New Germanate and Mixed Cobalt Germanate Salt Inclusion Materials: $[(Rb_6F)(Rb_4F)][Ge_{14}O_{32}] and$ $[(Rb_6F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O_{30}F_2]$

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Abstract

of Single crystals $[(Rb_{6}F)(Rb_{4}F)][Ge_{14}O_{32}]$ and two new germanates, $[(Rb_{6}F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O_{30}F_{2}]$, were synthesized via high temperature RbCl/RbF flux growth. Both compounds crystallize in the cubic space group $F\overline{4}3m$ and possess the germanium framework of the previously reported salt inclusion material (SIM), [(Cs₆F)(Cs₃AgF)][Ge₁₄O₃₂], related to the Ge₇O₁₆ zeolitic family. These materials demonstrate the ability to accommodate a variety of salt-inclusions, and exhibit chemical flexibility enabling modifications of the framework through incorporation of Co. Alteration of the salt-inclusion led to intrinsic luminescence of [(Rb₆F)(Rb₄F)][Ge₁₄O₃₂] while modification of the framework resulted in an unanticipated Rb/Co salt/inclusion in [(Rb₆F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O₃₀F₂]. Fluorescence measurements were performed on [(Rb₆F)(Rb₄F)][Ge₁₄O₃₂]. First-principles calculations in the form of density functional theory (DFT) were performed for [(Rb₆F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O₃₀F₂] to elucidate its electronic and magnetic properties, and stability at 0 K.

Introduction

Salt inclusion materials, or SIMs, have attracted significant attention as potential nuclear waste forms,¹ and although a significant number of structures and compositions have been reported to date,^{2–10} their synthesis/discovery is not readily predictable. Most SIMs have been grown as single crystals out of molten salt fluxes and it has been shown that the salt inclusion typically consists of elements from the molten flux itself. Hence, by choosing different alkali and halide species for the melt that has led to one SIM composition, it is possible to obtain different compositions in a fairly predictable way. Similarly, one can attempt to modify not the salt inclusion, but the framework itself, by adding other framework forming species to the melt. In both cases the energetics i.e., heats of formation, will ultimately determine the ability to create the target phase. While crystals obtained from fluxes need to be relatively thermodynamically stable, they may in fact be metastable structures or compositions whose formation was governed by kinetic and/or steric factors.

Our group recently discovered and structurally characterized a new germanate framework SIM, [(Cs₆F)(Cs₃AgF)][Ge₁₄O₃₂], containing Cs₆F and Cs₃AgF salt inclusion species located in the large channels of the framework.¹¹ To further investigate this structure type and to explore the extent to which the salt inclusion and/or the framework of the structure could be chemically modified, we explored the synthesis of other germanate SIMs via flux crystal growth. Specifically, we focused on modifying the salt inclusion and on modifying the framework. Interestingly, as discussed within, the salt inclusion could easily be modified and, surprisingly, so could the framework via Co incorporation. The change in the salt inclusion resulted in intrinsic luminescence, while the cobalt incorporation resulted in an unprecedented Rb/Co salt inclusion as well as in the presence of cobalt in the framework itself.

In our previous report,¹¹ we focused on investigating the promising ion exchange properties of the germanium framework-based salt-inclusion. The Cs structure was found to undergo successful ion exchange with both K and Rb. In the current effort, a family of this structure type is created with the successful synthesis of two new compositions: (1) a Rb salt-inclusion with all germanium framework and (2) a mixed Ge/Co framework with Rb/Co site mixing. The cubic germanate framework has now demonstrated room temperature luminescence in $[(Rb_6F)(Rb_4F)][Ge_{14}O_{32}]$ and the capacity to incorporate 3*d* transition metals in $[(Rb_6F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O_{30}F_2]$.

Experimental

Reagents

GeO₂ (99.999%, Alfa Aesar), RbCl (Alfa Aesar, 99.8%), and RbF (99.1%, Alfa Aesar) were used as received for the synthesis of both compounds. For the synthesis of [(Rb₆F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O₃₀F₂], CoF₂ (Alfa Aesar, anhydrous powder, 98%) was used as received.

Synthesis

Single crystals of [(Rb₆F)(Rb₄F)][Ge₁₄O₃₂] were grown out of a eutectic RbCl/RbF flux. To a cylindrical silver crucible welded shut on one end, 2 mmol of GeO₂ were added and covered by 1.70 g of RbCl and 1.30 g of RbF. The silver crucible was crimped shut and placed upright in a programmable box furnace. The reaction mixture was heated at 300 °C/h to 900 °C, held at this temperature for 12 h, slow cooled to 400 °C/h at a rate of 6 °C/h, and then rapidly cooled to room temperature by shutting the furnace off. The solidified flux matrix was dissolved in hot water, aided by sonication, and the products were collected by vacuum filtration and dried with acetone. Colorless, irregular block-shaped crystals approaching 1 mm³ in size were obtained in approximately 30% yield along with large amounts of AgCl, which was removed from the target phase using a concentrated sodium thiosulfate solution.

For the synthesis of $[(Rb_6F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O_{30}F_2]$, 1 mmol of CoF₂ and 4 mmol of GeO₂ were layered beneath 3.42 g of RbCl and 2.42 g of RbF in a cylindrical silver crucible. The reaction mixture was heated at 300 °C/h to 850 °C, maintained at this temperature for 24 h, slow cooled to 400 °C at 6 °C/h, and then rapidly cooled to room temperature by shutting the furnace off. Once cooled to ambient temperature, the solidified RbCl/RbF flux was dissolved in distilled water, aided by sonication, and the resulting products were isolated by vacuum filtration. Bright green, irregular block-shaped crystals approximately 0.3 x 0.2 x 0.15 mm³ in size were obtained in more than 25% yield together with large amounts of powder.

Single-Crystal X-ray Diffraction (SCXRD)

X-ray intensity data from suitable crystals of $[(Rb_6F)(Rb_4F)][Ge_{14}O_{32}]$ and $[(Rb_6F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O_{30}F_2]$ were collected at 301(2) K using a Bruker D8 QUEST diffractometer equipped with a PHOTON 100 CMOS area detector and an Incoatec microfocus source (Mo K α radiation, $\lambda = 0.71073$ Å) [1]. The raw area detector data frames were reduced and corrected for absorption effects using the SAINT+ and SADABS programs.^{12, 13} Initial structural

models were obtained with SHELXT.¹⁴ Subsequent difference Fourier calculations and full-matrix least-squares refinement against F^2 were performed with SHELXL-2018 using the ShelXle interface.¹⁵ The data collection covered 100% of reciprocal space to $2\theta_{max} = 66.19$ and 75.50°, with an average reflection redundancy of 16.4 and 128.3, for [(Rb₆F)(Rb₄F)][Ge₁₄O₃₂] and [(Rb₆F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O₃₀F₂], respectively, and $R_{int} = 0.047$ after absorption correction in both cases.

For both $[(Rb_6F)(Rb_4F)][Ge_{14}O_{32}]$ and $[(Rb_6F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O_{30}F_2]$ the pattern of systematic absences in the intensity data was consistent with *F*-centering, but showed no screw axes or glide planes. Of the five resulting space group candidates (*F*23, *Fm* $\overline{3}$, *F*432, *F*-43*m* and *Fm* $\overline{3}m$), *F* $\overline{4}3m$ (space group #216) was determined to be the best choice by the development of the structural model described below.

The structure of $[(Rb_6F)(Rb_4F)][Ge_{14}O_{32}]$ was refined to an R_1 value of 3.08 %. Any attempt to account for the trace of Ag that was detected by the elemental analysis (EDS) did not result in any improvement of the model, suggesting that silver particulates from the reaction vessel had adhered to the sample used for EDS analysis, making it appear that silver was present in the sample. Large positive and negative electron density peaks were observed near the Rb2 site, indicating disorder of this site. To account for the disorder, the positive electron density peak was assigned to Rb3 site, and site occupancies were restrained to add up to a single atom. After several refinement cycles, the occupancies of Rb2 and Rb3 refined to 0.28(3) and 0.16(8) and the R_1 value decreased from 3.08 to 1.85%. The Flack parameter after the final refinement cycle was 0.010(18), consistent with the correct absolute structure assignment and the absence of inversion twinning.

The structure of $[(Rb_6F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O_{30}F_2]$ is extensively disordered, involving split and mixed sites. Accurate determination of composition was limited because of the disorder, and the given compositional uncertainties are likely underestimated. Considerable difficulty was met in achieving a plausible structural model under elemental composition constraints. Semi-quantitative elemental analysis results showed Rb, Ge, Co, O and F as the only significant constituent elements in the crystals, and the structure was modeled using only those elements. Several refinement models were considered, none flawless; the best is presented here. The asymmetric unit in *F*-43*m* consists of one pure Rb site Rb(1A)/Rb(1B), one mixed Rb/Co site Rb(2A)/Co(2A), two mixed Ge/Co sites Ge(1)/Co(1) and Ge(2)/Co(2), one pure Ge site Ge(3),

three unique oxygen atoms O(1)-(O3), one mixed O/F site O(4)/F(4) and two pure F sites F(5) and F(6A)/F(6B). All cation sites have partial or mixed occupancy except for the (pseudo)octahedral germanium site Ge(3) (Wyckoff site 16e, .3m site symmetry). The Ge(3) site occupation factor (sof) refined to 1.003(5) and was fixed at 100% germanium. Sites Ge(1)/Co(1) (24f, 2.mm site symmetry) and Ge(2)/Co(2) (16e, .3m symmetry) were initially refined as germanium. The Ge(2)and Ge(3) sofs refined to significantly less than 100% Ge Ge(1) = 0.888(6) and Ge(2) = 0.953(5), accompanied by a reduction in the R_1 -factor from 0.027 to 0.020. This observation was attributed to mixing of Co(III) onto these sites. A mixed-site Ge/Co model was judged to be more likely than a Ge/vacancy model. The sites were constrained to full occupancy and refined to sof Ge(1)/Co(1)= 0.52(3)/0.48(3) and sof Ge(2)/Co(2) = 0.78(2)/0.22(2). Site Rb(1) is split over two positions, both on site 24g (2.mm symmetry) with refined occupancies Rb(1A) = 0.88(1) and Rb(1B) =0.10(1). Site Rb(2) was initially refined as pure Rb, but it was necessary to introduce cobalt onto this site cluster to account for a distance of 2.24 Å to the 100%-occupied site O(4)/F(4). This is certainly unreasonably short for a Rb-O/F distance and therefore this site must be occupied by a smaller atom for which Co is the most plausible choice. The Rb part of the site cluster Rb(2A)/Co(2A) is located on site 48h (...m symmetry), generating three equivalent atoms with occupancies sof Rb(2A) = 0.261(5). The Co part is located on site 16e (.3m symmetry), generating one site with an occupancy of sof Co(2A) = 0.224(15). The total Rb(2A)/Co(2A) site cluster occupancy therefore sums to 1.0 within experimental error, providing support for the unusual Co admixture model. Three oxygen atoms O(1) and O(2) on site 48h, O(3) on site 16e refined to 100% occupied within error. The sof for site O(4)/F(4) (site 16e) refined to greater than 100% oxygen but less than 100% fluorine and was refined as a 50/50 O/F mixture. Site F(5) (site 4d, -43m site symmetry) refined to significantly greater than 100% oxygen in trials but refined well as 100% F. The final anion site, F(6), is split over two positions F(6A) on site 4c with -43m symmetry and F(6B) on site 16e. The occupancies of these two sites refined to F(6A) = 0.39(4) and F(6B) =0.13(2). The above model generates an electroneutral composition of Rb_{4.51}Ge_{5.10}Co_{2.35}O₁₅F_{1.96} (assuming Co³⁺). All atoms were refined with anisotropic displacement parameters except for minor disorder component atoms Rb(1B), Co(2A) and F(6A)/F(6B). These were refined with a fixed isotropic displacement parameter of 0.025 Å² for refinement stability. The largest residual electron density peak and hole in the final difference map are +1.45 and -0.72 e⁻/Å³, located 1.53 Å from Ge(2)/Co(2) and 0.60 Å from Ge(1)/Co(1), respectively. The Flack after the final

refinement cycle was -0.017(8), consistent with the correct absolute structure assignment and the absence of inversion twinning. Charge balance for the compound may be achieved by assigning +1/+4/+3/-1/-1 to Rb/Ge/Co/O/F. Final unit cell parameters were determined by least-squares refinement of 9092 reflections. Crystallographic and refinement data and selected interatomic distances for both compounds are provided in Tables 1 - 3.

Powder X-ray diffraction (PXRD)

Powder X-ray diffraction data were collected on a Bruker D2 Phaser powder X-ray diffractometer using Cu K α radiation. The step scan covered the angular range 5-65° 2 θ in steps of 0.04°. The experimental and calculated PXRD patterns for [(Rb₆F)(Rb₄F)][Ge₁₄O₃₂] were found to be in good agreement (see Supporting information Fig. S1).

Energy-Dispersive Spectroscopy (EDS)

Elemental analysis was performed on single crystals using a TESCAN Vega-3 SBU SEM with EDS capabilities. The crystals were mounted on carbon tape and analyzed using a 20 kV accelerating voltage and an accumulation time of 1 min. As a qualitative measure, EDS confirmed the presence of Rb, Ge, O, F in [(Rb₆F)(Rb₄F)][Ge₁₄O₃₂] and additionally the presence of Co in [(Rb₆F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O₃₀F₂]. For EDS spectra, see supporting information Figs. S2 and S3.

Fluorescence Spectroscopy

Room temperature emission spectra were collected on a ground sample of $[(Rb_6F)(Rb_4F)][Ge_{14}O_{32}]$ at an excitation wavelength of 258 nm using a Perkin Elmer LS 55 fluorescence spectrometer. For emission measurements the ground sample was placed inside a 6 mm quartz sample holder. Emission scans were performed in the 300 – 800 nm range. Excitation spectra were collected in the 200 – 400 nm range at an emission wavelength of 500 nm.

First-principles Calculations

We performed first-principles calculations for $[(Rb_6F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O_{30}F_2]$ in the form of density functional theory (DFT) with an on-site Coulomb interaction, i.e., DFT+*U*, using the Vienna Ab-initio Package (VASP) code,^{16, 17} using the projector augmented wave (PAW) method^{18, 19} and generalized gradient approximation of Perdew, Burke and Ernzerhof (PBE).²⁰ To model the mixing and partial occupies in the studied compound, we generated super quasi-random structures (SQS) with the experimentally proposed composition $[(Rb_6F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O_{30}F_2]$, using the mcsqs code provided by the Alloy Theoretic Automated Toolkit (ATAT) toolkit.^{21–24} To see if the $[(Rb_6F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O_{30}F_2]$ system is thermodynamically stable, i.e., it breaks the Rb-Co-Ge-F-O convex hull, we compared its formation energy with respect to Open Quantum Materials Database $(OQMD)^{25, 26}$ convex hull. We used the OQMD calculations set-up: 520 eV cut-off energy for the plane wave basis set, 10^{-4} eV energy convergence criterion, $2 \times 2 \times 2$ **k**-point mesh for Rb_{4.51}Co_{2.35}Ge_{5.10}F_{1.96}O₁₅, and $U_{eff} = 3.3$ eV for the Co atoms. We considered the system to be spin-polarized, with high-spin ferromagnetic (FM) and antiferromagnetic (AFM) (0 µ_B magnetic moment) ordering of the Co atoms.²⁷ For calculating the electronic and optical properties we performed more rigorous calculations, using 520 eV cut-off energy for the plane wave basis set, 10^{-6} eV and 10^{-3} eV/Å energy and forces convergence criteria, respectively, and the same **k**-point mesh as the OQMD calculations. The ground state geometries were obtained by relaxing the cell volume, cell shape and atomic positions.

Results and Discussion

Synthesis

Crystals of [(Rb₆F)(Rb₄F)][Ge₁₄O₃₂] and [(Rb₆F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O₃₀F₂] were grown by an enhanced flux growth method using metal halide reagents and reaction vessels that minimized surface area to volume ratios that has been developed to encourage the synthesis of mixed anion compounds as well as other complex germanates, which we previously reported.^{2, 3,} ^{28–30} The use of mixed alkali halide fluxes at relatively high temperatures has proven successful for the discovery and crystal growth of numerous new oxyfluoride and salt-inclusion materials.^{31–} ³³ The specific conditions used in this work take advantage of the relatively low melting point of the eutectic RbCl/RbF system (540 °C) compared to the synthesis and presumed crystal nucleation temperatures of °C 850 900°C for $[(Rb_6F)(Rb_4F)][Ge_{14}O_{32}]$ and and [(Rb₆F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O₃₀F₂], respectively. It is important to notice that the synthesis of [(Rb₆F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O₃₀F₂] involves CoF₂ as a reagent, instead of a cobalt oxide reagent, which affects the rate of dissolution in the flux.

Crystal Structure

The title compounds are closely related to the previously reported salt inclusion phase $[(Cs_6F)(Cs_3AgF)][Ge_{14}O_{32}]^{11}$ as all share a large germanium framework and salt-inclusion components within the 3D channels. The nature of the salt inclusion in $[(Rb_6F)(Rb_4F)][Ge_{14}O_{32}]$

and the presence of fluorine in the germanium framework in $[(Rb_6F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O_{30}F_2]$, however, differentiate the new compositions from the previously reported $[(Cs_6F)(Cs_3AgF)][Ge_{14}O_{32}]$.

The germanium framework structure of $[(Rb_6F)(Rb_4F)][Ge_14O_{32}]$ is derived from GeO₆ octahedra that share edges with one another to form Ge₄O₁₆ tetramers. The Ge-O bond lengths for GeO₆ octahedra range from 1.826(3) Å – 1.958(3) Å, in good agreement for what is observed in $[(Cs_6F)(Cs_3AgF)][Ge_{14}O_{32}]$. These tetramers are further connected through edge sharing to intervening GeO₄ tetrahedra to form the overall 3D framework shown in Fig. 1. The Ge-O bond lengths in the GeO₄ tetrahedra range from 1.728(4) Å – 1.749(4) Å consistent with the bond lengths in $[(Cs_6F)(Cs_3AgF)][Ge_{14}O_{32}]$. The framework of $[(Rb_6F)(Rb_4F)][Ge_{14}O_{32}]$ contains large, interconnected channels running down the *a*-, *b*- and *c*-axis in which the salt inclusion is located. The salt inclusion consists of tetrahedral Rb₄F (Rb-F bond lengths of 2.446(16) Å) and octahedral Rb₆F units (Rb-F bond lengths of 2.974(3) Å) that are located in the channels and arranged to occupy alternating channels in an NaCl fashion, shown in Fig. 2. This impacts the size of the unit cell that consists of 8 cubes (2 x 2 x 2). In contrast, the presence of Ag in $[(Cs_6F)(Cs_3AgF)][Ge_{14}O_{32}]$ resulted in the salt inclusions consisting of octahedral Cs₆F and trigonal pyramidal Cs₃AgF units.



Fig. 1 Germanium oxide framework in [(Rb₆F)(Rb₄F)][Ge₁₄O₃₂] made up of Ge₄O₁₆ tetramers and GeO₄ polyhedra. Ge and O are shown in deep blue and red, respectively.



Fig. 2 Structure of [(Rb₆F)(Rb₄F)][Ge₁₄O₃₂] highlighting the arrangement of tetrahedral Rb₄F and octahedral Rb₆F salt inclusions in the channels. Disordered Rb atoms of the Rb₄F tetrahedra are not shown. Rb is shown in orange, F is shown in green, Ge is shown in deep blue, and oxygen is shown in red.

The structure of $[(Rb_6F)(Rb_3.1Co_0.9F_{0.96})][Co_3.8Ge_{10.2}O_{30}F_2]$ is distinct from both $[(Rb_6F)(Rb_4F)][Ge_{14}O_{32}]$ and $[(Cs_6F)(Cs_3AgF)][Ge_{14}O_{32}]$ in that it contains two fluorine atoms in the framework, causing it to be best described as an oxyfluoride framework. The structure is composed of mixed Ge(2)/Co(2)O₆ and Ge(3)(O/F)₆ octahedra. The M-O bond lengths for the Ge(2)/Co(2)O₆ octahedra range from 1.839(3) Å – 1.966(2) Å, which is in good agreement with Co-O bond distances taking into account partial site occupation by Ge.³⁴ The Ge-O bond lengths for the Ge(3)(O/F)₆ octahedra range from 1.827(2) Å – 1.9552 (2) Å consistent with the bond lengths for six-coordinated tetravalent Ge.³⁵ The Ge(2)/Co(2)O₆ octahedral tetramers to form tetramers that are connected in an alternating fashion to Ge(3)(O/F)₆ octahedral tetramers via corner-sharing through intervening Ge(1)/Co(1)O4 tetrahedra in all three dimensions of the crystal structure, creating large channels running down the *a*-, *b*- and *c*-axis. The Co/Ge-O bond lengths in the Ge(1)/Co(1)O4 tetrahedra range from 1.710(3) Å – 1.724(3) Å. The channels created by the Co/Ge polyhedral network are occupied by fluoride ions, F(5) and disordered F(6), pure rubidium

Rb(1) ions and, most strikingly, an unusual mixed Rb(2A)/Co(2A) clusters, shown in Fig. 3. The arrangement of the Co/Ge- and Ge-based tetramers in the framework columns is reminiscent of the *ABAB* stacking sequence in rock-salt. The complete pictorial representation of the crystal structure assembly for $[(Rb_6F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O_{30}F_2]$ is provided in Figs. 4 and 5.



Fig. 3 Representations of the (Rb₆F) octahedral and (Rb_{3.1}Co_{0.9}F_{0.96}) tetrahedral salt inclusions in $[(Rb_6F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O_{30}F_2]$ The F4 anion is part of the framework and directly bonded to Ge. F5 and F6 are part of the salt inclusion. Fluorine atoms are green, rubidium atoms orange and cobalt atoms blue.



Fig. 4 Assembly of (Co/Ge)O₆ and Ge(O/F)₆ based tetramers via corner-sharing with (Co/Ge)O₄ tetrahedra forming columns that constituent the cobalto-germanate framework in [(Rb₆F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O₃₀F₂]. The mixed Co/Ge and pure Ge octahedra are shown in green and deep blue, respectively. O is shown in red.



Fig. 5 Projection of the crystal structure of $[(Rb_6F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O_{30}F_2]$ down the *c*-axis. Pure fluorine and pure rubidium sites are depicted as green and orange spheres, respectively. The mixed green and white spheres represent the mixed Rb/Co clusters in the channels. The mixed Co/Ge and pure Ge octahedra are shown in green and deep blue, respectively. O is shown in red.

Table 1 Crystallographic and refinement data for $[(Rb_6F)(Rb_4F)][Ge_{14}O_{32}]$ and $[(Rb_6F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O_{30}F_2]$.

Empirical formula	FGe7O16Rb5	$Co_{2.35}F_{1.96}Ge_{5.10}O_{15}Rb_{4.51}$
Formula weight (g/mol)	1210.53	1171.63
Temperature (K)	301(2)	301(2)
Wavelength (Å)	0.71073	
Crystal system	Cubic	
Space group	<i>F</i> -43 <i>m</i>	
Unit cell dimensions (Å)	<i>a</i> = 15.34580(10)	<i>a</i> = 15.2686(14)
Volume (Å ³) and Z	3613.84(7) and 8	3559.6(10) and 8
Density (calculated) (mg/m ³)	4.450	4.373
Absorption coefficient (mm ⁻¹)	24.956	22.982
F(000)	4368	4250
Crystal size (mm x mm x mm)	0.04 x 0.04 x 0.04	0.20 x 0.18 x 0.16
Theta range for data collection (°)	2.30 to 33.10	2.310 to 37.752
Index ranges	-23<=h<=19, -	-26<=h<=26, -26<=k<=26, -
	23<= <i>k</i> <=21, -	26<= <i>l</i> <=26
	23<= <i>l</i> <=17	
Reflections collected	12156	69679
Independent reflections	740 [$R(int) = 0.0468$]	1006 [R(int) = 0.0474]
Data / restraints / parameters	740 / 0 / 46	1006 / 0 / 52
Goodness-of-fit on F^2	1.071	1.127
Final <i>R</i> indices [I>2sigma(I)]	$R_1 = 0.0185,$	$R_1 = 0.0200,$
	$wR_2 = 0.0445$	$wR_2 = 0.0514$
R indices (all data)	$R_1 = 0.0239,$	$R_1 = 0.0238,$
	$wR_2 = 0.0474$	$wR_2=0.0530$
Largest diff. peak and hole	$1.001 \text{ and } -0.797 \text{ e.Å}^{-3}$	$1.454 \text{ and } -0.718 \text{ e.Å}^{-3}$

Bond	Distance (Å)
F(5) - Rb(1) x6	2.974(3)
F(2) - Rb(2) x4	2.446(16)
Ge(1) - O(1) x3	1.958(3)
Ge(1) - O(6) x3	1.826(3)
Ge(2) - O(3) x3	1.842(3)
Ge(2) - O(4) x3	1.970(3)
Ge(3) - O(3) x2	1.728(4)
Ge(3) - O(6) x2	1.749(4)

Table 2 Select Interatomic Distances (Å) of [(Rb₆F)(Rb₄F)][Ge₁₄O₃₂].

Table 3 Select Interatomic Distances (Å) of $[(Rb_6F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O_{30}F_2]$. M(1) = Ge(1)/Co(1) = 0.52(3)/0.48(3); M(2) = Ge(2)/Co(2) = 0.78(2)/0.22(2).

Bond	Distance (Å)
M(1) – O(2)	1.710(3)
M(1) – O(2)	1.710(3)
M(1) - O(1)	1.724(3)
M(1) - O(1)	1.724(3)
M(2) – O(2) x3	1.839(3)
M(2) - O(3) x3	1.966(2)
Ge(3) - O(1) x3	1.827(2)
Ge(3) - F(4) x2	1.955(2)
Ge(3) - O(4)	1.955(2)

Photoluminescence

Fig. 6 shows the emission spectrum for $[(Rb_6F)(Rb_4F)][Ge_14O_{32}]$, which exhibits considerable luminescence at room temperature. This is in contrast to the previously reported germanium framework SIM, $[(Cs_6F)(Cs_3AgF)][Ge_14O_{32}]$, which does not luminesce at room temperature. The emission spectrum of $[(Rb_6F)(Rb_4F)][Ge_14O_{32}]$ consists of one broad peak with a maximum at approximately 509 nm, consistent with the yellow-green luminescence color observed in $[(Rb_6F)(Rb_4F)][Ge_14O_{32}]$ crystals. The absence of a color center in $[(Rb_6F)(Rb_4F)][Ge_14O_{32}]$ and the absence of luminescence in the other members of germanium framework SIMs makes it difficult to make a structural argument for the source of luminescence in this material that appears to be intrinsic to the framework in the case of $[(Rb_6F)(Rb_4F)][Ge_14O_{32}]$. One can postulate that the reactive nature of the fluoride salt in the flux, in this case toward the silver reaction vessel, creates defects in the $[(Rb_6F)(Rb_4F)][Ge_14O_{32}]$ structure that are too minor or too disordered to be detected by single crystal structure determinations, yet that lead to luminescent behavior.



Fig. 6 Emission spectrum at an excitation λ of 258 nm and optical image of $[(Rb_6F)(Rb_4F)][Ge_{14}O_{32}]$ crystals under UV-light.

Attempts were made to tune the luminescence of [(Rb₆F)(Rb₄F)][Ge₁₄O₃₂] by doping it with a small amount of Mn on any of the available Ge sites within the framework. Reactions were

performed as described in the experimental section with the addition of a small percentage of MnO₂. Orange crystals matching the morphology and unit cell of [(Rb₆F)(Rb₄F)][Ge₁₄O₃₂] were prepared, however, the presence of Mn was not detectable in the single crystal X-ray diffraction data and no room temperature luminescence was observed.

Electronic Structure

The DFT calculations for the [(Rb₆F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O₃₀F₂] structure indicate it relaxes to two ferromagnetic states with high spin (HS) and low spin (LS), having magnetic moments of 3.21 and 0.79 μ_B/Co atom, despite the starting antiferromagnetic configuration of the LS system. The calculated lattice parameters (Table 4) are very close to the experimental values, with an error < 1%. The LS system is more stable, having a 4.8 meV/atom more negative energy compared to the HS system. Both systems do not break the OQMD convex hull with the LS lying 45.8 meV above the convex hull, indicating that these SQS are metastable. Exploring different SQS might yield a slightly more stable state, however, considering that a purely random structure has much higher energy and very different DOS (see supplementary material Figure S4), we believe that a more stable structure would have minuscule effect on DOS and optical properties. The band gaps of the HS system are 1.18 and 1.45 eV in the spin-up and spin-down channel, respectively, while for the LS system are 1.08 and 1.38 eV in the spin up and spin down channel, respectively. From the projected DOS (Figs. 7a and 7b) the HS and LS system the states at the top of the valence band come predominantly from O, while the state at the bottom of the conduction band results from Co, indicating that this compound should behave more like a charge-transfer semiconductor. The different semiconductor character from CoO is caused by the presence of Co-F bonds from the Rb/Co clusters in the framework channels. The bonding with F pushes the Co orbitals to lower energies, decreasing the Co and increasing the O contribution to the states at the top of the valence band. The adsorption indexes of the HS and LS systems are almost identical for energies below 3 eV (Fig. 7d), which arises from the similarity of the HS and LS DOS proximal to their band gaps (Fig. 7c).



Table 4 DFT calculated crystallographic data of [(Rb₆F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O₃₀F₂]. The system relaxed to high spin (HS) and low spin (LS) states.

Fig. 7 Projected density of states (PDOS) of [(Rb₆F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O₃₀F₂] in: a) high spin (HS), and b) low spin (LS) state. The total DOS, Co, Cs, Si and O PDOS are shown in black, blue, green, orange and red, respectively. c) total DOS and d) absorption indexes of [(Rb₆F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O₃₀F₂] in HS and LS state, shown in blue and red, respectively.

Conclusion

 $[(Rb_{6}F)(Rb_{4}F)][Ge_{14}O_{32}]$ Germanate salt inclusion materials (1) and $[(Rb_6F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O_{30}F_2]$ (2) were synthesized as single crystals using a high temperature RbCl/RbF flux method. Both compounds have the germanate framework observed previously in [(Cs₆F)(Cs₃AgF)][Ge₁₄O₃₂]. Modification of the salt inclusion with Rb has led to intrinsic room temperature luminesce of the framework in [(Rb₆F)(Rb₄F)][Ge₁₄O₃₂]. Fluorescence measurements show a broad peak with a maximum of approximately 509 nm, consistent with the yellow-green color observed in the crystals under UV light. We have also now demonstrated the the ability to incorporate other elements into framework, in the case of [(Rb₆F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O₃₀F₂]. Incorporation of cobalt resulted in unprecedented Rb/Co mixing within the salt inclusions. First principles calculations for [(Rb₆F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O₃₀F₂] indicated that its a metastable phase, where the low spin (LS) state is slightly more stable than the high spin (HS) state. The DOS indicate that [(Rb₆F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O₃₀F₂] has higher charge-transfer semiconductor character, in contrast to CoO, due to the presence of fluorine in the structure.

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Supplementary information

The crystallographic CIF files for [(Rb₆F)(Rb₄F)][Ge₁₄O₃₂] and [(Rb₆F)(Rb_{3.1}Co_{0.9}F_{0.96})][Co_{3.8}Ge_{10.2}O₃₀F₂] were deposited into the CCDC data base with CCDC numbers of 2014420 and 1905088 (<u>https://www.ccdc.cam.ac.uk</u>).

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