





hyperfine interactions

(and how to do it in WIEN2k)

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my talks on YouTube http://goo.gl/P2b1Hs



Kohn-Sham equations



$$E = T_o[\rho] - \int V_{ext} \rho(\vec{r}) d\vec{r} - \frac{1}{2} \int \frac{\rho}{2}$$
 interacting with electron charge distribution

nuclear point charges

1-electron equations (Kohn Sham)

vary p

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

$$\int \frac{\rho(\vec{r})}{|\vec{r}' - \vec{r}|} d\vec{r} \qquad \frac{\partial E_{xc}(\rho)}{\partial \rho}$$

$$\frac{\partial E_{xc}(\rho)}{\partial \rho}$$

$$\rho(\vec{r}) = \sum_{\varepsilon_i \le E_F} |\Phi_i|^2$$

$$E_{xc}^{LDA} \propto \int \rho(r) \ \varepsilon_{xc}^{\text{hom.}}[\rho(r)] \ dr$$

$$E_{xc}^{GGA} \propto \int \rho(r) \ F[\rho(r), \nabla \rho(r)] \ dr$$

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$$E_{xc}^{GGA} \propto \int \rho(r) \ F[\rho(r), \nabla \rho(r)] \ dr$$
but approximately

exchange and correlation effects,

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

Definition:

hyperfine interaction

all aspects of the nucleus-electron interaction that go beyond the nucleus as an electric point charge.

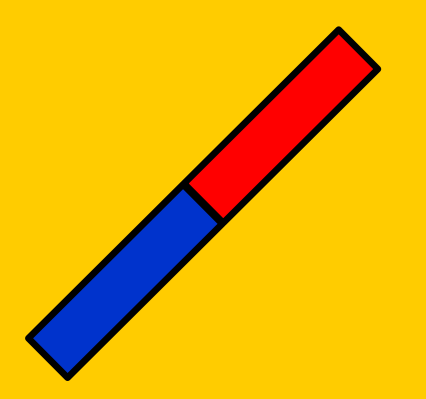
electric point charge

- volume
- shape

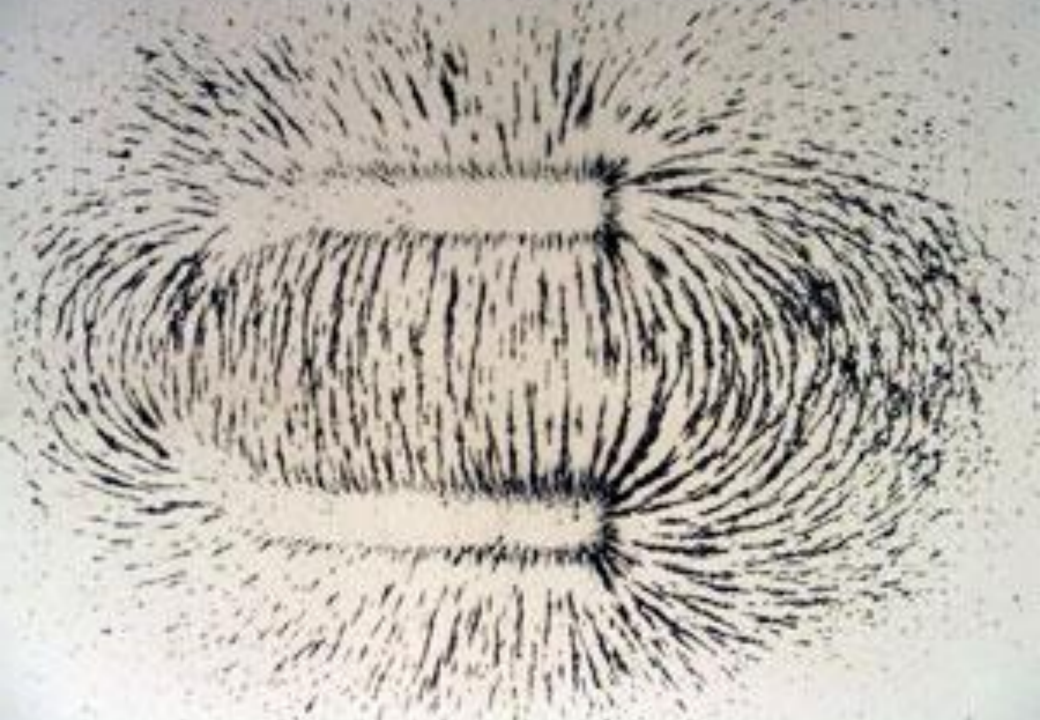
point volume shape magnetic moment magnetic hyperfine interaction

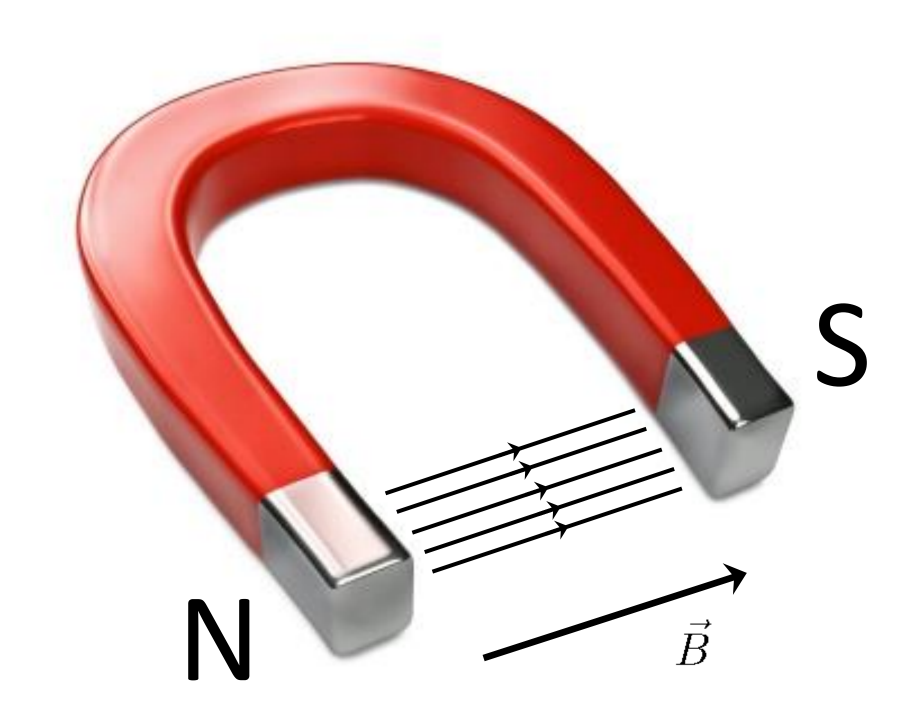
electric quadrupole interaction

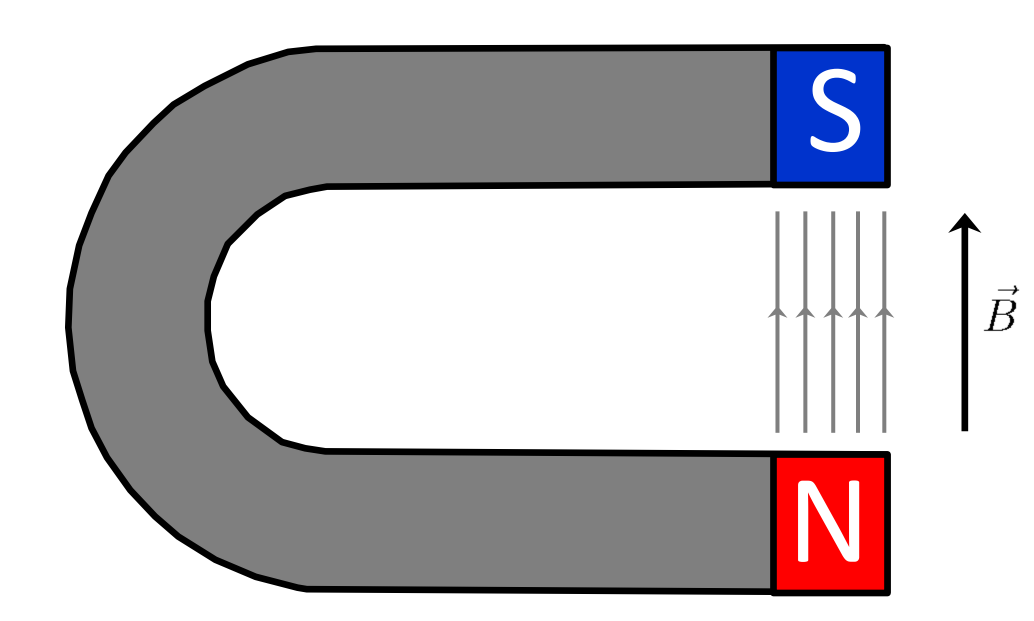
isomer shift

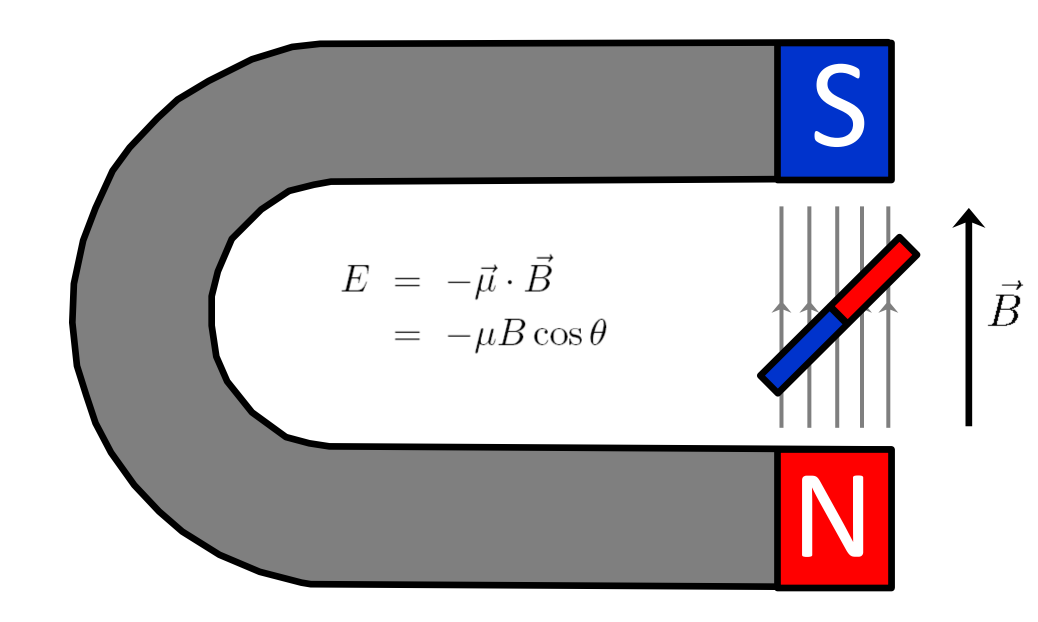


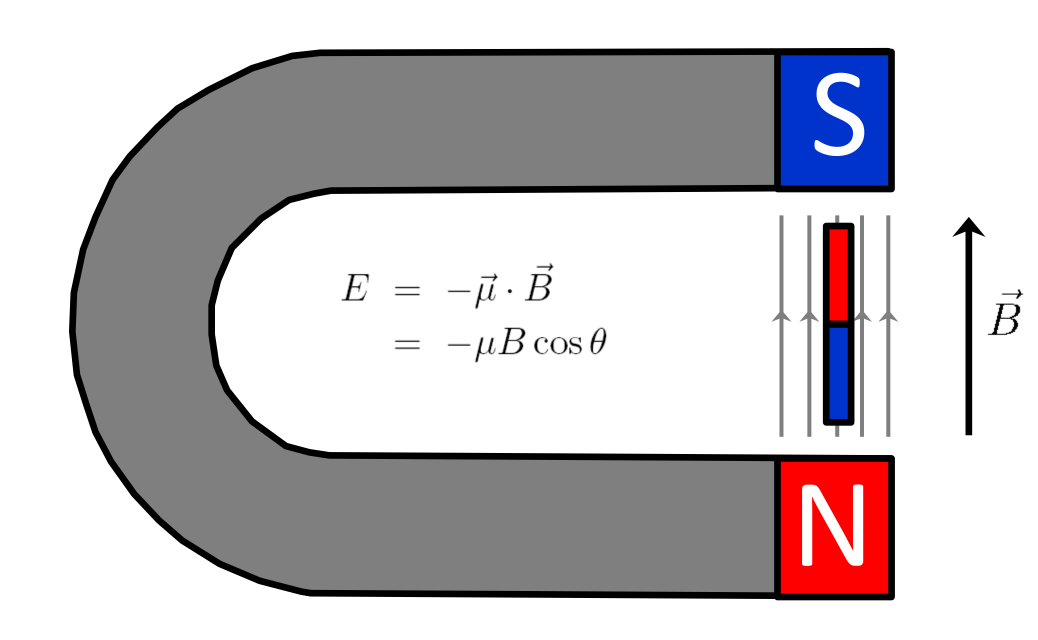


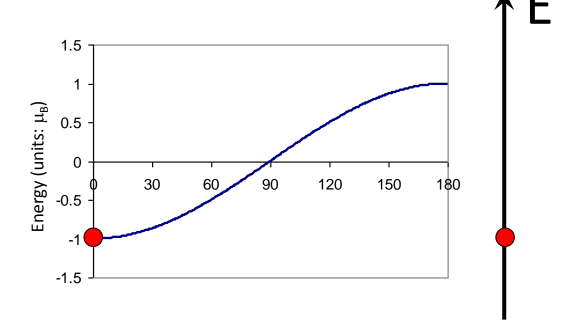




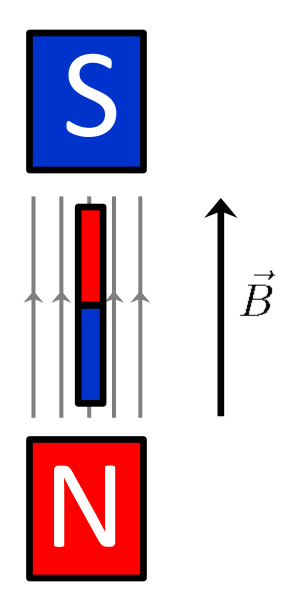


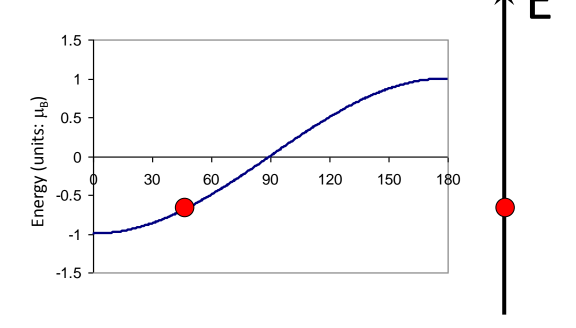




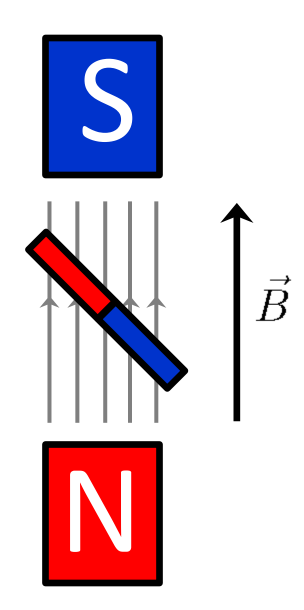


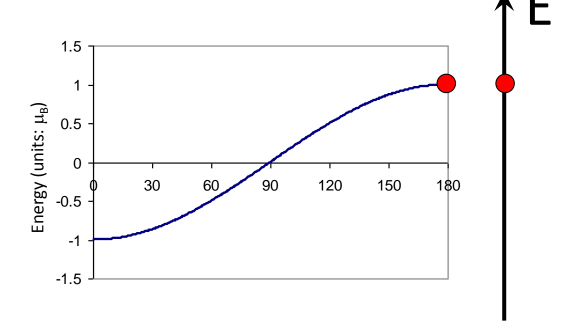
$$E = -\vec{\mu} \cdot \vec{B}$$
$$= -\mu B \cos \theta$$



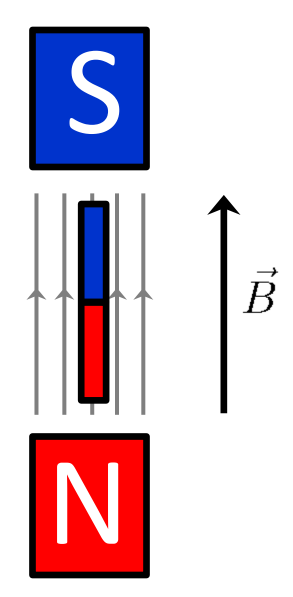


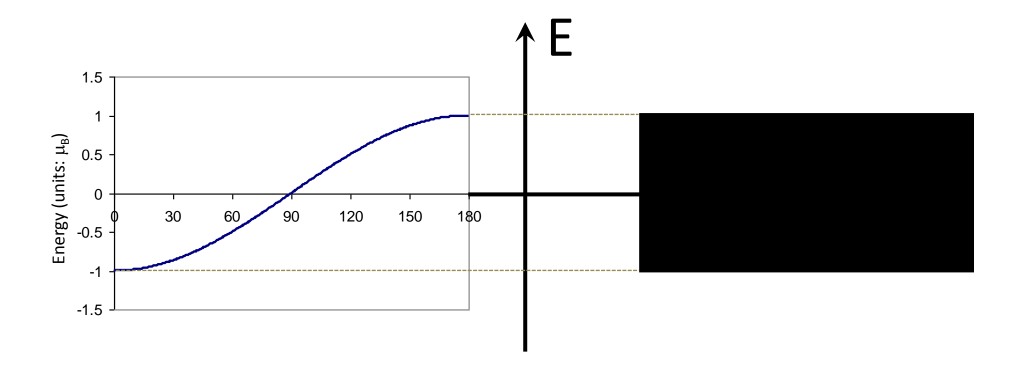
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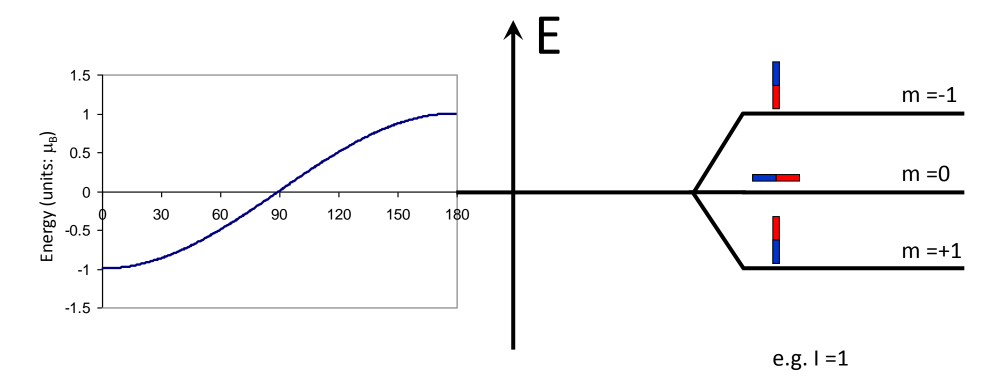




$$E = -\vec{\mu} \cdot \vec{B}$$
$$= -\mu B \cos \theta$$

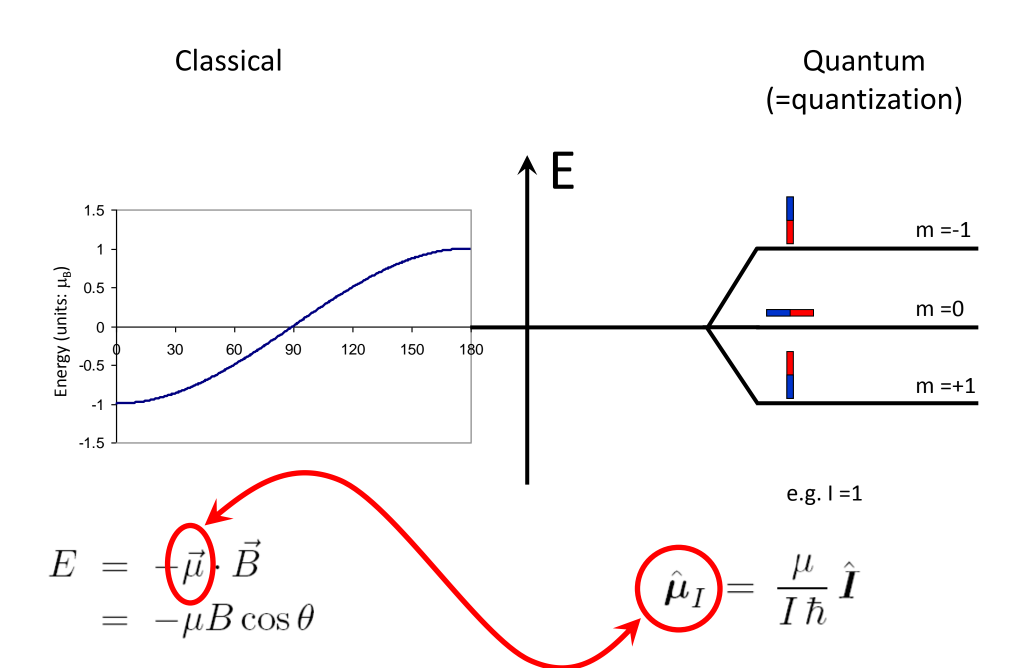
Classical

Quantum (=quantization)



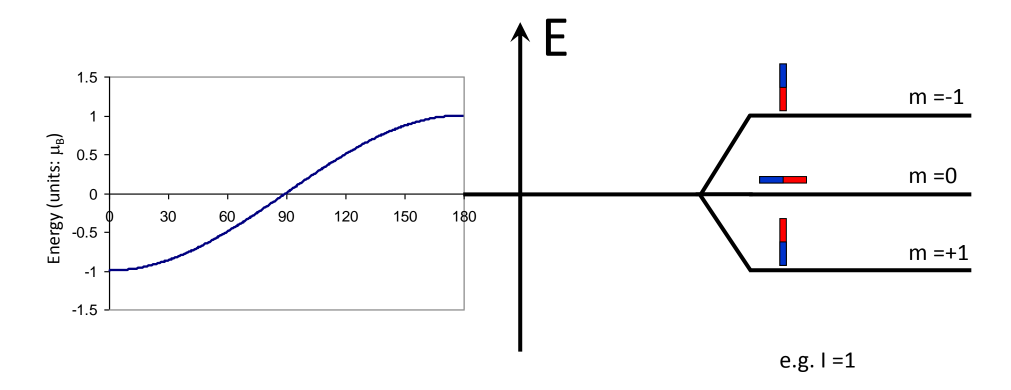
$$E = -\vec{\mu} \cdot \vec{B}$$
$$= -\mu B \cos \theta$$

$$\hat{\boldsymbol{\mu}}_{I} = \frac{\mu}{I \, \hbar} \, \hat{\boldsymbol{I}}$$



Classical

Quantum (=quantization)



Hamiltonian:

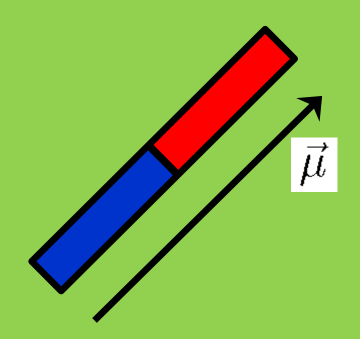
$$\hat{H} = -rac{\mu B}{I \hbar} \hat{I}_z$$

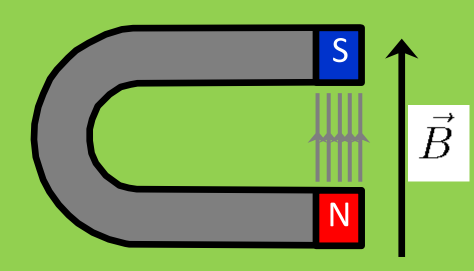
nuclear property

(vector)

electron property

(vector)





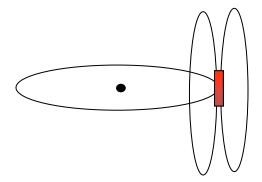
interaction energy (dot product):

$$E = -\vec{\mu} \cdot \vec{B}$$

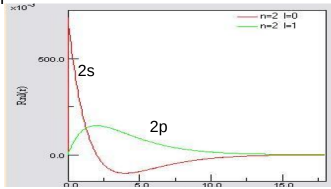
Source of magnetic fields at the nuclear site in an atom/solid

$$B_{tot} = B_{dip} + B_{orb} + B_{fermi} + B_{lat}$$

➤ B_{dip} = electron as bar magnet

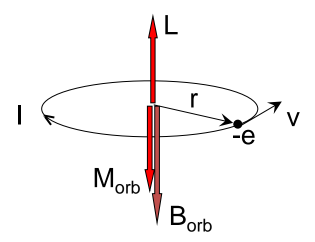


➤ B_{Fermi} = electron in nucleus

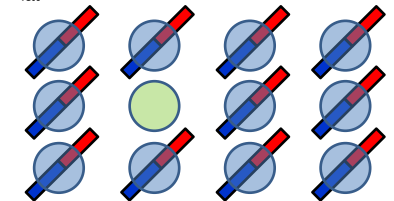


$$-\frac{2\mu_B\mu_0}{3}\left(\left|\psi_{e,\uparrow}(\mathbf{0})\right|^2-\left|\psi_{e,\downarrow}(\mathbf{0})\right|^2\right)$$

➤ B_{orb} = electron as current loop



 \triangleright B_{lat} = neighbours as bar magnets



How to do it in WIEN2k?

Magnetic hyperfine field

In regular scf file:

:HFFxxx

(Fermi contact contribution)

After post-processing with LAPWDM:

- orbital hyperfine field ("3 3" in case.indmc)
- dipolar hyperfine field ("3 5" in case.indmc)

```
in case.scfdmup
```

After post-processing with DIPAN:

lattice contribution

in case.outputdipan

more info: UG 7.8 (lapwdm) UG 8.3 (dipan)

How to do it in WIEN2k?

Magnetic hyperfine field

In regular scf file:

(Fermi conto https://youtu.be/L4t5ZAJAsoY

tutorial video

step-by-step

After post-processing with LAPWDM:

- orbital hyperfine field ("3 3" in case.indmc)
- · dipolar hyperfine field ("3 5" in case.indmc)

```
in case.scfdmup
```

:HFFxxx

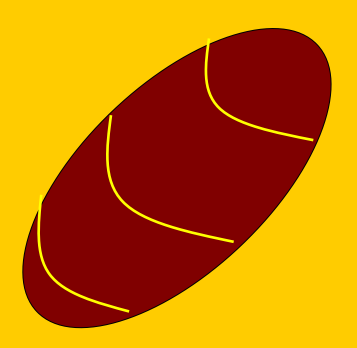
After post-processing with DIPAN:

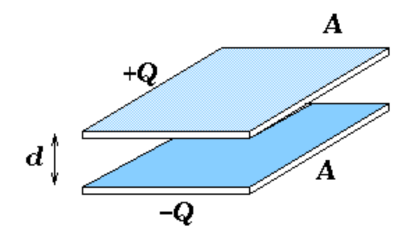
lattice contribution
 in case.outputdipan

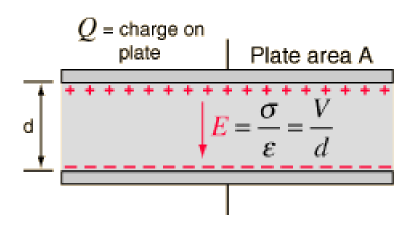
more info: UG 7.8 (lapwdm) UG 8.3 (dipan) magnetic hyperfine interaction

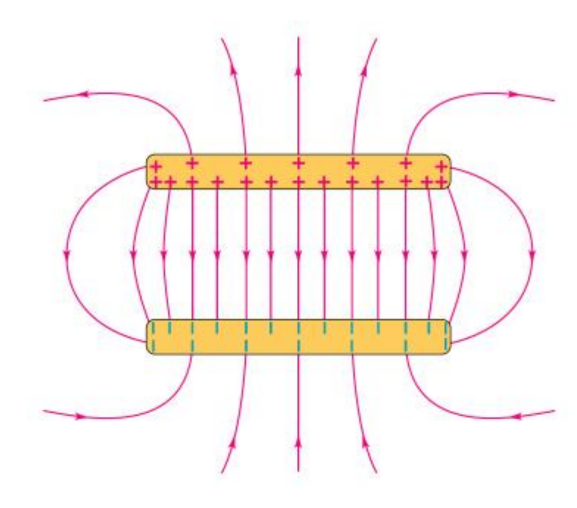
electric quadrupole interaction

isomer shift









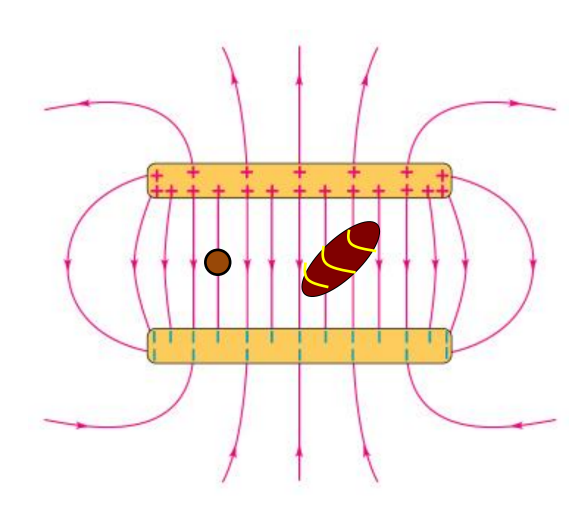
• Force on a point charge:

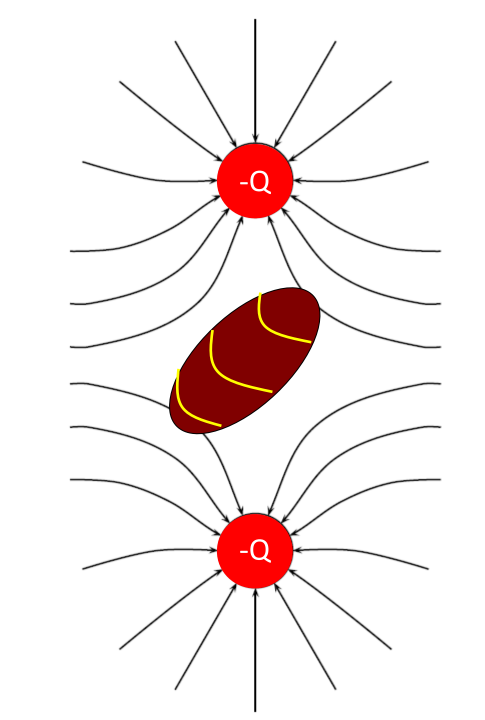
$$\vec{F} = Q\vec{E}$$

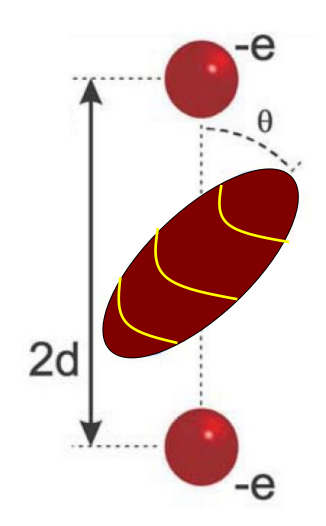
• Force on a general charge:

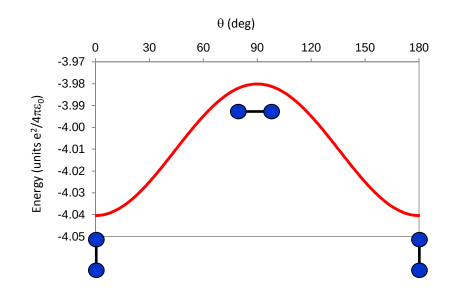
$$ec{F} = \int ec{E} \, dQ$$

$$= Q ec{E}$$



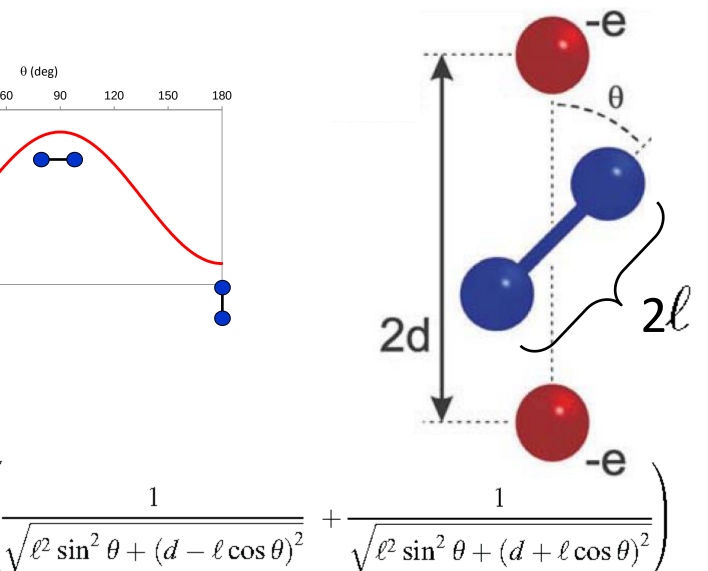


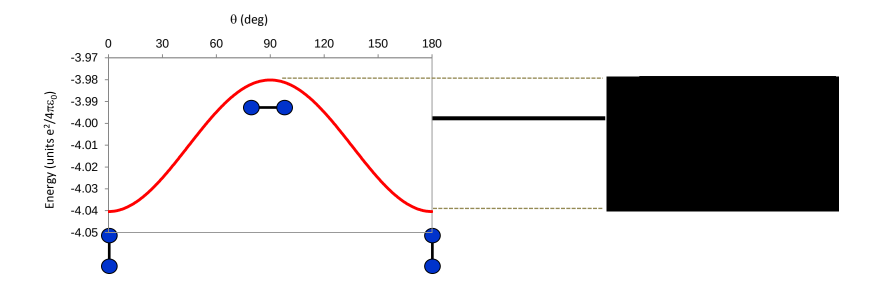


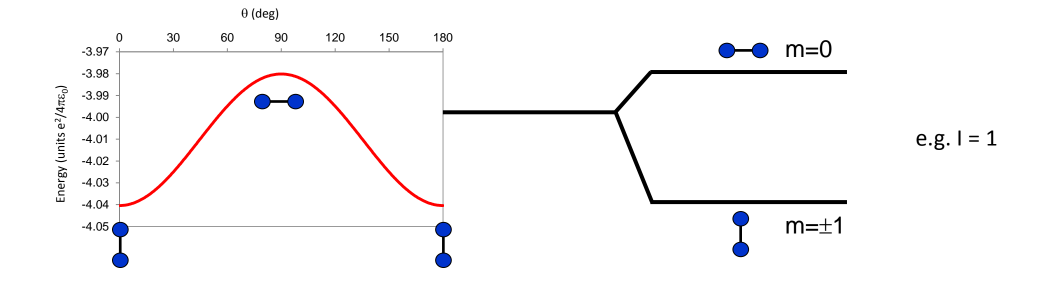


$$C = e^2/(4\pi\varepsilon_0)$$

$$E_0(heta) \; = \; - \; 2C \Biggl(rac{1}{\sqrt{\ell^2 \sin^2 heta + (d - \ell \cos heta)^2}}$$



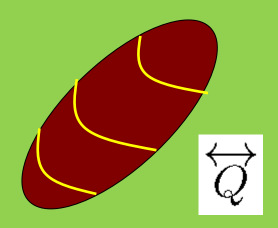




$$H_{qq}^{nuc} = \frac{e \, Q \, V_{zz}}{4 \, I (2I - 1) \, \hbar^2} \left[(3I_z^2 - I^2) + \frac{\eta}{2} \left(I_+^2 + I_-^2 \right) \right]$$

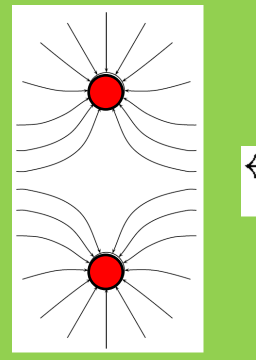
nuclear property

(tensor - rank 2)



electron property

(tensor – rank 2)





interaction energy (dot product):

$$E_Q \propto \overleftrightarrow{Q} \cdot \overleftrightarrow{V}$$

How to do it in WIEN2k?

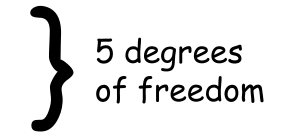
Electric-field gradient

In regular scf file:

:EFGxxx

:ETAxxx

Main directions of the EFG



Full analysis printed in case.output2 if EFG keyword in case.in2 is put (UG 7.6) (split into many different contributions)

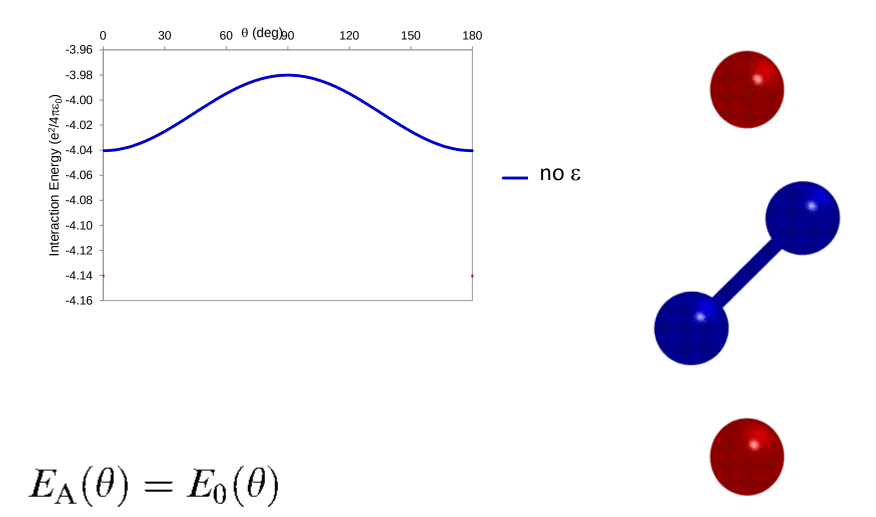
more info:

- · Blaha, Schwarz, Dederichs, PRB 37 (1988) 2792
- EFG document in wien2k FAQ (Katrin Koch, SC)

magnetic hyperfine interaction

electric quadrupole interaction

isomer shift



$$C = e^2/(4\pi\varepsilon_0)$$

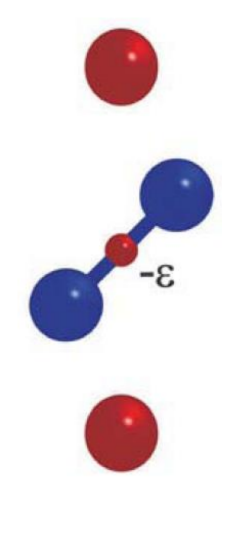
$$E_{\rm A}(\theta) = E_0(\theta) + \frac{-2\varepsilon C}{e\ell}$$

 $C = e^2/(4\pi\varepsilon_0)$

$$-\frac{2C}{e}\frac{\epsilon}{\ell} = -\frac{2C}{e}\frac{\epsilon}{\ell^3}\ell^2$$

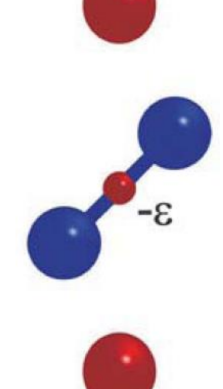
$$E_{A}(\theta) = E_{0}(\theta) + \underbrace{\frac{-2\varepsilon C}{e\ell}}_{E_{corA}}$$

$$C = e^{2}/(4\pi\varepsilon_{0})$$



$$\rho(0)$$
 $\langle R^2 \rangle$

$$\frac{2C}{\ell} \frac{\epsilon}{e} = -\frac{2C}{e} \left(\frac{\epsilon}{\ell^3}\right) \ell^2$$



$$E_{A}(\theta) = E_{0}(\theta) + \underbrace{\frac{-2\varepsilon C}{e\ell}}_{E_{corA}}$$

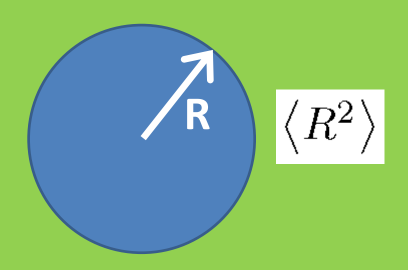
$$C = e^{2/(4\pi\varepsilon_{0})}$$

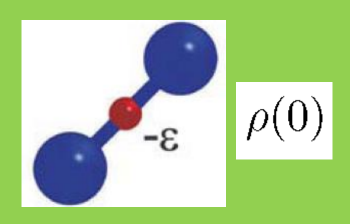
nuclear property

(scalar)

electron property

(scalar)





interaction energy (dot product):

$$E \propto \langle R^2 \rangle \cdot \rho(0)$$

How to do it in WIEN2k?

Isomer shift calculations

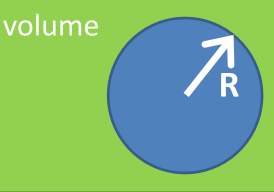
In regular scf file:

:RTOxxx

= electron density near the nucleus of atom xxx (i.e. at the first radial mesh point, typically 0.0005 au)

rank nuclear property • electron property

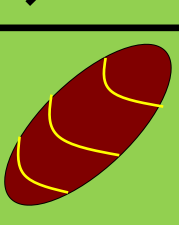
(dot product)

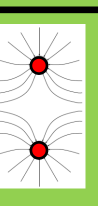


magnetic moment

shape







How to measure hyperfine interactions?

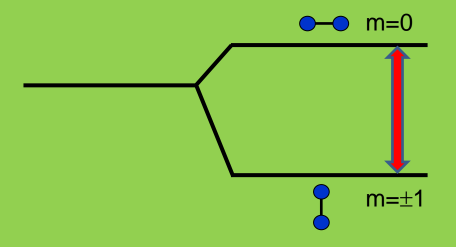


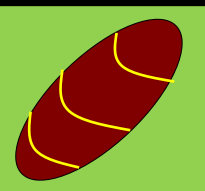
- NMR
- NQR
- Mössbauer spectroscopy
- TDPAC
- Laser spectroscopy
- LTNO
- NMR/ON
- PAD

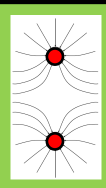


rank nuclear property • electron property

(dot product)

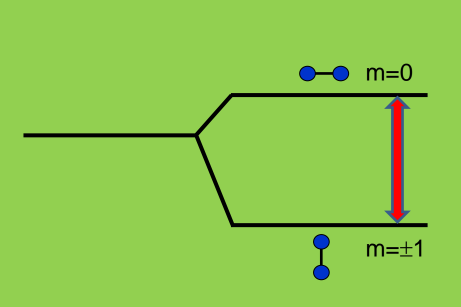


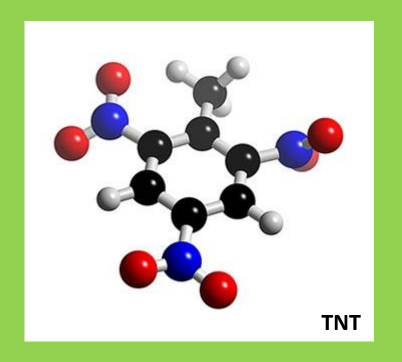




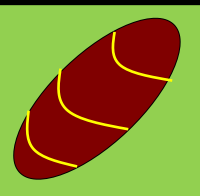
rank nuclear property • electron property

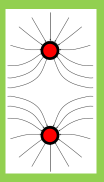
(dot product)











hyperfine interactions (and how to do it in WIEN2k)



want to read more ?

Katrin Rose and SC, Phys. Chem. Chem. Phys. 14 (2012) 11308-11317 http://dx.doi.org/10.1039/c2cp40740j

my talks on YouTube http://goo.gl/P2b1Hs