HIGH PRESSURE VAPOR TRANSPORT OF ZnGeP₂: I. PARAMETER EVALUATION

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Abstract

 $ZnGeP_2$ crystals and epilayers can be grown by chemical vapor transport using phosphorus or hydrogen chloride as transporting agent. The limiting factor in the transport rate is the volatility of germanium which has a seven order of magnitude smaller partial pressure than elemental zinc. Excluding oxygen using carbon films inside the growth vessels and employing a nitrogen shroud in the furnace tube also transport of $ZnGeP_2$ will be obtained when phosphorus is used as transport agent. Optical absorption measurements above show the presence of absorption band head systems which cannot be explained by the absorption lines of P_2 , P_4 , PO or GeO. These new bands are also present in the vapor above $ZnGeP_2$ but not above ZnP_2 . The experimental findings have been simulated by thermochemical calculations taking into account estimated values for the vapor species GeP.

I. Introduction

ZnGeP₂ crystallizes in the chalcopyrite structure which can be inferred from the zincblende lattice by doubling the unit cell of ZnS and ordering the cation sublattice. In the tetragonal unit cell a small distortion occurs in the direction of the c-axis which substantially affects the optical properties of this compound semiconductor. The material is of interest in context of its non-linear optical properpties and can be applied as optical parametric oscillator or for frequency mixing. For this reason, thick films or crystals of high transparency are in demand, which can be grown from the vapor phase.

The dissociation pressure of the ternary chalcopyrite $ZnGeP_2$ amounts to 3.5 bars at the melting point at 1295 K [2]. In the last contribution to this conference series [1] we directed attention to the sensitive change of this equilibrium pressure with composition of the condensed phase. Therefore, close control of the partial pressures of the constituting elements and their gaseous compounds during vapor growth is necessary.

In this paper we discuss vapor phase transport experiments of ZnGeP₂ in oxygen free environment using phosphorus or hydrogen chloride as transport agent. It is assumed that germanium as well as zinc could form volatile phosphorus species. To detect new phosphorus bearing gas species, optical absorption spectroscopy of the vapor phases over ZnGeP₂, ZnP₂ and GeP has been performed. The results are compared with thermochemical calculations using the free energy minimization method described by Ericson [3].

II. Transport Experiments

In order to distinguish the conditions of multiphase from those for single phase transport under exclusion of deleterious effects of residual oxygen contamination of the vapor atmosphere experiments

employing selected vapor composition have been performed using carbonized quartz tubes which were first etched in aqua regia, rinsed with distilled water, dried and finally baked under vacuum at 1223 K for 24 hours to minimize the concentration of water and OH-groups in the walls of the tubes. Prior to sealing under vacuum with a quartz plug the ampoules were filled with about 1 g pre-synthesized ZnGeP₂. To fix the phosphorus vapor pressure appropriate amounts of Zn₃P₂ and P were added. The ampoules had a diameter of 10 mm and a length of 150 mm. A temperature differences of $\Delta T = 1308-1243$ K was applied for 168 hours using a resistance heated furnace. After cooling, the ampoules were opened under N₂ atmosphere and the products characterized by x-ray diffractometry, scanning electron microscopy (SEM) and energy dispersive x-ray analysis (EDX). Typical results are given in table I.

Table I Results of transport experiments using P and HCl as transporting agents.

No.	Ingot	Ampoule	Transported Phases
III/2	ZnGeP ₂ (1.1631 g) Zn ₃ P ₂ (3.8 mg) P (47.1 mg)	heated, carborized HCl added (35 kPa)	$\frac{ZnGeP_2}{Zn_3P_2}$, ZnP_2 (5%) Zn_3P_2 (5%) GeP or Ge (1%) transport rate ~ 3 mg/hr
III/3	ZnGeP ₂ (1.12338 g) Zn ₃ P ₂ (2.9 mg) P (50 mg)	heated, carbonized HCl added (35 kPa)	$ \underline{ZnGeP_2} $, ZnP_2 (2%) $ Zn_3P_2 $: (5%) $ GeP \text{ or } Ge (2\%) $ $ transport rate ~ 3 mg/hr$
III/4	ZnGeP ₂ (0.935 g) Zn ₃ P ₂ (2.3 mg) P (47.3 mg)	heated, no graphite layer	ZnP ₂ , ZnGeP ₂ Zn ₃ P ₂ : Ge (GeP or Ge) transport rate ~ 2 mg/hr
III/5	ZnGeP ₂ (1.054 g) Zn ₃ P ₂ (2.6 mg) P (49.6 mg)	heated, carbonized double wall	$\frac{ZnP_2}{Zn_3P_2}$: $\frac{ZngP_2}{Ge}$ (GeP or Ge) transport rate ~2 mg/hr

In all experiments transport of ZnGeP₂ was observed. The main transported phases in experiments without HCl were ZnP₂, Zn₃P₂ and ZnGeP₂ The presence of Ge or GeP can be inferred from the XRD patterns (see fig. 1.a). Heating of the tube under vacuum and/or the presence of a carbon film inside the ampoule wall diminuishes the oxygen content so that no Ge_xO_x species can be formed. Nevertheless, the transport of ZnGeP₂ without halogen proceeds at a high rate. This points to the fact that other volatile Ge species have to be present in the vapor phase. The specific stoichiometry of the Ge_xP_y species that are responsible for the observed transport in the experiments III/4 and III/5 is not known at this time. The x-ray diffractograms of the transported material, shown in figs. 1.a and 1.b, can be indexed as ZnGeP₂ (JCPDS 33-1471), ZnP₂ (JCPDS 23-748), Zn₃P₂ (JCPDS 22-1021, Ge (4-545) and GeP (21-348). SEM images of a polished sample of experiment III/5 reveal that at least three phases are present (see figure 2). A dark phase (ZnP₂), a dark grey phase (ZnGeP₂) and a light grey phase (Zn₃P₂) containing 2 mole% Ge. The element concentrations were measured by microprobe analysis (EDX). To separately study the chemical transport behavior of GeP 10-2 mole Ge and P were placed in a BN-boat sitting in an evacuated and sealed quartz ampoule (I = 250 mm, ϕ = 20 mm) and treated in a temperature

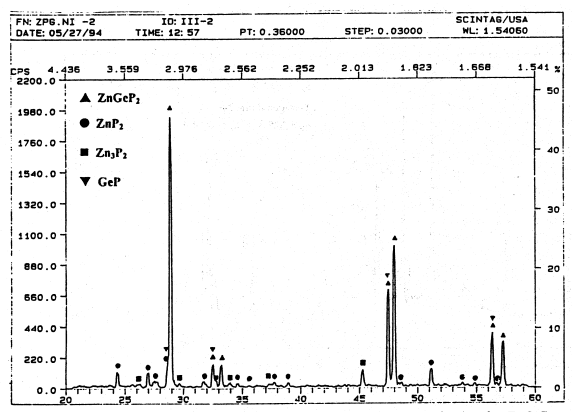


Figure 1.a: x-ray diffractogram of transported material from experiment III/2. Besides ZnGeP₂ small amounts of the phases ZnP₂, Zn₃P₂, Ge or GeP are present. (Cuk α radiation, λ = 1.5406 Å).

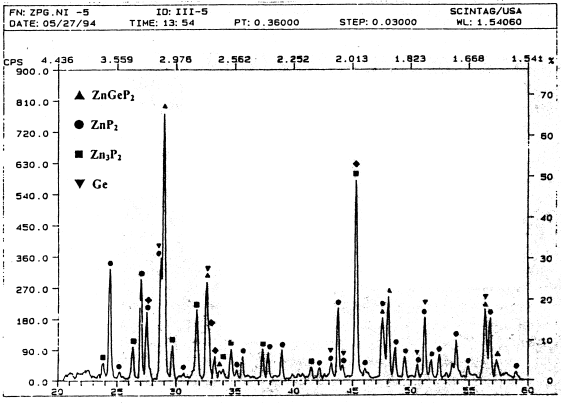


Figure 1.b.: X-ray diffractogram of transported material from experiment III/5. Main phases: Zn $_3P_2$, ZnP2, ZnGeP2 and Ge and GeP possible. (Cuk $_{\alpha}$ radiation, $\lambda = 1.5406$ Å).

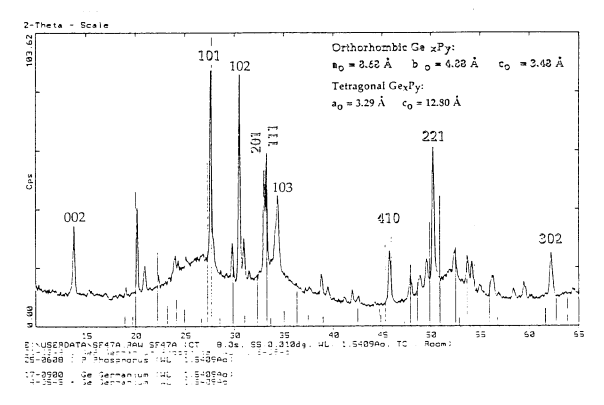


Figure 1.c: X-ray diffractogram of transported material from a chemical vapor transport experiment with GeP using phosphorus as transporting agent. Present phases: GeP, Ge and unknown other germanium phosphides of orthorhombic and tetragonal symmetry. (Cuk $_{\alpha}$ radiation, λ = 1.5409 Å).

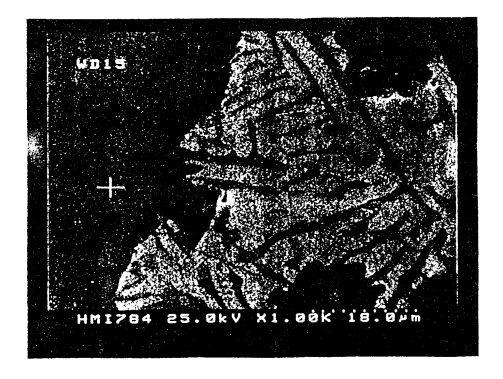


Figure 2: Scanning electron micrograph of a polished sample from transported material in experiment III/4. Black phase: ZnP_{2} , dark grey phase: $ZnGeP_{2}$, light grey phase : $Zn_{2,8}Ge_{0,2}P_{2}$.

difference from $\Delta T = 1173$ - 673 K. Within two days a black layer has been formed consisting of orthorhombic and tetragonal germanium phosphides (fig. 1.c).

Composition and structural analysis of the vapor species in the system Zn-Ge-P can be analysed by optical measurents in the vapor phase. In the following chapter we present results from optical absorption measurement.

III. Optical absorption spectroscopy

Optical emission and absorption spectroscopy is a method to study molecular structures of gas species and to obtain their thermochemical data. It can also be used to quantitatively measure element concentrations in the vapor phase. (e.g. by Atomic Absorption Spectrometry AAS). Although it is difficult to identify new gas species from such spectra the absorption or emission lines can be utilized to distinguish different gas species in the vapor phase. In our experiments the pressures in the analyzed gas volumes range from mbar up to several bars. We compare the absorption spectra of the elements with the absorption behaviour over the condensed binary and ternary compounds. In the wavelength range from 180 to 1000 nm vibronic spectra have been described, in which transitions occur from rotational/vibrational states of one electronic state to rotational/vibrational states of another electronic state [4].

The setup to measure the absorption bands in the vapor phases over ZnGeP2, ZnP2 and GeP is illustrated in figure 3. The equipment consists of a two zone furnace with integrated isothermal furnace liners (Dynatherm) to guarantee a uniform temperature profile for the quartz glass cuvettes of 50, 100 mm length and 10 mm diameter. The temperature was controlled by PID temperature controllers (Eurotherm 808), the temperature was monitored by two NiCr/Ni thermocouples. The cuvettes were mounted on top of a V-shaped rim which was mounted at the ends on holders fixed on a vibrational isolated table. To change the cuvettes the furnace was mounted on a slide which enabled a reproducible positioning of the cuvettes in every run. As light source a xenon lamp (ORIEL 7340, Xe 6253, 150 W) was employed. To guide the light through the heated quartz glass cuvettes (Hellma Inc.) and to focus the transmitted light onto the entrance slit of a monochromator (ISA HR 320) a system of quartz lenses (UV grade) was used. The light intensity was detected at the exit slit of the monochromator by a photomultiplier tube (Hamamatsu 522; operation voltage U = 1200 V using an Ortec HV supply) and amplified with a preamplifier (Keithley 602). To improve the signal to noise ratio the light was chopped with 334 Hz (Chopper: Standford Research SR 540) and phase sensitiv amplified by a lock-in amplifier (EG&G 5206). To overcome the so-called schlieren effect, which also contributes the noise level of the measurement by air turbulences in the heated furnace core, the cuvettes to be measured were placed between two empty quartz glass cuvettes. To investigate the absorption behavior, about 50 mg of red phosphorus, ZnGeP₂, ZnP₂ and GeP, respectively, were filled into the evacuated and sealed cuvettes. The results are shown in the figures 4.a to c. Figure 4.a shows a compilation of the transmission curves over ZnGeP, at 807 K, GeP at 881 K and ZnP2 at 773 K. The different temperatures of the cuvettes take into account the temperature dependence of the vapor pressures and were chosen to yield comparable absorption intensities.

The absorption curves of $ZnGeP_2$, GeP and ZnP_2 show a common minimum at about 245 nm and a transmission maximum in the wavelength range from 240 to 200 nm. At higher wavelengths a continuous increase of transmission is typical for all spectra. This general curve shape can be explained by the transmissivity of the quartz glass used (lenses and cuvettes) and the higher light intensity of the Xe-lamp at higher wavelengths. As reference the transmission curve of a 10 mm thick specimen of GE type 124 fused quartz is shown in figure 4.b. Additional minima in the curves are typical features of the vapor species over the related compounds. Due to the moderate resolution of oused monochromator (0.05 nm) compared to vacuum spectrographs with an resolution of 0.001 nm only band heads could be detected.

a) ZnP₂: two spectra have been measured at 773 and 723 K (figure 4.c). The transmission curve exhibits two strong absorption maxima at 290 and 296.7 nm and shows high absorption at wavelengths smaller than 260 nm for the 773 K spectum, which are not explained by absorption bands of gaseous Zn or Zn₂

[5, 6]. Since homogeneous evaporation of ZnP_2 has been described in the literature [7] volatile zinc phosphides exists in the vapor phase. However, the pronounced absorption maxima observed over ZnP_2 are not present in the vapor phase over $ZnGeP_2$ (not shown in figure 4.a and 4.f). This means that within the detection limit of our experiments no zinc phosphide species are present in the equilibrium above the chalcopyrite.

b) GeP: Comparing the number of absorption maxima of P, GeP and $ZnGeP_2$ in the wavelength range from 190 to 250 nm (figs. 4.d, 4.e and 4.f) GeP exhibits the most complex spectrum. The analysis of the line positions shows that this spectrum consists of absorption lines of P_2 [5, 8, 9] and at least of one additional vapor species (see Table II), whose line positions relate to the most prominent absorption maxima in the range from 180 to 245 nm.

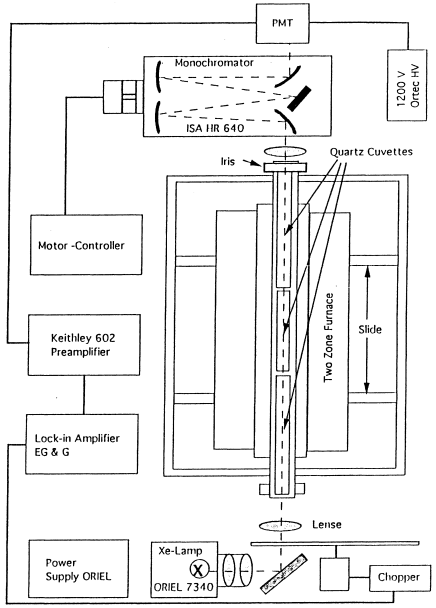


Figure 3: The experimental set-up to measure transmission curves of the equilibrium vapor over $ZnGeP_{2'}$ $ZnP_{2'}$ GeP and P (red).

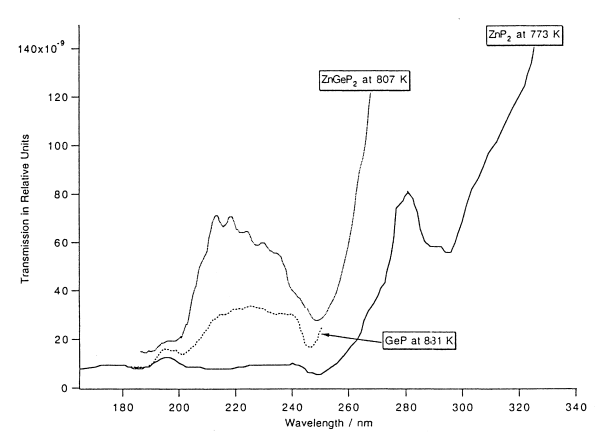


Figure 4.a: Transmission curves of the equilibrium vapor phases over ZnGeP2, ZnP2, GeP and P (red).

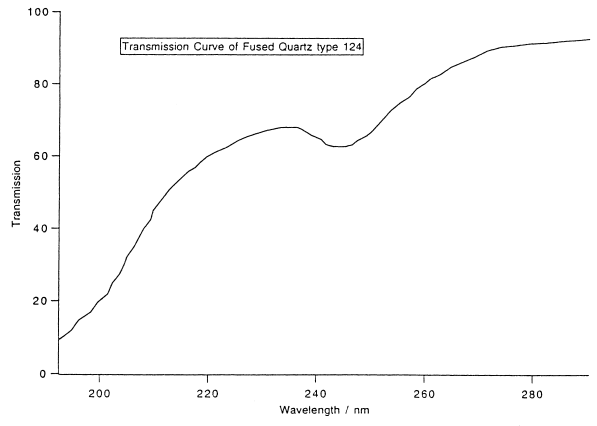
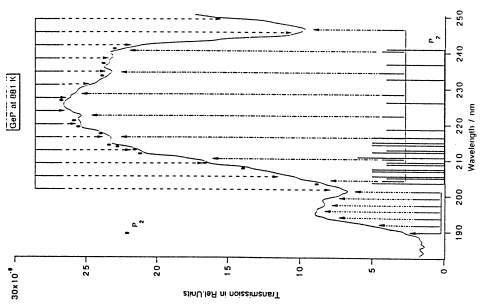


Figure 4.b: Transmission curve of a 10 mm thick fused quartz plate (GE Quartz, type 124)



Figure 4.d: Transmission curves of the equilibrium vapor phase over GeP.



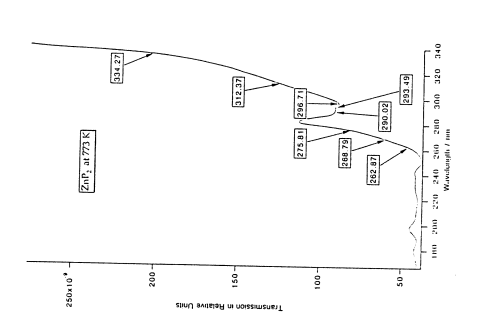


Figure 4.c: Transmission curve of the equilibrium vapor phase over ZnP2.

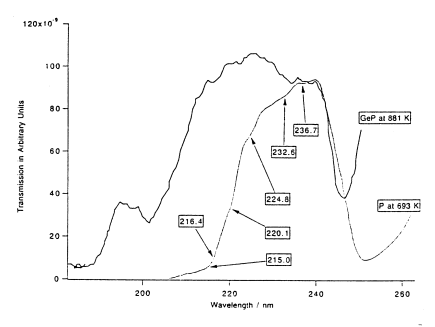


Figure 4.e: Transmission curves of the equilibrium vapor phase over GeP and P (red).

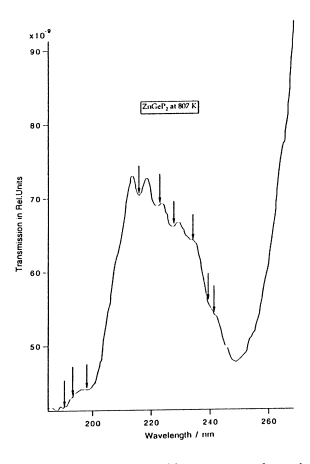


Figure 4.f: Transmission curves of the equilibrium vapor phase above ZnGeP2. The positions of the arrows mark the position of Ge_xP_y band heads.

In order to check the vibrational analysis of a band system the attempt has been made to set out the wave numbers of the absorption maxima in a Deslandres table [4]. Three progressions have been found which are depicted as line grids in figure 4.d together with the line positions of P_2 . The analysis is correct if the wave numbers of corresponding bands in different v' (excited states) and v'' (ground states) progressions, respectively, are constant. Tentatively the grid with narrow distances could be assigned as a v' progression for v'' = 0 while the other two grides belong to higher v' and v'' progressions. For thermochemical calculations the existence of the gaseous species GeP has been assumed.

Table II Positions of the absorption heads in figure 4.d with possible progressions I, I and III (P: P_2 lines).

Absorption lines	Absorption lines
of P ₂ in nm	of Ge _x P _y in nm
-	III: 190.3, 194.1, 195.4,
	197.0, 198.0, 200.0.
	201.5 I
203.6	
205	
205.5	205.35 I
206.9	
207.49	207.58 P
208.83	
209.43	209.8 I, P
210.81	211.07 P
	212.02 I
212.26	
212.86	212.98 P
	213.62
214.3	214.26 P
215.0	
	215.94 II
216.4	216.8 P
	217.78 I, II
218.64	218.74 P
220.14	220.34 I, P
	222.59 II
	223.55 I
226.16 I	226.11 I
	227.08 I
	228.37 II
	231.59 I
232.65	232.55
	234.16 I, II
236.76	236.73 P
	238.98 I
240.99	240.91 P, II
	241.56
	243.2
	244.51 I
	246.77 II

IV. Heat of formation of ZnGeP2

In the last contribution to preceding conference [1] we inferred the heats of formation of ZnP_2 , Zn_3P_2 and $ZnGeP_2$ from equilibrium vapor pressure measurements (Table III). The values for ZnP_2 and $ZnGeP_2$ differ by ~ 2 kcal/mole. This result is in agreement with Differential Thermal Analysis DTA measurements that we employed to study the reaction of ZnP_2 with Ge to form $ZnGeP_2$. Figure 5 shows a typical result. Mainly endothermic contributions to the reaction heat occur which are about 2 kcal/mole smaller than the heat of fusion of $ZnGeP_2$ ($\Delta H^m = 8.8$ kcal/mole).

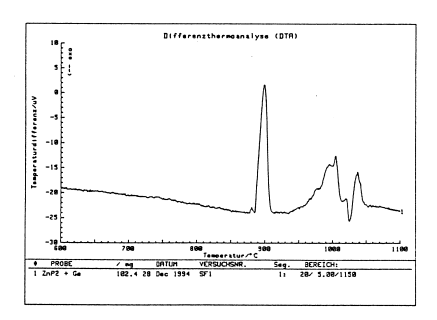


Figure 5. Differential Thermal Analysis DTA curve of reaction ZnP₂ + Ge = ZnGeP₂

V. Phase relations, predominance area diagram and vapor phase composition

Figure 6 depicts the Gibbs phase triangle for the system Zn-Ge-P. The GeP binary shows five condensed phases but only GeP and presumably GeP₂ are stable under normal pressure conditions. GeP₃ and GeP₅ can only be prepared under high pressure conditions (65 kbar) [10]. IV-V compounds of compsition AB₂ (A = Si, Ge; B = P, As) crystallize in two modifications: in a high pressure modification of pyrite structure [10] and an low pressure orthorhombic structure [11]. In the x-T diagram of Ge-P the existence of GeP₂ is not mentioned [10, 11]. In accord with the Gibbs phase rule, triangular subfields are defined in the Zn-Ge-P composition field. The solid phases coexisting in equilibrium for any composition in these subfields are given by the corner compounds and elements.

With the thermochemical data, summarized in Table III, thermochemical calculations using the free energy minimization technique were carried out. One result is the predominance area diagram depicted in figure 7. showing the stability field for ZnGeP₂ as a function of the partial pressures P₄ and Zn. The number of phases found in the XRD pattern from experiment III/3 (figure 1.a), especially the presence of Ge and/or GeP, shows that the system was not in equilibrium. This may be due to the fact that the heats of formation of ZnGeP₂ and ZnP₂ do not differ significantly so that Ge and ZnP₂ can occur as metastable

Table III Thermochemical data used for free energy minimization calculations. The dimensions of the standard heat of formation ΔH_f^0 , heat of fusion ΔH_f^0 heat of evaporation ΔH_g^0 are kcal/mole, for the standard entropies H_g^0 and molar heats H_g^0 0 cal/mole deg.

Element/	$\Delta II_{\mathbf{f}}^{\circ}$	S_{f}^{o}	$c_p = A+B$	·10-3·T+	$C \cdot 10^5 \cdot T^{-2}$
Compound	ΔΗ _{fu} ΔΗ _e		λ	В	С
P(red)	0.00	5.45	4.051	3.559	_
P4 (white)		39.28	18.28	15.12	_
P(1)	17.31	_	25.168	_	T>317 K
P4(g)	30.77	66.89	19.562	0.162	-3.213
P ₂ (g)	40.68	52.11	8.675	0.191	-0.994
Zn(s)	0.00	9.943	5.096	2.782	0.13
Zn (1)	1.75	-	7.495	-	-
Zn (g)	31.15	38.451	4.965	-	-
$ZnCl_2(s)$	-99.13	26.62	14.290	8.994	-
$ZnCl_2(1)$	2.474	-	2.588	-	-
ZnCl ₂ (g)	-63.839	66.185	14.74	-1.03	-
Ge(s)	0.00	7.425	5.577	0.931	-0.25
Ge(1)	8.82		6.596		T>1210 K
Ge (g)	89.34	40.1	7.038	-0.873	0.76
Gell ₄ (g)	21.68	51.89	14.939	5.276	-5.166
GeP(s)	-0.13	14.6	10.84	2.7	-1.25
GeP(g)	28.4	44.43	12.34	-	-
GeO(s)	-50.3	11.9	9.543	3.497	-
GeO (g)	-10.99	53.49	8.92	0.0239	9-1.3685
Ge ₂ O ₂ (g)-	112.	75.1	20.	-	_
Ge ₃ O ₃ (g) -	212.	99.3	32.		-
GeCl(g)	174.36	58.72	10.314	-0.532	-1.18
GeCl ₂ (g)	-40.84	70.63	13.818	0.052	-0.88
ZnP ₂	-24.15	14.4	17.01	3.997	0.65
Zn ₃ P ₂ -	-38.65	36.2	30.15	6.225	-0.36
ZnGeP ₂ -	-26.3	28.22	24.65	3.5	-4.0

phases. The predominance area diagram allows the equilibrium between ZnGeP₂, ZnP₂ and Zn₃P₂ as found in experiment III/5 (fig. 1.b). Calculating the phase equilibria under experimental conditions (Table I) the triple point was found 20 K below the experimental value of 1243 K. Above this temperature ZnGeP₂ and ZnP₂ were stable, the equilibrium pressure at 1240 K amounted to 3.4 bar. In the presence of HCl higher pressures have to be expected. They ranged from 6.8 to 5.8 bar. It was found that mainly ZnGeP₂ is be stable in presence of a small amount of Ge which is in agreement with the experimental findings.

The calculations employing estimated thermochemical data for gaseous species GeP (Table III) show a dominance in the formation of ZnGeP₂. and a small amount of ZnP₂. However, the experiment showed the transport of ZnGeP₂, Zn₃P₂ and ZnP₂ in comparable amounts. This points to the fact that the stability of the assumed vapor species GeP was over estimated. Further measurements such as mass spectrometric analysis are in progress to yield more insight in the vapor phase composition above GeP.

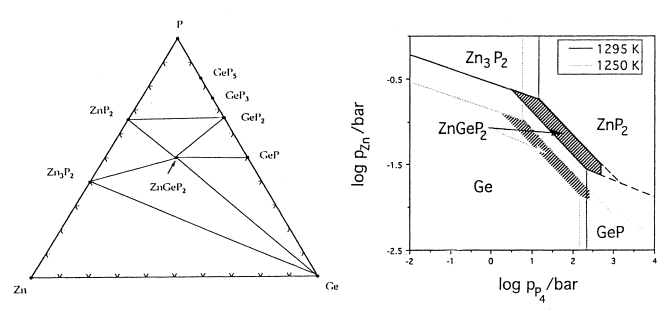


Figure 6. Gibbs phase triangle of the system ZnGeP₂.

Figure 7. Predominance area diagram for the system Zn-Ge-P

VI. Conditions of single phase ZnGeP₂ transport

he vapor composition and transport rate described in table I that lead to multiphase transport, we have performed experiments under slightly changed conditions resulting in single phase ZnGeP₂ transport. This requires working with smaller temperature gradients, typically < 4 K/cm ($\Delta T = 1300\text{-}1290 \text{ K}$) and short transport distances. A typical vapor composition for single phase transport is: [Zn] = 6.27 x 10⁻⁷ mol/cm³), [P] = (3.254 x 10⁻⁵ mol/cm³) and [HCl] = (1.3 x 10⁻⁵ mol/cm³)

VII. Concludig remarks

 $ZnGeP_2$ can be grown by chemical vapor transport using phosphorus as transporting agent. The presence of volatile Ge_xP_y species can be concluded from careful analysis of the absorption spectra measured in the vapor phases above $ZnGeP_2$ and GeP in the wavelength range from 190 to 260 nm. Due to the small

difference in the heats of formation between $ZnGeP_2$ and ZnP_2 a close space vapor transport geometry is recommended to grow crystals or layers of $ZnGeP_2$.

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References

- 1. S. Fiechter, R. H. Castleberry, G. Wood, K. J. Bachmann, H. T. Tran, K. Ito and J. S. Scroogs, "High pressure vapor transport of binary and ternary compound semiconductors," <u>Proceedings of the 6. International Symposium on Experimental Methods for Microgravity Materials Science</u>, R. Schiffmann (ed.), (TMS, Warrendale PA, 1994) 93-100.
- 2. A. S. Borshchevskii, and T. M. Shantsovoi, "Pressure and Composition of the Vapor above Semicoinductive ZnGeP₂," <u>Inorg Chem.</u>, 11 (1975), 1853-1855.
- 3. G. Ericson, and K. Hack "ChemSage A Computer Program for the Calculation of Complex Chemical Equilibria," Metallurgical Transactions B, 21B (1990), 1013-1023.
- 4. G. Herzberg, <u>The Spectra and Structures of Simple Free Radicals</u> (Dover Publications, New York, 1971).
- 5. R. W. B. Pearse, and A. G. Gaydon, <u>The Identification of Molecular Spectra</u> (John Wiley & Sons, New York, 1963).
- 6. R. C. Weast, and M. J. Astle, <u>CRC Handbook of Chemistry and Physics</u> (CRC Press, Boca Raton, Florida, 1981), Line Spectra of the Elements, E-205 to E-334.
- 7. V. B. Lazarev, V. J. Shevchenko, S. F. Marenkin, and G. Magomedgadghiev, "The Growth of Large Tetragonal CdP₂ and ZnP₂ Crystals," <u>J. Crystal Growth</u>, 38 (1977), 275-276.
- 8. A. Jakowlewa, "Fluoreszenz and Absorption des Phosphordampfes," Z. P., 69 (1931), 548-564.
- 9. G. Herzberg, L. Herzberg, and G. Milne, "On the Spectrum of the P₂ Molecule," <u>Canad. J. Res.</u>, A18 (1940) 139-143.
- 11. P. C. Donohue, and H. S. Young, "Synthesis, Structure and Superconductivity of New High Pressure Phases in the System Ge-P and Ge-As," J. Solid State Chem., 1 (1970) 143-149.
- 12. T. Wadsten, "The Crystal Structures of SiP₂, SiAs₂ and GeP," Acta Chem. Scand., 21 (1967) 593-594.
- 13. T. B. Massalski, <u>Binary Alloy Phase Diagrams</u> (American Society for Metals, Metals Park, Ohio, 1990) Vol. 2, p. 1978-1979.