# SERS spectra of $(NH_3)_2(CH_2)_2HXO_4$ (X = P, As) adsorbed in silver colloids

Daizy Philip, Antony Eapen and G Aruldhas\* Department of Physics, University of Kerala, Kariavattom, Trivandrum-695 581, India

Received 19 May 1994, accepted 27 July 1994

**Abstract** : SERS spectra of  $(NH_3)_3(CH_2)_3HPO_4$  and  $(NH_3)_3(CH_2)_3HAsO_4$  are recorded in two types of silver colloids. In the phosphate, two distinct adsorption sites are present which give rise to different enhanced bands in the two colloids. The ethylene diaminonium group is close to the silver surface in one colloid whereas the phosphate group is closer in the second colloid. In the arsenate, bands due to both ethylene diaminonium and arsenate groups are enhanced in one type of colloid whereas in the second colloid only the bands due to the arsenate ions are enhanced

**Keywords** : SERS spectra, silver colloid, adsorption sites, ethylene diammonium phosphate, ethylene diammonium arsenate

PACS No. : 33 20 -t

Surface Enhanced Raman Scattering (SERS) spectroscopy is a powerful tool for investigating the co-ordination and geometry of adsorbed molecules on metal surfaces. Phosphates being poor scatterers are expected to be good adsorbers compared to arsenates [1]. In continuation of our earlier study [2] on the IR and polarized Raman spectra of  $(NH_{3})_2(CH_2)_2HPO_4$ , ethylene diammonium phosphate (EDP), and  $(NH_3)_2(CH_2)_2HAsO_4$ , ethylene diammonium arsenate (EDA), we report here the results of their SERS spectra in silver colloids. Greaves and Griffith [1] have investigated the SERS spectra of certain vanadates, phosphates and arsenates and found larger enhancement factor for phosphate compared to arsenates. They also noticed that larger anions interact less effectively with the silver surface.

Two different silver colloids (i) greenish-yellow (colloid 1) and (ii) greenishgrey (colloid 2) have been used in the investigation. Colloid 1 prepared [3] from sodium

## 464 Daizy Philip, Antony Eapen and G Aruldhas

borohydride and silver nitrate has a strong absorption maximum at 390 nm. Colloid 2 is prepared [4] using silver nitrate and sodium citrate and has a broad absorption band around 430 nm. For recording SERS spectra, the samples are prepared by mixing equal volumes of colloid 1/ colloid 2 and  $10^{-4}$  M aqueous solution of the compound. Normal Raman spectra could not be obtained for aqueous solutions of the compounds. SERS spectra of the compounds in freshly prepared colloid 1 are also recorded. The chemicals have been procured from Sigma, USA and deionized water has been used throughout. Raman spectra were recorded on a Dilor GMBH Z 24 spectrometer using 514.5 nm radiation (power 200 mW) of an Ar<sup>+</sup> laser. Absorption spectra were recorded using a UV-Shimadzu UV-Visible recording spectrophotometer.

EDP and EDA have absorption bands around 340 and 400 nm respectively. The addition of  $10^{-4}$  M aqueous solution of EDP to colloid 1 changes the colour of the solution to light pink with an additional very broad absorption band around 620 nm. However in colloid 2, the absorption band around 430 nm becomes broadened with another extinction maximum around 700 nm. EDA in colloid 1 shows two broad bands around 650 and 395 nm. As in EDP in colloid 2, the band around 430 nm becomes more broadened.

In the SERS spectrum of EDP (Figure 1), a low frequency band is observed at 234 cm<sup>-1</sup> in colloid 1 and at 217 cm<sup>-1</sup> in colloid 2. Further, in colloid 1 bands corresponding to  $(NH_3)_3(CH_2)_2^{2+}$  are enhanced which are not observed in colloid 2 (Table 1). The symmetric

EDP in	EDP in	EDA in	EDA m	Assignment	
colloid 1	colloid 2	colloid 1	colloid 2		
1598 mbr		1627 m $\delta NH_3^+$		δNH <sup>+</sup>	
1476 m				$\delta CH_2$	
1387 mbr		1364 m		t CH <sub>2</sub> , w CH <sub>2</sub>	
		1163 m		r NH <sup>+</sup> 3	
	1053 mbr			$v_{as}$ PO <sub>3</sub> ( <i>E</i> )	
	952 s			$v_{\zeta} \operatorname{PO}_{3}(A_{1})$	
		812 vs	810 m	v, AsO₄-	
675 m				Combination	
	595 sh			$\delta_{as} PO_3(E)$	
	545 sh			$\delta_{1} \operatorname{PO}_{3}(A_{1})$	
468 s	444 s	470 s	472 vs	δC-CN	
440 s			455 vs	$\delta \operatorname{AsO}_4^{3-}$	
234 s		221 m	m v Ag–N		
	217 m			v Ag-O	

<b>Table 1.</b> SENS Spectral data (chi - ) and band assignment	Table 1	I. SERS sp	ectral data	(cm <sup>-1</sup> ) and	band ass	ignments
---	---------	------------	-------------	-------------------------	----------	----------

v-very, s-strong, m-medium, br-broad, sh-shoulder, t-twisting, w-wagging, r-rocking

and asymmetric stretching and bending modes of  $PO_3$  are not observed in colloid 1. However, these are observed at 952, 1053, 545 and 595 cm<sup>-1</sup> in colloid 2. These observations suggest that the  $(NH_3)_2(CH_2)_2^{2+}$  and  $HPO_4^{2-}$  groups are close to the silver surface in colloid 1 and colloid 2 respectively. In otherwords, there are two adsorption sites in EDP which is supported by the low frequency bands at 234 and 214 cm<sup>-1</sup> in colloid 1 and colloid 2 respectively. Bands in the SERS spectrum recorded in freshly prepared colloid 1 (Figure 1) are not intense. It seems that aged colloid is a better substrate for surface enhancement.



In the case of EDA, bands due to both ethylene diammonium and arsenate groups are enhanced in colloid 1. In addition, a low frequency band at  $221 \text{ cm}^{-1}$  is observed (Figure 2). No splitting is observed for the stretching and bending modes of the arsenate ion indicating a higher symmetry for the ion when compared with the phosphate. In otherwords, the symmetry of the ion is probably not lowered due to adsorption or the ion may be existing as AsO<sub>4</sub><sup>3-</sup> rather than HAsO<sub>4</sub><sup>2-</sup>. In colloid 2, only bands due to the arsenate ion are enhanced

#### 466 Daizy Philip, Antony Eapen and G Aruldhas

(Figure 2 and Table 1). Though the arsenate bands are enhanced in both the colloids, low frequency band which corresponds to silver-molecule stretching mode is observed only in colloid 1. Therefore, it is reasonable to assume that the band observed at  $221 \text{ cm}^{-1}$  is due to v Ag-N mode arising from the nearness of the ethylene diammonium group to the silver surface. The non-observation of the low frequency v Ag-O band in both the colloids suggests that the enhancement of the arsenate bands is probably due to the electromagnetic field contribution.



SERS spectrum recorded in fresh colloid 1 (Figure 2) showed two low frequency bands at 194 and 106 cm<sup>-1</sup>. Further, only the arsenate bands are prominent in the spectrum.

As the colloid is aged only a medium intense band around  $221 \text{ cm}^{-1}$  is observed for this low frequency Ag-molecule stretching mode and both ethylene diammonium and arsenate bands are enhanced.

### Acknowledgment

Daizy Philip thanks the Council of Scientific and Industrial Research, New Delhi for a Research Associateship.

#### References

- [1] SJ Greaves and W P Griffith J Raman Spectrosc 19 503 (1988)
- [2] D Philip and G Aruldhas J. Raman Spectrosc 21 211 (1990)
- [3] J A Creighton, C G Blatchford and M G Albrecht J Chem. Soc Faraday Trans 2 75 790 (1979)
- [4] PC Lee and D Meisel J Phys Chem. 86 3391 (1982)