

Further Applications

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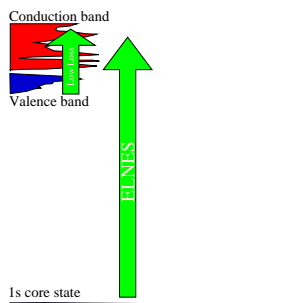
- Lecture IV: Further Applications
 - Core level and optical spectroscopies
 - Systematic prediction of new materials



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SPECTRAL PROPERTIES

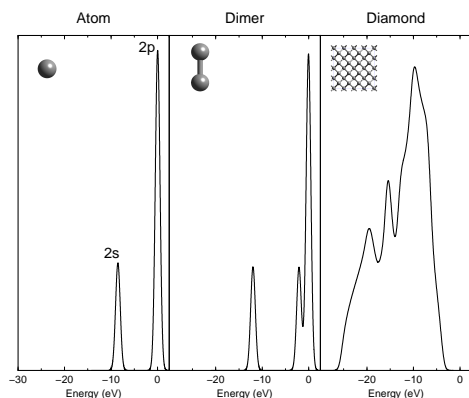
- Many techniques for extracting information about occupied and unoccupied electronic states
- Can be classified by:
 - between which states the transitions take place
 - how the transitions are induced
 - how the effect is measured
- For example, an Electron Energy Loss experiment



Pictorial representation of possible spectral processes for diamond.

A SIMPLE EXAMPLE

- In an isolated atom (e.g. C), the potential is spherically symmetric
- The energy levels are those of 1s, 2s, 2p ...
- As another atom approaches, the potential is disturbed and the levels split
- As more and more atoms arrive, the potential becomes further complicated, the levels become multiply split, and a better description is that of a continuum



Total densities of states for a carbon atom, dimer and diamond. These are the results of actual DFT pseudopotential calculations.

- Transitions caused by disturbances of the system
- A restriction to “perturbations” allows an interpretation through Fermi's Golden Rule

$$R_{i \rightarrow f} \propto |\langle \phi_i | \Delta \hat{V} | \phi_f \rangle|^2 \rho(E_f)$$

- Matrix element term $|\langle \phi_i | \Delta \hat{V} | \phi_f \rangle|^2$, probability that any transition will take place
- DOS term indicates how many such transitions are possible to a given energy
- $\Delta \hat{V}$ describes the perturbation

THEORY

What is the rôle of theory?

- To enable interpretation of highly nonintuitive experimental results
- In the building of a specialist intuition
- Use the theoretical understanding to suggest new routes to more information

How is this done?

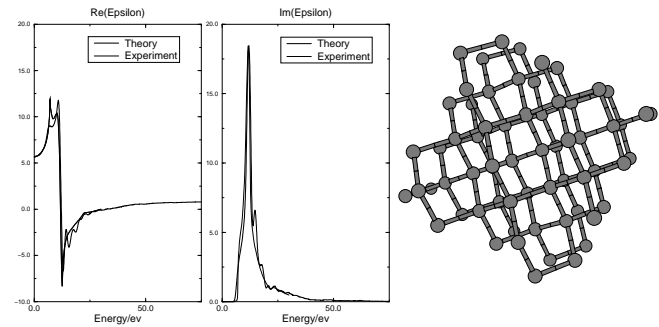
- Model experimental results at increasing levels of sophistication

BAND-STRUCTURE CALCULATIONS

- Kohn-Sham eigenstates do not formally correspond to single particle state required in spectral calculations
- However, it is nearly universal to interpret them in this way
- Very good agreement between theory and experiment has been found
- More accurate calculation of “quasiparticle” energies show a relation to Kohn-Sham energies

$$\varepsilon_2(\mathbf{q} \rightarrow \mathbf{0}_{\hat{\mathbf{u}}}, \hbar\omega) = \frac{2e^2\pi}{\Omega\varepsilon_0} \sum_{\mathbf{k}, v, c} |\langle \psi_{\mathbf{k}}^c | \hat{\mathbf{u}} \cdot \mathbf{r} | \psi_{\mathbf{k}}^v \rangle|^2 \delta(E_{\mathbf{k}}^c - E_{\mathbf{k}}^v - E),$$

- Similar to Fermi's Golden rule — ε_2 is related to real transitions
- ε_1 is obtained through a Kramers-Kronig transform — the response is causal
- There are many complications
 - Local field effects — the local electric field at any given site is not the external electric field
 - Excitonic effects are ignored
 - Kohn-Sham eigenvalues are used for quasiparticle energies
- Despite this — the dielectric properties of diamond are well reproduced



The complex dielectric function for diamond was calculated with a 0.5eV Gaussian smearing and a 0.7eV Scissor operator applied to the conduction bands.

PREDICTION OF NEW MATERIALS

- Prediction of new materials a central area of condensed matter science
- Aim to predict all possible crystal structures of a given composition
- Then screen for interesting/useful properties
- Synthesis may then be directed towards most promising candidates (properties and likely success)

POSSIBLE STRATEGIES

- “Brute force” methods, e.g. simulated annealing
 - either too time consuming for moderately complex structures if reliable models for the total energy are used
 - or, the models are not reliable (e.g. difficulties handling variety of bonding)
- Random approaches cannot prove all structures have been found
- A more elegant approach is presented ...

1. Graph theory used to enumerate all possible, topologically distinct trial structures
2. First principles quantum mechanics used to establish ...
 - stability
 - structural parameters
 - properties
 - density
 - bulk moduli
 - electronic structure

- sp^2 -bonding → each carbon atom bonded to 3 neighbours
- Framework → the structure is 3-dimensionally connected
- Graphite, fullerenes will not emerge
- Framework structures may be expected to have unusual properties — short strong bonds
- Considerable experimental effort in the synthesis of carbon based structures
- Maybe the predicted materials have been synthesised, but not recognised
- Or will inspire further work ...

STEP ONE - GRAPH THEORY

STEP TWO - FIRST PRINCIPLES QM

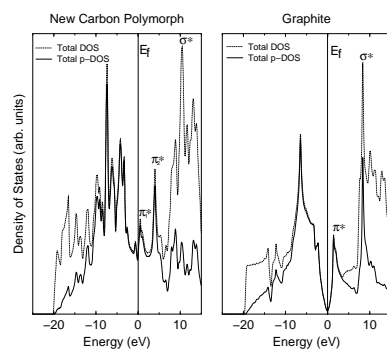
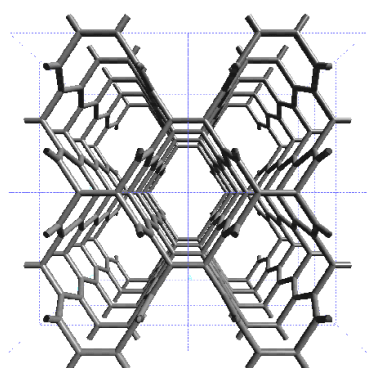
- Can be proved, Graph Theoretically, there are no sp^2 framework structures with 1,2,3, or 5 atoms per primitive cell
- Here, all with 4 and 6 are investigated
- A set of topologically distinct “trial structures” are passed on to the next stage – fourteen in this case

- Full geometry optimisation performed, with and without symmetry imposed
- Density Functional based total energy code, CASTEP
- Planewaves, Vanderbilt Ultrasoft pseudopotentials
- Generalised Gradient Approximation for exchange-correlation

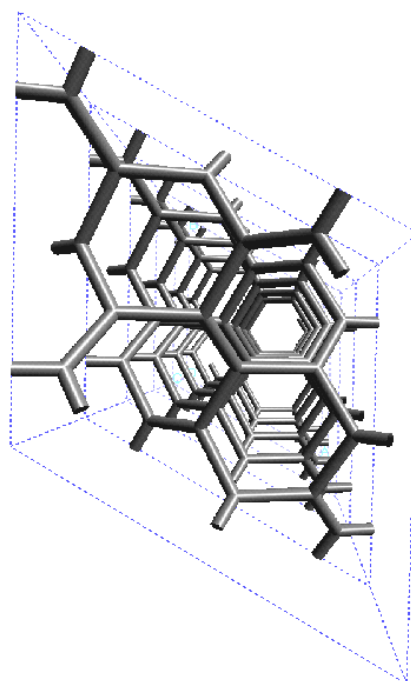
- All 14 structures could be relaxed
- Bond lengths between 1.34 and 1.48Å in all but one case — 1.55Å in a structure 4-fold coordinated
- Four membered rings — 6(3)3-26 — *not* unfavourable
- All structures less dense than diamond
- Around 0.4eV/atom less stable than diamond
- 6(3)1-10 the most stable, 6(3)4-25 and 6(3)2-22 close (within 0.03eV/atom)
- Bulk moduli — close to that of diamond, but 6(3)6-09 does not exceed it

- Electronically, the proposed polymorphs are found to have a wide variety of electronic properties
 - 6(3)2-22 is a 1.9eV semiconductor (GGA)
 - 6(3)1-10 is a semimetal, like graphite
 - 6(3)4-25 is metallic
- Further analysis of the DOS can lead to possible spectral signatures for these structures
- p-projected DOS are related to ELNES as could be measured in a transmission electron microscope — only small samples needed

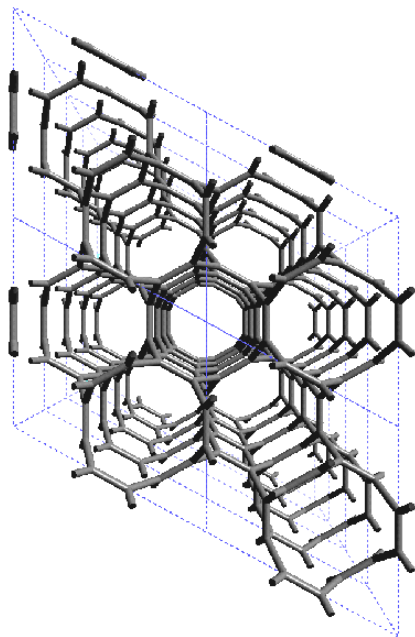
POLYMORPH 6(3)4-25



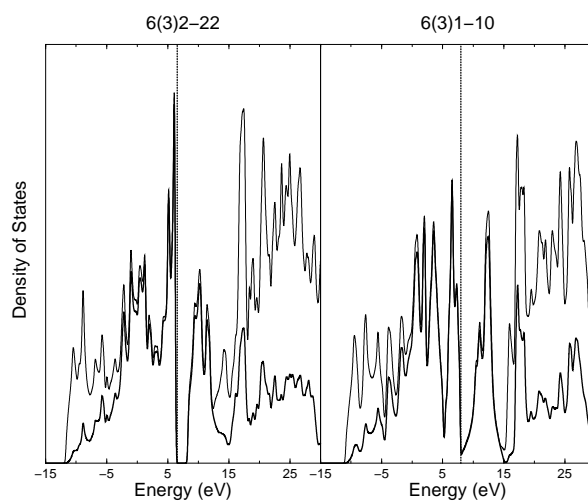
POLYMORPH 6(3)1-10



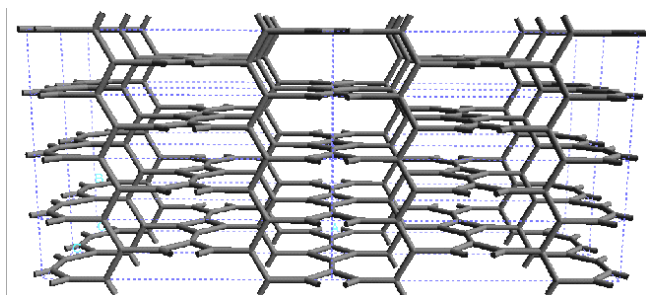
POLYMORPH 6(3)2-22



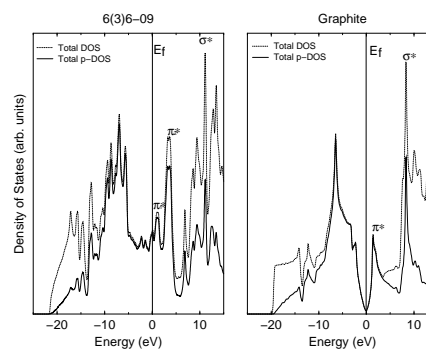
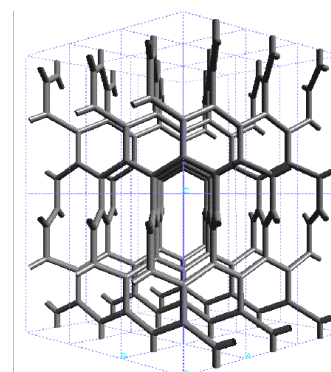
6(3)2-22 vs. 6(3)1-10



POLYMORPH 6(3)3-26



POLYMORPH 6(3)6-09



- The fertility of combining graph theory and quantum mechanics is demonstrated
- While some of the 14 structures have been previously known, the whole set was generated without prior knowledge, and leaves no gaps
- The method is open to extension — mixed coordination, species etc.
- New developments in QM total energy techniques can readily be applied
- We hope the structures themselves will be of interest — mechanical and electronic properties

B. Winkler, C.J. Pickard, V. Milman, W.E. Klee, and G. Thimm.

Prediction of a nanoporous sp²-carbon framework structure by combining graph theory with quantum mechanics.

Chemical Physics Letters, 312:536-541, 1999.