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From simple to complex crystal chemistry in the RE-Au-Tt systems (RE = La, Ce, Pr, Nd; Tt = Ge, Pb)

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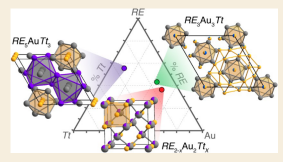
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ABSTRACT: Polar intermetallics are an intriguing class of compounds with complex relationships between composition and structure that are not fully understood. This work reports a systematic study of the underexplored ternary composition space RE–Au–Tt (RE = La, Ce, Pr, Nd; Tt = Ge, Pb) to expand our knowledge of the intriguing chemistry and diversity achievable with these metallic constituents. These composition spaces are particularly interesting because of the potential to find Au-bearing, highly polar intermetallic compounds. The elements were first reacted through arc welding under an inert atmosphere, followed by annealing at 850 °C. X-ray diffraction of the products identified seven unreported compounds ranging from the simple NaTl-type compounds La_{1.5}Au₂Pb_{0.5} and Nd_{2-x}Au₂Pb_x to the



more structurally complex La₅AuPb₃ in the Hf₅CuSn₃-type structure and Pu₃Pd₄-type RE₃Au₃Ge (RE = La, Ce, Pr, Nd). First-principles electronic structure calculations investigate the combination of Fermi surface—Brillouin zone interactions, electrostatic interactions, and delocalized metallic bonding that contributes to the formation of these phases. These calculations show that a mixture of electrostatic and metallic bonding plays a dominant role in these phases. The RE—Au—Tt composition space remains full of potential for discovering materials with relevant magnetic and quantum properties, provided the crystal chemistry can be comprehended.

KEYWORDS: Intermetallics, Au chemistry, polar interactions, rare earth chemistry, solid-state bonding, electronic structure

■ INTRODUCTION

Polar intermetallic compounds, a subset of metal-rich solids, have received much interest in academic laboratories and industrial research settings because of their complex crystal chemistry and potential for fascinating physical properties. Strictly defined, polar intermetallics are compounds that have a ratio of valence electrons per atom between $1.2 \le (e/a) \le 4$, existing between the limits of Hume-Rothery and Zintl phases, which contain $1 \le (e/a) \le 2$ and $(e/a) \ge 4$, respectively.¹⁻⁴ Varying the e/a ratio and chemical composition provides a diversity of crystal structures and distinct chemical bonding motifs ranging from networks to clusters and quasicrystals.⁵⁻⁷ The complexity of polar intermetallics can be justified because they contain structural and bonding features from both Hume-Rothery and Zintl type compounds. Unlike Hume-Rothery and Zintl phases, polar intermetallics suffer from complicated electron counting rules and the inability to assign definitive oxidation states, making predictions of crystal chemistry challenging.⁸ This is further convoluted by the wide range of compositions and stoichiometries that can be combined to form polar intermetallics, leaving large swathes of phase space unexplored. Therefore, to further our understanding, we must find new compounds that can help us unravel the intricacies of polar intermetallics, understand

their structural tendencies, and further interpret their chemical bonding and overall crystal chemistry.

Ternary rare-earth (RE) noble metal (M) tetrels (Tt) are an exciting composition space to conduct such a study because of the relatively large electron affinity differences between the constituent elements. These phases can be quickly synthesized by arc-melting followed by annealing at accessible temperatures. The products crystallize easily, and numerous ternary compounds have been identified and characterized. Indeed, these compounds are of great interest owing to their fascinating structural chemistries and physical properties, including superconductivity (e.g., RE₂Pt₃Ge₅, RE₂Ir₃Ge₅), 10 magnetocaloric effects (e.g., GdRu₂Ge₂, RE₂M₂Sn), 11,12 complex magnetic ordering (e.g., RECrGe₃, Nd₁₁₇Co₅₄Sn₁₁₁, RE₂Ni₂Pb), 13,14 and other quantum phenomena (e.g., RE_{n+1}M_nTt_{3n+1}). The magnetic properties stem from the interaction between the localized 4f electrons of a rare-earth atom and the d electrons of transition metal. However, there

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Table 1. Space Groups and Refined Cell Parameters for All Compounds from Powder X-ray Diffraction

	Space Group	a (Å)	b (Å)	c (Å)	α (deg)	β (deg)	γ (deg)
La ₃ Au ₃ Ge	R3 (148)	14.0300(4)	14.0300(4)	6.2015(6)	90	90	120
Ce ₃ Au ₃ Ge	$R\overline{3}$ (148)	13.865(2)	13.865(2)	6.150(1)	90	90	120
Pr ₃ Au ₃ Ge	$R\overline{3}$ (148)	13.8118(5)	13.8118(5)	6.1249(5)	90	90	120
Nd ₃ Au ₃ Ge	$R\overline{3}$ (148)	13.7075(4)	13.7075(4)	6.0935(5)	90	90	120
La ₅ AuPb ₃	$P6_3/mcm$ (193)	9.692(1)	9.692(1)	7.0087(9)	90	90	120
$La_{1.5}Au_2Pb_{0.5}$	$Fd\overline{3}m$ (227)	7.2252(1)	7.2252(1)	7.2252(1)	90	90	90
$Nd_{2-x}Au_2Pb_x$	$Fd\overline{3}m$ (227)	7.0906(6)	7.0906(6)	7.0906(6)	90	90	90

are no apparent mechanisms for controlling magnetism in these polar intermetallic compounds, even though some systems like RE–Ni–Ge have been exhaustively studied. 17–19

It is possible to add further complexity to the crystal chemistry of polar intermetallics by studying gold-containing compounds. Au has the highest electron affinity of any element outside of the halides, indicating that the most polarized intermetallics may exist in M = Au compositional spaces. However, many RE-Au-Tt ternary phase spaces remain underexplored, with most systems reporting only one or two compounds. One common thread is the REAuTt phase, which crystallizes in the rhombohedral LiGaGe structure type with space group $P6_3mc$ (No. 186). The [AuGe]ⁿ⁻ anionic sublattice in this compound features a strong polar-covalent bond between Ge-Au and Au-Au, generating a threedimensional polyanionic backbone. REAu2Ge2 phases in the tetragonal ThCr₂Si₂ structure type have also been identified in some RE systems, such as Ce and Pr, and are reported to order antiferromagnetically with Néel temperatures of 16 and 11.9 K, respectively. 19,23,24 More complex and unique topologies also pervade these systems, such as the orthorhombic Ce₃Pd₆Sb₅ type formed in the RE-Au-Sn systems (RE = Ce, Nd, Sm).²

This work identifies seven new compounds in the RE–Au–Tt (RE = La, Ce, Pr, Nd; Tt = Ge, Pb) ternary phase space through a systematic study of these composition spaces at 850 °C. These compounds adopt three structure types spanning simple to complex crystal chemistry across the Au-rich to Aupoor composition range. They include $RE_{2-x}Au_2Pb_x$ (RE = La, Nd) in the NaTl type, La_5AuPb_3 in the Hf_5CuSn_3 type, and RE_3Au_3Ge with RE = La, Ce, Pr, Nd in the Pu_3Pd_4 structure type. Density functional theory (DFT) calculations were performed to interpret the origin of structural favorability for these phases and examine the nature of the covalent, ionic, and metallic contributions to polar intermetallic bonding. The results of this work will further our fundamental understanding of these systems and gold-containing polar intermetallic phases in general.

■ EXPERIMENTAL SECTION

Synthesis

The starting metals—gold (splatter, 99.999%, Materion Advanced Chemical), tetrels Ge (powder, 99.99%, Alfa Aesar), and Pb (shot, W. H. Curtin & Co., ca. 1960), and the rare-earth elements La, Ce, Pr, and Nd filings (99.8%, HEFA Rare Earth), were weighed out in the desired stoichiometric ratios and ground together with an agate mortar and pestle to form a uniform sample with a total weight of 200 mg. The samples were cold-pressed into 6 mm pellets using a maximum pressure of 2500 psi and reacted under argon gas in a Centorr 5SA arc furnace with a current of 30–45 A. All samples were flipped and melted multiple times to ensure homogeneity. The weight loss for each sample after arc-melting was less than 2%. Each resulting ingot was subsequently sealed in an evacuated fused silica tube (less than 4×10^{-2} Torr) and annealed in a Thermo Scientific Lindberg

Blue M muffle furnace by heating to 850 °C over 12 h, holding for 200 h, and cooling back to room temperature over 24 h. The final products had a silvery metallic luster and were all brittle. The Lacontaining products showed some sensitivity to air fading to a dull black color over several days to weeks when left at ambient conditions. As a result, all products were stored in an argon atmosphere glovebox ($O_2 < 0.1$ ppm; $H_2O < 0.1$ ppm). The preparation of LaAuGe3, LaAuGe2, La4Au5Ge11, LaAu2Ge, LaAu3Ge, La2AuGe, and La2Au2Ge was also attempted following the same synthetic procedure; however, the products from these reactions were all multiphase based on powder X-ray diffraction and fully indexed to known compounds.

Powder X-ray Diffraction

X-ray powder diffractograms were collected at room temperature using a PANalytical X'Pert powder diffractometer equipped with Cu $K\alpha$ radiation (λ = 1.54183 Å). The samples were prepared by grinding the annealed ingots in the glovebox using an agate mortar and pestle. The fine powder was then spread on a zero-diffraction silicon plate and measured from 5° to 90° 2 θ . Le Bail refinements were carried out on the collected data of all compositions to confirm phase purity and determine the lattice parameters using the EXPGUI interface for the General Structure Analysis System (GSAS). ^{26,27} Refined unit cell parameters for all synthesized compounds are given in Table 1, and Le Bail refinement of powder X-ray diffraction data is shown in Figure S1. A shifted-Chebyshev function was used to model the background. All crystal structures were visualized using VESTA²⁸ and Diamond 3^{29}

Single-Crystal X-ray Diffraction

Single crystals were harvested from crushed samples of the Lacontaining phases and mounted on the end of pulled glass fibers. Crystals were silvery gray and lustrous and generally formed as small cubes or plates approximately $10-50~\mu m$ in size. Diffraction data sets were collected as omega scans $(0.5^{\circ}$ scan width at $60~\mu m$ detector distance) on a Bruker Apex II platform diffractometer equipped with a 4KCCD Apex II detector and Mo K α radiation source at 296 K. Peak hunting, indexing, data integration, and finalization were performed with the provided Bruker APEX II software. The structures of each crystal were solved using the shelXle program package for SHELX refined on $F_o^{2.30}$ Anisotropic atomic displacement parameters for each structural solution are given in Table S1. Solved structures have been submitted to the CCDC and can be obtained under deposition numbers 2129946-2129948. Crystal structures were visualized with the Crystal Impact Diamond 3 and VESTA software packages. 28,29

Electronic Structure Calculations

Density functional theory (DFT) calculations using the Vienna Abinitio Simulation Package (VASP), version 5.4.4, were performed to optimize the geometries with a convergence criterion of 1×10^{-3} eV/Å and calculate the wave functions and charge densities of each reported compound with a convergence criterion of 1×10^{-8} eV. $^{3\,I_332}$ DFT optimized total energies, unit cell parameters, and atomic positions are listed in Tables S2–S5. Calculations were done using the projector augmented wave (PAW) potentials with exchange and correlation described by Perdew–Burke–Ernzerhof (PBE) with a Γ-centered Monkhorst–Pack k-point grid containing at least 1000 k-points atom $^{-1}$. 33,34 Spin–orbit coupled (SOC) fully noncollinear magnetic calculations of the NaTl-type La_{1.5}Au₂Pb_{0.5} structure were

additionally performed.³⁵ A comparison of non-SOC and SOC calculations is shown in Figure S2, indicating that SOC effects do not substantially affect the electronic structures of these systems. Thus, they are not included in any further calculations. The density of states (DOS) and crystal orbital Hamilton population (COHP) calculations were carried out using the Local-Orbital Basis Suite toward Electronic Structure (LOBSTER).^{36,37} Bader charges on each atomic position were determined using the Bader program for VASP.^{38–40}

RESULTS AND DISCUSSION

Synthesis and Structural Characterization

Six ternary intermetallic systems containing a rare earth (RE) metal, gold, and a main group tetrel metal (Tt) were synthetically explored: La–Au–Pb, Nd–Au–Pb, La–Au–Ge, Ce–Au–Ge, Pr–Au–Ge, and Nd–Au–Ge. In each case, elements were mixed in stoichiometric ratios, arc welded under flowing Ar, and sealed in fused silica ampules under vacuum and annealed at 850 °C for 200 h. Structural characterization using single-crystal and powder X-ray diffraction of the resulting products indicated the formation of seven previously unknown compounds in three distinct classes: a NaTl-type structure type for RE_{2-x}Au₂Pb_x (La_{1.5}Au₂Pb_{0.5}, Nd_{2-x}Au₂Pb_x), followed by La₅AuPb₃ in the Hf₅CuSn₃ structure type, and the most complex RE₃Au₃Ge (La₃Au₃Ge, Ce₃Au₃Ge, Pr₃Au₃Ge, Nd₃Au₃Ge) in the Pu₃Pd₄ structure type.

The first class of compounds presented in this work are two newly discovered $RE_{2-x}Au_2Pb_x$ (RE=La, Nd) compounds adopting the NaTl structure type ($Fd\overline{3}m$): $La_{1.5}Au_2Pb_{0.5}$ and $Nd_{2-x}Au_2Pb_x$. These compounds, illustrated in Figure 1a, form an ordered superstructure of the simple body-centered cubic arrangement, in which Au atoms alternate with statistically mixed RE and Pb sites. The refined unit cell parameters of these compounds were determined through Le Bail refinement of powder X-ray diffraction, and the fit diffractogram for the $La_{1.5}Au_2Pb_{0.5}$ phase is shown in Figure 1b (refined cell

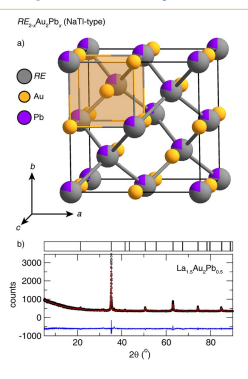


Figure 1. (a) Structure of $RE_{2-x}Au_2Pb_x$ in the cubic NaTl type and (b) the Le Bail refinement of $La_{1.5}Au_2Pb_{0.5}$.

parameters for all structures given in Table 1). For the La analog, the RE/Pb elemental mixing was determined through single-crystal X-ray diffraction to be 79(2)% La and 21(2)% Pb, leading to an overall composition of $La_{1.58(3)}Au_2Pb_{0.42(3)}$, which is in agreement with the nominally loaded $La_{1.5}Au_2Pb_{0.5}$ composition. Similar statistical mixing is expected for the Nd structure. Crystallographic and structural parameters for this structure solution are provided in Tables 2–4.

Despite lacking a band gap, NaTl-type phases are generally considered Zintl phases due to their octet-satisfied anionic networks. Many examples of binary alkali-main group metal NaTl-type compounds are known, but only a few Aucontaining ternary compositions such as Li₂AuSb, Li₂Au_{0.75}Si_{1.25}, and Li₂Au_{0.25}In_{1.75} are reported.⁴¹ A Paulinginspired calculation of the valence electrons per anionic atom in these phases confirms the classification of these phases as Zintl compounds with ratios approaching or above 4.00 e/a. For example, the most electron-poor of these species, Li₂Au_{0.25}In_{1.75}, lies just below the lower bound of the Zintl edge (3.75 e/a). RE-bearing NaTl-type phases are similarly rare, but compounds such as Mg_{0.75}Ce_{0.5}Zn_{0.75} (4.67 e/a) also form well into the Zintl regime. 42 By contrast, the Au-rich RE-Au-Tt phases presented here do not conform to the Zintl mold. For example, La_{1.5}Au₂Pb_{0.5} and La₃Au₃Ge form at valence densities of 2.20 and 2.50 e/a, respectively, and thus fall well short of Pauling's demarcation. Similarly, although the RE-rich La₅AuPb₃ forms at a higher e/a count of 4.50, its structure lacks the polyanionic network expected to satisfy the Zintl-Klemm concept. This is indicative that these compounds represent a step into a previously unexplored compositional region of non-Zintl NaTl-type phases.

The next compound identified is La_5AuPb_3 , which forms in the rhombohedral Hf_5CuSn_3 structure type $(P6_3/mcm)$. The structure of La_5AuPb_3 is shown in Figure 2a, and refined unit cell parameters from Le Bail powder (Figure 2b) and single-crystal diffraction are given in Tables 1 and 2, respectively. This structure type is, in itself, a derivative of the classic Mn_5Si_3 structure, ⁴⁴ which consists of two crystallographically distinct columns of face-sharing octahedra that run along the c axis. In the Hf_5CuSn_3 -type, noble metal atoms stuff the usually empty RE octahedra centered at the unit cell vertices, creating a one-dimensional chain of 91% occupied Au sites within a distance comparable to metallic bonding interactions. These $Au@La_6$ octahedra are tiled in the c axis and edges the in c0-plane.

The final and most complex series of compounds discovered crystallize in the Pu₃Pd₄ structure type (R3) with the generic formula RE_3Au_3Ge (RE = La, Ce, Pr, Nd). The Pu_3Pd_4 structure type is present in the Inorganic Crystal Structure Database (ICSD) for RE-Au binary systems, 45,46 including Nd₃Au₄ and Th₃Au₄. However, it has only once been reported in a ternary system in the form of Nd₃Pd_{3.79}Ru_{0.21}. Figure 3a shows the large rhombohedral unit cell of this structure type solved from single-crystal diffraction data for La₃Au₃Ge (crystallographic data provided in Tables 2-4), and Figure 3b shows the Le Bail refinement for this structure. The refined cell parameters for all members of the RE3Au3Ge family prepared here are additionally provided in Table 1. The solved La₃Au₃Ge structure is characterized by an AlB₂-like arrangement of RE octahedra centered by a disordered mixture of Au (43%) and Ge (57%). Forming a scaffolding layer between the octahedra is a Au-rich (89%) network of atoms forming hexagonal channels that run along the c-axis of the structure.

Table 2. Crystallographic Data for the Nominally Loaded La_{1.5}Au₂Pb_{0.5}, La₅AuPb₃, and La₃Au₃Ge from Single-Crystal X-ray Diffraction

Nominal formula	La _{1.5} Au ₂ Pb _{0.5}	La ₅ AuPb ₃	La ₃ Au ₃ Ge
Refined formula	La _{1.58(3)} Au ₂ Pb _{0.42(3)}	$La_5Au_{0.91(1)}Pb_3$	$La_3Au_{3.17(1)}Ge_{0.83(1)}$
Formula mass (g mol-1)	700.37	1494.83	1101.55
Space group; Z	$Fd\overline{3}m; 4$	<i>P</i> 6 ₃ /mcm; 2	$R\overline{3}$; 6
a, b (Å)	7.2315(4)	9.7121(3)	14.04(1)
c (Å)	7.2315(4)	7.0033(3)	6.220(5)
V (Å3)	378.17(6)	572.08(4)	1062(2)
$ ho_{ m calc}$ (g cm ⁻³)	12.3006	8.67731	6.88628
Absorption correction		Multi-scan	
Radiation		Μο Κα	
μ (mm ⁻¹)	113.162	73.701	83.445
θ range (deg.)	4.88 – 34.225	2.421 - 30.501	2.901 – 27.563
hkl ranges	$-10 \le h \le 9$	$-13 \le h \le 13$	$-18 \le h \le 17$,
	$-11 \le k \le 10$	$-11 \le k \le 13$	$-13 \le k \le 18$,
	$-11 \le l \le 9$	$-10 \le l \le 9$	$-8 \le l \le 7$
No. reflections; $R_{\rm int}$	743; 0.0124	6479; 0.0426	1691; 0.055
No. unique reflections	51	343	539
No. parameters	4	15	26
$R(F) \text{ for } F_0^2 > 2\sigma (F_0^2)^a$	0.0270	0.0331	0.0439
$R_{ m w}(F_{ m o}{}^2)$ b	0.0538	0.0791	0.1132
Goodness of fit	1.230	1.081	1.056
$(\Delta ho)_{ m max}$, $(\Delta ho)_{ m min}$ (e Å-3)	0.815, -1.969	2.455, –1.983	4.097, -2.838

 $R(F) = \sum ||F_o| - |F_c|| / \sum |F_o||$

 $p = [\max(F_o^2, 0) + 2F_c^2]/3$

Table 3. Refined Atomic Coordinates and Equivalent Isotropic Displacement Parameters for the Nominally Loaded Compounds (a) $La_{1.5}Au_2Pb_{0.5}$, (b) $La_5Au_2Pb_3$, and (c) La_3Au_3Ge

compound	atom	Wyck.	occ.	\boldsymbol{x}	y	z	$U_{\rm eq} (\mathring{\rm A}^2)^a$
(a) La _{1.5} Au ₂ Pb _{0.5}	Au1	8 <i>a</i>	1	1/8	5/8	5/8	0.0212(4)
	La1	8d	0.79(2)	3/8	7/8	7/8	0.0277(6)
	Pb1	8d	0.21(2)	3/8	7/8	7/8	0.0277(6)
(b) La ₅ AuPb ₃	Pb1	6g	1	0.6132(1)	1	3/4	0.0202(3)
	Au1	2b	0.907(9)	1	1	1/2	0.0310(7)
	La1	4d	1	1/3	2/3	1/2	0.0192(3)
	La2	6g	1	1	0.2619(2)	3/4	0.0213(4)
(c) La ₃ Au ₃ Ge	Au1	18f	0.89(1)	0.72301(7)	0.94251(7)	0.2178(1)	0.0174(3)
	Ge1	18f	0.11(1)	0.72301(7)	0.94251(7)	0.2178(1)	0.0174(3)
	La1	18f	1	0.4988(1)	0.8761(1)	0.4019(2)	0.0178(5)
	Au2	3 <i>a</i>	0.56(2)	1/3	2/3	2/3	0.037(1)
	Ge2	3 <i>a</i>	0.44(2)	1/3	2/3	2/3	0.037(1)
	Au3	3b	0.43(2)	1/3	2/3	1/6	0.041(2)
	Ge3	3 <i>b</i>	0.57(2)	1/3	2/3	1/6	0.041(2)

 $^{^{}a}U_{\mathrm{eq}}$ is defined as one-third of the trace of the orthogonalized U_{ij} tensor.

 $^{{}^{\}mathrm{b}}\,R_w(F_o^2) = \big[\sum [w(F_o^2 - F_c^2)^2] / \sum w F_o^4 \big]^{1/2} \,; \,\, w^{-1} = \left[\sigma^2 (F_o^{\ 2}) + (Ap)^2 + Bp \right] \qquad \text{where}$

Table 4. Selected Refined Bond Lengths [Å] for La_{1.5}Au₂Pb_{0.5}, La₅AuPb₃, and La₃Au₃Ge from Single-Crystal X-ray Diffraction

La _{1.5} A	.u ₂ Pb _{0.5}	La _S A	AuPb ₃	La ₃ Aı	1 ₃ Ge
Au1-Au1	3.131(1)	Pb1-La2	3.3206(7)	Au1-Au1	3.020(2)
Au1-La1	3.131(1)	Pb1-La2	3.413(2)	Au1-La1	3.026(2)
Au1-La1	3.6158(2)	Pb1-La1	3.4825(7)	Au1-La1	3.125(3)
La1-La1	3.1313(2)	Pb1-La1	3.4844(7)	Au1-Au1	3.170(2)
		Pb1-La2	3.7061(6)	Au1-La1	3.128(3)
		Au1-La2	3.087(1)	Au1-La1	3.231(3)
		La1-La1	3.5016(2)	Au1-La1	3.334(3)
		La1-La2	3.5017(2)	Au1-La1	3.385(3)
		La1-La2	4.034(1)	Au1-La1	3.455(2)
		La1-La2	4.035(1)	La1-Au3	3.059(2)
				Au2-Au3	3.110(2)

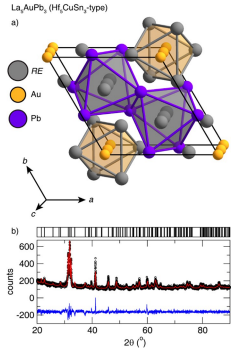


Figure 2. (a) Structure and (b) Le Bail refinement of Hf_5CuSn_3 -type La_5AuPb_3 .

This leads to an overall refined composition of $La_3Au_{3.17(1)}Ge_{0.83(1)}$ for this structure. Contact distances between Au atoms in this network range from 2.8 to 3.5 Å, which is within the standard ranges of aurophilic contacts. Interestingly, through a roughly 45° rotation, the scaffolding lattice and octahedral $La_8(Au/Ge)_{33}$ unit can be viewed together as a highly distorted CsCl-like fragment, in which cubes of Au and Ge house the RE atoms. The intergrowth of this cubic motif with octahedral units suggests that primarily polar interactions between the Au-rich structural units and RE domains stabilize the structural building blocks. These interfaces between the metallically delocalized electrons in the Au bonding network and Au-centered RE polyhedra also hint at some intriguing structural chemistry that can be investigated through DFT electronic structure calculations.

Electronic Structure Investigation with DFT

The electronic stability of polar intermetallic compounds is understood to be controlled by a complicated mixture of structurally related effects. Differing interpretations of the interatomic interactions in polar intermetallic compounds lead

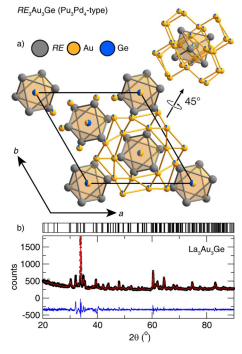


Figure 3. (a) Structure of Pu_3Pd_4 -type RE_3Au_3Ge (RE = La, Ce, Pr, Nd), highlighting the CsCl-like $RE_8(Au/Ge)_{33}$ cluster and (b) the Le Bail refinement of La_3Au_3Ge .

to disparate frameworks for understanding these structure-properties relationships. A classical Hume–Rothery approach would connect stability to valence electron count and Fermi surface–Brillouin zone interactions, ⁵³ whereas looking through the Zintl lens, charge transfer and filled octets are paramount. ⁵⁴ To parse the relative contributions of these effects in highly polar Au-containing RE–Au–Tt phases, density functional theory (DFT) calculations can be employed to investigate the electronic structure of these newly reported compounds directly.

Interactions between the Fermi surface and Brillouin zone can lead to structural stability in intermetallics via the electronic structure. To One way to visualize this is with a density of states (DOS) plot, such as those shown for $La_{1.5}Au_2Pb_{0.5}$, La_5AuPb_3 , and La_3Au_3Ge in Figure 4a. The band population for each structure is plotted against energy, with E_F marking the level below which bands are filled with electrons. In this picture, a peak in the DOS shows a high population of bands at a particular energy value, whereas a local minimum, or pseudogap, in the DOS represents an energy level with relatively few bands. For electronically stable systems, Fermi

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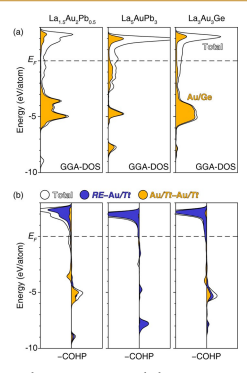


Figure 4. Electronic structure calculations on $La_{1.5}Au_2Pb_{0.5}$, La_5AuPb_3 , and La_3Au_3Ge reveal (a) density of states (DOS) and, (b) crystal orbital Hamilton population (–COHP) curves showing the total (black), RE–Au/Tt (blue), and Au/Tt–Au/Tt (gold) interactions.

surface—Brillouin zone interactions open up pseudogaps in the DOS at $E_{\rm F}$, which lowers the energies of the highest occupied crystal orbitals. Therefore, DOS plots that show the presence of a pseudogap at the Fermi level are correlated with energetically favorable phases, whereas $E_{\rm F}$ falling on a DOS peak is associated with electronic instability. However, the picture is much simpler for the three structures investigated here. The DOS plot near $E_{\rm F}$ is nearly flat for each compound, indicating few, if any, stabilizing effects coming from atomic orbital overlap in these crystal structures. This is further confirmed by investigating the crystal orbital Hamilton population (-COHP) curves. These calculations show the contributions to bonding (positive when plotted as negative COHP) and antibonding (negative when plotted as negative COHP) interactions plotted against energy. In the COHP curve shown for each structure in Figure 4b, the total COHP and specific RE-Au/Tt and Au/Tt-Au/Tt interactions show that very few bonding or antibonding interactions occur near the Fermi level. Thus, the DOS and COHP indicate that these structures are likely not stabilized primarily through orbital overlap.

Next, investigating the role of electrostatic interactions shows that significant differences in the electron affinity (EA) between RE elements and Au (EA_{La} = 48 kJ/mol, EA_{Au} = 222.8 kJ/mol) result in moderate charge transfer. Consequently, electrostatic interactions may have a significant role in the stability of the compounds reported here. To quantitatively assess this charge transfer, DFT-calculated Bader charges were determined for each atom in the La-containing structures. Bader charges are tabulated by assigning the DFT-calculated electron density within a structure to each atom according to the Bader method, which divides space at the zero flux surface of the electron density between atoms. The result of this

calculation for each structure is plotted in Figure 5, where black and white spheres represent cationic and anionic atoms,

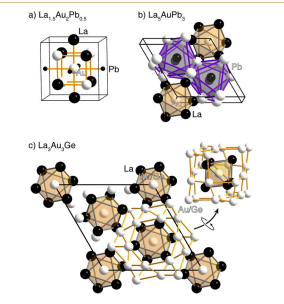


Figure 5. Bader ionicity calculations for (a) $La_{1.5}Au_2Pb_{0.5}$, (b) La_5AuPb_3 , and (c) La_3Au_3Ge . In this scheme, white spheres represent anionic atoms, and black spheres represent cations. The volume of each sphere is proportional to the magnitude of the charge.

respectively, and the volume of each sphere is proportional to the tabulated Bader charge (calculated Bader charges given in Table S5). Examining these Bader schemes for each Lacontaining structure shows that RE atoms are cationic and Au atoms are anionic, as would be expected from their respective electron affinities. In $La_{1.5}Au_2Pb_{0.5}$, the cubic Au coordination environment contains half Au contacts and half La/Pb disordered sites. The ability of La and Pb to occupy the same position in this structure is clarified with the observation that both atoms behave as cations relative to Au. This is not altogether unexpected, as Pb's electron affinity (EA_{Pb} = 35.1kJ/mol) is much closer to La than Au. Pb thus adopts a cationic character in this structure, which maximizes the favorable secondary electrostatic interactions. Intriguingly, the La_{1.5}Au₂Pb_{0.5} phase chooses to adopt the NaTl-type instead of the full Heusler structure, which would seem to be better optimized for maximizing favorable electrostatic interactions. This suggests that the shorter Au-Au contacts of the NaTltype are crucial to the structure preference and that electrostatic effects do not purely govern this compound. A full investigation of the structural preferences in these systems may be an exciting study for future work.

Despite containing the same combination of elements, the ionicity of Pb is inverted in the La₅AuPb₃ structure. In this phase, anionically centered Au@La₆ octahedra are intergrown with ionically opposite La@Pb₆ clusters. This arrangement of charges increases the favorable interactions between positive and negative ions within and between adjacent clusters. Considering that Pb behaves like a cation in La_{1.5}Au₂Pb_{0.5}, these results suggest that Pb atoms can behave amphionically in this compositional space, filling whatever ionic role is needed to suit neighboring La and Au atoms.

The La₃Au₃Ge structure contains the same Au@La₆ octahedra as those seen in La₅AuPb₃. However, without an oppositely charged cluster to satisfy this arrangement, the

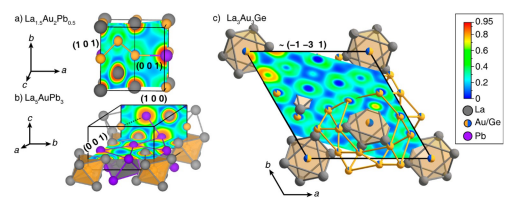


Figure 6. Electron localization function (ELF) slices through a) the (1 0 1) and (0 0 1) planes in $La_{1.5}Au_2Pb_{0.5}$, (b) the (1 0 0) and (0 0 1) planes in La_5AuPb_3 , and (c) the \sim (-1 -3 1) plane in La_3Au_3Ge .

outlying cationic La atoms are instead buffered by the scaffolding network of Au atoms. The atoms in this layer form distorted cubes around each La, housing each cation within an Au-rich coordination environment. A Bader analysis shows that each structure is arranged to create significantly favorable electrostatic interactions, especially between RE and Au atoms as seen in the octahedral clusters in La₅AuPb₃ and La₃Au₃Ge. This aligns with expectations given the electron affinities of these elements. However, other aspects of these phases, such as nearest-neighbor Au—Au contacts in La_{1.5}Au₂Pb_{0.5} and La₃Au₃Ge, seem to run counter to a purely electrostatically driven structural model. This discrepancy implies that other factors may contribute to the stability of these compounds, such as delocalized metallic bonding.

The final potential source of stability in these compounds stems from delocalized metallic bonding. Electron localization function (ELF) calculations are beneficial for identifying delocalized electrons within an intermetallic structure. 56 Figure 6 shows the structures of La_{1.5}Au₂Pb_{0.5}, La₅AuPb₃, and La₃Au₃Ge overlaid with slices of the calculated ELF surface. In these plots, values close to zero (deep blue) represent locations in the electronic structure with a very low probability of locating electrons, such as in the space between atoms. Values close to 1 (red), on the other hand, show locations of highly localized electrons, such as in core states, covalent bonds, or lone pairs. In between, ELF values of 0.5 correspond to fully delocalized electrons, such as those in an electron gas. The ELF plots for these structures all show a combination of these features. In the La_{1.5}Au₂Pb_{0.5} phase, delocalized bonding between Au atoms is readily apparent along the (101) lattice plane, while the (001) layer shows similar metallic character between Au and Pb. This delocalized bonding network, especially along the Au-Au contacts, hints at the favorability of the NaTl-type structure, which blends electrostatic interactions with metallic Au-Au bonding. La atoms in La_{1.5}Au₂Pb_{0.5} remain noninteracting, as seen in the ELF plot, with regions of deep blue ELF minima separating their electron density from their neighbors. A similar picture is seen for La₅AuPb₃ and La₃Au₃Ge, where the La atoms are also highly localized, and there is significant delocalization and interaction between Au and the Pb or Au/Ge scaffolding atoms in both structures. Thus, this supports our overall picture of these phases as polar intermetallics, containing a mixture of electrostatically driven interactions, such as those between RE and Au atoms, and delocalized metallic bonding, such as in the Au-Au and Au-Tt in all structures.

Altogether, an analysis of the electronic structure in these phases gives us a complete picture of the bonding interactions contributing to their stability. Au-containing polar intermetallics offer a step into a region of solid-state composition space where structures are not controlled by either Fermi Surface—Brillouin Zone interactions or by the filling of octets through complete charge transfer. The characteristic mixture of delocalized metallic bonding and favorable electrostatic interactions potentially presents a unique pathway for forming highly unusual metallic oxidation states and stabilizing structures only accessible through gold's extraordinary elemental properties.

CONCLUSION

A synthetic investigation into the underexplored RE-Au-Tt (RE = La, Ce, Pr, Nd; Tt = Ge, Pb) composition space revealed seven novel compounds in three structure types: NaTl-type La_{1.5}Au₂Pb_{0.5} and Nd_{2-x}Au₂Pb_x, Hf₅CuSn₃-type La₅AuPb₃, and Pu₃Pd₄-type RE₃Au₃Ge (RE = La, Ce, Pr, Nd). Each compound was synthesized through the direct hightemperature reaction of elements and subsequent annealing at 850 °C. These compounds are representative polar intermetallics, which are notable for their complex structural features originating from a combination of electronic, electrostatic, and metallic bonding schemes. Au-containing polar intermetallics are of particular interest due to Au's uniquely high electron affinity among nonhalide elements, which enables the stabilization of ternary structures that are not realizable in other systems. Deconvolution of the bonding contributions in these phases can be determined through a DFT investigation of each compound's electronic structure. The density of states and crystal orbital Hamilton populations reveal approximately optimized bonding interactions without a notable pseudogap at the Fermi level, indicating that Fermi surface-Brillouin zone interactions do not govern stability. Visualizations of the calculated Bader charges in these compounds show that electrostatic forces are a vital stabilizing force in these structures, where the Au and Tt bonding network maximizes the anionic-cationic interactions. To finally examine the metallic bonding, ELF calculations indicate delocalized electrons in the Au/Tt bonding networks, while RE atoms remain electronically isolated. RE-Au-Tt polar intermetallic compounds present a composition space ripe for exploring novel materials with complex crystalline and magnetic structures.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsorginorgau.1c00057.

Powder X-ray diffraction data, Anisotropic displacement parameters for La_{1.5}Au₂Pb_{0.5}, La₅Au₂Pb₃, and La₃Au₃Ge, DFT total energies, DFT optimized geometries, DFT-calculated Bader charges, and spin—orbit coupling DOS and COHP curve comparison for La_{1.5}Au₂Pb_{0.5} (PDF)

Accession Codes

CCDC 2129946–2129948 contain the supplementary crystallographic data for this paper. These data can be obtained free of charge via www.ccdc.cam.ac.uk/data_request/cif, or by emailing data_request@ccdc.cam.ac.uk, or by contacting The Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge CB2 1EZ, UK; fax: +44 1223 336033.

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Notes

The authors declare no competing financial interest.

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