Structural and Electronic Flexibility in Hydrides of Zintl Phases with Tetrel-Hydrogen and Tetrel-Tetrel Bonds

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Abstract: The hydrogenation of Zintl phases enables the formation of new structural entities with main group element-hydrogen bonds in the solid state. The hydrogenation of SrSi, BaSi and BaGe yields the hydrides SrSiH_{132}, BaSiH_{133}, and BaGeH_{134}. The crystal structures show a six-fold superstructure compared to the parent Zintl phase and were solved by a combination of X-ray, neutron and electron diffraction and the aid of DFT calculations. Layers of connected HSi$_r$ (HBa$_s$) tetrahedra containing hydride ions alternate with layers of infinite single- and double-chain polyanions, where hydrogen atoms are covalently bound to silicon and germanium. The idealized formulae AeTtH$_x$ (Ae = alkaline earth, Tt = tetrel) can be rationalized with the Zintl-Klemm concept according to (Ae$^{2+}$)$_2$[(TrH)$_3$]$(TtH^x$)$(H^x$)$_y$, where all Tt atoms are three-binding. The non-stoichiometry (SrSiH$_{132}$, x = 0.17(2); BaGeH$_{134}$, x = 0.10(3)) can be explained by additional π-bonding of the Tt chains.

The formation of covalent element-hydrogen bonds is a common theme in synthetic chemistry. Recently, new polymeric triel- and tetrel-hydrogen units were found in the solid state by hydrogenation of Zintl phases or by direct synthesis in hydrogen atmosphere.[1] Examples are triel based polyanions -$[(TtH)_{3}Tt]$ and -$[(TtH)Tt]$ forming puckered honeycomb-layers with hydrogen bound to the triel atom in AeTr$_2$H$_2$ and AeTrTtH (Ae = Sr, Ba; Tr = Al, Ga; Tt = Si, Ge, Sn), respectively,[2] and pseudo-alkane like moieties, e. g. neopentane like [Ga(Ga$_3$H$_6$)$_3$].[3,4] Hydrogen containing tetrel based anions comprise monomeric SiH$_3^-$ in ASiH$_3$ (A = K-Cs),[5,6] infinite single chains -$[SiH]$ in BaSiH$_{12}$,[6] and infinite triple chains -$[HTrH]^+$ in CaSiH$_{15}$, SrGeH$_{12}$, and BaSnH$_{12}$.[6] For the latter, the parent Zintl phases AeTt crystallize in the Cr6 structure type (space group Cmcm) and exhibit infinite polyanionic silicon and germanium zigzag chains -$[Tt^3]$ in agreement with the Zintl-Klemm concept. Since the hydrogenation of CaSi to CaSiH$_{15}$ is reversible, this system is a potential hydrogen storage material.[6,7] The hydrogenation of Zintl phases is a very promising yet barely explored synthetic pathway to new main group element-hydrogen entities in the solid state. In this contribution, we extend this approach and report on new main group-hydrogen structural units in the hydrides of SrSi, BaSi and BaGe. We will focus on the rationale behind the multitude of polyanionic structural motifs, the formation of new tetrel-tetrel and tetrel-hydrogen bonds in the solid and the hydrogen content of the hydrides.

The Zintl phases SrSi and BaGe were hydrogenated at 453 K and 473 K under hydrogen gas pressures of 9.0 and 5.0 MPa, respectively, forming SrSiH$_{132}$, x = 0.17(2) (1.27 wt-% hydrogen, elemental analysis: 1.16(6) wt-%), and BaGeH$_{134}$, x = 10.17(3) (0.74 wt-% hydrogen, elemental analysis: 0.73(3) wt-%, see Supplementary Information for details). The hydrides are gray powders sensitive against air and moisture. Exposed to air, they yield amorphous products. The color of BaGeH$_{134}$ changes to yellow in the course of this reaction, which is probably caused by hydrolysis to polygermane, (GeH$_3$)$_x$. A similar color change was seen for the moisture induced decomposition of SrAlGeH and BaAlGeH where the formation of Ge-H bonds was followed by IR spectroscopy.[8]

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Figure 1. Top left: SAED pattern of a SrSiD$_{132}$ crystal (inset does not scale linearly). Top right: Simulation using the final model (Pnma, sixfold superstructure regarding the parent Zintl phase), all observed reflections are explained. Bottom left: Simulation using the model with half-occupied hydrogen sites (Cmcm, threefold superstructure), rows of reflections are missing due to C centering. Bottom right: Simulation based on the CaSiH$_{15}$ structure type, additional row of reflections explained, but mismatch for position and quantity. For corresponding stereographic projection and further SAED patterns with simulations see supplementary information (Figs. S4, S5).
The crystal structures of SrSiH₅₋ₓ, x = 0.17(2) and BaGeHₓ₅₋ₓ, x = 0.10(3) were elucidated by a combination of electron, X-ray and neutron powder diffraction. The latter two clearly showed a threefold superstructure regarding the parent Zintl phase with $d_{\text{inter}} \approx 3 d_{\text{Zintl phase}}$, i.e., perpendicular to the $\sqrt[3]{2}$ zigzag chains. Structure solution and refinement first led to a model in space group type Cmcm with two fully and two ca. 50% occupied hydrogen sites. Selected-area electron diffraction (SAED) patterns of the Sr compound indicated a further cell doubling, i.e., a six-fold superstructure (Fig. 1). With the help of DFT relaxations of various crystal structure candidates, a model with completely ordered hydrogen distribution was found in space group Pnma (C₃₀ = 6 Pnma, Fig. 2). Joint Rietveld refinements using laboratory X-ray and neutron diffraction data of the deuterides revealed vacancies of up to ca. 20% on some of the hydrogen sites (for details of the structure solution and refinement cf. Supplementary Information). SAED patterns of a BaSi hydride sample suggested the presence of a phase isotypic to SrSiH₅₋ₓ and BaGeHₓ₅₋ₓ. The formerly unassigned side phase in BaSiHₓ samples [5] is thus identified as BaSiH₅₋ₓ (Figs. S8-S10 in Supplementary Information).

The hydrides of SrSi and BaGe are non-stoichiometric with respect to hydrogen and crystallize in space group type Pnma (SrSiH₅₋ₓ: hydride: $a = 15.1009(6) \, \AA$, $b = 3.9512(18) \, \AA$, $c = 24.6298(12) \, \AA$, deuteride: $a = 15.070(2) \, \AA$, $b = 3.9593(2) \, \AA$, $c = 24.642(2) \, \AA$; BaGeHₓ₅₋ₓ: hydride: $a = 15.6253(9) \, \AA$, $b = 2.4308(2) \, \AA$, $c = 25.8728(14) \, \AA$; deuteride: $a = 15.804(3) \, \AA$, $b = 2.4293(6) \, \AA$, $c = 25.855(4) \, \AA$). [16] Tab. S5, S7 and S8 in Supplementary Information). The crystal structures can be rationalized in their idealized composition $\text{AeTM}_2$ according to the Zintl concept as $(\text{Ae}^2)^+ (\text{TH}_x) (\text{T}^2\text{H}_2) (\text{H}_x)^2$, i.e., they contain hydride anions as well as hydrogen covalently bound to the polyanions. Having a formal charge of 1-, all tetrel atoms are expected to form three covalent bonds, either to three tetrel atoms or to two tetrel and one hydrogen atom. The polyanions show two structural motifs (Fig. 2). In the first one, the parallel $\sqrt[3]{2}$ zigzag chains, which are already present in the parent Zintl phase, are inclined towards each other and form additional tetrel-tetrel bonds: This results in double chains $\sqrt[3]{2}$, which run parallel to the double chain. The crystal structure is clearly stabilized by strong ionic bonding from additional hydride ions in $\text{Ae}_2$ tetrahedra (Fig. 2). Crystal structure (for both) and hydrogen content (for the BaSi hydrides) contradict an earlier report of hydrides of AesSi Zintl phases, where, however, hydrogen positions were not determined. [10] The crystal structure shows a close relationship to the parent Zintl phase. The chain direction of the polyanions ([001] in the Zintl phase, [010] in the hydride) shrinks by 2% upon formation of the hydride. A considerable expansion of more than 30% occurs in the direction perpendicular to the $\text{Ae}_2$ tetrahedra sheets ([010] in the Zintl phase, [100] in the hydride) (Fig. 2) due to the incorporation of hydrogen. Connecting the polyanions of the Zintl phase to double chains leads to a reduction of 15% in that direction ([100] in the Zintl phase, [001] in the hydride, accounting for the sixfold superstructure). These strongly anisotropic effects of hydrogenation certainly reflect the bonding situation, which was investigated by quantum-mechanical calculations (DFT).

In zigzag chain direction [010], tetrel-tetrel bonds are 2.41 Å [2.438 Å (Si) and 2.58 Å [2.633 Å (Ge)] (DFT results in brackets, Fig. 2, e1-3) while the additional chain-connecting bond (Fig. 2, e4) is longer with 2.59 Å (Si) and 2.62 Å (Ge). This is not the case for DFT calculated values of the hypothetical stoichiometric composition $\text{AeTM}_2$ (2.443 Å (Si) and 2.655 Å (Ge)). The tetrel-tetrel distances are comparable to those in the pristine Zintl phases, Si-Si-Si angles range from 96° to 111° (Ge-Ge-Ge: 101° to 107°) and are in a similar range as in the chair configuration of black phosphorus (P-P-P: 102.09° and 96.34°) [17] and gray arsenic (As-As-As: 96.64°) [18] respectively.

Refined distances Si-D / Ge-D (averaged d(Si-D) = 1.62 Å [1.650 Å], averaged d(Ge-D) = 1.58 Å [1.588 Å]; DFT results in brackets, Fig. 2, e1-3) indicate covalent interactions. Si-D distances are in between those found in $\beta$-KSiD₃ (1.537(8) - 1.545(6) Å) [19] and BaSiD₃ (1.641(5) Å), while Ge-D distances are somewhat longer than in molecular GeD₃ (1.517(3) Å at 5 K) [19] or SrGeD₃ (1.521(9) Å). [20] Sr-D and Ba-D distances reach from 2.35(4) to 2.57(4) Å and 2.41(6) to 2.73(6) Å, respectively, and are comparable to those in $\text{DAe}_2$ tetrahedra of the binary deuterides SrD (average 2.44 Å and 2.62 Å, respectively), [19] thus
underlining the ionic nature of bonding within the HAE₄ tetrahedra sheets.

Hydrogen atoms covalently bound to the polyanions do not show full occupation (85(2)% in BaGeHₓ₄, 77.2(8)% in SrSiHₓₓ₋₄ₓ). There is no hint to further ordering, not even at low temperature, as studied by neutron diffraction on BaGeHₓ₄ at 10 K (Supplementary information, S3). Such hydrogen-atom vacancies are an intrinsic feature of Zintl-phase hydrides AE₄TtHₓ (AE = Ca-Ba, Tt = Si–Sn, 1 < y < 2).[1-3] They may be related to additional π-bonding due to depopulation of π* bands upon the formation of hydride anions. The effect of π-bonding has been discussed in detail for the parent Zintl-phase silicides.[4-7] Additional π-bonding in non-stoichiometric samples can be assumed comparing DFT-derived bond lengths, which were calculated on idealized structures with full occupation. Experimentally observed tetrel-tetrel distances along the chain direction [010] are systematically shorter (π-bonded) while chain-connecting bonds are longer than calculated distances.

The density of states (DOS) (Fig. 3) was calculated for SrSiHₓₓ and BaGeHₓ₄ using the relaxed structures with an idealized full hydrogen site occupation. While the DOS of BaGeHₓ₄ hints at a band gap as expected for an electron-precise Zintl-phase hydride, SrSiHₓₓ has significant electron density at the Fermi level and shows only a pseudo-gap. This could explain the non-stoichiometry, which is more pronounced in the latter compound, by electronic effects.

BaGeHₓ₄₋₄ and SrSiHₓₓ₋₄ are considered the missing link between SrGeHₓ₄₋₁ and with three tetrel zigzag chains connected to form a triple-chain polyanion -[HGeHₓ⁺] and BaSiHₓ₋₂ with a single chain -[SiHₓ₋₂]⁻. They underline the rich structural chemistry of the hydrides of Zintl phases and represent new structural motifs of main-group element chemistry. Examining the hydrides of CrB type Zintl phases AE₄TtHₓ (AE = Ca–Sr, Tt = Si–Ge, Sn) reveals a remarkable compositional, structural and electronic flexibility. There are two main positions for hydrogen atoms, in AE₄ tetrahedra and attached to tetrel polyanions, both of which may be under-occupied, and there are various types of polyanions in the solid state. Single tetrel zigzag chains may be saturated by hydrogen atoms like in BaSiHₓ₋₂ or such chains may be connected to form double chains like in HGeHₓ₋₄ or triple chains like in CaSiHₓ₋₂, γ-SrGeHₓ₋₂, and BaSnHₓ₋₂. Alternatively, none of the above may occur for hydrogen-poor phases α- and β-SrGeHₓ₋₂ where the electronic balance is adjusted by the Ge–Ge bond length and the degree of π-bonding.

Thus, three types of structural and electronic compensation to the presence of hydrogen may be distinguished: (1) Increasing connectivity from two- to three-bonding by linking of single to double or triple chains, thus considering hydrogenation as an oxidation, e. g. [Si(2b)³⁺], + ½ H₂ = [Si(3b)⁺], + H; (2) covalent bonds to hydrogen, e. g. [SiH⁺], + H₂ = [SiH₂⁺], + H; (3) increasing π-bonding in the polyanionic chains by depopulating π* bands upon formation of hydride anions. The balance between (1) and (2) strongly depends on geometrical restrictions implied by the alkaline earth atoms. With their increasing size, binding hydrogen covalently (2) outweighs the linking of polyanions (1), because of its higher space requirements. The highest degree of condensation is found in CaSiHₓₓ₋₄ and BaSnHₓ₋₂ where the layers of hydride-filled AE₄ tetrahedra are strongly puckered to fit to the translational period of the polyanionic layers. Complete condensation to hydrogen-free layer-like polyanions -[SiHₓ₋₂]⁻ is thus unlikely due to geometrical restrictions in these AE₄Tt compounds. For smaller hydrogen contents, however, the electronic flexibility of the π electron reservoir seems sufficient and only some strengthening of existing Si-Si and Ge-Ge bonds are found instead of forming new ones. Yet another solution to the optimization of space requirements and electronic structure is found in NdGaHₓ₋₁, where hydrogen bridges two gallium zigzag chains approximately halfway between them.[16]

To conclude this geometric reasoning, the combination of large alkaline earth and small tetrel atoms favors high hydrogen contents and single polyanionic chains, e. g. -[SiHₓ₋₂]⁻ in BaSiHₓ₋₂, while small alkaline earth and large tetrel atoms should support the maximum degree of polyanion condensation. However, the tendency of the tetrel atoms to form stable covalent bonds to hydrogen limits this endeavor as seen for the inertness towards hydrogen (SrSn) or tendency to decomposition (SrPb, BaPb).[17] Varying the combinations of alkaline earth and tetrel atoms or substituting them by other elements gives ample opportunity to synthesize tetrel-tetrel and tetrel-hydrogen bonds in the solid. Such considerations may serve as a guide for the search for further compounds with new main group element-hydrogen entities or higher hydrogen content.

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[9] Further details on the crystal structure investigations may be obtained from the Fachinformationszentrum Karlsruhe, 76344 Eggenstein-Leopoldshafen, Germany (fax: (+49)7247-808-666; e-mail: crysdata@fiz-karlsruhe.de), on quoting the depository number numbers CSD-433047, and -433048.
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The Zintl phases SrSi, BaSi and BaGe form hydrides $\text{AeTtH}_{5-x}$, $\text{Ae} = \text{Sr}, \text{Ba}$; $\text{Tt} = \text{Si, Ge}$, with new single- and double-chain polyanions characterized by additional tetrel-hydrogen and tetrel-tetrel bonds.

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Structural and Electronic Flexibility in Hydrides of Zintl Phases from Tetrel-Hydrogen to Tetrel-Tetrel Bonds

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