First principles calculation of vibrational Raman spectra in large systems: signature of small rings in crystalline SiO$_2$

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We present an approach for the efficient calculation of vibrational Raman intensities in periodic systems within density functional theory. The Raman intensities are computed from the second order derivative of the electronic density matrix with respect to a uniform electric field. In contrast to previous approaches, the computational effort required by our method for the evaluation of the intensities is negligible compared to that required for the calculation of vibrational frequencies. As a first application, we study the signature of 3- and 4-membered rings in the Raman spectra of several polymorphs of SiO$_2$, including a zeolite having 102 atoms per unit cell.

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Vibrational Raman spectroscopy [1] is one of the most widely used optical techniques in materials science. It is a standard method for quality control in production lines. It is very effective in determining the occurrence of new phases or structural changes at extreme conditions (high pressure and temperature), where it is often preferred to the more difficult and less readily available x-ray diffraction experiments based on synchrotron sources [2]. Moreover, it can be used in the absence of long-range structural order as for liquid or amorphous materials [3, 4, 5]. The theoretical determination of Raman spectra is highly desirable, since it can be used to associate Raman lines to specific microscopic structures.

Density functional theory (DFT) [6] can be used to determine with high accuracy both frequencies and intensities of Raman spectra. Vibrational frequencies can be efficiently determined using first order response [3, 4]. Within this approach Raman intensities (RI) calculation is also possible, but requires a computational time significantly larger and is not practical for large systems. Thus, while many examples of frequency calculations have been reported so far [3, 4], RI were predicted from first-principles in a very limited number of cases involving systems with a small number of atoms [6, 7, 8]. In this Letter we show that it is possible to obtain RI in extended solids with a computational cost negligible with respect to that required for the frequency determination. The efficiency of our approach will lead ab-initio calculations to become a routine instrument for the interpretation of experimental Raman data. Our method is based on second order response to DFT. In particular, we compute the second order derivative of the electronic density matrix with respect to a uniform electric field, using pseudopotentials and periodic boundary conditions. As a first application we calculate Raman spectra of several SiO$_2$ polymorphs, including a zeolite having 102 atoms per unit cell [12].

In a Raman spectrum the peak positions are fixed by the frequencies $\omega_\nu$ of the optical phonons with null wavevector. In non-resonant Stokes Raman spectra of harmonic solids, the peak intensities $I^\nu$ can be computed within the Placzek approximation [4] as:

$$I^\nu \propto |\mathbf{e}_i \cdot \hat{\mathbf{A}}^\nu \cdot \mathbf{e}_f|^2 \frac{1}{\omega_\nu} (n_\nu + 1),$$

where $\mathbf{e}_i$ ($\mathbf{e}_f$) is the polarization of the incident (scattered) radiation, $n_\nu = (\exp(\hbar \omega_\nu/k_B T) - 1)^{-1}$, $T$ is the temperature, and

$$A^\nu_{lm} = \sum_{k \gamma} \frac{\partial^2 \mathcal{E}^{ei}}{\partial E_i \partial E_m \partial u_{k \gamma}} \frac{w^\nu_{k \gamma}}{\sqrt{M_\gamma}}.$$ 

Here $\mathcal{E}^{ei}$ is the electronic energy of the system, $E_i$ is the $l$th Cartesian component of a uniform electric field, $u_{k \gamma}$ is the displacement of the $\gamma$th atom in the $k$th direction, $M_\gamma$ is the atomic mass, and $w^\nu_{k \gamma}$ is the orthonormal vibrational eigenmode $\nu$.

Linear response [4, 8] can be used to determine $\omega_\nu$, $w^\nu$, and also the dielectric tensor $\mathcal{E}^{ei}$ defined as $\epsilon^{ei}_{lm} = \delta_{lm} - (4\pi/\Omega) \partial^2 \mathcal{E}^{ei}/(\partial E_i \partial E_m)$, where $\Omega$ is the cell volume. RI have been computed [4, 10] through Eq. (4), obtaining $\hat{\mathbf{A}}^\nu$ by finite-differences derivation of $\mathcal{E}^{ei}$ with respect to $u_{k \gamma}$. This approach requires $36N^3$ linear response calculations, where $N^3$ is the number of atoms. Thus, the scaling of the RI calculation is the same as that of the frequency calculation with a much larger prefactor. This has limited the applications of this approach to small systems. RI have also been computed from the dynamical autocorrelation functions of $\mathcal{E}^{ei}$ in a molecular dynamics (MD) run [11]. This approach also copes with liquids or anharmonic solids, but is very demanding, requiring the calculation of $\mathcal{E}^{ei}$ at each MD step.

Alternatively, RI can be obtained knowing the second order derivative of the DFT density matrix $\rho = \sum \gamma |\psi_\gamma\rangle \langle \psi_\gamma|$ being $|\psi_\gamma\rangle$ the normalized occupied Kohn-Sham (KS) eigenstates [3]. In fact, according to the well known Hellmann-Feynman theorem

$$\frac{\partial \mathcal{E}^{ei}}{\partial u_{k \gamma}} = 2 \text{Tr} \left\{ \rho \frac{\partial \mathcal{E}^{ext}}{\partial u_{k \gamma}} \right\},$$

where $\mathcal{E}^{ext}$ is the external energy of the system. This equation provides a simple solution to determine RI [9].
where $\text{Tr}\{O\}$ is the trace of the operator $O$, and $v^{\text{ext}}$ is the external ionic potential (the KS self-consistent potential is $V^{\text{KS}} = V^{\text{Hxc}} + v^{\text{ext}}$, where $V^{\text{Hxc}}$ is the sum of the Hartree and the exchange-correlation potential). Thus

$$\frac{\partial^3 E^{\text{cl}}}{\partial E_{\text{cf}} \partial E_{\text{cm}} \partial u_{k\gamma}} = 2 \text{Tr} \left\{ \left( \frac{\partial^2 \rho}{\partial E_{\text{cf}} \partial E_{\text{cm}}} \right) \frac{\partial v^{\text{ext}}}{\partial u_{k\gamma}} \right\}. \quad (3)$$

The $\partial^2 \rho/(\partial E_{\text{cf}} \partial E_{\text{cm}})$ calculation requires six second-order calculations, instead of the 36 $N^{3r}$ first-order calculations needed for the finite-differentiation $[10]$. Because of this better size-scaling, the $\hat{A}^r$ calculation through Eq. (3) is much more efficient and the time for RI calculation is negligible compared to that for $\omega_{\nu}$ in large systems.

The approach based on Eq. (3) has already been used in isolated molecules [13] but never in extended systems. Indeed, in solids the calculation of $\partial^2 \rho/(\partial E_{\text{cf}} \partial E_{\text{cm}})$ is not trivial because the position operator, required by the electric field perturbation, is ill-defined in periodic boundary conditions. Because of this, although a formalism to calculate derivatives of $\rho$ at any order was proposed by Gonze already in 1995 [6], only very recently Nunes and Gonze [14] were able to include perturbations due to macroscopic electric fields. To do that, they use the polarization-Berry phase formalism [15], arguing that this concept remains valid in the presence of finite electric fields. This approach has been applied so far to a one dimensional non-self-consistent model [14]. In the following we give an expression for the second derivative of $\rho$, that does not require the Berry phase formalism to cope with uniform electric fields, and we use it to compute $\hat{A}^r$ in real systems with a DFT self-consistent Hamiltonian.

The derivative of $\rho$ with respect to two generic perturbation parameters $\lambda$ and $\mu$ is:

$$\frac{\partial^2 \rho}{\partial \lambda \partial \mu} = \sum_v \left[ |P \eta^{(\lambda,\mu)}_v\rangle \langle \psi_v| + |P \partial \eta^{(\lambda,\mu)}_v/\partial \lambda \rangle \partial \psi_v/\partial \mu |P + \right.$$ \hspace{1cm} \left. \sum_{v'} |\psi_{v'}\rangle \langle \partial \psi_{v'}/\partial \lambda |P |P \partial \eta^{(\lambda,\mu)}_v/\partial \mu \rangle |\psi_v| \right] + cc, \quad (4)$$

where $P = (1 - \rho)$ is the projector on the empty state subspace, the sums over $v$ and $v'$ run over the occupied states, and $|\eta^{(\lambda,\mu)}_v\rangle$ are the second derivatives of the occupied KS-orbitals in the parallel-transport gauge [6]. According to our derivation:

$$|P \partial \psi_v/\partial \lambda \rangle = \hat{G}_v \left[ \frac{\partial V^{\text{KS}}}{\partial \lambda} \rho \right] |\psi_v\rangle, \quad (5)$$

$$|P \eta^{(\lambda,\mu)}_v\rangle = \hat{G}_v \left\{ \frac{\partial^2 V^{\text{KS}}}{\partial \lambda \partial \mu} \right\} + \left[ \frac{\partial V^{\text{KS}}}{\partial \lambda} \partial \rho / \partial \mu \right] + \left[ \frac{\partial V^{\text{KS}}}{\partial \mu} \partial \rho / \partial \lambda \right] |\psi_v\rangle. \quad (6)$$

Here,

$$\hat{G}_v = \sum_c \frac{|\psi_c\rangle \langle \psi_v|}{\epsilon_v - \epsilon_c}$$

The above commutator is well defined, bounded operators, since the density matrix is much more efficient and the time for RI calculation is negligible compared to that for $\omega_{\nu}$ in large systems.

TABLE I: Raman activity in Si computed with our approach ($\gamma_{\text{SOR}}$), and by finite differences ($\gamma_{\text{FD}}$). $N$ is the number of inequivalent k-points.

| $N$ | 2 | 10 | 28 | 60 | 110 | 182 |
|-----|---|----|----|----|-----|-----|
| $\gamma_{\text{SOR}}$ | 8.54 | 5.30 | 5.32 | 5.39 | 5.40 | 5.40 |
| $\gamma_{\text{FD}}$ | 18.99 | 7.09 | 5.69 | 5.45 | 5.41 | 5.40 |

is the Green function operator projected on the empty states $|\psi_c\rangle$, $[A, B] = AB - BA$, and the first derivative of the density matrix is:

$$\frac{\partial \rho}{\partial \mu} = \sum_v |P |P \partial \psi_v/\partial \mu \rangle |\psi_v| + cc. \quad (7)$$

Since $\partial V^{\text{KS}}/\partial \lambda$ and $\partial^2 V^{\text{KS}}/\partial \lambda \partial \mu$ depend on $\partial \rho/\partial \lambda$, $\partial \rho/\partial \mu$, and $\partial^2 \rho/\partial \lambda \partial \mu$, Eqs. (4-7), should be solved self-consistently.

The advantage of the present formulation, compared to that of Ref. [3], lies in the introduction of the commutators of Eqs. (4-7). Thanks to the commutators, all the quantities needed with our formalism are well defined in an extended insulator, even if the perturbation $\mu$ or $\lambda$ are the component $E_l$ of a uniform electric field, i.e., if $\partial V^{\text{KS}}/\partial \lambda = -e r_l + \partial V^{\text{Hxc}}/\partial E_l$, being $r_l$ the $l$th Cartesian component of the position operator $r$, and $e$ the electron charge. In particular, in an insulator, the commutators $[r, \rho]$ and $[r, \partial \rho/\partial \mu]$ in Eqs. (4-7) are well defined, bounded operators, since the density matrix is localized ($\langle r'|\rho|r'\rangle$ goes to zero exponentially for $|r' - r'| \to \infty$).

Finally, in a periodic system, the right-hand side of Eq. (6) can be easily computed in terms of $|\psi_k\rangle$, that are the periodic parts of the Bloch-wavefunctions $|\psi_k\rangle$ with reciprocal-lattice vector $k$, using the substitutions:

$$\langle \psi_k | r, \rho | \psi_k \rangle = i \sum_{v'} \langle u_{k'} | P_{k} \partial \rho/\partial E_m \rangle | u_{k'} | u_{k'} \rangle \quad (8)$$

$$\langle \psi_k | r, \partial \rho/\partial E_m | \psi_k \rangle = i \sum_{v'} \langle u_{k'} | P_{k} \partial \rho/\partial E_m \rangle | u_{k'} | u_{k'} \rangle \quad (9)$$

where $l$ and $m$ are Cartesian indexes, $c$ is an empty band index, $v$ and $v'$ are occupied band indexes, and $P_k$ is the projector on the empty subspace of the point $k$. In our implementation, the derivative with respect to $k_l$ in the right-hand side of Eq. (6) is computed numerically by finite-differences, using an expression independent from the arbitrary wavefunction-phase, as in Refs. [3, 5, 6].

We test our approach on Si in the diamond phase, where the Raman activity is determined by $\gamma = a \partial \epsilon_{111}^{\text{cm}}/\partial u$, where $a = 10.20$ $a.u.$ is the lattice spacing and $u$ the displacement of one atom along the $(1, 1, 1)$ direction. We compute $\gamma$ for various grids of $k$-points, using both our second order response method ($\gamma_{\text{SOR}}$) and by finite
In this class of materials, that includes the all-silica zeolites, the quartz, cristobalite, tridymite and coesite polymorphs of SiO$_2$, and vitreous silica (v-SiO$_2$), each Si atom is tetrahedrally coordinated to four O atoms and each O atom is bonded to two Si atoms. The properties of these systems can be effectively described in terms of the n-membered rings (n-MRs) of tetrahedra contained in their structure. E.g., a clear correlation between the presence of 3- and 4-MRs and the degradation of optical v-SiO$_2$ fibers under UV radiation has been observed. In the v-SiO$_2$ Raman spectra the two sharp peaks at 490 cm$^{-1}$ ($D_1$ line) and 606 cm$^{-1}$ ($D_2$ line), have been attributed to the breathing mode (BM) of the O atoms towards the ring center of 4-MRs and 3-MRs, respectively. This attribution has been confirmed by DFT vibrational frequency calculations. The attribution would be further supported by experimental measurements on well characterized crystalline polymorphs containing 3- and 4-MRs. However, the strong Raman peak at 520 cm$^{-1}$ in coesite, a phase that contains 4-MRs, is shifted by 30 cm$^{-1}$ with respect to the $D_1$ line in v-SiO$_2$, and no Raman measurements has been published on the H-ZSM-18 zeolite, that is the only known SiO$_2$ crystalline polymorph with 3-MRs. Interestingly this zeolite contains 4-MRs as well.

To clarify this topic, we compute the Raman spectra of α-quartz, coesite, α-cristobalite, and H-ZSM-18. In Fig. 1 we compare our results with the available experimental spectra. The vibrational frequencies are systematically underestimated by 5% by our calculation. To simplify the comparison with the experiments, in Fig. 1, the theoretical frequencies are multiplied by a scaling factor of 1.05. The ability of the method in reproducing quantitatively all the measured features is evident.

In order to associate Raman peaks of Fig. 1 to the small-ring BMs, we project the vibrational eigenmode $\mathbf{w}^\nu$ on the subspace generated by the BMs of a given kind of rings, $\mathcal{R}$, and on the corresponding complementary subspace, $\mathcal{R}^c$. We use the two resulting projected vectors to decompose $\hat{A}^\nu$ so that $\hat{A}^\nu = \hat{A}^\nu_{\mathcal{R}} + \hat{A}^\nu_{\mathcal{R}^c}$. Since $I^\nu$ is quadratic in $\hat{A}^\nu$, see Eq. (1), $I^\nu = I^\nu_{\mathcal{R}} + I^\nu_{\mathcal{R}^c} + I^\nu_{\text{overlap}}$, where $I^\nu_{\text{overlap}}$ is the term bilinear in $\hat{A}^\nu_{\mathcal{R}}$ and $\hat{A}^\nu_{\mathcal{R}^c}$. A Raman peak can be associated to a ring BM (i.e. the Raman activity is solely due to the BM) if, and only if, $I^\nu_{\mathcal{R}} \gg |I^\nu_{\text{overlap}}|$.

The structure of H-ZSM-18 contains two equivalent 3-MRs and two kinds of 4-MRs which we will call 4-MRs$_0$, and 4-MRs$_1$. In Fig. 2, we show the projected Raman spectra of the zeolite and the coesite. In the H-ZSM-18 spectrum, the peaks at 485, and 615 cm$^{-1}$ are very well described by the BM of 4-MRs$_0$ and 3-MRs, respectively. A direct analysis of the vibrational eigenmodes shows that both BMs are decoupled from other modes. The frequencies of the two peaks are very close to those of the measured $D_1$ and $D_2$ lines in v-SiO$_2$ (490, and 606 cm$^{-1}$), thus confirming that these lines are due to rings BMs. However, the presence of small-MRs in a structure does not guarantee, in general, the occurrence of completely decoupled BMs. This is the case of the 4-MRs in coesite and the 4-MRs$_1$ in the zeolite, whose BMs exhibit a large $|I^\nu_{\text{overlap}}|$, see Fig. 2. These over-

![FIG. 1: Vibrational Raman spectra of various SiO$_2$ polymorph powders. Measurements are from Refs. 22. Theoretical frequencies are rescaled by +5%, and the spectra are convoluted with a uniform Gaussian broadening having 4.0 cm$^{-1}$ width.](image-url)
lapses imply the existence of a coupling with other modes, that, in turn, explains the 30 cm⁻¹ difference between the 4-MRs frequency of coesite and that of the 4-MRs in the zeolite. A comparable frequency shift from the 4-MRs line of vitreous silica is observed, with opposite sign, for the 4-MRs in the zeolite.

In conclusion, with the aim of building an instrument for the routine interpretation of Raman spectra, we developed a method for the efficient calculation of Raman intensity. We computed the Raman spectra of SiO₂ polymorphs containing up to 102 atoms. We found that: i) not all the small-membered-rings have decoupled breathing modes, ii) the H-ZSM-18 zeolite provides decoupled breathing mode of 4- and 3-membered rings, whose frequencies nicely coincide with the D₁ and D₂ lines of vitreous silica. An experimental determination of the Raman spectra of this zeolite can thus provide an experimental calibration for the determination of the density of decoupled small membered rings in vitreous silica.

Calculations were performed at IDRIS supercomputing center. Our approach was implemented in the PWSCF code [24].

FIG. 2: Raman intensities projected on the breathing modes of various rings labeled 3-MRs, and 4-MRs. For clarity, the overlap intensity (I' \text{overlap} in the text) is shifted vertically.

\[ I \text{ of various rings labeled 3-MRs, and 4-MRs} \]

\[ \text{v-SiO}_2 \]

\[ D \]

\[ \text{the 4-MRs frequency of coesite and that of the} \]

\[ \text{lapses imply the existence of a coupling with other modes,} \]

\[ \text{that, in turn, explains the} 30 \text{ cm}^{-1} \text{ difference between the} 4-\text{MRs frequency of coesite and that of the} D_1 \text{ line of} \]

\[ \text{v-SiO}_2 \text{. A comparable frequency shift from the} D_1 \text{ line is observed, with opposite sign, for the} 4-\text{MRs}_1 \text{ BMs in the} \]

\[ \text{zeolite.} \]

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