Synthesis, Structure, and Properties of Anti-perovskite Nitrides Ca_3MN , M = P, As, Sb, Bi, Ge, Sn, and Pb

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A family of anti-perovskite nitrides of the formula Ca_3MN , where M is a Group IV or a Group V element, is reported. Ca_3BiN , synthesized by mixing and pressing powders of Ca_3N_2 and Bi into a pellet and subsequently heating the pellet at 1000°C in flowing, dry N_2 gas, is prototypical of this family of compounds. The bismuth in Ca_3BiN has an uncommon oxidation state of 3- as suggested by the properties (semiconducting and diamagnetic) of the compound. The anionic bismuth can be substituted by other trivalent anions Sb^{3-} , As^{3-} , and P^{3-} as expected. The structures of Ca_3AsN and Ca_3PN are distorted from the simple cubic anti-perovskite cell at room temperature because of the small size of As^{3-} and P^{3-} . More interestingly, Ca_3N_2 also reacts with Ge, Sn, and Pb at 1300°C to form similar anti-perovskites, resulting in metallic phases and the unusual open shell oxidation state of 3- for the Group IV elements. @ 1992 Academic Press, Inc.

Introduction

The perovskite structure with the formula ABX_3 is adopted by many oxide compounds. where A and B are cations and X is oxygen. The structure, which emphasizes the twelvefold coordination of the A atom, can be described by a cubic unit cell which contains an A atom at the center of a cube, B atoms at the corners, and X atoms at the center of the cell edges. Another common way to describe the structure is to shift the origin so that a B atom is at the center of the cube, A atoms at the corners, and X atoms on the face centers. The latter emphasizes the octahedral coordination of the B atom by six X atoms. Anti-perovskite structures, in which X and B interchange their roles, are also known (1); i.e., the X anion is at the cationic B site instead and becomes the center of an octahedron formed by six metal atoms.

Among nitride anti-perovskites two classes can be further distinguished. In the first class, nitrogen can be thought of as an interstitial atom which goes into the octahedral hole of an already existing metal framework with a slightly modified cell constant (2-5). Therefore, extensive metal-metal bonding continues to exist in the nitride. Examples of this class are Mn_4N (3.857 Å), Fe_4N (3.795 Å), and Fe_3PtN (3.857 Å), whose original compounds are fcc manganese (3.862 Å), fcc iron (3.666 Å), and Fe₃Pt (Cu₃Au structure) (3.727 Å), respectively (6). The cell constant is listed in the parenthesis following each compound for comparison. Note that the inclusion of interstitial nitrogen causes only a small change in the lattice parameter.

The new anti-perovskite nitrides reported in this paper, i.e., Ca_3MN , where M is a

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Group IV or Group V element, belong to the second class. In this class, the A and B sites are occupied by anions M^{3-} and N^{3-} , respectively, and the X sites are occupied by cations Ca^{2+} , thus the cation and anion positions are completely reversed from a normal oxide perovskite. Note that the Ca₃M intermetallics do not exist for the Group IV or V element, except for Ca₃Pb (7), in contrast to the first class of interstitial nitrides.

Oxide perovskites, though simple in structure, exhibit many interesting properties such as structural phase transitions, ferroelectricity, superconductivity (8), etc. Furthermore there are many oxide compounds which are often considered as derivatives of the perovskite structure, such as the K_2NiF_4 structure. The compounds in this new family of nitrides also exhibit interesting physical phenomena. The synthesis, characterization, and properties of these new phases are described in the following sections.

Sample Preparation

Since some of the reactants and products are air-sensitive, all sample manipulations were carried out in an Ar-filled glove box. Calcium nitride (Ca_3N_2) was first prepared by heating granular calcium ($\sim 0.3 \times 0.3 \times$ 0.3 cm^3) in an N₂ atmosphere at 900°C. The product was then ground into a fine powder and mixed with the third element (in powdered form) stoichiometrically. Pressed pellets of the mixtures were heated in N_2 at 1000°C for 2 days to make the P, As, Sb, Bi compounds. Special care was taken to synthesize the phosphorus compound by sealing the pellet in a quartz tube and preheating the pellet at 200°C for 5 hours to avoid producing white phosphorus.

For Ge, Sn, and Pb, however, a higher temperature, 1300°C, was necessary to make the cubic phase. To prevent losing materials at such a high temperature, the samples were sealed in niobium tubes $(\frac{3}{8}'')$

diameter, 0.022" thick wall, 3" long) by arc welding in an Ar atmosphere before the reaction. Since one nitrogen atom is released from every Ca₃N₂ unit when the compound is formed, the amount of sample that can be sealed in the tube is limited to approximately 0.001 mole so that the maximum nitrogen pressure, ~ 20 atm, did not blow up the tube. Noncubic germanium and tin phases were formed when the reaction temperature was reduced to 1000°C. Though a cubic Ca₃PbN was found at 1000°C, the lattice constant is smaller than that of the one made at high temperatures, and the X-ray diffraction peaks are rather broad, indicating some kind of disorder. Attempts to make the anti-perovskite analogues with carbon and silicon have not yet been successful.

The colors of the samples are yellow, red, gray, and black for the phosphorus, arsenic, antimony, and bismuth compounds, respectively. All the germanium, tin, and lead compounds have a dark metallic luster and become black powders after grinding. All samples are air-sensitive and readily decompose in moist air to release NH₃.

Structure Determination

X-ray powder diffraction data were collected on a Scintag XDS 2000 automated diffractometer using $CuK\alpha$ radiation. The $K\alpha_2$ lines were stripped using Scintag software. The powder samples were covered with a thin layer of Mylar, 0.5 mil. in thickness, to avoid air exposure when the data were taken. The patterns from the cubic phases of Ge, Sn, Pb, Sb, and Bi were indexed and solved by trial and error. The lattice parameters were refined by a leastsquare fit and are shown in Table I. Theoretical intensities based on the cubic anti-perovskite structure were calculated and matched well with the experimental integrated intensities (Tables II and III). No superlattice lines or splitting of peaks due to possible distortions from the perfect cubic

TABI	LΕ	I	

LATTICE CONSTANTS OF THE Ca₃MN SERIES

M = Bi, Sb, As, P		M = Pb, Sn, Ge		
Ca ₃ BiN Ca ₃ SbN Ca ₃ AsN Ca ₃ PN	4.8884(5) Å 4.8541(4) Å 4.77 Å ^a 4.73 Å ^b	Ca₃PbN Ca₃SnN Ca₃GeN	4.9550(7) Å 4.9460(6) Å 4.7573(5) Å	

^a Distorted, orthorhombic cell: a = 6.7249, b = 6.7196, c = 9.5336 Å; pseudo-cubic cell: a' = 4.77 Å $\sim a/\sqrt{2} \sim b/\sqrt{2} \sim c/2$.

^b Distorted, orthorhombic cell: a = 6.7091, b = 9.4518, c = 6.6581 Å; pseudo-cubic cell: a' = 4.73 Å ~ $a/\sqrt{2} \sim b/2 \sim c/\sqrt{2}$.

cell were detected within the limit of the machine (intensity limit: 0.3% of the strongest peak; typical peak width: 0.09° FWHM at 30° in 2θ).

The arsenic and phosphorus compounds, however, show clear splitting of the cubic diffraction lines and extra "superlattice" peaks (see Figs. 1 and 2). The true symmetry for Ca₃AsN at room temperature is orthorhombic, which was not known until the distorted structure was completely solved, because the "a" and "b" (Table I) are so close that we could not resolve them in our diffractometer. (The Miller indices used in Fig. 1 are therefore based on a pseudotetragonal cell.) The distorted structure of Ca₃AsN involves tilts of the nitrogencentered octahedra along three axes. It has been solved by Rietveld refinement of neutron and X-ray powder diffraction data, and will be reported separately (9).

Magnetic Susceptibility Measurement

The magnetic susceptibility of the Bi, Sb, As, and P compounds was measured between 320 and 4 K by the use of a Faraday balance (10). The susceptibility at room temperature was measured at different fields to subtract small ferromagnetic impurity signals from the data using the method of Owen and Honda (11). The extent of such ferromagnetic contaminations is very small, on the order of 10^{-6} g Fe, and is likely picked up as small particles when handling the sample in the glove box. Except for a small amount of paramagnetic impurities showing Curie-like behavior at low temperatures (<150 K), the susceptibilities of the samples are diamagnetic and temperature-independent below 320 K. The intrinsic susceptibility of the samples was obtained by subtracting the Curie susceptibility and is tabulated in Table IV. The theoretical value (12) is also presented in the table for comparison by assuming a simple ionic picture and the "additive rule."

The experimental values are smaller than the calculated ones due to two reasons: First, the ionic picture is not totally correct and covalency between the species produces positive susceptibility, i.e., Van Vleck paramagnetism. Alternatively, the covalency decreases the effective radius of the anions, especially rapidly for N^{3-} , and thus also reduces the core diamagnetism, since it is proportional to the square of the ion radius. Second, the phosphorous and arsenic compounds are distorted at room temperature, resulting in a stronger bonding between the P or As anion and the Ca cation, hence the susceptibility increases due again to the Van Vleck paramagnetism.

The susceptibility of the As compound was measured at high temperatures, up to 1200 K, by using a furnace to heat the sample at 100 K/hr (Fig. 3). It is likely that the decreasing susceptibility with increasing temperature indicates that the magnitude of the distortion is decreasing, and when the susceptibility is no longer temperaturedependent the compound is cubic. Diffraction data reported elsewhere shows that the distortion does in fact increase when the temperature is decreased from 300 to 10 K (9). From the data of Fig. 3 the transition to cubic symmetry would occur near 1025 K. The high temperature susceptibility is closer

	(Ge	Sn		Pb	
hkl	$I/I_{0,obs}$	I/I _{0,caic}	<i>I</i> / <i>I</i> _{0,obs}	I/I _{0,calc}	I/I _{0,obs}	$I/I_{0,\text{calc}}$
100	<1	1		18	51	50
110	14	13	40	37	73	69
111	100	100	100	100	100	100
200	67	65	61	62	62	60
210	1	1	14	11	30	29
211	4	5	20	16	35	32
220	44	42	45	42	40	42
300	2	<1]	7	1)	18	3]
221		1		5		13
310	2	2	8	7	18	15
311	38	39	44	41	44	43
222	17	14	12	14	15	14
320	<1	<1	5	3	7	8
321	3	3	8	8	16	17
400	7	6	7	6	6	7
410	<1	<1]	5	2]	11	5]
322		<1		2		5
330	1	<1	4	1	8	3)
411		1		3		6
331	16	16	16	17	17	19
420	18	19	19	19	17	20
421		—	3	2	9	8
332			_	—	5	5

TABLE II

Observed and Calculated Intensities of Ca_3MN , M = Ge, Sn, Pb

Note. Brackets indicate that the peaks overlap.

to the calculated value (see Table IV) presumably due to a decrease of the Van Vleck term. A similar transition occurs in the phosphorus compound at a slightly higher temperature, 1070 K (Fig. 4 and Table IV).

Electrical Conductivity Measurements

Four-probe conductivity measurements were performed at 13 Hz by lock in detection on pellets of Ca₃BiN and Ca₃PbN sintered at 1000°C. The sample dimensions were $\frac{1}{2}$ " in diameter and 0.050" in thickness. Four spring-loaded pins arranged in a colinear configuration were pressed against the pellet to serve as the contacts, which were shown to be ohmic from the linearity of the I–V characteristic up to the maximal current applied, \sim 70 mA. To prevent air exposure, the moisture-sensitive sample was mounted on a supporting stage and covered in the glove box by a Lucite cap, which also supports the four spring-loaded pins.

We calibrated the system by measuring the resistance of a solid molybdenum sample of the same thickness and diameter as the nitride samples, and a proportionality constant between the resistivity and the resistance of the Mo was obtained (13). This value was then used to convert the resistance to the resistivity for the sample. This conversion is valid if we measure the isotropic conductivity of a randomly oriented polycrystalline sample or a material of cubic symmetry.

The temperature dependence of the re-

TABLE III Observed and Calculated Intensities of $Ca_3MN, M = Sb, Bi$

	S	БЪ	Bi		
hkl	$I/I_{0,obs}$	$I/I_{0, calc}$	I/I _{0,obs}	I/I _{0,calc}	
100	16	19	54	51	
110	42	38	76	70	
111	100	100	100	100	
200	64	62	59	60	
210	12	11	31	30	
211	17	16	33	32	
220	41	42	39	41	
300	6	1]	17	3]	
221		5		13	
310	8	7	14	15	
311	38	41	40	43	
222	13	14	13	14	
322	3	3	8	8	
321	8	8	16	18	
400	6	6	6	7	
410	4	2]	10	5]	
322		2		5	
330	4	1)	8	3)	
411		3		6	
331	12	17	15	19	
420	15	19	16	20	
421	3	3	7	8	

Note. Brackets indicate that the peaks overlap.

sistivity of Ca_3BiN from 4 to 320 K is shown in Fig. 5. The magnitude of the resistivity indicates that Ca_3BiN is a semiconductor. However, the small positive temperature dependence suggests that there are conducting states due to overlap of the wave functions of the electrons, or holes induced by the impurities or defects in the sample; that is, this material is an "impurity band" conductor. The turn-around at 20 K probably comes from the "freeze-out" of some of those charge carriers.

The resistivity of Ca₃PbN is much smaller than that of Ca₃BiN, as expected. The room temperature value of $5 \times 10^{-4} \Omega$ cm is large for a nontransition metal compound and is near the minimum metallic conductivity expected for one carrier per unit cell. The small resistance ratio ($\rho(4.2 \text{ K}) = 1.7 \times 10^{-4} \Omega \text{ cm}$) indicates that the sample has a high impurity/defect level or perhaps that the intergrain resistivity of the polycrystal-line pellet is large.

The resistance of a small pellet of Ca₃SbN at room temperatures was measured by an ohmmeter; the value, 10 kohm, indicates that it is a semiconductor. The ohmmeter registered "infinity" for the insulating As and P compounds. At room temperature the two-point resistance of the Pb, Sn, and Ge nitride samples was lower than the ohmmeter resolution of 0.01 Ω .

Discussion

The formal oxidation state of the Group V atoms M in Ca₃MN is 3-. This anionic behavior of the Group V elements is seldom seen in oxides, but is common in intermetallic compounds of M with electropositive elements. For example, M also has 3 - oxidation state in $M'_{3}M$ (14), where M' is an alkaline metal and M is P, As, Sb, or Bi. The assignment of the 3 - oxidation state implies the following scheme for the band structure: The calcium 4s band is empty, but the bismuth 6p bands and nitrogen 2p bands should be full. Therefore, the solid has a filled valance band derived mainly from the Bi-6p orbital and empty conduction bands above the Fermi level derived mainly from the Ca orbitals. The calculated band structure shows that Ca₃BiN has a small band gap of 0.2 eV (15). This is consistent with the observation that Ca₃BiN is quite susceptible to impurities to form impurity "bands."

When the bismuth is substituted for by other trivalent anions, Sb^{3-} , As^{3-} , and P^{3-} the band gap is expected to increase in sequence, because the electronegativity of the anions increases, corresponding to a decrease of the energy of the *p* levels of the anions relative to the calcium levels. The color of the compounds (black, gray, red, and yellow for the Bi, Sb, As, and P,



TWO THETA (DEGREES)

FIG. 1. X-ray powder diffraction pattern of Ca₃AsN. The pattern is indexed on a tetragonal cell, but the true symmetry is orthorhombic with a and b almost the same (see Table I). Asterisk signs, *, are used to indicate impurities or second phases. The air-sensitive sample was covered by a thin layer of mylar for protection, but the mylar formed a semicircular arc over the sample so that there were no diffraction peaks from the mylar.

respectively) supports the above picture. More interestingly, Bi, Sb, and As can also be replaced by their left neighboring elements in the periodic table, namely, Pb, Sn, and Ge. Therefore, these Group IV metals also have an oxidation state of 3-, again an unknown state in oxide chemistry. These new phases should be metallic because the Group IV element has one electron less than the Group V neighbors so



FIG. 2. X-ray powder diffraction pattern of Ca₃PN. The pattern is indexed on an orthorhombic cell; the broad peak at 25° is from a flat mylar covering over the air-sensitive sample. Impurities or second phases are indicated by *.

that their p shell is not completely filled, resulting in a partially filled valence band if we apply the same band structure scheme mentioned above, i.e., we assume a rigid band approximation. The calculated band structure also shows that the Fermi level of Ca₃PbN lies below the top of the valence band (15). A search for the "electrondoped" analogue of the anti-perovskite by substituting Bi with Group VI elements such as Te has not yet been successful. However, we have been able to make a pseudo-quarternary compound Ca_3Bi_{1-x} Pb_xN (x = 0.5) with a lattice constant

TABLE IV MAGNETIC SUSCEPTIBILITIES OF THE Ca₃MN, M = Bi, Sb, As, and P

Compound	$\chi_{\rm M,exp}(10^{-6} {\rm ~emu/mole})$	$\chi_{M,calc}(10^{-6} \text{ emu/mole})$	
Ca ₃ BiN	- 153.5		
Ca ₃ SbN	- 120.4	- 191.9	
Ca ₃ AsN (300 K)	- 82.3		
Ca ₃ AsN (1150 K)	- 112.5	-156.9	
Ca ₃ PN (300 K)	-74.6		
Ca ₃ PN (1150 K)	-99.0	- 126.9	

intermediate between those of the Bi and the Pb phases (a = 4.925 Å).

Since the radii of As^{3-} and P^{3-} are smaller than that of Bi^{3-} , we should expect that cubic Ca₃AsN and Ca₃PN are not structurally stable from the geometrical argument that As^{3-} and P^{3-} are too small to completely fill the hole formed by their 12 calcium neighbors in O_h symmetry. The room temperature structure of these two members is in fact distorted from cubic as expected. Although we did not perform a high temperature X-ray study, the phase transitions detected at 1025 and 1070 K in the high temperature magnetic susceptibility for the arsenic and the phosphorus phases are likely the structural phase transitions corresponding to the change of the crystal structure from the distorted phase to the cubic anti-perovskite. At high temperatures, the effective radii of As^{3-} and P^{3-} are larger due to thermal vibrations, hence the cubic structure is expected to be the stable one. The phosphorus compound undergoes a similar transition at a higher temperature, as may also be expected from the smaller ionic radius of P^{3-} . Such transitions to the cubic phase at high temperatures in distorted oxide perovskites are rather common (8).

The diamagnetic susceptibility of the Bi, Sb, As, and P compounds, respectively, decreases from Bi to P because the radius of



FIG. 3. Magnetic susceptibility of Ca₃AsN at high temperatures. Two curves are shown: one was taken on heating (\odot) , the other one is a cooling curve (+); both were taken at a rate of 100 K/hr. A phase transition at 1025 K with a small hysteresis is implied for Ca₃AsN.



FIG. 4. High temperature magnetic susceptibility of Ca₃PN. A phase transition at 1070 K is also implied for Ca₃PN.



FIG. 5. Temperature dependence of the electrical resistivity of a polycrystalline sample of Ca₃BiN.

the trivalent anion decreases from Bi to P and the Larmor diamagnetism is proportional to $\langle r \rangle^2$, where $\langle r \rangle$ is the mean radius of the ionic species. One can roughly estimate the radius of Bi³⁻ from the accepted ionic radius of Ca^{2+} (1.0 Å (16)) and the lattice constant by assuming that the spherical Bi³⁻ just touches its 12 calcium neighbors. The value thus obtained is 2.4 Å, which is much larger than either that of atomic Bi, 1.5 Å, or that of Bi^{3+} (1.0 Å (16)), but is compatible with those of Bi^{3-} anions in the intermetallics of bismuth and alkali metals, i.e. 2.2, 2.2, 2.3, and 2.3 Å for the hexagonal BiLi₁, $BiNa_3$, BiK_3 , and $BiRb_3$, respectively, (again assuming known ionic radii, 0.76, 1.02, 1.38, and 1.52 Å for Li⁺, Na⁺, K⁺, and Rb⁺, respectively (16)), indicating that the Bi in this nitride compound is rather ionic and hence produces a large diamagnetism.

The average Ca–N bond distance, 2.45 Å, of those anti-perovskites is close to those in Ca_3N_2 (17), 2.47 Å, CaNiN (18), 2.51 Å, and Ca_2ZnN_2 (19), 2.48 Å. In the anti-perovskite compounds discussed in this paper, the nitrogen is in an octahedral hole formed by six calcium atoms. A distorted octahedral coordination of nitrogen by six metal atoms is frequently observed in the known ternary nitrides containing calcium. Examples are CaNiN (18), where the N is coordinated to two Ni and four Ca atoms; Ca_2ZnN_2 , where the N is surrounded by five Ca atoms and one zinc atom (19); CaGaN, where the N has six nearest neighbors, one Ga and five Ca, (20); and Ca₄In₂N, where the N is in an octahedral coordination by six calcium atoms (21). Even in Ca_3N_2 itself the N is coordinated by six calcium atoms in a distorted octahedron (17). This empirical observation is useful in building structural models to solve the structure of other ternary calcium nitrides. The only exception to the rule is the previously known Zintl phase CaGeN₂, where $(GeN_2)^{2-}$ adopts a β cristobalite structure, with the calcium ions "stuffed" into tetrahedral sites; each N is coordinated by two Ge with a very short Ge-N distance of 1.85 Å and by one Ca (22-24).

In summary, we have investigated a series of anti-perovskite nitrides that form with Ca, N, and a Group IV or V element. By varying the central anions, the properties change from metallic (Ca₃PbN, Ca₃SnN, Ca₃GeN) to semiconducting with small band gaps (Ca₃BiN, Ca₃SbN), to insulating with structural phase transitions in Ca₃AsN and Ca₃PN. Ca₃AsN appears to be the first example of a distorted anti-perovskite whose structure is known, since there are few antiperovskites in literature, and this work is reported in a separate paper (9).

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