Natural off-stoichiometry causes carrier doping in half-Heusler Þlled tetrahedral structures

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The half-Heusler Þlled tetrahedral structures (FTSs) are zinc-blende-like compounds, where an additional atom is Þlling its previously empty interstitial site. The FTSs having 18 valence electrons per formula unit are an emerging family of functional materials, whose intrinsic doping trends underlying a wide range of electronic functionalities are yet to be understood. Interestingly, even pristine compounds without any attempt at impurity/chemical doping exhibit intriguing trends in the free carriers they exhibit. Applying the Þrst principles theory of doping to a few prototype compounds in $\text{RdB}^{\times}C^{\text{IV}}$ and $A^{\text{IV}}B^{\text{IX}}C^{\text{V}}$ groups, we describe the key ingredients controlling the materialsÕ propensity for both intrinsic and extrinsic doping: (a) The spontaneous deviations from 1:1:1 stoichiometry reßect predictable thermodynamic stability of specificating phases. (b) Bulk ABC compounds containing 3elements in the position (ZrNiSn and ZrCoSb) are predicted to be naturally 3d rich. TheB = 3d interstitials are the prevailing shallow donors, whereas the potential acceptors (e.g., Zr vacancy and Sn-on-Zr antisite) are ineffective electron killers, resulting in an overall uncompensated character, even without any chemical doping. In these materials, the band edges are Ònatural impurity bandsÓ due to non-Daltonian off-stoichiometry, such as interstitials, not intrinsic bulk controlled states as in a perfect crystal. (c) BulkABC compounds containing & belements in the position (ZrPtSn, ZrIrSb, and TaIrGe) are predicted to be naturally rich and A poor. This promotes the hole-produciogn-A antisite defects rather than B-interstitial donors. The resultant type character (without chemical doping) therein is ÒlatentÓ for Sn and Sb; however, as the on-A hole-producing acceptors are rather deep and pheness is manifest only at high temperature or via impurity doping. In contrast, in TaIrGe (Ir, 5d), the prevailing hole-producing Ge-on-Ta antisite $(C\text{-on-A})$ is shallow, making it a read-type compound. This general physical picture establishes the basic trends of carriers in this group of materials.

DOI: [10.1103/PhysRevB.95.0852](https://doi.org/10.1103/PhysRevB.95.085201)01

I. INTRODUCTION: THE FAMILIES OF HALF-HEUSLER FILLED-TETRAHEDRAL STRUCTURE COMPOUNDS WITH 18 VALENCE ELECTRONS

NatureÕs most succinct ternary structures belong to the 1:1:1 equiatomicABCcompounds [EB]. They encompass two main families: (i) the eight valence-electron (octet) family EQ] containing the group&¹B^{II}C^V (e.g., LiZnP),A¹B^{III}C^{IV} (e.g., LiGaSi), $\mathsf{A}^{\mathsf{II}}\mathsf{B}^{\mathsf{II}}\mathsf{C}^{\mathsf{IV}}$ (e.g., MgSrSi), and $\mathsf{A}^{\mathsf{II}}\mathsf{B}^{\mathsf{III}}\mathsf{C}^{\mathsf{III}}$ (e.g., MgYGa) and (ii) the 18 valence-electron family $[$ DI4] that appears in a few chemical groups, such $AB^{\times}C^{\vee}$ (e.g., ScPtSb) $A^{\text{IV}}B^{\text{X}}C^{\text{IV}}$ (e.g., ZrNiSn), $A^{\text{IV}}B^{\text{IX}}C^{\text{V}}$ (e.g., TiCoSb), andA^VB^{IX}C^{IV} (e.g., TaIrGe [2Ð14]). These are $\dot{\text{O}}$ multifunctional materials $\dot{\text{O}}$ \parallel , due to their capacity to host a rich variety of 3d, 4d, and 5d transition metal elements and because of their appearance in numerous (over 40) crystal structure types, as summarized in $Ré2$, based on the Inorganic Crystal Structure Database (ICSD5). The combination of structures and elements provides opportunities for tuning and designing electronic band structures, spin-orbit

B. The chemical stability Þeld reveals tendencies towards off-stoichiometry and identiÞes the leading charge neutral defects

The formation of ABC ternary under a given set of chemical potential condition with μ _{A,B,C} < 0 requires that μ $_A$ + μ $_B$ + μ $_C$ = H (ABC), where H (ABC) is the compound formation energy. Projection of this equation onto the μ A, μ B plane leads to a chemical potential triangle frame μ _A + μ _B > H (ABC), with μ $_{A,B}$ <

IV. RESULTS: DEFECT CALCULATIONS ON ABC COMPOUNDS

A. Outline of the salient features of the method: Defect formation energy, charge transition energy, equilibrium Fermi energy, and carrier concentrations

The central quantities calculated here are (i) the defect formation energies H $(O,q,\mu$, $E_F)$ for various charge states q, chemical potentialp, and Fermi energy value_F and (ii) the defect charge transition level $(D,q | q)$ between charge statesq andq .

The formation energ $\forall I$ (D,q,µ, E_F) for defect D in charge statesq depend linearly on the atomic chemical potential $\{ \mu \}$

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FIG. 9. Isosurface plot of the majority carrier concentrations as a function of excess chemical potertials dB elements within the allowed stability triangle region for four half-Haussler compounds: ZrNiSn (a) and (b), ZrPtSn (c) and (d), ZrCoSb (e) and (f), and ZrIrSb (g) and (h). The results are shown at two temperatures: at growth condlition 850 C) and at room temperature from quench after growth. The letters R and P denote the chemical potential condition, where high and low majority carrier concentration occurs, respectively. R and P represent, respective \mathbb{B}_r -rich andB-poor condition forB = 3d compoundsÑZrNiSn and ZrCoStG-rich andC-poor conditions fo $B = 5d$

functional (mixing 25% of exact exchange) predicts a higher ionization level for Ni interstitial, above the CBM $[Fi6(a)]$ $[Fi6(a)]$, due to effect from enhanced Hartree-Fock exchange. However, the 25% of exact exchange is only an approximation used to eliminate self-interaction corrections. We feel that our uncertainty for the Ni $(D+1)$ charge transition level can be 0.1 eV, considering the effect of exchange on band gap opening and on eigenvalues of in-gap states. We, therefore, consider two computational scenarios allowing for 0.1 eV uncertainties, as shown by poinR and L in Fig. $6(a)$. The two scenarios lead to completely different physical behavior in temperature dependence of carrier density. In case (i), the Ni interstitial completely ionizes and its $(1/0)$ transition level is above the CBM [theR point in Fig. $6(a)$]. Thus, then-type carrier density remains nearly constant with temperature (\times 10²⁰ cm^{S3}, the dashed red line in Fig.0. While in case (ii), the Ni interstitial $(+1/0)$ transition level lies slightly below the CBM [e.g., E_{CBM} Š0.07 eV, theL point in Fig. $6(a)$], the Ni interstitial only partially ionizes due to thermal excitations (10% of Ni interstitials ionize at room temperature from simulation); therefore, the carrier density increases almost linearly with temperature (the solid red line in Fig.). When the donor is resonant, we predict a high carrier density in ZrNiSn (10^{19} cm^{83} at room temperature). This result from case (iii) is consistent with the observed line \overline{a} rdependence of the electrical conductivity in undoped ZrNiSn samples, 66 , so it is the preferred scenario.

4. Magnetism of ionized interstitial Ni

Ni interstitials have the potential to create magnetic moments in both fully ionized and partially ionized cases. As a magnetic impurity, the Ni interstitial possesses a localized moment of 0.8 μ _B from the x² Š y² 3d orbital. For magnetism in the dilute doping limit, it was known that p

2. The Sn-doped ZrCoSb (p type)

HSE results show that Sn-on-Sb is the dominant defect, having the lowest formation energy, but its acceptor transition level ($0'$ 1Ð) is rather deep (Fi[g.](#page-8-0) Hence, the resulting hole concentration is low at room temperature and relatively high only at growth condition, 850C. This case illustrates latent p-type compound dominated by an uncompensated, deep, hole producer with low formation energy (Sn-on-Sb antisite defect), which can only generate small amount of hole carriers at room temperature (see Appendix for details).

E. Effect of A atom selection inABC compounds with either $B = 3d$ or $B = 5d$ elements (the IV-X-IV and IV-IX-V groups)

Although we expect an approximately similar carrier doping behavior in the group on IV NiSn , AIV PtSn,AIV CoSb, andA^{IV} IrSb compounds withA^{IV} = Ti and withA^{IV} = Zr group, we are aware of the difference among the two groups. The chemical stability Þeld of the^{IV} = Ti group, as shown in the Þrst column of Fig 3 , appears to be larger in area and more extended to the -rich condition than its $\mathbf{A}^{IV} = \mathbf{Z}$ r and Hf counter parts. Recently, Wambach al. [36] measured the thermopower factor in the Ti-Ni-Sn compositional space using thin Þlm growth technique with combinatorial approach.

FIG. 12. Fundamental band gaps (in electron volt from HSE calculation) of a group of 18 valence-electron half-Heusler compounds in four prototype chemical groups^{IV} B^XC^{IV}, A^{IV} B^{IX}C^V, A^{III} B^XC^V, andA^V B^{IX}C^{IV}. The ternary compounds with cubic structure (θ) are nonmetal (shaded in green), whereas those in noncubic structures are metal (shaded in blue). Those that are predicted unstable are denoted l minus sign $\check{\mathfrak{G}}$).

50 cubic half-Heusler compounds reveals that thand structures are energetically unfavorable, about 0.7 eV per atom higher than the ground state (Fig.), indicating their non-existence.

2. Compound formation enthalpies and competing phases

The Þtted elementary reference energy method (FERE [73]) in conjunction with DFT has been successfully applied to compute compound formation enthalpies in cubic half-Heusler compounds and in their computing phases ([Tab](#page-17-0)les and [III\)](#page-18-0). The results show signibcant improvement over the plain GGA calculations, and are subsequently used in carrier doping studies to constrain the chemical potential stability Þeld of the host compound.

3. Concept and computational formula for defects and doping

Defect formation energis deÞned as the energy cost to create a point charged defect in an inÞnite lattice space (i.e., at dilute limit) through exchanging an atom and electron with the chemical reservoir and Fermi sea, respectively. To extract the defect formation energy from a Þnite supercell calculation, one uses the following formula

H (D,q,\mu, E_F) = { E(D,q) Š E_H}
$$
\pm
$$
 $\mu^0 + \mu$

$$
+ q(E_V + E_F) + H_{corr}
$$
 (A1)

where H $(D,q,\mu$, E_F) is the formation energy of defe α in charge stateq under the condition of elemental chemical potentialµ and the parametric electronic Fermi

TABLE III. Calculated formation enthalpies in units of electron volt per atom of Þve half-Heusler FTS (ZrNiSn, ZrCoSb, ZrPtSn, ZrIrSb, and TaIrGe) and their major competing ph[as](#page-19-0)es.

10. Doping trends in Þve ABC compounds

(Sn-on-Zr f - acceptors pins E_F^{eq} closer to VBM (E_V + 0.6 eV), thus introducing holes into the host. In $E2q$, the hole carrier density in Sn-doped ZrCoSb shows maximum at theor the dominant donors and acceptors in ZrNiSn, ZrCoSb, Co-poor and Sb-poor condition (the lower right corner in the ZrPtSn, ZrIrSb, and TaIrGe. The characteristics of the domstability Þeld), because the Co-poor condition suppresses Co-n ten, Ennes, and tan set the share construction of their primary opponents are interstitial donors which kill holes, while Sb-poor condition is beneÞcial to the creation of Sn-on-Sb substitution. An example consequence in conservation carriers, which serves as the key for the variation of charged impurity concentration as function of chemical potential condition is discussed in Fig. which is analogous to the total impurity concentration diagram Fermi level condition.
in Fig. 4 in Fig. 4. We summarize in $Fig23$ our defect computational results described in terms of defect formation energy, transition levels, a_{doping} information for the Þve compounds. The formation energy and carrier density are described at the equilibrium

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