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# Experimental and Theoretical Study of the Vibrational Spectra of 12-Crown-4-Alkali Metal Cation Complexes

S. Al-Rusaese, A. A. Al-Kahtani, and A. A. El-Azhary\*, and A. A. El-Azhary\*,

Chemistry Department, Faculty of Science, King Saud University, P.O. Box 2455, Riyadh 11451, Kingdom of Saudi Arabia, and Chemistry Department, Faculty of Science, Cairo University, Giza, Egypt

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The vibrational, Raman, and IR, spectra of the five 12-crown-4 (12c4) complexes with Li<sup>+</sup>, Na<sup>+</sup>, K<sup>+</sup>, Rb<sup>+</sup>, and Cs<sup>+</sup> alkali metal cations were measured. Except for a small shift of the position of some bands in the vibrational spectra of the Li<sup>+</sup> complex, the vibrational spectra of the five complexes are so similar that it is concluded that the five complexes exist in the same conformation. B3LYP/6-31+G\* force fields were calculated for six of the eight predicted conformations in a previous report (J. Phys. Chem. A 2005, 109, 8041) of the  $12c4-Li^+$ , Na<sup>+</sup>, and K<sup>+</sup> complexes that are of symmetries higher than the  $C_1$  symmetry. These six conformations, in energy order, are of  $C_4$ ,  $C_s$ ,  $C_s$ ,  $C_{2\nu}$ ,  $C_{2\nu}$ , and  $C_s$  symmetries. Comparison between the experimental and calculated vibrational frequencies assuming any of the above-mentioned six conformations shows that the five complexes exist in the  $C_4$  conformation. This agrees with the fact that the five alkali metal cations are larger than the 12c4 ring cavity. The B3LYP/6-31+G\* force fields of the C4 conformation of the Li<sup>+</sup>, Na<sup>+</sup> and K<sup>+</sup> complexes were scaled using a set of eight scale factors and the scale factors were varied so as to minimize the difference between the calculated and experimental vibrational frequencies. The root-mean-square (rms) deviations of the calculated frequencies from the experimental frequencies were 7.7, 5.6, and 5.1 cm<sup>-1</sup> for the Li<sup>+</sup>, Na<sup>+</sup>, and K<sup>+</sup> complexes, respectively. To account for the earlier results of the Li<sup>+</sup> complex that the  $C_s$  conformation is more stable than the  $C_4$  conformation by 0.16 kcal/mol at the MP2/ 6-31+G\* level, optimized geometries of the complex were calculated for the  $C_4$  and  $C_s$  conformations at the MP2/6-311++G\*\* level. The  $C_4$  conformation was calculated to be more stable than the  $C_s$  conformation by 0.13 kcal/mol.

#### Introduction

Recently there has been a wide interest shown in the chemistry of crown ethers. This interest is due to the ability of crown ethers to form strong complexes with ionic species, especially alkali metal cations, with high selectivity. Consequently, crown ethers have found a large number of industrial and medical applications. Crown ethers are used in nuclear waste disposal,<sup>1–5</sup> membrane transport,<sup>6–8</sup> anion activation,<sup>9,10</sup> formation of soluble and insoluble polymers,<sup>11–14</sup> and macrocyclic liquid crystals.<sup>15</sup>

Because of the importance of the applications of crown ethers, our objective was set to study this class of molecules to determine in which conformation, or conformations, these large ring flexible molecules exist in. The methodology we use to achieve this goal is through the conformational analysis and vibrational study of free crown ethers and some of their complexes. Although our attention was attracted first to the most important crown ether, 18-crown-6 (18c6), due to its large size, our effort was turned to the smaller and easier to study 12c4.

In a previous report, <sup>16</sup> a full conformational search of the possible conformations of free 12c4 was performed at the MM3 level. The search was performed using an efficient method of conformational search of cyclic molecules, the CONFLEX method. <sup>17–20</sup> The method, as implemented in the CAChe

program,<sup>21</sup> has the advantage of being fully programmed. This minimizes the human interference and consequently minimizes human error. This is compared to other conformational search methods that can be termed as half programmed-half manual. The conformational search of 12c4 resulted in the prediction of 180 conformations. Energy order of the predicted conformations was calculated at levels of theory as high as the MP2/6-31+G\* level. Similar to previous reports, 22-24 the study predicted that the  $S_4$  conformation is the lowest energy conformation of 12c4. At the MP2/6-31+G\* level, the  $S_4$ conformation is more stable than the  $C_i$  conformation by 2.61 kcal/mol. In a following vibrational study of free, or rather uncomplexed, 12c4, it was shown for the first time that 12c4 in the solid, liquid, and considered solution phases exists in the  $C_i$  conformation.<sup>25</sup> The case is similar for the larger 18c6. It is known experimentally, using the vibrational spectra<sup>26,27</sup> and X-ray,  $^{28,29}$  that 18c6 exists in the  $C_i$  conformation. A similar conformational search, as that performed for 12c4, predicted for the first time that the lowest energy conformation of free 18c6 is a  $S_6$  conformation. At the MP2/6-31+G\* level, the  $S_6$ conformation is more stable than the experimentally known  $C_i$ conformation by 1.84 kcal/mol.<sup>30</sup> It was shown that the stability of the lowest energy  $S_4$  conformation of 12c4 and the lowest energy  $S_6$  conformation of 18c6 is due to a combination of both more number of oxygen atoms participating in hydrogen bonding and at distances shorter than that of any of the other conformations. 16,30

In a continuing study, conformational analysis of 12c4—alkali metal cation complexes was performed.<sup>31</sup> The study predicted eight possible conformations of these complexes. It was

<sup>\*</sup> Corresponding author. Present address: Chemistry Department, Faculty of Science, King Saud University, P.O. Box 2455, Riyadh 11451, Kingdom of Saudi Arabia. E-mail:azhary60@hotmail.com. Telephone: (9661) 467 4367. Fax: (9661) 467 5992.

King Saud University.

<sup>‡</sup> Cairo University.

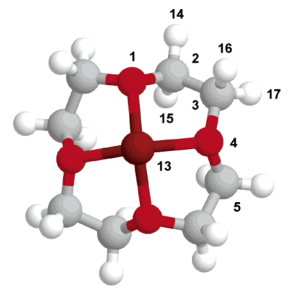
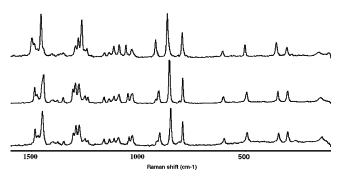


Figure 1. Structure and atom numbering of the  $C_4$  conformation of the 12c4-alkali metal cation complexes.



**Figure 2.** Raman spectra of the solid phase of the three 12c4-Li<sup>+</sup> (top), Na<sup>+</sup> (middle) and K<sup>+</sup> (bottom) complexes.

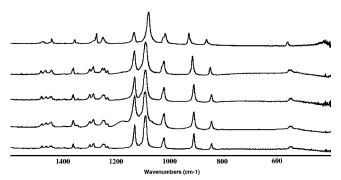


Figure 3. IR spectra of the solid phase of the 12c4-Li<sup>+</sup> (top), Na<sup>+</sup> (second from the top), K<sup>+</sup> (third from the top), Rb<sup>+</sup> (fourth from the top) and Cs+ (bottom) complexes.

concluded that the  $C_4$  conformation is the lowest energy conformation of the 12c4-Na<sup>+</sup>, K<sup>+</sup>, Rb<sup>+</sup> and Cs<sup>+</sup> complexes. For the  $12c4-Li^+$  complex, a  $C_s$  conformation is the lowest energy conformation, although, at the MP2/6-31+G\* level, the  $C_4$  conformation is less stable than this  $C_s$  conformation by only 0.16 kcal/mol. The best agreement between the experimental and calculated binding energies is obtained assuming the  $C_4$  or C<sub>s</sub> conformation of the 12c4-Li<sup>+</sup> and Na<sup>+</sup> complexes and the  $C_4$  conformation of the 12c4-K<sup>+</sup> complex. For the 12c4-Rb<sup>+</sup> and  $-Cs^+$  complexes, poor agreement is obtained between the experimental and calculated binding energies assuming  $C_4$ conformations of both complexes. Instead, the best agreement is obtained assuming a  $C_{2v}$  structure of both complexes. Since this disagrees with the energy order mentioned above, Feller

TABLE 1: Geometry of the 12c4-Alkali Metal Cation Complexes at the B3LYP/6-31+G\* and MP2/6-31+G\* Levelsa

Levels"					
coordinate	Li <sup>+</sup>	Na <sup>+</sup>	$K^+$	$Rb^+$	$Cs^+$
$O_1 - C_2$	1.437	1.434	1.431	1.429	1.428
	1.440	1.437	1.433	1.432	1.431
$C_2-C_3$	1.524	1.525	1.524	1.525	1.525
	1.516	1.517	1.516	1.516	1.516
$C_1-O_4$	1.440	1.437	1.433	1.431	1.430
	1.444	1.441	1.437	1.435	1.434
M-O	2.120	2.334	2.747	2.986	3.226
	2.018	2.394	2.708	2.930	3.144
$C_2-H_{14}$	1.098	1.099	1.100	1.100	1.101
	1.097	1.099	1.099	1.100	1.100
$C_2-H_{15}$	1.099	1.098	1.098	1.098	1.098
2 .5	1.099	1.097	1.097	1.097	1.097
$C_3-H_{16}$	1.093	1.094	1.095	1.095	1.095
	1.093	1.093	1.094	1.094	1.094
$C_3 - H_{17}$	1.096	1.097	1.098	1.099	1.099
- 5 17	1.096	1.098	1.098	1.099	1.099
$O_1C_2C_3$	106.0	107.8	108.5	108.8	108.9
- 1 - 2 - 9	105.3	107.1	107.3	107.5	107.5
$C_2C_3O_4$	109.0	111.8	112.6	112.9	113.1
2 2 .	109.4	111.2	111.5	111.8	112.0
$C_3O_4C_5$	116.7	115.9	115.4	115.3	115.3
	115.2	114.4	114.0	113.9	113.9
$MO_4C_3$	109.4	109.2	109.9	110.2	110.4
	109.5	110.0	110.8	111.0	111.2
$MO_4C_5$	109.1	113.2	118.6	120.8	122.7
	109.2	115.1	119.7	121.9	123.7
$O_1C_2H_{14}$	110.2	110.0	110.0	110.0	110.1
	110.1	109.9	110.1	110.2	110.2
$C_3C_2H_{14}$	110.5	109.7	109.4	109.2	109.2
	111.0	110.2	110.0	109.8	109.8
$O_1C_2H_{15}$	110.2	110.3	110.4	110.4	110.5
	110.2	110.3	110.4	110.5	110.5
$O_3C_2H_{15}$	111.4	111.0	110.7	110.6	110.5
	111.5	111.1	110.9	110.8	110.8
$C_2C_3H_{16}$	110.3	110.1	109.9	109.9	109.8
	110.3	110.1	110.0	109.8	109.8
$O_4C_3H_{16}$	105.7	105.6	105.7	105.7	105.7
	105.6	105.6	105.7	105.7	105.7
$C_2C_3H_{17}$	111.1	110.2	109.8	109.6	109.5
	111.6	110.7	110.5	110.3	110.3
$O_4C_3H_{17}$	111.0	110.7	110.7	110.7	110.8
	110.7	110.5	110.7	110.7	110.7
$O_1C_2C_3O_4$	-50.9	-56.1	-58.4	-58.9	-59.4
	-52.8	-56.9	-57.9	-58.5	-58.9
$C_2C_3O_4C_5$	-91.5	-84.0	-81.0	-80.2	-79.7
	-88.8	-83.0	-81.6	-80.8	-80.4
$C_3O_4C_5C_6$	168.1	164.7	163.5	162.9	162.7
	168.7	165.6	165.0	164.6	164.4
$MO_1C_2C_3$	43.5	37.5	29.9	26.3	23.2
	45.1	36.7	30.5	27.1	24.0

<sup>a</sup> See Figure 1 for atom numbering. Bond lengths in Å and angles in degrees. For each coordinate, the first line corresponds to the B3LYP/ 6-31+G\* level and the second line corresponds to the MP2/6-31+G\* level. M refers to alkali metal cation.

and co-workers argued that spectral measurement of the binding energy of the 12c4-Rb+ and -Cs+ complexes samples both complexes in this high energy  $C_{2\nu}$  conformation rather than the lowest energy  $C_4$  conformation.  $^{32-37}$ 

Surprisingly, as the number of studies of the vibrational spectra of free 12c4 is quite limited, <sup>38–44</sup> the number of studies of the vibrational spectra of its alkali metal cation complexes, to the best of our knowledge, is very scarce. 38,40 The most detailed study of the vibrational spectra of 12c4-alkali metal cation complexes was reported by Fukushima and Tamaki in 1987.40 The authors reported the Raman spectra of the 12c4- $Li^+$ ,  $-Na^+$ ,  $-K^+$ ,  $-N\hat{H}_4^+$ ,  $-Mg^{2+}$ ,  $-Ca^{2+}$ ,  $-Sr^{2+}$ ,  $-Ba^{2+}$ , and −Pb<sup>2+</sup> complexes in the region below 1000 cm<sup>-1</sup>. The assignment of the fundamental vibrations was aided by frequencies calculated using an empirical force field. The study predicted a  $D_{2d}$  structure of the 12c4-Li<sup>+</sup>, -Na<sup>+</sup>, -K<sup>+</sup>, -NH<sub>4</sub><sup>+</sup>, and  $-\mathrm{Mg}^{2+}$  complexes and a  $C_{2\nu}$  structure of the 12c4 $-\mathrm{Ca}^{2+}$ ,  $-Sr^{2+}$ , and  $-Ba^{2+}$  complexes. This result contradicts an earlier

TABLE 2: FT-Raman and FT-IR Frequencies  $(cm^{-1})$  and Assignment of the  $12c4-Li^+$  Complex<sup>a</sup>

	Ran	nan		IR		
	$MeOH^b$		$H_2O^c$	$\overline{\text{MeOH}^b}$		
solid	MeOH	H <sub>2</sub> O	$H_2O$	solid	ref 40	assgnt
155	158					(v <sub>39</sub> )
167	167	164				$(v_{61})$
176	182	185				
	199	201				$(v_{20})$
229	214	221				$(v_{38})$
250		221			252	(038)
266					262	
282	285		292			$v_{60}$
305	305	300	307		302	$v_{19}$
224	311	313	310		312	
324	328 341	338	324 340			$v_{18}$
355	351	352	348		355	$v_{59} = v_{17}$
		374	374			- 17
	400	401	399			$(v_{37})$
	415	415		422		$v_{58}$
501	500	499	499	5.00	504	$v_{36}$
				562	579	$v_{57}$
603	603		604	607	319	$v_{16}$
795	786	788	785	007	792	$v_{35}$
	802	803	800		805	$v_{56}$
815	816	816	818		818	
0.4	0.48	0.48	0.40	0.4	848	
865	862	862	860	865	865	$v_{15}$
912 920	907 917	908 916	900 917		902	$v_{14}$
929	928	925	926	931		$v_{34} = v_{55}$
		944	939	, , ,		- 33
1024		1022	1022	1020		$v_{13}$
1033		1029	1029	1027		$v_{54}$
1048		1049	1049	1046		
1059 1078	1084	1059 1085	1060 1085	1083		
1078	1090	1083	1083	1065		
1070	1102	1102	1102			
	1113	1113	1111	1111		$v_{11}$
1116	1120			1117		$v_{32}$
1120	1128	1120	1126	1107		
1138 1159	1137 1161	1138 1161	1137 1158	1137		$v_{52}$
1242	1241	1244	1243	1246		$v_{31} = v_{10}$
1253	1252	1211	1254	1253		V 10
1266	1261	1260	1262			$v_{30}$
		1269	1269			
1281	1277	1286	1280	1278		
1296	1288 1296	1294 1296	1288 1298	1289		
1349	1350	1352	1350	1348		$v_8$
1367	1366	1361	1361	1360		$v_{49}$
1376	1381	1369	1372			- 47
1397	1392	1396	1388			
1405	1400		1400			
1410	1411	1307	1407	1.4.4.6		
1446 1457	1442 1454	1442	1455	1446		$v_{47}$
1437	1434	1454 1464	1433	d		$v_{26}$
1472	1479	1475	1473	d		$v_{25}$
1487	1486		1488	d		$v_{46}$
1499				d		$v_5$
2876		2875	<b>a</b> 00 -	2875		
2886		2885	2885	2882		
2902		2010		2020		
2923 2942		2919 2945		2920		
2970		2943	2974	2966		
2991		-21.		2990		
3002		3003	3001			
<sup>a</sup> Me0	OH stands	for metl	hanol. as	sgnt is the	fundamenta	ıl band
		1 .	.1 .			. 1

 $^a$  MeOH stands for methanol. assgnt is the fundamental band assignment. For bands in parentheses, their assignment as fundamentals was not certain, and they were not used in the scaling of the force field.  $^b$  The 12c4—alkali metal cation complex synthesized using methanol as a solvent.  $^c$  The 12c4—alkali metal cation complex synthesized using H<sub>2</sub>O as a solvent.  $^d$  There is one broad band in this region.

IR and NMR study of free 12c4 and its  $Li^+$  complex that both species adopt a square  $C_4$  conformation.<sup>38</sup>

TABLE 3: FT-Raman and FT-IR Frequencies (cm $^{-1}$ ) and Assignment of  $12c4-Na^+$  Complex<sup>a</sup>

ASSI	nment	01 12	C4-N	a Col	прієх"				
		Raman	l			IR			
	$MeOH^b$		H	$I_2O^c$	$MeOH^b$		$_{2}\mathrm{O}^{c}$		
solid	MeOH	H <sub>2</sub> O	solid	MeOH	solid	solid	MeOH	ref 40	assgnt
150	131	138	129	124					$(v_{40})$
153 171	148 171		148 173						$v_{39}$
183	181		180	188					$v_{61}$
193	196	193	198	***					$v_{20}$
209 215	204 215	202	210	204 212					$v_{38}$
213	248	240	246	243					$v_{60} = v_{19}$
254		257	260						• • • • • • • • • • • • • • • • • • • •
260 280	285	264	268 276	273					**
304	302	303	302	301				303	$v_{59} = v_{18}$
347	348	350	347	346				349	$v_{17}$
355 366	368	363		352 369					21
421	414	416	415	309					$v_{58} = (v_{37})$
494	494	496	494	493				495	$v_{36}$
602	601	598	603	599	553	553		601	$v_{57}$
796	794	795	792	794				796	$v_{16} = v_{35}$
	802	803	808		808				$v_{56}$
815	814	815	810 825	815	922	825	825		
821 833	827	829	833	823 830	823 830	831	830	828	
855	856	853	855	855	852	853	852	858	$v_{15}$
879	881	875		875	900	907			
896 903	895 906	896 904	905	898 905	899	897		907	$v_{14} = v_{34}$
918	920	916	919	918	918	918	919		$v_{55}$
1020		1028	1028		1025	1026			$v_{13}$
1029 1049		1033 1041	1031 1049		1032	1030			$v_{54} = v_{33}$
					1055				- 55
1068 1093	1098	1064 1095	1072 1092	1097	1093	1093	1098		$v_{12}$
1093	1106	1111	1092	1103	1093	1093	1098		$v_{53}$
1115	1115	1115	1115	1110	1113	1113			$v_{11}$
1126 1140	1141	1124 1140	1124	1123 1139	1125	1125	1120		$v_{32}$
1162	1160	1165	1138 1162	1158	1135	1135	1138		$v_{52} = v_{31}$
1238	1236	1239	1237	1241	1236	1237	1237		$v_{10}$
1251	1249	1251	1250	1254	1248 1256	1248 1256	1248 1255		$v_{51}$
1273	1268	1269	1279	1266	1230	1230	1233		$v_{30} = v_{29}$
					1289	1289	1290		- 27
1293 1307	1291	1293 1307	1296 1307	1291	1293	1293	1293		$v_{50}$
1307	1305	1307	1307	1306	1304	1304 1348	1305		$v_9$
1351	1354	1353	1353	1353		1355			$v_8$
	1367	1363	1264	1366	1265	1362	1365		
	1307	1303	1304	1300	1365 1368	1365 1367	1303		$v_{49}$
1371		1371	1371		1375	1375			
1380	1383	1386	1380	1379	1380 1387	1379 1387			$v_{28}$
1396	1396	1396	1396		1395	1395			$(v_7)$
1400			1402	1402	1399	1399			$(v_{27})$
1408 1446	1410	1410	1408 1444	1413	1405	1405			$v_{48}$
1440			1		1447	1447			$v_{26} = v_{47}$
		1453	1452						$v_6$
1473			1470 1476		1469	1469			$v_{25}$
1493		1486	1470		1485	1485			$v_{46}$ $v_{5}$
					2964	2965			-
2871		2881	2870 2887		2875 2878	2875			
		2001	2894		2891				
****		2907	2910		2907	2910			
2922 2930		2933	2925 2943		2922 2936	2921 2936			
2950		2953	2943		2957	2957			
					2966	2969			
			2975		2975				

 $<sup>^{</sup>a,b,c}$  See corresponding footnotes in Table 2.

There are two reports of X-ray studies of the structure of  $12c4-Na^+$  complex. A5,46 Both studies show that the complex in the solid phase has a  $C_4$  structure. Using X-ray,  $C_s$ ,47  $C_b$ 48

and  $C_4^{49}$  conformations have been reported for the 12c4-Cu<sup>2+</sup>,  $-Mg^{2+}$ , and  $-Ca^{2+}$  complexes, respectively.

In view of these conflicting results about the conformation, or conformations, assumed by 12c4-alkali metal cation complexes and the absence of a detailed vibrational study of these complexes, it was felt that such a study would be necessary. This would be also in line with our previous studies of the conformational and vibrational analysis of crown ethers. The aim of the present report is then to measure the vibrational, Raman, and IR spectra of the five 12c4-alkali metal cation complexes. In addition, to compare between the experimental and calculated vibrational spectra of these complexes assuming any of the possible conformations of the complexes. This is in an effort to determine in which conformation these complexes exist in. Finally, aided by accurately calculated scale quantum mechanical (SOM) B3LYP force field, the fundamental vibrational frequencies of the complex are assigned.

### **Experimental Details**

12c4-alkali metal cation complexes were synthesized by heating for 2 h, under reflex in methanol or in aqueous solution, a slight excess of an equimolar amount of the alkali metal chloride salts and 12c4.26 The solvent was allowed to evaporate by placing the mixture in a closed desiccator under vacuum whereby crystals of the complex were formed. Some of the complexes were obtained in an emulsified form that did not form a precipitate. To these complexes, ether was added. Upon evaporation of the ether, crystals of the complexes were obtained.

FT-IR spectra were measured using a Thermo Nicolet Nexus 870 FT-IR instrument. The instrument uses a KBr beamsplitter and an InGaAs detector. Typically, between 128 or 256 scans were collected with a resolution of 1 cm<sup>-1</sup>. Solid samples were measured as KBr pellets. Solution samples, in methanol, were measured using a fixed path length cell equipped with KBr windows. The path length of the cell was varied using Teflon spacers of different thicknesses.

The FT-Raman spectra were measured using a Thermo Nicolet Nexus FT-Raman spectrometer. The instrument uses an air-cooled Nd:YVO4 laser source which emits continuouswave laser energy at a wavelength of 1064 nm, a XT-KBr beamsplitter, a 180° sample configuration and an InGaAs detector. Between 1024 and 4096 scans at a resolution of 4 cm<sup>-1</sup> were collected. To maximize the signal-to-noise ratio and due to the weak scattered light from the complex samples, a laser power of up to 1.0 W was used.

### **Computational Details**

Conformational analysis of the five 12c4—alkali metal cation complexes, to predict the possible conformations of the complexes, was reported in a previous publication.<sup>31</sup> Cartesian coordinate force fields were calculated, at the corresponding optimized geometries, for six of the eight predicted conformations. The other two conformations have  $C_1$  symmetry and were not considered. The six conformations considered have symmetries of, in energy order,  $C_4$ ,  $C_s$ ,  $C_s$ ,  $C_{2v}$ ,  $C_{2v}$ , and  $C_s$ , as detailed in ref 31. Notice that for the Li<sup>+</sup> complex in all levels considered in ref 31, except at the HF/3-21G level, the energy order of the first  $C_4$  and second  $C_s$  conformations is reversed and the third  $C_s$  conformation collapses to the second  $C_s$ conformation. The force fields were calculated at the B3LYP level due to its known excellent accuracy to CPU time ratio.50 As was detailed in ref 31, the 6-31+G\* basis set was used for all atoms except for the K, Rb, and Cs atoms. 31,51-56 For these,

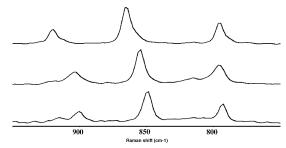


Figure 4. Raman spectra of the solid phase of the three 12c4-Li<sup>+</sup>-(top), Na<sup>+</sup> (middle) and K<sup>+</sup> (bottom) complexes in the 950–750 cm<sup>-1</sup> region. The Figure shows a shift of the position of some bands of the 12c4-Li+ complex compared to the corresponding bands of the Na+ and K+ complexes.

the Hay and Wadt's 10-valence-electron effective core potential (ECP) with a (5s4p)/[3s2p] valence basis set was used.<sup>57</sup> The basis set has an additional six-term d-type polarization functions with exponents of  $\alpha_d = 0.48$  for K,  $\alpha_d = 0.24$  for Rb, and  $\alpha_d$ = 0.19 for Cs. For simplicity, this basis set will be termed the 6-31+G\* basis set. The additional diffuse function in the 6-31+G\* basis set was used to minimize the basis set superposition error and for its known necessity for accurate computation of the properties of the metal cation complexes. The basis set used for the K, Rb, and Cs atoms is the same as that used by Feller el al. for similar computations of 12c4and 18c6-alkali metal cation complexes. 33,37,38,61

All ab initio computations were performed using the Gaussian 98W<sup>62</sup> and Gaussian 03W<sup>63</sup> programs. The Gaussian program default parameters of geometry optimization and force field calculations were used. The force fields and dipole derivative tensors were calculated analytically. Since analytical polarizability derivatives are not available in the current version of the Gaussian program, these were calculated numerically.

The selection of the internal coordinates, 64-66 conversion of the Cartesian coordinate force fields to internal coordinate force fields, 65,67 scaling of the internal coordinate force fields 65,68,69 and refinement of the scale factors were done using the SCALE2 program<sup>70</sup> and were performed as was detailed elsewhere.<sup>25</sup> They are mentioned here for completeness. The SCALE2 program has the advantage of minimizing human interference and consequent error. In the first step of the program, from the frequency job archive of the Gaussian program output file, the program generates four files of the molecule Cartesian coordinates, Cartesian coordinate force field, dipole derivative tensors and polarizability derivatives. In the second step, the program generates the internal coordinates that are used to convert the Cartesian coordinate force field, dipole derivative tensors and polarizability derivatives to their internal coordinate counterparts. In the third step of the program, the internal coordinate force field is scaled according to the equation

$$F_{ij}^{\text{ scaled}} = F_{ij}^{\text{ theo}} (c_i c_j)^{1/2}$$

where  $c_i$  and  $c_j$  are the scale factors of the internal coordinates i and j, respectively. The reproducibility of the frequencies calculated by the Gaussian program is checked using a scale factor of 1.0. To aid in the initial assignment of the fundamental vibrations from the experimental spectra, the internal coordinate force fields are scaled with an initial scale factor of 0.963.71-73 The experimental fundamental vibrations were then assigned to the calculated frequencies in frequency order taking into consideration the IR and Raman intensities. As the number of assigned bands increased, the number of scale factors was increased and the scale factors were varied, to minimize the

TABLE 4: FT-Raman and FT-IR Frequencies  $(cm^{-1})$  and Assignment of  $12c4-K^+$  Complex<sup>a</sup>

	Raman		II		
Me	$eOH^b$	$H_2O^c$	MeG	$OH^b$	
solid	MeOH	solid	MeOH	solid	assgnt
140	136	141			$v_{39}$
191	189	188			$v_{60}$
208	202 217	212			$(v_{38})$
231	239	234			$v_{19}$
269	264	265			$v_{59}$
301	304	301			$v_{18}$
343	349	344			$v_{17}$
356	359	360			$v_{58}$
375 397	376 398	376			210=
429	425	413		419	$v_{37}$
492	497	492		417	$v_{36}$
				549	$v_{57}$
599	597	600			$v_{16}$
793	795	793		791	$v_{35}$
808	804	810		808	$v_{56}$
814 849	816 848	849	848	846	41
900	902	901	040	040	$v_{15} = v_{34}$
915	910	914	916	913	$v_{55}$
				1025	$v_{13}$
1032		1030		1031	$v_{54}$
1045		1045			$v_{33}$
1059	1007	1055		1004	$v_{12}$
1097 1116	1097 1112	1094 1115		1094 1111	<i>v</i> <sub>53</sub>
1116	1112	1113		1111	$v_{11} = v_{52}$
1162	1163	1162		1151	$v_{31}$
1238	1243	1237		1237	$v_{10}$
			1250	1248	$v_{51}$
1253	1256	1250			$v_{29}$
1067		1050		1254	$v_{30}$
1267 1278	1277	1258 1278		1259 1278	
1278	1277	1278	1291	1278	$v_{50}$
1304	1304	1305	1306	1303	$v_9$
1349	1353	1350		1348	$v_8$
1368	1368	1371		1365	$v_{49}$
1379	1377	1379		1378	$v_{28}$
1392	1386	1391		1388	$v_7$
1401 1410	1400 1407	1401 1410		1399 1405	$v_{48}$
1410	1407	1410		1449	$v_{27} \ v_{47}$
1450		1450		1117	$v_{26}$
				1467	$v_{46}$
1471		1471		1472	$v_6$
1485		1485		1483	$v_5$
2871		2871		2865	
2886		2886		2882 2908	
2906 2929		2906 2928		2908 2926	
2938		2939		2934	
				2949	
				2956	

*a,b,c* See corresponding footnotes in Table 2.

difference between the calculated and experimental vibrational frequencies, as indicated by the rms deviation of the difference between the calculated and assigned experimental vibrational frequencies.

#### **Results and Discussion**

#### Structure of the 12c4—Alkali Metal Cation Complexes.

Figure 1 shows the structure and atom numbering of the  $C_4$  conformation of the 12c4—alkali metal cation complex. Table 1 shows the geometry of this conformation at the B3LYP and MP2 levels using the 6-31+G\* basis set. The details of the MP2 computations are as given in ref 31. It can be noticed from the data in Table 1 that there is an increase of the displacement of the metal cation out of the ring plane, M—O bond length, where M refers to alkali metal cation, by the increase of the metal

cation size. While the C–C bond length is unchanged for the five complexes, the C–O bond length of the Cs<sup>+</sup> complex is shorter by 0.01 Å than that of the Li<sup>+</sup> complex. Also, the two CCO bond angles increased by about 4 degrees. There is a large increase of the  $MO_4C_5$  bond angle by about 14 degrees, although the other  $MO_4C_3$  bond angle increased by less than 2 degrees. In addition, there is a large decrease of the  $MO_1C_2C_3$  dihedral angle by about 20 degrees, as the large  $Cs^+$  cation is displaced out of the ring plane compared to that of the smaller Li<sup>+</sup> cation. These previous differences are monotonic in going from the smallest Li<sup>+</sup> cation to the largest  $Cs^+$  cation.

While the C-O bond lengths calculated at the MP2 and B3LYP levels are close to each other within 0.003 Å, the C-C bond lengths calculated at the B3LYP level are too short by not more than 0.01 Å, compared to those calculated at the MP2 level. The M-O bond length of the Li<sup>+</sup> complex is calculated to be equal at the MP2 and B3LYP levels, while for the Na<sup>+</sup> complex it is calculated to be too long by about 0.06 Å at the MP2 level compared to that at the B3LYP level. Using the effective core potential basis set for the K<sup>+</sup>, Rb<sup>+</sup>, and Cs<sup>+</sup> cations for the corresponding 12c4 complexes, the M-O lengths are calculated to be too short by about 0.06 Å at the B3LYP level compared to that at the MP2 level. There are two reports of the X-ray experimental geometry, to the best of our knowledge, of only the Na<sup>+</sup> complex. 45,46 The calculated geometries at the B3LYP and MP2 levels in Table 1 are in good agreement with these X-ray experimental geometries.

Experimental Vibrational Spectra of the 12c4—Alkali Metal Cation Complexes. The Raman spectra in the 100-1600 cm<sup>-1</sup> region of the Li<sup>+</sup>, Na<sup>+</sup>, and K<sup>+</sup> complexes in the solid phase and the IR spectra in the 400–1600 cm<sup>-1</sup> region of the five 12c4—alkali metal cation complexes of the solid phase are shown in Figures 2 and 3, respectively. No reasonable Raman spectra could be obtained for the Rb<sup>+</sup> and Cs<sup>+</sup> complexes, even using samples synthesized using either methanol or water as a solvent, as will be detailed shortly. Tables 2-4 list the Raman vibrational frequencies of the solid and methanol solution phases of the Li<sup>+</sup>, Na<sup>+</sup>, and K<sup>+</sup> complexes, in addition to the Raman frequencies of the water solution phase of the Li<sup>+</sup> and Na<sup>+</sup> complexes. The IR vibrational frequencies of the three Li<sup>+</sup>, Na<sup>+</sup>, and K<sup>+</sup> complexes of the solid phase are also added in Table 3. The assignment of the fundamental vibrations is included in Tables 2-4. The Raman vibrational frequencies of the methanol solution reported in ref 40 are added in Tables 2 and 3 for the Li<sup>+</sup> and Na<sup>+</sup> complexes, respectively.

It has been reported that the solvent used in the synthesis of the crown ether-metal complexes affects the stability of the formed complexes.<sup>26</sup> In the current study, the complexes were synthesized using methanol as a solvent. To study the effect of the solvent used in the synthesis of the complexes, the Li<sup>+</sup>, Na<sup>+</sup>, and K<sup>+</sup> complexes were also synthesized using water as a solvent. The Raman vibrational frequencies of the aqueous phase of the Li<sup>+</sup> complex, the solid and aqueous phase of the Na<sup>+</sup> complex and the solid phase of the K+ complex are included in Table 2. In addition, the IR vibrational frequencies of the solid and methanol solution phases of the Na<sup>+</sup> complex are added to Table 3. In the IR spectra of the Na<sup>+</sup> complex of the solid phase, Table 3, the difference between the frequencies of the corresponding bands, when methanol and water were used as solvents in the synthesis of the complex, is about 1 or 2 cm<sup>-1</sup>, but it can be as large as 6 cm<sup>-1</sup> for very weak bands. For the Raman spectra of the solid phase, the difference is also 1 or 2 cm<sup>-1</sup>, but for some bands, the difference is as large as 6 cm<sup>-1</sup>, such as for  $v_{29}$ . These small band position differences in the

TABLE 5: Vibrational Frequencies (cm<sup>-1</sup>), in Energy Order, of the  $C_4$ ,  $C_s$ ,  $C_s$ ,  $C_{2v}$ ,  $C_{2v}$  and  $C_s$  Conformations of the  $12c4-K^+$  Complex. The Vibrational Frequencies Are Scaled with a Frequency Scale Factor of 0.963<sup>71</sup>

		(	C <sub>4</sub>		$C_{\rm s}$		C <sub>s</sub>		Z <sub>2v</sub>		$\overline{C}_{2v}$		C <sub>s</sub>
no.	exp						freq						
1	F	В		A"		A"	66	$A_2$	47	$A_2$	63	A"	-26
2		В	112	A'	103	A'	94	$B_2$	88	$A_1$	105	A'	57
3		E	122	A' $A''$	116 129	A' $A'$	109 131	$A_1$ $A_2$	114 139	$B_2 B_1$	106 130	A'' $A'$	106 111
5	140	B	135	$A^{\prime\prime}$	149	$A^{\prime\prime}$	140	$A_1$	142	$B_2$	131	$A^{\prime\prime}$	131
6 7	189	$\stackrel{A}{E}$	149 184	A' $A'$	156 168	A'' $A'$	145 149	$B_2 B_1$	154 162	$A_1$ $A_2$	141 153	A' $A'$	136 162
8	109	L	104	A'	187	A'	180	$B_2$	191	$A_1$	153	A'	175
9	(217)	B	195	A''	208	A' $A''$	213	$A_1$	194	$A_1$	201	A''	198
10 11	239 269	$\stackrel{A}{E}$	234 259	A' $A''$	249 250	A''	235 249	$A_1$	222 242	$B_2 B_1$	206 219	A' $A''$	204 243
12				A'	266	A'	254	$A_2$	248	$B_1$	268	A'	266
13 14	304 349	$A \\ A$	282 331	$A^{\prime\prime}$ $A^{\prime}$	267 320	A' $A''$	304 306	$B_2 B_1$	289 299	$A_2$ $A_1$	292 302	A'' $A'$	274 331
15	356	E	346	$A^{\prime\prime}$	342	A'	321	$A_1$	337	$B_2$	316	$A^{\prime\prime}$	340
16 17	398	В	391	A' $A'$	361 440	A' $A''$	386 394	$A_1$	411 422	$A_1$ $B_1$	380 396	A' $A'$	354 442
18	497	B	472	$A^{\prime\prime}$	482	A'	505	$B_2$	500	$B_1$	506	$A^{\prime\prime}$	484
19 20	549	E	528	A' $A'$	498 567	$A^{\prime\prime} A^{\prime\prime}$	507 541	$A_2$	507 549	$B_2$ $A_1$	509 515	A' $A''$	499 543
21	597	A	570	$A^{\prime\prime}$	575	$A^{\prime\prime}$	544	$A_1 B_1$	570	$A_1$	534	A'	551
22	795	В	763 782	A'	757 780	A'	781	$B_1$	753	$B_2$	791 796	A'	759
23 24	808	E	182	A'' $A'$	786	$A^{\prime\prime}$ $A^{\prime}$	785 796	$A_1 B_2$	757 801	$A_1 B_1$	802	A'' $A'$	792 796
25	848	A	825	A'	832	A'	822	$A_1$	833	$A_1$	811	A'	817
26 27	902	$\frac{B}{A}$	879 882	A'' A''	873 879	$A^{\prime\prime}$ $A^{\prime\prime}$	866 888	$A_2$ $A_2$	866 879	$A_2 B_1$	873 891	A'' A''	866 888
28	915	E	893	$A^{\prime\prime}$	891	A'	897	$B_1$	887	$A_2$	899	A'	899
29 30	1024	A	998	A' $A'$	893 994	$A^{\prime\prime}$ $A^{\prime\prime}$	901 984	$B_2 B_1$	889 985	$B_2 B_1$	906 986	A'' A''	901 993
31	1030	E	1002	$A^{\prime\prime}$	1002	A'	1001	$A_1$	1002	$A_1$	1002	A'	999
32 33	1045	В	1019	A' $A'$	1014 1019	A' $A'$	1030 1031	$B_2$	1025 1032	$B_2$	1021 1030	A' $A'$	1014 1023
34	1043	A	1019	A''	1019	A''	1031	$A_1$ $A_2$	1032	$A_1$ $A_2$	1050	A''	1023
35	1095	E	1072	A''	1081	$A'_{II}$	1067	$B_2$	1078	$B_2$	1069	$A'_{II'}$	1071
36 37	1116	A	1092	$A' \\ A''$	1086 1095	$A^{\prime\prime}$ $A^{\prime\prime}$	1085 1093	$A_2 B_1$	1089 1095	$A_2 B_1$	1086 1109	A" A"	1085 1089
38		B	1100	$A^{\prime\prime}$	1112	$A^{\prime\prime}$	1112	$B_1$	1112	$B_2$	1117	A'	1108
39 40	1135	E	1114	$A' \\ A''$	1114 1118	A' $A''$	1121 1123	$\frac{A_2}{B_2}$	1115 1125	$A_2$	1127 1128	A" A"	1110 1134
41	1162	B	1136	A'	1142	A'	1143	$A_1$	1148	$A_1$	1133	A'	1143
42 43	1237 1250	$\frac{A}{E}$	1218 1229	A'' A'	1223 1228		1211 1224	$A_2 B_1$	1227 1227	$A_2 B_2$	1201 1212		1215 1226
44	1230	E		$A^{\prime\prime}$	1234	$A^{\prime\prime}$	1232	$B_2$	1243	$B_1$	1218	$A^{\prime\prime}$	1235
45	1250 1254	B	1240		1242		1250	$A_1$	1244	$A_1$	1227	A'	1237
46 47	1291	$\frac{B}{E}$	1249 1272	A'	1246 1267		1251 1264	$A_2 B_2$	1247 1265	$A_2 B_1$	1245 1254	A'' $A'$	1251 1266
48			1206	$A^{\prime\prime}$	1273	$A^{\prime\prime}$	1266	$B_1$	1295	$B_2$	1257	A''	1269
49 50	1306 1348	$A \\ A$	1286 1338	A''	1301 1337	A' $A''$	1277 1330	$A_1$ $A_2$	1303 1334	$A_1$ $A_2$	1268 1324	A' $A''$	1353 1334
51	1365	E	1354	$A^{\prime\prime}$	1355	$A^{\prime\prime}$	1347	$B_1$	1351	$B_1$	1338	A''	1348
52 53	1377	В	1369	$A'_{A''}$	1359 1369		1356 1370	$B_2 A_2$	1360 1372	$B_2 A_2$	1351 1355	$A'_{A''}$	1355 1369
54	1386		1380	A'	1375	A'	1374	$B_1$	1378	$A_1$	1377	A'	1378
55 56	1400	E	1386	A' $A''$	1387 1388	A''	1379 1400	$\frac{A_1}{B_2}$	1379 1403	$B_2 B_1$	1386 1395		1390 1398
57	1407	B	1389	$A^{\prime\prime}$	1402	A'	1411	$A_1$	1406	$A_1$	1405	A'	1402
58 59	1450 1449	$\frac{B}{E}$	1453 1454	A''	1443 1455		1452 1458	$\frac{A_2}{B_2}$	1447 1447	$A_2$	1453 1453	A''	1440 1453
60	1447	L		$A^{\prime\prime}$	1459		1458	$A_2$	1456	$B_1$	1471	$A^{\prime\prime}$	1456
61	1471	A	1455	A'	1459 1467		1463	$B_1$	1456	$A_2$	1473		1464
62 63	1467	$\frac{B}{E}$	1459 1468	A''	1467		1469 1473	$A_1$ $B_2$	1456 1474	$A_1$ $B_1$	1477 1479		1464 1473
64				A'	1482		1479	$B_1$	1474	$B_2$	1488		1476
65 66	1485	$\frac{A}{B}$	1482 2913		1491 2909		1487 2873	$A_1$ $A_2$	1492 2905	$A_1$ $A_2$	1491 2849		1490 2844
67		E	2919	A'	2914	A'	2878	$B_1$	2910	$B_1$	2849	A'	2853
68 69		A	2922	$A^{\prime\prime}$ $A^{\prime}$	2915 2919		2912 2918	$B_2$ $A_1$	2911 2915	$B_2 A_1$	2855 2856		2916 2918
70		B	2938	$A^{\prime\prime}$	2931	A'	2933	$A_2$	2934	$B_2$	2931	A'	2930
71 72		E	2941	A' $A''$	2932 2940		2935 2939	$B_2 B_1$	2934 2936	$A_1$ $A_2$	2932 2932		2931 2939
73		A	2944	A'	2943	$A^{\prime\prime}$	2939	$A_1$	2937	$B_1$	2934	A'	2941
74 75		$B_{\Gamma}$	2953	$A'_{A''}$	2948		2949	$B_2$	2941	$B_2$	2942		2954
75 76		E	2957	$A^{\prime\prime}$	2949 2958		2949 2954	$A_2 B_1$	2944 2954	$A_2 B_1$	2943 2946		2955 2970
77		A	2964	A'	2967	A''	2956	$A_1$	2955	$A_1$	2947	A'	2978
78 79		$\frac{B}{E}$	3004 3005	$A^{\prime\prime}$ $A^{\prime}$	2996 2996		2993 2993	$\frac{A_2}{B_2}$	2997 2997	$A_2$	2992 2992		2998 3000
80				$A^{\prime\prime}$	3000	A''	2998	$B_1$	3000	$B_1$	2993	A''	3001
81		A	3006	A'	3002	A'	2998	$A_1$	3000	$A_1$	2994	A'	3002

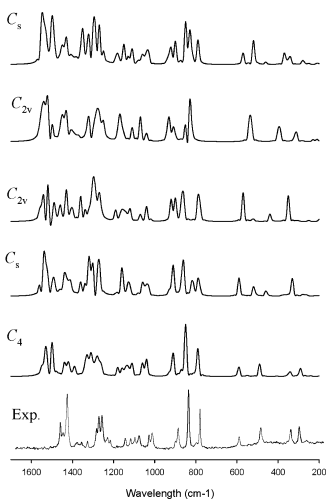
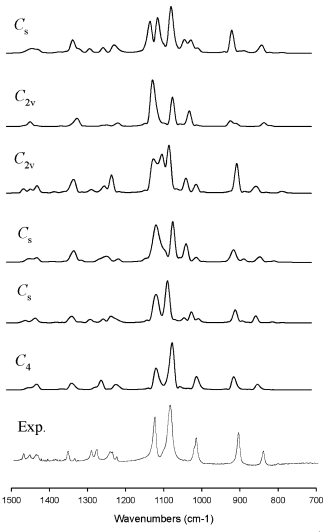


Figure 5. Experimental and calculated Raman spectra of the 12c4-K<sup>+</sup> complex using unscaled frequencies. The Raman intensity of the third lowest energy  $C_s$  conformation was not calculated. See text.

solid phase spectra are most probably due the crystal structure and orientation. In addition, these small band position differences are not associated with intensity differences and exist only in the Raman spectra and not in the IR spectra. Consequently, it is reasonable to assume that these small position differences in the Raman spectra are only due spectral reasons rather than being due to conformational change. The case is similar for the Li<sup>+</sup> and K<sup>+</sup> complexes. It can be concluded, for the 12c4alkali metal cation complexes, that the solvent used in the synthesis of the complexes has no effect on the conformational stability of the formed complexes. It is probable that this effect might be significant for the larger 18c6-alkali metal cation complexes with a larger and more flexible ring and consequently less energy gap between the possible conformations.<sup>26,30,73</sup>

All bands reported in ref 40 of the Raman spectrum of the Na<sup>+</sup> complex in the methanol solution phase, Table 3, are observed in the current study, also in the Raman spectrum for the same methanol solution phase, and at a position difference of not more than 2 cm<sup>-1</sup>. This is also observed for the Li<sup>+</sup> complex, Table 2, except for bands at 252, 262, and 579 cm<sup>-1</sup> that are not observed in the current study. Instead, the 252 and 262 cm<sup>-1</sup> bands are observed only in the Raman spectrum of the solid phase, while the 579 cm<sup>-1</sup> band is not observed in any of the measured spectra in the current study. Unlike the case of the Na<sup>+</sup> complex, the position difference between the bands observed in the current study and that in ref 40 of the Li<sup>+</sup> complex is about 4 cm<sup>-1</sup> and as large as 6 cm<sup>-1</sup>, for the Raman spectrum measured in same methanol solvent.



**Figure 6.** Experimental and calculated IR spectra of the 12c4-K<sup>+</sup> complex using unscaled frequencies.

It is clear from Figures 2 and 3 that the vibrational spectra of the five 12c4-alkali metal cation complexes are so similar that it is reasonable to conclude that the five complexes exist in the same conformation. As will be seen shortly, it is concluded that the five complexes exist in the  $C_4$  conformation. There is a shift of the position of some bands of the Li<sup>+</sup> complex, compared to those of the other four complexes. Figure 4 compares between the Raman spectra of the solid phase of the 12c4-Li<sup>+</sup>, -Na<sup>+</sup>, and -K<sup>+</sup> complexes in the region of 950-750 cm<sup>-1</sup>. For example,  $v_{55}$  of the 12c4-Li<sup>+</sup>, -Na<sup>+</sup>, and -K<sup>+</sup> complexes is at 931, 918, and 915 cm<sup>-1</sup>, respectively, while  $v_{56}$  of the same complexes is at 802, 802, and 808 cm<sup>-1</sup>, respectively. The shift of the position of some bands of the Li<sup>+</sup> complex is not necessarily toward higher energy. For example,  $v_{53}$  of the 12c4-Li<sup>+</sup>, -Na<sup>+</sup>, and -K<sup>+</sup> complexes is at 1082, 1098, and 1095  $\text{cm}^{-1}$ , respectively.

The similarity between the vibrational spectra of the five 12c4—alkali metal cation complexes, as presented by the data in Figures 2 and 3 and Tables 2–4, suggests that the five alkali metal cation complexes exist in the same conformation. Comparison was made between the experimental and calculated vibrational frequencies for six of the eight predicted conformations reported in ref 31 of the 12c4— $Li^+$ ,  $-Na^+$ , and  $-K^+$  complexes that are of symmetries higher than the  $C_1$  symmetry. These six conformations, in energy order, are of  $C_4$ ,  $C_s$ ,  $C_s$ ,  $C_{2v}$ ,  $C_{2v}$ , and  $C_s$  symmetries. Because of the doubly degenerate

E representation of the  $C_4$  point group, the comparison concludes that the three complexes exist in the  $C_4$  conformation. Table 5 shows such comparison for the 12c4-K<sup>+</sup> complex. The vibrational frequencies are scaled with a one frequency scale factor of 0.963.71 Although no Raman spectra could be obtained for the Rb<sup>+</sup> and Cs<sup>+</sup> complexes, the similarity between the IR spectra of the five complexes clearly shows that the five complexes exist in the same conformation. The existence of the five complexes in the  $C_4$  conformation is a reflection of the fact that the five alkali metal cations are larger than the ring cavity of 12c4. Scaling of the force field was then conducted using the B3LYP/6-31+G\* force field of the  $C_4$  conformation. The conclusion that the  $Na^+$  complex has a  $C_4$  structure is in agreement with X-ray result that this complex exists in the  $C_4$ conformation. 45,46 This is the only complex of the 12c4—alkali metal cation complexes for which, to the best of our knowledge, an X-ray result has been reported. Figures 5 and 6 compare between the experimental and calculated Raman and IR, respectively, spectra of the 12c4-K<sup>+</sup> complex. Because of an unsolved problem in the computations, no Raman intensities were calculated for the third lowest  $C_s$  conformation. Both figures, Figures 5 and 6, show a better agreement between the experimental and calculated Raman and IR spectra assuming a  $C_4$  conformation of the complex.

Table 6 compares between the calculated, unscaled, vibrational frequencies of the five 12c4-alkali metal cation complexes of the  $C_4$  conformation. The data in Table 6 reflect the similarity between the calculated vibration spectra of the five complexes as observed experimentally, Figures 2 and 3 and Tables 2-4. This is only with the exception of the region below 400 cm<sup>-1</sup>. This region is characterized by the vibrational modes involving the alkali metal cation, where the difference between the calculated geometries of the five alkali metal cations, described before, is reflected by the difference between the calculated spectra in this region. For example, for the 12c4-K<sup>+</sup> complex, the calculated TED shows a large contribution of the K<sup>+</sup>-O stretching mode for the  $v_{18}$ ,  $v_{20}$ ,  $v_{38}$ ,  $v_{39}$ ,  $v_{59}$ , and  $v_{61}$ vibrational modes. Notice also that these bands, compared to the other intense bands, are of modest intensity to have a significant influence on the measured experimental vibrational spectra.

Scaling of the Force Fields of the C<sub>4</sub> Conformation. 12c4 alkali metal cation complexes are large ring molecules with 81 normal modes of vibration. While free 12c4 has no redundant internal coordinates,<sup>25</sup> due to symmetry and the additional alkali metal cation, 12c4-alkali metal cation complexes have 90 redundant internal coordinates that were generated by the SCALE2 program. This set of 90 internal coordinates was reduced to 81 nonredundant internal coordinates. The nine vibrations corresponding to the redundant coordinates were calculated to be zero. A set of eight scale factors were used in the scaling of the force fields. The values of the scale factors were estimated after comparison between those determined for free 12c425 and those determined by Rauhut and Pulay as average scale factors of a set of 20 molecules.<sup>71</sup> Using this initial set of scale factors, without varying the scale factors, the rms deviations of the difference between the calculated and experimental vibrational frequencies for the Li<sup>+</sup>, Na<sup>+</sup>, and K<sup>+</sup> complexes were 11.0, 6.4, and 5.9 cm<sup>-1</sup>, respectively.

The definition of the internal coordinates, used in the conversion of the internal coordinate force fields to the Cartesian coordinate force fields, the initial and final values of the eight scale factors of the force fields of the three  $12c4-Li^+$ ,  $-Na^+$ , and  $-K^+$  cation complexes, and the corresponding rms devia-

TABLE 6: Comparison between the Calculated, Unscaled, Vibrational Frequencies (cm<sup>-1</sup>) and IR and Raman Intensities for the Five 12c4-Alkali Metal Cation Complexes of the C<sub>4</sub> Conformation<sup>a</sup>

	Li <sup>+</sup>			Na <sup>+</sup>				K <sup>+</sup>			Rb <sup>+</sup>		Cs	+	
no.	sym.	freq	IR	Ra	freq	IR	Ra	freq	IR	Ra	freq	IR	Ra	freq	IR
1	A	3141	24	38	3129	37	46	3120	58	48	3116	69	49	3112	79
2		3087	0	308	3079	5	91	3077	13	50	3075	16	59	3072	18
3 4		3079 3037	6 82	313 226	3067 3039	15 68	539 199	3056 3034	24 54	637 148	3051 3030	29 50	651 129	3045 3024	35 48
5		1542	12	9	1539	11	11	1539	9	13	1540	8	13	1540	7
6		1512	1	9	1511	2	11	1510	3	12	1510	3	13	1509	3
7		1432	0	1	1434	0	1	1432	1	1	1434	2	1	1434	2
8		1385	2	2	1388	2	2	1389	2	2	1390	2	2	1391	2
9		1327	18	4	1333	20	4	1335	21	5	1337	20	5	1339	21
10 11		1256 1136	6 5	5 3	1263 1133	4 0	4 1	1264 1141	4 0	4 2	1266 1144	5 0	4 2	1267 1148	5 0
12		1057	5	0	1060	0	4	1057	0	4	1057	0	4	1058	0
13		1026	63	1	1032	72	1	1036	76	1	1038	77	1	1039	79
14		920	0	0	915	0	7	912	0	8	910	0	8	910	0
15		871	33	20	861	38	17	857	43	15	854	44	14	854	46
16 17		603 384	2 44	3 5	596 354	1 9	3 4	592 344	1 8	3 2	593 340	1 7	3 1	593 337	0
18		322	31	1	310	1	2	292	0	3	289	ó	3	287	7
19		311	0	0	244	6	1	243	2	0	247	1	0	247	1
20		205	0	1	195	22	0	155	21	0	110	10	0	88	7
21	B	3140	0	40	3128	0	39	3119	0	39	3115	0	39	3110	0
22 23		3086 3076	0	66 115	3074 3062	0	56 97	3066 3050	0	42	3062 3046	0	35 81	3057 3039	0
23 24		3076	0	115 28	3002	0	50	3030	0	86 82	3046	0	96	3039	0
25		1525	0	2	1517	0	2	1515	0	1	1513	0	2	1513	0
26		1511	0	27	1509	0	27	1509	0	27	1508	0	26	1507	0
27		1436	0	1	1440	0	0	1442	0	0	1444	0	0	1445	0
28 29		1418	0	2	1420 1293	0	3	1422 1297	0	4	1422	0	4	1423	0
30		1287 1275	0	10 19	1293	0	26 6	1297	0	33 2	1300 1289	0	34 1	1301 1289	0
31		1173	0	2	1177	0	2	1180	0	2	1181	0	2	1181	0
32		1128	0	1	1132	0	3	1133	0	3	1135	0	3	1137	0
33		1064	0	3	1062	0	0	1068	0	0	1071	0	0	1073	0
34		928	0	7	915	0	0	916	0	0	915	0	1	916	0
35 36		782 496	0	8 4	786 493	0	9 4	792 490	0	9 4	793 488	0	9 3	795 487	0
37		403	0	0	409	0	0	406	0	0	406	0	0	406	0
38		234	0	0	216	0	0	202	0	0	198	0	0	194	0
39		171	0	0	159	0	0	140	0	0	136	0	0	136	0
40		108	0	0	124	0	0	117	0	0	112	0	0	103	0
41 42	E	63 3140	0 5	0 93	69 3129	0 8	1 91	56 3119	0 14	1 88	47 3115	0 17	1 86	45 3111	0 19
43	L	3086	63	20	3075	65	44	3069	57	61	3067	51	63	3062	46
44		3077	13	23	3064	31	30	3053	61	33	3048	79	37	3042	94
45		3030	2	4	3033	3	2	3030	5	4	3027	6	7	3021	7
46		1532	12	6	1525	12	4	1524	11	4	1523	11	4	1524	11
47 48		1511 1436	14	1 1	1510 1438	14	1 1	1509 1438	14	1 1	1509 1440	13	1 2	1508 1441	12
46 49		1402	1 27	0	1405	2 29	0	1436	2 31	1	1440	3 32	1	1441	3 33
50		1314	30	7	1320	28	6	1321	27	7	1323	25	7	1324	25
51		1263	24	1	1273	24	1	1276	24	2	1279	24		1280	24
52		1154	55	1	1154	69	1	1156	88	1	1157	98	2 2 2	1159	108
53		1102	245	4	1108	215	3	1113	187	2	1116	175	2 2	1118	162
54 55		1037 943	21 43	3 0	1039 930	16 50	2	1041 927	14 49	2	1042 925	13 50	0	1043 925	12 48
56		810	1	0	810	0	0	812	0	0	812	0	0	814	0
57		559	18	0	551	13	0	548	12	0	549	12	0	549	12
58		439	78	0	409	0	0	406	0	0	406	0	0	406	0
59		351	6	0	290	14	0	269	5	0	264	4	0	260	3
60		286	3	0	214	20	0	191	18	0	193	16	0	192 75	16
61		176	9	0	183	1	0	127	9	0	94	5	0	75	4

<sup>&</sup>lt;sup>a</sup> IR and Ra are the IR and Raman intensities, respectively. No Raman intensity was calculated for the Cs<sup>+</sup> complex.

tions are given in Table 7. Since none of the vibrations corresponding to the C-H stretching mode and most of the vibrations involving the Na butterfly mode were not assigned, the scale factors corresponding to the C-H stretching and Na butterfly modes were held fixed. The total number of the experimental frequencies used in the determination of the final scale factors are 50, 60, and 56 vibrational frequencies for the 12c4−Li<sup>+</sup>, −Na<sup>+</sup>, and −K<sup>+</sup> complexes, respectively.

The experimental vibrational frequencies used in the scaling of the force fields and the corresponding calculated vibrational frequencies and Raman and IR intensities of the  $C_4$  conformation of three 12c4-Li<sup>+</sup>, -Na<sup>+</sup>, and -K<sup>+</sup> cation complexes are given

TABLE 7: Natural Internal Coordinate Scale Factors<sup>a</sup>

				final	
coordinate	description	initial	Li <sup>+</sup>	Na <sup>+</sup>	K <sup>+</sup>
$R_1-R_4$	C-C stretching	0.970	0.9858	0.9977	0.9785
$R_5 - R_{12}$	C-O stretching	1.022	1.0013	1.0106	1.0092
$R_{13}$	M-O stretching	0.940	0.8927	0.8477	1.0900
$r_{13}-r_{28}$	C-H stretching <sup>b</sup>	0.920	0.9200	0.9200	0.9200
$\alpha_1 - \alpha_{32}$	CH <sub>2</sub> bending	0.940	0.9454	0.9417	0.9391
$\zeta_1 - \zeta_9$	ring bending <sup>c</sup>	1.040	1.0426	1.0130	1.0494
$\tau_1 - \tau_9$	ring Torsion	1.030	0.8968	1.0634	0.9784
$ au_{10}$	MO buterfly <sup>b</sup>	0.940	0.9400	0.9400	0.9400
rms	·		7.7	5.6	5.1

<sup>&</sup>lt;sup>a</sup> Nonredundant internal coordinates. M refers to alkali metal cation. <sup>b</sup> Fixed. See text. <sup>c</sup> Including those which contain the M metal cation.

in Tables  $8\!-\!10$ . Bands for which their assignments as fundamental was not certain, and were consequently not used in the scaling of the force fields, are included between parentheses in Tables  $8\!-\!10$ . In the scaling of the force fields, the experimental vibrational frequencies of the methanol solution, whenever available, were used, in most of the cases. Since the vibrational spectra of the three  $12c4\!-\!Li^+$ ,  $-Na^+$ , and  $-K^+$  complexes are quite similar, Figures 2 and 3, the assignment of the experimental to the calculated vibrational frequencies was revised so that the same strong features in the spectra of the three complexes would correspond to the same mode in the calculated spectra.

A few remarks should be mentioned about the assignment of the experimental to the calculated vibrational frequencies for the three 12c4-Li<sup>+</sup>, -Na<sup>+</sup>, and -K<sup>+</sup> complexes. Except for  $v_{17}$  and  $v_{19}$  of the Li<sup>+</sup> complex and  $v_{17}$  and  $v_{18}$  of the Na<sup>+</sup> and K<sup>+</sup> complexes, which are observed as strong bands in the Raman spectra at about 350 and 305 cm<sup>-1</sup>, respectively, most of the other vibrational bands in the 150-400 cm<sup>-1</sup> region are very weak and difficult to assign. In addition, only Raman spectral data are available in this region. Bands in the 100-150 cm<sup>-1</sup> region are also broad and due to the overlap of bands in this region are not possible to assign. In the vibrational spectra of the 12c4-Na<sup>+</sup> complex, three bands are observed in the 790-820 cm<sup>-1</sup> region, although, using the scaled force field, two bands are predicted to be in this region. The band observed at 794 cm<sup>-1</sup> was selected as a fundamental since it is observed and predicted as intense band in the Raman spectra. Two other bands are observed at 802 and 814 cm<sup>-1</sup>. The 802 cm<sup>-1</sup> band was selected, rather than the 814 cm<sup>-1</sup> band, as a fundamental in agreement with the scaled B3LYP vibrational frequency, at 800 cm<sup>-1</sup>. The 1340-1500 cm<sup>-1</sup> region is the most difficult region to assign. It can be divided into two regions, the first is at  $1340-1410 \text{ cm}^{-1}$  and the second is at  $1440-1500 \text{ cm}^{-1}$ . Either region has six bands, two of which are of E symmetry. It was easier to assign the second 1440-1500 cm<sup>-1</sup> region, but the difference between the calculated, using the scaled B3LYP force field, and experimental frequencies in this region was as high as  $21 \text{ cm}^{-1}$ . On the other hand, it was not simple to assign the first 1340–1410 cm<sup>-1</sup> region, especially since some bands were weak. Consequently, two calculated bands were preferred to be left unassigned in this region.

For the three 12c4–Li<sup>+</sup>, -Na<sup>+</sup>, and -K<sup>+</sup> complexes, the values of the scale factors corresponding to the C–C stretching, C–O stretching, and CH<sub>2</sub> bending modes, Table 7, are close to each other, within 0.02 at the highest, and those of the ring bending mode are close to each other, within 0.04. For the M–O stretching mode, the values of the scale factors for the three complexes are different, within 0.24, although this can be rationalized in large part due to the three different M alkali metal cations, Li<sup>+</sup>, Na<sup>+</sup>, and K<sup>+</sup>. The values of the scale factors

TABLE 8: Experimental and Calculated Vibrational Frequencies (cm $^{-1}$ ) and Raman and IR Intensities ((4 $\pi\epsilon_{\mathring{a}}$ ) $^2$ Å $^4$ amu $^{-1}$ ) of the  $C_4$  Conformation of the 12c4–Li $^+$ Complex $^a$ 

Compi	icx ·			cal	lcd	
		exptl		IR	Ran	nan
no.	sym	freq	freq	int	int	depo
1	A		3016	23.9	37.9	0.07
2 3			2964	0.4	307.0	0.15
4			2957	6.2	314.2 225.6	0.01
5		1499	2916 1499	82.5 11.8	8.7	0.00 0.75
6		1477	1470	1.3	8.7	0.64
7			1400	0.0	1.3	0.04
8		1350	1345	2.1	2.2	0.32
9 10		1289 1241	1292 1222	18.7 6.7	3.9 5.2	0.27
11		1113	1112	7.8	3.2	0.36 0.00
12		1048	1048	1.3	0.2	0.75
13		1020	1020	64.5	1.3	0.00
14		907	908	1.2	0.4	0.18
15 16		862 603	865 605	27.7 1.3	20.1 2.2	0.07 0.10
17		351	362	45.5	5.0	0.10
18		324	320	32.0	1.4	0.08
19		305	298	0.2	0.2	0.26
20	D.	(199)	199	0.1	0.7	0.08
21 22	B		3015 2963	0.0 0.0	40.5 66.1	0.75 0.75
23			2954	0.0	114.6	0.75
24			2903	0.0	28.2	0.75
25		1479	1482	0.0	2.3	0.75
26		1454	1468	0.0	26.9	0.75
27 28			1400 1378	0.0 0.0	1.3 1.9	0.75 0.75
29			1255	0.0	10.1	0.75
30		1261	1241	0.0	18.5	0.75
31		1159	1146	0.0	3.6	0.75
32 33		1120	1123	0.0	1.0 2.7	0.75
33 34		1059 917	1057 918	0.0 0.0	7.7	0.75 0.75
35		786	771	0.0	7.6	0.75
36		500	501	0.0	4.3	0.75
37		(400)	393	0.0	0.1	0.75
38 39		(229) (158)	232 164	0.0 0.0	0.3 0.3	0.75 0.75
40		(136)	104	0.0	0.3	0.75
41			60	0.0	0.5	0.75
42	E		3015	5.2	93.2	0.75
43 44			2963	63.1	19.6	0.75
45			2955 2909	13.3 2.1	22.8 4.4	0.75 0.75
46		1487	1489	11.8	5.8	0.75
47		1446	1469	13.7	1.3	0.75
48		1260	1398	1.1	1.0	0.75
49 50		1360 1277	1362 1281	26.6 34.2	0.2 6.6	0.75 0.75
51		1252	1229	29.5	1.3	0.75
52		1137	1131	119.1	2.0	0.75
53		1083	1092	175.3	2.7	0.75
54 55		1027	1030	21.1	2.7	0.75
55 56		931 802	934 799	41.1 0.2	0.3 0.0	0.75 0.75
57		562	563	20.3	0.0	0.75
58		415	413	73.4	0.2	0.75
59		341	345	4.3	0.0	0.75
60 61		282 (167)	281 170	4.8 9.5	0.1 0.1	0.75 0.75
01		(107)	1/0	7.3	0.1	0.75

<sup>a</sup> Here, sym, exptl, calcd, freq, int, and depo stand for symmetry, experimental, calculated, frequency, intensity and depolarization ratio, respectively. For the bands in parentheses, their assignment as fundamentals was not certain, and they were not used in the scaling of the force field.

corresponding to the ring torsion mode are also different, within 0.17. Notice that in the vibrational analysis of free 12c4, for most of the conformations considered in that study, a value of about 1.05 was obtained for the scale factor corresponding to the same ring torsion internal coordinate.<sup>25</sup> This value of 1.05 is the closest to that of the Na<sup>+</sup> complex, in the current study,

TABLE 9: Experimental and Calculated Vibrational Frequencies (cm<sup>-1</sup>) and Raman and IR Intensities  $((4\pi\epsilon_{\hat{a}})^2)$  $\rm \AA^4~amu^{-1}$ ) of the  $C_4$  Conformation of the 12c4-Na $^+$ Complex<sup>a</sup>

calcd Raman exptl IR freq int int depo no. sym freq 0.03 A3005 37.3 46.1 2 2956 4.8 91.0 0.47 3 14.9 2945 540.4 0.00 4 2918 67.6 197.5 0.01 5 1486 1494 11.0 11.2 0.75 1453 1467 2.0 10.5 0.66 (1396)1396 0.2 1.1 0.05 8 1354 1348 3.1 2.2 0.39 9 1305 1297 21.2 3.9 0.35 10 1237 1227 5.3 4.3 0.35 11 1115 1118 8.7 3.0 0.00 12 1064 1057 0.7 0.2 0.32 13 1028 70.7 0.01 1028 1.63 14 895 899 0.2 0.3 0.71 15 852 861 34.2 16.1 0.07 1.2 16 601 594 3.1 0.082.9 17 347 352 99 0.00 3.1 18 302 300 0.3 0.04 19 248 0.5 248 7.6 0.01 20 196 191 20.4 0.1 0.64 21 22 3004 0.75 В 0.0 38.7 2952 0.0 55.7 0.75 23 24 2940 0.0 97.1 0.75 2905 0.0 50.0 0.75 25 1470 1473 1.7 0.75 0.0 26 27 1444 1465 0.0 0.75 26.7(1400)1399 0.75 0.0 0.3 28 1380 1378 0.02.7 0.75 29 27.5 1268 1258 0.00.75 30 1256 1251 0.0 3.5 0.75 31 3.5 0.75 1160 1158 0.0 32 1124 1133 0.0 1.1 0.75 33 1049 1051 0.0 3.4 0.75 34 7.8 906 907 0.0 0.75 35 794 777 0.08.5 0.75 36 494 494 0.0 3.6 0.75 37 (414)410 0.0 0.1 0.75 38 204 210 0.0 0.2 0.75 39 0.5 0.75 148 155 0.0 40 121 0.1 (131)0.0 0.75 41 71 0.0 0.8 0.75 42 Е 3004 8.4 91.5 0.75 43 2953 64.8 43.4 0.75 44 2942 0.75 31.4 29.8 45 2912 2.0 0.75 3.1 46 1476 1480 11.7 4.2 0.75 47 1447 1466 13.4 1.0 0.75 48 1410 1399 2.2 1.2 0.75 49 1363 27.5 0.4 0.75 1365 50 5.9 1290 1284 35.1 0.75 51 1248 1239 31.6 1.4 0.75 52 53 1138 1140 145.9 2.4 0.75 1.8 1098 1100 132.8 0.75 54 1033 1033 14.3 2.3 0.75 55 918 920 45.0 0.3 0.75 56 0.75 802 800 0.1 0.2 57 553 552 12.9 0.1 0.75 58 364 0.75 368 0.1 6.6 59 285 282 10.8 0.00.75 215 60 215 25.9 0.2 0.75 181 0.0 0.75

which has the largest number of bands corresponding to the ring torsion mode assigned. This value of 1.05 is most different from that of the Li<sup>+</sup> complex. In a different calculation the scale factors of the Li+ complex were varied after all of the low energy vibrational bands whose assignment as fundamentals were not certain, and thus are included between parentheses in Table 9, were included in the refinement of the scale factors. The rms deviation became better, 6.8 cm<sup>-1</sup>, but the values of the scale factors were close to those in Table 7, within 0.004.

Frequencies (cm<sup>-1</sup>) and Raman and IR Intensities  $((4\pi\epsilon_{\hat{a}})^2)$  $A^4$  amu<sup>-1</sup>) of the  $C_4$  Conformation of the 12c4-K<sup>+</sup> Complex<sup>a</sup>

TABLE 10: Experimental and Calculated Vibrational

no.         sym         exptl freq freq freq int         IR int         Raman int         depo           1         A         2996         58.1         47.7         0.02           2         2954         13.1         50.3         0.61           3         2934         23.8         637.0         0.01           4         2913         53.9         148.2         0.03           5         1485         1491         8.4         13.3         0.75           6         1471         1464         2.9         11.8         0.67           7         1206         1202         14.8         14.9         14.8         14.8
no.         sym         freq         freq         int         int         depo           1         A         2996         58.1         47.7         0.02           2         2954         13.1         50.3         0.61           3         2934         23.8         637.0         0.01           4         2913         53.9         148.2         0.03           5         1485         1491         8.4         13.3         0.75           6         1471         1464         2.9         11.8         0.67
2     2954     13.1     50.3     0.61       3     2934     23.8     637.0     0.01       4     2913     53.9     148.2     0.03       5     1485     1491     8.4     13.3     0.75       6     1471     1464     2.9     11.8     0.67
3 2934 23.8 637.0 0.01 4 2913 53.9 148.2 0.03 5 1485 1491 8.4 13.3 0.75 6 1471 1464 2.9 11.8 0.67
4 2913 53.9 148.2 0.03 5 1485 1491 8.4 13.3 0.75 6 1471 1464 2.9 11.8 0.67
5 1485 1491 8.4 13.3 0.75 6 1471 1464 2.9 11.8 0.67
6 1471 1464 2.9 11.8 0.67
7 1386 1392 1.1 1.4 0.04 8 1348 1347 3.8 2.8 0.38
9 1306 1297 22.5 4.4 0.36
10 1237 1226 5.1 4.3 0.31
11 1116 1118 7.9 2.7 0.00 12 1059 1056 1.8 0.4 0.05
13 1024 1029 76.0 1.7 0.02
14 902 0.0 0.4 0.72
15 848 852 36.9 14.3 0.07 16 597 592 0.7 2.9 0.09
17 349 344 7.9 2.6 0.00
18 304 297 0.4 2.2 0.07
19 239 240 2.0 0.5 0.02 20 159 20.9 0.1 0.51
21 B 2995 0.0 39.3 0.75
22 2944 0.0 42.3 0.75
23 2929 0.0 86.2 0.75 24 2903 0.0 81.7 0.75
25 1468 0.0 1.8 0.75
26 1450 1463 0.0 26.3 0.75
27 1407 1398 0.0 0.3 0.75 28 1377 1378 0.0 3.6 0.75
28 1377 1378 0.0 3.6 0.75 29 1254 1259 0.0 34.2 0.75
30 1250 1251 0.0 0.7 0.75
31 1162 1158 0.0 3.4 0.75 32 1138 0.0 1.7 0.75
32 1138 0.0 1.7 0.75 33 1045 1048 0.0 3.6 0.75
34 902 898 0.0 8.1 0.75
35 795 782 0.0 8.8 0.75
36 497 497 0.0 3.5 0.75 37 398 403 0.0 0.2 0.75
38 (217) 208 0.0 0.1 0.75
39 140 141 0.0 0.4 0.75
40 115 0.0 0.1 0.75 41 56 0.0 1.0 0.75
42 E 2995 13.6 88.1 0.75
43 2947 57.0 60.6 0.75
44 2931 61.7 32.8 0.75 45 2909 4.6 3.8 0.75
46 1467 1477 11.0 3.8 0.75
47 1449 1463 12.8 0.9 0.75
48 1400 1397 2.4 1.3 0.75 49 1365 1362 29.2 0.6 0.75
50 1291 1284 35.6 6.3 0.75
51 1250 1239 33.1 2.0 0.75
52 1134 1141 177.2 2.9 0.75 53 1094 1100 88.2 1.1 0.75
54 1030 1033 14.9 2.3 0.75
55 913 914 43.7 0.3 0.75
56 808 803 0.2 0.2 0.75 57 549 551 11.1 0.2 0.75
58 359 360 6.5 0.1 0.75
59 269 272 6.0 0.1 0.75
60 189 188 17.0 0.0 0.75 61 130 8.8 0.2 0.75

<sup>&</sup>lt;sup>a</sup> See corresponding footnote in Table 6.

It is reasonable then to assume that the values of the scale factor corresponding to the ring torsion mode of the Li<sup>+</sup> complex are not in error. In addition, this difference in the values of the scale factor, between that of the Li<sup>+</sup> and Na<sup>+</sup> complexes, is most probably too large to be attributable to a wrong assignment.

The rms deviation, of the difference between the experimental and calculated vibrational frequencies, of the Li<sup>+</sup> complex is higher than that of the other two Na<sup>+</sup> and K<sup>+</sup> complexes, Table 7. This larger rms deviation of the Li<sup>+</sup> complex is attributed

<sup>&</sup>lt;sup>a</sup> See corresponding footnote in Table 6.

mainly to seven bands, out of the 50 bands assigned for this complex. In a different calculation, these seven bands were excluded from the assignment. The rms deviation became 4.1 cm<sup>-1</sup> and the values of the scale factors were almost unchanged, within 0.009, including that of the ring torsion mode. Since the assignment of these seven bands was reliable, they were included in the final calculations, as presented in Table 8.

Relative Energies of the  $C_4$  and  $C_s$  Conformations of the Li<sup>+</sup> Complex. The vibrational study preformed in this report predicts a C<sub>4</sub> structure of the 12c4-Li<sup>+</sup> complex, as well as for the other four alkali metal cation complexes. As was mentioned above, in the previous conformational analysis report of the 12c4-alkali metal cation complexes,<sup>31</sup> the  $C_4$  conformation was predicted to be the most stable conformation for the four 12c4-Na<sup>+</sup>, -K<sup>+</sup>, -Rb<sup>+</sup>, and -Cs<sup>+</sup> complexes. For the 12c4-Li<sup>+</sup> complex, a  $C_s$  conformation was calculated to be more stable than the  $C_4$  conformation by 0.16 kcal/mol at the MP2/6-31+G\* level. The calculated binding energy predict a  $C_4$  or  $C_s$  conformation of the 12c4–Li<sup>+</sup> complex but could not differentiate whether the complex exists in the  $C_4$  or  $C_s$ conformation. This is because the calculated binding energies of the  $C_4$  and  $C_s$  conformations are too close to each other, -93.0 and -89.9 kcal/mol, at the MP2/6-31+G\* level, respectively. This is accompanied by a large uncertainty of the experimental binding energy, -90±12 kcal/mol. To further clarify whether the  $C_4$  or  $C_s$  conformation is the more stable conformation of the 12c4-Li<sup>+</sup> complex, optimized geometry of the complex was calculated for the  $C_4$  and  $C_5$  conformations at the MP2/6-311++G\*\* level. The MP2 computations were performed with the fixed core option. The  $C_4$  conformation was calculated to be more stable by 0.13 kcal/mol than the  $C_s$ conformation. This is compared to an energy difference of 0.16 kcal/mol at the MP2/6-31+G\* level, with the  $C_s$  conformation being more stable than the  $C_4$  conformation. These energy differences, at either the MP2/6-31+G\* or MP2/6-311++G\*\* level, are too small compared to the forces which are experienced in the solid or the solution phase. It is then concluded that other forces in the solid or the solution phase are the stabilizing force of the  $C_4$  conformation over the  $C_s$  conformation for the Li<sup>+</sup> complex.

#### Conclusion

This is the first time, to the best of our knowledge, that a full and detailed vibrational analysis of the spectra of any of the five 12c4-alkali metal cation complexes is reported. The similarity between the measured vibrational spectra of the five complexes indicates that the five 12c4-alkali metal cation complexes exist in the same conformation. Comparison between the experimental and calculated vibrational frequencies of the possible conformations of the 12c4-alkali metal cation complexes, predicted in a previous conformational analysis study,<sup>31</sup> indicates that the five 12c4—alkali metal cation complexes exist in the  $C_4$  conformation. This conclusion is in agreement with the previous conformational analysis study that the  $C_4$  conformation is the lowest energy conformation of the 12c4-Na<sup>+</sup>,  $-K^+$ ,  $-Rb^+$ , and  $-Cs^+$  complexes. For the Li<sup>+</sup> complex, the  $C_s$  conformation was calculated to be more stable than the  $C_4$ conformation by only 0.16 kcal/mol, at the MP2/6-31+G\* level.31,61 To further investigate this point, optimized geometries were calculated, in the present work, for the  $C_s$  and  $C_4$ conformations of the Li<sup>+</sup> complex at the MP2/6-311++G\*\* level. The  $C_4$  conformation was calculated to be more stable than the  $C_s$  conformation by 0.13 kcal/mol.

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