Optical to UV spectra and birefringence of SiO\textsubscript{2} and TiO\textsubscript{2}: First-principles calculations with excitonic effects

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Abstract

A first principles approach is presented for calculations of optical – ultraviolet (UV) spectra including excitonic effects. The approach is based on Bethe-Salpeter equation calculations using the NBSE code combined with ground-state density-functional theory calculations from the electronic structure code ABINIT. Test calculations for bulk Si are presented, and the approach is illustrated with calculations of the optical spectra and birefringence of \(\alpha\)-phase SiO\textsubscript{2} and the rutile and anatase phases of TiO\textsubscript{2}. An interpretation of the strong birefringence in TiO\textsubscript{2} is presented.

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I. INTRODUCTION

As in many subfields in condensed matter physics, the application of density-functional theory (DFT)\textsuperscript{12,13} has significantly influenced studies of material optical properties. However, treatments based on Kohn-Sham dipole transitions alone only provide a beginning for modeling optical spectra, since a ground-state one-electron approach cannot account for a range of excited-state and optical effects. Thus quantitative calculations of optical spectra require the integration of DFT with a number of many-body techniques. Local field corrections from the random phase approximation (RPA) alone are inadequate to reproduce near-gap optical spectra, because they do not take into account self-energy and excitonic effects. Consequently a quantitative theory and interpretation must also include quasi-particle self-energy effects based e.g., on the $GW$-method\textsuperscript{4,5} and particle-hole interactions based on the Bethe-Salpeter equation (BSE)\textsuperscript{6} The success of the BSE method has effectively settled the long-running discussion of the origin of many low energy spectral features\textsuperscript{7,8} For example, for the rutile phase of TiO$_2$, early studies could only speculate about the relative importance of excitonic effects versus band structure topology in the dominant threshold absorption\textsuperscript{9} whereas we show here that these features are dominated by excitonic effects.

While the BSE is often necessary for accurate calculations of optical spectra, the approach has had limited applications, largely due to a lack of efficient computational methods for general systems. To address this need, we have developed an approach based on an interface, here dubbed AI2NBSE, between the BSE code NBSE developed at National Institute of Standards and Technology (NIST)\textsuperscript{10} and the general purpose DFT electronic structure code ABINIT\textsuperscript{11} One of the objectives of this work is to improve the availability of BSE codes. Our interface also provides a comparison to the BSE codes EXC\textsuperscript{12}, and EXCITING\textsuperscript{13} and other recent codes\textsuperscript{14} and also to approaches which model electron-hole interactions within time-dependent density-functional theory (TDDFT)\textsuperscript{15} The latter calculations are usually simpler than BSE ones, but in practice are limited by an incomplete knowledge of the exchange-correlation functional and the neglect of damping effects. The overall strategy of our interface differs from these approaches in several respects. For example our AI2NBSE interface achieves efficiency in the BSE calculations through the use of the Hybertsen-Levine-Louie\textsuperscript{16,17} dielectric screening, and can also treat finite momentum transfer. Also, the interface requires only generic input, and thus can be adapted to other
The interface is tested on bulk Si, yielding results in good agreement with other approaches. In particular we find that the calculations based on ABINIT are in excellent agreement with those from the optimal basis function (OBF) code. As initial applications, we report optical spectra and anisotropic optical properties of the common rutile and anatase phases of TiO$_2$, and for the $\alpha$-phase of SiO$_2$. These are important materials for many applications. However, we are not aware of earlier calculations for TiO$_2$ that include electron-hole interactions, which are needed to reproduce their rich dependence on polarization and phase.

In the remainder of this paper we briefly summarize the key formulas describing excitonic effects and other features of NBSE. We then briefly describe our integration of ABINIT and NBSE, using bulk Si as an illustrative test case, with some further details in an Appendix. Finally, calculations are presented for the birefringent spectra of the $\alpha$-phase SiO$_2$, and similarly for the rutile and anatase phases of TiO$_2$, followed by a summary and conclusions.

II. BSE AND EXCITONIC EFFECTS

A. The Exciton Secular Equation

In this paper we only briefly summarize the BSE formalism, following the notation and theoretical developments of Shirley et al. Formally the BSE provides a complete theory for optical spectra through a hierarchy of equations derived from the two-particle Green’s function. However with certain approximations to the electron-hole interaction, the BSE can be reduced to an eigenvalue problem of an effective particle-hole Hamiltonian:

$$H|f\rangle = [H_{1e} + H_{2e}]|f\rangle = \Omega_f|f\rangle,$$

where the eigenstates $|f\rangle$ are given by a superposition of particle-hole basis states $|nn'k(q)\rangle$, i.e.,

$$|f\rangle = \sum_{nn'k} \psi_f(nn'k(q)) |nn'k(q)\rangle$$

and throughout this paper we use atomic units ($e=\hbar=m=1$). Here $\psi_f(nn'k(q))$ is the amplitude of a given particle-hole (or excitonic) state with Bloch crystal momentum index $q$. The operators $a_{nk}$ and $a_{nk}^\dagger$ act on bare particle states $|0\rangle$. 

$$|nn'k(q)\rangle \equiv a_{nk}^\dagger a_{n'k+q}|0\rangle.$$
and momentum transfer $q$, the index $n$ runs over all occupied valence bands, the index $n'$ runs over unoccupied bands, and $|0\rangle$ denotes the many-body ground-state with energy $\Omega_0 = 0$. For optical absorption $q$ is usually negligible. However for inelastic x-ray scattering and other spectroscopies and for computational purposes, it is desirable to retain the explicit momentum-transfer dependence. The single-particle contribution to the Hamiltonian for a particle-hole pair is diagonal in the quasi-particle/hole basis, so that one has

$$H_{1e}|nn'k(q)\rangle = (E_{n'k+q} - E_{nk})|nn'k(q)\rangle,$$

where the quasi-particle energies $E_{nk}$ are Kohn-Sham eigenvalues $\varepsilon_{nk}$ plus quasiparticle self-energy corrections

$$E_{nk} = \varepsilon_{nk} + \Sigma_{nk}.$$  

Here $\Sigma_{nk}$ is the self-energy calculated in the $GW$ approximation for which efficient approximations have been developed.\textsuperscript{20,25}

The electron-hole interaction contribution includes both “direct” and “exchange” couplings $H_{2e} = V_D + V_X$,

$$H_{2e}|nn'k(q)\rangle = \sum_{n''n'''} [V_D(nn'k, n''n'''k'; q) + V_X(nn'k, n''n'''k'; q)] |n''n'''k'(q)\rangle,$$

with matrix elements defined explicitly below. Once the BSE secular equation is solved, the optical constants may be obtained formally using a Fermi golden rule expression in terms of the excitonic final states coupled to the current-like operator $J'_\mu$. In the NBSE code, however, these properties are calculated using resolvent techniques. In particular the imaginary part of the dielectric tensor is given in terms of resolvents

$$\text{Im} \epsilon_{\mu\nu}(\omega) = -4\pi \text{Im} \left[ \langle 0|J'_\mu[\omega - H + i\eta]^{-1}J'_\nu|0\rangle - \langle 0|J'_\nu[-\omega - H - i\eta]^{-1}J'_\mu|0\rangle \right].$$

In terms of the particle-hole states, the current-like operator coupling to the ground-state is given approximately by

$$J'_\mu|0\rangle \approx \sum_{nn'k} |nn'k(q)\rangle \frac{\langle \psi_{n'k+q}|J_\mu|\psi_{nk}\rangle}{\varepsilon_{n'k+q} - \varepsilon_{nk}}.$$  


where $J_\mu$ is the $\mu$-th component of the current operator and $\psi_{nk}$ are approximated as Kohn-Sham one-particle states. For small $q$, the matrix elements of $J_\mu$ are approximated by

$$\frac{1}{\omega} \langle \psi_{n'k+q} | J_\mu | \psi_{nk} \rangle \approx \left( \frac{\epsilon_{n'k+q} - \epsilon_{nk}}{\omega q_\mu} \right) \langle \psi_{n'k+q} | e^{i\mathbf{q} \cdot \mathbf{r}} | \psi_{nk} \rangle$$

$$\approx \frac{1}{q_\mu} \langle \psi_{n'k+q} | e^{i\mathbf{q} \cdot \mathbf{r}} | \psi_{nk} \rangle.$$  \hspace{1cm} (8)

where $\mathbf{q} = q\hat{\mu}$. In the NBSE code these resolvents in the above expressions for the dielectric tensor are calculated using an efficient iterative Lanczos algorithm.  

B. Electron-hole Interaction

The interaction kernel $H_{2e}$ of Eq. (5) accounts for two processes which scatter an electron from band $n'$ to band $n'''$, and a hole from band $n$ to band $n''$. The first is the attractive direct screened Coulomb interaction between the electron and the hole, and the second is the repulsive unscreened exchange interaction. Each of these contributions can be written in terms of two-particle integrals between electron and hole Kohn-Sham orbitals $\psi_{nk}(\mathbf{x})$ and $\psi_{n'k+q}(\mathbf{x})$. Matrix elements of the direct term are given by

$$V_D(nn'k, n''n'''k'; \mathbf{q}) \approx - \int d^3 x \, \psi_{n'''k'+q}(x) \, \psi_{n'k+q}(x)$$

$$\times \int d^3 x' \, \psi_{nk}^*(x') \, \psi_{n''k'}(x') \, W(x, x'; \omega = 0).$$  \hspace{1cm} (9)

Here the electron-hole interaction $W(x, x'; \omega)$ is the screened Coulomb attraction as mentioned above, i.e., $W = \epsilon^{-1}(\omega)V$, which in an exact theory should include the frequency dependence of the dielectric response. However the NBSE code approximates the screening with the static, spatially dependent Hybertsen-Levine-Louie dielectric function. This model maps the local density in real solids to the density dependence in a homogeneous system. These calculations require the ground-state density, whose Fourier coefficients are

$$\rho(\mathbf{G}) = \frac{2}{N} \sum_{nk, G'} f_{nk} \psi_{nk}(\mathbf{G} + \mathbf{G'}) \psi_{nk}^*(\mathbf{G'}),$$

where $f_{nk} = \theta(\epsilon_F - \epsilon_{nk})$ is the occupation of state $nk$, $N$ is the number of unit cells in the ABINIT calculation, which is the same as the number of $k$ points in the calculation,
and spin degeneracy is assumed. Here $\psi_{nk}(G)$ are the Fourier coefficients of the Bloch wave expansion\cite{footnote17} in reciprocal lattice vectors

$$\psi_{nk}(x) = e^{i k \cdot x} \sum_G \psi_{nk}(G) e^{i G \cdot x}. \quad (11)$$

Because the dielectric function is modeled locally, the exciton amplitudes must be transformed from the Bloch basis to a local basis with coordinates $x$ and $x'$. These local coordinates can be considered as transform analogs of the band and wave-vector indices $n$, $n'$, and $k$\cite{footnote27}.

Similarly matrix elements of the exchange term are

$$V_X(nn'k,n''nk';q) \approx +2 \int d^3x \psi_{n''k'+q}^*(x) \psi_{n'k'}(x) \times \int d^3x' \psi_{nk}(x') \psi_{n'k+q}(x') |x - x'|. \quad (12)$$

As noted, for example by Hybertsen and Louie, the exchange term in the BSE should be unscreened.

The matrix dimension of the electron-hole interaction $H_{2e}$ is generally very large. For example, for the calculations for SiO$_2$ presented below, 216 $k$ points are sampled, and there are 24 doubly degenerate valence (hole) states and 26 conduction (electron) states. For this case, there are over $10^5$ electron-hole pairs, each requiring a representation in both periodic and local bases.

C. AI2NBSE Interface

The above theory has been implemented in a modular code which uses the output of ground-state density-functional theory calculations from the electronic structure code ABINIT as input to Bethe-Salpeter equation calculations using the NBSE code. The interface serves as a driver for both ABINIT and NBSE, starting from a single input file, and also constructs the various physical quantities and arrays needed in the calculations. No explicit changes in the structure or coding of either ABINIT or NBSE are used. Both the interface and documentation are available from the authors\cite{footnote28}. Additional details are given in the Appendix.
III. OPTICAL SPECTRA OF SILICON

As a quantitative test, illustrative results from AI2NBSE for bulk Si are presented in Fig. 1. The two spectra compared are each calculations of the imaginary part of the dielectric function for bulk Si using NBSE: in one case the ground-state quantities above are calculated with the optimal basis function (OBF) pseudopotential, plane-wave code – for which NBSE was originally designed – and in the other case the same quantities are calculated from ABINIT. In both calculations twenty valence and conduction bands were included with an eight Hartree cutoff criteria for the vectors $\mathbf{G}$. The close quantitative agreement between OBF and ABINIT thus serve as a quality check on the various theoretical and algorithmic approximations used in our interface.

Our AI2NBSE interface can also be applied to finite momentum transfer calculations. For example, Fig. 2 illustrates the spectrum for a momentum transfer of magnitude $q = 0.8 a_0^{-1}$, where $a_0$ is the Bohr radius. Results from a recent TDDFT calculation are also plotted.

IV. OPTICAL SPECTRA OF $\alpha$-QUARTZ

The excitonic character of the quartz optical spectra had been recognized even before first-principles investigations became possible. The large band gap and small static dielectric
FIG. 2: Calculated $\epsilon_2(q, \omega)$ for Si with impulse magnitude of $q = 0.8 \ a_0^{-1}$ along the [1, 1, 1] direction. AI2NBSE’s result (solid line), and for comparison the result from a recent TDDFT calculation (dashed line) are plotted.\(^{29}\)

constant of $\alpha$-quartz–and the prominent low-energy features in the spectrum–suggested a strong excitonic role in the optical properties, which was subsequently confirmed by first-principles calculations.\(^{24}\) Our calculations, like those of Ref. 24, reproduce the measured spectra in detail, with excellent agreement in oscillator strength and calculated feature positions corresponding with those measured to within tenths of an eV. The results can be seen in Fig. 3.

The calculations of Ref. 24 were performed for photon polarization within the hexagonal plane (\textit{i.e.}, the “ordinary ray”), noting that this is the most commonly measured spectrum. This material is however uniaxial, possessing an independent optic axis normal to the hexagonal plane. We have also carried out calculations for photon polarization perpendicular to the hexagonal plane (\textit{i.e.}, the “extraordinary ray”). As illustrated in Fig. 3, quartz exhibits some anisotropy, but it is less pronounced than for rutile. This result is consistent with the known role of the SiO\(_4\) tetrahedron versus the anisotropic crystal structure, and the structural phase insensitivity of the optical functions in quartz.\(^{30}\)

While birefringence in quartz is a well-studied effect, it is small relative to TiO\(_2\) crystals discussed in the the next section. Below the interband transition energies, but above the lattice response, we find static indices of refraction of 1.52 for the ordinary polarization, and 1.53 for the extraordinary at 0.7 eV, in precise agreement with the measured values.\(^{32}\)
FIG. 3: Imaginary part of the dielectric function for SiO$_2$ with in-plane polarization (top), and out-of-plane polarization (bottom). Theoretical spectra from ABINIT and OBF (solid line and dashed line) spectra are plotted for both polarizations. The calculated spectra use a 28 Hartree plane-wave cutoff and included a 216-point Brillouin zone sampling and the 26 lowest-lying, degenerate bands above the Fermi level. The experimental spectrum, reproduced from Ref. 24 is also shown (crosses) for the ordinary ray.

V. OPTICAL SPECTRA OF RUTILE AND ANATASE

As one of the simplest transition-metal oxides, TiO$_2$ exhibits a variety of natural crystal structures and presents a fundamentally interesting system for first-principles electronic structure methods. This material is also an important component in various ultraviolet (UV) applications. It has been demonstrated that *ab initio* methods can describe various physical properties of TiO$_2$.$^{18,19}$ There have been a few *ab initio* studies of UV dielectric spectra for rutile and anatase phases. Glassford and Chelikowsky$^{33}$ reported calculations for the rutile phase using a plane-wave pseudopotential approach; Mo and Ching$^{34}$ used a linear-combination-of-atomic orbital method for both phases (and brookite); and Asahi *et al.*$^{35}$
studied the anatase phase with a linearized augmented plane-wave method. However, none
of these studies included excitonic effects. A theoretical treatment that includes excitonic
states may contribute to understanding its optical properties. Toward this end, we present
calculations of the spectra of TiO$_2$, with polarization dependence, for both the rutile and
anatase phases using our AI2NBSE interface.

Each of the two TiO$_2$ phases is tetragonal, and Mo and Ching$^{34}$ and Fahmi et al.$^{36}$
have reviewed their structural properties including crystal structures, space groups, and
differences in bond lengths and angles. These two crystal structures can be considered as
arrangements of slightly distorted oxygen octahedral elements with a titanium atom at the
center of each, so that each titanium has an oxygen coordination of six, and each oxygen
has a titanium coordination of three. The relationship between the two structures has been
described in terms of varied orientation among the octahedral chains. The two polymorphs
studied here can be generated with six-atom unit cells corresponding to two TiO$_2$ units.
Each unit possesses two inequivalent bonds of “apical” and “equatorial” character, such
that each titanium sees two apical and four equatorial bonds, while and each oxygen sees
one apical and two equatorial bonds.

Structural similarities in rutile and anatase lead to similarities in their electronic struc-
ture. In Fig. 4 we plot the ABINIT-calculated densities of states for for the two materials.
The local-density approximation (LDA) gives the uppermost valence-band width of about
5 eV for anatase and 6 eV for rutile, and this band is regarded to be dominated by O
$2p$ character. The anatase LDA band gap is greater than that of rutile by about 0.2 eV,
corresponding with the measured difference.$^{37}$ Because the oxygen bonding environment is
planar in both phases, there is a decomposition of the O $2p$ into $p_\pi$ and $p_\sigma$ states.$^{35,38}$

The first conduction band is dominated by Ti $3d$ character, with the lower half regarded
as $t_{2g}$-like, and the upper half regarded as $e_g$-like.$^{35,38}$ As can be seen in Fig. 4 and from
the band structures,$^{18}$ these sub-bands are reasonably resolved energetically. Our calcu-
lations were performed with semicore states treated as core using Troullier-Martins-type
pseudopotentials$^{33,39}$ and as valence using Teter-type pseudopotentials.$^{18,40}$ While the semi-
core states have spatial localizations comparable to oxygen $2p$ states, their energies are many
eV below the Fermi level. Our calculations suggest that the optical and UV spectra are not
highly sensitive to the treatment of Ti semicore states, and that the primary source of the
discrepancy between theoretical and experimental spectra is the neglect of the excitonic
FIG. 4: Density of states $\rho(E)$ with respect to the Fermi energy $E_F$ for the uppermost valence and lowest conduction bands of TiO$_2$ in the rutile (solid line) and anatase (dashed line) phases. No band gap corrections are included. The LDA gap is calculated to be 2.0 eV for rutile and 2.2 eV for anatase.

effects in the low-energy region.

A. Rutile Phase

The spectra calculated for rutile with the AI2NBSE interface (Fig. 5) demonstrate significant excitonic effects, as can be seen by comparison with calculations (Fig. 6) which neglect electron-hole interactions and are calculated within the RPA. One signature of the excitonic effects is the sharp onset in our BSE calculations, which better reproduces experimental spectra. Our results for the polarization dependence in TiO$_2$ show a much more prominent anisotropic optical response than in SiO$_2$. This results in strong birefringence, as seen in Fig. 7. Our calculated static indices of refraction differ by 0.30, in agreement with the observed birefringence.\textsuperscript{9}

Two pronounced anisotropic features in our calculations are consistent with experimental spectra.\textsuperscript{9,41} First, the primary onset absorption feature assumes a doublet structure for the ordinary ray which is not evident for the extraordinary ray. These low-energy excitonic features at 4.0 and 5.3 eV are clearly evident in our calculation and are expected to involve $t_{2g}$-like final states.\textsuperscript{34} Second, the onset of absorption for the extraordinary ray is significantly stronger than for the ordinary. For the extraordinary ray, a single feature is measured at 4.1 eV, which is stronger, broader, and more symmetric than for the ordinary ray. However, a
FIG. 5: Imaginary part of the dielectric function for rutile with in-plane polarization (top) and out-of-plane polarization (bottom). Theoretical from ABINIT and OBF (solid line and dashed line) spectra are plotted for both polarizations. The calculated spectra use a 32 Hartree plane-wave cutoff and included a 216-point Brillouin zone sampling and the 26 lowest-lying, degenerate bands above the Fermi level. For comparison the experimental spectrum from Ref. 9 (crosses) is also plotted.

The third anisotropic feature of our calculations, namely the much stronger absorption peak for the extraordinary ray above 6 eV, is not clearly seen in the experimental reflectivity spectra. It should be noted that the peak at 7.4 eV is more prominent in the x-ray absorption spectra than in the reflectivity data, and is expected to involve $e_g$-like conduction states. To summarize, our BSE calculation gives a stronger, less structured $t_{2g}$-like absorption band for the extraordinary ray than for the ordinary, in accord with experiment.
FIG. 6: Imaginary part of the dielectric function for rutile with in-plane polarization (top) and out-of-plane polarization (bottom) calculated without core-hole interactions ($V_D = 0$). Two spectra are plotted corresponding to a treatment of semicore states with Teter-type Ti pseudopotentials (solid line), and with Troullier-Martins pseudopotentials where the semicore states are pseudized (dashed line).

B. Anatase Phase

Theoretically, both the valence-band and conduction-band densities of states are similar to that of rutile, as shown in Fig. 4. However, their experimental absorption spectra show more variation (Fig. 8.). Strikingly, the measured spectrum of anatase for the ordinary ray (in-plane polarized) shows a clear low-energy shoulder below the primary absorption feature.\textsuperscript{41,43} This feature is not seen in rutile, and calculations neglecting the electron-hole interactions in anatase do not clearly resolve it.\textsuperscript{34,35} Aside from this low-energy feature, the spectral structure and anisotropy of anatase, both calculated and measured, have some similarities to that of rutile. Thus features in the spectra have been analyzed\textsuperscript{35,43} and conclusions regarding the relevant bands are reminiscent of results for rutile.\textsuperscript{33} For the $e_g$-like
FIG. 7: Ordinary (dashed line) and extraordinary (solid line) indices of refraction $n$ of rutile-TiO$_2$ as a function of photon energy. At 589 nm, which is below the band gap but well above the lattice resonances, the calculated difference between the two indices is 0.30 while the measured value is 0.29.

absorption band, the experimental spectrum for anatase does show a stronger absorption for the extraordinary ray, similar to what is calculated in both the rutile and anatase phases, but not seen in experimental rutile spectrum.

C. Interpretation of Polarization Dependence

Molecular orbital interpretations of the TiO$_2$ spectra have been reported widely and provide a qualitative picture of the electronic structure in this system. Such analyses illustrate the hybridization between the Ti 3$d$ and O 2$p$ states. However, immediately above and below the Fermi level, the hybridization has been shown to be weak. The calculations of Sorantin and Schwartz demonstrate that the valence band for rutile just below the Fermi level is primarily $\pi$-bonding O 2$p$ character, while lower in energy is primarily $\sigma$-bonding O 2$p$ character. Just above the Fermi level, the states are, in energetic order, primarily Ti $t_{2g}$ and $e_g$.

A molecular-orbital analysis, considered in conjunction with measured x-ray spectra, allowed an empirical identification of features in the optical spectra of Harbeke and Cardona with specific electronic transitions. Although selection rules, inferred from the approximate atomic states, were invoked in that work, no effort was made to analyze the polarization dependence of the selection rules. Some of the most prominent features could not be identified
FIG. 8: Imaginary part of the dielectric function for anatase with in-plane polarization (top) and out-of-plane polarization (bottom). Theoretical spectra from ABINIT and OBF (solid and dashed lines) are plotted for both polarizations. The calculated spectra use a 32 Hartree plane-wave cutoff and include a 216-point Brillouin zone sampling and the 26 lowest-lying, degenerate bands above the Fermi level. For comparison the experimental spectrum reproduced from Ref. 9 (crosses) also plotted.

at all, and were attributed to the unaccounted excitonic effects. Assignments of the same experimental features were made, for ordinary and extraordinary rays independently, from a band-structure perspective by Glassford and Chelikowsky.33 But the calculated spectra again did not treat the excitonic interactions, and hence did not reproduce the experimental features to the level of agreement reported here. Also, local electronic structure was not emphasized in the peak assignment analysis.

Perhaps the most conspicuous anisotropic characteristic in the rutile spectra is the stronger onset absorption for the extraordinary ray. We observe that the first eV below the Fermi level is dominated by the O 2pπ states, and further, that the oxygen bonding
planes are all defined by in-hexagonal-plane normals. This implies that, for some in-plane polarization, there is a single dipole allowed channel from one of the two $\pi$-state orientations (corresponding to the two oxygen-bonding planes), and the three $t_{2g}$ states, while for the extraordinary polarization, there are two dipole-allowed channels. This is consistent with the calculated and measured stronger threshold resonance for the extraordinary ray. In connection with this point, we emphasize that while the $t_{2g}$ and $e_g$ symmetry labels are approximate because of angular and bond-length distortions within the octahedra, the selection rules above are unmodified from the above even when the local, $D_{2h}$ symmetry group is considered, while the actual crystallographic group in rutile is of even higher symmetry.

VI. SUMMARY AND CONCLUSIONS

In summary, we have developed a first principles approach for calculations of various optical spectra, including finite momentum transfer in crystals. The method combines ground state electronic structure calculations from ABINIT with BSE calculations from the NBSE code. The method is tested on bulk Si, yielding results in good agreement with other methods. Calculations are presented for the macroscopic dielectric spectra and its orientation dependence in $\alpha$-quartz, rutile TiO$_2$, and anatase TiO$_2$. Our quartz spectra for the ordinary ray reproduce the strong excitonic character and are in good agreement with experiment and earlier theoretical work. The anisotropy of the rutile and anatase phases of TiO$_2$ are more significant than for quartz. The static indices of refraction for the two polarizations of rutile differ by more than 10%, in agreement with experiment. Also our calculated absorption at low energies reproduces experiment better than previous theoretical results that neglect excitonic effects. In particular our calculations reveal an additional low-energy feature in anatase also found in experiment. Also we are able to interpret the anisotropy in the threshold behavior for rutile in terms of $\pi$ to $t_{2g}$ selection rules.

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APPENDIX: AI2NBSE INTERFACE

In order to calculate the optical spectra, the AI2NBSE interface first obtains Kohn-Sham energies and wave functions from the self-consistent ground-state electronic structure code ABINIT. The single-particle eigenenergies are modified to include self-energy corrections.
according to the $GW$ approximation. Subsequently the interface constructs several quantities needed for the NBSE calculations. These include current–operator matrix elements between Kohn-Sham states and the ground-state charge density for calculating Hybertsen-Levine-Louie screening. Thus our interface does not take advantage of ABINIT’s dielectric function capability.

Typical AI2NBSE calculations are divided into four modular stages and require a single input file which contains all parameters needed to define both the system and its ground-state and excited state one-electron properties. Briefly the modular operations are as follows after the input file is read and stored:

1) ABINIT calculation: ABINIT inputs are generated and ABINIT is run. These calculations supply the Kohn-Sham eigenvalues $\varepsilon_{nk}$ and eigenfunctions $\psi_{nk}$ both for the occupied and unoccupied states. Currently the interface uses a standard serial distribution of ABINIT. However, calculations for large systems can still be performed by means of consecutive runs which are automatically set up by the interface, with minimal input or intervention from a user.

2) Density components: Fourier components of the ground-state density $\rho(G)$ are generated using the eigenfunctions from stage 1.

3) Dipole matrix elements: Dipole matrix elements in Eq. (8) are calculated, and the eigenfunctions are converted to the format used by NBSE.

4) NBSE calculation: All quantities required for the Bethe-Salpeter calculation are collected and NBSE is run. The output includes various optical spectra and optical constants.