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Electronic properties of TlFe_2Se_2 using density functional theory

Doping and anion height are found to drastically affect the electronic structures of superconductors [1-2]. In this study, we investigated the composition-dependence of the electronic properties of the newly discovered intercalated TlFe_2Se_2 superconductor using density functional theory. We calculated the electronic structures of $\text{Tl}_x\text{Fe}_2\text{Se}_2$ with various Tl concentrations ($x = 1.00, 0.75$ and 0.50) using a $2 \times 2 \times 1$ supercell. Fractional coordinate z_{Se} of Se which essentially controls the anion height are taken from “relaxed” nonmagnetic and magnetic configurations of TlFe_2Se_2 which have values of $z_{\text{Se}} = 0.341 \text{ \AA}$ and 0.348 \AA , respectively. We also used the experimental value of $z_{\text{Se}} = 0.357 \text{ \AA}$ taken from [3], which is higher than the simulated ones. We also added a hypothetical value of $z_{\text{Se}} = 0.364 \text{ \AA}$ which is the highest among the fractional coordinate values used. All calculations of the electronic structures are done using QUANTUM ESPRESSO [4]. Generalized gradient approximation of Perdew-Burke-Ernzerhof [5] is used for exchange-correlation potentials.

The density of states (DOS) for various Tl content has shown metallic properties where states near the Fermi energy (E_F) are mostly from Fe- d states. This is consistent with the typical features of iron-based superconductors. For various fractional coordinates, z_{Se} , the DOS have also shown similar characteristics. Band structure calculations on the other hand revealed different results. For $x = 1.00$, no pocket is found around the zone center (Γ point) of the First Brillouin zone. For $x = 0.75$ and 0.50 on the other hand, hole-like pockets are being observed around the Γ point which is typical of Fe-based superconductors. The appearance of the hole-like pockets might be due to the shifting of the E_F towards lower energy when Tl content is reduced. This indicates a possible doping effect in this material (i.e. hole doping).

Furthermore, we also studied the possible three dimensionality of $\text{Tl}_x\text{Fe}_2\text{Se}_2$ with various Tl content by observing the Z point of the first Brillouin zone. As Tl content is reduced, a shift of bands is observed towards higher energies which resulted to an appearance of a shallow electron-like pocket around the Z point when $x = 0.75$. The electron-like pocket is mainly of Fe- $d_{xz}+d_{yz}$ character. This is not consistent though with the experiments where the observed electron-like pocket has Fe- $d_{xy}+\text{Se}-p_z$ character [6]. Looking at the band structure calculations for various z_{Se} , the electron-like band found around the Z-point above E_F for $z_{\text{Se}} = 0.341 \text{ \AA}$ seems to shift towards lower energies as the z_{Se} increases, which eventually crossed the E_F for $z_{\text{Se}} = 0.357 \text{ \AA}$. This electron-like pocket is now consistent with the experimental results where the orbital character is found to be of Fe- $d_{xy}+\text{Se}-p_z$. Increasing further the z_{Se} to 0.364 \AA , the bands split and a hole-like band is formed below E_F .

In conclusion, these results suggest that the Tl content plays a significant role in tuning the electronic properties of $\text{Tl}_x\text{Fe}_2\text{Se}_2$ where doping effect might occur with the appropriate value of Tl concentration. In addition, the anion height shows strong control of the band topology of this material.

References

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