Band gap tuning from an indirect EuGa$_2$S$_4$ to a direct EuZnGeS$_4$ semiconductor: syntheses, crystal and electronic structures, and optical properties†

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Four isostructural europium chalcogenides, EuZnGeS$_4$ (1), EuGa$_2$S$_4$ (2), EuIn$_2$S$_4$ (3) and EuIn$_2$Se$_4$ (4), have been synthesized using a high-temperature solid-state method. Single-crystal X-ray diffraction analysis indicates that they crystallize in the orthorhombic space group Fddd with $Z = 32$. Their structures feature an MQ$_4$ (M = Zn, Ge, Ga and In; Q = S and Se) tetrahedrally constructed 3D network, and Eu$^{2+}$ ions occupy the bicapped trigonal prismatic cavities. Compounds 1–3 have optical band gaps of 2.26, 2.32 and 2.37 eV, respectively. Theory calculations indicate that 2, with an indirect band gap, can be tuned to 1, with a direct band gap, via simple chemical substitution.

Introduction

Rare-earth chalcogenides have been extensively explored in the past twenty years because of their rich structures and potential applications in the fields of magnetism,$^1$ thermoelectricity,$^2$ photoluminescence,$^3$ second-order nonlinear optics,$^4$ and so on. Among them, europium chalcogenides have been paid much attention as Eu ions can be divalent or trivalent. It is well known that the Eu$^{2+}$ cation has a similar ionic radius to alkali-earth metal cations, so many Eu$^{2+}$ compounds have alkali-earth metal analogues. Recently, many alkali-earth metal chalcogenides have been studied as second-order nonlinear optical materials as they have high nonlinear optical coefficients and laser induced damage thresholds, such as Ba$_2$Sn$_4$I$_{15},^5$ Ba$_2$Sn$_3$I$_{16},^6$ Ba$_2$Sn$_n$Se$_{3+},^7$ BaHgSe$_2,^8$ BaHgS$_2$ [ref. 9] and BaCdSnSe$_{3+},^9$ which have encouraged many chemists and material scientists to explore them in the field of laser frequency conversion.

When checking the database, it was discovered that there are a large group of ternary or quaternary compounds can be uniformly written as II–III$^0$ (II–III$^0$)-I$^0$ or quaternary II–II$^0$(I$^1$)–IV–Ch$_4$ (I = Mg, Ca, Sr, Ba, Eu and Yb; II$^0$ = divalent transition metal; I$^0$ = Li, Na, Cu and Ag; III$^0$ = B, Al, Ga and In; III$^1$ = Yb, Sb and Bi; IV = Ge and Sn; Ch = S and Te).$^{10}$ The representative members of this family are listed in Table 1. What’s more, there are also many related II–III$^0$–Ch$_4$ compounds containing transition-metal reported, where II = divalent transition-metal Zn, Cd, Hg and Mn, III = Ga and In, and Ch = S and Se.$^{11}$ If take EuGa$_2$S$_4$ as the parent compound for this series, the sites of Eu and S can be also occupied by divalent alkali-earth metal or Yb, and Se or Te, respectively. The Ga sites can be replaced by the same group element B, Al and In. Specially, two Ga sites can be also replaced by two different elements to form quaternary compound II–II$^0$–IV–Ch$_4$ or II–I$^1$–II$^0$–IV–Ch$_4$, in which one site is occupied by tetravalent cations like Ge and Sn, the other site occupied by divalent transition-metal cations or monovalent coinage or alkali metal cations.

Stimulated by their versatile structure types and potential applications and our rich experience in rare-earth chalcogenides,$^{12}$ we recently synthesized four such compounds, EuZnGeS$_4$ (1), EuGa$_2$S$_4$ (2), EuIn$_2$S$_4$ (3) and EuIn$_2$Se$_4$ (4). Compounds 1, 3 and 4 can be viewed as the derivants of 2. Compound 1 is firstly obtained, 3 and 4 are new phases of EuIn$_2$S$_4$ and EuIn$_2$Se$_4$, respectively, different from the known Ccmm ones.$^{13}$ Compound 2 was only characterized using powder X-ray diffraction data.$^{14}$ Here, we studied their syntheses, crystal and electronic structures, and optical properties.

Experimental section

Materials

Eu$_2$O$_3$ (99.9%, Aladdin), Zn (99.99%, Aladdin), Ga$_2$O$_3$ (99.999%, Aladdin), In$_2$O$_3$ (99.9%, Aladdin), Ge (99.999%, Aladdin), S (99.95%, Aladdin), Se (99.999%, Aladdin), B powder (99%, Aladdin) and KI (99.0%, China Sinopharm Chem.).

Syntheses and analyses

All starting materials were used as received without further purification. Single crystals of the title compounds were...
Table 1 Known structure types of ternary II-II₃ (II-III)₄-Ch₄, or quaternary II-II₄(II₄)-IV₄-Ch₄ (II = Mg, Ca, Sr, Ba, Eu and Yb; II' = divalent transition metal; I = Li, Na, Cu and Ag; III = B, Al, Ga and In; III' = Yb, Sb and Bi; IV = Ge and Sn; Ch = S, Se and Te) compounds. All these data are from ICSD or Pearson’s crystal data.

| Compound         | Space group | Crystal system  | Pearson code | Phase prototype |
|------------------|-------------|-----------------|--------------|-----------------|
| MgAl₂S₄         | Pnma        | Orthorhombic    | oP28         | Mg₂SiO₄         |
| MgAl₂S₄         | R3m         | Trigonal        | hR7          | ZnIn₂S₄         |
| MgAl₂S₄         | R3m         | Trigonal        | hR7          | ZnIn₂S₄         |
| MgGa₂S₄         | C2/c        | Monoclinic      | mS84         | MgGa₂S₄         |
| MgIn₂S₄         | Fd3m        | Cubic           | cF56         | MgAl₂O₃         |
| MgIn₂Te₄        | I4/mcm      | Tetragonal      | tI14         | Cu₂HgI₄         |
| SrB₂S₄          | P4/nnc      | Tetragonal      | tP28         | SrB₂S₄          |
| SrB₂S₄          | R3          | Trigonal        | hR63         | SrB₂S₄          |
| SrAl₂S₄         | Cccm        | Orthorhombic    | oS28         | BaGe₄Se₄        |
| SrAl₂S₄         | Fddd        | Orthorhombic    | oF224        | EuGa₄S₄         |
| SrGa₂Te₄        | I4/mcm      | Tetragonal      | tI14         | CaIn₄Te₄        |
| SrCu₂GeSe₄      | Amma2       | Orthorhombic    | oS32         | SrCu₂GeSe₄      |
| SrCu₂SnS₄       | P3₁         | Trigonal        | hP24         | SrCu₂SnS₄       |
| SrCu₂SnS₄       | P3₁.21      | Trigonal        | hP24         | SrCu₂SnS₄       |
| BaB₂S₄          | Cc          | Monoclinic      | mS28         | BaB₂S₄          |
| BaAl₂S₄         | Pa3         | Cubic           | cP84         | BaAl₂S₄         |
| BaAl₂Se₄        | P4/nnc      | Tetragonal      | tP28         | BaAl₂Se₄        |
| BaAl₂Te₄        | P4/nmc      | Tetragonal      | tP14         | BaAl₂Te₄        |
| BaSb₂B₄         | Pnma        | Orthorhombic    | oP28         | KBaP₀₄          |
| BaBi₂S₄         | C2/m        | Monoclinic      | mS28         | BaBi₂S₄         |
| BaAg₂GeS₄       | I4/m         | Tetragonal      | tI16         | K₃VO₄          |
| BaAg₂SnS₄       | Fd22        | Orthorhombic    | oF16         | BaAg₂SnS₄       |
| BaAu₂SnS₄       | P2₁2/1       | Orthorhombic    | oP32         | BaAu₂SnS₄       |
| BaCdSnS₄       | Fddd        | Orthorhombic    | oF224        | BaCdSnS₄       |
| BaHgSnS₄       | Pnn2        | Orthorhombic    | oP28         | BaHgSnS₄       |
| EuNa₂SiSe₄      | R3c         | Trigonal        | hR48         | EuNa₂SiSe₄      |
| EuNa₂GeSe₄      | I4/m         | Cubic           | cI16         | Tl₃V₄S₆        |
| YbCaIn₄S₄       | Pnma        | Orthorhombic    | oP28         | Mg₂SiO₄         |

* As there are so many members can be classified to this family, only one compound was chosen as a representative for each type of structure. All these data are checked results from Inorganic Crystal Structure Data (ICSD) and Pearson’s Crystal Data (PCD).

obtained by solid-state reactions with KI as flux. The starting materials are stoichiometric mixture of Eu₂O₃, Zn, Ge and S for 1, Eu₂O₃, Ga₂O₃ and S for 2, Eu₂O₃, In₂O₃ and S for 3, and Eu₂O₃, In₂O₃, and Se for 4. A certain amount of boron powder and KI were added to each sample as reducing reagent and flux, respectively. Each sample has a total mass of 500 mg and 400 mg KI additional. The mixtures of starting materials were ground into fine powder in an agate mortar and pressed into pellets, followed by being loaded into quartz tubes. The tubes were evacuated to be 1 × 10⁻⁴ torr and flame-sealed. The samples were placed into a muffle furnace, heated from room temperature to 573 K in 5 h and equilibrated for 10 h, followed by heating to 923 K in 5 h and equilibrated for another 10 h, then heated to 1223 K in 5 h and homogenized for 10 days, finally cooled down to 573 K in 5 days and powered off. All the crystals of 1–4 stable in moisture and air were obtained and hand-picked under a microscope since the yields were not high, then washed using ethanol and water under ultrasonic wave, whose purities were confirmed by powder X-ray diffraction (PXRD) study. The PXRD patterns were collected with a Bruker D8 Advance diffractometer at 40 kV and 100 mA for CuKα radiation (λ = 1.5406 Å) with a scan speed of 5° min⁻¹ at room temperature. The simulated patterns were produced using Mercury v2.3 program provided by the Cambridge Crystallographic Data Center and single-crystal reflection data. A representative PXRD pattern of 3 corresponds well with the simulated one (Fig. 1), indicating it is phase-pure of the selected crystals. Semiquantitative microscope element analysis on the as-prepared single crystals was performed on a field-emission scanning electron microscope (FESEM, HITALCHI S-4800II) equipped with an energy dispersive X-ray spectroscopy (EDS, Bruker, Quantax), which confirmed the presence of each element with the approximate compositions of 1–4. The exact compositions were established from X-ray structure determination.

**Structure determination.** The intensity data set was collected on Bruker D8 QUEST X-ray diffractometer with graphite-monochromated MoKα radiation (λ = 0.71073 Å). The structures were solved by direct methods and refined by full-matrix least-squares techniques on F² with anisotropic thermal parameters for all atoms. All the calculations were performed with Siemens Shelleit-97 crystallographic software package.

The final refinement included anisotropic displacement parameters for all atoms and a secondary extinction correction. The crystallographic data, atomic coordinates and equivalent isotropic displacement parameter, and bond lengths are listed in Tables 2, S1 and S2, respectively. Their CIF documents have also been deposited with Fachinformationszentrum Karlsruhe, 76344 Eggenstein-Leopoldshafen, Germany [fax: +49-7247-808-666; e-mail: crysdata@fiz.karlsruhe.de] with depository number CSD-432057 for 1, CSD-432055 for 2, CSD-432056 for 3 and CSD-432058 for 4.

**UV-Vis-NIR diffuse reflectance spectroscopies.** The diffuse reflectance spectra were recorded at the room temperature on...
Table 2  Crystal data and structure refinement parameters of 1–4

| Chemical formula | EuZnGeS₄ (1) | EuGa₂S₄ (2) | EuIn₂S₄ (3) | EuIn₂Se₄ (4) |
|------------------|-------------|-------------|-------------|-------------|
| F₁₀              | 418.16      | 419.64      | 509.84      | 697.44      |
| T (K)            | 296         |             |             |             |
| Crystal system   | Orthorhombic|             |             |             |
| Space group      | Fddd        |             |             |             |
| a (Å)            | 12.234(4)   | 12.209(11)  | 13.016(3)   | 13.490(1)   |
| b (Å)            | 20.398(6)   | 20.445(18)  | 20.772(5)   | 21.748(1)   |
| c (Å)            | 20.682(6)   | 20.680(19)  | 21.076(5)   | 21.859(1)   |
| V (Å³)           | 5161(3)     | 5162(8)     | 5699(2)     | 6412.9(6)   |
| Dₑ/calcd (g cm⁻³) | 4.305      | 4.320       | 4.754       | 5.779       |
| μ (mm⁻¹)         | 19.076      | 19.048      | 16.166      | 31.487      |
| f(000)           | 6048        | 6048        | 7200        | 9504        |
| θ range (°)      | 2.18 to 25.50 | 2.18 to 25.49 | 2.08 to 25.48 | 2.01 to 27.50 |
| Meas. rens       | 5315        | 2843        | 9320        | 7650        |
| Indep. rens/Rint | 1203/0.283  | 1171/0.0303  | 1329/0.0581  | 1844/0.0403  |
| R₁ | R₂ | (I > 2σ(I))% | 0.0483/0.1245 | 0.0342/0.0841 | 0.0428/0.0878 | 0.0493/0.0666 |
| R₁ | R₂ | (all data)% | 0.0653/0.1279 | 0.0421/0.0894 | 0.0426/0.0878 | 0.0493/0.0666 |
| GOF on F²        | 1.129       | 1.145       | 1.090       | 1.093       |
| Δρmax/Δρmin c Å⁻³| 2.571/−1.815 | 3.368/−2.110 | 1.627/−2.332 | 2.042/−1.457 |

* R₁ = [|F₁₀| − |F₁|]/|F₁₀|; wR₂ = [w(F₂ − F₁)²]/[w(F₁)²]²/2.

Results and discussion

Crystal structure

Compounds 1–4 crystallize in the orthorhombic space group Fddd. The crystal structure of 1 is illustrated as a representative as they are isostructural. There are three Eu, one Zn, one Ge and four S atoms in the crystallographically independent unit. Its coordination geometry is shown in Fig. 2. All the three Eu atoms coordinated with eight neighboring S atoms to form EuS₈ bicapped trigonal prisms, Zn and Ge are fourfold-coordinated with four S atoms to form ZnS₄ or GeS₄ tetrahedron.

The structure can be viewed as layered if no consideration of the connection between Eu and S atoms (Fig. 3a). The layers parallel to the ac plane are constructed by the connection between ZnS₄ and GeS₄ tetrahedra (Fig. 3b). It can be observed that either ZnS₄ or GeS₄ tetrahedra form dimers via sharing edges, respectively, namely, [Zn₂S₆]₈⁻ or [Ge₂S₆]₈⁻ units. Both of which are isolated and alternately linked to form the 2D structure. Take the whole structure consideration, the structure of 1 can be described as Eu–S bonds constructed 3D structure, and Zn²⁺ and Ge⁴⁺ cations occupy the tetrahedral cavities (Fig. 4a). It is necessary to understand how the EuS₈ bicapped trigonal prisms connect if we want to know how the 3D structure forms. The Eu atoms arrange regularly along the c direction. Eu(1) atoms arrange in a line along the c direction, while Eu(2) and Eu(3) atoms alternatively arrange in another line along the c direction. As shown in Fig. 4b, each EuS₈ units has six neighboring EuS₈ units, four of them connect with the central one via sharing edges, while the other two connected via sharing

Fig. 2  Coordination geometry of 1.
corners. The former and latter have the Eu–Eu distances of 5.208(2) and 5.947(1) Å, respectively.

The Zn–S and Ge–S bond lengths in 1 are in the range of 2.247(4)–2.303(5) and 2.246(4)–2.315(4) Å, respectively, which are in good agreement with 2.243(2)–2.305(2) Å of Ga–S bonds in 2. These values are consistent with 2.242(1)–2.342(1) Å of Ga–S bonds in (K₃I)[SmB₁₂(GaS₄)₃]. The Eu–S bonds in 1 have the distances of 3.070(4)–3.112(5) Å, similar with those discovered in RbEuGeS₄.

As compounds 2–4 have more similar compositions. Here only the comparison is performed between 1 and 2. There are two Ga sites in 2, which can be substituted by one divalent and one tetravalent cations, respectively. So far, there are no quaternary II-III₂ (III-III⁰)-Ch₄ or II-II⁰ (I₂)-IV-Ch₄ compounds reported with Fddd structure. Compound 1 represents the first one with this structure. It is interesting to study which one of the two Ga sites will be replaced by divalent cation. First, it has to be mentioned that both of the two Ga atoms occupy 32h sites. Second, both Zn and Ge positions are close to that of Ga in periodic table of the elements, so it is almost impossible to distinguish them by atomic weights when solving the structure.

Third, bond-valence calculation was performed on 2 to help determine which site will be occupied by Zn²⁺ ion, and the other one by Ge⁴⁺ ion. If Ga(1) and Ga(2) atoms are replaced by Zn and Ge atoms, respectively, their bond valences calculated to be 2.368 and 3.267, respectively. If Zn and Ge exchange their sites, their bond valences are 2.329 and 3.331, respectively. Even the calculation precision is considered, it is still difficult to determine that Ga(1) site is occupied by Zn or Ge atom. The most reasonable solution is that Zn and Ge atoms statistically co-occupy both of two Ga sites according to the bond-valence calculation results. For simplicity and convenience, the structure of 1 is described above as Zn and Ge atom completely occupy one Ga site, respectively. Definitely, the molar ratio of Zn and Ge should be 1 : 1 to maintain electric neutrality.

**Optical properties**

The UV-Vis-NIR diffuse reflectance spectra of 1–3 are shown in Fig. 5. Their optical band gaps are determined to be 2.26, 2.32 and 2.37 eV, respectively, which are consistent with their orange or yellow colors. Comparing these values, it may be proposed that there will be no big band gap change when the Ga atoms in
EuGa$_2$S$_4$ are replaced by II + IV two types of atoms or by heavier III atoms like In atoms. This supposition may help design II-III$_2$(III-III)$_0$-Ch$_4$ or II-II$_0$(I$_2$)-IV-Ch$_4$ compounds with tunable band gaps.

**Theory investigation**

To investigate the electronic structures of 1 and 2, their band structures together with densities of states (DOS) computations based on the DFT theory were performed using Material Studio software. The calculated band structures along high symmetry points of the first Brillouin zone are shown in Fig. 6, from which it can be seen that the band gaps of 1 and 2 are calculated to be 1.32 and 2.69 eV (Table 3), respectively, the former one is underestimated but reasonable in view of the calculation precision. Both the lowest conduction band (CB) and highest valence band (VB) of 1 are located at $I$ point, indicating that 1 has a direct band gap. Different from 1, the CB and VB of 2 are located at $I$ and $X$ points, respectively, showing that 2 has an indirect band gap.

The total and partial densities of states (DOS and PDOS) of 1 and 2 are plotted in Fig. 7. For 1, the highest VB is mainly constituted of S-3p, and the lowest CB is mainly composed of Ge-4s and S-3p orbitals. The VBs ranging from 2.5 to 7.5 eV are primarily consisted of Ge-4p, and minor Zn-4s and S-3p orbitals. Briefly, the band gap of 1 is determined by the S-3p and Ge-4s orbitals, the respective valence orbitals of S$^{2-}$ and Ge$^{4+}$ ions. For 2, the highest VB is mainly constituted of S-3p, and the lowest CB is mainly composed of Ga-4s, S-3p and Ga-4p orbitals, so its band gap is determined by valence orbitals of S$^{2-}$ and Ga$^{3+}$ ions. The VBs

**Table 3** Experimental and calculated optical band gaps of 1–3

|                  | EuZnGeS$_4$ (1) | EuGa$_2$S$_4$ (2) | EuIn$_2$S$_4$ (3) |
|------------------|-----------------|-------------------|-------------------|
| Band gap (eV)    |                 |                   |                   |
| Experimental     | 2.26            | 2.32              | 2.37              |
| Calculated       | 1.32            | 2.69              | —                 |

EuGa$_2$S$_4$ are replaced by II + IV two types of atoms or by heavier III atoms like In atoms. This supposition may help design II-III$_2$(III-III)$_0$-Ch$_4$ or II-II$_0$(I$_2$)-IV-Ch$_4$ compounds with tunable band gaps.
from 0 to ~4.0 eV are primarily consisted of S-3p, Eu-4f and minor Ga-4p orbitals. Comparing the PDOS of 1 and 2, it can be concluded that the optical absorptions of 1 and 2 are mainly ascribed to the charge transitions from S-3p to valence orbitals of Ge$^{4+}$ or Ga$^{3+}$ cations.

It is well known that direct or indirect semiconductors have different applications in the optoelectronics research field. From EuGa$_2$S$_4$ to EuZnGeS$_4$, two Ga$^{3+}$ sites are replaced by one Zn$^{2+}$ and one Ge$^{4+}$ cations, which is a simple substitution but the parent structure EuGa$_2$S$_4$ can be maintained and the measured band gap have a little change. Together with the above discussion, it should be interesting to understand why semiconductor type can be changed while crystal structure and measured band gap keep stable. Besides, it seems the optical absorptions of 1 and 2 have little relationship with the valence orbitals of Eu$^{2+}$, which may give us some rules to explore II-III$_2$ (III-III) compounds containing divalent transition-metal, CdGa$_2$Se$_4$ and HgGa$_2$Se$_4$, have direct band gaps of 2.40 and 1.93 eV, respectively, close to our sulides. Besides, it is interesting to find that the calculated band gaps for compounds EuZnGeS$_4$, CdGa$_2$Se$_4$ and HgGa$_2$Se$_4$ with direct band gaps are much lower than their experimental values.

Conclusions

Four isostructural europium chalcogenides were synthesized using solid-state reactions. Their measured band gaps indicate that they are semiconductors. Interestingly, the band gap type can be tuned from indirect to direct when using one divalent Zn$^{2+}$ and one tetravalent Ge$^{4+}$ cations to substitute the two trivalent Ga$^{3+}$ cations sites in EuGa$_2$S$_4$. This study may give some experience how to tune structures and band gaps of II-III$_2$ (III-III) compounds containing divalent transition-metal, CdGa$_2$Se$_4$ and HgGa$_2$Se$_4$, have direct band gaps of 2.40 and 1.93 eV, respectively, close to our sulides. Besides, it is interesting to find that the calculated band gaps for compounds EuZnGeS$_4$, CdGa$_2$Se$_4$ and HgGa$_2$Se$_4$ with direct band gaps are much lower than their experimental values.

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