



Applied Spectroscopy Reviews

ISSN: 0570-4928 (Print) 1520-569X (Online) Journal homepage: https://www.tandfonline.com/loi/laps20

Investigation of L(+)-Ascorbic acid with Raman spectroscopy in visible and UV light

Rolf W. Berg

To cite this article: Rolf W. Berg (2015) Investigation of L(+)-Ascorbic acid with Raman spectroscopy in visible and UV light, Applied Spectroscopy Reviews, 50:3, 193-239, DOI: 10.1080/05704928.2014.952431

To link to this article: https://doi.org/10.1080/05704928.2014.952431

© 2015 The Author(s). Published with license by Taylor & Francis© Rolf W. Berg



0

Published online: 20 Aug 2014.

|--|

Submit your article to this journal 🗹

Article views: 4790



🜔 View related articles 🗹

View Crossmark data 🗹



Citing articles: 8 View citing articles 🗹



Investigation of L(+)-Ascorbic Acid with Raman Spectroscopy in Visible and UV Light

ROLF W. BERG

Department of Chemistry, Technical University of Denmark, Lyngby, Denmark

Abstract: Raman spectroscopy investigations of L(+)-ascorbic acid and its mono- and di-deprotonated anions (AH^{-} and A^{2-}) are reviewed and new measurements reported with several wavelengths, 229, 244, 266, 488, and 532 nm. Results are interpreted, assisted by new DFT/B3LYP quantum chemical calculations with 6-311++G(d,p) basis sets for several conformations of ascorbic acid and the anions. Raman spectra were measured during titration with NaOH base in an oxygen-poor environment to avoid fluorescence when solutions were alkaline. The ultraviolet (UV) absorption band for ascorbic acid in aqueous solution at \sim 247 nm was found to cause strong resonance enhancement for the ring C-C stretching mode (called **B**) at ~1692 cm⁻¹. The ascorbate mono-anion absorbs at \sim 264.8 nm giving Raman resonance enhancement for the same ring C–C bond stretching, downshifted to ~1591 cm⁻¹. Finally, for the ascorbate dianion, absorption was found at ~298.4 nm with molar absorptivity of ~7,000 L mol⁻¹ cm^{-1} and below ~220 nm. With UV light (244 and 266 nm), strongly basic solutions gave pronounced Raman resonance enhancement at $\sim 1556 \text{ cm}^{-1}$. Relatively weak preresonance enhancement was seen for A²⁻ when excitation was done with 229 nm UV light, allowing water bands to become observable as for normal visible light Raman spectra.

Keywords: Raman spectroscopy, ascorbic acid, ascorbate, X-ray structure, DFT calculation, conformation

Introduction

Recently, when studying ultraviolet (UV)-Raman spectra of food and beverages, we discovered interesting details in the resonance Raman spectra of the L-enantiomer of ascorbic acid, commonly also called vitamin C: No matter what kind of fruit juice, beer, or wine studied with 229 nm excitation, we did not observe anything other than the ascorbic acid, either naturally present or added for preservation. This nonlinear behavior with respect to Raman spectral intensity motivated the present study of ascorbic acid, $C_6H_8O_6$ or AH_2 .

Ascorbic acid is a cyclic lactone (ester) containing an endiol group, -C(OH)=C(OH)-, as shown in Figure 1. The compound is acidic, as a consequence of electron delocalization in the ring system, making the OH group—labeled 3—prone to giving off its proton when in contact with neutral or basic solutions (1). Ascorbic acid is also a strong antioxidant, with a high reactivity toward oxygen. The acid participates

Address correspondence to Rolf W. Berg, Department of Chemistry, Technical University of Denmark, Building 207, DK-2800 Kgs. Lyngby, Denmark. E-mail: rwb@kemi.dtu.dk

Color versions of one or more of the figures in the article can be found online at www.tandfonline. com/laps.

[©] Rolf W. Berg



Figure 1. L-Ascorbic acid, C6H8O6 (systematic name (5R)-5-((1S)-1,2-dihydroxyethyl)-3,4dihydroxyfuran-2(5H)-one), or here just AH₂. The endiol group is the -C(OH)=C(OH)- part of the ring with the double bond between C2 and C3. Other atom numbering systems have been used in the literature (4, 24–26, 43). The dihedral (torsional) angles are defined as shown for atoms A–B–C–D, with A nearest the observer, who is looking down the B–C bond. The angle is that between the projected bonds AB and CD (+ clockwise).

in many redox reactions in natural biological systems and is commonly found in small concentrations in fruits and vegetables (2–6). The acid is soluble in water and is an important nutrition component for humans, primates, and guinea pigs, which all are incapable of synthesizing it from glucose due to their lack of the enzyme gulonolactone oxidase in the glucoronic acid pathway (7). A deficiency of ascorbic acid in a person's diet causes a hemorrhagic condition (scurvy) of the skin and gums with attack on collagen in binding tissues. The acid is sometimes added to food as a preservative; for example, it helps to decrease the browning of white wines (8) and to protect human skin from degradation in UV light (9).

The reducing behavior of AH₂—associated with the endiol group in the ring—limits the stability in solution and in the solid state under ambient environments. At physiological pH values, the ascorbic acid AH_2 , depending on the concentration, splits off a proton to form the ascorbate ion AH⁻. Chemical evidence, X-ray structure solutions, and ab initio studies indicate that the deprotonation involves the proton H3 on the O3 site; see Figure 1. The OH group on C3 has an acidity constant of $pK_a \sim 4.1$ to ~ 4.25 , forming a highly stable AH⁻ mono-anion of ascorbic acid (4, 10–17). Other tautomers are not so stable; for example, the hydrogen on O2 can be moved to O1, forming a tautomer of ascorbic acid (4), but this is not the only possibility. The so-called iso-ascorbic acid, a stereoisomer of the ascorbic acid, occurs with inversion of the OH group at C5 but behaves in a quite similar way. The ionization of ascorbic acid is accompanied by a distortion of the planarity of the ring (18). In strongly alkaline solutions a further proton is lost to form one or more forms of the ascorbate di-anion A^{2-} (13, 19). The most stable di-anion in water solution seems to be obtained by deprotonation of the O2 site or eventually the O6 or O5 sites of the O3 deprotonated ascorbate mono-anion (4); see Figure 1. Formations of these di-anions are rare under physiological conditions requiring a highly basic environment because pK_a for the ascorbate mono-anion is ~11.5 to ~11.79 (11, 14, 16, 17, 19). At even higher basic pH values isomerization at C4 may take place (20). The acid/base and oxidation/reduction reactions of ascorbic acid are coupled, and in highly basic solutions and under the presence of oxygen the molecules are easily oxidized to various radicals, forming in the end salts of the dehydroascorbic acid, $C_6H_6O_6$ (ascorbic acid that has lost H2 and H3 protons and two electrons) (19, 21–23). Quite stable free radicals of ascorbic acid seem to be formed via ionization of the O3 or O2 anions, and also a stable di-radical seems to exist via ionization of the O3 O2 di-anion (4).

Structural Determinations

The crystal structure of AH₂ is known after complete solutions by Hvoslef (24, 25), based on X-ray and neutron diffraction data, and a redetermination at 120 K was later obtained (26). The crystal structures of several salts of AH₂ have been obtained, including the sodium salt NaAH (10), as well as calcium (27–29), strontium (30), lithium (31), thallium (32), and other ascorbates (33). The crystal structures were often found to contain various amounts of hydrate water in addition to one or more ascorbate ions that look much like the AH₂ depicted in Figure 1, although without H3. Interestingly, in, for example, the calcium dihydrate and thallium salts, two different ascorbate ions are present in the unit cell and—as in AH₂—these ions assume quite different conformations (different torsion angles χ_1 and χ_3 around C2–O2 and C4–C5 bonds) (27–29, 32). In addition, we note that the hydrogen-bonding tendencies of O2 and O3 in certain ascorbate salts seem to favor the formation of anion dimers in solution (1, 12, 28, 34–37). No salt of the rather unstable A^{2–} ion is at hand, and no crystal structure thereof seems to have been solved.

Raman Vibrational Spectroscopy

In 1943 Edsall and Sagall (38) used filtered light from a mercury arc (the blue e and violet k lines at 435.8 and 404.7 nm, respectively) to excite Raman spectra of aqueous solutions. From comparing three different ascorbic-like compounds, assignments of bands due to AH_2 and AH^- were made, and it was concluded that the presence of the ring in the structures gave rise to strong signals in the Raman spectra and that signals from the side chains were not nearly as strong. In 1971 Hvoslef and Klæboe used a red helium-neon laser to study AH₂ as crystals and in aqueous dilution to supplement infrared absorption experiments. This pioneering work also included deuteration studies (39). Structure clarifications on L-ascorbic acid were later made by infrared (IR) and UV spectroscopy by, for example, Falk and Wojcik (40), Lohmann et al. (41, 42), and Ferrer and Baran (43). The acid and many salts were studied as solids and in aqueous solutions (also in D₂O between AgCl windows) by use of IR and ¹³C-NMR spectroscopy (17, 34–37, 41–44). Panicker et al. (45) obtained Fourier transform infrared (FTIR) and FT-Raman spectra including surface-enhanced Raman spectra studies with Ag colloids (probably excited with a 1,064 nm Nd-YAG laser), and many of the vibrational modes of the acid were confirmed or reassigned. The influence of high pressures (up to 5.9 GPa) on the hydrogen bonding in AH_2 crystals has been studied (46), and extreme experiments have been performed with simple equipment in cold environments and at high altitudes (47). The temperature dependence (from 15 to 418 K) of the AH₂ Raman spectrum has been investigated, and a crystal phase change in the temperature range 200-270 K has been identified (48). (It is interesting that this phase change, detected by short time scale spectroscopy, did not influence the average structure—based on long timescale diffraction experiments—compare (24)-(26)-but such behaviors have been seen for others crystals, as discussed in, for example, (49–51).) Infrared spectra of AH_2 in the polycrystalline state and in aqueous solution have been presented by Bichara et al. (52). Most recently, IR and Raman spectra of ascorbic acid were given and their vibrational origin discussed by Singh et al. (53) and Yadav et al. (54) (the last report repeats much of the work in the first, including some printing errors).

Many Raman spectra in the published literature are incomplete or of minor quality and the ascorbate salts have not been well studied. Consequently, we found it necessary to critically review the literature and perform new measurements as reported in the following sections.

Quantum Chemical Modeling

Early ab initio calculations for AH_2 and AH^- were reported in 1976 by Carlson et al. (55), who started from the molecular geometry of the crystallographic data. Later, Al-Laham et al. (56) performed an STO-3G conformational analysis of the acid by forcing the ring geometry to be constant and optimizing only the conformation of the side chain. In 1997, Milanesio et al. (26), apparently not aware of the previous works, made extensive Hartree-Fock (HF) and density functional theory (DFT) calculations with B3LYP/6-311+G(2d,2p) basis sets. They described several conformers of the isolated AH₂ molecule, including a global minimum conformation called G, characterized by four intramolecular hydrogen bonds. Apparently independent of this, in 1998 Mora and Melendez (57) optimized 36 conformers of the AH_2 molecule at different levels of theory (RHF/6-31G, RHF/6-31G(d,p), RHF/6-311+G(d,p), and MP2/6-31G(d,p)) and described several optimized molecular geometries of low energy, including one close to the so-called crystallographic B structure of the solid (note that their results are given for the mirror image D-AH₂). O'Malley (58) modeled the effect of hydrogen bonding by considering up to three water molecules in the first solvation shell. In 2003, Juhasz et al. (15), based on DFT/B3LYP/6-31G(d) type calculations, elaborated on the effects of conformation on the acidity of the AH2 acid and found a global minimum with a minimum energy of -684.755581553 Ha in the gas phase. Later, in 2006, Dimitrova (59), not citing these previous works, reported results based on SCF/6-31G(d,p), B3LYP/6-31G(d,p), and B3LYP/6-31++G(d,p) models, and she obtained vibrational frequencies of L-AH₂ (alone and with five nearby water molecules) and correlated her frequencies with the experimental results of Hvoslef and Klæboe (37). Because the ascorbic acid has several hydrogen bond accepting and donating sites, these few water molecules are not sufficient for a good model, and more water molecules should be necessary to complete the first solvation shell. Geometry optimization of species in aqueous solution by use of a continuum model offers an attractive alternative. Accordingly, Allen et al. (4) have made a comprehensive study of ascorbic acid conformers, including some tautomers, some anions, and radical species in an aqueous conducting polarization continuum model and in the gas phase, using the DFT/B3LYP/6-311++G(d,p) high level method. Costanzo et al. (23) have added a further study of much the same content in the gas phase but including molecular dynamics on AH^- and the radical anion A^{*-} modeled in aqueous solution. Shimada et al. (46) also calculated the Raman spectrum of AH₂ starting from the established crystal structure, and Bichara et al. (52) reported further DFT calculations without referencing much of the previous work. Finally, Singh et al. (53) and Yadav and others (54) have reported comprehensive structural and vibrational studies of AH_2 and the AH_2^- and AH_2^+ radical ions using DFT methods and reporting optimized geometric structural, experimental vibrational spectra (harmonic frequencies along with IR intensities and Raman activities and depolarization ratios), as well as thermodynamic properties.

By reviewing these reports, it seems that many of the researchers have not read most of the prior literature on the subject. This is reflected in the many different numbering schemes for the atoms in the ascorbic acid and is why we are motivated to give a preferred definition of the atom numbers—based on Hvoslef's work (24, 25)—as proposed in Figure 1. The numbering problem makes it rather cumbersome to compare the many different previous reports, but in general the researchers have obtained rather consistent results.

To complement the many previous calculations we found it necessary to perform a comparative study on the AH^- and A^{2-} anions, as well as similar recalculations on the AH_2

molecule. It appeared that the Gaussian 03W program (60) could give reasonably reliable results at the DFT/B3LYP/6-311++G(d,p) level (61).

UV Resonance Raman Spectroscopy

In 1946, Harrand and Lennuier (62)—when exciting some yellow crystals with yellow light—noticed an intimate connection between the absorption and an enhancement of certain vibrational transition bands related to the chromophore. The effect was later named resonance Raman (RR) and a short review was recently given (63). The enhancement means that much more intensive spectra can be obtained and applied analytically but an inevitable problem is that samples get heated and often destroyed because of the absorption; however, certain samples can intentionally be moved, thereby avoiding decomposition (64–66).

It has been known for long time that ascorbic acid absorbs in deep UV; the UV spectra of ascorbic acid/ascorbate in aqueous solution, for example, show $\pi \rightarrow \pi^*$ absorption peaks at ~243.5 and ~265.5 nm (11) while being transparent in other ranges. Large variations in peak position and absorptivity have been reported, with values depending on concentration and pH, due to dissociation of the acid and the reactivity with oxygen. The molar absorptivity at these wavelengths is perhaps about ~9,560 and ~14,560 L mol⁻¹ cm⁻¹, respectively (9, 11, 14, 67). The absorption bands have been used for analytical purposes in many investigations (22, 68, 69).

In a search for UV absorbers it was recently discovered that divalent ascorbate anions embedded in a calcium carbonate lattice could be used (16). The filter solid was prepared by dripping aqueous solutions of CaCl₂ into Na₂CO₃ solutions containing dissolved ascorbic acid that is transformed into divalent ascorbate anions by the base. The filter material gains stability due to the absence of O₂ molecules inside CaCO₃. The optical absorption was seen in the UV B region (280–315 nm) and was ascribed to the absorption of A^{2-} ions at 292 nm in the CaCO₃ lattice and at 297.5 nm in aqueous solution (16).

Interestingly, there has been little specific mention in the literature that RR spectroscopy could be useful for studies on ascorbic acid compounds. According to the strong UV light absorptions, AH_2 , AH^- , and A^{2-} should exhibit strong RR enhancements when excitation is done with wavelengths within the UV range. The close connection between the absorption process and the resonance enhancement has been examined comprehensively by, e.g., Albrecht (70) and Hassing and Sonnich Mortensen (71).

By resonance Raman spectroscopy it may be possible to detect the presence of very small amounts of compounds due to the strong signal enhancement. For instance, Tuschel et al. (72) have shown that small amounts of explosives could be detected by using resonance-enhanced Raman spectroscopy with a 229 nm deep ultraviolet (DUV) laser. However, it must be done with care because the RR effect may destroy the normal Beer-Lambert-Bouguer type of relation. The signal needs not be linearly related to the concentration of the resonating chromophore (63, 73–75) and the resolution of UV Raman spectra may sometimes be limited (76). Raman spectroscopy performed with DUV laser lines is known to have the possibility to give spectra with little fluorescence (77). Thus, Loppnow et al. (78) reported several UV Raman spectra of aromatic hydrocarbons with very little fluorescence for compounds like toluene and naphthalene excited within their absorption bands. These aromatic hydrocarbons exhibit bands at about 1600 cm⁻¹, assignable to the C=C vibration or ring stretching, and these modes are strongly active in RR (78).

Therefore, it seemed worthwhile to examine ascorbic acid by DUV Raman spectroscopy.



Figure 2. Raman cell for titration experiments in oxygen-free pure argon atmosphere.

Experimental

The compounds L(+)-ascorbic acid (C₆H₈O₆, CAS Reg. No. 50-81-7) and L(+)-sodium ascorbate (NaC₆ H_7O_6 , CAS Reg. No. 134-03-2) were obtained in highly pure states from Sigma Aldrich (St. Louis, MO, USA), and spectra were obtained directly on the solid salts or on solutions in cuvettes. Solutions of AH₂ in H₂O were prepared by use of freshly boiled cold deionized and pure normal or heavy water ($\sim 90\%$ D₂O). Control of pH was obtained by titration of AH₂ solutions with 1.0 M and 4.2 M oxygen free NaOH solutions, which were standardized with 1.0 M HCl solution. A tight titration apparatus (illustrated in Figure 2) was needed to prevent oxygen from the air penetrating into the solution and forming radicals due to oxidation. Suprasil quartz cuvettes with stems were fitted via rubber tubing to Pyrex glass tube T-connections with two entrances. One entrance was fitted to a valve for evacuation and oxygen-free argon atmosphere refilling. The second entrance was a closed rubber septum that allowed for injection of solutions of ascorbic acid, HCl, or NaOH, respectively. In this way titrations in highly oxygen-poor environments could be performed (Figure 2) by adding ascorbic acid and/or NaOH solutions through the syringe and intensively shaking to obtain solutions of L(+)-sodium ascorbate and L(+)-disodium ascorbate (Na₂C₆H₆O₆, CAS Reg. No. 21547-69-3).

Visible light Raman measurements were carried out directly on the powders or on solutions in quartz cuvettes. We used a DILOR-XY Raman spectrometer with a macroscopic sampling stage from Horiba-Jobin-Yvon, Ltd. (Villeneuve d'Ascq, France) (79) and a green 532 nm wavelength Spectra-Physics (Santa Clara, CA, USA) Millennia module based on a 1,064 nm Nd:YVO₄ laser and a crystalline LiB₃O₅ frequency-doubling device for the excitation. A Supernotch-Plus filter (Kaiser Optical Systems, Inc., Ann Arbor, MI, USA) was used to filter off the Rayleigh scattered light. The spectrometer grating $10 \times 10 \text{ cm}^2$ had 1,800 grooves per millimeter and spherical mirrors in Czerny-Turner configuration (76) to disperse the remaining light onto a charge-coupled device detector cooled to 140 K with liquid nitrogen. The entrance slit was set to 200 μ m to obtain a resolution of about 8 cm⁻¹. Raman lines of liquid cyclohexane were used for calibration (80).

Ultraviolet Raman experiments were carried out with several UV laser lines and an InVia Reflex UV-Raman spectrometer from Renishaw plc (Gloucestershire, UK). The excitation light was filtered in a quartz prism monochromator with several slits and sent into the InVia instrument and through the attached microscope equipped with a traditional $15 \times$ UV lens objective. The laser output power was adjusted to about 5–15 mW, of which about 20% or less reached the sample surface. UV excitation light of 229 and 244 nm was obtained by frequency-doubling of continuous 457.9 and 488.0 nm light from a Lexel Ar⁺ gas laser (95-SHG-QS from Cambridge Laser Laboratories Inc., Fremont, CA) working by intracavity second harmonic generation in a β -Ba₂O₄ borate crystal (81). A solid-state FQCW 266-50 laser from CryLaS GmbH (Berlin, Germany) was used to obtain 266 nm. This laser emitted a highly stable continuous wave beam with a fixed wavelength generated as the fourth harmonic from a strong 1,064 nm laser (an Nd:YVO₄ crystal system in single frequency longitudinal mode operation, pumped by 808 nm from solid-state laser diodes and doubled in two stages). The light scattered from the sample was conducted from the microscope, through UV dielectric filters with long wavelength bandpass, dispersed in a single-stage spectrograph and detected with a UV-enhanced charge-coupled device detector, thermoelectrically cooled to a low temperature (-40° C). The entrance slit width was set to 40 μ m to obtain a resolution of ~8 cm⁻¹ (76). The acquisition time was 2 × 20 s with automatic removal of cosmic spikes. The spectra were scaled but not further corrected for monochromator and detector efficiencies. Other experimental details are given elsewhere (81).

The UV absorption spectra of the solutions were measured through Suprasil quartz cuvettes by use of a UV-Vis recording double beam spectrophotometer (UV-2401PC) from Shimadzu Corporation (Tokyo, Japan). The thickness *b* of the sample layer in the cuvette was 5 mm. The absorbance *A* was measured during titration and was used to obtain the molar absorptivity ε by applying Beer's law (A = ε *b c*, where *c* is the concentration) and taking the stepwise dilution into account.

Results and Discussion

The calculations on the structures and the corresponding IR and Raman spectra are presented in this section and then the spectral measurements follow. We also describe what occurs with the spectra when AH_2 molecules react with base.

Calculations—Structures

Data from solved crystal structures are listed in Table 1 together with the model calculation results. Selected bond lengths, bond angles, and dipole moments are given with obtained corresponding minimum energies at the DFT/B3LYP/6-311++G(d,p) level (the lower the minimum energy, the more stable the configuration). The calculations were performed for the AH₂ molecule, the AH⁻ anion, and the A²⁻ di-anion, all assumed to be in free gas-phase states (vacuum) as well as in hypothetical dissolved isotropic aqueous situations simulated by use of the polarizable continuum model (PCM) as implemented in the Gaussian program (60). Different starting point conformations were tried for the AH₂ species: First the average geometry of the Hvoslef crystal structure (10, 25) and then the "most stable molecule" identified by several researchers (e.g., Milanesio et al. (26), Singh et al. (53), and Yadav et al. (54)). For the AH⁻ and A²⁻ anions, the optimizations started from similar geometries, after leaving out the H3 proton for AH⁻ as well as the H2 proton for A²⁻, as shown in the Figures 3–10.

	ling			In water	(PCM) ^e	12	Figure 9	Figure 14	1.234	1.317	1.294	1.416	1.444	1.428	1.433	1.398	1.445	1.518	1.544	1.534		107.5	111.4	106.4	109.2	105.4	117.1	131.5	124.2	129.4	130.9	119.9	1135
$d A^{2-}$	²⁻ Di-ion model	Minimum	conformers	In water	(PCM)	11		Figure 8	1.225	1.302	1.351	1.383	1.447	1.402	1.432	1.373	1.482	1.496	1.547	1.546		108.1	111.5	104.1	111.9	104.2	119.1	129.4	123.6	132.3	130.2	117.9	112.4
2, AH ⁻ , an	¥.			In gas	phase	10			1.221	1.283	1.368	1.394	1.446	1.387	1.435	1.381	1.506	1.492	1.551	1.555		108.1	111.2	103.2	112.6	104.6	118.6	130.2	123.3	133.5	129.9	117.5	114.6
sults for AH	nodeling	unu	mers ^c	In water	(PCM) ^e	6	Figure 5	Figure 13	1.229	1.381	1.268	1.389	1.443	1.434	1.429	1.381	1.419	1.532	1.536	1.529		108.0	109.9	110.6	105.4	106.0	120.5	129.5	122.7	126.7	133.0	121.6	112.8
alculated ^a re	AH ⁻ Ion 1	Minir	confor	In gas	phase	8			1.226	1.383	1.264	1.395	1.439	1.433	1.431	1.383	1.421	1.538	1.535	1.531		107.4	110.0	110.9	104.6	106.9	121.2	128.8	120.5	128.5	134.7	120.7	112.3
ed to DFT c			NaAH or LiAH crvstal	X-ray	structure ^f	7			1.233(6)	1.384(5)	1.287(5)	1.358(5)	1.448(5)	1.410(5)	1.423(5)	1.373(6)	1.416(6)	1.516(6)	1.536(6)	1.503(6)	(。	108.0(3)	110.6(4)	109.5(4)	105.8(4)	105.2(4)	120.4(4)	129.0(4)	121.6(4)	128.7(4)	131.3(4)	122.9(4)	116.1(4)
ures compar		unu	mers	In water	(PCM) ^e	6	Figure 4	Figure 12	1.216	1.355	1.334	1.366	1.449	1.419	1.426	1.347	1.454	1.500	1.541	1.533	Angles (109.6	109.2	108.0	109.6	103.6	121.8	129.0	124.8	127.2	132.0	118.4	114.2
rrystal struct	lecule	Minir	confor	In gas	phased	5			1.204	1.355	1.343	1.377	1.450	1.413	1.424	1.340	1.457	1.499	1.543	1.538		109.5	108.3	108.8	109.5	103.9	123.6	128.1	123.1	128.1	131.3	119.3	114.9
a in solved c	AH_2 mo	er like	stal ^c	In water	(PCM)	4	Figure 3	Figure 11	1.205	1.355	1.350	1.364	1.445	1.419	1.428	1.338	1.464	1.502	1.540	1.525		109.7	108.6	108.5	109.3	104.0	124.5	126.9	122.9	128.6	131.2	119.5	114.4
structure dat		Conform	in crys	In gas	phase	3			1.205	1.355	1.350	1.364	1.445	1.419	1.428	1.338	1.464	1.502	1.540	1.525		109.7	108.6	108.5	109.3	104.0	124.5	126.9	122.9	128.6	131.2	119.5	114.4
Selected s			AH,	crystal	structure ^b	2			1.216(2)	1.361(2)	1.326(3)	1.355(2)	1.444(2)	1.427(3)	1.431(4)	1.338(2)	1.452(3)	1.493(3)	1.521(4)	1.521(3)		109.1(2)	109.5(2)	107.8(2)	109.5(2)	104.0(2)	121.4(3)	129.1(2)	124.6(2)	127.5(2)	133.5(2)	117.1(2)	114.8(2)
						Column 1	Structure spectra	Bond distances (Å)	C1-01	C2-02	C3-O3	C1-04	C4-04	C5-05	C6-06	C2-C3	C1-C2	C3-C4	C4-C5	C5-C6		C4-04-C1	04-C1-C2	C1-C2-C3	C2-C3-C4	C3-C4-04	04-C1-01	01-C1-C2	C1-C2-02	02-C2-C3	C2-C3-03	03-C3-C4	C3-C4-C5

 Table 1
 Table 1

					is set	++G(d n) hav	'B3LYP/6-311	n of the DET/	c hv annlicatic	in this worl	^a Calculated
0.0000104	0.0000100	0.0000057	0.0000056	0.0000186		0.0000041	0.0000041	0.0000117	0.0000058		Root mean square 3radient norm
-684.0658267	-684.0558907	-683.7741241	-684.5608318	-684.4801816		-685.0346989	-684.9945050^{d}	-684.9891598	-684.9891598		Total energy (Ha)
13.527	9.123	5.251	2.731	2.060		5.565	3.505 ^d	3.224	3.225		Dipole moment (D)
					162.1 (Li salt)					-151.60	C5-C6-06-H ₀ 6
-49.4	28.0	25.5	48.8	42.4	164.3	84.3	65.5	-168.4	-168.5	170.40	Angle χ_6
					–176.1 (Li salt)					64.75	C4-C5-C6-06
73.1	-159.3	-159.2	-177.4	-173.6	165.2	-72.4	-78.6	61.1	61.1	66.50	Angle χ_5
					– /0.0 (LI salt)					01.66-	с0н-со-со-но
31.2	28.3	28.8	39.5	36.7	-61.4	74.4	94.4	-71.0	-70.9	-67.60	Angle χ_4
					62.6 (Li salt)					50.80	C3-C4-C5-O5
-52.8	-49.6	-52.6	-59.6	-57.7	55.9	58.9	48.9	57.2	57.2	66.60	Angle X ₃
	$156.2)^{h}$	$160.7)^{h}$								-4.10	C2-C3-O3-H3
No H3	No H3 (H ₀ 5	No H3 (H _o 5	No H3	No H3	No H3	1.9	4.0	-3.5	-3.4	-3.80	Angle χ_2
					135.2 (Li salt)					-35.10	C1-C2-02-H2
No H2	No H2	No H2	0.3	0.2	-110.5	16.2	1.4	-0.6	-0.6	-2.90	Angle χ_1
					igle ^g (°)	Dihedral ar					
112.3	107.5	106.8	110.2	109.6	108.5(3)	114.0	111.4	107.3	107.3	108.0(2)	C5-C6-O6
110.6	109.3	108.8	109.7	109.2	108.5(4)	112.1	110.4	106.3	106.3	106.9(2)	D5-C5-C6
113.5	113.1	113.5	114.1	114.3	110.1(4)	114.2	111.9	111.6	111.6	112.7(2)	24-C5-C6
108.1	107.9	109.6	108.3	108.6	113.4(4)	111.7	111.6	112.8	112.8	111.7(2)	24-C5-O5
111.5	114.7	114.3	111.8	112.1	110.3(4)	111.1	110.3	110.7	110.7	110.4(2)	04-C4-C5

molecules, calculated by us (note that his coordinates were given in a left-handed coordinate system). ^cStarting from Hvoslef's average structures (Hvoslef (10) or Hvoslef (25)).

⁴Our values starting from most stable gas molecule. Similar results were found (with other definitions of atom numbers and signs) by Singh et al. (53) and Yadav et al. (54), who used Gaussian 03 and DFT/B3LYP with the smaller $6-31++G^{**}$ basis set. Their energy and dipole moment were given as -684.99328 Ha and 3.325 Debye.

^eThe global minima in the water-phase model (most stable conformations) are shown in boldface.

¹Diffraction results by Hvoslef (10). Dihedral angles were calculated by us (note that his coordinates were given in a left-handed coordinate system). Dihedral angles in italics are for the analogous Li⁺ salt that also has only one individual AH⁻ ion in its structure (31).

^gDihedral angles are defined in Figure 1.

^hHo5 is bound to O3 and the angle given is $\chi_2 = C2-C3-O3-H_05$.



Figure 3. Ascorbic acid molecule, AH_2 , as optimized by us starting from Hvoslef's average crystal structure (25). Calculations done with the Gaussian minimization procedure using DTF/B3LYP/G6-311++(d,p) with the water as solvent in the PCM solvation modeling. This is a local but not the global minimum of the single-molecule.

The values for calculated geometries, bonds, and angles (see Table 1) differ somewhat from the crystal structure values and depend on the kind of model used and the starting point. The problem of the calculation ending in a nonglobal minimum also needs to be addressed, as pointed out by many of the previous researchers.

For the AH_2 molecule optimizations, starting from an average conformation as in the crystal structure, there was hardly any influence of the kind of model (gas phase versus PCM). Specifically, the obtained minimum geometry (see Figure 3) did not depend on the presence or absence of the solvent (Table 1, columns 3–4), but the result did not give the global minimum. The most stable molecule could only be obtained starting from other guessed conformations. Moreover, the resulting minimum conformation was quite dependent on the kind of modeling used, the environment being either a vacuum or a PCM continuum; see Table 1, columns 5–6. As seen in Figure 4, the most stable AH_2 conformer (the global minimum) has five hydrogen bond interactions and does not look like the molecules in the crystal structure. The molecule in the "crystal" conformation, when alone as seen in Figure 3, forms only three intramolecular hydrogen bond interactions and is only in a local minimum state. In the crystal the stable phase in reality. Data for the obtained most



Figure 4. Most stable structure of the ascorbic acid molecule, AH_2 , as optimized by Gaussian minimization with the DTF/B3LYP/G6-311++(d,p) procedure and using water as the solvent in the PCM solvation modeling. The structure is reminiscent of that found in the gas phase by Milanesio et al. (26), Singh et al. (53), and Yadav et al. (54).

Intramol	scular nyul	ogen pond	uistances l	OT AH2, A	н, апа А					manons	
	AH ₂ m	olecule			AH ⁻ (anion			A ²⁻ ai	nion	
Conforn in crv	ner like _{stal} b	Other mi	inimum	Startin	g from Per like	Other mi	nimum ^b	Starting	g from ter like	Other mir	himum bare ^b
m ci)	Stat	COLLION		CONTOL		COLLOI		COIIIOIII		COLLOLI	1013
In	In	In	II	$Hvoslef^{b}$	Yadav ^c	In	II	$Hvoslef^{b}$	$Yadav^{c}$	In	In
gas	water	gas	water	In water	In water	gas	water	In water	In water	gas	water
phase	(PCM)	phase ^c	(PCM) ^d	(PCM)	(PCM)	phase	(PCM) ^d	(PCM) ^d	(PCM)	phase	(PCM)
2	3	4	5	9	7	8	6	10	11	12	13
	Figure 3		Figure 4				Figure 5	Figure 9			
	Figure 11		Figure 12	Figure 6	Figure 7		Figure 13	Figure 14	Figure 10		Figure 8
2.484	2.484	2.516	2.655	2.528	2.525	2.398	2.515	No H2	No H2	No H2	No H2
2.725	2.725	2.714	2.751	No H3	No H3	No H3	No H3	No H3	No H3	No H3	No H3
3.083	3.083			1.826		1.795	1.873	1.727		1.008	1.059
				0.982				0.993		1.687	1.473
				2.724				2.658		2.632	2.487
		2.944	2.676		2.852						
		2.170	2.697	2.095	2.061			2.009	1.918		
						2.187	2.319			1.800	1.889
											0.982
											2.600
		2.178	2.541		2.274						
								0.969			
-684.9891598	-684.9891598	-684.9945050	-685.0346989	-684.5596177	-684.5588461	-684.4801816	-684.5608318	-684.0658267	-684.0613658	-683.7741241 -	-684.0558907
in this work n Hvoslef's	by applicati average stru	on of the DH totures; see H	FT/ B3LYP/ Hooslef (10)	5-311++G(or Hvoslef	d,p) basis se (25).	st.					
	Intramold Conform in cry in cry in cry in cry in cry ass phase 2 2 2.725 3.083 3.083 3.083 3.083 1.003 1.003	Intramolecular hydiAH2 mConformer likeInInInInInIngaswaterphase(PCM)232323233.0833.0833.0833.0839.0833.08310 this work by applicatiin this work by applicatein Hvoslef's average strue	Intramolecular hydrogen bondAH2 moleculeAH2 moleculeConformer likeOther mInInInInInInInIngasphaseconfophase(PCM)phasephase(PCM)phase234Figure 3Figure 3A2.4842.5162.7252.7252.7143.0833.0833.0832.4842.4842.1702.4842.3032.1702.4842.7252.7142.4842.7252.7143.0833.0833.083a.0833.0832.170in this work by application of the DFm Hvoslef's average structures; see F	Intramolecular hydrogen bond distances 1AH2 moleculeAH2 moleculeConformer in crystalbConformerInInInInin crystalbOuther minimumin crystalbOuther minimumin crystalbOuther minimumin crystalbOuther minimumin crystalbOuther minimumgaswatergaswaterphase(PCM)phase(PCM)phase(PCM)phase(PCM)gas3.0833.0833.0833.0833.0833.0833.0833.0833.0832.7442.5162.7552.7142.7512.6762.7532.1702.6842.644991598-684.9945050in this work by application of the DFT/ B3LYP(M)m Hvoslef's average structures; see Hvoslef (10)	Intramolecular hydrogen bond distances for AH2, A AH2 molecule AH2 molecule Conformer like Other minimum starting Conformer conformer conformer In In In In woster Starting in crystal ^b 0 ther minimum starting conformer in crystal ^b 0 the minimum starting conformer 2 3 4 5 6 Figure 11 Figure 12 Figure 6 2.724 2.735 2.735 2.714 2.724 2.724 3.083 3.083 3.083 0.882 0.882 2.744 2.444 2.676 2.724 2.724 2.735 2.714 2.697 2.095 2.724 2.170 2.697 2.097 2.095 2.724 2.173 2.170 2.697 2.095 2.724 2.44 2.644 2.644 2.744 2.744 2.944 2.697 2.095 2.724 <td>Intramolecular hydrogen bond distances for AH2, AH², and AAH2AH2and AAH2AH2moleculeAH⁻¹AH2AH2moleculeAH⁻¹Conformer likeOther minimumStarting fromin crystal^bconformer likeNadav⁶gaswaterBasNaderphase(PCM)ProvideProvidRemoterRemoterphase(PCM)ProvideAH42.4842.5162.6552.5232.7352.7142.751NoH33.0833.0833.0832.9442.6762.4842.751NoH3NoH33.0833.0833.0832.7242.7442.4842.6762.5282.5252.7452.7142.751NoH3NoH3NoH3NoH3NoH3NoH33.0833.0833.0832.0742.8552.1702.0972.0952.0612.1702.6972.0952.0742.1782.5412.7342.73418.001598-684.9991598-684.9991598-684.9891598-684.9991598-684.999150916.44.9891598-684.9991504-685.634609916.44.9891598-684.9991504-685.634609916.44.9891598-684.9991504-685.634609916.44.9891598-684.9991508-684.9991508-684.9891598-684.9945050-685.634609916.44.9891598-684.9945050-685.</td> <td>Intramolecular hydrogen bond distances for AH2, AH⁻, and A⁻ molecultAltramolecular hydrogen bond distances for AH2, AH⁻, and A⁻ molecultAltramoleculeAH⁻ moleculeAltramoleculeAth- anionConformer likeOther minimumIn<td>$\begin{tabular}{ c c c c c c c c c c c c c c c c c c c$</td><td>$\begin{array}{$</td><td>$\begin{tabular}{ c c c c c c c c c c c c c c c c c c c$</td><td>$\begin{array}{ c c c c c c c c c c c c c c c c c c c$</td></td>	Intramolecular hydrogen bond distances for AH2, AH ² , and AAH2AH2and AAH2AH2moleculeAH ⁻¹ AH2AH2moleculeAH ⁻¹ Conformer likeOther minimumStarting fromin crystal ^b conformer likeNadav ⁶ gaswaterBasNaderphase(PCM)ProvideProvidRemoterRemoterphase(PCM)ProvideAH42.4842.5162.6552.5232.7352.7142.751NoH33.0833.0833.0832.9442.6762.4842.751NoH3NoH33.0833.0833.0832.7242.7442.4842.6762.5282.5252.7452.7142.751NoH3NoH3NoH3NoH3NoH3NoH33.0833.0833.0832.0742.8552.1702.0972.0952.0612.1702.6972.0952.0742.1782.5412.7342.73418.001598-684.9991598-684.9991598-684.9891598-684.9991598-684.999150916.44.9891598-684.9991504-685.634609916.44.9891598-684.9991504-685.634609916.44.9891598-684.9991504-685.634609916.44.9891598-684.9991508-684.9991508-684.9891598-684.9945050-685.634609916.44.9891598-684.9945050-685.	Intramolecular hydrogen bond distances for AH2, AH ⁻ , and A ⁻ molecultAltramolecular hydrogen bond distances for AH2, AH ⁻ , and A ⁻ molecultAltramoleculeAH ⁻ moleculeAltramoleculeAth- anionConformer likeOther minimumIn <td>$\begin{tabular}{ c c c c c c c c c c c c c c c c c c c$</td> <td>$\begin{array}{$</td> <td>$\begin{tabular}{ c c c c c c c c c c c c c c c c c c c$</td> <td>$\begin{array}{ c c c c c c c c c c c c c c c c c c c$</td>	$\begin{tabular}{ c c c c c c c c c c c c c c c c c c c$	$ \begin{array}{ $	$\begin{tabular}{ c c c c c c c c c c c c c c c c c c c$	$ \begin{array}{ c c c c c c c c c c c c c c c c c c c$

¢ Table 2

^eOur values starting from most stable gas molecule. Quite similar results were found (with other definitions of atom numbers and signs) by Singh et al. (53) and Yadav et al. (54), who used Gaussian 03 and DFT/B3LYP with the $6-31++G^{**}$ basis set. Their energy and dipole moment were given as -684.99328 Ha and 3.325 Debye. ^dThe global minima in the water-phase model (most stable conformations) are shown in boldface.



Figure 5. Optimized structure of the AH^- ion in the PCM water model as obtained after Gaussian minimization based on our guess. This global minimum conformer of AH^- has three internal hydrogen bonds (Table 2).

stable hydrogen bonds and the corresponding energies are summarized in Table 2 (columns 2–5).

For the AH⁻ anion it was possible in a similar way among the calculated results to find low energy conformers with 3–4 intramolecular hydrogen bond interactions (see Figures 5–7 and Table 2). In particular, the global minimum energy conformer (Figure 5) has an interesting geometry with protons bound as $O1\cdots$ H2–O2, $O3\cdots$ Ho5–O5, and O5…Ho6–O6. This conformation deviates markedly from the geometry found in the crystal structures of several known ascorbate salts. A brief summary of the geometries of these crystal structures is given in Table 3. The reason for the different values found for the



Figure 6. Optimized structure of the AH^- ion in the PCM water model as obtained after Gaussian minimization based on the crystal structures solved by Hvoslef (10). This local minimum conformer of AH^- has three internal hydrogen interactions but the energy is not as low as that of Figure 5 (Table 2).

	0	N N N N N N	se	lected crystal si	tructures ^a		I		20
Dihedral angles in	Ascorbate	Lithium	Sodium						
degrees, defined in	minimum	ascorbate	ascorbate	Calcium di-	-ascorbate	Strontiu	n diascorbate	Thallium	ascorbate
Figure 1	conformer	(31)	(10)	dihydrate	(27–29)	dihy	drate (30)	(3	(2)
χ ₁ C1–C2–O2–H2	0.3	135.2	-110.5	-154.8	-118.3	-142.3	-120.7	Missing	Missing
χ ₃ C3–C4–C5–O5	-59.6	62.6	55.9	169.9	-69.2	170.4	-69.9	55.6	61.1
χ ₄ C4–C5–O5–H ₀ 5	39.5	-76.6	-61.4	-39.9	-47.6	-44.5	-58.7	Missing	Missing
χ ₅ C4-C5-C6-O6	-177.4	-176.1	165.2	-74.4	-73.2	-70.6	-72.5	166.5	64.0
χ ₆ C5–C6–O6–H ₀ 6	48.8	162.1	164.3	Missing	141.2	97.9	135.6	Missing	Missing
^a The angles were calc	sulated by us us	ing data from	the CCSD data	base and their M	lercury program	I. Data for som	e hydrogen atoms :	are missing in th	e database

Table 3	dihedral (torsion) angles χ_1 , $\chi_3 - \chi_6$ for the ascorbate ion minimum energy conformer (Figure 5) compared with similar 3	selected crystal structures ^a
	Calculated dihedral	



Figure 7. Optimized structure of the AH^- ion in the PCM water model as obtained after Gaussian minimization based on the results of Singh et al. (53) and Yadav et al. (54) for AH_2 . This local minimum conformer of AH^- has four internal hydrogen interactions but the energy is not as low as that of Figure 5 (Table 2).

torsional angles (defined in Figure 1) is that the realized energy minimum in each crystal depends on the particular interactions in that crystal. In contrast, the global minimum conformation of our single molecular AH^- anion looked different, independent of whether the environment was modeled in the gas phase or by the PCM (compare columns 8–9 in Table 1). This puts a limit on the applicability of our simple one-molecule modeling results with respect to predicting the excact properties of the different crystals.

Finally, for the A^{2-} di-anion, low energy conformers with intramolecular hydrogen bond interactions were found in the gas phase and in the aqueous PCM environment models. In the gas phase, the global minimum energy conformation was not particulary distinct: Different conformations gave energies not much different (an example is given for the gas phase in Table 2, column 12). When the modeling was done via PCM, the situation



Figure 8. Optimized structure of the A^{2-} ion in the PCM water model as obtained after Gaussian minimization based on a guess. This local minimum conformation of A^{2-} , rather similar to a similar gas-phase conformer (Table 1), has two internal hydrogen interactions but the energy is not as low as that in Figure 9 (Table 2).



Figure 9. Optimized structure of the A^{2-} ion in the PCM water model as obtained after Gaussian minimization starting from a guessed structure derived from the crystal results by Hvoslef (10, 25). This global minimum conformer of A^{2-} has two internal hydrogen bonds (Table 2).

was about the same. Selected results are shown in Table 1 (columns 10–12) and Table 2 (columns 10–13) and corresponding structures are depicted in Figures 8–10. One PCM minimum conformer (depicted in Figure 8, and rather similar to the gas-phase conformer; see Table 1, columns 10–11) has protons bound as $O3\cdotsHo5-O5$ and $O5\cdotsHo6-O6$ (Table 2, column 13), but this is not the global minimum conformer; that honor must be shifted to the conformer shown in Figure 9. The conformer in Figure 9 presumably owes its stability to its two short internal hydrogen bond interactions $O3\ldots.Ho5\ldots.O5$ and $O4\ldots.Ho6\ldots.O6$. In addition, the conformer in Figure 9 is more stable than the conformer depicted in Figure 10, which only has a single $O4\cdotsHo6-O6$ interaction and some additional stabilization from its Ho5 placed centrally above the resonance ring system, at a distance of 2.38 Å from C3.



Figure 10. Optimized structure of the A^{2-} ion in the PCM water model as obtained after Gaussian minimization based on a guessed structure derived from the results for AH2 of Singh et al. (53) and Yadav et al. (54). This local minimum conformer of A^{2-} has one internal hydrogen interaction and the Ho5 is interacting with the π electrons above C3, but the stabilization is not as good as in the conformer of Figure 9 (Table 2).



Figure 11. Raman and infrared spectra calculated for the conformer of ascorbic acid AH₂ simulated in water solution for the local minimum conformation as shown in Figure 3.

When trying to understand the behavior of ascorbate salts in solution it is worth remembering that the O2 and O3 charges seem to favor the formation of anion dimers via hydrogen bonding, and this tendency tends to complicate things further, as discussed in the literature (1, 12, 28, 34–37). These features have not been studied here.

Calculations—Spectra

Infrared absorption and Raman scattering spectra were calculated for the different conformations of AH_2 , AH^- , and A^{2-} . Selected calculated spectra are shown in Figure 11–14 and the data are summarized in Table 4, to facilitate mutual comparisons. Selected modeling results for ions AH^- and A^{2-} are detailed in Table 5 with emphasis on the dihedral angles and the O-H modes.

By comparing spectra in Figures 11 and 12, calculated for the water medium (PCM) as an example, we clearly see differencies: Strong Raman bands are predicted for the "crystal conformation" (in Figure 3), at OH⁻ stretchings at 3855, 3796, and 3752 cm⁻¹, whereas analogous bands are predicted at 3500, 3359, and 3307 cm⁻¹ for the global minimum conformer (see Figure 4). Accordingly, the AH₂ spectra are significantly conformer dependent, as also expected. The same was true for the gas-phase results (not shown). The reason is that differences in the conformation geometry are substantially reflected in the couplings among the vibrational modes, leading to noticeable differencies in frequencies and intensities. This is similar to what has been found for other molecules such as amphetamine (82, 83) or for ionic liquids (84, 85).

					Tab	le 4						
Selected wavenumber	r data ^a for]	IR absorp	tion and Ra	aman scatte y the Gauss	ring stron sian 03/DH	g bands of FT/B3LYP/	AH ₂ , AH ⁻ (6-311++C	, and A ^{2–} J(d,p)/PCI	compounds M	s dissolved i	in water. D	ata were
Modeling			V	H ₂				-HA			A^{2-}	
	Confo	rmer like in c	rystal	Global r	ninimum conf	ormer ^b	Global	minimum cor	former	Global r	minimum confe	ormer
Total energy (Ha)		-684.9891598			-685.0346989			-684.5608318			-684.0658267	
	Str Sp	ructure Figure bectra Figure 1	1	Str	ucture Figure bectra Figure 1	2	N N	ructure Figure pectra Figure	5	Str	ucture Figure 9 ectra Figure 14	
Assignment ^b Brief description of normal modes	Mode no. (cm ⁻¹) ^c	IR (km/mol)	Raman (Å ⁴ /AMU)	Mode no. (cm ⁻¹)	IR (km/mol)	Raman (Å ⁴ /AMU)	Mode no. (cm ⁻¹)	IR (km/mol)	Raman (Å ⁴ /AMU)	Mode no. (cm ⁻¹)	IR (km/mol)	Raman (Å ⁴ /AMU)
O6–H str	54: 3861	57	122	54: 3505 ^d	250 ^d	165 ^d	50: 3771	83	105	48: 3708	260	145
O5–H str	53: 3842	33	63	53: 3491	209	126	49: 3455	536	179	47:3206	1003	201
O3-H str	52: 3796	101	100	51: 3307	563	286						
O2–H str	51: 3753	120	81	52: 3360	515	300	51:3776	117	210			
C6–H asym str	50: 3072	26	50									
C4–H str	49: 3069	ю	131									
C6–H str				50: 3075	29	267	48:3104	36	147	46:3086	71	246
C6–H sym str	48: 3023	41	141	49: 3013	50	574						
C5-H str	47: 3012	16	91	48: 2994	1	253						
C4-H str				47: 2978	3	412						
C4-H & C5-H str ooph							46: 3043	50	172			
C6–H str							45:3010	48	284			
C4-H, C5-H, C6-H str										45:3016	241	823
C4-H, C5-H, C6-H str										44: 3005	75	582
C4-H, C5-H, C6-H str										43:2997	39	287
C1 C2 C3 ring breath	46: 1829	367	18	46A: 1784	266	6	44A: 1746	203	57	42A: 1699	602	66
C1-01, C2-C3 str	45: 1775	432	98	45B: 1699	1610	469						
C3-C4, C1-C2 str, ring OH	42: 1440	48	7	43C: 1433	103	27						
bend												
C1-01, C2-C3, C2-02 str,							43 <i>B</i> : 1604	2426	68	41:1550	454	62
03-H-05 str												
C-H bend	44: 1521	9	7				41:1474	307	37			
C6H ₂ sci, O3–H–O5 bend										40: 1498	454	101
C5-C6 str, C-H bend	43: 1458	Э	1									
C-H, O-H bend, C-C str	41: 1399	7	1							39B: 1495	1522	273
C5-C6 str, C6-O6-H bend										38: 1432	40	12
C2-02 & C3-03 str							39 <i>C</i> : 1415	568	112			
											(сопппиеа о	n nexi page)

Selected wavenumbe	r data ^a for	IR absorp calculat	tion and R ed by the (aman scatte Gaussian ()	Tab Pring stron 3/DFT/B3	le 4 g bands of LYP/6-311	AH ₂ , AH ⁻ ++G(d.p))	, and A ^{2–} PCM (<i>co</i>)	compound: <i>utinued</i>)	s dissolved	in water. D	ata were
Modeling			, ,	VH ₂				_HA	Ì		A^{2-}	
	Confe	ormer like in c	rystal	Global	minimum conf	former ^b	Global	minimum cor	former	Global	minimum conf	ormer
Total energy (Ha)		-684.9891598			-685.0346989			-684.560831			-684.0658267	
	S SI	ructure Figure	1	SS	ructure Figure pectra Figure 1	4 0	S S	ructure Figure pectra Figure	5	Str Sp	ucture Figure ectra Figure 1	6 4
Assignment ^b Brief description of normal modes	Mode no. (cm ⁻¹) ^c	IR (km/mol)	Raman (Å ⁴ /AMU)	Mode no. (cm ⁻¹)	IR (km/mol)	Raman (Å ⁴ /AMU)	Mode no. (cm ⁻¹)	IR (km/mol)	Raman (Å ⁴ /AMU)	Mode no. (cm ⁻¹)	IR (km/mol)	Raman (Å ⁴ /AMU)
C3–C4 str, C–H bend	40: 1392	16	8							36C: 1369	560	100
C1-C2, C3-C4 str, ring OH	39: 1364	20	4									
bend												
C2-O2, C4-C5 str	38: 1331	103	13	37: 1330	43	4						
C3-O3 str, H4 def	37: 1309	161	1									
C-H, O-H bend	36: 1295	9	10	36: 1295	514	16				37: 1396	32	20
C-H, O-H bend	35: 1284	58	17	35: 1285	134	9	35: 1324	150	1	34: 1321	355	109
C1-C2, C2-O2 & C3-O3										35: 1345	389	8
str												
C3-C4 str, C-H bend							33: 1255	81	52	32: 1272	294	88
C-H, O-H bend	34:1248	8	4	34: 1247	16	42				33: 1304	222	12
C–H bend	33: 1185	<i>LT</i>	1	33: 1207	65	10				30:1193	112	17
C1-C2, C3-O3, C3-C4 str	32: 1171	141	6	32: 1169	253	30				31:1232	83	17
C1-04, C3-C4, C2-02 str	31: 1126	195	8	31:1110	446	51						
C4-C5, C5-O5, C4-O4 str	30: 1118	90	6	29:1090	95	13				27:1067	5	11
C2-02, C4-04 & C1-04	29: 1075	65	1				29:1089	278	18			
str												
C1-C2, C4-C5, C5-C6 str										29:1151	106	18
C6H ₂ rock, C4-C5 str										28:1103	271	15
04-C4, C5-C6, C6-O6 str	28:1067	55	4	30:1099	242	20						
C1-04, C3-C4, C6-06 str	27:1043	76	ŝ									
O3–H–O5 bend												
C5-C6 str	26:1005	11	5									
C6-O6 & C5-O5 str							26:1036	201	7	24:1028	232	4
C4-04 & C5-05 str	25:991	41	2	27: 1036	159	ε	25:1030	353	7	26:1057	111	ε
C4-04 & C6-06 str				26: 962	47	S				25:1032	164	15
C4-C5 & C2-02 str C5-05 str	24: 866	15	б				23 · 860	76	Ŷ	22 · 871	6	15
							10.000	2	,	1.0.11	`	2

ŝ									
S							20: 759	151	2
	19: 632	10	21				19: 729	14	9
13	18:601	15	10				18: 716	15	26
	17:572	23	14				17: 688	25	13
							16: 607	16	б
	15:403	384	ю	13: 420	145	2			
							15: 578	20	49
	14: 377	98	1						
1	12: 355	169	7				11: 391	39	
							9: 351	30	5
1									
							7: 324	8	19
	5: 149	200	2	7:217	123	1			
							5: 223	35	5
	5: 149	200	7	1: 217	173	-	5: 223	35	
le 6-311++	-G(d,p) basis	set. For st	ructures se	e Table 1.	- - -	•	-	-	
g	1 1 6-311++	14: 377 1 12: 355 1 5: 149 6-311++G(d,p) basis	14: 377 98 1 12: 355 169 1 5: 149 200 6-311++G(d,p) basis set. For st	14: 377 98 1 1 12: 355 169 2 1 5: 149 200 2 6-311++G(d,p) basis set. For structures set	14: 377 98 1 1 12: 355 169 2 1 5: 149 200 2 7: 217 6-311++G(d,p) basis set. For structures see Table 1.	14: 377 98 1 1 12: 355 169 2 1 5: 149 200 2 7: 217 123 6-311++G(d,p) basis set. For structures see Table 1. 6.31 1 1	14: 377 98 1 1 12: 355 169 2 1 12: 355 169 2 5: 149 200 2 7: 217 123 6-311++G(d,p) basis set. For structures see Table 1. 6 6 6	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$\begin{array}{cccccccccccccccccccccccccccccccccccc$

"Abbreviations for approximate vibrations: asym = asymmetric, bend = bending, breath = breathing, def = deformation, iph = in phase, ooph = out of phase, rock = rocking, sci = scissoring, str = stretching, sym = symmetric, tors = torsion (rotation). For modes A, B, and C see text and Figure 25. "See also values calculated by Shimada et al. (46) (reported in their table 2, column 5)." ^oFor example, for mode 54 (03–H3 stretching) in the gas medium, Singh et al. (53) and Yadav et al. (54) got a wavenumber of 3798 and IR and Raman signals

of 109 km/mol and 101 Å⁴/AMU. We got identical values for that medium.

 Figures Figures Dihedral angle χ1 C1-C2-O2-H2 Dihedral angle χ2 C2-C3-O3-H3 Dihedral angle χ3 C3-C4-C5-O5 Dihedral angle χ5 C4-C5-C6-O6 Dihedral angle χ6 C5-C6-O6-H06 Dipole moment 	Local minimum, starting from Hvoslef ^c Structure Figure 6 1.2 No H3 -54.9 35.4 75.5 -52.0 2.258	Selected model result AH ⁻ ion conformers Local minimum, starting from Yadav ^d Structure Figure 7 -0.2 No H3 52.0 85.6 -74.1 57.4 7.006	s for ions AH ⁻ and A Global minimum Structure Figure 5 Spectra Figure 13 0.3 No H3 -59.6 39.5 -177.4 48.8 2.731	 ⁽²⁻ in water (PCM)^{al,} Global minimum, starting from Hvoslef ^c Structure Figure 9 Spectra Figure 9 Spectra Figure 9 Spectra Figure 9 Spectra Figure 9 31.2 31.2 73.1 -49.4 13.527 	A ²⁻ di-ion conformers A ²⁻ di-ion conformers Local minimum, starting from Yadav ^d Structure Figure 10 No H2 No H3 53.0 -26.3 -63.1 34.2 16.076	Other local minimum Structure Figure 8 No H2 No H3 (Ho5 156.2)° -49.6 -49.6 28.3 -159.3 28.3 -159.3 9.123
(Debye) Total energy (Ha) Root mean square gradient norm	-684.5 <u>59</u> 6177 0.0000061	-684.5588461 0.0000022	-684.5 <u>60</u> 8318 0.0000056	$-684.0\underline{658}267$ 0.0000104	-684.0 <u>61</u> 3658 0.0000197	-684.0558907 0.0000100

Table 5

OH str mode no.	51:	51:	51:	48:	48:	48:
Wavenumber	3778	3780	3776	3708	3708	3708
Intensity	IR 118, Ra 199	IR 111, Ra 207	IR 117, Ra 210	IR 260, Ra 146	IR 255, Ra 408	IR 260, Ra 145
Assignment ^f	02–H2 str	$05-H_{0}5$ str +	O2-H2 str	06-H ₀ 604	$05-H_05\dots 03 \text{ str}^{g}$	O6-H5 str
		02-H2 str		str^g		
OH str mode no.	50:	50:	50:	47:	47:	47:
Wavenumber	3760	3766	3771	3206	3673	3206
Intensity	IR 182, Ra 124	IR 95, Ra 77	IR 83 Ra 105	IR 1003 Ra 210	IR 384 Ra 169	IR 1003 Ra 201
Assignment ^f	$06-H_0604+\pi$	$05-H_{0}5$ str +	O6-H ₀ 6 str	05-H ₀ 503	$06-H_0604+\pi$	O5-H5 str
	str^{g}	$06-H_0604+\pi$		str^g	$\mathrm{Str}^{\mathrm{g}}$	
		Str ^g				
OH str mode no.	49:	49:	49:	No H2 present	No H2 present	No H2 present
Wavenumber	3432	3734	3455			
Intensity	IR 611, Ra 179	IR 200, Ra 145	IR 536, Ra 179			
Assignment ^f	05-H ₀ 503	$06-H_0604+\pi^g$	05-H ₀ 503			
	str^g		str ^g			
^a Calculated in this	work by application of t	he DFT/B3LYP/6-311++	-G(d,p) model. For stru	icture data see Table 1		

	fa
_:	ple
a)	ĕ
Ŕ	Ē
[a]	5
ີ່ວ່	F
š	ģ
en En	S
at	Б
p	a
E	es
Ë	lu
ğ	va
Ä	Я
÷	н
£	Ξ
Ξ	Ξ
[e]	Ē.
ö	Ξ
Ξ	ba
$\widehat{\mathbf{a}}$	lo]
Ŧ	Ο
ž	s
4	ee
T	Ĕ
는	ĩ
2	ž
Ľ.	÷Ξ
×	E
Κ	8
1	ĕ
8	50
Ξ	aı
£	al
D	dr.
ē	e
th	1
JC	Ч.
ã	-
. <u>e</u>	Ŀ
at	5
E:	Ē
ď	
ap	Ξ
N	Ğ
-0	jĽ,
겊	ef
20	ç
s	μĘ
Ē	S
1 tl	<u>ľ</u>
Ξ.	p
Вd	ar
atƙ	al
Ë	đ
<u>c</u>	<u>j</u> e
а	

^bDihedral angles are defined in Figure 1. Dihedral angles are in degrees. Oroval multiplication μ_{ab} , μ_{ab}

^gInfluenced by hydrogen bonding.



Figure 12. Raman and infrared spectra calculated for the most stable ascorbic acid AH_2 simulated in water solution for the conformation as shown in Figure 4.

Considerable changes also appeared in the spectra when the proton H3 was removed to simulate the ascorbate anion formation (see Figures 12 and 13). We compared the global minima conformers found for the water medium (PCM modeling) shown in Figures 4 and 5, trying to understand this quite complex process. When the H3 is lost, apparently Ho5 after energy minimization takes over the interaction with O3 (see Table 2). In Table 4 for AH₂ we see that three strong OH stretching Raman bands at 3500 cm⁻¹ (from O5 and O6), 3359 cm⁻¹ (from O2), and 3307 cm⁻¹ (from O3) are transformed to two strong OH stretching bands for the AH⁻ ascorbate anion: A high-frequency band at 3772 cm⁻¹ (from O2 and O6) and one at 3455 cm⁻¹ (from O5; see also Table 5).

This quite complicated situation is carried further on when another proton (H2) is removed (from the ascorbate anion to form the ascorbate di-anion); see the A^{2-} global minimum conformer shown in Figure 9. The spectrum, shown in Figure 14, now contains only two strong OH stretching Raman bands, predicted at 3708 cm⁻¹ (from O6) and 3206 cm⁻¹ (from O5) in the PCM water solution simulation (see Tables 4 and 5).

Comparisons between Experimental Measurements and DFT Results

Reference Raman spectra for AH_2 and NaAH solids were measured for comparison with both the previous literature and the calculations. As always, the spectra consist of strong, medium, and weak intensity peaks corresponding to the vibrational transitions in the molecules (shown in Figure 15). In Table 6 the measured Raman shifts are compared to literature values and the bands are assigned to transitions between various levels corresponding to the respective stretching, bending, and deformation vibrations. As one can see in Table 6, band positions of the literature spectra and the measured values are often in



Figure 13. Raman and infrared spectra calculated for the ascorbate ion AH⁻ simulated in water solution for the conformation as shown in Figure 5.



Figure 14. Raman and infrared spectra calculated for the ascorbate di-anion A^{2-} simulated in water solution for the conformation as shown in Figure 9.

	Ascorbic	acid			Sodium	ascorbate	
Hvoslef and Klæboe (39)	Panicker et al. (45)	De Gelder et al. (44)	Shimada et al. (46)	Saraiva et al. 300K (48)	This work Figure 15, Figure 19	Hvoslef and Klæboe (39)	Assignments
			3526	3535			O-H str
			3410	3419	3304 m		O-H str
				3324	3247 w		O-H str
							O-H str
3002 m	3004 w		3003	3010			0-H + C-H str
2978 m				2985	2979 m	2978 s	C-H str
					2962 m	2965 m	C-H str
						2954 m	
					2950 m	2940 w	C-H str
2916 vs	2919 s		2917	2924	2924 vs	2925 vs	C-H str
2903 vs							
2866 w	2879 wsh			2870	2892 w		C-H str
1754 w	1758 w			1755	1703 s	1704 vs	$C=C+C=O \operatorname{str} A$
1667 vs	1661 vvs	1667 vs	1667	1674	1598 s	1598 vsbr	$C=C+C=O \operatorname{str} E$
1654 vs		1653 vs	1652	1659			C=C str
1498 m	1484 m	1498 m	1497	1501			CH ₂ sci
1487 w				1464	1483 w	1482 s	CH bend
					1359 vw	1360 m	
1450 w	1452 w			1452	1330 w	1330 s	CH ₂ bend (sci)
1372 w	1371 w		1371	1375	1305 w	1305 s	ring def, CH ₂ wag
1344 vw							C-O-H bend
1321 s	1323 s	1319 s		1324	1277w	1275 s	CH ₂ bend (wag) C

Table 6

1247 wbr	1236 m C–O–H bend (twi)	1156 m	1127 m C–C(–O)–O str	C–O–C str, ring def		1077 s C-O-C str, C-O-H	bend	1048 s C–O–C str	1015 s ring O–H bend	C–H and O–H bend	935 s	888 s C4–C5 ring str	832 vs C–C ring str	OH oopl def	757 m OH oop1 def/ring	def		719 m OH oopl def/ring def	707 sbr OH oopl def/ring def	653 vs OH oopl def/C–C	604 vw ring str	(continued on next page)
1252 vw	1236 vw	1155 vw	1126 w			1078 w		1048 m	1024 w		937 m	888 w	$829\mathrm{m}$		758 vw			718 w	704 w	653 m		
1300	1261	1227	1202	1133	1121	1073		1037	1029	995		876	825	745	713			631	590	570		
	1256			1129					1025			870	821					628	588	566		
1296 m	1256 s			1130 vs					1025 msh				820 s	710 m			696 m	629 s	588 m	566 s		
	1258 s		1193 w	1113 s		1081 w		1048 m		984 w		871 m	823 m	742 sh			693 m	621 s	581 w	564 m		
1297 m	1258 s	1226 w	1200 m	1131 vs	1114 vw	1066 m		1039 vw	1026 s	993 w		872 m	822 s	742 w	711 m		694 w	629 s	589 s	567 s		
1295 w	1256 s	1226 w	1199 w	1128 s		1066 w			1028 m	992 w		871 w	820 m	742 vw	706 m br			629 s	588 w	566 m		

		Ascorbic	acid			Sodium asc	corbate	
This work Figure 15, Figure 16	Hvoslef and Klæboe (39)	Panicker et al. (45)	De Gelder et al. (44)	Shimada et al. (46)	Saraiva et al. 300K (48)	This work Figure 15, Figure 19	Hvoslef and Klæboe (39)	Assignments
492 vw	490 w					588 w	585 m	
476 vw	477 m	468 w			478	497 vw	496 w	C-O ipl def/ring def
						427 vw	427 m	
448 w	448 s	452 w	448		451	385 vw	383 vw	O-H wag
								C–O ipl def
362 w	363 m		363		365	366 vw	369 vw	O-H wag
343 w	345 m		344		348	339 vw	338 s	O-H wag
						304 vw	304 m	
295 vw	295 m		296		297	267 vw	266 m	C–O bend
271 vw	273 w		256		267	225 vw	227 w	O wag (ring)
257 w	259 m				256	190 vw	194 vw	
224 w	223 m		223		224	175 vw	177 vw	C–OH bend
210 w	208 m				209	160 vw	163 vw	Lattice
181 w	180 w				178			Lattice
163 w	163 m				164			Lattice
150 vw	148 m				98	140 m	141 m	Lattice
140 vw	138 m				87	116 w	116 m	Lattice
124 m	122 w				LL		103 vw	Lattice
114 w	113 w							Lattice
	$91\mathrm{m}$						94 s	Lattice
	81 s							Lattice
	73 ssp				67		75 wbr	Lattice
	43 vw						58 w	Lattice



Figure 15. Reference Raman spectra for AH₂ and NaAH solids. Measurement details: Laser, 532 nm. Power level, ~ 200 mW. Slit width, ~ 8 cm⁻¹. Accuracy, ± 1 cm⁻¹. Assignments are given in Table 6. Insert shows details of the range from 1800 to 1400 cm⁻¹.

excellent accordance, although minor disagreements can be seen in some cases. Explanations for this could be that the known low crystal symmetries may make the spectra dependent on the crystallite orientations, and double refraction of the nonisotropic material may further influence the spectral appearances (86). This is probably why the careful and reliable results obtained by Hvoslef and Klæboe in 1971 (39) sometimes differ from certain newer measurements; in some cases our results are reminiscent of these old data but sometimes we obtained very good agreement with newer data; for example, those of Saraiva et al. (48) (except their instrument was probably miscalibrated to give wavenumbers consistently some 5 ± 4 cm⁻¹ too high; see Table 6). Our general explanation for these deviations must be that the spectra depend heavily on wavenumber scale calibration, polarization, anisotropy, and crystallite orientation, as previously seen for many other crystal cases (87–89).

In Figure 15, it is clearly seen that ascorbic acid shows a weak band at 1746 cm⁻¹ and a broad doublet band at 1667–1653 cm⁻¹. The doublet band is not present in sodium ascorbate, which only has a strong band at 1703 and another strong one at 1598 cm⁻¹ (see the inset in Figure 15). According to the modeling for AH₂ and AH⁻, these bands originate from coupled modes of C1=O1 and C2=C3 stretchings (Table 6). The splitted doublet in AH₂ most probably is due to intermolecular couplings between C1=O1 and C2=C3 stretchings in two adjacent AH₂ molecules. This doublet band of AH₂ (at ~1660 cm⁻¹)



Figure 16. Measured and calculated Raman spectra for ascorbic acid. Top: AH_2 in concentrated aqueous solution, measured with 488 and 532 nm laser lines. The spectrum of water is included for reference (532 nm). Middle: AH_2 powder measured with excitation wavelength 532 nm. Bottom: Calculated spectrum for the most stable conformer (see Figure 4) as found by Gaussian modeling within a PCM water model. Assignments are given in Table 7.

can be used analytically; for example, to examine whether neutral vitamin C tablets contain mixtures of $AH_2/NaAH$ crystals or a mixture of the acidic AH_2 and a base such as a carbonate.

Measured Raman spectra of AH₂ dissolved in water and as crystal powder are shown in Figure 16, together with the Raman spectrum calculated for the minimum conformation simulated within the PCM aqueous environment (geometry as in Figure 4). As seen in Figure 16, the spectrum in the aqueous solution (top) is comparable to that of the crystalline phase (middle), except for the broad OH stretching feature at \sim 3650 to \sim 3070 cm⁻¹, as already discussed by Hvoslef and Klæboe (39). This means that no drastic rearrangement of the structure occurs by dissolution in water compared with the crystal, although the conformer equilibrium might shift. The experimental spectrum in solution is, of course, dominated by the presence of H_2O molecules giving rise to the water bands, with an envelope depending on temperature, pH, and presence of solutes (90, 91). We note that H₂O molecules were not directly incorporated into the modeling in Figure 16 (bottom) and therefore the OH stretching band envelope is absent from the calculated spectrum. In addition, there are AH_2 spectral differences and shifts between the aqueous solution and the crystal, indicating that the vitamin C molecules must be strongly affected by hydrogen bonding to the water molecules. Most distinctive in this respect are the Raman shifts from 2917 and 1667–1653 cm^{-1} for the solid, compared to 2945 and 1692 cm^{-1} for the acid

in solution (Table 7). These peaks represent the side chain C–H stretchings and the ring C1=O1 and C2=C3 stretchings, respectively, and it therefore seems natural that these peaks change upon dissolution in water.

No distinct sign was observed of any dissociation reaction of ascorbic acid to ascorbate, caused by the protolytic equilibrium with the water (1):

$$C_6H_8O_6(aq) + H_2O(l) \rightarrow C_6H_7O_6(aq) + H_3O^+(aq).$$
 (1)

The solution spectrum in Figure 16 did not seem to change significantly when we added HCl, as also seen previously (39). The wavenumber values are given in Table 6 (solid) and Table 7 (solution). The overall similarity between the spectra—from powder, solution, and as calculated—indicates that the PCM aqueous environment model works reasonably well. Note that entirely correct wavenumber positions cannot be expected to result from the modeling, due to its simplicity and the fact that only a single molecular entity was considered. It has often been the practice in similar cases to "calibrate" the wavenumber scales by multiplying by a rather arbitrary constant near one (61), but we preferred not to do so (an extra parameter would, of course, make a better fit). Still, the accordance between the DFT calculations with water as solvent and the experimental Raman spectrum of AH_2 in real water solution seems rather remarkable (see Figure 16).

Visible laser lines of 532 (green) and 488 nm (blue) were used for the Raman measurements, giving quite identical results. The dependence on polarization was also studied (see Figure 17). Some of the bands (arrows) were highly polarized, and the observations were used to confirm the assignments by comparison to the calculated polarization values. Raman spectra of deuterated and normal aqueous AH₂ solutions are shown and compared in Figure 18, complementing the results of Hvoslef and Klæboe (39). The comparison reveals that many bands from isotopically undisturbed AH2 molecules are still present in the freshly made D₂O solution. Thus, the exchange of hydroxyl protons with deuterium in D₂O solution is incomplete, in support of the previous conclusion by Hvoslef and Klæboe (39) based on neutron diffraction and deuteration IR absorption spectra. In particular, hydrogen-bonded O-H...O stretching bands seem to show their presence in Figure 18 at around 3440 cm⁻¹, in addition to the expected O-D...O stretching bands from D₂O seen at \sim 2400 cm⁻¹. The broad 3440 cm⁻¹ band shape is interpreted as due to the many O–H...O microstates of hydrogen bonding interactions between AH₂ and the solvent OD₂. The AH₂ oxygen-bound hydrogens (or at least some of them) are not exchanged fast with deuterium. The O-D...O stretching band is broad as in ordinary water, and for the same reasons. The fact that the O-H... O stretching band envelope occurs at higher wavenumbers in D_2O solution than in ordinary water (at 3440 cm⁻¹ compared to 3410 cm⁻¹) shows that it is not just the hydrogen content in our ($\sim 90\%$) D₂O/H₂O solvent mixture, although some part of the band may have that origin. The 1692 cm^{-1} band assigned to the ring C1=O1 and C2=C3 stretchings (mode B) is shifted to ~ 1685 cm⁻¹, and this shift—not surprisingly—shows that the C1=O part of the ring must also be interacting with the solvent.

Sodium ascorbate results are shown in Figure 19 and the wavenumber data are included in Table 7. The only Raman data available in the literature on ascorbate salts are those of Hvoslef and Klæboe (39).

To sum up, our observed bands for AH_2 and AH^- compare quite well with other spectral modeling results (4, 15, 23, 26, 46, 52–59) and with the available experimental bands in the literature (39–45). We note, however, that the AH_2 sample of Panicker et al. (45), presumably solid, seems to have been not entirely pure, because the Raman spectrum

Ascorbic acid	1 solution	Sodium asco	chate solution	Disodium di-ascorbate solution	
This work		This work		This work	
Figure 16	Hvoslef and	Figure 19	Hvoslef and	Figure 21	
Figure 21	Klæboe (39)	Figure 21	Klæboe (39)	Figure 26	Assignments
2945 m	2944 mbr			2922 s br	C–H str
2904 sh	2904 msp	2939 s	2933 s		C–H str
	2863 vw	2895 br sh	2905 s		C–H str
1760 w	1762 mbr	1718 m	1717 vs	1701 w	C=C + C=0 str A
1692 vs	1693 vsbr	1591 s	1594 vsbr	1556 s	C=C + C=O str B
1650 br sh	1503 w	1436 vw	1434 vw?		C=C str
1470 w	1470 m	1472 w	1360 mbr	1470 vw	$CH_2 sci$
1353 w, pol	1355 w		1325 w	1343 w br	CH_2 bend
1299 w	1299 m	1292 w	1295 s	1292 w sp	ring def, CH ₂ wag, C–O–H
			1280 s		bend C ?
1277 vw	1273 vw		1242 m	1228 vw w	
1220 vw sh	1218 mbr	1143 w	1142 m		C–O–H bend (twi)
		1113 w	1114 m	1117 w	C-C(-0)-0 str
1149 m	1150 m			1151 m sp	C–O–C str, ring def
1119 vw	1119 w		1086 vw	1116 vw	
1085 vw	1087 vw		1065 w	1070 vw sh	C–O–C str, C–O–H bend
1051 m, pol	1052 m	1041 m	1043 s	1043 w	C-O-C str
1020 w	1022 w	1020 w	1018 m	1021 vw sh	ring O–H bend

Measured Raman spectral bands (in cm⁻¹) and assignments for aqueous solutions of ascorbic acid and sodium ascorbates, compared to literature Table 7

935 w 937 mbr 929 m 931 s 937 vw 875 wbr 873 mbr 869 w 869 m 875 vw C4-C5 ring st 875 wbr 873 mbr 869 w 869 m 875 vw C4-C5 ring st 766 vw 797 vw 81 m 832 vs 825 m sp C4-C5 ring st 766 vw 797 vw 764 wbr 764 vw 014 opl def/rin 730 vw 788 vw 757 vw 764 vw 014 opl def/rin 695 wbr, pol 697 s 724 w 762 w 764 vw 634 m 669 s 764 vw 764 vw 014 opl def/rin 664 m 669 s 669 s 669 w 014 opl def/rin 654 w 633 s 660 s 640 s 640 s 014 opl def/rin 654 vw, pol 565 wbr 660 s 640 s 640 s 040 s 014 opl def/rin 87 vw 580 w 430 w 640 s 640 s 040 s 014 opl def/rin 87 vw 580 w 610 w 631 s 050 w 014 opl def/rin 87 vw 336 wbr 360 wbr <t< th=""><th>979 w</th><th>981 w</th><th></th><th>975 wbr</th><th>981 vw</th><th>C–H and O–H bend</th></t<>	979 w	981 w		975 wbr	981 vw	C–H and O–H bend
875 whr 873 mhr 869 w 869 m 875 ww C4-C5 ring st 825 s, pol 826 vs 831 m 832 vs 825 m sp C-C ring st 766 vw 797 vw 757 vw 762 w 764 vw 0H oopl def/rin 695 whr, pol 697 s 724 w 725 whr 762 w 764 vw 0H oopl def/rin 664 m 664 m 707 vs 714 vw 0H oopl def/rin 664 m 663 w 669 s 669 w 0H oopl def/rin 663 w 669 s 669 w 0H oopl def/rin 877 vw 589 m 640 w 640 w 640 s 669 w 0H oopl def/rin 877 vw 589 m 640 w 640 w 640 s 660 s 660 w 0H oopl def/rin 877 vw 445 mhr 436 vw 437 m 300 vs 0-H wag C-O ipl def/rin 3306 vw 305 whr 304 vw sh 309 vs 0-H wag 0-H wag	935 w	937 mbr	929 m	931 s	937 vw	
825 s, pol 826 vs 831 m 832 vs 825 m sp C-C ring st 766 vw 797 vw 757 vw 762 w 764 vw 0H oopl def/rin 804 wbr 730 vw 757 vw 757 vw 762 w 764 vw 0H oopl def/rin 695 wh of 697 s 724 w 725 wbr 714 vw 0H oopl def/rin 634 m 633 s 668 w 669 s 669 w 0H oopl def/rin 634 m 633 s 668 w 669 s 669 w 0H oopl def/rin 565 wbr 630 w 640 w 640 s 642 vw 0H oopl def/rin 430 w 258 vw 936 wbr 330 vw 335 wbr 330 vw 335 wbr 330 vw 335 wbr 330 vw 935 wbr 300 vs 0H oopl def/rin 330 vw 0D wbr 330 vw 0D wbr 300 vs	875 w br	873 mbr	869 w	869 m	875 vw	C4–C5 ring str
766 vw 797 vw 804 wbr OH oopl def/rin 730 vw 768 vw 757 vw 762 w 764 vw 0H oopl def/rin 695 w br, pol 697 s 724 w 752 wbr 764 vw 0H oopl def/rin 695 w br, pol 697 s 724 w 725 wbr 764 vw 0H oopl def/rin 664 m 664 m 706 m 707 vs 714 vw 0H oopl def/rin 634 m 669 w 669 s 669 w 0H oopl def/rin 634 m 633 s 668 w 669 s 640 w 587 w 589 m 640 w 640 s 640 w 564 vw, pol 565 wbr 603 w 605 vs 610 w 487 vw 488 w 430 m 0H onpl def/rin 322 vw 396 wbr 304 vw sh 309 vs 0.04 wag 306 vw 306 vw 309 vs 0.04 vwsh 0.04 wag	825 s, pol	826 vs	831 m	832 vs	825 m sp	C-C ring str
730 vw768 vw757 vw762 w764 vw 697 s 72 w 762 w 764 vw 0 H oopl def/rin 664 m 697 s 724 w 725 wbr 0 H oopl def/rin 664 m 603 s 669 s 669 s 669 w 0 H oopl def/rin 634 m 633 s 668 w 669 s 669 w 0 H oopl def/rin 587 w 589 m 640 w 640 s 669 s 669 w 0 H oopl def/rin 587 w 589 m 640 w 640 s 640 s 642 vw 0 H oopl def/rin 587 w 589 m 640 w 640 s 640 s 610 w 0 H oopl def/rin 587 w 589 m 640 s 640 s 640 s 0.40 w 0.1 wag 570 w 380 w 396 wbr 437 m 330 vw 0.1 wag 0.1 wag 306 vw 306 wbr 304 vw sh 309 vs 0.1 wag 0.1 wag	766 vw	wv 797 vw		804 wbr		OH oopl def
695 w br, pol 697 s 724 w 725 wbr OH oopl def/rin 664 m 664 m 706 m 707 vs 714 vw OH oopl def/rin 664 m 664 m 706 m 707 vs 714 vw OH oopl def/rin 634 m 663 s 669 s 669 w 0H oopl def/rin 587 w 589 m 640 w 640 s 640 s 0H oopl def/rin 587 w 589 m 640 w 640 s 640 s 0H oopl def/rin 587 w 589 m 640 w 640 s 640 s 0H oopl def/rin 487 vw 488 w 490 m 610 w C-0 ipl def/rin 487 vw 443 vw 437 m 350 vw 0-H wag C-0 ipl def/ring 396 wbr 306 vw 309 vs 00 ws 0-H wag 306 vw 305 wbr 309 vs 0-H wag 0-H wag	730 vw	768 vw	757 vw	762 w	764 vw	
664 m 664 m 706 m 707 vs 714 vw OH oopl def/rin 634 m 633 s 668 w 669 s 669 w 0H oopl def/rin 587 w 589 m 640 w 669 s 669 w 0H oopl def/rin 587 w 589 m 640 w 669 s 640 s 0H oopl def/rin 587 w 589 m 640 w 640 s 640 s 0H oopl def/rin 587 w 589 m 640 w 640 s 640 s 040 m 564 vw, pol 565 wbr 603 w 605 vs 610 w 0H oopl def/rin 487 vw 488 w 430 m 610 w 0.1H wag C-0 ij 0.1H wag C-0 ij 392 vw 396 wbr 304 vw sh 309 vs 0.0H wag 0.0H wag	695 w br, pol	697 s	724 w	725 wbr		OH oopl def/ring def
634 m 633 s 668 w 669 s 669 w OH oopl def/rin 587 w 589 m 640 w 640 s 640 s 640 s 0H oopl def/rin 564 vw, pol 565 wbr 603 w 605 vs 610 w OH oopl def/rin 487 vw 488 w 603 w 605 vs 610 w C-O ipl def/rin 433 vw 436 wbr 436 mbr 437 m 0-H wag C-O ipl def/ring 322 vw 396 wbr 304 vw sh 309 vs 0-H wag C-O ipl def/ring	664 m	664 m	706 m	707 vs	714 vw	OH oopl def/ring def
587 w 589 m 640 w 640 s 640 s 642 vw OH oopl def/C-C 564 vw, pol 565 wbr 603 w 605 vs 610 w OH oopl def/C-C 487 vw 488 w 603 w 605 vs 610 w C-O ipl def/ring 487 vw 488 w 430 m 630 w 0.04 wag C-O ij 322 vw 396 wbr 304 vw sh 309 vs 0.04 wag C-O ij	634 m	633 s	668 w	669 s	669 w	OH oopl def/ring def
564 vw, pol 565 wbr 603 w 605 vs 610 w 487 vw 488 w 490 m C-O ipl def/ring 487 vw 488 w 430 m C-O ipl def/ring 443 vw 445 mbr 436 vw 437 m 0-H wag C-O ipl def/ring 392 vw 396 wbr 304 vw sh 309 vs 00 m br 350 vw 0-H wag	587 w	589 m	640 w	640 s	642 vw	OH oopl def/C-C ring str
487 vw 488 w 490 m C-O ipl defring 443 vw 445 mbr 436 vw 437 m 0-H wag C-O ipl defring 392 vw 396 wbr 304 vw sh 309 vs 00 vs 00 -H wag	564 vw, pol	565 wbr	603 w	605 vs	610 w	
443 vw 445 mbr 436 vw 437 m 0-H wag C-O i 392 vw 396 wbr 400 m br 350 vw 0-H wag 306 vw 305 wbr 304 vw sh 309 vs 0-H wag	487 vw	488 w		490 m		C–O ipl def/ring def
392 vw 396 wbr 400 m br 350 vw 0-H wag 306 vw 305 wbr 304 vw sh 309 vs 0-H wag	443 vw	445 mbr	436 vw	437 m		O-H wag C-O ipl def
306 vw 305 wbr 304 vw sh 309 vs 0-H wag	392 vw	396 wbr		400 m br	350 vw	O–H wag
	306 vw	305 wbr	304 vw sh	309 vs		O–H wag



Figure 17. Dependence of Raman spectra of AH_2 solution on polarization, as obtained with 532 nm excitation on the DILOR-XY instrument. VV and VH refer to vertically (V) polarized incoming beam being analyzed with a sheet polarizer (V and H) after 90° of horizontal (H) scattering. The temperature was 24°C.

showed signs of ascorbate ions at \sim 1484 cm⁻¹ and their spectral resolution was low (doublet at 1667–1653 cm⁻¹ not resolved).

Titration Experiments

To investigate more closely the behavior of AH_2 and the deprotonated forms AH^- and A^{2-} in water, we conducted titrations by adding small aliquots of NaOH solution. Oxygenpoor conditions were maintained inside quartz cuvettes as described in the Experimental section. During the titrations, typical Raman spectra were obtained using 488 and 532 nm laser excitation, as shown in Figures 20 and 21. The bottom spectra clearly show the dissolved AH_2 with the characteristic ~1692 cm⁻¹ band present before the start of titration (pH ~2). Vibration bands occurred at ~1757, ~1692, and ~1353 cm⁻¹, respectively, for both the visible and DUV spectra, which is comparable with the literature (39). Then, along with addition of NaOH, the spectrum gradually changed to that of AH⁻, developing the characteristic bands at ~1719 and ~1591 cm⁻¹ (pH = ~9, Figure 21). A weak vibration band was observed at 1430 cm⁻¹ in the UV spectra and in the visible spectra, which also occurred between 1430 and 1440 cm⁻¹. As expected, there was no indication of any intermediate species. By further titration beyond the AH⁻ state (pH > 12), a strong



Figure 18. Raman spectra of normal and deuterated ($D_2O/H_2O = \sim 90\%$) aqueous AH₂ solutions, as obtained with 532 nm on the DILOR-XY instrument. Solutions were freshly made and recorded at 24°C within 1 h after preparation.

fluorescence soon appeared and it became increasingly difficult to obtain good spectra, especially with the green laser line.

To avoid the fluorescence we were motivated to use deep UV excitation. Three different laser lines (229, 244, and 266 nm) were available, allowing for a study of the influence of these short wavelengths on the appearance of the spectra during the titration. Hence, spectra were recorded during the titration at times when the compositions corresponded to the presence of AH_2 , AH^- , and A^{2-} in water solution. The results obtained with these wavelengths are illustrated in Figures 22–24 and are discussed in the following.

AH_2

The results for AH_2 are given in Figure 22. The water bands around 3440 cm⁻¹ are clearly observable in spectra obtained with the visible laser line at 448 nm (as with other visible laser lines). In contrast, when the three UV wavelengths were used, the water bands were absent or not remarkable. The reason for this is that the UV light is absorbed and the occurrence of significant absorption-associated resonance Raman enhancement: The RR spectrum simply becomes so strong that the spectrum of water is not seen. The spectrum of the ascorbic acid is observed and must hence be enhanced to a considerable extent.

As mentioned an intense band occurs in all of the AH₂ spectra at $\sim 1692 \text{ cm}^{-1}$. This band, due to a mode called B, is also visible in Figure 12, at 1699 cm⁻¹ and is assigned to the ring C1=O1 and C2=C3 in-phase stretching mode, no. 45. This mode and two



Figure 19. Comparison of ascorbate AH⁻ Raman spectra, obtained from an aqueous solution, from a crystalline powder sample, and as calculated.

other interesting vibrations called A and C, involving the ring in resonance, are depicted in Figure 25. The respective calculated wavenumbers, as well as the IR and Raman signal strength for the modes called A, B, and C, are given in Table 8. Mode 45 (B) is quite polarized (polarization ratio calculated to 0.21) in accordance with what is also observed; see Figure 17. Modes A and C are not as intense as mode B for AH_2 in water, probably because the three vibrations involve different variations of symmetry couplings between the electronic transition under absorption and the vibrations of C1–O1, C2–O2, and C3–O3, as shown in Figure 25. Mode C is not only related to the ring but also involves the side chain.

AH^{-}

By the titration with OH⁻, a single proton can be removed from AH₂ to form AH⁻ and water (pH ~9). Four such typical experiments are shown in Figure 23. Again, very distinct water bands were seen near 3440 cm⁻¹ for measurements done with visible wavelengths. The water bands disappeared from the spectra when UV excitation wavelengths 266 and 244 nm were used; the rest of the spectrum of the ascorbate was observed and must hence be enhanced to a considerable extent. For measurements excited with the 229 nm line, water bands became perhaps weakly observable. The mode B vibration gave a very intense signal, occurring at a wavenumber shift of ~1591 cm⁻¹ (as for all measurements done on AH⁻). Compared to AH₂, vibration B for the deprotonated form, AH⁻, is shifted to a slightly lower wavenumber. Vibration A occurred at a wavenumber shift of ~1717 cm⁻¹, also the



Figure 20. Raman spectra versus titration experiments on ascorbic acid dissolved in water. Bottom: Neat solution of ascorbic acid in oxygen poor water ($pH = \sim 2$). Middle and top: After addition of the calculated amounts of NaOH solution, NaAH, and Na₂A solutions were formed (middle, $pH = \sim 9$, and top, pH = > 12). Laser excitation: 488 nm. Spectrometer: DILOR-XY.

same for all four excitation lines. The only signal not accounted for is the one occurring at 1652 cm^{-1} between bands for vibrations A and B when the excitation wavelength was 266 nm. This signal is probably due to a resonance enhancement of some overtone or combinational mode and is enhanced only for the 266 nm excitation.

 A^{2-}

When the titration was continued, one more proton was withdrawn from AH⁻ to form A²⁻. The study of A²⁻ in a strong alkaline environment (pH > 12) is complicated due to fast reaction with even small amounts of O₂. The A²⁻ ion is only stable for a short amount of time after addition of NaOH to a AH⁻ solution. To exclude O₂ as much as possible the method described in the Experimental section was absolutely necessary. We assume that A²⁻ ions react with electrons to form radicals (19, 23), and the radicals emit fluorescence, which makes it impossible to obtain Raman spectra with visible laser lines shortly after the NaOH addition. It was, however, possible for some minutes to obtain Raman data for the A²⁻ ion before the solution was ruined by fluorescing radicals.

Characteristic measurements on such basic solutions are shown in Figure 24. It appears that the water bands were distinct with visble (488 nm) and very deep UV excitation (229 nm), whereas they were not visible when excition was done with the 266 and 244 nm lines. Again, the reason is ascribed to the resonance phenomena and absorption. The



Figure 21. Titration experiments on ascorbic acid dissolved in water after increasing amounts of NaOH solution, done with 532 and 488 nm laser excitation and the DILOR-XY instrument. Bottom: Neat solutions of ascorbic acid in water (pH \sim 2). Top: Spectra versus increasing additions of base (to pH = >12).

experimentally measured Raman spectrum of A^{2-} is depicted in Figure 26, which also shows the DFT calculated spectrum with water as solvent. We note a good agreement between the measured and calculated spectra for A^{2-} . It seems that the resonance Raman spectra are reminiscent of much of the normal Raman spectra except that they are much more intense.

The OH stretching mode calculated at ca. 3708 cm^{-1} (Figure 26, bottom) in practice is so weak and broad (hydrogen bonding) that it is not observable under the broad water OH-stretching band (top). The rest of the theoretical spectrum is not very far from the observations. Taking into account that the solution probably is an ensemble of different conformations, the fitting seems surprisingly good. In our opinion this shows that the measured bands characteristic for A^{2-} can be satisfactorily assigned using the DFT calculated Raman spectrum.

Vibration modes A, B, and C for AH_2 and AH^- can also be identified for A^{2-} . For visible excitation the Raman bands occur at about 1697, 1556, and 1353 cm⁻¹, respectively. Measurements of the A^{2-} solutions show that several of the DUV spectral signals are slightly altered and do not occur at excactly the same positions and the band shapes do not correspond entirely to the results obtained with visible laser light (compare spectra in Figure 24 in the fingerprint range 1600–500 cm⁻¹).

The explanation is that different modes are more or less in resonance for different excitation wavelengths.



Figure 22. Experimental Raman spectra of ascorbic acid in aqueous solution versus excitation wavelength (pH = \sim 2).



Figure 23. Experimental Raman spectra of ascorbate in aqueous solution versus excitation wavelength ($pH = \sim 9$).



Figure 24. Experimental Raman spectra of A^{2-} in aqueous solution versus excitation wavelength (pH = >12).

The measurements with visible excitation wavelengths stand out as reference nonresonance-enhanced Raman spectra.

Discussion of Resonance Raman Spectra and UV-Vis Absorption Spectra

UV-Vis absorption spectra are needed to fully understand the differences between visible and UV Raman spectra for the ascorbate compounds. As pointed out ealier, absorption plays an important role in relation to the excitation wavelength chosen. Accordingly, we remeasured the absorption spectra during the titration, as shown in Figure 27, and the details are given in Table 9. As expected, the absorption spectra varied according to the different ascorbic species formed. The acid itself (AH₂) was found to have its absorption band positioned in deep UV, at a wavelength between ~ 210 and ~ 290 nm, and with a maximum at \sim 247 nm; see Figure 27. The position of the band maximum depends somewhat on the concentration, so the values in Table 9 must be taken with caution. The ascorbate mono-ion (AH⁻) was found to absorb light between \sim 230 and \sim 295 nm with a maximum at \sim 264.8 nm. The ascorbate di-ion (A²⁻) gave an absorption band between \sim 260 and \sim 330 nm with a maximum at about \sim 298.4 nm. In addition to this band, A²⁻ was found to have a deep UV band below ~ 220 nm. A reasonable consistency is seen between our absorption data reported in Table 9 and the values given in the literature (no data seem to exist for A^{2-}). We note that the absorption maxima show a trend toward lower absorption energy (longer wavelength) along the series $E(AH_2) > E(AH^-) > E(A^{2-})$.



Figure 25. Vibrational modes labeled A, B, and C of ascorbic acid molecules AH₂, AH⁻, and A²⁻.

Resonance Raman effects typically occur when the excitation wavelength lies at a wavelength where the molecule absorbs. What happened in the present case versus shifts in excitation wavelength is illustrated in Figures 22–24.

The Raman spectrum of AH_2 obtained with 229 nm lies inside the absorption range between 210 and 290 nm, and accordingly the water bands (appearing from about 3000 to 3500 cm⁻¹) are completely drowned out and only the strongly resonant AH_2 bands appear. The Raman spectrum of AH^- lies just on the edge where absorption starts for AH^- (between 230 and 295 nm; Figure 23). Only measurements of the three species conducted with the visible laser lines have significant water bands. Absorption is apparently enough to make the water signal disappear and the resonance is strong enough to let the Raman spectrum stay essentially persistent. The Raman spectrum of A^{2-} recorded with an excitation wavelength of 229 nm has a significantly intensive water band compared to AH_2 and AH^- . The absorption band of A^{2-} , between 260 and 330 nm (see Figure 24) is so far away from the excitation wavelength at 229 nm that no significant absorbance occurs and resonance enhancement for the A^{2-} molecule ion is hence absent.

The UV wavelengths 244 and 266 nm are within the range where absorption takes place, making the water Raman signal disappear in all cases (Figures 22 and 23). In contrast, measurements conducted with visible wavelengths occur without absorbance and

Table 3	8
---------	---

Wavenumber values (cm⁻¹) and calculated Raman intensity (int, in units of Å²/amu) of vibration bands A, B, and C for AH₂, AH⁻, and A²⁻ in water. Observed wavenumber values (Obsd.) from Table 7. For intensity abbreviations see Table 6

Mode see Figure 25	А	В	С
Approx. assignment	C1–C2–C3 (iph)–sym str	C1 = O1 ooph C2 = C3 asym str	Ring def, OH bend
AH_2 (H_2O)	Mode 46	Mode 45	Mode 43
	1784, Int = 9	1699, Int = 469	1433, $Int = 27$
	Obsd. = 1760 w	Obsd. = 1692 vs	Obsd. = 1299 w ?
$AH^{-}(H_2O)$	Mode 44	Mode 43	Mode 39
	1746, Int = 57	1604, Int = 89	1415 Int = 112
	Obsd. = 1718 m	Obsd. = 1591 s	Obsd. = 1292 w?
$A^{2-}(H_2O)$	Mode 42	Mode 39	Mode 36
	1699, Int = 99	1495, $Int = 273$	1369 Int = 100
	Obsd. = 1701 w	Obsd. = 1556 s	Obsd. = 1292 w
			sp?

RR enhancement. The same experiments conducted with UV laser lines showed no sign of the water band for AH_2 and AH^- . The A^{2-} measurement with 229 nm excitation was the only UV experiment that showed a clear water band, whereas when conducted with the 244 nm laser line no sign of the water band was seen. This is a consequence of absorption in combination with the RR phenomenon.

Discussion of Modes A, B, and C

We have dealt with three specific vibrational modes called A, B, and C, illustrated in Figure 25 (data given in Table 8). The vibration A in the AH₂ molecule in solution (mode no. 46) was approximately described as C1 = O1 + C2 = O2 + C2 = C3 in-phase ring stretching or briefly as C1-C2-C3 ring breathing. For AH⁻ and A²⁻ similar motions occurred as mode nos. 44 and 42. For A (and B) only the atoms in the π -bonded lactone ring system moved, but not the atoms in the side chain. Because the absorption bands are due to $\pi \to \pi^*$ transitions in the ring, it is to be expected that modes A and B are exhibiting the strongest resonance Raman effects. According to Table 4, the A modes were calculated at 1784, 1746, and 1699 cm⁻¹ in the DFT/B3LYP/6-311++G(d,p)/PCM. The second vibration (mode B) is the C2 = C3 stretching coupled out-of-phase with the C1 = O1 stretching and with the C2–O2–H2 and C3–O3–H3 bending for AH_2 or only with C3–O3 stretching in-phase for AH⁻. For A²⁻mode B in the ring was not very distinctly seen in the spectra and was coupled to OH motions of the side chain. Table 4 gives calculated modes B (number 45, 43, and 39, respectively) at 1699, 1604, and 1495 cm^{-1} . The third vibration (mode C) is an in-plane deformation of the lactone ring in combination with the C2–O2–H2, C3–O3–H3, and ring–C4–H4 angle bendings in the ring, coupled to side chain hydrogen angle deformations (see Figure 25). Modes C (number 43, 39, and 36 for the



Figure 26. Raman spectra of A^{2-} : Experimental results in freshly made solution compared to the DFT calculated result for ion in water solvent (PCM).

AH₂, AH⁻, and A²⁻, respectively) were calculated to occur at 1433, 1415, and 1369 cm⁻¹ (Figure 25 and Table 4).

Because of the deprotonation along the series $AH_2 > AH^- > A^{2-}$ (when protons successively split off), the coupling patterns of the atomic motions vary and hence the



Figure 27. Typical absorption spectra as obtained during titration.

	Band wavelength ^a	Molar	Literature values f w	for $\pi \to \pi^*$ K band in vater ^a
Species	nm	$L \text{ mol}^{-1} \text{ cm}^{-1}$	nm	$L \text{ mol}^{-1} \text{ cm}^{-1}$
AH ₂	~247.0	~6,800	243.5 (11)	9,600–10,000 (11)
			243 (14)	$9,560 \pm 350$ (14)
			248 (65)	7,189 in CH ₃ CN (64)
			245 (66)	
			240 in CH ₃ CN (64))
AH^{-}	~ 264.8	$\sim \! 16,\! 500$	265.5 (11)	14,800 (11)
			265 (14)	$14,560 \pm 450$ (14)
			260 (65)	
A^{2-}	~ 298.4	$\sim 7,000$		
	~212.6	~21,000		

Table 9		
Ultraviolet absorption data of ascorbic acid species; see I	Figure 2	27

^aDependent on concentration and protolytic dissociation (14).

normal modes and their eigenvalues, but recognizable vibrations A, B, and C occurred in all of the species. The trend is an irregular shift to lower band wavenumbers values.

Conclusion

Figure 27 shows the UV-Vis absorption spectra of aqueous solutions of AH₂, AH⁻, and A²⁻. AH₂ absorbs between 210 and 290 nm with a maximum at \sim 247 nm for concentrated solutions, AH⁻ absorbs between 230 and 295 nm with a maximum at \sim 265 nm, and A²⁻ absorbs between 260 and 330 nm with a maximum at about 298.4 nm.

The RR phenomena can be documented in Figures 22–24 showing the Raman spectra of AH₂, AH⁻, and A²⁻ recorded with excitation wavelengths 229 and 488 nm. The spectra can be rationalized in combination with the UV-Vis spectrum of the three compounds: The spectra recorded with 229 nm excitation wavelength gave resonance enhancements for AH₂ and AH⁻ but not for A²⁻. This fits with the fact that AH₂ and AH⁻ absorb light at 229 nm but A²⁻ absorbs little light at 229 nm. Accordingly, the UV-Vis spectra hint at which excitation wavelengths are most useful for resonance enhancement of the specific compounds.

On the other hand, the presented results show what excitation wavelengths should *not* be used if one wants to see what an ascorbic acid-containing sample contains: In the early stages of this work we realized that no matter what kind of fruit juice, beer, or wine studied with 229 nm excitation we did not observe anything other than the ascorbic acid, naturally present or added for preservation. This effect of nonlinearity in Raman spectral intensities, here exemplified in the disappearance of the water signals, has been seen before, as recently discussed in a case where up to high concentrations of heptane could not be seen in 229 nm excited Raman spectra when toluene was present (63).

Acknowledgments

The author thanks Christian Kistrup Hansen for carefully doing most of the experiments in this research. The author also thanks Peter Wood from CCSD (http://www.ccdc.cam.ac.uk/free_services/mercury; Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge CB2 1EZ, UK) for confirming the late Jan Hvoslef's use of left-handed coordinate systems (10, 25).

References

- 1. Tolbert, B.M., Downing, M.R., Carlson, W., Knight, M.K., and Baker, E.M. (1975) Chemistry and metabolism of ascorbic acid and ascorbate sulfate. *Ann. New York Acad. Sci.*, 258: 48–69.
- Neta, P., Huie, R.E., Mosseri, S., Shastri, L.V., Mittal, J.P., Maruthamuthu, P., and Steenken, S. (1989) Rate constants for reduction of substituted methylperoxyl radicals by ascorbate ions and N,N,N,N-tetramethyl-p-phenylenediamine. *J. Phys. Chem.*, 93: 4099–4104.
- Zapata, S. and Dufour, J.-P. (1992) Ascorbic ascorbic, dehydroascorbic and Isoascorbic, acid simultaneous determinations by reverse phase ion interaction HPLC. J. Food Sci., 57 (2): 506–511.
- 4. Allen, R.N., Shukla, M.K., Reed, D., and Leszczynski, J. (2006) Ab inito study of the structural properties of ascorbic acid (vitamin C). *Int. J. Quantum Chem.*, 106 (14): 2934–2943.
- Amorati, R., Pedulli, G.F., and Valgimigli, L. (2011) Kinetic and thermodynamic aspects of the chain-breaking antioxidant activity of ascorbic acid derivatives in non-aqueous media. *Org. Biomol. Chem.*, 9: 3792–3800.
- 6. Schlueter, A.K. and Johnston, C.S. (2011) Vitamin C: overview and update. *J. Evid. Base. Compl. Alternative Med.*, 16 (1): 49–57.
- Padayatti, S.J., Katz, A., Eck, P., Lee, J., Chen, S., Corpe, C., Dutta, S.K., and Levine, M. (2003) Vitamin C as an antioxidant: evaluation of its role in disease prevention. *J. Am. Coll. Nutr.*, 22: 18–35.
- Peng, Z., Duncan, B., Pocock, K.F., and Sefton, M.A. (1998) The effect of ascorbic acid on oxidative browning of white wines and model wines. *Aust. J. Grape Wine Res.*, 4 (3): 127–135.
- 9. Darvin, M.E., Sterry, W., and Lademann, J. (2010) Resonance Raman spectroscopy as an effective tool for the determination of antioxidative stability of cosmetic formulations. *J. Biophoton.*, 3 (1–2): 82–88.
- 10. Hvoslef, J. (1969) Changes in conformation and bonding of ascorbic acid by ionization. The crystal structure of sodium ascorbate. *Acta. Crystallogr.*, B25: 2214–2223.
- 11. Ogata, Y. and Kosugi, Y. (1970) Ultraviolet spectra of L-ascorbic acid and cupric ascorbate complex. *Tetrahedron*, 26: 4711–4716.
- Forsberg, O., Johansson, K., Ulmgren, P., and Wahlberg, O. (1973) Equilibrium studies of lascorbate ions. *Chem. Scripta*, 3: 153–158.
- Berger, S. (1984) ¹³C, ¹³C spin coupling constants as a probe for mesomeric structures: vitamin C. J. Chem. Soc., Chem. Comm., 1252–1253.
- 14. Karayannis, M.I., Samios, D.N., and Gousetis, C.P. (1997) A study of the molar absorptivity of ascorbic acid at different wavelengths and pH values. *Anal. Chim. Acta*, 93: 275–279.
- Juhasz, J.R., Pisterzi, L.F., Gasparro, D.M., Almeida, D.R.P., and Csizmadia, I.G. (2003), The effect of conformation on the acidity of ascorbic acid: a density functional study. *J. Mol. Struct.*, 666–667: 401–407.
- Sato, H. and Ikeya, M. (2004) Organic molecules and nanoparticles in inorganic crystals: vitamin C in CaCO₃ as an ultraviolet absorber. *J. Appl. Phys.*, 95 (6): 3031–3036.
- Köse, D.A. and Zümreoglu-Karan, B. (2009) Complexation of boric acid with vitamin C. New J. Chem., 33 (9): 1874–1881.
- Kanters, J.A., Roelofsen, G., and Alblas, B.P. (1977) Changes in conformation and bonding of D-isoascorbic acid by ionisation. The crystal structure of sodium d-isoascorbate monohydrate. *Acta Crystallogr.*, B33: 1903–1912.

- 19. Njus, D. and Kelley, P.M. (1993) The secretory-vesicle ascorbate-regenerating system: a chain of concerted H⁺/e⁻ transfer reactions. *Biochim. Biophys. Acta*, 1144 (3): 235–248.
- Brenner, G.S., Hinkley, D.F., Perkins, L.M., and Weber, S. (1964) Isomerization of the ascorbic acids. J. Org. Chem., 29: 2389–2392.
- 21. Njus, D. and Kelley, P.M. (1991) Vitamins C and E donate single hydrogen atoms in vivo. *FEBS Lett.*, 284 (2): 147–151.
- 22. Washko, P.W., Welch, R.W., Dhariwal, K.R., Wang, Y., and Levine, M. (1992) Review: ascorbic acid and dehydroascorbic acid analyses in biological samples. *Anal. Biochem.*, 204: 1–14.
- Costanzo, F., Sulpizi, M., Vandevondele, J., Della Valle, R.G., and Sprik, M. (2007) Ab initio molecular dynamics study of ascorbic acid in aqueous solution. *Mol. Phys.*, 105 (1): 17–23.
- Hvoslef, J. (1968) The crystal structure of L-ascorbic acid, "vitamin C." I. The X-ray analysis. Acta Crystallogr., B24: 23–35.
- Hvoslef, J. (1968) The crystal structure of L-ascorbic acid, "vitamin C." II. The neutron diffraction analysis. *Acta Crystallogr.*, B24: 1431–1440.
- Milanesio, M., Bianchi, R., Ugliengo, P., Roetti, C., and Viterbo, D. (1997) Vitamin C at 120 K: experimental and theoretical study of the charge density. J. Mol. Struct. (Theochem), 419: 139–154.
- 27. Hearn, R.A. and Bugg, C.E. (1974) Calcium binding to carbohydrates: crystal structure of calcium ascorbate dihydrate. *Acta Crystallogr.*, B30: 2705–2711.
- Hvoslef, J. and Kjellevold, K.E. (1974) The crystal structure of calcium ascorbate dihydrate. Acta Crystallogr., B30: 2711–2716.
- Abrahams, S.C., Bernstein, J.L., Bugg, C.E., and Hvoslef, J. (1978) Comparison of two independent determinations of the structure of calcium L-ascorbate dihydrate. *Acta Crystallogr.*, B34: 2981–2985.
- Ståhl, K., Andersen, J.E.T., and Christgau, S. (2006) Strontium diascorbate dihydrate. Acta Crystallogr, 62: m144–m149.
- 31. Koziol, A.E., Stepniak, K. and Lis, T. (1992) The crystal structure of lithium L-ascorbate dihydrate. *Carbohydr. Res.*, 226: 43–48.
- 32. Hughes, D.L. (1973) Crystal structure of thallium(I) L-ascorbate. J. Chem. Soc. Dalton Trans., 2209–2215.
- McClelland, B.W. (1974) Crystal and molecular structure of barium 2-O-sulfonato-L-ascorbate dihydrate. *Acta Crystallogr.*, B30: 178–186.
- Tajmir-Riahi, H.A. (1990) Coordination chemistry of vitamin C. Part I–III. J. Inorg. Biochem., 40: 181–188.
- Tajmir-Riahi, H.A. (1991a) Coordination chemistry of vitamin C. Part II. J. Inorg. Biochem., 42: 47–55.
- Tajmir-Riahi, H.A. (1991b) Coordination chemistry of vitamin C. Part III. J. Inorg. Biochem., 44: 39–45.
- Tajmir-Riahi, H.A. and Boghai, D.M. (1992) Coordination chemistry of vitamin C. Part IV. J. Inorg. Biochem., 45: 73–84.
- Edsall, J.T. and Sagall, E.L. (1943) Raman spectra of L-ascorbic acid, tetronic acid and related compounds. J. Am. Chem. Soc., 65 (7): 1312–1316.
- 39. Hvoslef, J. and Klæboe, P. (1971) Vibrational spectroscopic studies of L-ascorbic acid and sodium ascorbate. *Acta Chem. Scand.*, 25: 3043–3053.
- Falk, M. and Wojcik, M.J. (1979) Effect of temperature and hydrogen-bond strength on the i.r. absorption of isolated OH oscillators: crystalline L-ascorbic acid. *Spectrochim. Acta*, 35: 1117–1123.
- 41. Lohmann, W., Pagel, D., and Penka, V. (1984) Structure of ascorbic acid and its biological function. Determination of the conformation of ascorbic acid and isoascorbic acid by infrared and ultraviolet investigations. *Eur. J. Biochem.*, 138 (3): 479–480.
- 42. Lohmann, W., Beinhauer, K., and Sapper, H. (1984) Proton- and carbon-13-NMR spectra of ascorbate and isoascorbate. *Naturwissenschaften*, 71 (9): 477–478.
- 43. Ferrer, E.G. and Baran, E.J. (2001) Reduction of vanadium(V) with ascorbic acid and isolation of the generated oxovanadium(IV) species. *Biol. Trace Elem. Res.*, 83: 111–119.

- 44. De Gelder, J., De Gussem, K., Vandenabeele, P., and Moens, L. (2007) Reference database of Raman spectra of biological molecules. *J. Raman Spectros.*, 38: 1133–1147.
- 45. Panicker, C.Y., Varghese, H.T., and Philip, D. (2006) FT-IR, FT-Raman and SERS spectra of vitamin C. *Spectrochim. Acta*, 65: 802–804.
- 46. Shimada, H., Nibu, Y., and Shimada, R. (2008) Pressure effects on inter- and intramolecular vibrations in hydrogen-bonded L-ascorbic acid crystal. *J. Raman Spectros.*, 39: 32–39.
- 47. Jehlička, J., Vítek, P., and Edwards, H.G.M. (2010) Raman spectra of organic acids obtained using a portable instrument at −5°C in a mountain area at 2000 m above sea level. J. Raman Spectros., 41: 440–444.
- Saraiva, G.D., Lima, J.A., Jr., de Sousa, F.F., Freire, P.T.C., Mendes Filho, J., and Souza Filho, A.G. (2011) Temperature dependent Raman scattering study of L-ascorbic acid. *Vib. Spectros.*, 55: 101–106.
- 49. Adams, D.M. and Berg, R.W. (1976) Low-temperature far-infrared spectra of some tetrachloropalladate(II) and tetrachloroplatinate(II) salts. J. Chem. Soc. Dalton Trans., 1976: 52–58.
- 50. Krebs Larsen, F. and Berg, R.W. (1977) A neutron diffraction study of the crystal structure of deuterated ammonium tetrachloropalladate(II) at low and ambient temperature. *Acta Chem. Scand.*, A31: 375–378.
- Krebs, B., Greiwing, H., Brendel, C., Taulelle, F., Gaune-Escard, M., and Berg, R.W. (1991) Crystallographic and ²⁷A1-NMR study on premelting phenomena in crystals of sodium tetrachloroaluminate. *Inorg. Chem.*, 30: 981–988.
- Bichara, L.C., Lanus, H.E., Nieto, C.G., and Brandan, S.A. (2010) Density functional theory calculations of the molecular force field of L-ascorbic acid, vitamin C. J. Phys. Chem., A144: 4997–5004.
- 53. Singh, P., Singh N.P., and Yadav, R.A. (2010) Study of the optimized molecular structures and vibrational characteristics of neutral L-ascorbic acid and its anion and cation radical using density functional theory. *J. Chem. Pharm. Res.*, 2 (5): 656–681.
- 54. Yadav, R.A., Rani, P., Kumar, M., Singh, R., Singh, P., and Singh, N.P. (2011) Experimental IR and Raman spectra and quantum chemical studies of molecular structures, conformers and vibrational characteristics of L-ascorbic acid and its anion and cation radical. *Spectrochim. Acta*, A84 (1): 6–21.
- 55. Carlson, G.L., Cable, H., and Pedersen, L.G. (1976) An ab initio study of ascorbic acid. *Chem. Phys. Lett.*, 38 (1): 75–78.
- Al-Laham, M.A., Petersson, G.A., and Haake, P. (1991) Ab initio study of ascorbic acid conformations. J. Comput. Chem., 12(1): 113–118.
- 57. Mora, M.A. and Melendez, F.J. (1998) Conformational ab initio study of ascorbic acid. *J. Mol. Struct. (Theochem)*, 454: 175–185.
- O'Malley, P.J. (2001) Density Functional calculations modelling the spin density distribution, hyperfine couplings, and hydrogen bonding environment of the ascorbate (vitamin C) free radical. *J. Phys. Chem.*, B105: 11290–11293.
- Dimitrova, Y. (2006) Theoretical study of the changes in the vibrational characteristics arising from the hydrogen bonding between vitamin C (L-ascorbic acid) and H₂O. *Spectrochim. Acta*, A63: 427–437.
- Frisch, M.J., Trucks, G.W., Schlegel, H.B., Scuseria, G.E., Robb, M.A., Cheeseman, J.R., Montgomery, J.A., Jr., Vreven, T., Kudin, K.N., Burant, J.C., Millam, J.M., Iyengar, S.S., Tomasi, J., Barone, V., Mennucci, B., Cossi, M., Scalmani, G., Rega, N., Petersson, G.A., Nakatsuji, H., Hada, M., Ehara, M., Toyota, K., Fukuda, R., Hasegawa, J., Ishida, M., Nakajima, T., Honda, Y., Kitao, O., Nakai, H., Klene, M., Li, X., Knox, J.E., Hratchian, H.P., Cross, J.B., Bakken, V., Adamo, C., Jaramillo, J., Gomperts, R., Stratmann, R.E., Yazyev, O., Austin, A.J., Cammi, R., Pomelli, C., Ochterski, J.W., Ayala, P.Y., Morokuma, K., Voth, G.A., Salvador, P., Dannenberg, J.J., Zakrzewski, V.G., Dapprich, S., Daniels, A.D., Strain, M.C., Farkas, O., Malick, D.K., Rabuck, A.D., Raghavachari, K., Foresman, J.B., Ortiz, J.V., Cui, Q., Baboul, A.G., Clifford, S., Cioslowski, J., Stefanov, B.B., Liu, G., Liashenko, A., Piskorz, P., Komaromi, I., Martin, R.L., Fox, D.J., Keith, T., Al-Laham, M.A., Peng, C Y., Nanayakkara, A., Challacombe, M., Gill,

P.M.W., Johnson, B., Chen, W., Wong, M.W., Gonzalez, C., and Pople, J.A. (2004) *Gaussian* 03 W, Rev. E.01. Gaussian, Inc.: Wallingford, CT.

- 61. Fairchild, S. Z., Bradshaw, C. F., Su, W. and Guharay, S. K. (2009) Predicting Raman spectra using density functional theory. *Appl. Spectros.*, 63 (7): 733–741.
- 62. Harrand, M. and Lennuier, R. (1946) Exaltation of the intensity of certain bands in the Raman spectra emitted from solids with an absorption band near the excitation wavelength. *Compt. Rend. Acad. Sci.*, 223: 356–357 (in French).
- 63. Liu, C. and Berg, R.W. (2013) Nonlinearity in intensity versus concentration dependence for the deep UV resonance raman spectra of toluene and heptane. *Appl. Spectros. Rev.*, 48: 425–437.
- 64. Shriver, D.F. and Dunn, J.B.R. (1974) The backscattering geometry for Raman spectrometry of colored materials. *Appl. Spectros.*, 28 (4): 319–323.
- 65. Asher, S.A., Ludwig, M., and Johnson, C.R. (1986) UV resonance Raman excitation profiles of the aromatic amino acids. *J. Am. Chem. Soc.*, 108: 3186–3197.
- Asher, S.A. (2002) Ultraviolet Raman spectroscopy. In *Handbook of Vibrational Spectroscopy*, vol. 1. Chalmers, J.M. and Griffiths, P.R. Eds. J. Wiley & Sons: New York, pp. 557–571.
- Wittine, K., Gazivoda, T., Markus, M., Mrvos-Sermek, D., Hergold-Brundic, A., Cetina, M., Ziher, D., Gabelica, V., Mintas, M., and Raic-Malic, S. (2004) Crystal structures, circular dichroism spectra and absolute configurations of some L-ascorbic acid derivatives. *J. Mol. Struct.*, 687 (1–3): 101–106.
- Davey, M.W., Bauw, G., and Van Montagu, M. (1996) Analysis of ascorbate in plant tissues by high-performance capillary zone electrophoresis. *Anal. Biochem.*, 239: 8–19.
- Kimoto, E., Terada, S., and Yamaguchi, T. (1997) Analysis of ascorbic acid, dehydroascorbic acid, and transformation products by ion-pairing high-performance liquid chromatography with multiwavelength ultraviolet and electrochemical detection. *Meth. Enzymol.*, 279: 3–12.
- 70. Albrecht, A.C. (1961) On the theory of Raman intensities. J. Chem. Phys., 34: 1476–1484.
- Hassing, S. and Sonnich Mortensen, O. (1980) Kramers-Kronig relations and resonance Raman scattering. J. Chem. Phys., 73 (3): 1078–1083.
- 72. Tuschel, D.D., Mikhonin, A.V., Lemoff, B.E., and Asher, S.A. (2011) Deep ultraviolet resonance Raman excitation enables explosive detection. *Appl. Spectros.*, 64 (4): 425–432.
- 73. Vickers, T.J., Mann, C.K., Zhu, J., and Chong, C.K. (1991) Quantitative resonance Raman spectroscopy. *Appl. Spectros. Rev.*, 26 (4): 341–375.
- McHale, J.L. (2002) Resonance Raman spectroscopy. In *Handbook of Vibrational Spectroscopy*, vol. 1. Chalmers, J.M. and Griffiths, P.R., Eds. J. Wiley & Sons: New York, pp. 534–556.
- Efremov, E.V., Ariese, F. and Gooijer, C. (2008) Achievements in resonance Raman spectroscopy. Review of a technique with a distinct analytical chemistry potential. *Anal. Chim. Acta*, 606: 119–134.
- Liu, C. and Berg, R.W. (2012) Determining the spectral resolution of a charge-coupled device (CCD) Raman instrument. *Appl. Spectros.*, 66 (9): 1034–1043.
- Asher, S.A. and Johnson, C.R. (1984) Raman spectroscopy of a coal liquid shows that fluorescence interference is minimized with ultraviolet excitation. *Science, New Series*, 225 (4659): 311–313.
- Loppnow, G.R., Shoute, L., Schidt, K.J., Savage, A., Hall, R.H., and Bulmer, J.T. (2004) UV Raman spectroscopy of hydrocarbons. *Phil. Trans. R. Soc. Lond.*, A362: 2461–2476.
- Berg, R.W., Ferré, I.M., and Schäffer, S.J.C. (2006) Raman spectroscopy evidence of 1:1:1 complex formation during dissolution of WO₃ in a melt of K₂S₂O₇:K₂SO₄. *Vib. Spectros.* 42: 346–352.
- 80. Berg, R.W. and Nørbygaard, T. (2006) Wavenumber calibration of CCD detector Raman spectrometers controlled by a sinus arm drive. *Appl. Spectros. Rev.*, 41: 165–183.
- 81. Liu, C. (2012) *Implementation of Deep Ultraviolet Raman Spectroscopy*, Ph.D. Thesis, Printed at DTU Chemistry: Lyngby, Denmark.
- 82. Berg, R.W., Nørbygaard, T., White, P.C., and Abdali, S. (2011) Ab initio calculations, Raman and SERS spectral analyses of amphetamine species. *Appl. Spectros. Rev.*, 46 (2): 107–131.

- Berg, R.W., Shim, I., White, P.C., and Abdali, S. (2012) ROA and Raman spectra of amphetamine species—quantum chemical model calculations and experiments. *Am. J. Anal. Chem.*, 3: 410–421.
- Berg, R.W., Deetlefs, M., Seddon, K.R., Shim, I., and Thompson, J. (2005) Raman and ab initio studies of simple and binary 1-alkyl-3-methylimidazolium ionic liquids. J. Phys. Chem. B, 109 (40): 19018–19025.
- 85. Berg, R.W. (2007) Raman spectroscopy and ab-initio model calculations on ionic liquids: a review. *Monatsh. Chem. Chem. Mon.*, 138: 1045–1075.
- Turrell, G. (1984) Analysis of polarization measurements in Raman microspectroscopy. J. Raman Spectros., 15 (2): 103–108.
- 87. Porto, S.P.S., Giordmaine, J.A., and Daman T.C. (1966) Depolarization of Raman scattering in calcite. *Phys. Rev.*, 147: 608–611.
- 88. Porto, S.P.S. and Krishnan, R.S. (1961) Raman effect of corundum. J. Chem. Phys., 47: 1009–1012.
- 89. Dawson, P. (1972) Polarisation measurements in Raman spectroscopy. *Spectrochim. Acta*, 28: 715–723.
- Brooker, M.H., Hancock, G., Rice, B.C., and Shapter, J. (1989) Raman frequency and intensity studies of liquid water (H₂O, H₂¹⁸O and D₂O). *J. Raman Spectros.*, 20 (10): 683–694.
- Baschenko, S.M. and Marchenko, L.S. (2011) On Raman spectra of water, its structure and dependence on temperature. *Semiconductor Physics, Quantum Electronics and Optoelectronics*, 11 (1): 77–79.