Multi-gap absorption in CaCu$_3$Ti$_4$O$_{12}$ and the predictivity of ab initio methods

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We report the electronic properties of the quadruple perovskite CaCu$_3$Ti$_4$O$_{12}$ as obtained via several density-functional based methods, and propose a new interpretation of optical experiments to the effect that four distinct transitions (centered around 0.7, 1.5, 2.5, and 3.5 eV) contribute to the spectrum. The comparison with experiment is satisfactory, especially after we account for the effects of spin disorder, which does not close the fundamental gap but suppresses the transition intensity. We find that some of the methods we employ tend to overestimate considerably the gaps for standard values of the respective adjustable parameters.

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I. INTRODUCTION

The popular line “If it’s been measured, why are you calculating it?” attributed to Volker Heine emphasizes the need for electronic structure theory to harness its predictive and interpretive potential. In this paper, we revisit the low-energy optical properties of the quadruple perovskite CaCu$_3$Ti$_4$O$_{12}$ (CCTO henceforth, risen to popularity a decade ago because of its anomalous dielectric response), heeding the advice implicit in Heine’s remark in two distinct respects. Firstly, theoretical predictions predated reliable experiments, and here we provide a new and improved interpretation of the latter. Secondly, electronic structure theory often revisits known results to provide additional insight and to validate its predictive power in retrospect. In this spirit we apply to CCTO several density-functional-theory (DFT) state-of-the-art methods, which yield a mixed bag of good and bad news. Some advanced methods appear to be struggling, while others yield satisfactory agreement with experiment.

A. Motivation

Early experimental reports [10] on CCTO circa 2002 had suggested a fundamental gap in excess of 2.5 eV. Values as low as 0.2 eV obtained in DFT local-density-approximation (LDA) calculations were, not unreasonably, discounted in view of the known gap underestimation problem of local and semilocal functionals. Looking at the LDA bands, however, we realized that the lowest gap might be a low-energy transition between localized and predominantly Cu-like states, rather than the natural dipole-allowed transition between O p valence and Ti d conduction bands, a situation analogous to other Mott-like insulators. Therefore, in 2006 we used self-interaction corrected LDA (PSIC) known by then to reproduce quite accurately the gaps in many Mott-like cuprates [11] to find out how beyond-LDA bands would look like in the Cu-dominated gap region. We found that the fundamental gap (indirect, forbidden, and between mostly Cu-like states) was only about 0.6 eV, moderately increased in absolute value over the LDA value. This surprising result seemed to point to smaller-than-usual correlation effects in the nearly filled 3d Cu(II) shell; put differently, the on-site interaction, which self-interaction corrections largely restore to its correct size, appeared to be rather more screened at Cu sites in CCTO than in other magnetic Cu oxides.

Systematic experiments (see Sec. III C below) first appeared in 2008, when Kant et al. inferred from optical conductivity an electronic structure qualitatively matching that suggested by LDA and, to a somewhat larger extent, by PSIC with weak Cu-dominated transitions starting at about 0.5-0.7 eV. In 2011, a different picture was proposed [12] to the effect that the “Cu-Cu” transition would start at about 1.8 eV, based on reflectivity measurements interpreted via GGA+U (Generalized Gradient Approximation +U). The calculations used a U–J parameter much smaller than the usual value for Cu oxides (reminding us of the low-correlation argument), yet it produced a fundamental gap much larger than previously obtained by PSIC. This suggested that it would be a good idea to revisit and expand our previous investigation applying further advanced methods to CaCu$_3$Ti$_4$O$_{12}$ to help sort out the matter and provide a robust interpretation.

In this work, we discuss the electronic properties of CCTO based on results from several different DFT-based methods. We eventually propose an interpretation of experiments, as well as further experimental tests, based on one of them, the variational PSIC method (VP-SIC henceforth). Our conclusion in summary is that CCTO has a multifold interband absorption due to its unusual Cu-induced upper-valence and lower-conduction band structure, and that the fundamental transition peaks around 0.7-0.8 eV (1500 nm), while the absorption peaking at 1.8 eV (700 nm) is an O p valence band to Cu d conduction band transition, at variance with the pre-
More intense absorptions between 2.5 to 4 eV are due to transitions into the higher-lying Ti $d$ conduction band. The fundamental gap is a Mott-like gap in the sense that the system has an odd electron count, the gap open between strongly localized and spin-polarized states, and would not exist in the absence of magnetic moments. We account for spin disordering in the paramagnetic (PM) phase in which most measurements are performed: the fundamental gap survives unscathed the breakdown of magnetic order, but the intensity of the transition across that gap is suppressed. As a test of the suggested picture, we point out features that should be observable in low- vs high-temperature optical and energy-loss spectroscopy experiments.

Besides VPSIC and GGA/LDA, we calculate the electronic structure using GGA+U, hybrid functionals, and many-body perturbation theory. Our theory-experiment comparison indicates that hybrid and GGA+U end up quite far from experiment, overestimating severely the gaps, whenever standard values are used for the adjustable parameters they depend on, a conclusion that has obvious methodological implications. Also, the corrections to the eigenvalues of local or semilocal functionals provided by VPSIC are close to those of non-self-consistent one-shot GW, suggesting that the "beyond-local" correlation is described by VPSIC with similar accuracy.

II. METHODS

As usual in the business of ab initio optical properties, we elect to interpret the eigenvalues and eigenvectors of Kohn-Sham equations as quasiparticle energies and states. This is justified by the Kohn-Sham equations being formally identical to Hedin-Lundqvist quasiparticle equations if the self-energy $\Sigma$ is identified with the exchange-correlation potential; for LDA, e.g., $\Sigma_{\text{LDA}}(\mathbf{r},\mathbf{r}^\prime,E) = \delta(\mathbf{r}-\mathbf{r}^\prime)\delta(E) + V_{\text{xc}}^{\text{LDA}}(\mathbf{r})$, and similarly for functionals containing some degree of non-locality and implicit energy dependence such as hybrids or PSIC. It is obviously interesting, therefore, to compare results obtained by different exchange-correlation functionals. This is done in Sect. IV in particular IV A. In the same spirit, we also discuss in Sect. IV C "many-body" corrections to semilocal functionals, both empirical and based on $G_0W_0$ non-self-consistent many-body perturbation theory.

In recent years, methods going beyond the local or semi-local approximation have become more affordable, and we are in the position to evaluate their relative merits under the assumption stated above. The optical conductivity, extinction coefficient, and electron-energy-loss function are extracted from the dielectric function $\tilde{\varepsilon}(\omega)$ calculated within the random phase approach from the joint density of states obtained with the variational version of pseudo self-interaction-corrected LDA (VPSIC). We also calculate gaps and transitions with Ceperley-Alder LDA Perdew-Bekke-Ernzerhof GGA, the Dudarev version of the GGA+U, the Heyd-Scuseria-Ernzerhof (HSE) hybrid functional. To avoid bias due to changes in volume, we use the cubic magnetic primitive cell (40 atoms) at the experimental lattice constant of 7.38 Å with internal coordinates optimized with GGA, imposing a threshold of 0.01 eV/Å on force components. Since CCTO is cubic, the positions of Ti, Cu and Ca are fixed by symmetry; Ti-O octahedra and Cu-O plaquettes are "rigid" and all identical geometrically (see Fig. 1), so the O positions are determined by just the Ti-O and Cu-O distances (1.959 Å and 1.963 Å, respectively).

![FIG. 1. (Color online) Crystal structure of CCTO.](image)
FIG. 2. LDA vs VPSIC bands of CCTO. Arrows indicate schematically the four transitions discussed in the text.

III. VPSIC RESULTS VS. EXPERIMENT

This Section compares the VPSIC optical functions of CCTO with two distinct sets of experimental data. In Sec. IV we will examine and discuss the gaps provided by the other methods. We will be switching units several times to ease the comparison with experiment.

A. Band structure

CCTO is G-type antiferromagnetic (AF) on the Cu lattice with a Neél temperature of 25 K. Its band structure obtained by LDA and VPSIC is displayed in Fig. 2. The LDA and VPSIC bands are rather similar in structure and energy separation. The near-gap bands connected by the transition marked ‘1’ in Fig. 2 are dominated by O-hybridized Cu-like spin-polarized states. The top valence band and the bottom conduction band are fully spin-polarized, and as can be seen in the orbital- and site-projected density of states (DOS) as obtained by VPSIC in Fig. 3, their projections on any given Cu site have opposite polarization. As usual, the removal of self-interaction tend to increase all the gaps. The largest increases are found for the transitions marked ‘3’ and ‘4’ in Fig. 2 to the upper conduction band of predominantly Ti character, i.e. for the standard charge transfer gaps. The local orbital character of the near-gap spin-polarized states is completely determined by Cu $d$ and ligand O’s in each plaquette (see Fig. 4), in accordance with the DOS of Fig. 3.

As far as optical absorption is concerned, the band structure in Fig. 2 suggests that four distinct relevant absorptions are expected, marked ‘1’ to ‘4’. The first transition is between O-hybridized Cu-like bands, between 0.5 and 0.9 eV (2000-1000 nm, 4000-7000 cm$^{-1}$). A large joint DOS is expected due to extended parallel-band sectors especially around the X point; also, despite the similar character of the initial and final states, the matrix elements should not be suppressed, because this is an intersite transition (intralattice transitions are forbidden by the spin conservation selection rule). The second transition is between valence O $p$ and low-conduction Cu $d$ bands in the range 1.4-1.9 eV (800-650 nm, 11000-15000 cm$^{-1}$), which is not expected to be suppressed selection-rule-wise. The third absorption is Cu $d$ upper-valence to Ti $d$-O $p$ conduction at 2.6-3.0 eV (500-400 nm, 21000-24000 cm$^{-1}$), which is expected of average intensity; finally, an intense O $p$-Ti $d$ transition should start at 3.4 eV (350 nm; 27500 cm$^{-1}$).

FIG. 3. (Color online) DOS of CCTO from VPSIC in the near-gap region, projected on orbitals of the Cu and O atoms of a plaquette in the $xy$ plane (see Fig. 1).

The lower conduction band of Cu character is affected only weakly by self-interaction corrections, and accordingly the lower-energy O $p$-Cu $d$ and Cu $d$-Cu $d$ transitions ‘1’ and ‘2’ change moderately compared to LDA. This feature is key to our interpretation, and, as we will see in Sec. IV, it is not shared by other methods. The small magnetic fundamental gap ‘1’ may be labeled as Mott-like, since it depends on the interplay of spin polarization, Hund coupling, and on-site repulsion, and it is coherent with the textbook definition $U-c_{t}$, i.e. it includes (thanks to self-interaction removal) the cost U of adding an electron in the empty $d$ state as well as the hopping $t$, which is included in the band width. Since even LDA finds this gap (albeit barely), the on-site correlation acting on these states must not be especially strong. Further, the stronger O $p$-Cu $d$ hybridization plays a role in reducing the gap in CCTO compared to...
e.g. in YBa$_2$Cu$_3$O$_6$, whose gap$^2$ is 1.2 eV.

![Image of a gap; a similar conclusion was drawn earlier for MnO](image)

A related result

![Image of a gap; a similar conclusion was drawn earlier for MnO](image)

FIG. 4. (Color online) Magnetization density in CCTO seen along the z axis. All three kinds of Cu-O plaquettes are visible. The Cu onto which the DOS in Fig 3 is projected is the one at bottom center, with the $xy$-orbital-shaped density.

B. Paramagnetic vs antiferromagnetic phase

Before delving into the comparison, we need to discuss the role of spin disorder. The experiments we consider are done well above the Néel temperature $T_N=25$ K of the AF phase of CCTO, and therefore probe the paramagnetic (PM) phase. Since PM CCTO is insulating experimentally, it is most likely a collection of thermally disordered, randomly oriented Cu moments, rather than a zero-moment Pauli-type PM (in the latter phase, CCTO is found to be a metal). The fundamental transition, labeled ‘1’ in Figs 2 and below, is between Cu-like spin-polarized states (see Fig 2 and Fig 3) and is spin-selective in the sense that spin-allowed transitions only occur with matrix elements involving same-spin sites of the Cu lattice. In the PM, we expect the intensity of transition ‘1’ to be reduced, because spin mixtures will be involved. To expect a good match with experiment, this suppression should be assessed and accounted for.

We do not aim at sampling in detail the PM configurations (which is outside our scope and well beyond “naive” sampling techniques); rather we need to show a) that a gap survives in the misaligned-spin, i.e. non-AF-ordered, system when moments are non-zero, and b) that the intensity of the transition between the spin-polarized states is suppressed. Point a) is worth making directly; the notion that ab initio methods can obtain good gaps and magnetism in correlated systems where LDA or GGA fail (as the PSIC does, e.g. for YBCO, CuO, LaTiO$_3$, LaNiO$_3$/LaAlO$_3$ superlattices, etc.) is generally considered with suspicion because of the almost ubiquitous assumption of magnetic order in such systems. Our simplified disordered-moments PM, in fact, turns out to have a gap; a similar conclusion was drawn earlier for MnO$^2$ based on essentially the same electronic-structure technique (and a much better spin-disorder sampling). As to point b), in the PM the spin states are mixtures, i.e. spin projections are no longer just unity or zero, referring to a given quantization axis. Intersite transitions, that were spin-conserving in the AF, will thus be suppressed in the PM, whereas on-site transitions between formerly opposite-spin states will gain non-zero amplitude. While the latter are expected to be suppressed by the dipole selection rule, it is appropriate to explore how these two effects play out quantitatively.

To assess the degree of intensity suppression of absorption ‘1’ in the PM phase, we perform non-collinear-magnetization calculations whereby the six Cu moments in the primitive cell are oriented randomly, but directionally constrained to give a total magnetic moment of zero—that is, mimicking in effect one of the thermally accessible configurations of the disordered paramagnet. The non-collinear spins in the PM model are constrained by a penalty function: if that penalty is turned off, the AF ground state is recovered. Spin-orbit coupling is not included in these calculations. Since the low energy bands obtained with semi-local-functionals and self-interaction corrections are quite similar (see Fig 2), we use GGA to access the non-collinear and penalty-function features of VASP. As we are only interested in the effect on the lowest gap, we display the low energy portion of the GGA absorption for the PM (aligned to match the VPSIC gap) together with the VPSIC absorption.

The key result, as can be seen from the bands in Fig 5, is that the gap remains non-zero in the disordered PM, and close to the AF value. This should help dispel the myth that the distraction of magnetic order will lead to metallicity in ab initio calculations. Indeed, it does not, as long as magnetic moments survive.$^2$ A related result relevant to our interpretation below is that, as we surmised, the fundamental absorption is indeed suppressed in the PM compared to the AF, whereas the rest of the spectrum is practically unchanged. This improves agreement with experiments done at high temperature, as we discuss in the next Section. We report only the low energy portion of the PM optical constants (up to about 1.5 eV), since spin disorder only affects the spin-polarized transition ‘1’. We note in passing that the metallic Pauli-PM phase (not shown) shows typically metallic optical features such as a Drude peak at low-frequency, which are not observed in any of the experiments.

C. Comparison with experiments

Based on diffuse reflectance measurements, Ref.$^{10}$ suggests as lowest-energy transition an indirect-gap absorption peaking at 700 nm, and attributes it to transitions from the mainly Cu-like upper valence states to the mainly Cu-like first conduction band, i.e. to transition ‘1’ of our band structure in Figs 2 and 5. Higher-energy intense absorptions are attributed to O $p$-Ti $d$ dipole
FIG. 5. GGA bands for the CCTO cell with non-collinear spins mimicking the disordered PM.

This interpretation is based on GGA+U calculations with U–J=6.5 eV, a rather low value for Cu oxides, which nevertheless pushes transition ‘1’ up to the needed 1.7-1.8 eV (see the discussion in Sec.IV).

The fundamental transition, which connects the upper valence and bottom conduction Cu-like states and is labeled ‘1’ in Figs.2 and 3, is instead at lower energy, peaking at about 1500 nm in the AF phase. However, in the PM the intensity of this absorption is suppressed. As no experimental data were reported in this wavelength region, it is probable that no significant signal was detected. Account for spin disorder resolves the potential discrepancy. Conversely, our result suggests that similar experiments in the AF phase at low temperature (and pure, untwinned, magnetically ordered crystals) should reveal this low-energy transition, providing a direct experimental countercheck of our interpretation.

FIG. 6. Extinction coefficient for PM and AF CCTO calculated with VPSIC. Compare with Fig.7b (inset) of Ref.10.

To compare with this experiment directly, we calculate the extinction coefficient (Fig.6) and conductivity (Fig.7 and 9; see the discussion below) from the dielectric function \(\tilde{\varepsilon}(\omega) = \varepsilon_1 + i\varepsilon_2\) (the strongest dependence is on the imaginary part \(\varepsilon_2\)). Based on these results, and in particular the extinction coefficient displayed in Fig.6 (to be compared e.g. with the inset of Fig.7b of Ref.10), we propose a different interpretation than that just outlined: the peak at 700 nm is the O p-Cu d labeled ‘2’ in Fig.2 and 3, which is also an indirect transition; the shoulder at 400-450 nm is the O p-Cu d labeled ‘3’; and finally the main peak at 300 nm is due to the main interband O p-Ti d transition, labeled ‘4’.

We now come to wide-range dynamic conductivity measurements which also seem to suggest a multigap spectrum. Tauc extrapolation at low energy is difficult due to low intensity and the probable indirect character of the transition, but a very weak indirect transition starting at about 5000 cm\(^{-1}\) can be inferred\(^8\).\(^9\) Another more intense transition follows at about 1.5-1.7 eV and finally the intense allowed absorption peaks at 3 eV.

These features are reasonably well reproduced by our calculation for the PM in Fig.7, as can be seen comparing with the experimental data in Fig.8 where the Tauc fits suggest an onset (i.e. a minimum gap) at about 0.6-0.9 eV. This assignment is only tentative as there is no clear linear behavior over an extended frequency range.

Comparing AF and PM results, it appears that the seemingly strongly forbidden character of the fundamental transition is mainly a token of spin disorder, rather than of interband matrix element suppression. Indeed, intrasite d-d transitions would be suppressed by Laporte’s selection rule, but here they are effectively intersite, because of the spin structure of the material; the fundamental transition in the AF is in fact quite prominent (Fig.5 and Fig.6). Thus the same measurements below \(T_N\) should show a marked intensity enhancement in...
FIG. 8. Experimental Tauc relation for CCTO, to be compared with Fig. 7. Lines are possible fits for indirect forbidden transitions whose intercepts with the frequency axis identify the minimum gap. Figure by courtesy of P. Lunkenheimer.

the 5000-8000 cm$^{-1}$ (0.5-1 eV) range, providing another countercheck on our interpretation. The same applies to the conductivity, displayed in Fig. 9, whose behavior for the PM compares favorably with Fig. 7 of Ref. 8.

FIG. 9. Dynamical conductivity for CCTO calculated with VPSIC. Compare with Fig. 7 of Ref. 8.

A related result from Ref. 8 is that DC conductivity is Arrhenius-activated with a 0.2 eV characteristic energy. We attribute this simply to thermal carriers excitation across the fundamental gap. The latter is 0.6 eV in VPSIC, but we have estimated from a single GGA calculations including spin-orbit (not shown in the Figures) that the spin-orbit splitting of the upper valence and lower conduction bands (both having sizable Cu character) will reduce the gap to about 0.3 eV.

In closing this Section, we point out that beside low-temperature optical absorption another possible countercheck on our suggestions is electron-energy-loss spectroscopy, again in the 0.5-1 eV range. As shown in Fig. 10, the energy-loss function $\text{Im}[1/\epsilon(\omega)]$ has a marked peak at 0.9 eV in the AF (i.e., at low temperature) which is strongly suppressed in the PM (i.e., at high temperature) because of spin disorder. The sharp main plasmon at 13.5 eV (Fig. 10 inset) is the same in the AF and PM.

FIG. 10. Energy-loss function of AF and PM CCTO. The inset shows the main plasmon in the AF.

IV. RESULTS WITH OTHER FUNCTIONALS

Having obtained a satisfactory interpretation of the electronic structure of CCTO using VPSIC, we examine and compare the main band gaps obtained by GGA+U and the HSE hybrid functional. As mentioned earlier, GGA+U in the version employed here depends on the U–J parameter; we apply to the Cu $d$ shell a U–J ranging from 0 to 8 eV (a value of 8 or 9 eV is standard for Cu oxides). The HSE hybrid depends on the fraction $\alpha$ of screened Hartree-Fock exchange and the screening cut-off wavevector $\mu$; we consider $\alpha=0, 0.1, \text{ and } 0.25$, the latter being the proper HSE recipe (while varying $\alpha$, we keep the standard $\mu=0.2$ Å$^{-1}$); then we explore values of $\mu$ from 0.1 to 0.5 Å$^{-1}$, at the standard $\alpha=0.25$ (large $\mu$ means strongly screened Fock exchange at all wavevectors, recovering GGA as $\mu\to\infty$). The HSE standard value has a theoretical foundation in the formulation of the functional, and has the merit of being system-independent. That said, we deem this exploration worthwhile, as the $\alpha$ and $\mu$ parameters have been, on occasion, adjusted away from their standard value to cure various different issues in cuprates and titanates.

Finally we discuss quasiparticle corrections from GGA-based $G_0W_0$ many-body perturbation theory; $G_0W_0$ has no adjustable parameter per se, but uses the GGA bands to evaluate the Green’s function and screened interaction, whence the quasiparticle energies, and is not self-
A. Parameter dependence of main gaps in HSE and GGA+U

In this Section we discuss the electronic structure of AF CCTO as function of the relevant adjustable parameters of the various methods. We consider the transitions defined in Fig.2 namely the fundamental gap, i.e. the ‘1’ transition; the main charge transfer gap, i.e. the ‘4’ transition; the upper valence-upper conduction gap, i.e. the ‘3’ transition; and the minimum gap between the Cu-like lower conduction band and the Ti-like upper conduction band, i.e. roughly the difference of the ‘4’ and ‘2’ transitions, labeled ‘4–2’. These are shown for GGA+U as function of the U–J parameter in Fig.11, for HSE as function of the mixing parameter \( \alpha \) in Fig.12 and of the screening parameter \( \mu \) in Fig.13. By construction, plain GGA (which is quite similar to LDA in Fig.2) is recovered in each of the limits of vanishing U–J, \( \alpha \), and \( \frac{1}{\mu} \). The energies reported are edge-to-edge eigenvalue differences at the X point (for ‘1’ and ‘3’) and Γ point (for ‘2’ and ‘4’).

As expected, all valence-to-conduction gaps increase rapidly with U–J, \( \alpha \), and \( \frac{1}{\mu} \). The ‘1’ transition increases fastest in all cases, and the conduction-conduction gap diminishes. This means that, relative to the valence band, the Cu-like lower conduction bands are pushed up in energy more than the Ti-like upper conduction bands, opposite to what is observed in VPSIC.

This effect is especially strong in GGA+U, so much that the ‘4–2’ transition becomes negative, i.e. the empty Cu band is pushed into the Ti band. This is because the U correction acts efficiently on the Cu-like bands by enhancing both spin polarization and orbital polarization. The former widens the ‘1’ gap; the latter cleans up the \( O_p \) valence from Cu-like character, pushing it down and widening somewhat the apparent charge transfer gaps. We do not apply a U on Ti. That would only have minor, if any, effects: Ti orbitals are not spin-polarized, and U would only leverage orbital polarization, i.e. it would purge the valence and Cu-like states of what little Ti orbital content they have, thereby widening modestly, if at all, the relative gaps (at least for physically sensible values: of course one may always hope to get some effect for unphysically large U’s). In conclusion, to obtain a ‘1’ transition in the vicinity of 1 eV or less, U–J should be between 2 and 3, which is tiny by Cu-oxide standards.

In HSE, all gaps increase linearly with \( \alpha \) (Fig.12), with the ‘1’ transition increasing faster than the others. As Fig.13 shows, all the gaps decrease as \( \mu \) increases, i.e. when the Hartree-Fock exchange is gradually screened away. At standard \( \alpha \) the HSE functional gives a fairly good ‘4’ transition, 4.1 eV at Γ. The ‘1’ transition is overestimated hugely at over 3 eV, whereas, depending on which experiments one believes and whatever uncertainties one may attach to them, that gap is in the range of 0.5-1.5 eV at most. To obtain such a value one should
either use a small $\alpha$ in the vicinity of 0.05, or, probably better, a more or less standard value like 0.15 to 0.25 with a larger than usual $\mu$ (as large as 1.6 Å for $\alpha=0.25$). One way of stating this is that, compared to the upper bands, the Cu $d$ bands get too large a “correlation” correction (in the commonly-used, if questionable, sense of “any correction needed beyond semilocal DFT”) from standard HSE, and that the correction should be more “screened” than it is, consistently with the smaller-than-usual $U$–$J$ mentioned above for GGA+U. An additional issue enhancing the sensitivity of Cu states to U-like corrections may have to do with the fact that the band structure of CCTO is dominated by long range hoppings. Whatever the final answer, this overcorrection cannot be attributed (not straightforwardly, anyway) to self-interaction removal, which operates –although in different guises– in both HSE and VPSIC, and has moderate effects on the Cu states in the latter.

It is worth reiterating that the parameters in common use in the literature are those at the high end of the range considered here. $U$–$J$ up to 8 or 9 eV is quite usual in Cu oxides and $\alpha=0.25, \mu=0.2$ Å$^{-1}$ is the standard HSE recipe. For those values, both GGA+U and HSE produce an electronic structure whereby the Cu-like conduction states are way too high in energy, and the gap is too large by a factor of at least 3. Used in their default setting, GGA+U and HSE would predict a fundamental gap $\Delta^1$ of 2.2 eV and 3.2 eV respectively, whereas experiments and VPSIC agree that CCTO has a gap of less than 1 eV. Besides, the multiple absorptions involved in the CCTO spectra are not reproduced, as they end up being squashed by the overcorrection into a single high-onset-energy transition.

### B. Discussion

There are some general conclusion to be drawn from the results just discussed. At the very bottom, GGA+U, HSE, and VPSIC are all semiempirical methods, in that they depend on some sort of parameter. One recognizes, though, that these parameters intervene very differently in each method.

By construction, GGA+U is the most directly affected by its internal parameters. These can be estimated to some degree on a non-empirical basis from atomic quantities or from linear response but in all cases they are externally-determined system-dependent inputs (occasionally even dependent on internal parameters or external conditions within the same system), and not self-consistent and internal, so that in the end they are simply regarded as adjustable by most practitioners. Whether this is admissible or desirable is as much a philosophical as an operational question that depends on the specific objectives of an investigation. In the present case we found that the consolidated parameter recipe simply does not seem to function. One may (we don’t) elaborate further about which atom and shell the correction should be applied to (e.g. would U’s on Ti or O $p$ make a difference: we argued above that they would not cure the problem), adding more parameters: this would probably bring us no nearer to a solution.

As mentioned, HSE’s standard recipe has a theoretical foundation in the formulation of the functional, and has the merit of being system-independent (beside the practicality of including screening at the functional formulation level, and not a posteriori as in other hybrids based on, again, empirical estimates or models of the screening). That said, the $\alpha$ and $\mu$ parameters have been adjusted away from their standard value to cure various different issues (structure, electronic properties, magnetism, etc.) in many occasions, among which cupric oxide CuO.

It is only fair to discuss in this context the parametric dependence built into VPSIC. For a detailed treatment we defer to the original work and to a recent review which also discusses in detail the analogies and differences with GGA+U and hybrids. In short, the screening of atomic self-interaction corrections by the environment is described by a single parameter $\alpha_s=1/2$, a value based on a Slater transition-state argument. The dependence on $\alpha_s$ of relevant quantities in solids has been studied and the result is that $\alpha_s=1/2$ is indeed the optimal value on average over a vast class of materials. We systematically use that value hence effectively we do not regard $\alpha_s$ as a parameter at all. In a case where a detailed comparison has been carried out VPSIC has been found to perform similarly to HSE; discrepancies (and controversy) did occur in other cases, however, especially on cupric oxide CuO.

Since we are dealing with a titanate that is strongly characterized by cuprate-like electronic features, it is appropriate to recall the solid success record of VP-SIC on the electronic and structural properties of Cu oxides of various composition and dimensionality. It describes correctly the magnetic and insulating (anywhere from semiconducting to high-insulator) character of YBa$_2$Cu$_3$O$_{7}$ monoclinic CuO, GeCuO$_3$, Ca-doped YCuO cuprate, all of which are metals and non magnetic in GGA or LDA. This is further strong circumstantial evidence supporting the use of VPSIC as reference for the other methods in CCTO, even if one were to gloss over the experimental evidence discussed above.

### C. Quasiparticle corrections

It has long been customary to estimate quasiparticle energies as density-functional eigenvalues supplemented by “self-energy corrections”. In many materials, these corrections are dominated by a “scissor operator”, i.e. a $k$- and energy-independent relative shift of conduction and valence bands. A simple empirically-determined scissor correction is $\Delta=\frac{\Delta_0}{\varepsilon_\infty} \varepsilon_\infty$ eV. CCTO has a high-frequency dielectric constant $\varepsilon_\infty=\varepsilon_1(\omega=0)=12.6$, so the correction is $\Delta\sim0.7$ eV. The resulting total minimum gap
is roughly 0.85 eV, essentially in the VPSIC (and experimental) ballpark. Thus, the VPSIC and empirical scissor give similar corrections to local functionals, despite being completely unrelated.

Next we calculate the same sort of correction using \( G_0 W_0 \) non-selfconsistent quasiparticle energies. The latter calculation is rather difficult to converge in general, and particularly for this large system. We use a softer O potential enabling a cutoff of 280 eV, which does not seem to affect the eigenvalues much. The use of the specialized potential PAW sets provided with VASP for GW calculations is prevented by their large energy cutoff; this should not be a serious problem, as the standard PAWs we use do contain high-energy projectors and should perform rather well in the low-energy range we deal with here.) To assess convergence in \( k \) and in the number of bands for the virtual-transitions summation we used \( 2 \times 2 \times 2 \) and \( 4 \times 4 \times 4 \) \( k \)-grids and between 256 and 4092 bands. For a typical bulk material the latter choice would be overkill, but our system has of order 130 occupied bands per spin channel, so this setting seems necessary. Note also that that energy convergence in GW is more critical than in the standard joint-DOS calculations in the previous Sections: in the latter, unoccupied bands are only used in the Kronig-Kramers relation, whereas in GW they enter the evaluation of all energy-dependent parts of the self-energy.

V. SUMMARY

We examined the electronic structure of \( \text{CaCu}_3\text{Ti}_4\text{O}_{12} \) as obtained via several different density-functional based methods, and proposed a new interpretation of experiments to the effect that four distinct transitions contribute to the spectrum. The comparison of results from VPSIC calculations with experiment is satisfactory, especially after we account for the effects of spin disorder, which does not close the fundamental gap but suppresses the intensity of the fundamental transition. GGA+U and HSE at the standard values of their internal parameters overestimate drastically the fundamental gap, hence the conclusion that their corrections to the position of the flat Cu \( d \) bands should be more “screened” than they are. On the other hand, the corrections to local- or semilocal- functional eigenvalues provided by VPSIC are close to those of non-self-consistent one-shot GW, suggesting that much of the ”beyond-local” correlation is indeed provided by VPSIC with similar accuracy, at least in this material.

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The 'Mott' label is used here in the restricted sense of an odd-electron-count insulators (typically undergoing magnetic order-disorder transitions) to describe which one must describe reasonably electron interactions across energy scales from direct interaction, of order 1-10 eV, to exchange (Hund, super, double) of order 0.1-1 eV, and structural distortions, of order 0.01-0.1 eV. Within first-principles approaches, only many-body perturbation-theory, hybrid-density-functional, and self-interaction-corrected density-functional methods seem able to reach this level of accuracy (of course, dynamical mean field theory does also, although it generally uses empirical inputs in the self-energy and, especially, about structure); all of these methods remove self interaction from local density functionals—hence our interest in them in the present context.