

Identification of Potential Photovoltaic Absorbers Based on First-Principles Spectroscopic Screening of Materials

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There are numerous inorganic materials that may qualify as good photovoltaic (PV) absorbers, except that the currently available selection principle—focusing on materials with a direct band gap of ~ 1.3 eV (the Shockley-Queisser criteria)—does not provide compelling design principles even for the initial material screening. Here we offer a calculable selection metric of “spectroscopic limited maximum efficiency (SLME)” that can be used for initial screening based on intrinsic properties alone. It takes into account the band gap, the shape of absorption spectra, and the material-dependent nonradiative recombination losses. This is illustrated here via high-throughput first-principles quasiparticle calculations of SLME for ~ 260 generalized $I_p\text{III}_q\text{VI}_r$ chalcopyrite materials. It identifies over 20 high-SLME materials, including the best known as well as previously unrecognized PV absorbers.

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Most currently used solar-absorbing photovoltaic materials such as Si, GaAs, and CuInSe₂ have been discovered accidentally, and were subsequently improved incrementally over tens of years, at significant research and development cost. Indeed, databases listing all previously made inorganic crystals contain a vast number of candidates [e.g., $\sim 140\,000$ entries in the inorganic crystal structure database (ICSD) [1]], yet almost none were ever systematically screened for potential PV significance. A contributing factor to this state of affairs is the absence of suitable “selection metric” which can systematically drive materials screening. Heuristically established selection criteria of good absorbers have generally relied on favoring direct-gap materials over indirect gap materials, a distinction based on the wave vector momentum of the initial and final states across the gap. This important distinction, motivated by the need to have strong absorption that enables the use of a small amount of material (i.e., thin films), is insufficient. Indeed, it fails to recognize the fact that some direct-gap materials might have a dipole-forbidden (DF) direct transition lower in energy than the dipole-allowed (DA) direct transition [2–5], so being direct does not guarantee good absorption. Likewise, indirect gap materials with properly positioned higher energy DA transitions might prove efficient. The classic and almost universally used predictor of PV cell efficiency due to Shockley and Queisser (SQ) [6] depends only on material’s band gap, offering but a very rough selection criterion for good PV materials—an optimal gap of ~ 1.3 eV, no matter whether it is direct or indirect. This criterion alone has proven over the years to be insufficient as numerous materials with this gap are poor PV absorbers.

PV device efficiency (η) represents a complex convolution of thermodynamic, defect structure and optical characteristics, in addition to considerations such as fabrication, element abundance, toxicity, and cost. Given the vast number ($\sim 10^5$) of inorganic materials that may need to be eventually screened, we look for an initial filter based on the intrinsic spectroscopic and thermodynamic properties of the absorber, postponing scrutiny of imperfections (defects, fabrication-induced effects) and economic factors until after the initial field of candidates has been significantly narrowed down. We use the “spectroscopic limited maximum efficiency (SLME)” selection metric, which accounts, within the thermodynamic “detailed balance” approach for (i) the existence of various energetic sequences of DA, DF and indirect band gaps (denoted by E_g^{da} , E_g^{df} , and E_g^i , respectively), (ii) the specific shape of the absorption near the threshold, and (iii) the dependence of radiative recombination losses on the energy separation between the minimum gap E_g (allowed or not) and E_g^{da} . The SLME captures the leading physics of absorption, emission, and recombination characteristics, resolving a spread of different efficiencies for materials having the same gap. The spectroscopic quantities used in these three factors are obtained from first-principles quasiparticle calculations. It is illustrated here for ~ 260 generalized $I_p\text{III}_q\text{VI}_r$ chalcopyrite materials in ICSD, revealing over 20 high-SLME materials including the currently recognized best solar absorbers, yet adding a few previously unrecognized and potentially good PV absorbers.

We first classify materials into four “optical types,” based on the relative order of the direct-allowed, direct-forbidden, and indirect transitions, as illustrated

schematically in Fig. 1. “Optical type 1 (OT1)” is the case when the DA direct gap is the lowest energy transition and the next direct-but-forbidden transition is above it (i.e., $E_g^{da} \leq E_g^{df}$). “Optical type 2 (OT2)” is the case where the lowest transition is direct-but-forbidden, i.e., when $E_g^{da} > E_g^{df}$. Accordingly, “optical type 3 (OT3)” and “optical type 4 (OT4)” are two types of indirect gap materials, corresponding to cases with $E_g^i < E_g^{da} \leq E_g^{df}$ and $E_g^i < E_g^{df} < E_g^{da}$, respectively. Each of these four optical types has different characteristic absorption profile near threshold, depending on the order and energy separation between allowed and forbidden states. Subsequent, quantitative spectroscopic calculations (below) will demonstrate this classification on a class of materials.

The SLME is generalized from the SQ limiting efficiency. The power conversion efficiency [7] of a thin film solar cell depends on the fraction of the radiative electron-hole recombination current (f_r) and the photon absorptivity [$a(E)$]. SLME improves upon the SQ efficiency formula in the description of both f_r and $a(E)$. SQ efficiency assumes $f_r = 1$; i.e., the radiative recombination is the only recombination process for all optical types of materials. This could be a good approximation for OT1 materials such as GaAs [8] where radiative recombination dominates. However, for other types of materials where E_g^{da} is not the minimum band gap, the nonradiative recombinations (e.g., Auger recombination) is frequently much more significant [9,10] (i.e., $f_r \ll 1$). Here in SLME, we approximate $f_r = e^{-\Delta/kT}$, where k is the Boltzmann constant, T is the temperature, and $\Delta = E_g^{da} - E_g$, near absorption threshold of a pure semiconductor [11]. This form we chose is inspired by the Boltzmann formula for the relative number densities of atoms in the different excited energy states in thermal equilibrium with a black-body radiation field. Clearly for OT1, $\Delta = 0$ and $f_r = 1$ as in SQ limit, while for other types of materials, $\Delta > 0$ and f_r decreases exponentially as Δ . It is expected that the radiative recombinations happen mainly across E_g^{da} . When E_g^{da} shifts up by Δ , the radiative recombination rate is reduced by a factor of $e^{-\Delta/kT}$, but it still has to balance the incoming light in equilibrium. Therefore, a

smaller f_r here actually is an indicator of higher nonradiative recombination loss. On the other hand, it also means that the more radiative recombination loss (larger f_r) relative to nonradiative recombination loss, the better for PV absorber [12].

For photon absorptivity, SQ efficiency assumes a step-function absorptivity [i.e., $a(E) = 1$ for $E \geq E_g$ and 0 for $E < E_g$] for all materials. In SLME, we take $a(E) = 1 - e^{-2\alpha(E)L}$, where L is the thickness of the thin film with a zero-reflectivity front surface and unity-reflectivity back surface [13]. The $\alpha(E)$ is the calculated absorption coefficient from first principles. Thus, different optical types also manifest different $a(E)$ through their absorption coefficient $\alpha(E)$. In addition, SLME also uses the standard AM1.5G flat-plate solar spectrum at 25 °C [14]. The way we combine detailed balance with the fraction f_r of radiative recombination loss and the absorptivity $a(E)$ to get SLME is described in the Supplemental Material [15].

The required inputs for SLME calculations are the band gaps and absorption spectrum. These quantities are calculated (see details in [15]) based on *GW* approximation [16] for the electron’s self-energy. The method has been widely and successfully applied in first-principles quasiparticle electronic-structure calculations for many materials [17–19]. It enables direct comparison with experimental photoemission or inverse-photoemission measurements. Out of many *GW* schemes, we choose to apply *GW* approximation perturbatively on the top of the wave functions and energy eigenvalues calculated from a generalized Kohn-Sham scheme with the hybrid exchange-correlation functional HSE06 [20], i.e., $G_0W_0 + \text{HSE06}$ [21]. Within this scheme, it has been shown that for a variety of materials (even those with shallow *d* states), the calculated excited-state properties such as band gaps agree well with experiment [21,22]. Our $G_0W_0 + \text{HSE06}$ predicted minimum band gaps for some I-III-VI (I = Cu, Ag) compounds have an average error of less than 12% with respect to experiments (see Fig. S1a [15]).

We illustrate our foregoing ideas by considering generalized I-III-VI chalcopyrite group materials, i.e., $I_p\text{III}_q\text{VI}_r$, where we use I = Li, Na, K, Rb, Cs, Cu, Ag; III = B, Al, Ga, In, Tl, Sc, Y; VI = O, S, Se, Te; and any stoichiometric ratios ($p:q:r$) reported in ICSD. This group includes the well-known PV absorbers such as CuInSe_2 , CuGaSe_2 , and their solid solution Cu(In, Ga)Se_2 . We consider here 256 reported compounds of this group [15], covering most of stoichiometries and structure types that have been documented in ICSD. So far, most studies of ordinary chalcopyrites have focused on compounds with (1:1:2) stoichiometry [23]. Figure S2 [15] shows the distribution of all integer stoichiometries reported in ICSD for this group, indicating that in addition to the most popular (1:1:2), some other stoichiometries, like (3:1:3), (1:3:5), etc., are also rather common but their physical material properties are mostly unknown [24]. As will be seen below,

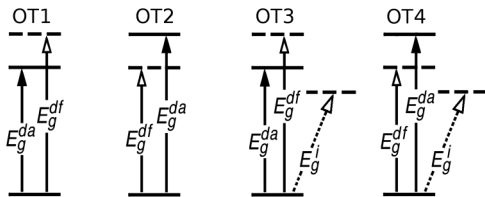


FIG. 1. Schematic diagrams of different optical types. Electric-dipole-allowed (forbidden) direct optical transition is denoted by a line with an arrow pointing to solid (dashed) horizontal line. Indirect states are shown as laterally displaced dashed lines.

some compounds with non-(1:1:2) stoichiometry could be also good for PV absorbers.

The calculated *GW* band gaps of considered 256 compounds are given in the Supplemental Material, Table S1 [15]. Figure 2 shows the *GW* gaps of 215 compounds [25] classified into four optical types. Some clear trends emerge here. (i) Within the same structure type, the band gap of materials decreases with increasing atomic number of one atom when the other two atoms are held fixed. For instance, for OT1 materials, $E_g^{da}(\text{LiAlSe}_2) > E_g^{da}(\text{LiGaSe}_2) > E_g^{da}(\text{LiInSe}_2)$. (ii) The optical types can change if the stoichiometry changes within the same element set. For example, Cu_3TlSe_2 (OT3) \rightarrow Cu_5TlSe_3 (OT4) \rightarrow Cu_7TlSe_4 (OT1) \rightarrow CuTlSe_2 (OT2). (iii) For the same compound, the minimum band gap (E_g) may vary by more than 2 eV in different crystal structures, whether or not the optical type changes. For example, for NaTiO_2 , $E_g = 0.07$ eV in the space group of #225 (OT4) and 2.27 eV in #166 (OT2). For LiInO_2 (OT3), $E_g = 0.19$ eV and 4.05 eV, respectively, in two structures of the same space group (#141). (iv) All reported $\text{I}_3\text{III}_1\text{VI}_3$ materials have small differences (less than 0.2 eV) between E_g^{df} , E_g^{da} , and E_g^i , except Li_3BO_3 which has a 0.43 eV difference between E_g^{da} and E_g^{df} .

Different optoelectronic applications may require different optical types. For example, transparent conductors benefit from a large transparency band gap (E_g^{da}), while the gap that decides dopability (the minimum gap E_g , whether allowed or not) can be much lower [26,27]. Thus, OT2 with small E_g^{df} and large E_g^{da} , and indirect gap materials (OT3 or OT4) with a small E_g^i and large enough E_g^{da} are preferred. Table S2 [15] lists 26 such potential transparent conductors with $3 \text{ eV} < E_g^{da} < 5 \text{ eV}$ and $E_g^{da} - E_g > 0.5 \text{ eV}$ found in Fig. 2, including well-known transparent conductors such as $\text{Cu}_1\text{III}_1\text{O}_2$ (III = Al, Ga, In) [28,29]. For light emitter and scintillator, OT1 with large dipole matrix element across E_g^{da} is preferred.

The SLME definition shows that it depends also on thin film thickness (L). Figure 3 illustrates that SLME increases as L increases. At very small L , due to weak absorptivity, the SLME of AgInTe_2 and Cu_7TlS_4 and their SLME difference are also small. At very large L , the SLME of these two OT1 materials approach the same SQ efficiency limit since they have the same E_g , and hence the SLME difference due to different characteristic absorption spectra also disappears. Therefore, to include the effect of the material-dependent spectroscopic properties in SLME, a reasonable size of L should be used and we choose here $L = 0.5 \mu\text{m}$.

Figure 4 shows calculated SLME for generalized I-III-VI thin film materials with thickness $L = 0.5 \mu\text{m}$ resolved into ‘‘optical types.’’ The SQ efficiency limit under AM1.5G solar spectrum is shown as the solid line and depends universally only on E_g for all optical types,

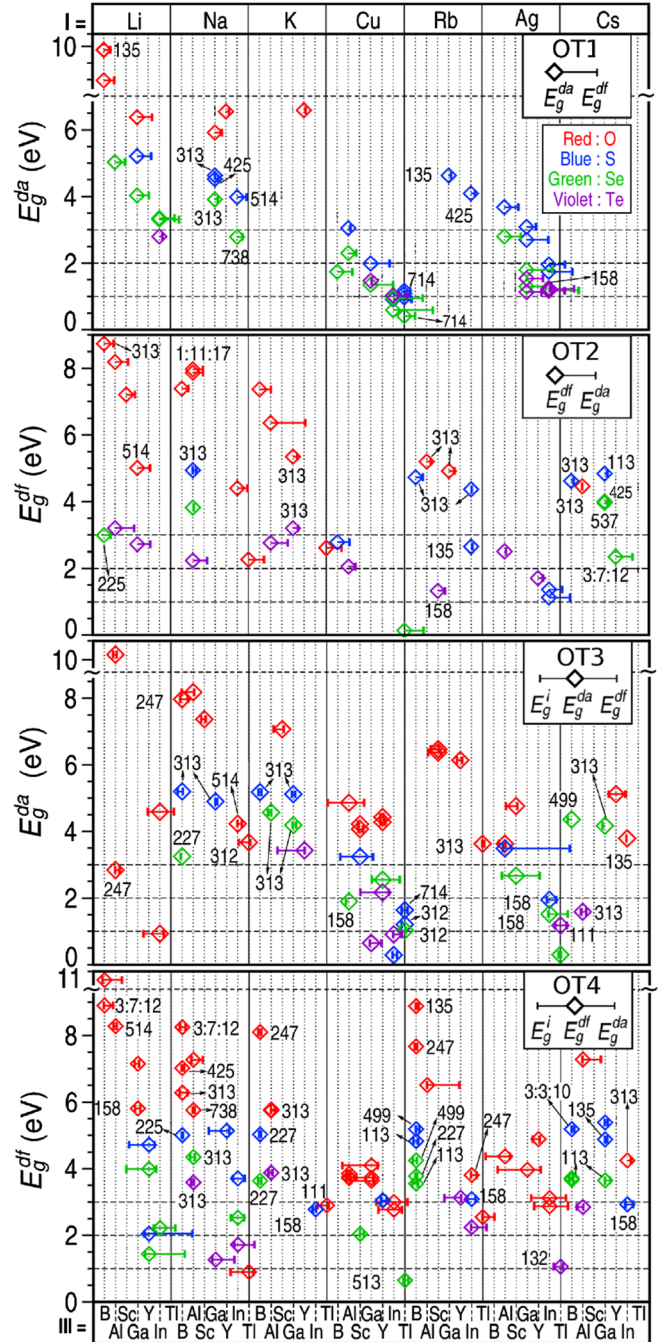


FIG. 2 (color online). *GW* band gaps of generalized I-III-VI chalcopyrite materials. At each point, the right horizontal bar represents $|E_g^{da} - E_g^{df}|$, and the left horizontal bar corresponds to $E_g^{da} - E_g^i$ (OT3) or $E_g^{df} - E_g^i$ (OT4). Each block separated by vertical dotted lines has a width of 0.5 eV. All stoichiometric ratios other than (1:1:2) are marked. Different materials with the same chemical formula are not distinguished.

predicting that the best gap for a PV absorber is 1.34 eV, at which $\eta_{\text{SQ}} = 33.7\%$. Our approach reveals instead a broad distribution of η values even around the same E_g , depending on the ‘‘optical types’’ and absorption spectra. For instance, AgInTe_2 , Cu_7TlS_4 , and CuYTe_2 have almost

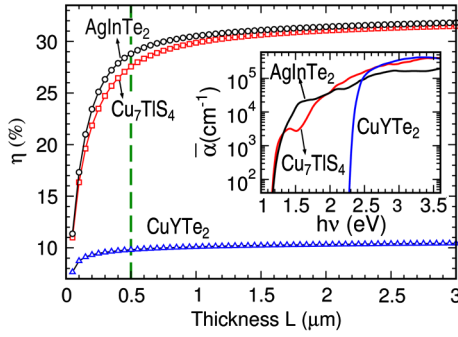


FIG. 3 (color online). The SLME as a function of thin film thickness for AgInTe_2 , Cu_7TlS_4 , and CuYTe_2 . The vertical dashed line indicates the thickness adopted in Fig. 4. The inset shows their absorption spectra.

the same minimum gap (1.17 eV), but their SLMEs vary significantly, being 27.6%, 22.6%, and 7.5%, respectively. From the inset of Fig. 3, it can be seen that the SLME difference between AgInTe_2 and Cu_7TlS_4 originates from different onset absorption spectra. For CuYTe_2 , the E_g^{da} is about 1 eV larger than E_g , i.e., $\Delta = 1$ eV, and hence the nonradiative recombination loss dominates. This large Δ leads to a much smaller overlap between absorption spectrum and solar spectrum. Hence, although the absorption near E_g^{da} (2.2 eV) is very strong in CuYTe_2 , the SLME is still rather small. Therefore, a material with the minimum gap being around 1.0–1.5 eV does not necessarily mean it is a good PV absorber.

From Fig. 4, we can find that there are about 25 materials with SLME higher than 20% (see Table S3 [15] for details). These high-SLME materials have the band gaps ranging from 0.8 to 1.75 eV. Most of them—18 out of 25—are OT1.

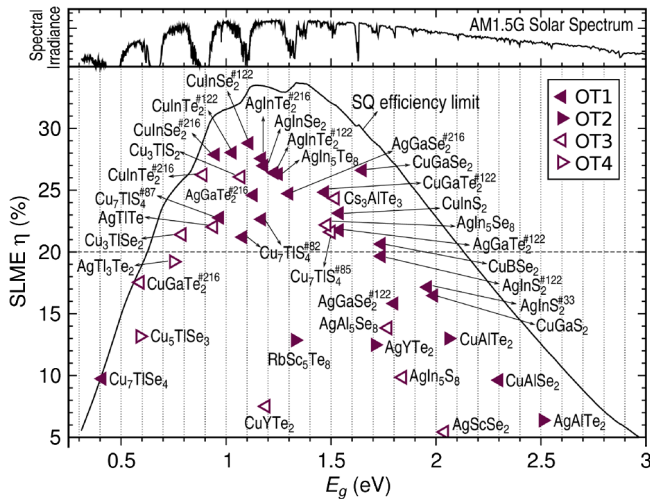


FIG. 4 (color online). SLME (η) vs the minimum gap E_g for generalized I-III-VI chalcopyrite materials at $L = 0.5 \mu\text{m}$. The compounds with SLME $< 5\%$ are not shown. The shown space group number (superscript) is used to distinguish different materials with the same chemical formula.

Seven of them are OT3. None of them have been found to be OT2 or OT4. The common character among these OT3 materials is that E_g^i is only slightly smaller than E_g^{da} , i.e., small Δ . For example, $\Delta = 0.07$ eV for Cs_3AlTe_3 , and 0.14 eV for Cu_3TlS_2 . Relative to OT1, this small Δ may cause higher joint density of states (DOS) at energies near E_g^{da} and good dispersive bands around the gap edges, and hence could lead to stronger onset optical absorption. OT2 and OT4 materials, both with $E_g^{\text{df}} < E_g^{\text{da}}$, are least favorable for PV absorbers.

Our predicted high-SLME materials in Fig. 4 include current best thin film solar absorber materials used in industry such as CuInSe_2 , CuGaSe_2 , and CuInS_2 . The top candidate is CuInSe_2 [30]. Its SLME is 28.8% at $L = 0.5 \mu\text{m}$, which exceeds the highest efficiency (20%) currently reported in experiments [31–33]. It is gratifying that the SLME flashes out in addition to CuInSe_2 , CuGaSe_2 , and CuInS_2 , also seven other high-SLME materials, namely, CuInTe_2 [34], CuGaTe_2 [35], AgInS_2 [36], AgInSe_2 [37], AgInTe_2 [38], AgGaSe_2 [39], and AgGaTe_2 [40], that have been found experimentally to be reasonable solar absorbers but are much less studied. Most of these previously recognized absorber materials within this group have (1:1:2) stoichiometry. Here we find that materials with other stoichiometries (e.g., AgIn_5Se_8 , Cs_3AlTe_3) also have high SLME.

Interestingly, it is found that all six high-SLME Cu-Tl-VI materials (i.e., four Cu_7TlS_4 , one Cu_3TlS_2 , and one Cu_3TlSe_2) are in non-(1:1:2) stoichiometry and contain only Tl of +1 oxidation state. Since Tl is highly toxic, these Tl-containing materials might be specifically disfavored in practical application. However, it indeed suggests that other similar high-SLME materials may be derived by replacing Tl^{1+} with other nontoxic elements in the 1+ oxidation state. Table S4 [15] summarized our identified $\text{Cu}_p(\text{I}, \text{II})_q\text{VI}_r$ materials that have SLME more than 20%, not involving Tl.

In summary, the strategy adopted in this work involved (i) recognizing a broad partitioning of materials into different “optical types”; (ii) developing a generalized selection metric SLME that considers different optical types and material-dependent nonradiative recombination loss; and (iii) testing the idea on a couple of hundred generalized I-III-VI chalcopyrites using high-throughput first-principles spectroscopy calculations and identifying potential PV absorber materials that from the point view of absorption are comparable to CuInSe_2 in the same group. Our identified high-SLME materials within this group include almost all currently using PV absorber materials as well as those that have been testified to be promising in experiment. It suggests that as an initial filter, our proposed SLME is indeed very effective for selecting good potential PV absorber materials [41]. The strategy of combining advanced selection metric with high-throughput first-principles evaluation of the spectroscopic input data could enable

identification of hitherto overlooked, promising candidate materials in different optoelectronic technology areas.

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