Infrared spectra of potassium citrate monohydrate single crystals

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The infrared spectra of potassium citrate monohydrate single crystals (001, 010 plates) have been recorded in the region 400-4000 cm⁻¹ using a Perkin-Elmer, infrared spectrophotometer, model 337 with a double grating dispersing unit. The free state frequencies of the citrate ion were taken from the reported works of Rao & Swamy (1970, 1971, 1972a, 1972b) and the observed frequencies in the present spectra were assigned to different modes of vibration of the citrate ion in the unit cell. The spectra were analysed according to the standard procedures of Halford (1946) and Hornig (1948). The splittings of the frequencies revealed that the citrate ion must have an approximate C_8 site symmetry in the unit cell. Thus sixty one out of the ninety six expected fundamental frequencies of the citrate ion in the potassium citrate monohydrate unit cell are identified in the spectra and assigned to their respective species. The hydrogen bond frequency range has been identified according to the details given by Hadzi (1965). The effect of orientation of the ions in the unit cell has been discussed with reference to the spectra of different crystal plates.

1. INTRODUCTION

The work done on the molecular structure of the citrate ion is very meagre and hence the authors have taken up this work. So far the authors have reported the tentative analysis of the infrared spectra of anhydrous citric acid (Rao & Swamy 1970), sodium citrate and ammonium citrate (Rao & Swamy 1971), lithium citrate tetrahydrate (Rao & Swamy 1972a), and potassium citrate monohydrate (Rao & Swamy 1972b). As a continuation the authors now have recorded the infrared spectra of potassium citrate monohydrate single crystals and an assignment of the observed frequencies is reported in this paper. Prior to the present work there is only one paper by Duval (1962) on the infrared spectrum of potassium citrate monohydrate which did not report any analysis. The common frequencies in the polycrystalline spectra of these compounds were taken as the free state frequencies of the citrate ion. Using this set of frequencies

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of the citrate ion, the analysis of the frequencies in the single crystal spectra of potassium citrate monohydrate has been carried out.

2. EXPERIMENTAL

Potassium eitrate monohydrate of B.P. grade was procured from Eagle Pharmaceuticals and purified by repeated recrystallization. Single crystals were grown from a saturated solution of the substance in distilled water, at room temperature. The crystal has a cleavage plane identified to be (001). Accordingly, (001) specimen plates were obtained by cleaving and grinding the cleaved plate over a slightly undersaturated aqueous solution of the salt. The (010) face is grown naturally which is identified from the morphological details given by Groth (1910). The axes identification was confirmed by taking X-ray oscillation rotation photographs of (010) and (001) plates. Then the infrared spectra of (001) and (010) plates were recorded which are reproduced in figures 1, 2 and 3. The observed frequencies with assignments are given in table 2 and the citrate ion frequencies for comparison are given in table 1.



Fig. 1. Infrared spectrum of potassium citrate monohydrate-single crystal (010) plate spectrum. 4000-400 cm⁻¹ region using PEIRS, Model 337-Double beam instrument Grating dispersing unit.



Fig. 2 Infrared spectrum of potassium citrate monohydrate-single crystal spectrum b-axis parallel to slit of the spectrophotometer—4000-400 cm⁻¹ region using PEIRS, model 337-Double beam instrument-Grating dispersing unit.

3. RESULTS AND DISCUSSION

It is known from Burns & Iball (1957) and Love *et al* (1960) that potassium eitrate monohydrate crystallizes in the monoclinic system with four molecules in its unit cell of dimensions a = 7.06A, b = 11.72A, c = 13.69Å, and $\beta = 112.0^{\circ}$ and has a space group C_{2A}^{5} . Full structural details are not available for this compound to know anything about the structure of the citrate ion in the unit cell. However, the structure of the citrate ion in magnesium citrate decahydrate crystal is known from the details given by Johnson (1965). According to this, citrate ion has only C_1 site symmetry in the crystal unit cell. Data on the bond lengths and bond angles of different bonds in the citrate ion from the details given by Johnson has been quoted in our earlier paper (Rao & Swamy 1972b). The C_1 point group symmetry of the citrate ion could be inferred from the spectroscopic observations as follows.



Fig. 3 Infrared spectrum of potassium citrate monohydrate-single crystal (001) plate spectrum—a-axis parallel to slit of the spectrophotometer-4000-400 cm⁻¹ region using PEIRS, model 337-Double beam instrument—Grating dispersing unit.



Fig. 4a Magnesium tricitrate unit cell-viewed along the b-axis.



Fig. 4b Hydrogen bonding and molecular-pack ing in the magnesium tricitrate unit cell viewed along the b-axis.



Fig. 5 Structure of the tricitrate ion $(C_6H_5O_7)^{-\frac{1}{1}}$

Table 1. Infrared frequencies of the citrate ion in the free state (taken from polycrystalline studies) in comparison to the frequencies of the citrate in the solid state from single crystal spectra

Polyorystalline		Frag	Single	Crystal	A			
Freq. cm ⁻¹	Int.	number	Freq. cm ⁻¹	Species	- Assignment			
2937	VS	٧ ₃	2940 2928	b _u a _u	3H ₂ asymm. stretch.			
2895	B	ν1	2900 2878	au bu	CH ₂ symm. stretch.			
1572	m.	٧ ₈	$\begin{array}{r} 1574 \\ 1566 \end{array}$	bu COO asymm. stretc au				
1549	m.	ν ₉	$\begin{array}{c} 1555\\ 1548 \end{array}$	$a_u \\ b_u$				
1532	m.	<i>v</i> 10	1541 b _u 1533 a _u					
1501	m.	<i>ν</i> 17	1502 a 1490 å	au bu	CH_2 bends.			
1480	m.	ν_{19}	1487 1477	$a_u \\ b_u$				
1465	m.	ν5	1472 1464	$a_u \\ b_u$	COO symm. stretches.			
1445	m.br.	νe	1455 1444	a _ո Ես				
1420	m.	¥7	1433 1420	$a_u \\ b_u$				
1406	m.sh.	V13	1413 1395	$a_u \\ b_u$	CH_2 bends.			
1340	m.br.	ν ₃₈	1346 1333	au bu	CH ₂ deformations.			
1285	m.br.	V36	1295	100 0000-0				
1182	w.	<i>v</i> ₁₁	1189 1183	au bu	C-OH stretch.			

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Assignment	Crystal	Single	E.o.o.	Polycrystalline		
	Freq. Species cm ⁻¹		- rroq numbor	Int.	Freq. cm ⁻¹	
C-OH oscillations.	bu au	1075 1069	V ₂₇	m	1071	
	b _u a _u	$1065 \\ 1057$	P28	w.	1065	
C-C stretches.	b _u	97 0	v_{12}	W	969	
	$b_u \\ a_u$	950 945	ν_{13}	w	950	
COO deformations	$a_u \\ b_u$	917 911	V21	я	914	
	au	900	ν_{22}	ŝ	900	
	$b_u a_u$	860 856	ν_{23}	m	856	
	$a_u \\ b_u$	849 842	V ₂₄	m	842	
CH ₂ rockings.		777 764	ν ₄₀	w	770	
		749 744	ν_{41}	vw	747	
		535 528	$ u_{2artheta}$			
C–C bendings.		523 511	$\nu_{3^{\circ}}$			
		505 495	ν ₃₁			
COO out of plane bendings.		468 462	V.17			
-		460 453	v ₃₈			
C-C torsions.		428 419	V42			
		420 411	Pau			

Table 1 contd.

From the available data on the infrared and Raman spectra of sodium citrate, the authors observed that many of the groups of frequencies are occuring in both the spectra. This is possible only when the citrate has C_{2v} , C_1 etc., point groups. Apart from this the observed infrared frequencies in the spectra of potassium citrate monohydrate single crystals could be satisfactorily explained only when we take C_1 point group for the free citrate ion. Thus we were lead by the above facts to the conclusion that C_1 must be the point group of the free

The citrate ion has eighteen atoms and must have forty eight frecitrate ion. The number of frequencies of different bond stretchings, bendings etc., auencies. were calculated according to the procedure of Decius (1949). The common infrared frequencies in the spectra of sodium citrate potassium citrate, ammonium citrate and lithium citrate tetrahydrate were noted. These frequencies should belong to the eitrate ion having C_1 point group symmetry. The frequencies are assigned to their respective groups using the tables for the charac-The frequencies toristic group frequencies. are numbered $\nu, \nu_{12}, \ldots, \nu_{42}$. In assigning the frequencies to ν_1 etc., there was some difficulty regarding assignment whenever two groups of frequencies are very near and this difficulty was overcome by observing the single crystal spectra of potassiam citrate monohydrate. Thus the free state citrate ion frequencies have been identified and are given in table 1. We have identified thirty one fundamental frequencies out of forty eight that are theoretically expected. Using this set of frequencies of the citrate ion, the infrared spectra of potassium citrate monohydrate single crystals were subjected to an analysis as given below.

From the available structural data on (Burns & Iball 1954) potassium citrate monohydrate and the symmetry of the free citrate ion, we can conclude that the site of symmetry of citrate ion in the potassium citrate monohydrate crystal must be C_1 . According to the details given by Johnson (1965) on the structure of the citrate ion in the unit cell of magnesium citrate decahydrate, all the atoms of the ion are arranged on two planes which are nearly perpendicular to each other. The fine structure and specially the splitting of the fundamental frequencies in the spectrum reveal the fact that the citrate ion may be considered to have an approximate site symmetry C_s *i.e.*, the citrate ion may be considered approximately to consists of two C_s groups (considering the atoms on each plane forming a C_s group and the two groups being perpendicular to each other the interaction will be negligible). The observed frequencies are assigned accordingly and discussed below. The correlation table between C_1 (the point group), C_d (site group) and C_{2b} (space group) is given in table 2.

According to the orientations of the citrate ions in the unit cell as reported by Johnson (1965) for magnesium tricitrate it is concluded that T_x , T_y , T_z , the dipole moment transition vectors, are parallel to c, a and b crystallographic axes respectively, and T_x , T_y are of b species and T_z is of a species. According to the selection rules a and b species are only permitted in the infrared. Therefore in the (010) plate spectrum it is expected that only b species should be present and in the (001) plate spectrum both a and b species should be present.

A free state fundamental like ν_1 will split into four components due to the presence of four molecules in the unit cell. According to the correlation table 2 only two frequencies of a and b species appear in the infrared, out of the four

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	pocies Assignment	O–H stretch water of crystallisation.	b. va.CH, asymm. stretch.	a. v. CH, symm. stretch.	pa realized and	OH stretches lowered due to hvdrogen bonding.		OH stretches lowered due to hydrogen bonding.	O-H-O bend. water of crystallization.	bu vs. v2, v10, COO asymm. bu stretches. bu	a. ν ₁₇ , ν ₁₈ , CH ₂ bends.
	Int. S	vvw vvw s.br	σ	c 8	8	1	M A	MA MA	nó nú	ک ە مە	a2
	Single crystal I.R. freq.cm ⁻¹ (001) plate a.axis	3675 3650 3400	0106	0000	2878		2248	1950 1880 1855	1648 1630	1575 1570 1550 1540	1520 1503 1497
	Species			ar	'nv					2 a a 3 a 3 a 3 a 3 a 3 a 4	a _u a _u
	Int.	VS VS	1	84	84	* *	M	л. Ж.	ø	30 07 07	ന ത ത
	Single crystal I.R. freq.cm ⁻¹ (001) plate b-axis	3425 3300		2928	2900	2380 2350 2322	2299 2275 2248	$1952 \\ 1860 \\ 1852$	1651 1648 1638	1566 1555 1533	1502 1499 1487
	Species			b_u	p"				solved	111	
	Int.	vrw w.br w.br br.ab	ł	ø	at:	88	8	\$	not re		
	Single crystal I.R. freq.cm ⁻¹ (010) plate	3779 3687 3672 3390	1	2940	2878	2360 2325	2293 2275 2250	1950 1908 1848	1650 1335		
o moorde	Int.	85	vs.br	67	ø	a a	8	8A NB	8A 8A	60	80
•	I.R. freq.cm ⁻¹ KBr pellet spectrum due to authors	3663 3560	3125	2937	2895	2370 2350	2295	1765 1760	1/29 1645 1630	1572 1549 1530	1501 1498 1480

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	Assignment	¹ 8, ¹ 8, ¹ 7, COO symm. stretches.	ν_{19} . CH ₂ bend.	¹²³ CH2 deformations. 136	^{ν11} .C-OH stretch.	¹ 27 C-OH oscillations. 1 ²⁸	V12 C-C stretches.	P21	¹ 22 COO deformations. ¹ 23 124	vio CH2 rocking.	²²⁸ C-C bends.	 V3T COO out of plane bends. V3B 	v43 C-C torsion.
	Species	Ф.Ф. а В. и В. и В. и В. и В. и В. и В. и В. и	* 0	1	ł		b_u^{u}	ł	a ⁿ	; 		11	I
	Int.	80	ø		w.br.	m.br.	мл	w.br.	₩	w.br	8 8	8 B	мл
td	Single crystal I.R. freq.cm ⁻¹ (001) plate c.axis	1473 1467 1467 1450 1442 1425	1415	1294	1188 1182	1070 1060	970 950	915	006	775	529 522	460 455 450	420
ole 2 cont	Species	3 3 3 8 8 8	Gu	a.	au	рк С К	bu Gu	n n n	а а а 9 9 9 6			I	
Tal	Int.	as as as	æ	æ	м	m.br.	a v	*	w w.br	w.br w	***	A	₿
	Single crystal I.R. Freq.cm-1 (001) plate b-axis	1472 1455 1433	1413	1346	1189	1072 1058	970 947	916	900 857 849	768 743	528 519 498	458	419
	Species		I		<i>b</i> "	ь. Ъ.	ъ. С.		b	11	I		1
	Int.		I	s.br.	* *	* *	w w w	MAA	w vw.br	₿	8	8	1
	Single crystal I.R. freq.cm ⁻¹ (010) plate		I	1325 1280	1186 1183	1075 1065	970 950	917	900 860 	748	520	464	1
	Lat.	80	ati	m.br.ab. region	▶	ww WVW	* *	8	E	***	m.br. m.br.	≱	₿
	I.R. freq.cm ⁻¹ KBr Pellet Spectrum due to authors	1465 1451 1420	1406	1348 1280	1182	1071 1059	969 950	914	900 856 842	770 747	530 510	454	411

cont
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Table

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split frequencies mentioned above. Of these two, the a component could be identified first since it should be present in (001) plate spectrum and should be absent in (010) plate. When once the a component was identified the remaining split component should be the b component. Since we have recorded the spectra of (001) plate with a and b axis parallel to the slit of the spectrophotometer, the assignments of the species have been done with ease. But in the 1300-1600 cm⁻¹ region none of the single crystal spectra are resolved very well and so in this region assignments have been offected with the help of the KBr pellet spectrum. In the case of the spectrum of an unoriented specimen the symmetric species will usually appear with high intensity compared to the antisymmetric species of the same group or bond. This principle was used while making the assignments with the KBr pellet spectrum. It was not possible to assign the split components of some of the frequencies like v_{40} , v_{41} etc., to their species because of the fact, that the b-axis spectrum is taken with random orienta-Thus the split components of the available fundamentals of the citrate tion. ion are assigned and the assigned frequencies are discussed below :

CH₂ Frequencies :

The stretchings of this group appear in the region 2937-2895 cm⁻¹; the bendings at 1501, 1480, 1406 cm⁻¹ have split up as expected and their species are assigned. Four CH₂ bending frequencies (free state) are expected but we could identify only three in the KBr spectrum and this region is not resolved well in the single crystal spectra. The deformations appear at 1340, 1285 cm⁻¹ and the rocking frequencies appear at 770, 747 cm⁻¹. The CH₂ group according to the crystallographic details on magnesium citrate, is known to be situated out of the back-bone plane and the frequencies pertaining to this group must appear in the spectra of both the C_8 groups, as was observed.

COO Group Frequencies :

Stretchings of this group appear at 1572. 1549 and 1532 cm⁻¹, 1465 1445 and 1420 cm⁻¹ and the deformations appear at 914, 900, 856, 842 cm⁻¹. The out of plane bendings appear at 453, 460 cm⁻¹. From the crystallographic details it is known that there are two COO groups in (010) plane which is the back-bone plane and the other COO group is in (001) plane which is perpendicular to the back-bone plane. These two planes represent two C_s groups. This group frequencies clearly proved that the point group C_1 consists of two C_s groups. The stretching region is not well resolved but the other regions are in conformity with a C_s site symmetry.

C-OH Frequencies :

The frequencies at 1182, 969, 1071, 1065 cm⁻¹ are attributed to the vibrations of this group. The 1182 cm⁻¹ is identified to be the stretching frequency of this group. The other frequencies are due to the bending modes of this group. According to the structural details, this group vibrations should appear only in (001) plate spectrum, since this group is in the (001) plane. But C-OH group frequencies may also appear in (010) plate spectrum, since the O-H-O type of bonds attached to some C-atom also will have C-OH group vibrations.

C-C Frequencies :

The frequencies of this group appear at 950, 530, 523, 508 cm⁻¹ and the split components are identified and duly assigned.

C-C Torsional Vibrations :

These are due to torsional vibrations of the molecular frame and identified at 420 and 411 cm⁻¹.

Hydrogen Bond Frequencies :

The frequency regions of this group were reported by us in the previous paper on potassium citrate at 2370, 1760, 1645 cm⁻¹. The O-H-O frequencies of the polycrystalline spectrum have split up into a number of components in the present single crystal spectra. Due to the lack of full details regarding the number and arrangement of hydrogen bonds, we were unable to derive maximum useful information from the spectra and could only identify the regions.

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