

Factors influencing the purity of electronic grade phosphine delivered to MOCVD tools

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ABSTRACT

Increasing mobility of InP films with usage time of one PH₃ cylinder prompted an investigation into factors influencing the purity of delivered PH₃. The presence of hygroscopic H_xPO_y residues in a delivery system greatly increases the dry-down time compared to that of a clean system. Static delivery system tests show increasing H₂O concentration with time and twice the increase in PH₃ versus N₂ over 48 h indicating reaction of metal oxides in components with PH₃ to generate H₂O. Gas purity may also vary during cylinder usage. Depletion of a high-purity PH₃ cylinder shows consistently low gas phase H₂O levels before phase-break but increasing levels after phase-break, as the cylinder depressurizes. The results highlight the importance of using pure PH₃, employing rigorous cycle-purging procedures to prevent H_xPO_y contamination, switching out cylinders in good time and using purification technology to control H₂O.

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1. Introduction

Minimizing incorporation of impurities such as oxygen can be critical to the performance of III–V compound semiconductor devices [1–3]. Oxygen is readily captured from the environment by aluminum and forms deep recombination centers in Al-containing materials [4]. Although it incorporates less in non-Al-containing materials such as InP, effects are still observed. Munns and coworkers reported a significant influence of hydride gas purity on InP. Increased mobility (77 K) was observed as H₂O was decreased from 2 μmol mol⁻¹ (ppm) to 100 nmol mol⁻¹ (ppb) level in the PH₃ source [5], but the mechanism for this is unclear. In molecular beam epitaxial growth of InP, oxygen can be bound to Be acceptors and to a lesser extent to Si donors [6,7]. Further, in vapor phase epitaxial InP, Iwata and Inoshita [8] have reported that oxygen suppresses Si incorporation. Since Si is the main residual background n-dopant in InP, oxygen may affect Si dopant levels and hence the film properties. Consequently, control of oxygenated impurities such as H₂O in gases such as PH₃ is very important.

PH₃ gas is the most commonly used phosphorus source for InP and associated alloys and is manufactured to a 99.9999% purity level. However, even though the gas purity is confirmed with state-of-the-art instrumentation [9–11], there are many factors that may still influence device performance. Ultimately,

device growth and performance tests such as low temperature photoluminescence (PL) and Hall mobility are used to provide complementary information that relate the atomic composition to critical parameters such as gas purity. In this paper we present PL results of InP films grown with high and lower purity PH₃ and data that show increasing mobility of InP films with run time, for growth with the same PH₃ cylinder. This prompted an investigation of factors that influence the purity of PH₃ delivered to MOCVD tools, particularly with respect to H₂O impurity. Issues such as the influence of H_xPO_y contamination in the delivery system on H₂O levels, generation of H₂O as a result of reaction of PH₃ with metal oxides in stagnant lines, and variable H₂O in the PH₃ delivered from cylinders are considered. Point-of-use purification for H₂O control upstream of the MOCVD tool is also discussed.

2. Experiments

2.1. InP MOCVD, mobility and low temperature PL measurements

9 μm InP films were grown using trimethyl-indium (TMI, Optograde, Dow Chemical, Andover, MA) and two PH₃ cylinders (Matheson Tri-Gas, New Johnsonville, TN). Cylinders 1 and 2 contained PH₃ with 99.9999% and 99.9997% purity, respectively, based on gas chromatography and cavity ring-down spectroscopy (CRDS) analysis. MOCVD was performed in a vertical quartz reactor at 10.13 kPa and 650 °C. The PH₃ flow rate was

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300 sccm, the V/III ratio 300, and the growth rate 11 \AA s^{-1} . The substrates were semi-insulating InP oriented approximately 2° off (1 0 0) towards a (1 1 0) direction. Prior to the sequence of InP growths, the MOCVD tool was routinely cleaned and its performance confirmed with a control PH_3 source. After cylinder installation the connection line was cycle purged before initiating the flow of PH_3 . For each PH_3 cylinder, two InP layers were grown successively; growths were performed with Cylinder 1, then Cylinder 2. Between the first and second growths, the PH_3 cylinder was kept connected in line. Hall mobility of InP layers were measured at 77 and 300 K [12]. The measuring error of the Hall coefficient was estimated to be $\pm 4\%$. PL measurements were made on the second of the two InP growths from each of the two cylinders, at a temperature of 10 K and using a He–Ne laser at 633 nm and an excitation power of 2 W cm^{-2} .

2.2. H_xPO_y contamination

Cylinder 2 was analyzed multiple times using the CRDS method in [9] to investigate the behavior of H_2O in the PH_3 delivered through the cylinder valve (Fig. 1A). The wet-up and dry-down of a clean and contaminated sampling manifold (Fig. 1B), was also compared. PH_3 containing $2.6 \mu\text{mol mol}^{-1} \text{ H}_2\text{O}$ or $<9 \text{ nmol mol}^{-1} \text{ H}_2\text{O}$ was sequentially introduced into the manifold via by-pass or purifier (Nanochem PHX, Matheson Tri-Gas, Longmont, CO) and CRDS was used to measure how rapidly the H_2O level equilibrated.

2.3. SEM study of intentionally generated H_xPO_y contamination

A 316 L stainless steel (SS) sub-manifold was designed to generate H_xPO_y (Fig. 1C). The entire manifold was initially purged with N_2 and then valves V_p and V_m were closed and V_s was opened to let air (atmospheric pressure) fill the second half of the tubing ($90 \text{ cm} \times 6.25 \text{ mm}$ outside diameter). Valve V_s was then closed and V_p was opened to allow PH_3 gas at 40 psig to fill the first half of the tubing (also $90 \text{ cm} \times 6.25 \text{ mm}$ outside diameter). The

cylinder valve and V_p were then closed and V_m was opened to let PH_3 and air mix and react for 10 min in the second half of the tube. The procedure was repeated twice more and after purging, the tubing was removed and sectioned. Samples were stored in a N_2 atmosphere to minimize exposure to atmosphere and were analyzed the same day on a JEOL model JSM-7000F scanning electron microscope (SEM-EDS).

2.4. Water measurements with static manifold/analyzer conditions

To study potential H_2O generation from the reaction of PH_3 with metal oxides, purified N_2 at 200 sccm was used to dry down the SS delivery system and CRDS analyzer in Fig. 1A to single digit nmol mol^{-1} levels. The whole system was then closed and left under static conditions for ~ 2 days and the H_2O concentration monitored. The manifold was then dried down again with purified N_2 and the same experiment repeated with purified PH_3 .

2.5. Phosphine cylinder depletion

The set-up shown in Fig. 1A was used to track the gas phase H_2O level during depletion of a PH_3 cylinder until empty. The full cylinder contained 16.33 kg Ultima 6 N purity PH_3 .

3. Results and discussion

3.1. Low temperature photoluminescence of MOCVD InP

Table 1 shows the impurity profiles of the two PH_3 cylinders. PL measurements were performed to further confirm the gas purities since all impurities except for N_2 and H_2O were below the detection limit of the gas analysis methods. The PL spectrum of InP is strongly temperature dependent [13,14]. We selected 10 K for comparing the exciton spectra of InP films because at this temperature, high-purity n-type InP exhibits significant upper and lower polariton branches of the free exciton. InP grown from PH_3 Cylinder 1 shows a typical exciton spectrum of high-purity

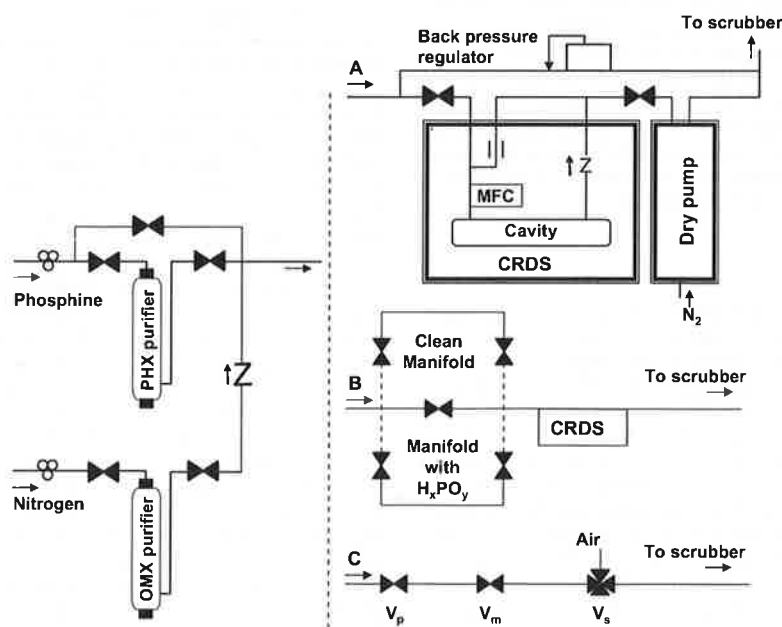


Fig. 1. Diagram of experimental set-up for (A) H_2O analysis and depletion, (B) delivery system contamination and (C) H_xPO_y generation studies.

Table 1
Impurity concentrations in PH₃ Cylinders 1 and 2.

Cylinder no.	Purity (%)	Gas Phase Impurity (nmol mol ⁻¹)										
		CO ₂	CO	O ₂	CH ₄	N ₂	Ar	H ₂ S	AsH ₃	GeH ₄	SiH ₄	H ₂ O
1	> 99.9999	< 16	< 14	< 14	< 7	27	< 9	< 15	< 3	< 0.6	< 4	< 50
2	> 99.9997	< 16	< 14	< 14	< 7	63	< 9	< 15	< 3	< 0.6	< 4	180

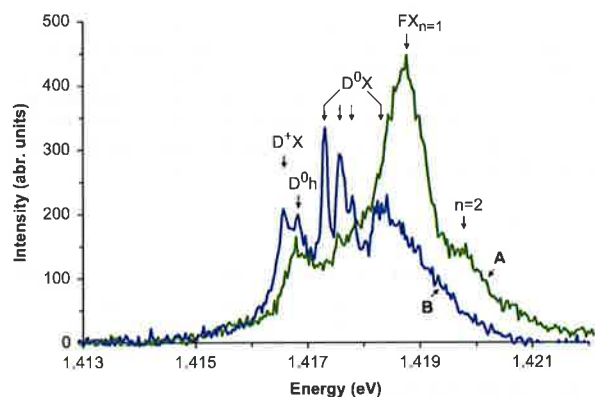


Fig. 2. Low temperature PL of InP grown with PH₃ Cylinder 1 (A) and Cylinder 2 (B); excitation wavelength 633 nm; excitation power 2 mW.

InP (curve A in Fig. 2). The upper ($h\nu_c(\text{FX})=1.41848$ eV) and lower ($h\nu_c(\text{FX}_{n=2})=1.41974$ eV) polariton branches of the free exciton are clearly observable, indicating high-purity n-type InP. At this temperature, the radiative decay of free holes with electrons bound to neutral donors ($h\nu_c(D^0h)=1.41661$ eV) is clearly visible while the radiative decay lines of neutral donor-bound excitons ($h\nu_c(D^0X)_n$ from 1.41692 to 1.41784 eV) are very weak. In the PL spectrum of InP grown from PH₃ Cylinder 2 (curve B in Fig. 2), the first evidence of high-purity InP (the line of lower polariton branch of the free exciton) disappears. Further, the upper polariton branch is not visible because of the higher intensity of $(D^0X)_n$. A new line appears for the radiative decay of an exciton bound to an ionized donor (D^+X), while the four (D^0X) lines are clearly visible compared with the PL profile of high-purity InP at the same low temperature. Clearly, compared to curve B, curve A in Fig. 2 shows a high-purity InP, indicating that Cylinder 1 contains PH₃ of higher purity than Cylinder 2.

3.2. Mobility of InP

In P mobility, especially at low temperature, is closely related to the purity of PH₃ at high V/III ratios. With high-purity PH₃, the average mobility of InP films grown under optimized conditions at 580–600 °C and V/III ratio of 450–800 (77 K) is 171,000 cm²(V s)⁻¹. The highest mobility reported in the literature is 234,000 cm²(V s)⁻¹ using high-purity PH₃ and TMI [15]. For typical n-type InP with a mobility of > 100,000 cm²(V s)⁻¹ the carrier concentration is about 10¹⁴ cm⁻³. Normally, as the mobility increases, the net carrier concentration (N_D-N_A) decreases, indicating that electrons are the dominant carriers. Table 2 shows the mobility of InP films grown with a V/III ratio of 300, using the two PH₃ cylinders. In particular we noted that the mobility increased significantly with the MOCVD run times. The mobility increases were 12% from run 1 to run 2 for Cylinder 1 and 25% from run 1 to run 2 for Cylinder 2. Similar trends, with mobility increases up to 36%, have been

Table 2
Mobility of InP layers grown in duplicate from Cylinders 1 and 2.

Cylinder	Growth	Mobility (cm ² /V s)		Carrier conc. (cm ⁻³)/10E14	
		77 K	300 K	77 K	300 K
Cylinder 1	1st run	137,000	4200	2.12	2.85
	2nd run	153,000	4600	1.53	2.12
	Mobility increase (%)	11.68	9.52		
Cylinder 2	1st run	114,000	4100	3.98	4.74
	2nd run	143,000	4300	2.76	2.93
	Mobility increase (%)	25.44	4.88		

observed with other similar high-purity PH₃ cylinders. As there are many factors that can influence the purity of layers, the exact cause of the increased mobility in the second run from each cylinder is unclear. However, since the tool performance was verified by running a PH₃ control prior to running the sample cylinders, the observation may be linked directly or indirectly to progressively decreasing H₂O concentration as PH₃ is flowed over time through the delivery system, as discussed in the following sections. Results in Ref. [5], showing the effect of H₂O impurity in PH₃ on mobility of InP, support our inferences in this regard.

3.3. Phosphorus oxide/oxyacid formation and characterization

Factors that can influence the delivery of high-purity gases have been discussed previously [16,17]. For PH₃, cylinder connection or disconnection cycle purge techniques are extremely important, because if not undertaken correctly, traces of air can remain and then react with PH₃ to form solid residues in the delivery system. Previous studies indicate that the residue is a mixture of lower oxides and oxyacids of phosphorus such as H₃PO, H₂PO, HPO, PO and PO₂ intermediates [18] that can go on to form hypophosphorous, phosphorous and phosphoric acids [19]. However, the presence and effects of these contaminants in gas delivery systems has not been discussed in detail in the literature.

To mimic what happens in an incompletely purged delivery line, PH₃ at 40 psig was intentionally introduced into a SS tube that contained air at atmospheric pressure. After removing any residual PH₃, the deposited H_xPO_y residues were analyzed by SEM-EDS (Fig. 3). The yellow colored residue in Fig. 3A had a composition of PO₂ based on EDS (data for H could not be measured), was polycrystalline in nature and had a high density and surface area. The average size of the particles on the surface of the layer was about 0.5 μm. A red-brown residue, with a composition PO, was also formed on a steel rod (Fig. 3B) inserted into the delivery tubing. The SEM image shows this material also had a high porosity and surface area. The two deposits are chemically and structurally different probably because of the different locations of each sample in the tube and how the reaction took place. The rod was positioned farthest from where the PH₃ entered the tube, whereas the yellow deposit was taken from the inside wall of the tube closest to the point of

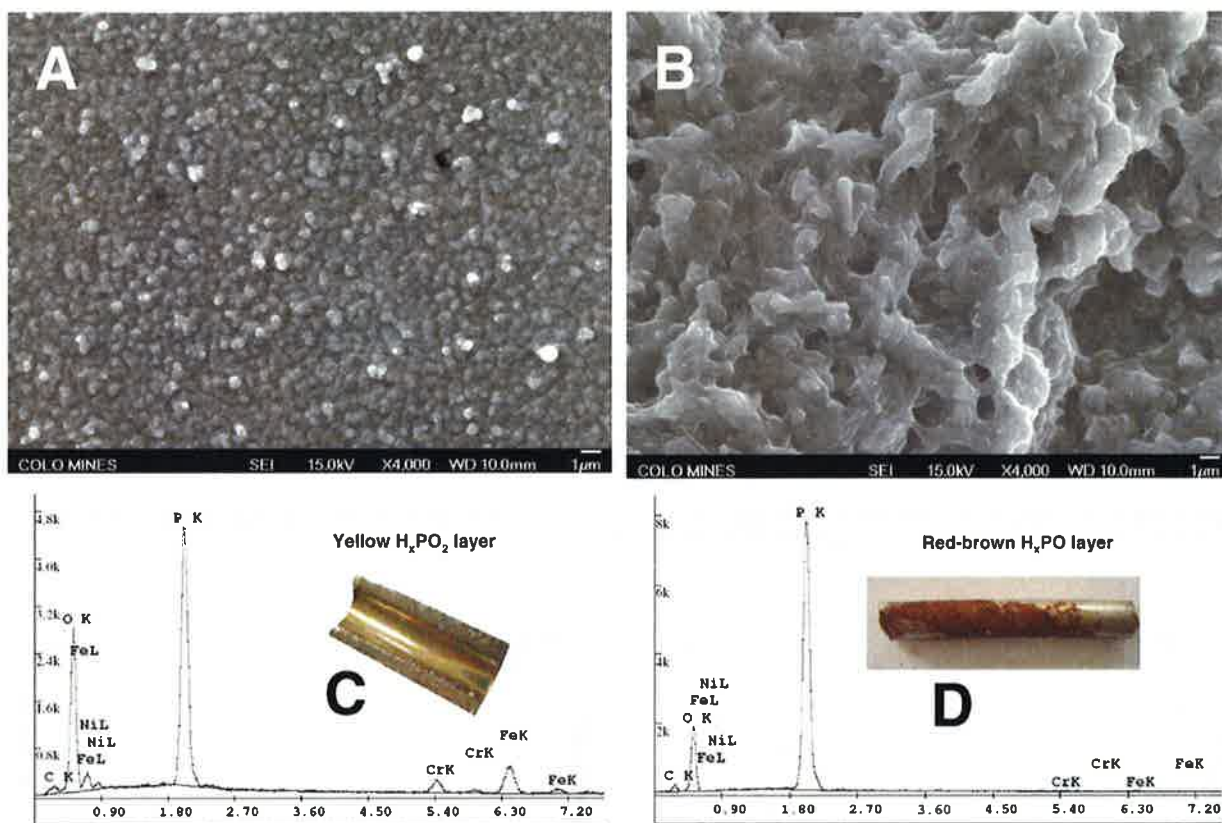


Fig. 3. SEM image of yellow H_xPO_2 on inner tube surface (A) with EDS analysis (C) and photograph (inset); SEM image of red-brown H_xPO on steel rod (B) with EDS analysis (D) and photograph (inset).

entry of the PH_3 . The higher oxygen content of the yellow deposit indicates that the PH_3 quenched much of the O_2 as it entered the second tube. Interestingly, although the samples were kept in a N_2 atmosphere, when the red-brown sample was loaded into the SEM vacuum chamber for analysis, the SEM shut down because the chamber pressure criterion could not be met. This indicates that the residue outgassed large amounts of H_2O and is evidence of the hygroscopic nature of H_xPO_y contamination.

3.4. Effect of H_xPO_y on PH_3 cylinder valve and delivery system

We studied the water vapor behavior of PH_3 Cylinder 2, which was returned from the field after observing the mobility increase discussed above. After routine cycle-purging of the cylinder connection and sampling manifold, over $3 \mu\text{mol mol}^{-1}$ H_2O was detected in the PH_3 , the first time it was analyzed. The cylinder was kept connected in-line with the cylinder valve closed and the entire sampling system was purged with purified N_2 for over 24 h. When PH_3 was flowed again from the cylinder a second time, the H_2O had decreased but was still $\sim 500 \text{ nmol mol}^{-1}$. After a further 24 and 48 h of purging the cylinder fittings and the delivery system with purified N_2 , water measurements from the PH_3 cylinder were close to the original concentration of $180 \text{ nmol mol}^{-1}$ (Fig. 4). The slow H_2O dry-down was attributed to H_xPO_y contamination in the cylinder valve. Therefore, even though the gas in the cylinder may meet the H_2O specification, if the cylinder valve contains H_xPO_y residues, dry-down times can be extensive and H_2O in the delivered PH_3 may be elevated for some time and affect devices.

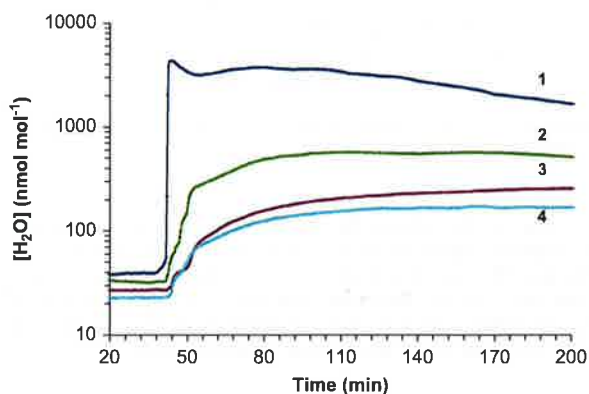


Fig. 4. H_2O concentration during sequential analysis of PH_3 Cylinder 2.

Fig. 5 compares the wet-up and dry-down characteristics of a clean PH_3 delivery system and a system contaminated with H_xPO_y . Curve A in Fig. 5 shows the rapid H_2O equilibration in the clean delivery line when the H_2O level in the PH_3 was changed by passing the PH_3 through the purifier or purifier bypass line. The dry-down time to reach 95% of the final water concentration from 2500 to $100 \text{ nmol mol}^{-1}$ was about 20 min. In contrast, curve B in Fig. 5 shows a very slowly equilibrating H_2O profile in a contaminated line, taking over 230 min to reach 95% of the final H_2O concentration.

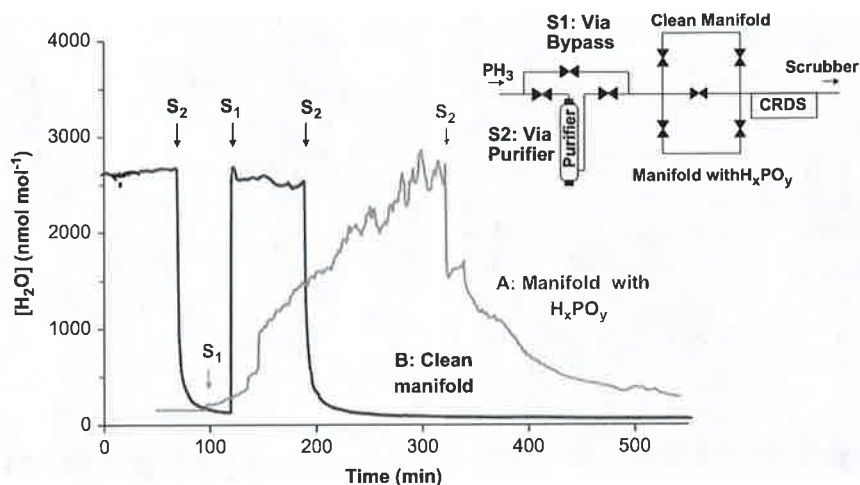


Fig. 5. Wet-up and dry-down trend of manifolds with (A) and without (B) H_xPO_y contamination. In each case wet and dry PH₃ was flowed through the manifolds. S1 and S2 are the points at which PH₃ is switched via purifier bypass and PH₃ purifier, respectively.

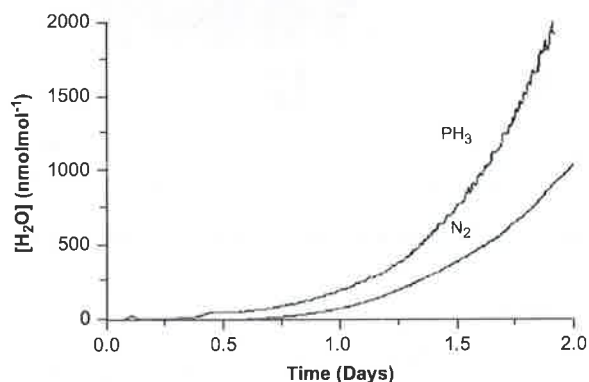


Fig. 6. H₂O concentration increase in PH₃ and N₂ under stagnant conditions over the course of several days.

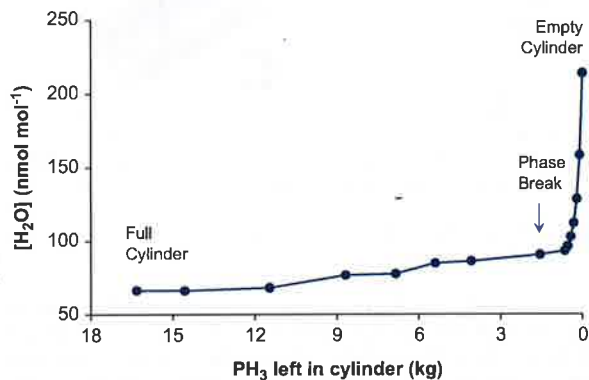


Fig. 7. H₂O concentration profile as PH₃ gas is withdrawn from a high-purity PH₃ cylinder from full to empty condition.

These results indicate that H_xPO_y contamination, can absorb H₂O if dry or release H₂O if wet. As shown in Fig. 4, release of H₂O into dry PH₃ can elevate the H₂O level in the delivered gas to μmol mol⁻¹ levels. Therefore even though the gas source itself may be highly pure, the purity at point-of-use may be substantially lower and cause performance variations of III–V devices.

3.5. H₂O impurity generation by reaction of PH₃ with metal oxides

Another factor that may influence the purity of delivered PH₃ gas is the generation of H₂O by reaction of hydride gases with metal oxides on the exposed steel surfaces of the gas delivery system [20].



In this work we studied the trend in water vapor in PH₃ under stagnant conditions (no flow). Fig. 6 shows the build-up of H₂O in PH₃ is much higher than that in N₂ under the same static conditions. The H₂O increased from low nmol mol⁻¹ levels to about 610 nmol mol⁻¹ in PH₃ and 420 nmol mol⁻¹ in N₂ after 36 h. After 48 h it increased to 1900 nmol mol⁻¹ in PH₃ and

900 nmol mol⁻¹ in N₂. In the N₂ case, the water is due to outgassing from the inner metal surfaces. In the case of PH₃, water not only continues to outgas but is also generated by reaction (1). The results show that in manifold design or use, it is important to keep process or purge gas under dynamic flow conditions rather than static conditions.

3.6. Water vapor profile during PH₃ gas consumption

Previous cylinder depletion studies on hydride gases have shown that gas phase H₂O can vary during gas withdrawal from the cylinder [3]. In this work we tracked the water vapor profile during depletion of a high-purity PH₃ cylinder (Fig. 7). The full cylinder (with ~16.33 kg PH₃) contained ~66 nmol mol⁻¹ H₂O. From the full condition to liquid phase-break (~1.56 kg PH₃ left), the H₂O level remained approximately constant and below the 100 nmol mol⁻¹ specification. As flow continued and the cylinder pressure finally dropped to ~137.9 kPa, the H₂O concentration increased to 214 nmol mol⁻¹. This result shows the importance of using a high-purity PH₃ source with low H₂O, switching out

cylinders in good time and/or the use of purification as discussed below, to ensure that water vapor is kept in control during III–V device fabrication.

Since water vapor can come from several sources and the delivery system may be difficult to purge and dry down due to the presence of even traces of H_xPO_y contamination, point-of-use purification is recommended close to the tool. A dry-down experiment with a ~ 3 m delivery line showed that as Nanochem PHX purified PH_3 was introduced into the line, H_2O was reduced from $200 \text{ nmol mol}^{-1}$ to single digit nmol mol^{-1} levels in < 100 min.

4. Conclusion

Increasing mobility in InP films with run time has highlighted the importance of H_2O contamination issues in PH_3 delivery systems. The purity of delivered PH_3 is dependent on several factors. As with other gases, the source purity is important. Impurities such as H_2O may rise as gas is withdrawn from the cylinder, particularly close to or after phase-break. However additional issues exist for PH_3 . Water may be generated by reaction of the PH_3 with surface metal oxides. Further H_xPO_y solids can slowly build up in delivery systems due to incomplete purging of atmospheric oxygen which reacts with the PH_3 . H_2O can be emitted from these H_xPO_y solids and significantly lengthen dry-down times. To consistently control impurities, high-purity PH_3 and at-tool point-of-use purification is recommended.

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