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Ab initio description of quasiparticle band structures and optical near-edge absorption of transparent conducting oxides

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Many-body perturbation theory is applied to compute the quasiparticle electronic structures and the optical absorption spectra (including excitonic effects) for several transparent conducting oxides (TCOs). We discuss $HSE+G_0W_0$ results (based on the hybrid exchange-correlation functional by Heyd, Scuseria, and Ernzerhof, and quasiparticle corrections from approximating the electronic self energy as the product of the Green's function and the screened Coulomb interaction) for band structures, fundamental band gaps, and effective electron masses of magnesium oxide, zinc oxide, cadmium oxide, tin dioxide, tin oxide, indium (III) oxide and silicon dioxide. The Bethe–Salpeter equation (BSE) is solved to account for excitonic effects in the calculation of the frequency-dependent absorption coefficients. We show that the $HSE+G_0W_0$ approach and the solution of the BSE are very well suited to describe the electronic structure and the optical properties of various TCOs in good agreement with experiment.

I. INTRODUCTION

Transparent conducting oxides (TCOs) combine high transparency in the visible spectral range with high electrical conductivity under ambient conditions.^{1,2} Posttransition-metal compounds such as zinc oxide (ZnO), indium (III) oxide (In_2O_3) , and tin dioxide (SnO_2) are typical TCOs as they have large fundamental band gaps rendering these materials transparent into the ultraviolet (UV) spectral range. Due to their very large gaps, especially magnesium oxide (MgO) and silicon dioxide (SiO₂) are transparent into the far UV. The gaps can be modified, e.g., not only by alloying ZnO with MgO or cadmium oxide (CdO) (see Ref. 3 and references therein), but also by varying their chemistry, for instance when going from SnO₂ to SnO.⁴ Free carriers, introduced by intentional as well as unintentional doping, are the reason for the remarkable conductivities of the TCOs.⁵ Prominent examples are aluminum-doped ZnO,⁶ tin-doped indium oxide,⁷ anti-mony-doped SnO₂,^{8,9} or even zinc–indium–tin oxide.¹⁰ Bulk TCO materials attract great attention due to their

Bulk TCO materials attract great attention due to their outstanding optical,¹ electrical,^{11–13} and electrochemical¹⁴ properties combined with excellent hardness and environmental stability.¹ This renders them highly interesting for applications as transparent front contacts for solar

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cells,^{2,11,15,16} as next-generation gate oxides for Si-based electronics,¹⁷ and in electrodes for photocatalytic water splitting.¹⁸ Surfaces of the TCOs are highly interesting since on these an electron accumulation^{19–21} as well as an electron depletion²² have been observed. Exploring the atomic structure and the termination of surfaces of such oxides is still subject of current research.^{23–25} In addition, the properties of the TCOs are drastically modified when they are prepared as nanobelts,²⁶ nanotubes,²⁷ nanoribbons,²⁸ nanowires,²⁹ and nanoparticles.³⁰

Even though the TCOs are highly interesting for these reasons, their most desirable widespread application in semiconductor electronics, for instance as (light-emitting) diodes, is currently hampered by the lack of stable and reproducible *p*-type TCOs. For other photovoltaic or optoelectronic devices, a deep understanding of intrinsic key properties is necessary. In this context, especially the electronic band structures around the fundamental gaps or the effective electron masses have to be thoroughly described. Also the optical absorption and emission properties near the band-edge and their dependence on the light polarization (due to dipole selection rules) need to be understood. For the TCOs, a plethora of experimental results is available,^{31,32} however, their interpretation can be challenging, for instance, due to sample-quality issues that depend on the manufacturing technique, the substrate temperature during growth, or the post deposition treatment.³³ In the case of In_2O_3 , growth can be phase-selective and can lead to both cubic as well as rhombohedral (rh)

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polymorphs with band gaps differing by as much as 0.7 eV.³⁴ In the case of ZnO (see Ref. 35 and references therein), and also for SnO₂ (see Refs. 36–38) the details of the valence-band ordering and the band symmetries are not unequivocally determined from experiment.

Due to the continuous development of sophisticated methods and algorithms as well as the increasing power of today's supercomputers, the ability to precisely describe theoretically various properties of such materials has made substantial progress during the last few years. The accurate parameter-free description of the quasiparticle (QP) electronic structures and the frequency-dependent dielectric functions of complex materials (see the reviews in Refs. 39-42 and references therein) has become possible and leads-hand in hand with modern experimental techniques-to a deep understanding. Many-body perturbation theory is used to take the excitation aspect of important experimental techniques into account and can be applied to compute the electronic and spectral properties of bulk oxides in a given ideal crystal structure^{43–47} but also for alloys, $^{3,48,49}_{,52-54}$ oxide systems with defects, $^{50,51}_{,50,51}$ and *n*-type TCOs.

In Sec. II of the present article, we summarize the theoretical and computational approaches that we used to compute QP band structures and the optical absorption of various oxide materials. We present the corresponding band structures and provide detailed information about band gaps and effective masses in Sec. III. The line-shape and the oscillator strength of the optical absorption edges are discussed in Sec. IV. Section V summarizes this review article.

II. THEORETICAL AND COMPUTATIONAL APPROACH

The present review article focuses on the description of excited state properties, such as the QP electronic structure and the optical absorption spectrum, for different oxides by means of an ab initio framework. Their ground state properties and, in particular, determining the equilibrium atomic geometries, e.g., within density functional theory (DFT),^{55,56} are not part of the present work. For more information about that as well as about all computational details we want to point the reader to Table I, which

TABLE I. For each material the references are given that contain more information on the calculations of the ground state and the excited state properties that are reviewed in this work.

Material	Ground state	Excited state	
MgO, ZnO, CdO	Refs. 45, 46, and 57	Refs. 43, 45, and 46	
SnO ₂	Ref. 44	Ref. 44	
SnO	Refs. 24 and 58	Refs. 24 and 58	
In ₂ O ₃	Ref. 59	Ref. 59	
$Mg_xZn_{1-x}O, Cd_xZn_{1-x}O$	Refs. 45, 46, and 48	Refs. 3, 45, and 46	
SiO ₂	Ref. 60	Refs. 60 and 61	

indicates the reference(s) in which the respective calculations are presented in more detail.

A. QP electronic structures

Important experimental techniques such as (inverse) photoelectron spectroscopy or x-ray absorption and emission studies probe excited state properties of a material, since during these processes an electron is added to or removed from the system. Hence, the ground state description provided by a Kohn-Sham (KS) scheme within DFT is not sufficient; instead, QP effects have to be taken into account.⁶² Interpreting the KS eigenvalues as excitation energies, typically leads to too small band gaps and wrong band dispersions,^{63,64} since the KS eigenvalues neglect the excitation aspect that is characteristic for the experimental techniques that are used to measure these quantities.

Electronic excitations can accurately be treated by correctly describing the electronic self energy Σ . Expressing Σ as the product of the Green's function of the electrons G and the screened Coulomb potential W, as it was introduced by L. Hedin in 1965,^{65,66} is an essential simplification of the description by neglecting vertex corrections. Nevertheless, the fully self-consistent solution of the resulting QP equation is very demanding from a computational point of view. For that reason, it is common to rely on perturbation theory to compute QP energies in practice: In the G_0W_0 scheme first-order QP corrections are calculated for the eigenvalues of some starting electronic structure. It is immediately clear that any initial electronic structure must not be too far from the final results for first-order perturbation theory to be sufficient.

We found that the range-separated HSE06 hybrid functional, $^{67-69}$ but with a range-separation parameter of 0.15 a.u.⁻¹ = 0.3 Å⁻¹, reasonably fulfills this requirement for several oxides.^{43–46,70,71} In this work, we review results for different TCOs that rely on the HSE functional to obtain the starting electronic structures for the G_0W_0 calculations; the entire approach is called HSE+ G_0W_0 in the following. To keep the computational effort reasonable and since spin-orbit induced shifts are smaller than the QP corrections, we assume that the influence of the spin-orbit interaction on the QP corrections is negligible. Therefore, we take the spin-orbit interaction into account⁷² only when calculating the HSE electronic structure and then apply the QP corrections calculated for spin-paired electrons.^{71–73} Overall, we expect the QP energies calculated by the scheme outlined above to be converged within about 0.1 eV.

A shortcoming of the HSE+ G_0W_0 approach is its large computational cost. This becomes particularly problematic for the calculation of optical spectra, i.e., when significantly more *k*-points have to be taken into account for converging, for instance, the macroscopic dielectric function around the absorption edge. In these cases, we pursue a different approach by approximating the results of the HSE+ G_0W_0 calculations via a DFT+ $U+\Delta$ method,^{3,41,43–46,70} where the term "DFT" indicates that either the local-density approximation (in the case of SnO₂ and SnO) or the generalized-gradient approximation (for MgO, ZnO, and CdO) is used. Therein, *U*, which denotes the additional Coulomb interaction term within the DFT+U approach,^{74,75} is adjusted such that the energy position of the *d* bands matches the HSE+ G_0W_0 result. In addition, Δ describes a scissors operator⁷⁶ that rigidly shifts all conduction bands so that the fundamental gap becomes identical to the HSE+ G_0W_0 value.

The HSE+ G_0W_0 as well as the DFT+ $U+\Delta$ calculations are carried out using the Vienna Ab initio Simulation Package (VASP).^{77–79} The wave functions are expanded into plane waves and the electron-ion interaction is described via the projector-augmented wave method.^{80,81}

B. Optical absorption

In optical absorption experiments, excitation energies are studied that are not high enough to remove electrons from the system. Instead, upon the irradiation with light, an electron that gets excited from the valence bands into the conduction bands, leaves behind a hole in the valence bands. Due to their having opposite charge, the photocreated hole and the excited electron interact via the screened Coulomb attraction, leading to the formation of excitons in the material. Excitonic effects due to the electron-hole interaction can, for instance, be taken into account in the ab initio description by solving a Bethe– Salpeter equation (BSE) for the optical polarization function.^{82,83}

For its numerical solution, the BSE is typically rewritten into an eigenvalue problem for the electron-hole pair Hamiltonian.⁸³ We use the KS eigenstates from the DFT+ $U+\Delta$ calculation to compute the matrix elements of the statically screened Coulomb attraction of the electrons and the holes as well as of the unscreened exchangelike term that accounts for local-field effects.^{41,84} The DFT+ $U+\Delta$ KS eigenvalues are used to describe the transition energies of noninteracting electrons and holes. The corresponding optical transition-matrix elements are computed using the longitudinal approximation.⁸⁵ After constructing the excitonic Hamiltonian, its eigenstates and eigenvalues can be used to calculate the optical properties of the system.^{86–88}

The converged calculation of the optical absorption spectrum in the vicinity of the band edge requires a large number of k points.^{43,89} Since for some of the materials discussed in this work, the lowest optical transitions are confined to the vicinity of the Γ point, this part of the Brillouin zone (BZ) can be sampled more densely to

obtain convergence for the absorption edge⁸⁹; we use hybrid *k*-point meshes⁸⁹ to achieve that goal. However, the computational cost of diagonalizing the resulting excitonic Hamiltonian matrices with ranks *N* of up to $\approx 100,000$ is much too high. Therefore, in this work an efficient time-evolution scheme⁹⁰ is used to calculate the dielectric function $\varepsilon(\omega)$ from the excitonic Hamiltonian.

When very dense *k*-point meshes are used, the large computational cost prohibits taking enough conduction bands into account for including high-energy optical transitions. This, however, becomes necessary to obtain converged results for the real part of $\varepsilon(\omega)$ at low photon energies due to the Kramers–Kronig relation^{91,92} between the real and imaginary parts. Therefore, as described in detail in Refs. 43, 45, and 46, we also solve a BSE using a more coarse *k*-point mesh to include optical transitions at intermediate energies. In addition, the optical transitions lying above this BSE cutoff are included (up to 200 eV) on the KS level. Finally, the absorption coefficient $\alpha(\omega)$ is calculated from the resulting dielectric function.

III. QP ELECTRONIC STRUCTURES

The QP band structures calculated using the $HSE+G_0W_0$ approach are depicted for MgO, ZnO, CdO, SnO₂, SnO, In₂O₃, and SiO₂ in Fig. 1. For the oxides studied in this work the uppermost valence bands are derived from O 2*p* states and the lowest conduction-band states originate from *s* states of the respective metal cation.⁹³

In addition, from Fig. 1 it is obvious that MgO, ZnO, SnO₂, In₂O₃, and SiO₂ are direct semiconductors with the fundamental band gap at the Γ point; the two monoxides CdO and SnO are found to be indirect semiconductors. While in CdO the conduction-band minimum occurs at the Γ point and the maximum of the valence bands occurs away from Γ , the situation is different for SnO. In this material the valence-band maximum is located at Γ and the conduction band shows a pronounced minimum at the M point of the BZ, whereas Γ is an M_1 saddle point for the lowest conduction band (see Fig. 1). The direct and the indirect QP band gaps of these oxides are compiled in Table II. These results for the electronic structures have been carefully compared with experimental data^{44,59,105-107} (see also Table I for the corresponding references) and the agreement is found to be rather good.

From Table II it also becomes clear that all the oxides studied in this work, with the exception of rocksalt- (*rs*) CdO and litharge- (*lt*) SnO, have fundamental band gaps that are large enough for visible light not to be absorbed by the intrinsic materials. Even though the direct gaps at the Γ point are as large as 1.81 eV (*rs*-CdO) or even 3.21 eV (*lt*-SnO) the indirect gaps of these two materials are much smaller. Hence, phonon-assisted optical absorption can occur in the visible spectral range for samples of both

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FIG. 1. QP band structures in the vicinity of the fundamental gap of (a) MgO, (b) ZnO, (c) CdO, (d) SnO₂, (e) SnO, (f) In_2O_3 , and (g) SiO₂ calculated using the HSE+ G_0W_0 approach. The valence-band maximum has been set to zero energy.

materials. In addition, in the case of *lt*-SnO the smallest *direct* gap of only 2.68 eV occurs at the *M* point of the BZ.

The HSE+ G_0W_0 results are used to derive the effective electron masses along different directions in the BZ via parabolic fitting to the QP band structures in the close vicinity of the Γ point (the *M* point in the case of SnO). We refer the reader to the literature for the effective masses that have been derived for the *rs* polymorphs of MgO and CdO,^{45,46} *rt*-SnO₂,⁴⁴ *rh*-In₂O₃,⁵⁹ or the wurtzite (*wz*) polymorphs of MgO, ZnO, and CdO.

TABLE II. Direct $[E_g^{\text{dir}}(\Gamma)]$ and indirect (E_g^{indir}) QP band gaps as well as optical band gaps for ordinary $(E_g^{\text{abs},\perp})$ and extraordinary $(E_g^{\text{abs},\parallel})$ light polarization are given in eV. Harmonic mean values for the effective electron masses m_e^* , derived from the HSE+ G_0W_0 band structure (HSE band structure for In₂O₃⁵⁹), are given in units of the freeelectron mass. Experimental values for $E_g^{\text{dir}}, E_g^{\text{indir}}$, and, if available, for m_e^* are given along with the corresponding reference in parenthesis.

	$E_g^{\rm dir}(\Gamma)$	E_g^{indir}	$E_g^{\ \mathrm{abs},\perp}$	$E_g^{\mathrm{abs},\parallel}$	m_e^*
rs-MgO	$7.49(7.67^{94})$		6.95	6.95	0.38
vz-ZnO rs-CdO	$3.21(3.4^{50})$ $1.81(2.28^{97})$	 0.68 (0.84 ⁹⁷)	2.96 1.73	3.00 1.73	$0.30(0.28^{50})$ $0.21(0.21^{98})$
rt-SnO ₂	3.65 (3.699)		4.13	4.75	0.23 (0.26 ¹⁰⁰)
t-SnO	$3.21(2.7^{101})$	$0.16(0.7^{101})$	1.79	2.29	0.42
$h-\ln_2O_3$	$3.25 (3.5^{102})$ 8 76 (8 8 ¹⁰³)				$0.22 (0.3^{102})$ 0.57 (0.58 ¹⁰⁴)
5-5102	0.70 (0.0)				0.57 (0.56)

For beta-cristobalite- (β) SiO₂, effective electron masses of $m_L^*(\Gamma_{\text{CBM}}) = 0.57$, $m_K^*(\Gamma_{\text{CBM}}) = 0.58$, and $m_X^*(\Gamma_{\text{CBM}}) = 0.57$ are obtained and in the case of *lt*-SnO, the conduction-band minimum at the *M* point of the BZ leads to electron masses of $m_A^*(M_{\text{CBM}}) = 0.37$ and $m_\Gamma^*(M_{\text{CBM}}) = 0.49$ (all in units of the free-electron mass). We also use these values to derive the harmonic mean values of the electron masses for all the oxides (see Table II).

In the case of rs-MgO^{45,46} we find a small anisotropy (indicating that the bands are not strictly parabolic within the k range used for the fit) and the two 2-fold degenerate valence bands are heavy-hole related (see Table III). Their masses are of the same order of magnitude in all three high-symmetry directions and are, roughly, one order of magnitude larger than the values for the light-hole band. Also the lowest CB turns out to be almost isotropic. As with rs-MgO, the two uppermost valence bands in rs-CdO are heavy-hole bands and the third one is of light-hole type.^{45,46} Since for *rs*-CdO the valence-band maximum occurs away from the Γ point in the BZ, the values for the effective mass of the uppermost two valence bands are negative along certain directions in k space (see Table III). This has been traced back to the fact that the p-d hybridization is forbidden at Γ due to the rs lattice symmetry.^{45,46,57} In this material holes are expected to occur at the valence-band maximum between K and Γ . We observe that the effective electron masses for β -SiO₂ are very isotropic; the anisotropy of the ones for lt-SnO (at the Mpoint of the BZ) is slightly larger. Comparison to experimental or other theoretical results is given in Refs. 44, 45, and 59.

From Table II it becomes clear that the electron masses for the different oxides decrease by almost a factor of two going along the row MgO, ZnO, SnO₂, In₂O₃, and CdO. Contrary, the values of the electron masses for SnO and SiO₂ are larger than for the other oxides. Again, we want to emphasize that the parabolic approximation is only fulfilled in the close vicinity around the Γ point. From our calculations of the effective electron masses we conclude that the electron mobilities should be large in *rs*-CdO,

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TABLE III. Effective masses m^* (in units of the free-electron mass m) at the BZ center along the $\Gamma - X$, $\Gamma - K$, and $\Gamma - L$ directions for rs-MgO and rs-CdO as derived from the HSE+SOC+ G_0W_0 scheme. Values are given for the lowest conduction band and the uppermost valence bands.

	rs-MgO	rs-CdO	
$m_{\chi}^{*}(\Gamma_{6c}^{+})$	0.36	0.19	
$m_{K}^{*}(\Gamma_{6c}^{+})$	0.42	0.25	
$m_L^*(\Gamma_{6c}^+)$	0.36	0.19	
$m_X^*(\Gamma_{8v}^{-})$	1.85	4.85	
$m_{K}^{*}(\Gamma_{8v}^{-})$	4.53	-1.35	
$m_L^*(\Gamma_{8v}^{-})$	3.21	-1.98	
$m_X^*(\Gamma_{8v}^{-})$	1.61	2.33	
$m_{K}^{*}(\Gamma_{8v}^{-})$	1.65	3.52	
$m_{L}^{*}(\Gamma_{8v}^{-})$	2.37	-3.63	
$m_X^*(\Gamma_{6v}^{-})$	0.44	0.36	
$m_{K}^{*}(\Gamma_{6v}^{-})$	0.44	0.38	
$m_L^*(\Gamma_{6\nu})$	0.36	0.24	

rh-In₂O₃, and rutile- (rt) SnO₂ with the ones of the other oxides being up to a factor of 2.7 smaller.

In addition, we used the electronic structures derived from the HSE+ G_0W_0 approach also for the derivation of the charge-neutrality levels and, hence, the band alignment for different materials.^{45,61,108,109} Furthermore, the HSE+ G_0W_0 approach has successfully been applied to describe nitride systems,^{39,110–114} antiferromagnetic transition-metal oxides,^{41,84} biaxially and uniaxially strained ZnO,^{45,73} and the nonequilibrium *wz* polymorphs of MgO and CdO.^{45,71}

IV. OPTICAL ABSORPTION

In Fig. 2 we give an overview of how QP corrections as well as excitonic effects influence the imaginary part of the dielectric function in the case of rs-MgO. From Fig. 2(a) it becomes clear that the QP shifts, simulated by a scissors shift Δ , induce a blueshift of the optical spectrum. If actual QP energies were used (instead of the scissors approximation), the shifts for the states at higher energies would be larger.⁴⁵ Figure 2(b) shows the impact of excitonic and local field effects: We observe a redshift of the BSE curve toward lower photon energies, that does not compensate the blueshift arising from the QP effects discussed above. Aside from the redshift, we observe a strong enhancement of the peak intensities and plateau heights due to the Coulomb interaction. In addition, the BSE curve shows a remarkable feature that dominates the absorption onset, which is attributed to a bound, Wannier-Mott-like electron-hole-pair state. Finally, the comparison with a measured curve¹¹⁵ in Fig. 2 shows how well the results obtained within the BSE approach agree with experiment and clearly points out the importance of including excitonic effects in the calculations.

The optical absorption coefficients calculated from the solution of the BSE for *rs*-MgO, *wz*-ZnO, *rs*-CdO,



FIG. 2. Imaginary parts of the dielectric function of *rs*-MgO calculated using the GGA (dotted black curve), the GGA + Δ (dashed red curve), and the BSE approach (solid blue curve). Experimental results (black circles) from Ref. 115 are shown for comparison.

rt-SnO₂, and *lt*-SnO are depicted in Fig. 3. We found that the inclusion of excitonic effects in the description is critically important for all these TCOs as they (i) dominate the spectral features around the absorption onset, $^{43-45,70,73}$ and (ii) strongly influence the overall spectral shape. 43,45

As a consequence, a pronounced excitonic peak is visible around the absorption onset, in particular, of rs-MgO and rt-SnO₂ [see Figs. 3(a) and 3(d)]. Such a peak also occurs at the onset of the absorption for wz-ZnO even though this is not as obvious from Fig. 3(b) due to the scale of this plot. However, the influence of such an excitonic bound state is significantly smaller for rs-CdO and lt-SnO [see Figs 3(c) and 3(e)]. One reason is that these two materials have indirect fundamental band gaps and, in addition, the electronic dielectric screening is significantly stronger for these two semiconductors, which is why the excitonic effects are reduced.

The influence of the electron-hole interaction on the overall line shape is referred to as redistribution of oscillator strength in the literature. For all the oxides, peaks at higher photon energies are red-shifted due to the excitonic effects by about 1-2 eV.^{43–45} In addition, a strong enhancement of the peak intensities and plateau heights due to the Coulomb interaction occurs at low photon energies.

Comparing the dielectric functions calculated from the solution of the BSE to experimental results has shown a very good agreement, for instance, for MgO, ZnO, and CdO,^{43,45} but also for SnO₂.⁴⁴ Peak positions and relative peak heights are very well described by the calculations for these materials. We also find good agreement of our results with other computational work on the same theoretical level, for instance, for *rs*-MgO¹¹⁶ or *wz*-ZnO.^{117,118} We want to point out that the scissors operator Δ that is used to mimic the QP gaps does not reproduce the energy dependence of the self-energy operator Σ . Hence, we find a general trend of a slight underestimation of the energy positions of the higher-lying CBs.⁴⁵ In addition, the



FIG. 3. Absorption coefficient (in 10^5 cm⁻¹) around the fundamental band gap for different oxide semiconductors as calculated from the solution of the BSE. Solid black (dashed red) curves represent ordinary (extraordinary) light polarization.

absolute value of the absorption coefficient turns out to be underestimated in the calculations (e.g., for ZnO⁵²), which can be explained, for instance, by imperfect samples, whereas our calculations describe defect-free crystals.

By linearly extrapolating the slope of the absorption curves (see Fig. 3) around the half maximum of the onset to zero absorption, we obtained the optical gaps of the oxides compiled in Table II. We want to point out the strong (noticeable) optical anisotropy of *lt*-SnO and *rt*-SnO₂ (wz-ZnO). In the case of rs-MgO and wz-ZnO the optical gaps are significantly smaller than the QP gaps also given in Table II. For these two materials this difference arises mostly due to the binding energies of the lowest bound excitonic states^{45,73,89} that reduces the optical gaps with respect to the QP gaps. In addition, the Lorentzian broadening of 0.1 eV, which is used to account for temperature, instrumental, and lifetime effects, further leads to a reduction of the optical band gap in our calculations. Both the broadening effect and the exciton-binding energy also play a role for rt-SnO₂. However, as elucidated in Ref. 44, the optical transitions between the uppermost valenceband and the lowest conduction-band states are dipole forbidden in this material; consequently, the optical gap is found to be much larger than the QP band gap (see Table II).

In the case of *rs*-CdO and *lt*-SnO the optical gap is found to be in between the lowest indirect and the lowest direct QP gap. For these two materials the dielectric screening is much larger which is the reason why there occurs no pronounced peak due to a bound excitonic state at the absorption onset. Nevertheless, excitonic effects are responsible for the optical gap being smaller than the lowest indirect gap for CdO, in addition to the broadening (see above). In the case of *lt*-SnO, the LDA+ $U+\Delta$ description of the uppermost valence band at the *M* point of the BZ differs by 0.73 eV from the HSE+ G_0W_0 result. This explains why the optical band gap found in our calculations turns out to be underestimated by this amount; also an influence on the line shape of the spectrum cannot be excluded and will be further investigated in the future.

The results for the dielectric functions obtained from the solution of the BSE have also been used to compute reflectivities as well as the electron energy loss functions for MgO, ZnO, and CdO.^{43,45} For these materials we identified valence and conduction-band levels (as well as their atomic origin) that are involved in the transitions that cause the main peaks of the spectra. The lowest eigenstates of the excitonic Hamiltonian have been calculated for MgO⁸⁹ as well as (strained and unstrained) ZnO.⁷³ In the case of aluminum nitride (AlN) and calcium oxide (CaO), the BSE approach helped to identify the importance of van Hove singularities and of the excitonic effects for the absorption spectra.¹¹⁹

V. SUMMARY

In this article, the QP band structures of MgO, ZnO, CdO, SnO₂, SnO, In₂O₃, and SiO₂, calculated using the $HSE + G_0 W_0$ approach, have been presented. For these different oxides, we conclude that our results agree well with experimental findings, regarding, for instance, fundamental band gaps and band dispersions or effective electron masses. In addition, we presented the optical absorption spectra of MgO, ZnO, CdO, SnO₂, and SnO. We showed that for these oxides the impact of excitonic effects is significant and has to be taken into account in the calculations. We find that this task can be successfully accomplished by solving the BSE. For SnO we conclude that the LDA+ $U+\Delta$ approach does not reproduce the HSE+ G_0W_0 results well enough, hence, a reliable parameter-free description of the optical properties requires future effort.

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