# Ab initio molecular dynamics: Basic Theory and Advanced Methods

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#### Wannier functions

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#### Examples: Theoretical infrared spectra

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# An Introduction to Maximally-Localized Wannier Functions

- The electronic ground state of a periodic system is usually described in terms of extended Bloch orbitals, simultaneous eigenstates of the periodic Hamiltonian and of the direct lattice translations.
- An alternative representation in terms of localized orbitals has been introduced by Gregory Wannier in 1937
- powerful tool in the study of the electronic and dielectric properties of materials: they are the solid-state equivalent of "localized molecular orbitals" and provide a picture of the nature of chemical bonding

- the charge center of a MLWF provides a kind of classical correspondence for the "location of an electron" (or electron pair) in a quantum-mechanical insulator.
- $\mathsf{E}.\mathsf{g}.:$  the ethylene molecule



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one of four C-H bond	One of two double bonds	
The single C-C bond	One of two <i>p</i> -like orbitals	
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#### E.g.: Platinum wire





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### Position operator in periodic systems

The position operator is ill defined in periodic boundary conditions. Indeed for a single particle wavefunction in periodic boundary conditions:

$$\phi(x+L) = \phi(x) \tag{1}$$

but  $x\phi(x)$  is not a periodic.

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For a non periodic system we can define the operator:

$$\langle X \rangle = \left\langle \Psi_0 | \hat{X} | \Psi_0 \right\rangle = \int x n(x) dx$$
 (2)

where n(x) is the one particle density,  $\hat{X} = \sum_{i=1}^{N} x_i$  and  $x_i$  the position operator of the *i*th particle in one dimension.

In periodic boundary condition we can on the other end define:

$$\langle X \rangle = \frac{L}{2\pi} \ln \ln \left\langle \Psi_0 | \exp \left[ i \frac{2\pi}{L} \hat{X} \right] | \Psi_0 \right\rangle$$
(3)

NOTE: this is not simply the expectation value of an operator, since it contains the imaginary part of the logarithm. The main ingredient is the operator  $\exp\left[i\frac{2\pi}{L}\hat{X}\right]$ , which is a many body operator.

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The electronic polarization corresponding to this definition of the position operator is:

$$P^{el} = \lim_{L \to \infty} \frac{-e}{2\pi} \ln \ln \left\langle \Psi_0 \right| \exp \left[ i \frac{2\pi}{L} \hat{X} \right] \left| \Psi_0 \right\rangle \tag{4}$$

where -e is the electron charge.

For a system of non-interacting particles, e.g. a crystal of lattice constant a, where periodic boundary conditions over a M linear cells (L = ma) are imposed, the Bloch vectors in the reciprocal cell are:

$$q_s = \frac{2\pi}{Ma}s,$$
  $s = 0, 1, ...., M-1$  (5)

The one-body orbitals can be chosen to have the Bloch from:

$$\phi_{q_s,m}(x+\tau) = \exp[1q_s\tau]\phi_{q_s,m}(x) \tag{6}$$

where  $\tau = la$  and m is a band index. The wavefunction can be written

$$|\Psi_0\rangle = \mathcal{A} \prod_{m=1}^{N/M} \prod_{s=0}^{M-1} \phi_{q_s,m}(x) \tag{7}$$

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#### The expectation value:

$$\langle X 
angle = rac{L}{2\pi} \operatorname{Im} \ln \left\langle \Psi_0 
ight| \exp \left[ i rac{2\pi}{L} \hat{X} 
ight] |\Psi_0 
ight
angle$$

can be recasted as:

$$\langle X \rangle = -\frac{L}{2\pi} \operatorname{Im} \ln \det \mathbf{S}$$
 (8)

where

$$S_{sm,s'm'} = \int_0^L \phi_{q_s,m}^* \exp\left[-i\frac{2\pi}{L}x\right] \phi_{q'_s,m'} dx \tag{9}$$

Using the  $\Gamma\text{-point}$  only (equivalent to M=1) the matrix  $S_{sm,s'm'}$  is now defined as

$$S_{n,m} = \int_0^L \phi_n^* \exp\left[-i\frac{2\pi}{L}x\right] \phi_m dx \tag{10}$$

and  $\phi_m$  are the Kohn-Sham orbitals.

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# Localization functionals

Wannier functions are defined in term of an unitary transformation of occupied Bloch orbitals.

The problem is their non-uniqueness.

Marzari and Vanderbilt<sup>1</sup> proposed a way to solve this by requiring that the total spread of the localized function should be minimal. In one dimension the functional for the spread is

$$\Omega = \langle x^2 \rangle - \langle x \rangle^2 = \frac{1}{(2\pi)^2} \ln |z|^2 \tag{11}$$

where

$$z = \int_0^L \exp\left[i\frac{2\pi x}{L}\right] |\phi(x)|^2 dx$$
 (12)

<sup>1</sup>Phys Rev B 56, 12847 (1997)

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## Localization methods

The mathematical problem which defines the localization procedure is to find the unitary transformation  ${\bm U}$  of the orbitals

$$|\tilde{\phi_n}\rangle = \sum_i U_{in} |\phi_i\rangle \tag{13}$$

that simultaneously minimize the spread functional  $\Omega$ .

The simplest approach to parameterize the unitary transformation is the Jacobi optimization procedure and consists in a direct product of elementary plane rotations of couples of orbitals. The transformed expectation values are obtained as

$$\tilde{z}_{I,i} = \cos(\gamma) z_{I,i} + \sin(\gamma) z_{I,j}$$
(14)

$$\tilde{z}_{I,j} = -\sin(\gamma)z_{I,i} + \cos(\gamma)z_{I,j}$$
(15)

To obtain the optimal angle:

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## Molecular properties from WF

Maximally localized Wannier functions are used as a tool to understand the nature of the chemical bonding.

In the case of a pseudopotential approach the localized orbitals only refers to the valence electrons in the effective charge of the nuclei. In the frame of the Kohn-Sham approach the polarization of a system is defined as

$$M_{\alpha}^{el} = \frac{2e}{|\mathbf{G}_{\alpha}|} \operatorname{Im} \ln \det \mathbf{Z}_{\alpha}$$
(17)

where  $\alpha$  denotes the Cartesian components and where the matrix:

$$(\mathbf{Z}_{\alpha})_{kl} = <\phi_k \exp[-i\mathbf{G}_{\alpha} \cdot \mathbf{r}]|\phi_l>$$
(18)

is defined in terms of the Kohn-Sham orbitals  $\phi_k$ .

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For the analysis of a liquid or solid  $M^{el}$  can be approximated as a sum of molecular dipole moments  $\mu_{I}^{el}$ .

$$M^{el} \approx \sum_{I} \mu_{I}^{el} \tag{19}$$

The expectation value of the position operator is given:

$$\mathbf{r}_k = -\frac{L}{2\pi} \operatorname{Im} \ln \mathbf{z}_k \tag{20}$$

which is just the center of a given maximally localized Wannier function (k labels the different electrons).

With this definition the electronic part of of the total dipole moment of the supercell can be approximated by

$$M^{el} \approx -2e \sum_{k} \mathbf{r}_{k}$$
 (21)

and molecular dipole moments can be defined as:

$$\mu_l^{el} = -2e \sum_{k \in I} \mathbf{r}_k \tag{22}$$

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## Infrared absorption spectra

The dipole autocorrelation function needed to calculate infrared spectra

$$C(t) = \langle \mathbf{M}(t) \cdot \mathbf{M}(0) \rangle \tag{23}$$

can be decomposed as

$$C(t) = C_{SS}(t) + C_{MM}(t) + C_{SM}(t)$$
 (24)

where

$$C_{SS}(t) = \sum_{II}^{solvent} < \mu_I(t) \cdot \mu_J(0) >$$
(25)

$$C_{MM}(t) = < \mu_0(t) \cdot \mu_0(0) >$$
 (26)

$$C_{SM}(t) = \sum_{I}^{SM(t)} (\langle \mu_{I}(t) \cdot \mu_{0}(0) \rangle + \langle \mu_{0}(t) \cdot \mu_{I}(0) \rangle)$$
(27)

After Fourier transform,  $C_{SS}(t)$  yields the IR spectra of the solvent, while  $C_{MM}(t)$  and  $C_{SM}(t)$  contribute to the spectrum of the solvent.

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- Whether or not the cross correlation term C<sub>SM</sub>(t) should be included in the analysis of the solvent spectrum has been answered differently by different authors.
- Care has to be taken to correct for nuclei quantum effects which have been neglected so far.

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## Atomic charges

It is possible to calculate atomic charges from Wannier centers, using some approximations.

The approach is closely related to the RESP approach.

Assuming neutral molecules

$$\sum_{A}^{N_{at}} Z_A - Mq_w = 0 \tag{28}$$

We want a set of charges  $q_A$  which reproduce the electrostatic potential V.

$$V(\mathbf{r}) = \sum_{A} \frac{Z_{A}}{|\mathbf{R}_{A} - \mathbf{r}|} - \sum_{a} \frac{q_{w}}{|\mathbf{r}_{a} - \mathbf{r}|}$$
(29)

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The potential is sampled on many positions  $\mathbf{r}_i$  outside the molecule and optimized with respect to  $q_A$ . Reference charges  $q_A^0$  are also included.

$$\Omega(\{q_A\},\lambda,\{\epsilon_\alpha\}) = \sum_i \left(\sum_A \frac{Z_A}{|\mathbf{R}_A - \mathbf{r}_i|} - \sum_a \frac{q_w}{|\mathbf{r}_a - \mathbf{r}|} - \sum_A \frac{q_A}{|\mathbf{R}_A - \mathbf{r}_i|}\right)^2 + w \sum_A (q_A - q_A^0)^2 - \lambda \sum_A q_A - \sum_{\alpha = x, y, z} \epsilon_\alpha \left(\sum_A q_A r_A^\alpha - \mu_\alpha\right)$$

The conditions:

$$\frac{\partial\Omega}{\partial q_A} = 0, \qquad \frac{\partial\Omega}{\partial\lambda} = 0, \qquad , \frac{\partial\Omega}{\partial\epsilon_{\alpha}} = 0$$
 (30)

reduce to a system of linear equation of dimension  $N_{at} + 4$ .

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## Infrared Spectrum of Aqueous Uracil

from M.-P. Gaigeot and M. Sprik, JPC B 107, 10344 (2003)

- The spectrum is obtained by FT of the time correlation function of total dipole moment.
- Here, unlike in classical model studies, there no separation between permanent and induced dipole moments (all is contained in the total electronic polarization.
- The spectroscopy of aqueous solutions adds further challenges. Because of the intense IR activity of water, the signal of the hydrated solute is usually completely obscured. This is a problem in experiment, as well as computation.

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Motivations: Nucleic acids are exposed to water solvent in a number of biological structures. The hydration of nucleic acids of great interest and extensively investigated by infrared spectroscopy.



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IR spectrum in vacuo (red) and in aqueous solution (blue) from CPMD at 20 and 310 K, respectively. Inset: experimental IR from ref <sup>2</sup> <sup>2</sup>Aamouche, A.; Berthier, G.; Cadioli, B.; Gallinella, E.; Ghomi, M. J. Mol. Struct. (THEOCHEM) 1998, 426, 307.

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► The 1400-1200 cm<sup>-1</sup> range is predominantly δ(N-H) and δ(CH) bending.

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## The IR spectrum in solution

- Substantial enhancement of the molecular dipole from 4.50 D of the isolated molecule to 7.50 D in solution (0.65 increase.)
- NO evidence of a reverse effect of the solute on the solvent. Dipole moments, even for molecules in the first hydration shell, were essentially identical to the pure liquid water values.

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On the other hand, the number and positions of infrared active bands is completely different in the 1500-1000 cm<sup>-1</sup> Vacuum: only two IR bands. Solution: five can be distinguished Marialore Sulpizi Ab initio molecular dynamics: Basic Theory and Advanced Met

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## Comparison to Experiment

- Overall patterns are very similar
- Theory and experiment are in remarkable good agreement for the main band generated by C-O and C-C stretching
- overall red shift in the calculation can be the BLYP functional, as well as the value of the fictitious electron mass used in the Car-Parrinello dynamics.
- However, the band calculated around 1330 cm-1 is too broad (assigned to N3-H3 bending motion).

(i) dynamical charges of the amide N3-H3 group are overestimated

(ii) H-bonds formed by this group to the water molecules are too strong.

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## Infrared Spectrum of N-methylacetamine<sup>3</sup>



Motivations:Structural motifs such as  $\alpha$ -helices or  $\beta$ -sheets can be identified by characteristic changes in position and shape of linear infrared or Raman bands, in particular the amide bands are used as structural probes The

amide I band shape is strongly affected by formation of intermolecular hydrogen bonds with the solvent.

<sup>3</sup>M-P Gaigeot et al JCTC 2005, 1, 772.

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Gas-phase trans

The most intense IR band located at 1609  $\rm cm^{-1}$  is related to the amide I mode.

In the domain 1500 to 1300  $\text{cm}^{-1}$  are 4 active bands of much lower intensity.

More intense active band at 1189  $\rm cm^{-1}$  is due to amide III.

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Isolated Cis-NMA Amidel &C-H), v(N-331 <sup>‡03</sup> 606 Aqueous Cis-NMA

328-1339-1365

407-1382 470-1430 560-1534

methyl rocking

600

#### Gas-phase cis

Infrared Spectrum of Aqueous Uracil

Infrared Spectrum of N-methylacetamine

The cis-NMA spectrum is totally different from the trans-NMA spectrum: in particular, the number of active bands is reduced from six to four.

The most intense active band is associated with the amide I mode.

The amide I band is peaked in the two spectra (cis, trans) trans- and cis-NMA at the same frequency, and for both molecules, this band has the most intense activity of the whole spectrum.

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#### Aqueous trans

Red-shift of the amide I and blue-shift of amide II band reducing the 150 cm<sup>-1</sup> frequency gap observed in the gas phase to 40-50 cm<sup>-1</sup> in solution.

blue-shift of 40  $\rm cm^{-1}$  for Amide III band.

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#### Aqueous cis

Amide I is  $80 \text{ cm}^{-1}$  down-shifted by the presence of the solvent, whereas amide III is blue-shifted by the same amount. The other bands keep more or less the same position.

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#### **Comparison to experiments**

Although band positions are systematically red-shifted (mainly due to our use of the BLYP functional), the main features, including band shapes and relative bands gaps, can be satisfactorily reproduced, the set of the se

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A factor of 1.064 used to adjusts exactly the position of the amide III band to the exp value of 1266 cm<sup>-1</sup>. Doing so, the theoretical amide I band is close to the correct position (its intensity is overestimated). The intermediate region between 1300 and 1600 cm<sup>-1</sup> is only approximately reproduced, but with correct features.

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Frequencies have been uniformly scaled by a factor of 1.117 in order to align the amide III band with the experimental peak at 1318 cm<sup>-1</sup>. The broad and intense double band in the 1500-1700 cm<sup>-1</sup> range due to the mixing of amide I and amide II is very well reproduced.