IOPscience

Home Search Collections Journals About Contact us My IOPscience

A density functional study of the high-pressure chemistry of $MSiN_2(M = Be, Mg, Ca)$: prediction of high-pressure phases and examination of pressure-induced decomposition

This article has been downloaded from IOPscience. Please scroll down to see the full text article. 2009 J. Phys.: Condens. Matter 21 275407 (http://iopscience.iop.org/0953-8984/21/27/275407)

View the table of contents for this issue, or go to the journal homepage for more

Download details: IP Address: 129.252.86.83 The article was downloaded on 29/05/2010 at 20:30

Please note that terms and conditions apply.

J. Phys.: Condens. Matter 21 (2009) 275407 (9pp)

A density functional study of the high-pressure chemistry of $MSiN_2(M = Be, Mg, Ca)$: prediction of high-pressure phases and examination of pressure-induced decomposition

S Rebecca Römer¹, Peter Kroll² and Wolfgang Schnick¹

¹ Department Chemie und Biochemie, Lehrstuhl für Anorganische Festkörperchemie, Ludwig-Maximilians-Universität München, Butenandtstraße 5-13 (D), D-81377 München, Germany

² Department of Chemistry and Biochemistry, University of Texas at Arlington,

700 Planetarium Place, Arlington, TX 76019-0065, USA

E-mail: wolfgang.schnick@uni-muenchen.de

Received 22 December 2008, in final form 15 April 2009 Published 12 June 2009 Online at stacks.iop.org/JPhysCM/21/275407

Abstract

Normal pressure modifications and tentative high-pressure phases of the nitridosilicates MSiN₂ with M = Be, Mg, or Ca have been thoroughly studied by density functional methods. At ambient pressure, BeSiN₂ and MgSiN₂ exhibit an ordered wurtzite variant derived from idealized filled β -cristobalite by a C1-type distortion. At ambient pressure, the structure of $CaSiN_2$ can also be derived from idealized filled β -cristobalite by a different type of distortion (D1-type). Energy-volume calculations for all three compounds reveal transition into an NaCl superstructure under pressure, affording sixfold coordination for Si. At 76 GPa BeSiN₂ forms an LiFeO₂-type structure, corresponding to the stable ambient-pressure modification of LiFeO₂, while MgSiN₂ and CaSiN₂ adopt an LiFeO₂-type structure, corresponding to a metastable modification (24 and 60 GPa, respectively). For both BeSiN₂ and CaSiN₂ intermediate phases appear (for $BeSiN_2$ a chalcopyrite-type structure and for $CaSiN_2$ a $CaGeN_2$ -type structure). These two tetragonal intermediate structures are closely related, differing mainly in their c/aratio. As a consequence, chalcopyrite-type structures exhibit tetrahedral coordination for both cations (M and Si), whereas in CaGeN₂-type structures one cation is tetrahedrally (Si) and one bisdisphenoidally (M) coordinated. Both structure types, chalcopyrite and CaGeN₂, can also be derived from idealized filled β -cristobalite through a B1-type distortion. The group–subgroup relation of the BeSiN₂/MgSiN₂, the CaSiN₂, the chalcopyrite, the CaGeN₂ and the idealized filled β -cristobalite structure is discussed and the displacive phase transformation pathways are illustrated. The zero-pressure bulk moduli were calculated for all phases and have been found to be comparable to compounds such as α -Si₃N₄, CaIrO₃ and Al₄C₃. Furthermore, the thermodynamic stability of BeSiN₂, MgSiN₂ and CaSiN₂ against phase agglomerates of the binary nitrides M₃N₂ and Si₃N₄ under pressure are examined.

S Supplementary data are available from stacks.iop.org/JPhysCM/21/275407

1. Introduction

Silicon nitride and related nitridosilicates (e.g. $\alpha/\beta/\gamma$ phases of Si₃N₄ [1–4], M₂Si₅N₈ (M = Ca, Sr, Ba, Eu) [5–7], MSiN₂ (M = Be, Mg, Ca) [8–10]) exhibit interesting and useful physical properties. Good wear resistance, high decomposition temperature, exceptional oxidation stability [11, 12], luminescence [13, 14] and nonlinear optical behaviour [15] render them interesting for industrial applications.

Throughout the last two decades, an increasing number of dense high-pressure oxosilicates have been synthesized and their material properties have been characterized [16, 17]. The high technological impact of these high-pressure silicate phases is vividly illustrated by the discovery of the extraordinary mechanical hardness of stishovite (rutile-type SiO₂) and post-stishovite (α -PbO₂ SiO₂) [18, 19]. Material properties of nitrides are often superior to those of the corresponding oxides, attributed to a higher covalency and degree of cross-linking in nitride structures [20]. Indeed, the discovery of spinel-type γ -Si₃N₄ triggered broad research efforts targeting new high-pressure nitrides and nitridosilicate with coordination behaviours of Si beyond that of a tetrahedron [3, 4].

The high-pressure behaviours of the $MSiN_2$ compounds (M = Be, Mg, Ca, Sr, Ba) is a worthwhile endeavour, since these compounds find application in several commercial/industrial processes. Moreover, they are of significant importance for the high-pressure chemistry of the prominent phosphor host lattices $M_2Si_5N_8$ (M = Ca, Sr, Ba) [21, 22]. MgSiN₂ is a thermally stable ceramic (up to 1400 °C [23]), and due to its wurtzite-analogous structure, discussed to substitute for AlN [24]. It is also widely used as a sintering agent and additive for Si₃N₄ ceramics [23, 25, 26]. CaSiN₂, besides possessing promising properties for ceramics applications [27], is an auspicious phosphor host lattice. Doped with Eu²⁺ or Ce³⁺ it is considered for use in LEDs [28, 29]. CaSiN₂:Ce³⁺ is one of the rare phosphor materials that can be excited by yellow-green light [29].

In this work we report on density functional calculations of ground-state structures and of several possible high-pressure phases of BeSiN₂, MgSiN₂ and CaSiN₂. We will illustrate their structural properties, calculate transition pressures and access their stability against decomposition into binary nitrides at elevated pressure. Results of the higher homologues, SrSiN₂ and BaSiN₂, will be reported elsewhere, as their ambientpressure phases are structurally quite different from those of BeSiN₂, MgSiN₂ and CaSiN₂ [10].

2. Method

Density functional calculations [30] are performed with the Vienna *ab initio* simulation package (VASP) to obtain optimized structures, total energies and properties. VASP combines the total energy pseudopotential method with a planewave basis set [31–33]. The generalized gradient approximation (GGA) [34], as well as the local density approximation (LDA) for the electron exchange and correlation energy, is used

Table 1.Calculation details.			
	Formula units / cell	k-point mesh	
α -CaSiN ₂	16	$4 \times 2 \times 2$	
α -MSiN ₂ (M = Be, Mg)	4	$4 \times 4 \times 4$	
β MSIN (M - Pa Ca)	4	5 ~ 5 ~ 5	

α -MSiN ₂ (M = Be, Mg)	4	$4 \times 4 \times 4$
β -MSiN ₂ (M = Be, Ca)	4	$5 \times 5 \times 5$
(chalcopyrite-type)		
γ -CaSiN ₂ , β -MgSiN ₂	3	$7 \times 7 \times 2$
(m-LiFeO ₂ -type)		
γ -BeSiN ₂	4	$5 \times 5 \times 4$
(s-LiFeO ₂ -type)		

and the projector-augmented-wave (PAW) method [35] is employed. We use 500 eV for the cutoff energy for the expansion of the wavefunction into the plane-wave basis set. Atomic forces are converged below 5×10^{-3} eV Å⁻¹ and stresses below 0.1 GPa. The Brillouin zone is integrated by the scheme developed by Monkhorst and Pack [36].

We made a comprehensive search of ABX_2 structure types within the international crystal structure database (ICSD). Targeting on structures with cation coordination of four and larger only, we selected more than a dozen candidate structures. These were then decorated according to the composition MSiN₂ and subsequently optimized in all parameters. We also invented additional structures by creating an ordered variant of AX types with high cation coordination. In total, we investigated about 20 structures—and table 1 contains only the most relevant types. While this approach is not a parameterfree global search, we are nevertheless confident that we investigate a wide variety of structural motives. Most of them are indeed observed in similar compounds (oxides).

Structural optimizations are obtained through relaxation of all structural parameters, positions as well as cell parameters. Detailed information on the cells used for calculation and the k-point meshes are given in table 1.

The GGA functional significantly better describes relative energies of structures with different coordination of the atoms. See, for instance, the pressure for the β - to γ -Si₃N₄ transition: 6 GPa are estimated using the LDA and 12.5 GPa within the GGA—with an experimental value of about 13 GPa [3]. Therefore, GGA is the better choice in comparison to the LDA for studying structural transformations at high pressure. Consequently, while we also performed all calculations using the LDA, all data reported in this work, enthalpy differences and transition pressures, are based on GGA calculations unless otherwise noted.

To obtain the zero-pressure bulk modulus, we vary the volume around the zero-pressure volume V_0 . We then fit the resulting E-V data to Murnaghan's equation of state (EOS) [37] to extract the bulk modulus B_0 of a structure. We note that using Birch's or Vinet's equations of state [38, 39] give similar values. From the fitted EOS the pressure p is calculated by numerical differentiation and the enthalpy H is calculated via H = E + pV. Neglecting entropy effects, we choose the enthalpy difference ΔH as an appropriate measure to compare the relative stability of solid-state structures under pressure.

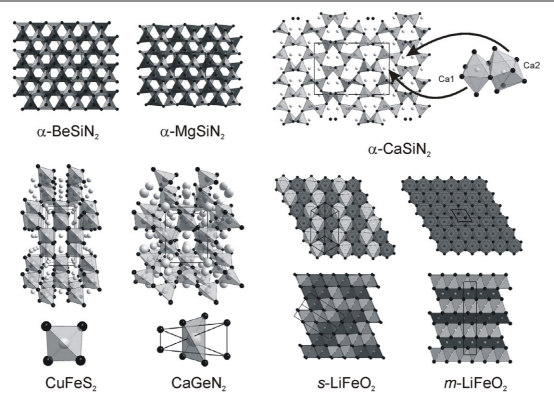


Figure 1. MSiN₂ structures: (1) structure of α -BeSiN₂ and α -MgSiN₂ (view along [001], MN₄ tetrahedra are depicted dark grey, SiN₄ tetrahedra light grey); (2) structure of α -CaSiN₂ (view along [100]; Ca atoms are depicted light grey, Si atoms dark grey, N atoms black); (3) structures of β -BeSiN₂ (CuFeS₂, c/a ratio 2.01) and β -CaSiN₂ (CaGeN₂, c/a ratio 1.36) (view along [010] with SiN₄ tetrahedra drawn; bottom BeN₄ tetrahedron and CaN₈ bisdisphenoid, respectively); (4) structures of γ -BeSiN₂ (s-LiFeO₂, top: octahedra layer, view along [0.5 0.5 0.25]; bottom: view along [-0.5 - 0.5 0]) and β -MgSiN₂ and γ -CaSiN₂ (m-LiFeO₂, top: octahedra layer, view along [001]; bottom: view along [010]) (M = Be, Mg, Ca atoms are depicted light grey, Si atoms dark grey, N atoms black, MN₆ octahedra are depicted light grey, SiN₆ octahedra dark grey).

3. Results and discussion

3.1. BeSiN₂

 α -BeSiN₂ crystallizes in an ordered wurtzite-type structure (space group $Pna2_1$, no. 33) [8], which can also be described as a C1-type distortion of the idealized filled C9 structure of β -cristobalite [40] (figure 1). We identified two candidate high-pressure structures: (I) β -BeSiN₂, which exhibits a chalcopyrite-like structure with tetrahedrally coordinated Be and Si (space group $I\bar{4}2d$, no. 122) [41]. This structure can be derived from β -cristobalite as well, however, in this case through a B1-type distortion [40] (figure 1). Therefore, both α - and β -BeSiN₂ are three-dimensional MN₄ tetrahedral networks with identical bonding topology, built up of cornersharing tetrahedra. (II) γ -BeSiN₂, which adopts an LiFeO₂ structure (further on called s-LiFeO₂ as it corresponds to the stable low-temperature modification of LiFeO₂) with tetragonal space group $I4_1/amd$ (no. 141) [42]. It is a rock salt superstructure (doubled unit cell) and both Be and Si are octahedrally coordinated by N (figure 1). In the supplementary data (available at stacks.iop.org/JPhysCM/21/275407) we detail the crystallographic data of the three polymorphs of $BeSiN_2$ (as well as those of all later described $MSiN_2$ phases) and compare them with available experimental data. Further information on bond lengths in BeSiN₂ (as well as MgSiN₂

Table 2. E_0 , V_0 , B_0 and ρ_0 for α -, β - and γ -BeSiN₂.

	$E_0/\mathrm{f.u.}~(\mathrm{eV})$	$V_0/f.u.$ (10 ⁶ pm ³)	B_0 (GPa)	$ ho_0 ({ m g}{ m cm}^{-3})$
α -BeSiN ₂	-30.897	33.87	220	3.19
β -BeSiN ₂		33.82	222	3.20
γ -BeSiN ₂		28.29	244	3.82

and $CaSiN_2$) is given in the supplementary data (available at stacks.iop.org/JPhysCM/21/275407).

Referring to our ambient-pressure GGA calculations, α -BeSiN₂ has the lowest energy of the four polymorphs (-30.902 eV/f.u.) and also the lowest density, ρ = 3.19 g cm⁻³ (exp. value 3.24 g cm⁻³ [8]). β -BeSiN₂ is only about 0.005 eV/f.u. higher in energy and about 0.3% denser (computed 3.20 g cm⁻³). This result is not surprising given the close resemblance of the two modifications. γ -BeSiN₂ then exhibits the highest density. With ρ = 3.82 g cm⁻³ this octahedral structure is about 20% denser than both tetrahedral structures, but also about 2.1 eV per formula units higher in energy (table 2).

The zero-pressure bulk moduli of the three polymorphs are 220 GPa for α -BeSiN₂, 222 GPa for β -BeSiN₂ and 244 GPa for γ -BeSiN₂. Shaposhnikov *et al* [43] and Petukhov *et al* [44] report LDA values of B_0 of 264 GPa for α -BeSiN₂ and of 242

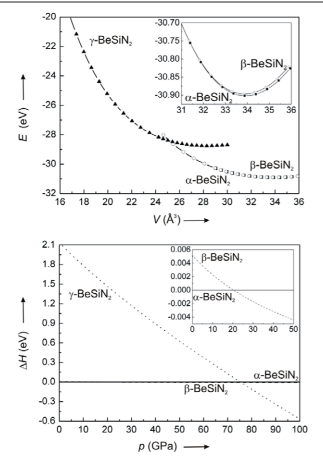


Figure 2. E-V (top) and enthalpy–pressure (bottom) diagram for all three considered phases of BeSiN₂ (E-V data points connected by a spline fit; enthalpy–pressure diagram from Murnaghan EOS evaluation).

and 240 GPa for β -BeSiN₂, respectively, but did not consider a γ -BeSiN₂. Their values match our LDA results for the bulk modulus (241 GPa for both structures). With respect to the bulk moduli, both β -BeSiN₂ and γ -BeSiN₂ are likely to be hard materials, ranking between B₄C (200 GPa) [45], α -Si₃N₄ (229 GPa) [46] and SiC (248 GPa) [45].

In figure 2 the E-V and the enthalpy-pressure curves of the three structures of BeSiN₂ are displayed. Apparently, α -BeSiN₂ is the most stable polymorph of BeSiN₂ for pressures up to 20 GPa, at which it will transform to β -BeSiN₂. Given the small enthalpy differences between α - and β -BeSiN₂, the actual value of the transition pressure has to be taken with care (see the inset in figure 2, bottom diagram). However, once β -BeSiN₂ is formed, a tetrahedral structure will remain stable up to about 76 GPa. At this pressure the structure of γ -BeSiN₂ with octahedral coordination becomes the most stable polymorph of BeSiN₂. The sequence of structures together with transition pressures and density increases is illustrated in figure 3.

$3.2. MgSiN_2$

 α -MgSiN₂ crystallizes in the orthorhombic space group *Pna*2₁ (no. 33) and exhibits the same wurtzite-type structure as

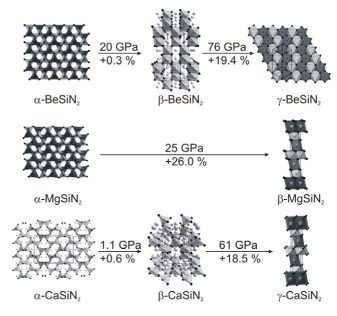


Figure 3. Sequence of structures of $BeSiN_2$, $MgSiN_2$ and $CaSiN_2$ together with transition pressures (in GPa) and values for the density increase (in percentage).

Table 3. E_0 , V_0 , B_0 and ρ_0 for α - and β -MgSiN₂.

	<i>E</i> ₀ /f.u. (eV)	$V_0/f.u.$ (10 ⁶ pm ³)	B_0 (GPa)	$ ho_0 ({ m g}{ m cm}^{-3})$
$\begin{array}{l} \alpha \text{-MgSiN}_2\\ \beta \text{-MgSiN}_2 \end{array}$		43.35 34.42	172 223	3.08 3.88

 α -BeSiN₂ [9] (figure 1). The search for high-pressure polymorphs revealed only β -MgSiN₂ (figure 1) exhibiting yet another LiFeO₂-type structure (space group $R\bar{3}m$, no. 166, denoted m-LiFeO₂: this type corresponds to the metastable low-temperature modification of LiFeO₂ [47]). The m-LiFeO₂ structure can be derived from the CdCl₂ structure (also $R\bar{3}m$), if the unoccupied octahedral sites in CdCl₂ are filled with a second sort of cation. Hence, both s- and m-LiFeO₂ are superstructures of the rock salt structure, differing only in the ordering of cations.

Within the zero-pressure GGA calculations, the ground state of α -MgSiN₂ has an energy of -28.495 eV/f.u. and a density of 3.08 g cm⁻³ (exp. value: 3.13 g cm⁻³ [9]). β -MgSiN₂ is about 1.1 eV higher in energy than α -MgSiN₂ and approximately 26% denser (computed 3.88 g cm⁻³) (table 3). The bulk modulus is 172 GPa for α -MgSiN₂, well within the range of previously calculated (182 GPa [48], 174 GPa [49]) and measured values (146 GPa [50], 184 GPa [51]). For β -MgSiN₂ we calculate a bulk modulus of 223 GPa, which is comparable to the bulk modulus of α -Si₃N₄ (220 GPa) [46].

Figure 4 then shows both the energy–volume and the enthalpy–pressure phase diagrams. Accordingly, α -MgSiN₂ will transform into β -MgSiN₂ at 25 GPa, increasing the coordination for both Mg and Si from four (tetrahedral) to six (octahedral) (figure 3).

Fang *et al* [48] reported a high-pressure transition of α -MgSiN₂ into a CsICl₂-type structure ($R\bar{3}m$, no. 166)

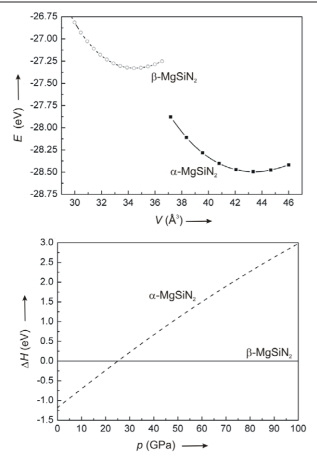


Figure 4. E-V (top) and enthalpy–pressure (bottom) diagram of the two considered phases of MgSiN₂ (E-V data points connected by a spline fit; enthalpy–pressure diagram from Murnaghan EOS evaluation).

and proposed a transition pressure of 16.5 GPa using LDA calculations. The lower transition pressure is explained by the choice of the different functional that artificially favours higher coordination. The structure type of CsICl₂ is, however, misleading. Both CsICl₂ and m-LiFeO₂ (our choice) adopt the same space group, but have very different structural parameters. In a hexagonal setting, the c/a ratios are 1.9269 for CsICl₂ [52] and 4.9899 for m-LiFeO₂ [47] (in a rhombohedral setting the rhombohedral angle is very different). As a consequence, the coordination environments are different, eightfold in CsICl₂ and sixfold in m-LiFeO₂-type. Since Fang *et al* report their optimized structure of β -MgSiN₂ having octahedral coordination, it is better addressed as being the m-LiFeO₂ structure type.

3.3. CaSiN₂

Ambient-pressure α -CaSiN₂ crystallizes in the orthorhombic space group *Pbca* (no. 61) [10]. Corner sharing SiN₄ tetrahedra form a three-dimensional network. The structure is also related to the β -cristobalite structure (D1-type distortion of idealized C9 structure of β -cristobalite) [40]. Ca is in sixand eightfold coordination to nitrogen, respectively (figure 1). A first candidate high-pressure polymorph, β -CaSiN₂, is

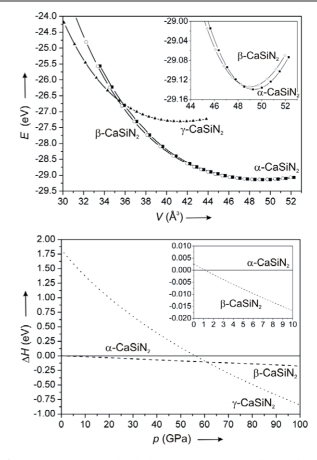


Figure 5. E-V (top) and enthalpy–pressure (bottom) diagram for all three considered phases of CaSiN₂ (E-V data points connected by a spline fit; enthalpy–pressure diagram from Murnaghan EOS evaluation).

isostructural to CaGeN₂ [53] (B1-type distortion of the idealized filled C9 structure of β -cristobalite), which in turn is closely related to the chalcopyrite structure [41]. It crystallizes in the tetragonal space group $I\bar{4}2d$ (no. 122). Si is tetrahedrally coordinated by N. Ca exhibits a 4 + 4 bisdisphenoidal coordination. The main difference between chalcopyrite-type and CaGeN₂-type structures is a different c/a ratio, while the crystallographic position of atoms are quite similar. Both cations in chalcopyrite are tetrahedrally coordinated. CaGeN₂, on the other hand, has one cation tetrahedrally coordinated and the other in bisdisphenoidal coordination (figure 1). The second candidate structure, γ -CaSiN₂, again adopts an m-LiFeO₂-type structure (figure 1) [47].

The energy of α -CaSiN₂ is computed as -29.140 eV per formula unit and its density $\rho = 3.24 \text{ g cm}^{-3}$ (exp. value 3.30 g cm⁻³ [10]). β -CaSiN₂ is only slightly higher in energy, some 0.006 eV, and about 0.6% denser. γ -CaSiN₂ with octahedral coordination of all atoms exhibits the highest density. With $\rho = 3.87 \text{ g cm}^{-3}$ it is about 19% denser than both α - and β -CaSiN₂. The energy of γ -CaSiN₂ is some 1.8 eV higher than that of α -CaSiN₂. The zero-pressure bulk moduli of the three phases are 131 GPa, 126 GPa and 189 GPa for α -, β - and γ -CaSiN₂, respectively (table 4). This places them among compounds as Zr₂InC (127 GPa) [54], Al₄C₃ (130 GPa) [55] and CaIrO₃ (180 GPa) [56]. Along

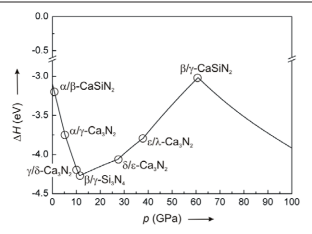


Figure 6. Enthalpy–pressure diagram for the formation of CaSiN₂ from Ca₃N₂ and Si₃N₄. The phase transition of β - into γ -Si₃N₄ as well as the proposed phase transformations of Ca₃N₂ (α -, γ -, δ -, ε - and λ -Ca₃N₂) have been taken into account (the pressure for each transition is indicated by a circle).

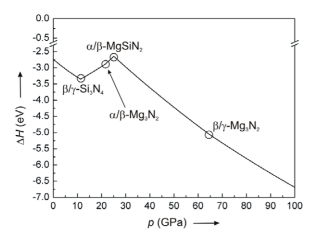


Figure 7. Enthalpy–pressure diagram for the formation of MgSiN₂ from Mg₃N₂ and Si₃N₄. The phase transition of β - into γ -Si₃N₄ as well as the proposed phase transformations of Mg₃N₂ (α -, β - and γ -Mg₃N₂) have been taken into account (the pressure for each transition is indicated by a circle).

the sequence $BeSiN_2$, $MgSiN_2$ and $CaSiN_2$, therefore, $CaSiN_2$ phases show the lowest bulk moduli.

Interestingly, when calculating CaSiN₂ in the MgSiN₂type structure [9], we obtained an even lower energy than that computed for α -CaSiN₂. The energy difference, a tiny 0.001 eV, however, is very small, and well within possible entropy effects or even within a systematic error of DFT calculations. Employing LDA calculations, on the other hand, places the α -CaSiN₂ 0.032 eV below the MgSiN₂ type.

Energy–volume and enthalpy–pressure phase diagrams of CaSiN₂ are shown in figure 5. Accordingly, we find a first transition of α -CaSiN₂ into β -CaSiN₂ already at 1.1 GPa. β -CaSiN₂ will be the most stable polymorph up to 61 GPa, when the structure of γ -CaSiN₂ with all-octahedral coordination is adopted. The sequence of the structures is illustrated in figure 3.

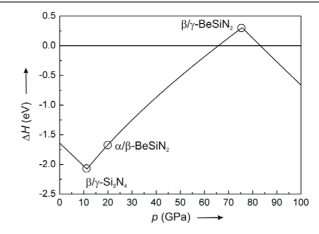


Figure 8. Enthalpy–pressure diagram for the formation of $BeSiN_2$ from Be_3N_2 and Si_3N_4 . The phase transition of β - into γ -Si₃N₄ has been taken into account (no phase transformations of Be_3N_2 are found up to 100 GPa) (the pressure for each transition is indicated by a circle).

Table 4. E_0 , V_0 , B_0 and ρ_0 for α -, β -, γ -CaSiN₂ and CaSiN₂ in the MgSiN₂ structure.

	<i>E</i> ₀ /f.u. (eV)	$V_0/f.u.$ (10 ⁶ pm ³)	B ₀ (GPa)	$ ho_0 ({ m g}{ m cm}^{-3})$
α -CaSiN ₂		49.26	131	3.24
β -CaSiN ₂		48.96	126	3.26
γ -CaSiN ₂		41.31	189	3.87
'MgSiN ₂ '	-29.141	50.64	121	3.15

3.4. Thermodynamic stability of MSiN₂ at high pressure

Finally, we access the thermodynamic stability of $MSiN_2$ compounds versus a phase agglomerate of M_3N_2 (M = Ca, Mg) and Si₃N₄. To achieve this, we took all known highpressure phases of Ca₃N₂ [57], Mg₃N₂ [58] and Si₃N₄ [3, 4] into account (figures 6 and 7). We find that both the ternary phases of CaSiN₂ and MgSiN₂ are always lower in enthalpy than a mixture of the two binary phases M_3N_2 (M = Ca, Mg) and Si₃N₄. This opens up another possible synthesis route for CaSiN₂ and MgSiN₂ and its high-pressure phases, as they should be attainable under high pressure starting from Si₃N₄ and Ca₃N₂ or Mg₃N₂, respectively. High temperatures should accompany the high-pressure syntheses to overcome high activation barriers due to strong bonds in and low interdiffusion coefficients of Si₃N₄ and Ca₃N₂/Mg₃N₂.

BeSiN₂, on the other hand, is stable against decomposition only up to 66 GPa, at which pressure a phase agglomerate of Be₃N₂ and Si₃N₄ is more favourable. Interestingly, above 83 GPa BeSiN₂ should reappear, adopting the γ -BeSiN₂ structure (figure 8). Consequently, γ -BeSiN₂ will be attainable only above 83 GPa, possibly from the binary nitrides.

4. Conclusion

The structures of α -MSiN₂ (M = Be, Mg, Ca), β -BeSiN₂ and β -CaSiN₂ are related to the filled β -cristobalite structure (filled C9 structure) by group–subgroup relations (figure 9). They

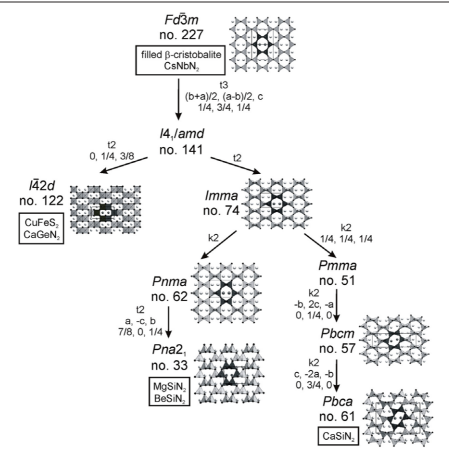


Figure 9. Group–subgroup schema for the relation of the α -MSiN₂ (M = Be, Mg, Ca) and the β -MSiN₂ (M = Be, Ca) structures to the filled β -cristobalite structure.

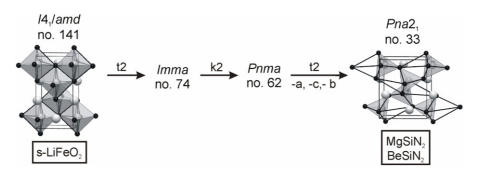


Figure 10. Group–subgroup scheme for s-LiFeO₂ and $BeSiN_2/MgSiN_2$. Both structures are presented in equivalent detail and the polyhedra of the partner structure are enhanced by thick black lines, respectively.

derive from the parent structure (space group $Fd\bar{3}m$, no. 227) by concerted rotations of tetrahedra (for a detailed discussion on these rotation patterns see the works of Thompson *et al* [40] and O'Keeffe and Hyde [59]). Hence, displacive phase transition pathways are conceivable for α -BeSiN₂ (*Pna2*₁) into β -BeSiN₂ (*I*42*d*) and for α -CaSiN₂ (*Pnma*) into β -CaSiN₂ (*I*42*d*). This might well result in quite low activation energy barriers for these two phase transitions.

The structures of γ -BeSiN₂ ($I4_1/amd$, s-LiFeO₂), β -MgSiN₂ ($R\bar{3}m$, m-LiFeO₂) and γ -CaSiN₂ ($R\bar{3}m$, m-LiFeO₂) are all related to the rock salt structure. The difference between the s- and m-LiFeO₂-type structures is a different

ordering of the two types of cations on octahedral sites. In m-LiFeO₂ layers of condensed LiO₆ octahedra alternate with layers of condensed FeO₆ octahedra. In s-LiFeO₂ structures every layer is occupied half by Li and half by Fe. The s-LiFeO₂-type structure seems to be preferred for compounds, with both cations having approximately the same size as in BeSiN₂ ($r(Be^{2+}) = 59$ [60], 31 pm [61]; $r(Si^{4+}) = 54$ [60], 29 pm [61]). If the two cations differ substantially in size as in MgSiN₂ and CaSiN₂ ($r(Mg^{2+}) = 86$ [60], 70 pm [61]; $r(Ca^{2+}) = 114$ [60], 105 pm [61]), the m-LiFeO₂-type structure is favoured, as the height of the octahedral layers can be adjusted to the size of the cation occupying it.

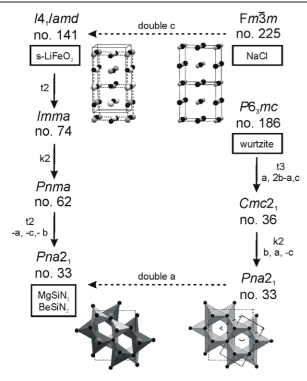


Figure 11. Relation of the NaCl structure to the s-LiFeO₂ structure and of the wurtzite (ZnS) structure to the BeSiN₂ and MgSiN₂ structure (pictures: s-LiFeO₂: double unit cell of NaCl drawn in addition to the tetragonal unit cell; wurtzite: unit cell of wurtzite drawn as well as of wurtzite reduced to $Pna2_1$ and of BeSiN₂ and MgSiN₂).

The s-LiFeO₂ structure adopted by BeSiN₂ (γ -phase) is related to α -BeSiN₂ (and α -MgSiN₂) through a groupsubgroup relation (figure 10). Simply put, by compressing α -BeSiN₂ along two unit cell axes and simultaneously elongating the third (corresponding to the c axis in tetragonal s-LiFeO₂), the distorted hexagonal close packing (hcp) of the anions in α -BeSiN₂ is transformed into a cubic close packing (ccp, distorted as well) and the cations are transferred from the tetrahedral sites in hcp to the octahedral sites in ccp by only a small movement. This transformation is analogous to the wellknown wurtzite-rock salt transition [62]. α -BeSiN₂ exhibits an ordered wurtzite structure and s-LiFeO₂ the corresponding ordered rock salt structure (compare figure 11), only that the ordering of the cations results in a doubled unit cell for s-LiFeO₂ and α -BeSiN₂ if compared to the rock salt and wurtzite structure, respectively. However, as wurtzite-type BeSiN₂ might transform first into a CuFeS₂-type structure according to the enthalpy-pressure phase diagram (figure 2), which is not related to the s-LiFeO2-type structure, it is unclear whether a displacive phase transition along the wurtziterock salt transformation pathway will take place. The m-LiFeO₂-type structure, while also exhibiting an ordered rock salt superstructure, is not related to either of the discussed structures.

It has to be noted that the s-LiFeO₂ structure $(I4_1/amd)$ is not related to the filled β -cristobalite structure, even though the space group sequence $I4_1/amd$ to $Pna2_1$ is identical to that of filled β -cristobalite $I4_1/amd$ to $Pna2_1$. In filled

 β -cristobalite, reduced to $I4_1/amd$, the X atoms of ABX₂ occupy the 8c positions whereas in s-LiFeO₂ they occupy the 8e positions.

5. Summary

We investigated the pressure-dependent phase diagrams of the nitridosilicates $MSiN_2$ (M = Be, Mg, Ca). For $BeSiN_2$ two candidate structures, a CuFeS₂-type (β -BeSiN₂) and a s-LiFeO₂-type (γ -BeSiN₂) were found to become lower in enthalpy at high pressure than the α phases. For MgSiN₂, an m-LiFeO₂-type (β -MgSiN₂) will appear at high pressures. For CaSiN₂ we found two potential high-pressure phases, a CaGeN₂-type (β -CaSiN₂) and a m-LiFeO₂-type (γ -CaSiN₂). In each system, the phases most stable at very high pressure— γ -BeSiN₂, β -MgSiN₂ and γ -CaSiN₂—exhibit ordered rock salt structures with octahedral coordination of Si. The calculated bulk moduli indicate an increase in hardness for these high-pressure phases. We found that CaSiN₂ and MgSiN₂ are thermodynamically stable against decomposition into the binary nitrides up to 100 GPa. BeSiN₂ exhibits a 'stability gap' between 66 and 83 GPa. In this pressure region a phase assemblage of Be₃N₂ and Si₃N₄ is more favourable.

Acknowledgments

Financial support by the Deutsche Forschungsgemeinschaft, (priority programme SPP 1236, project SCHN 377/13 and Kr 1805/10 and Heisenberg-programme Kr 1805/9) as well as the Fonds der Chemischen Industrie (FCI), Germany, is gratefully acknowledged. The authors would further like to thank the Leibniz Rechenzentrum, Munich for computational resources on the Linux Cluster System, as well as the Texas Advanced Computing Center at Austin.

References

- [1] Ruddlesden S N and Popper P 1958 Acta Crystallogr. 11 465
- [2] Hardie D and Jack K H 1957 *Nature* **180** 332
- [3] Zerr A, Miehe G, Serghiou G, Schwarz M, Kroke E, Riedel R, Fue
 β H, Kroll P and Boehler R 1999 Nature 400 340
- [4] Schwarz M, Miehe G, Zerr A, Kroke E, Poe B T, Fuess H and Riedel R 2000 Adv. Mater. 12 883
- [5] Schlieper T and Schnick W 1995 Z. Anorg. Allg. Chem. 621 1037
- [6] Schlieper T, Milius W and Schnick W 1995 Z. Anorg. Allg. Chem. 621 1380
- [7] Huppertz H and Schnick W 1997 Acta Crystallogr. C 53 1751
- [8] Eckerlin P 1967 Z. Anorg. Allg. Chem. 353 225
- [9] David J, Laurent Y and Lang J 1970 Bull. Soc. Fr. Minéral. Cristallogr. 93 153
 Wintenberger M, Tcheou F, David J and Lang J 1980 Z. Naturf.
- B **35** 604 [10] Gál Z A, Mallinson P M, Orchard H J and Clarke S J 2004
- Inorg. Chem. 43 3998
- [11] Hampshire S 1994 Materials Science and Technology vol 11, ed R W Cahn, P Haasen and E J Kramer (Weinheim: Wiley–VCH)
- [12] Nordberg L-O, Nygren M, Käll P-O and Shen Z 1998 J. Am. Ceram. Soc. 81 1461

- [13] Mueller-Mach R, Mueller G, Krames M R, Höppe H A, Stadler F, Schnick W, Juestel T and Schmidt P 2005 Phys. Status Solidi a 202 1727
- [14] Jüstel T, Nikol H and Ronda C 1998 Angew. Chem. 110 3250
 Jüstel T, Nikol H and Ronda C 1998 Angew. Chem. Int. Edn 37 3084
- [15] Lutz H, Joosten S, Hoffmann J, Lehmeier P, Seilmeier A, Höppe H A and Schnick W 2004 J. Phys. Chem. Solids 65 1285
- [16] Finger L W and Hazen R M 2001 *Rev. Mineral. Geochem.* **41** 123
- [17] Hazen R M, Downs R T and Finger L W 1996 Science 272 1769
- [18] Dubrovinskaia N and Dubrovinsky L S 2001 Mater. Chem. Phys. 68 77
- [19] Leger J M, Haines J, Schmidt M, Petitet J P, Pereira A S and da Jornada J A H 1996 Nature 383 401
- [20] Schnick W 1993 Angew. Chem. 105 846
 Schnick W 1993 Angew. Chem. Int. Edn Engl. 32 806
- [21] Römer S R, Braun C, Oeckler O, Schmidt P, Kroll P and Schnick W 2008 Chem. Eur. J. 14 7892
- [22] Römer S R 2009 *Dissertation* Ludwig-Maximilians-University Munich
- [23] Lences Z, Hirao K, Sajgalik P and Hoffmann M J 2006 Key Eng. Mater. 317/318 857
- [24] Peng L, Xu L, Zhicheng J, Zhang J, Yang J and Qian Y 2008 Commun. Am. Ceram. Soc. 91 333
- [25] Peng G, Jiang G, Li W, Zhang B and Chen L 2006 J. Am. Ceram. Soc. 89 3824
- [26] Peng G, Jiang G, Zhuang H, Li W and Xu S 2005 Mater. Res. Bull. 40 2139
- [27] Groen W A, Kraan M J and de With G 1994 J. Mater. Sci. 29 3161
- [28] Lee S S, Lim S, Sun S S and Wager J F 1997 Proc. SPIE—Int. Soc. Opt. Eng. 3241 75
- [29] Le Toquin R and Cheetham A K 2006 Chem. Phys. Lett. 423 352
- [30] Hohenberg P and Kohn W 1964 Phys. Rev. B 136 864
- [31] Kresse G and Hafner J 1993 *Phys. Rev.* B **47** 558 Kresse G and Hafner J 1994 *Phys. Rev.* B **49** 14251
- [32] Kresse G and Furthmüller J 1996 Comput. Mater. Sci. 6 15
- [33] Kresse G and Furthmüller J 1996 *Phys. Rev.* B **54** 11169
- [34] Perdew J P 1991 Electronic Structures of Solids '91 ed P Ziesche and H Eschrig (Berlin: Akademie Verlag)
- [35] Kresse G and Joubert J 1999 *Phys. Rev.* B **59** 1758
- [36] Monkhorst H J and Pack J D 1976 *Phys. Rev.* B **13** 5188
- [37] Murnaghan F D 1944 Proc. Natl Acad. Sci. USA **30** 244
- [38] Birch F 1952 J. Geophys. Res. 57 227
- [39] Vinet P, Ferrante J, Smith J R and Rose J H 1987 Phys. Rev. B 35 1945
 - Vinet P, Rose J H, Ferrante J and Smith J R 1989 J. Phys.: Condens. Matter 1 1941
- [40] Thompson J G, Withers R L, Palethorpe S R and Melnitchenko A 1998 J. Solid State Chem. 141 29
- [41] Pauling L and Brockway L O 1932 Z. Kristallogr. Kristallgeom. Kristallphys. Kristallchem. 82 188

- [42] Cox D E, Takei W J and Shirane G 1962 Am. Crystallogr. Assoc. 1962P 6
- [43] Shaposhnikov V L, Krivosheeva A V, D'Avitaya F A, Lazzari J-L and Borisenko V E 2008 Phys. Status Solidi b 245 142
- [44] Petukhov A G, Lambrecht W R L and Segall B 1994 Phys. Rev. B 49 4549
- [45] Ahuja R and Dubrovinsky L S 2002 J. Phys.: Condens. Matter 14 10995
- [46] Kruger M B, Nguyan J H, Li Y M, Caldwell W A, Manghnani M H and Jeanloz R 1997 Phys. Rev. B 55 3456
- [47] Douakha N, Holzapfel M, Chappel E, Chouteau G, Croguennec L, Ott A and Ouladdiaf B 2002 J. Solid State Chem. 163 406
- [48] Fang C M, Hintzen H T and de With G 2004 Appl. Phys. A 78 717
- [49] Huang J Y, Tang L-C and Lee M H 2001 J. Phys.: Condens. Matter 13 10417
- [50] Groen W A, Kaan M J and de With G 1993 J. Eur. Ceram. Soc. 12 413
- [51] Bruls R J 2000 PhD Thesis University of Eindhoven
- [52] Wyckoff R W G 1920 J. Am. Chem. Soc. 42 1100
- [53] Maunaye M, Guyader J, Laurent Y and Lang J 1971 Bull. Soc. Fr. Minéral. Cristallogr. 94 347
- [54] Manoun B, Saxena S K, Liermann H P, Gulve R P, Hoffman E, Asrsoum M W, Hug G and Zha C S 2004 Appl. Phys. Lett. 85 1514
- [55] Solozhenko V L and Kurakevych O O 2005 Solid State Commun. 133 385
- [56] Martin C D, Chapman K W, Chupas P J, Prakapenka V, Lee P L, Shastri S D and Parise J B 2007 Am. Mineral. 92 1048
- [57] Römer S R, Schnick W and Kroll P 2009 J. Phys. Chem. C 113 2943
- [58] Römer S R, Kroll P and Schnick W 2009 Phys. Status Solidi b at press (doi:10.1002/pssb.200945011)
- [59] O'Keeffe M and Hyde B G 1976 Acta Crystallogr. B 32 2923
- [60] Shannon R D 1976 Acta Crystallogr. A 32 751
- [61] Baur W H 1987 Crystallogr. Rev. 1 59
- [62] Desgreniers S 1998 *Phys. Rev.* B 58 14102
 Sharma S M and Gupta Y M 1998 *Phys. Rev.* B 58 5964
 Perlin P, Jauberthie-Carillon C, Itie J P, Miguel A S, Grzegory I and Polian A 1992 *Phys. Rev.* B 45 83
 - Xia H, Xia Q and Ruoff A L 1993 *Phys. Rev.* B **47** 12925 Ueno M, Yoshida M, Onodera A, Shimomura O and
 - Takemura K 1994 *Phys. Rev.* B **49** 14
 - Ueno M, Onodera A, Shimomura O and Takemura K 1992 *Phys. Rev.* B **45** 10123
 - Perlin P, Jauberthie-Carillon C, Itie J P, San Miguel A, Grzegory I and Polian A 1991 *High Pressure Res.* **7** 96
 - Perlin P, Gorczyca I, Porowski S, Suski T, Christensen N E and Polian A 1993 *Japan. J. Appl. Phys. Suppl.* **32** 135
 - Uehara S, Masamoto T, Onodera A, Ueno M, Shimomura O and Takemura K 1997 J. Phys. Chem. Solids 58 2093