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Spectrochimica Acta Part A 57 (2001) 959-969

SPECTROCHIMICA ACTA PART A

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$Na_{3}(PO_{2}NH)_{3}.4H_{2}O, [C(NH_{2})_{3}]_{3}(PO_{2}NH)_{3}.H_{2}O \text{ and} (NH_{4})_{4}(PO_{2}NH)_{4}.4H_{2}O$

Annamma John^a, Daizy Philip^b, Norbert Stock^c, Wolfgang Schnick^c, S. Devanarayanan^{d,*}

^a Department of Physics, St. John's College, Anchal, Kollam 691 306, India
 ^b Department of Physics, Mar Ivanios College, Thiruvananthapuram 695 015, India
 ^c Laboratorium fur Anorganische Chemie der Universitat, D-95440 Bayreuth, Germany
 ^d Department of Physics, University of Kerala, Kariavattom, Thiruvananthapuram 695 581, India

Received 31 July 2000; accepted 9 August 2000

Abstract

FT IR and FT Raman spectra of $Ag_3(PO_2NH)$, (Compound I), $Na_3(PO_2NH)_3$.H₂O (Compound II), $Na_3(PO_2NH)_3$.H₂O (Compound III), $[C(NH_2)_3]_3(PO_2NH)_3$.H₂O (Compound IV) and $(NH_4)_4(PO_2NH)_4$.H₂O (Compound V) are recorded and analyzed on the basis of the anions, cations and water molecules present in each of them. The PO₂NH⁻ anion ring in compound I is distorted due to the influence of Ag⁺ cation. Wide variation in the hydrogen bond lengths in compound II is indicated by the splitting of the v_2 and v_3 modes of vibration of water molecules. The NH₄ ion in compound V occupies lower site symmetry and exhibits hindered rotation in the lattice. The correlations between the symmetric and asymmetric stretching vibrations of P–N–P bridge and the P–N–P bond angle have also been discussed. © 2001 Elsevier Science B.V. All rights reserved.

Keywords: Infrared spectrum; Raman spectrum; Trimetaphosphimate; Tetrametaphosphimate; Hydrogen bonding

1. Introduction

Many salts of trimetaphosphimic acid and tetrametaphosphimic acid were known more than 100 years ago [1,2]. These compounds have been mainly characterized by IR spectroscopy [3,4] and X-ray powder diffraction [5]. Extensive amount of work has been done on the study of the characteristic features of the coordination of metaphosphimate anions by vibrational spectroscopy, involving the calculation of wavenumbers and forms of the normal vibrations [6–16]. The trimetaphosphimate anion is a cyclic system formed by three regularly alternating slightly distorted PO_2NH tetrahedra. The confirmation of the ring in the chair form with symmetry close to

^{*} Corresponding author. Tel.: +91-471-418920; fax: +91-471-307158.

E-mail address: phy@univker.ernet.in (S. Devanarayanan).

 C_{3v} has been established by X-ray structure analysis [17] and from the vibrational spectra of alkali metal trimetaphosphimates [9]. From a study of the vibrational spectra it is known that the tetrametaphosphimate anion in solution has S_4 symmetry and that in salts it has two kinds of structures, viz., C_{2h} symmetry and S_4 symmetry [14].

The present authors are interested in the study of the Raman and IR spectra of some newly synthesized phosphate compounds [18-20]. In this chapter a report of the analysis of both the IR and Raman spectra of Ag₃(PO₂NH)₃ (hereafter referred as compound **I**), Na₃ (PO₂NH)3.H₂O (compound II), Na₃(PO₂NH₃) .4H₂O (compound III), [C(NH₂)₃]₃(PO₂NH)₃.H₂O (compound IV) and (NH₄)₄(PO₂NH)₄.4H₂O (compound V) is made. The Raman spectra of these samples in aqueous solution are also recorded to clear the ambiguity regarding certain vibrations.

2. Experimental

The starting material Na₂(PO₂NH)₃.4H₂O (compound **III**) was prepared by the method reported by Olthof et al. [17]. To an acidified solution of this compound, an aqueous solution of AgNO₃ is added to get Ag₃(PO₂NH)₃ (compound **I**) [21]. Na₃(PO₂NH)₃.H₂O (compound **II**) was obtained by vapour diffusion of CH₃CH₂OH at 363 K into a 3×10^{-4} M aqueous solution of the starting material [22]. [C(NH₂)₃]₃(PO₂NH)₃.H₂O (compound **IV**) was obtained by the addition of [C(NH₂)₃]₂CO₃ to a freshly prepared solution of H₃(PO₂NH)₃ [23]. For the preparation of

Table 1 Crystallographic datas of the five compounds

Compounds	Crystallo- graphy	Space group	Z ^b	Refs.
I	Monoclinic	$P2_1/c(C_{2h}^5)$	4	[21]
П	Monoclinic	$C2(C_{2}^{3})$	2	[22]
Ш	Monoclinic	$P2_1/n(C_{2h}^5)$	4	[17]
IV	Orthorhombic	$Pbca(D_{2h}^{15})$	8	[23]
V	Monoclinic	$P2_1/n(C_{2h}^5)$	2	[24]

 $(NH_4)_4(PO_2NH)_4.4H_2O$ (compound V), the starting material $H_4(PO_2NH)_4.2H_2O$ was obtained according to the literature [6] by hydrolysis (333 K, 48 h) of (PNCl₂), in dioxane. Then an aqueous solution of ammonia (10%) is added to a stoichiometric amount of the acid followed by precipitation with acetone [24].

The IR and Raman spectra were recorded using a Bruker IFS 66V FT-IR/FT-Raman Spectrometer. The Raman spectra were recorded with a resolution of 4 cm⁻¹, using an Nd:YAG laser at wavelength 1.064 μ m and power 200 mW, over the range 50–3500 cm⁻¹. The IR spectra are obtained in the range 400–4000 cm⁻¹ with the samples as KBr pellets using a globar source.

3. Factor group analysis

The compounds I, II, III and V crystallizes in the monoclinic system and compound IV is orthorhombic. The crystal symmetry data of all the five compounds used in the present work are tabulated in Table 1.

The cations, anions and water molecules in all the compounds occupy general position. The trimetaphosphimate anion $(PO_2NH)_3^{3-}$ and tetrametaphosphimate anion (PO₂NH)₄⁴⁻ exhibit chair conformation, but in compound I the anion is markedly distorted because of the interaction between Ag⁺ and one nitrogen atom of the anion [21]. The $(PO_2NH)_3^3$ ring has C_{3y} symmetry [13] and the $(PO_2NH)_4^4$ ring has C_{2h} symmetry [15]. The cyclic anion rings in trimetaphosphimates are linked to one another by N-H---O hydrogen bonds to form pairs. These units are interconnected by O-H---O hydrogen bonds through water molecules forming columns. In compound V the tetrametaphosphimate rings are finked by N-H---O bonds forming column, along [100]. These columns are interconnected by O-H---O and N-H---O hydrogen bonds through water molecules and ammonium ions [21-24]. All H atoms are involved in hydrogen bonding. In II and III the sodium ions are approximately octahedrally coordinated by six oxygen atoms, either from the phosphonate groups or from the water

Compound I	$54A_{g}(R) + 54B_{g}(R) + 53A_{u}(IR) + 52B_{u}(IR)$
Compound II	62A(R,IR) + 61B(R, IR)
Compound III	$90A_{g}(R) + 90B_{g}(R) + 89A_{u}(IR) + 88B_{u}(IR)$
Compound IV	$144 \ddot{A}_{g}(R) + 14 \ddot{4} B_{1g}(R) + 144 B_{2g}(R)$
	$+ 144B_{3g}(R) + 144A_{u}(O) + 143B_{1u}(IR)$
	$+ 143B_{2u}(IR) + 143B_{3u}(IR)$
Compound V	$78A_g(R) + 78B_g(R) + 77A_u(IR) + 76B_u(IR)$

4. Results and discussions

The IR and Raman spectra recorded for all the five compounds under investigation are shown in Figs. 1-3. The observed spectral lines and their assignments are given in Table 2.

4.1. Vibrations of PO_2NH^- anion

Analysis of the spectra obtained shows that vNH modes are highly characteristic. These are obtained as weak bands in Raman and medium to strong in IR around 3170 cm⁻¹ in all the five compounds. The multiplicity of bands observed in compound I may be due to the participation of the NH groups in intramolecular and intermolecular hydrogen bonding [13]. The frequency shift observed in compound IV may be attributed to the overlapping of the vNH vibrations with that of vNH₂ vibrations. The NH deformations are observed in the region 1390–1322 and 781–723 cm⁻¹.

As expected the asymmetric and symmetric stretching vibrations of PO₂ group are observed in the region 1241–1141 and 1118–1039 cm⁻¹ respectively [26]. The asymmetric modes are very strong in the IR spectrum and the symmetric modes are very strong in the Raman spectrum. Both the asymmetric and symmetric stretching modes of PO₂ in compound V appear a little lower in frequency when compared to all the

other four compounds. This is due to the greater value of the average P–O bond length in compound V than that in all the other compounds [24]. The bending modes of PO₂ group are observed around 600 cm⁻¹. The wagging, rocking and twisting modes of PO₂ group are also assigned (Table 2).

The ring vibrations of the anion — the antisymmetric and symmetric stretches of P-NH-P bridges are expected in the region 1000-800 cm^{-1} and the ring breathing vibration around 400 cm⁻¹ [12,15]. In compound I the asymmetric stretching gives very weak Raman bands at 971 and 937 cm^{-1} and the symmetric stretching bands are observed at 831 and 793 cm⁻¹. The corresponding IR bands are singlets observed at 933 and 813 cm⁻¹ respectively. The large splitting (≈ 138 cm⁻¹) of the symmetric mode observed in the Raman spectrum confirms the distortion of the PO_2NH^- anion ring due to the influence of the stronger Ag⁺ cation on the anion structure [21]. In the Raman spectrum of compound V, a weak band is observed at 1008 cm⁻¹. Sukova et al. [15], in their studies on lanthanum and lanthanide tetrametaphosphimate had assigned the strong IR band around ≈ 1005 cm⁻¹ to v_{as} P–NH–P along with its components in the region 980-900 cm⁻¹. Accordingly the Raman band at 1008 cm^{-1} , in the present study, is assigned to $v_{as}P-NH-P$. Its components are the very weak bands at 980-910 cm^{-1} .

The corresponding IR. bands are very strong at 992 and 955 cm⁻¹. The weak to medium intensity Raman bands around 493 cm⁻¹ are due to the ring breathing vibrations. The IR spectra show medium to strong intensity bands in this region. The external deformation mode is assigned to the weak bands around 225 cm⁻¹ [9,12].

4.2. Vibrations of NH_4^+ cation

The normal modes of vibration of free NH₄⁺ ion under perfect T_d symmetry have frequencies of 3033, 1685, 3134 and 1397 cm⁻¹ for $v_1(A_1)$, $v_2(E)$, $v_3(F_2)$ and $v_4(F_2)$ modes, respectively [26].

Table 2 Spectral dé	ata (cm ⁻¹) a:	nd band assig	gnments of <i>i</i>	Ag ₃ (PO ₂ NH) ₃ ,	, Na ₃ (PO ₂ NF	H) ₃ .H ₂ O, Na ₃ (PO	D ₂ NH) ₃ .4H ₂	O, [C(NH ₂) ₃] ₃	PO2NH)3.H	$_2 O$ and $(NH_4)_4 (PO_2NH)_4.4 H_2 O^a$
Ag ₃ (PO ₂ N	H) ₃	Na ₃ (PO ₂ NI	H) ₃ .H ₂ O	Na ₃ (PO ₂ NH	0,3.4H ₂ O	[C(NH ₂) ₃] ₃ (PC	$(1)^{2}$ $(1)^{3}$	(NH ₄) ₄ (PON	$\rm H)_{4.}4H_{2}O$	Assignments
Raman	IR	Raman	IR	Raman	IR	.H ₂ 0 Raman	IR	Raman	IR	
	3472wbr				3455m		3586m 3418s	3377vw	3571m 3353sh	Combination
		3214w	3234s	3300wbr	3310m 3310m	3352w	3350sh	3301wbr	3265sh 3213sh	v_3H_2O v_1H_2O v_3NH_4
3273w 3233w	3223w 3151m	3173w	3168s	3173wbr	3173m	3198wbr	3126s		3162vs	$v \mathrm{NH}/v \mathrm{NH}_2$
3086VW	3049Sh							3058vwbr	3018sh	v_1 NH ₄
	2684w 2596vw		2678w 2659w		2704vvw 2644vvw				2716sh	combination v NHN(O ⁻)
			2575w 2321vw						2359vw	Combinations and overtones
2119vw		2124vw	2185w	208 Sumbr					2145m	
			1720w	10 M A COOZ						
	1643w		1686sh	1670vwbr	1694w 1661w 1603w	1649vw	1649sh	1688vw	1653m 1653m	$v_2 H_2 O$
1471w			1545w			1588vw	1663vs 1442sh			v _{as} CN δNH ₂ Combinations
								1420vwbr	1461s 1419sh 1402s	$v_4 \mathrm{NH}_4$
1390vvw 1369vvw	1345s		1355sh 1335vs		1355m 1326m		1322m		1385sh 1362sh 1343s	δNH
					1281m 1241s		1296sh			90H-Об
	1226vs	1222w	1211vs	1238m 1198m	1225s 1192m	1234vw	1210s	1192sh	1201vs	$v_{ m as}{ m PO}_2$

Table 2 (C	Jontinued)										
$Ag_3(PO_2N)$	H ₃	Na ₃ (PO ₂ NI	H) ₃ .H ₂ O	Na ₃ (PO ₂ NF	H) ₃ .4H ₂ O	$[C(NH_2)_3]_3(PO_2)$	NH) ₃ .H ₂ O	(NH ₄) ₄ (PONH) ₄ .4H ₂ O	Assignments	
Raman	IR	Raman	IR	Raman	IR	Raman	IR	Raman	IR		
1167w		1185w	1187s	1182m		1174vw		1180w 1148w	1141s		
1094vs	1111m	1107vs	1101s	1107vs	1118w	1106m	1102m	1080vs		$v_{\rm s} { m PO}_2$	
1040w	1054vs	1059w 1039w	1060vs 1043vs	1054m 1047m	1066vs 1054s	1049vw 1007vs	1046vs 1006w		1063s		
971 vw								1008w 980vw	992s	$q_{\rm -}HN-q_{\rm co}$	
937vw	933vs	955w	952vs	955vw 933vw 845w	960s 931s 860vw	921 vvw 853vw	946sh 929s	910vw 859vvw	955vs	Pe	
831 vw 793m	813m		826s		821m				818m	v _s P–NH–P	
		803m		806w	802m		l			H ₂ O liberation	
755s	774sh 731m	658vw	781m 662m	726m	740m 650sh	723wbr	779sh	745s		HNõ	
610s	602m 540s	607m	599m 542s	601m	608m	607 vw	608m	596vvw 560w	589m 556m	δPO_2	
		524m			53.6m	531m	515w			δCN ₃	
512w 470w	504s		513s 480m	517m 478m	504m		493w 492m	474w	506s	Ring breathing	
402		374m		377sh 360s		385sh 351w	4248n	423W 368s	421VW	ωPO ₂	
345w		- 010									
334w 297m		3188h 295m		297s		292w 277w		332m 288m		ρPO_2	
258vw		231vw		247w		5 1		228m 209m		Ring deformation	
167sh				157vw				166m		$^{1}\mathrm{PO}_{2}$	
124m		131w		129vw		134s		137m		0H_Nu	
040		1 14sh 84		04		109sh 02		105m 84m		VU-HU External mode	
\$40		04111		04111		шсо		04111			
с. С.			-								

^a Relative intensities: v, very; w, weak; m, medium; s, strong; sh, shoulder; br, broad.



Fig. 1. FT IR spectra of (a) $Ag_3(PO_2NH)_3$, (b) $Na_3(PO_2NH)_3$.H₂O (c) $Na_3(PO_2NH)_3$.4H₂O, (d) $[C(NH)_3]_3(PO_2NH)_3$.H₂O and (e) $(NH_4)_4(PO_2NH)_4$.4H₂O in the region 400–4000 cm⁻¹.

All the four modes are Raman active whereas v_1 and v_2 are IR inactive. In the crystal, the ion occupies a lower site symmetry, which may lead to the activation of inactive modes, splitting of degenerate modes and shifting of fundamental modes.

In the Raman spectrum of compound V, the weak broad band at 3058 cm^{-1} is assigned to the symmetric stretching mode v_1 . The corresponding IR band appears as a shoulder at 3018 cm⁻¹. The broadening (≈ 250 cm⁻¹) of the v_1 mode indicate the participation of the NH₄ ions in the formation of different types of hydrogen bonds N-H---N and N-H---O of varying strength [27]. The struc-tural investigation of compound V [24] reports that the NH_4 ions take part in extended hydrogen bonding by bridging, via, the lone pairs of OW2, N4-H42---OW2---H31-N3 (N---O---N 2.801, 2.858 Å). The observation of strong IR band extending from about 3700 to 2700 cm⁻¹ (FWHM \approx 720 cm⁻¹) also supports this. The absorption shoulders at

3265 and 3213 cm⁻¹ are due to the triply degenerate asymmetric stretching mode v_3 . The v_2 mode is observed as weak absorption band at 1699 cm⁻¹. The very weak Raman band at 1688 cm⁻¹ is the counterpart of this mode. The absorption triplet at 1461, 1419 and 1402 cm⁻¹ are due to the triply degenerate v_4 mode of the ammonium ion.

The splitting of the v_3 and v_4 mode and the activation of the IR inactive modes v_1 and v_2 supports the lowering of site symmetry (C₁) of the ion. All the four modes of the NH₄ ion are found to be shifted from the free state values. This observation, in agreement with the structural reports, shows that the ion is distorted from the ideal tetrahedral geometry [24].



Fig. 2. FT Raman spectra of (a) $Ag_3(PO_2NH)_3$, (b) $Na_3(PO_2NH)_3$. H_2O (c) $Na_3(PO_2NH)_3$. H_2O , (d) $[C(NH_2)_3]_3(PO_2NH)_3$. H_2O and (e) $(NH_4)_4(PO_2NH).4H_2O$ in the region 50–3500 cm⁻¹.



Fig. 3. FT Raman spectra of the aqueous solution of (a) $Na_3(PO_3NH)_3$, H_2O (b) $[C(NH_2)_3]_3(PO_2NH)_3$, H_2O and (c) $(NH_4)_4(PO_2NH)_4$, $4H_2O$ in the region 50-2000 cm⁻¹.

In a freely rotating NH₄ ion, the v_3 and v_4 modes are triply degenerate. However if it is in the locked-in position in the crystal lattice, the degeneracy is expected to be lifted [28]. In the present study, the removal of degeneracy of the v_3 and v_4 modes suggests that the ion is probably in a locked in position in the crystal lattice. The fundamental v_6 could not be identified independently as it merges in the region of the bending mode of PO₂. However, its combinations $v_2 + v_6$ and $v_4 + v_6$ are not observed. Hence it may be inferred that the NH₄ ion is executing hindered rotation in the lattice [29–32].

4.3. Vibrations of $[C(NH_2)_3]^+$ cation

In the structural studies of triguanidinium tri- μ imido-cyclotriphosphate monohydrate (compound **IV**) Stock et al. have assigned highly intense IR band at 1662 cm⁻¹ to the asymmetric CN stretching vibrations. The IR spectrum of tetraguanidinium tetra-µ-imido-cyclotetraphosphate tetrahydrate $[C(NH_2)_3]_4(PO_2NH)_4.4H_2O$ also shows a highly intense IR band at 1659 cm⁻¹ due to v_{as} CN [28]. In the present investigation, the very strong absorption band at 1663 cm⁻¹ is assigned to the asymmetric CN stretching vibrations. The very high intensity of this band differentiate it from the v_2 mode of water molecules, which are usually weak to moderate in IR. The symmetric CN stretching is very strong in Raman at 1007 cm⁻¹. Its corresponding IR band is weak at 1006 cm⁻¹. The high intensity of this band in the Raman spectra and the low intensity in the IR spectra, differentiate it from the $v_{as}P-NH-P$ mode, which is usually weak in Raman and intense in IR. The Raman spectrum of the aqueous solution also shows, this intense band without much shift (1010 cm^{-1}), which support our assignment of v_s CN. The deformation of CN₃ gives a weak band at 515 cm⁻¹. The vNH_2 mode is masked in the broad bands of vNH. However, the bands of NH₂ deformation are clearly identifiable. The weak Raman band at 1588 cm^{-1} and the shoulder at 1542 cm⁻¹ in the IR spectrum are due to the δNH_2 vibration.

4.4. Vibrations of water molecules

The v_1 and v_3 modes of water molecules are observed in the region 3472-3214 cm⁻¹ and the bending mode v_2 around 1661 cm⁻¹. The existence of broad overlapping bands in the stretching region shows the presence of both weakly and strongly bonded molecules of water of crystallization and of hydrogen bridges with different bond energies. The absorption band at 3472 and 1643 cm^{-1} in the unhydrated compound I are due to the water molecules adsorbed during pelletization. In the IR spectrum of compound III the v_3 mode of H₂O appear as a doublet at 3455 and 3372 cm⁻¹ and the v_2 mode as triplet at 1694, 1661 and 1639 cm⁻¹. This splitting of the fundamental vibrations are due to the wide variation in the hydrogen bond lengths, where the O---O ranges from 2.821 to 4.144 Å [17]. The librational modes of H₂O are observed as weak to medium intense bands in the region 806-802 cm⁻¹. This assignment is confirmed by the broad bands observed about this region in their solution spectra.

4.5. Hydrogen bonding

Structural investigations report that, in all the five compounds investigated, two anion rings are linked to each other by N–H---O hydrogen bonds forming pairs [17,21–24]. These units are interconnected by O–H---O hydrogen bonds through water molecules.

As a result of the comparison of the spectra of tetrametaphosphimates with NH group in the ring, with the spectrum of octa silver salt Ag_g (PO₂N)₄, Lunkwitz and Steger [7] were able to distinguish two bands at 3075 and 2675 cm⁻¹ corresponding only to NH groups and bridging hydrogen bonds NH---N(O⁻), respectively. The weak absorption bands in the region 2716–2575 cm⁻¹ are accordingly assigned to NH...N(O⁻) vibrations.

4.6. Correlation between the P-N-P stretching vibrations and the P-N-P angle

The correlation between the P–O–P angle and the asymmetric and symmetric stretching frequencies of the P–O–P bridge are well known [33–37]. The correlation exhibits a definite pattern of relationship. Several authors have used these correlations to calculate the unknown P–O–P angles in pyrophosphates [36,37] and cyclohexaphosphates [19]. Lazarev, [37] has used similar type of correlations in pyrosilicates and determined the Si–O–Si angles.

In the present investigation, a similar type of correlation is attempted for the P-N-P group. The structural and spectral data used are listed in Table 3. The curves (Fig. 4) are plotted for the two relationships

1. $v_{as}PNP - v_{as}PNP = f_1(\angle PNP)$

2. $(v_{as}PNP - v_{s}PNP)/(v_{as}PNP + v_{s}PNP)$

 $= f_2(\angle PNP)$

The curves show a definite pattern as in the case of POP bridge.

Table 3 Correlation between the PNP bridge angle and its stretching vibrations

Compound	$v_{as}PNP$ (cm ⁻¹)	$v_{\rm s} {\rm PNP}$ (cm ⁻¹)	$v_{\rm as} - v_{\rm s}$ (cm ⁻¹)	$(v_{\rm as} - v_{\rm s})/(v_{\rm as} + v_{\rm s})$	∠PNP (°)	Refs.
Ag ₃ (PO ₂ NH) ₃	933	813	120	0.0687	125.5	[21]; present work
Na ₃ (PO ₂ NH) ₃ .H ₂ O	952	826	126	0.0709	124.3	[22]; present work
$Na_3(PO_2NH)_3.4H_2O$	960 931	860 821 802	100 110 129	0.0549 0.0628 0.0744	121.0 122.6 124.4	[17]; present work
[C(NH ₂) ₃] ₃ (PO ₂ NH) ₃ .H ₂ O	921	853	68	0.0373	122.1	[23]; present work
$(NH_4)_4(PO_2NH)_4.4H_2O$	992 955	818	137 174	0.0773 0.0961	128.61 130.1	[24]; present work
Rh(PO ₂ NH) ₃ (NH ₃) ₃ .H ₂ O	935	870	65	0.03601	121	[16]
$Na_{4}\{Cu[(PO_{2}NH)_{3}]_{2}\}.10H_{2}O$	960 930	820 800	110 160	0.0628 0.0909	122 133	[15]
$[C(NH_2)_3]_4(PO_2NH)_4.4H_2O$	976 941	820 773	121 156	0.0687 0.0869	123 130	[23]



Fig. 4. Relationship between (a) the angle PNP and $v_{as}PNP - v_sPNP$ and (b) the angle PNP and $(v_{as}PNP - v_sPNP)/(v_{as}PNP + v_sPNP)$.

5. Conclusions

The large splitting of vs. P–NH–P modes in compound I indicates distortion of the PO₂NH⁻ anion ring due to the influence of Ag⁺ cation. NH₄⁺ ion exhibits hindered rotation in the lattice. The lifting up of the degeneracy of v_3 and v_4 mode and the activation of the IR inactive modes indicates lower site symmetry of the NH, ion. Splitting of the v_2 and v_3 modes of vibration of water molecules in compound III indicates the wide variation in the hydrogen bond lengths. The correlation between angle P–N–P and the stretching vibrations of the P–N–P bridge exhibit a definite pattern.

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