Can treat two principal QN n for each azimuthal QN l
 (e.g. 3p and 4p)
- Corresponding states are strictly orthogonal
 - (e.g.semi-core and valence)
- Tail of semi-core states can be represented by plane waves
- Only slightly increases the basis set (matrix size)

D.J.Singh, Phys.Rev. B 43 6388 (1991)





E.Sjöstedt, L.Nordström, D.J.Singh, *An alternative way of linearizing the augmented plane wave method,* Solid State Commun. 114, 15 (2000)

- Use APW, but at fixed E_{I} (superior PW convergence)
- Linearize with additional local orbitals (lo) (add a few extra basis functions)

$$\Phi_{k_n} = \sum_{\ell m} A_{\ell m}(k_n) u_{\ell}(E_{\ell}, r) Y_{\ell m}(\hat{r})$$

$$\Phi_{lo} = [A_{\ell m} u_{\ell}^{E_1} + B_{\ell m} \dot{u}_{\ell}^{E_1}] Y_{\ell m}(\hat{r})$$

optimal solution: mixed basis

- use APW+lo for states, which are difficult to converge: (f or d- states, atoms with small spheres)
- use LAPW+LO for all other atoms and angular momenta







SES (sodium electro solodalite)

K.Schwarz, P.Blaha, G.K.H.Madsen, Comp.Phys.Commun.**147**, 71-76 (2002)

Atomic partial waves

LAPW

$$\Phi_{k_n} = \sum_{\ell m} [A_{\ell m}(k_n) u_{\ell}(E_{\ell}, r) + B_{\ell m}(k_n) \dot{u}_{\ell}(E_{\ell}, r)] Y_{\ell m}(\hat{r})$$

APW+lo

$$\Phi_{k_n} = \sum_{\ell m} A_{\ell m}(k_n) u_{\ell}(E_{\ell}, r) Y_{\ell m}(\hat{r})$$

Plane Waves (PWs)

$$e^{i(\vec{k}+\vec{K}_n).\vec{r}}$$

- match at sphere boundary
 - LAPW
 - value and slope $A_{\ell m}(k_n), B_{\ell m}(k_n)$

 $A_{\ell m}(k_n)$

APW

value

plus another type of local orbital (lo)







E.Sjöststedt, L.Nordström, D.J.Singh, SSC 114, 15 (2000)

- Use APW, but at fixed E_{I} (superior PW convergence)
- Linearize with additional lo (add a few basis functions)

optimal solution: mixed basis

- use APW+lo for states which are difficult to converge: (f- or d- states, atoms with small spheres)
- use LAPW+LO for all other atoms and angular momenta

A summary is given in

K.Schwarz, P.Blaha, G.K.H.Madsen, Comp.Phys.Commun.**147**, 71-76 (2002)



The WIEN2k authors





An Augmented Plane Wave Plus Local Orbital Program for Calculating Crystal Properties

> Peter Blaha Karlheinz Schwarz Georg Madsen Dieter Kvasnicka Joachim Luitz

November 2001 Vienna, AUSTRIA Vienna University of Technology



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Europe: A, B, CH, CZ, D, DK, ES, F, FIN, GR, H, I, IL, IRE, N, NL, PL, RO, S, SK, SL, SI, UK (ETH Zürich, MPI Stuttgart, FHI Berlin, DESY, RWTH Aachen, ESRF, Prague, IJS Ljubjlana, Paris, Chalmers, Cambridge, Oxford)

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<u>far east:</u> AUS, China, India, JPN, Korea, Pakistan, Singapore,Taiwan (Beijing, Tokyo, Osaka, Kyoto, Sendai, Tsukuba, Hong Kong)



100 industries (Canon, Eastman, Exxon, Fuji, Hitachi, IBM, Idemitsu Petrochem., Kansai, Komatsu, Konica-Minolta, A.D.Little, Mitsubishi, Mitsui Mining, Motorola, NEC, Nippon Steel, Norsk Hydro, Osram, Panasonic, Samsung, Seiko Epson, Siemens, Sony, Sumitomo,TDK,Toyota).

mailinglist: 10.000 emails/6 years



The first publication of the WIEN code

FULL-POTENTIAL, LINEARIZED AUGMENTED PLANE WAVE PROGRAMS... FOR CRYSTALLINE SYSTEMS

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Institut für Technische Elektrochemie, Technische Universität Wien, A-1060 WIEN, Austria

and

S.B. TRICKEY

Quantum Theory Project, Depts. of Physics and of Chemistry, University of Florida, Gainesville, FL 32611, USA

PROGRAM SUMMARY

Title of program: WIEN

Computer Physics Communications 59 (1990) 399-415









In the Heart of EUROPE







 Book published by Shinya Wakoh (2006)

和光システム研究所 著

WIEN2k 入門

WIEN-code は 1980 年ごろから, グループの指導者である Karlheinz Schwarz によって書き 始められ, 1990 年に最初の copyrighted version の WIEN が発表された. その後 UNIX version となり, WIEN93, WIEN95, WIEN97 を経て, Fortran90 対応の WIEN2k へと改良・拡張 されてきた^{*1}. 基礎となるシュレーディンガー方程式はコーン・シャム方程式であり, バンド計算 法は主として FLAPW 法, ポテンシャルは LSDA, GGA などである. 最新の WIEN2k では, APW+lo も取り入れられており, ポテンシャルとしては電子相関が強いときに必要であると云わ れている補正 +U も扱えるようになっている. また, 並列計算機を使えば, 極めて複雑な結晶も 計算の対象とすることができる.





Authors of WIEN2k

P. Blaha, K. Schwarz, D. Kvasnicka, G. Madsen and J. Luitz

Other contributions to WIEN2k

- C. Ambrosch-Draxl (Free Univ. Berlin) optics
- E. Assmann (Vienna) Wannier functions
- F. Karsai (Vienna) parallelization
- R. Laskowski (Singapore), non-collinear magnetism, NMR chemical shifts, BSE
- L. Marks (Northwestern, US) , various optimizations, new mixer
- P. Novák and J. Kunes (Prague), LDA+U, SO
- B. Olejnik (Vienna), non-linear optics,
- C. Persson (Uppsala), irreducible representations
- V. Petricek (Prague) 230 space groups
- O. Rubel (McMaster Univ. Hamiton, ON) Berry phases
- M. Scheffler (Fritz Haber Inst., Berlin), forces
- D.J.Singh (NRL, Washington D.C., Oak Ridge), local oribtals (LO), APW+lo
- E. Sjöstedt and L Nordström (Uppsala, Sweden), APW+lo
- J. Sofo (Penn State, USA) and J. Fuhr (Barriloche), Bader analysis
- F. Tran (Vienna) Hartree Fock, DFT functionals
- B. Yanchitsky and A. Timoshevskii (Kiev), space group
- and many others



A series of **WIEN workshops** were held



 1st 	Vienna	April	1995	Wien95	
2nd	Vienna	April	1996		
3rd	Vienna	April	1997	Wien97	
 4st 	Trieste, Italy	June	1998		
5st	Vienna	April	1999		
6th	Vienna	April	2000		
7th	Vienna	Sept.	2001	Wien2k	
8th	Esfahan, Iran	April	2002		
	Penn State, USA	July	2002		
9th	Vienna	April	2003		
10th	Penn State, USA	July	2004		
11th	Kyoto, Japan	May	2005		
	IPAM, Los Angeles, USA	Nov.	2005		
12th	Vienna	April	2006		
13th	Penn State, USA	June	2007	2800	
14th	Singapore	July	2007		NOL
15th	Vienna	March	2008	2200 -	INZK
16th	Penn State, USA	June	2009	2000 -	
17th	Nantes, France	July	2010	g 1600 -	
18th	Penn State, USA	June	2011	2 1400 - 1200 -	
19th	Tokyo, Japan	Sept	2012	1000 -	/
20th	Penn State, USA	Aug.	2013	800 -	
21th	Nantes, France	July	2014	400 -	
	Warsaw, Poland	Oct.	2014	200 - 0 -	
22nd	Singapore	Aug.	2015	2002 2004 2	2006
23rd	McMaster, Canada	June	2016		









APW + local orbital method (linearized) augmented plane wave method

Total wave function

$$\Psi_k = \sum_{K_n} C_{k_n} \phi_{k_n} \quad \text{n...50-100 PWs /atom}$$

Variational method:

$$< E >= \frac{< \Psi / H / \Psi >}{< \Psi / \Psi >} \quad \frac{\delta < E >}{\delta C_{k_n}} = 0$$

upper bound minimum

Generalized eigenvalue problem: H C = E S C

Diagonalization of (real or complex) matrices of size 10.000 to 50.000 (up to 50 Gb memory)







Irreducible BZ (IBZ)

- The irreducible wedge
- Region, from which the whole BZ can be obtained by applying all symmetry operations
- Bilbao Crystallographic Server:
 - <u>www.cryst.ehu.es/cryst/</u>
 - The IBZ of all space groups can be obtained from this server
 - using the option KVEC and specifying the space group (e.g. No.225 for the fcc structure leading to bcc in reciprocal space, No.229)







- In order to solve $H\Psi = E\Psi$ we need to know the potential V(r)
- for V(r) we need the electron density ρ(r)
- the density $\rho(r)$ can be obtained from $\Psi(r)^*\Psi(r)$
- ?? $\Psi(\mathbf{r})$ is unknown before $H\Psi = E\Psi$ is solved ??







Band structure of fcc Cu





Program structure of WIEN2k



init_lapw

- initialization
- symmetry detection (F, I, Ccentering, inversion)
- input generation with recommended defaults
- quality (and computing time) depends on k-mesh and R.Kmax (determines #PW)

run_lapw

- scf-cycle
- optional with SO and/or LDA+U
- different convergence criteria (energy, charge, forces)
- save_lapw tic_gga_100k_rk7_vol0
 - cp case.struct and clmsum files,
 - mv case.scf file
 - rm case.broyd* files









WIEN2k: P. Blaha, K. Schwarz, G. Madsen, D. Kvasnicka, and J. Luitz



- individual FORTRAN programs linked by shell-scripts
- the output of one program is input for the next
- lapw1/2 can run in parallel on many processors







- + robust all-electron full-potential method (new effective mixer)
- + unbiased basisset, one convergence parameter (LDA-limit)
- + all elements of periodic table (comparable in CPU time), metals
- + LDA, GGA, meta-GGA, LDA+U, spin-orbit
- + many properties and tools (supercells, symmetry)
- + w2web (for novice users)
- ? speed + memory requirements
 - + very efficient basis for large spheres (2 bohr) (Fe: 12Ry, O: 9Ry)
 - less efficient for small spheres (1 bohr) (O: 25 Ry)
 - large cells, many atoms (n³, but new iterative diagonalization)
 - full H, S matrix stored \rightarrow large memory required
 - + effective dual parallelization (k-points, mpi-fine-grain)
 - + many k-points do not require more memory
- no stress tensor
- no linear response



w2web GUI (graphical user interface)



Structure generator

- spacegroup selection
- import cif file
- step by step initialization
 - symmetry detection
 - *automatic input generation*
- SCF calculations
 - Magnetism (spin-polarization)
 - Spin-orbit coupling
 - Forces (automatic geometry optimization)
- Guided Tasks
 - Energy band structure
 - DOS
 - Electron density
 - X-ray spectra
 - Optics



Execution >> StructGen[™] initialize calc. run SCF single prog. optimize(V,c/a) mini. positions

Tasks >>

Files >> struct file(s) input files output files SCF files

Session Mgmt. >> change session change dir change info

Configuration

Usersguide html-Version pdf-Version

Idea and realization

Session: TiC

/area51/pblaha/lapw/2005-june/TiC

StructGen™

You have to click "Save Structure" for changes to take effect!				
Save Structure				
Title: TiC				
Type: F				
P ^				
F				
B Spacegroups from CXY Difference Compt Semicro				
Blibao Cryst Server				
CXZ				
n H				
1_P1 <u> </u>				
Lattice parameters in 🔺 👱				
a=4.3280000386 b=4.3280000386 c=4.3280000386				
$\alpha = 90.000000$ $\beta = 90.000000$ $\gamma = 90.000000$				
Inequivalent Atoms: 2				
Atom 1 [•] Ti 7=22.0 RMT=2.0000 remove ato				
Pos 1: x=0.00000000 y=0.00000000 z=0.00000000 remove				
add position				
Atom 2: C Z=6.0 RMT=1.9000 remove ato				
add position				


 $P4_2/mnm$ D^{14}_{4h}

No. 136

 $P 4_2/m 2_1/n 2/m$

Structure given by:

spacegroup lattice parameter positions of atoms (basis)

Rutile TiO₂: P4₂/mnm (136) a=8.68, c=5.59 bohr Ti: (0,0,0) 2a

O: (0.304,0.304,0) Wyckoff position: x, x, 0













- NaCl structure (100) plane
- Valence electrons only
- plot in 2 dimensions
- Shows
 - charge distribution
 - covalent bonding
 - between the Ti-3d and C-2p electrons
 - e_g/t_{2g} symmetry





Select plot type: 3D-plot 🛞	Contur-plot 🔾	with labels 🔄	
Min -0.5 Max 2 C	Delta 0.1		
plot electron density			
edit TiC.in2 reset EMIN			







BALSAC

+

pd,

636mRyd

 Δ_{5}

(100) plane







TiC, bonding and antibonding states









antibonding C_p -Ti_d σ

bonding C_p -Ti_d σ







Atomic form factors for Ti and C



(b)





Reproducibility in density functional theory calculations of solids Kurt Lejaeghere *et al. Science* **351**, (2016); DOI: 10.1126/science.aad3000



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Reproducibility in density functional theory calculations of solids Kurt Lejaeghere *et al. Science* **351**, (2016); DOI: 10.1126/science.aad3000



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		l i	i ₁ AE						
_		<∆> ерезече	Elk	exciting	FHI-aims/tie	FLEUR	s+i+l/olfi	RSPt	WIEN2k/acc
-	Elk	0.6		0.3	0.3	0.6	1.0	0.9	0.3
	exciting	0.5	0.3		0.1	0.5	0.9	0.8	0.2
	FHI-aims/tier2	0.5	0.3	0.1		0.5	0.9	0.8	0.2
A	FLEUR	0.6	0.6	0.5	0.5		0.8	0.6	0.4
	FPLO/T+F+s	0.9	1.0	0.9	0.9	0.8		0.9	0.9
	RSPt	0.8	0.9	0.8	0.8	0.6	0.9		0.8
_	WIEN2k/acc	0.5	0.3	0.2	0.2	0.4	0.9	0.8	
PAW	GBRV12/ABINIT	0.9	0.9	0.8	0.8	0.9	1.3	11	0.8
	GPAN09/ABINIT	1.4	1.3	13	13	1.3	1.7	15	13
	GPAN09/GPAN	1.6	1.5	15	15	1.5	1.8	17	15
	JTH02/ABINIT	0.6	0.6	0.6	0.6	0.6	0.9	0.7	0.5
	PS11b100/QE	0.9	0.9	0.8	0.8	0.8	1.3	11	0.8
-	VASPGW2015/VASP	0.6	0.4	0.4	0.4	0.6	1.0	0.8	0.3
	GBRV14/CASTEP	1.1	1.1	11	10	1.0	1.4	13	10
L L L	GBRV14/QE	1.1	1.0	10	0.9	1.0	1.4	13	10
S	OTFG9/CASTEP	0.7	0.4	0.5	0.5	0.7	1.0	10	0.5
	SSSP/QE	0.5	0.4	0.3	0.3	0.5	0.9	0.8	0.3
	Vdb2/DACAPO	6.3	6.3	6.3	6.3	6.3	6.4	6.5	6.2
	FHI98pp/ABINIT	13.3	13.5	13.4	13.4	13.2	13.0	13.2	13.4
	HGH/ABINIT	2.2	2.2	22	22	2.0	2.3	22	21
d d	HGH-NLCC/BigDFT	1.1	1.1	11	11	1.0	1.2	11	10
С С	MBK2013/OpenMX	2.0	2.1	21	21	1.9	1.8	18	20
Z ONCVPSP (PD0.1) / ABINIT		0.7	0.7	0.7	0.7	0.6	1.0	0.8	0.6
	oncvpsp (SG15) 1/QE	1.4	1.4	13	13	1.3	1.6	15	13
	oncvpsp (SG15) 2/Castep	1.4	1.4	14	14	1.3	1.6	15	14

The most accurate code

AE: all electron

Fig. 4. Δ values for comparisons between the most important DFT methods considered (in millielectron volts per atom). Shown are comparisons of all-electron (AE), PAW, ultrasoft (USPP), and norm-conserving pseudopotential (NCPP) results with all-electron results (methods are listed in alphabetical order in each category). The labels for each method stand for code, code/specification (AE), or potential set/code (PAW, USPP, and NCPP) and are explained in full in tables S3 to S42. The color coding illustrates the range from small (green) to large (red) Δ values. The mixed potential set SSSP was added to the ultrasoft category, in agreement with its prevalent potential type. Both the code settings and the DFT-predicted EOS parameters behind these numbers are included in tables S3 to S42, and fig. S1 provides a full Δ matrix for all methods mentioned in this article.



Precision of DFT calculations illustrated for Si





Fig. 1. Historical evolution of the predicted equilibrium lattice parameter for silicon. All data points represent calculations within the DFT-PBE framework. Values from literature (data points before 2016) (*15, 16, 18, 56–62, 63–65*) are compared with (i) predictions from the different codes used in this study (2016 data points, magnified in the inset; open circles indicate data produced by older methods or calculations with lower numerical settings) and (ii) the experimental value, extrapolated to 0 K and corrected for zero-point effects (red line) (*26*). The concepts of precision and accuracy are illustrated graphically.





- "real" potentials contain the Coulomb singularity -Z/r
- the wave function has a cusp and many wiggles,
- chemical bonding depends mainly on the overlap of the wave functions between neighboring atoms (in the region between the nuclei) →







Theory vs. experiment: Agreement or disagreement: What can cause it?

Structure model:	Quantum mechanics:	Convergence:	Other effects:
unit cell supercell surface	mean field (DFT) many body theory ground vs. excited states	basis sets k -points	temperature T>0 K pressure
stoichiometry disorder impurities, defects	electron core-hole satellites all electron relativistic effects	vacuum supercell { quantum n.	average vibrations

These aspects need to be considered when comparing **theory** with **experiment**.



Vienna, city of music and the Wien2k code









1997 DFT Conference in Vienna *Walter Kohn Main speaker*

1998 Nobel Prize for Chemmistry

2001 invited lecturer

My 60th birthday



Kungliga Svenska Vetenskapsakademien har den 13 oktober 1998 beslutat att med det **NOBELPRIS** som detta år tillerkännes den som gjort den viktigaste-kemiska upptäckten eller förbättringen mad ena hälften belöna **Walter Kohn** för hans utveckling av täthetsfunktionalteorin . • stockholm den in december 1998



International Conference on Applied Density Functional Theory January 14-17, 2001 Vienna/Austria

dedicated to Karlheinz Schwarz on occasion of his 60th birthday Honorary Chairman: Walter Kohn

16.1. 2001

Walter Kohn Reter & Carlande Sky licky



Walter Kohn's 80 Ecole Normale Supérieur





World of Interacting Electrons and Nuclei **2** due walter <a>kohn



2k