Vibrational Spectrum and Normal Coordinate Analysis of *p*-Toluidino-*p*-chlorophenylglyoxime

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A complete vibrational assignment of the solid-state IR spectrum of p-toluidino-p-chlorophenylglyoxime was performed on the basis of normal coordinate analysis of a single molecule, assuming C_s point group symmetry. The calculated normal frequencies were in good agreement with the experimental one.

Key Words: Vic-dioxime, FTIR spectra, Valence force fields calculation, Normal coordinate analysis.

INTRODUCTION

Vic-dioximes have an important place among coordination compounds. The early studies on oxime complexes started in 1905 as mentioned in review of Chakravorty¹ where the reactions of dimethylglyoximes with Ni(II) ion were examined. Later in 1907, Tschugaff² isolated dimethylglyoxime complexes of Co(III) which played an important role as a model compound in the elucidation of some biological and biochemical mechanisms. Several studies were concentrated on vic-dioximes complexed with Co atom in the following years^{3, 4}. They were used as model compounds to explain the structure of vitamin B12 and coenzyme B12 since these big molecules play important roles in the biological systems.

At present, new interesting properties of *vic*-dioximes are under examination. In some studies, its Pt complexes were used as anti-tumor agents in chemotherapy of some certain types of cancer. Some were used in the production of semi-conductors. In some other studies, liquid crystal properties of some *vic*-dioximes were being examined⁵. They were also used as column packing materials in chromatographic separation of nucleotides and nucleosides after bonding to natural resins as functional groups⁶.

Vibrational spectra of dioximes were not examined before but the some metal complexes of the glyoxime molecule were studied by using only its IR spectrum⁷. Since these classes of compounds have a tendency of fluorescing, it is difficult to take their Raman spectra and use it in normal coordinate analysis (NCA).

Based upon the literature search, a complete vibrational analysis of this type of vic-dioximes is not established so far. Having such important properties of

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1816 Arslan et al. Asian J. Chem.

vic-dioxime derivatives, p-toluidino-p-chlorophenylglyoxime (pTpCPG), a complete vibrational analysis using FTIR measurement data along with the result of normal coordinate analysis are presented in this paper.

EXPERIMENTAL

p-Toluidino-p-chlorophenylglyoxime (pTpCPG) was synthesized as previously described in the literature⁸. Spectroscopically pure chemicals were obtained from Aldrich Chemicals, U.S.A. and used as such for recording spectra. The solid-state infrared spectrum of the pTpCPG studied were recorded using in the form of KBr pellet by BOMEN MB102 FTIR instrument in 4000–200 cm⁻¹ frequency ranges; the resolution was 1 cm⁻¹. The Raman spectrum of the compound in a spinning cell was excited using the 488.0 nm line of Ar⁺ gas laser and recorded on a Jobin-Yvon U 1000 spectrometer, which was calibrated against the laser plasma emission lines.

Normal coordinate analysis

Fig. 1 shows the structure of a p-toluidino-p-chlorophenylglyoxime molecule with the labeling of atoms. No structural studies are available for pTpCPG in the literature. Therefore, the structural parameters have been taken from related small molecules $^{7.9-15}$. The molecular parameters used for the calculations are reported in Table-1. Owing to its structure, the molecule pTpCPG belongs to the C_s symmetry point group. The 99 normal modes of pTpCPG are distributed between the two species (A' and A'') of the C_s point group as: 66A' + 33A''. All of the species are both IR and Raman active.

$$H_{20}$$
 C_{19}
 C_{10}
 C_{9}
 C_{10}
 C_{8}
 C_{8}
 C_{8}
 C_{18}
 C_{11}
 C_{11}
 C_{8}
 C_{18}
 C_{11}
 C_{11}

Fig. 1. Structure of p-toluidino-p-chlorophenylglyoxime

TABLE-1 MOLECULAR PARAMETERS USED FOR THE CALCULATIONS

Bond Lengths (Å)			**
$C_1 - C_2$	1.435	N ₃ —H ₁₄	1.008
C_2-N_3	1.472	O ₁₃ —H ₁₅	0.956
C ₁ —N ₄	1.290	C ₇ —H ₁₇	1.084
$C_1 - C_6$	1.510	C9—Cl ₁₉	1.700
C ₆ —C ₇	1.397	N ₃ —C ₂₂	1.426
N ₄ —O ₁₂	1.346	C_{25} — C_{32}	1.510
Bond Angles (°)			
$C_2-C_1-N_4$	125.42	C ₆ —C ₇ —H ₁₇	120.00
$C_2-C_1-C_6$	117.79	C ₈ —C ₉ —Cl ₁₉	120.00
C ₂ —N ₃ —H ₁₄	120.55	N ₄ —O ₁₂ —H ₁₆	97.95
C_2 — N_3 — C_{22}	118.90	N ₃ —C ₂₂ —C ₂₇	120.00
C ₁ -N ₄ -O ₁₂	121.45	C_{25} — C_{32} — H_{33}	109.47
C ₁ —C ₆ —C ₇	120.00	H ₃₄ —C ₃₂ —H ₃₅	109.47

In order to ascertain the amount of mixing among the internal coordinates and to obtain a more accurate description of the fundamental vibrations of pTpCPG, a normal coordinate calculation has been undertaken. The calculations carried out using Simple General Valence Force Field (SGVFF) in Wilsons GF matrix method with the computer program originally written by Schachtschneider¹⁶ and developed (for OS/2 -IBM) by Fischer et al. 17 under the name of SPSIM (SPectrum SIMultation).

The initial set of valence force constant and the corresponding off diagonal constant were transferred from related systems 11-13, 18-28. A zero order calculation with the transferred force constant was performed except for some deformational and longitudinal modes. The results indicated the reasonable agreement between the calculated and the observed frequencies. The initial set of force constant was refined by the method of least square technique by keeping some interaction force constants fixed throughout the refinement process. The final values of force constants and their description are given in Table-2. The calculated and observed wavenumber values are compared in Table-3, which also give the significant values of the potential energy distribution (PED) for each mode of vibration.

RESULTS AND DISCUSSION

The vibrational analysis of the pTpCPG under examination was performed in the foregoing study. FTIR part of the vibrational spectra of the compound were the only spectral source because of the difficulty of taking dispersive Raman spectra of the compound excited with a visible Laser source. A broad photoluminescence band along the IR wavelength region from 200 to 3500 cm⁻¹ prevented, obtaining a reasonable Raman signals. This was not a serious problem in the compound because of both A' and A" modes are active in both Raman and infrared in C_s point group symmetry. The IR spectrum of the p-toluidino-pchlorophenylglyoxime molecule is given in Fig. 2.

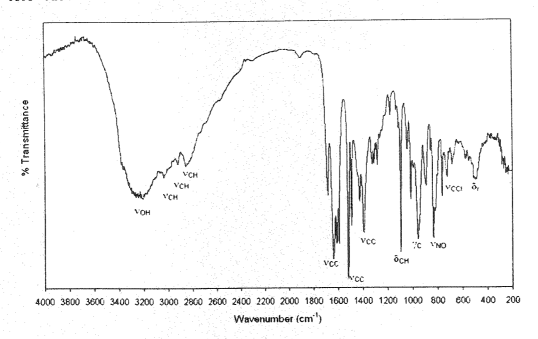


Fig. 2. Infrared spectrum of p-toluidino-p-chlorophenylglyoxime

TABLE-2
VALENCE FORCE CONSTANTS OF p-TOLUIDINO-pCHLOROPHENYLGLYOXIME†‡§

No Force constant Internal coordinate 1 $K(C_{ox}-C_{ox})$ 5.240 C_1-C_2 2 $K(C=N)$ 9.549* C_1-N_4 3 $K(C_{ox}-C_x)$ 5.101 C_1-C_6	
2 $K(C=N)$ 9.549* C_1-N_4 3 $K(C_{ox}-C_x)$ 5.101 C_1-C_6	
3 $K(C_{ox}-C_x)$ 5.101 C_1-C_6	
그리다 그 사이 바다에 가고 있다는 사람들이 되었다면 그를 가면서 되었다면 하는 사람들이 되었다.	
그리고 있다면 그 아래에 가장 요즘 가장 하는 사람이 얼마나 가장 얼마를 들어 있는 것이 되었다.	
4 $K(C_{ox}-N)$ 6.116 C_2-N_3	
5 K(N—H) 6.247* N ₃ —H ₁₄	
6 $K(N-C_1)$ 4.500 N_3-C_{22}	
7 K(N—O) 4.010 N ₄ —O ₁₂	
8 K(C—C) _{x, o} 6.415 C ₆ —C ₇	
9 $K(C-C)_{x, m}$ 6.727 C_7-C_8	
10 $K(C-H)_x$ 5.076* C_7-H_{17}	
11 $K(C-C)_{x,p}$ 6.760 C_8-C_9	
12 K(C—CI) 3.733 C ₉ —Cl ₁₉	
13 K(O—H) 6.103* O ₁₂ —H ₁₆	
14 $K(C-C)_1$ 6.415 $C_{22}-C_{23}$	par a state of
15 $K(C-H)_t$ 4.999* $C_{23}-H_{28}$	
16 K(C—CH ₃) 4.681 C ₂₅ —C ₃₂	
17 K(C _r —H) 4.588* C ₃₂ —H ₃₃	
18 $H(C_{ox}-C_{ox}=N)$ 2.651 $C_2-C_1-N_4$	
19 $H(C_{ox}-C_{ox}-C_{x})$ 0.350 $C_{2}-C_{1}-C_{6}$	
20 $H(N-C_{ox}-C_{x})$ 2.651 $N_{4}-C_{1}-C_{6}$	

No	Force constant		Internal coordinate
21	$H(C_{ox}-C_{ox}-N)$	0.350	C ₁ —C ₂ —N ₃
22	$H(C_{ox}-C_{ox}=N)$	2.651	C_1 — C_2 — N_5
23	$H(N-C_{ox}=N)$	1.654*	N_3 — C_2 — N_5
24	$H(C_{ox}-N-H)$	0.901	$C_2-N_3-H_{14}$
25	$H(C_{ox}-N-C_t)$	1.177	C_2 — N_3 — C_{22}
26	$H(H-N-C_t)$	0.901	H_{14} — N_3 — C_{22}
27	$H(C_{ox}-C-C)_x$	0.716	C_1 — C_6 — C_7
28	$H(C-C-C)_x$	1.242	C ₇ —C ₆ —C ₁₁
29	$H(CCC)_{x,o}$	0.989	C ₆ —C ₇ —C ₈
30	$H(C-C-H)_x$	0.508	C ₆ —C ₇ —H ₁₇
31	$H(C-C-C)_{x, m}$	0.865	C_8 — C_9 — C_{10}
32	H(C—C—CI)	0.828	C ₈ —C ₉ —Cl ₁₉
33	$H(C_{ox}=N-O)$	1.180	C_1 — N_4 — O_{12}
34	H(N—O—H)	0.682*	N ₄ —O ₁₂ —H ₁₆
35	$H(N-C_1-C_1)$	0.760	N ₃ C ₂₂ C ₂₃
36	$H(C-C-C)_{t}$	1.248	C_{23} — C_{22} — C_{27}
37	H(CC-C) _{t, o}	1.028	C_{22} — C_{23} — C_{24}
38	$H(C-C-H)_{i}$	0.515	C ₂₂ —C ₂₃ —H ₂₈
39	H(C—C—CH ₃)	0.754	C_{24} — C_{25} — C_{32}
40	$H(C-C_1-H_1)$	0.642	C ₂₅ —C ₃₂ —H ₃₃
41	$H(H-C_t-H)$	0.535	H ₃₃ —C ₃₂ —H ₃₄
42	$P(C_{ox1})$	0.432	C_1
43	$P(C_{ox2})$	0.602	C ₂
44	P(N)	0.106	N_3
45	$P(C)_x$	0.545	C ₆
46	$P(C)_{x, o}$	0.432	C ₇
47	$P(C)_{t}$	0.653	C ₂₂
48	$P(C)_{t, o}$	0.432	C ₂₃
49	$P(C)_{t, p}$	0.502	C ₂₅
50	$T(C_{ox}-C_{ox})$	0.085	C ₁ —C ₂
51	$T(C_{ox}=N)$	0.491	C ₁ N ₄
52	$T(C_{ox}-C_x)$	0.695	C_1 — C_6
53	$T(C_{ox}-N)$	0.037 ⋅	C ₂ —N ₃
54	T(N—C _d)	0.695	N ₃ —C ₂₂
55	T(N-O)	0.062	N ₄ —O ₁₂
56	T(C—C) _x	0.278	C6C7
57	$T(C-C)_t$	0.272	C_{22} — C_{23}
58	T(C—CH ₃)	0.211	C ₂₅ —C ₃₂
59	$F(C_{ox}-C_{ox}/C_{ox}=N)$	0.003	$C_1 - C_2 / C_1 - N_4$

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No	Force constant		Internal coordinate
60	$F(C_{ox}=N/C_{ox}-C_{x})$	0.065	$C_1 - N_4/C_1 - C_6$
61	$F(C_{ox}=N/C_{ox}=N-O)$	0.055	C ₁ —N ₄ /C ₁ —N ₄ —O ₁₂
62	$F(C_{ox}-N/N-C_t)$	-0.175	C_2 — N_3/N_3 — C_{22}
63	$F(C_{ox}-N/C_{ox}-N-C_t)$	1.602	C_2 — N_3 / C_2 — N_3 — C_{22}
64	$F(C_x-C_x/C_x-C_x)$	1.276*	C_6 — C_7 / C_6 — C_{11}
65	$F(C_x-C_x/C_{ox}-C_x-C_x)$	-0.066*	$C_6-C_7/C_1-C_6-C_7$
66	$F(C_x-C_x/C_{ox}-C_x-C_x)$	-0.086	$C_6-C_7/C_1-C_6-C_{11}$
67	$F(C_x-C_x/C_x-C_x-C_x)$	-0.609*	C ₆ —C ₇ /C ₇ —C ₆ —C ₁₁
68	$F(C_x-C_x/C_x-C_x-H)$	0.101*	C ₈ —C ₉ /C ₉ —C ₈ —H ₁₈
69	$F(C_x-C_x/C-C-Cl)$	0.197	C ₈ —C ₉ /C ₈ —C ₉ —Cl ₁₉
70	$F(C_x-C_x/C-CI)$	0.410	C ₈ —C ₉ /C ₉ —Cl ₁₉
71	F(C—C/C—C—CI)	0.654	C9-Cl ₁₉ /C ₈ C9Cl ₁₉
72	F(C—CH ₃ /C—C—CH ₃)	0.175*	C_{25} — C_{32} / C_{24} — C_{25} — C_{32}
73	F(C—CH ₃ /C—C ₁ —H ₁)	0.266*	C ₂₅ —C ₃₂ /C ₂₅ —C ₃₂ —H ₃₃
74	$F(C_{ox}/C_x)$	0.083	C ₁ /C ₆
75	$F(C_x/C_x)$	-0.065	C ₇ /C ₈
76	$F(C_x/C_x-C_x)$	-0.082*	C ₇ /C ₆ —C ₇
77	$F(C_x/C_t-C_t)$	-0.087	C_{25}/C_{24} — C_{25}
78	$F(C_x-C_x/C_x-C_x)$	-0.036	C ₆ C ₇ /C ₇ C ₈
79	$F(C_x-C_x/C_x-C_x)$	0.890*	C9-C10/C6-C11
80	$F(C_x-C_x/C_x-C_x)$	0.279	C ₆ —C ₇ /C ₉ —C ₁₀
81	$F(C_t-C_t-C_t/C_t-C_t-CH_3)$	-0.064	C_{23} — C_{24} — C_{25} / C_{24} — C_{25} — C_{32}
82	$F(C_t/C_t)$	-0.069	C ₂₅ /C ₂₆
83	$F(C_{r}-H_{t}/C_{r}-H_{t})$	0.073	C ₃₂ —H ₃₃ /C ₃₂ —H ₃₄
84	$F(C_{ox}-N-H/H-N-C_l)$	0.167*	$C_2-N_3-H_{14}/H_{14}-N_3-C_{22}$
85	$F(C_{ox}-N-H/C_{ox}-N-C_t)$	-0.574*	
86	$F(C_{ox}-C_{ox}/C_{ox}-N)$	0.432*	C_1 — C_2 / C_2 — N_3
87	$F(C_{ox}-C_{ox}/C_{ox}-C_{x})$	-0.051*	$C_1 - C_2/C_1 - C_6$
88	$F(C_1-C_1/C_1-C_1)$	-0.122*	C ₂₄ —C ₂₅ /C ₂₅ —C ₂₆
The	orce constants are given by V (atmosthin	a) II (in also	

[†]The force constants are given by K (strecthing), H (in-plane bending), P (out-of-plane bending), T (torsion), F (interaction).

The frequencies observed in the IR spectra along with their relative intensities, the corresponding calculated frequencies together with the respective potential; energy distribution (PEDs) and mode assignments are collected in Table-3. Here, the normal mode description following each fundamental in the last column is due to Wilson²⁹. As seen from Table-3, the agreement between experimental and calculated frequencies for pTpCPG is good. The calculated frequencies do not

[‡]Bond strechting and bond-bond interaction constants are in mdyn/Å, angle bending force constants are in mdyn Å/rad², and bond-angle interaction constants are in mdyn/rad.

[§] x: chlorobenzene group, t: methylbenzene group, ox: oxime group, o: ortho, m: meta, p: para * Included in the fit.

OBSERVED AND CALCULATED PLANAR AND NON-PLANAR FUNDAMENTALS (cm⁻¹) OF pTpCPG*

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	1											66	8	~	$\widetilde{\Xi}$	
												4)][96	1)1199	76][(20 H(16 K(
												(CH)][99	(CH)][99	CH)][97) + 20 H(+ 16 K(
	[(H)]	H)]		(x)) _x]	f) _x]	7	Ρ().J	P(c [v(CH)][99	c [v(CH)][99	[v(CH)][97	-C ₂) + 20 H(=N) + 16 K(
(9	-H)]	F	Ē		-H),,]	—H),,]	-H),J	-H).1	FÁ-	-H)2]	-H)(H-	etric [v(CH)][99	etric [v(CH)][99	ric [v(CH)][97	N—C,) + 20 H(C=N) + 16 K(
), %)	[(H—N)	K(0-H)]	K(0-H)]	-H),J	(C—H) _x]	(C—H) _x]	((C—H) _x]	(C—H) _[]	(C—H) _[]	(C—H) _[]	(C—H)(J	nmetric [v(CH)][99	nmetric [v(CH)][99	metric [v(CH)][97	1-N-C ₂) + 20 H(K(C=N) + 16 K(
PED, %)	9K(N—H)]	∞ K(0—H)]	00 K(0—H)]	ζ(c—H) _x]	9 K(C—H) _x]	9 K(C—H) _x]	9 K(C—H) _x]	8 K(C—H),]	9 K(C—H),]	9 K(C—H) _[]	9 K(C—H),]	symmetric [v(CH)][99	symmetric [v(CH)][99	symmetric (v(CH))[97	H(H—N—C ₀) + 20 H([53 K(C=N) + 16 K(
ts (PED, %))][99K(N—H)])][100 K(0—H)])][100 K(0—H)]	98 K _{(C—H)x}])[[99 K(C—H) _x])[[99 K(C—H) _x])] [99 K(C—H) _x])[98 K(C—H),]	J[99 K(C—H),J)[99 K(C—H)(]][99 K(C—H)[]	 asymmetric [v(CH)][99 	 asymmetric [v(CH)][99 	 symmetric [v(CH)][97 	21 H(H—N—C ₂) + 20 H(N)][53 K(C=N) + 16 K(
nents (PED, %)	NH)][99K(N—H)]	OH)][100 K(0—H)]	OH)][100 K(O—H)]	H][98 K _{(C—H)x}]	CH)][99 K(C—H)x]	CH)][99 K(C—H)x]	CH)] [99 K(C—H) _x]	CH)][98 K(C—H),]	CH)][99 K(C—H)[]	CH)][99 K(C—H)[]	CH)][99 K(C—H)[]	thyl), asymmetric [v(CH)][99	thyl), asymmetric [v(CH)][99	thyl), symmetric [v(CH)][97	H)][21 H(H—N—C) + 20 H(C=N][53 K($C=N$) + 16 K(
gnments (PED, %)	[v(NH)][99K(N—H)]	[v(OH)][100 K(O—H)]	[v(OH)][100 K(O—H)]	[VCH][98 K(C—H) _x]	[v(CH)][99 K(C—H) _x]	$[v(CH)][99 K(C-H)_x]$	[v(CH)] [99 K(C—H) _x]	[v(CH)][98 K(C—H),]	[v(CH)][99 K(C—H) ₁]	[v(CH))[99 K(C—H),]	[v(CH)][99 K(C—H),]	(methyl), asymmetric [v(CH)][99	(methyl), asymmetric [v(CH)][99	(methyl), symmetric [v(CH)][97	(NH)][21 H(H—N—C ₁) + 20 H([v(C=N)][53 K(C=N) + 16 K(C=N)]
Assignments (PED, %)	tch [v(NH)][99K(N—H)]	tch [v(OH)][100 K(O—H)]	tch [v(OH)][100 K(O—H)]	tch [vcH][98 K(c—H) _x]	tch [v(CH)][99 K(C—H) _x]	ch [v(CH)][99 K(C—H)x]	cch [v(CH)] [99 K(C—H)x]	cch [v(CH)][98 K(C—H) _[]	tch [v(CH)][99 K(C—H)t]	ch [v(CH)][99 K(C—H)t]	ch [v(CH)][99 K(C—H) _L]	ch (methyl), asymmetric [v(CH)][99	ich (methyl), asymmetric [v(CH)][99	Ich (methyl), symmetric [v(CH)][97	d [v(NH)][21 H(H—N—C ₂) + 20 H($\frac{(C=N)[53 \text{ K}(C=N) + 16 \text{ K})}{(C=N)}$
Assignments (PED, %)	stretch [v(NH)][99K(N—H)]	stretch [v(OH)][100 K(OH)]	stretch [v(OH)][100 K(O-H)]	stretch [v _{CH}][98 K _{(C—H)_x]}	stretch [v(CH)][99 K(C—H) _x]	stretch [v(CH)][99 K(C—H) _x]	stretch [v(CH)] [99 K(C—H) _x]	stretch [v(CH)][98 K(C—H),]	stretch [v(CH)][99 K(C—H) _{[1}]	stretch [v(CH)][99 K(C—H)t]	stretch [v(CH)][99 K(C—H)t]	stretch (methyl), asymmetric [v(CH)][99	stretch (methyl), asymmetric [v(CH)][99	stretch (methyl), symmetric [v(CH)][97	cend [v(NH)][21 H(H—N—C ₂) + 20 H(tretch [v(C=N)][53 K(C=N) + 16 K(
Assignments (PED, %)	VH stretch [v(NH)][99K(N—H)]	OH stretch [v(OH)][100 K(O—H)]	OH stretch [v(OH)][100 K(O—H)]	7H stretch [vcH][98 K(c_H),]	CH stretch [v(CH)][99 K(C—H) _x]	TH stretch [v(CH)][99 K(C—H) _x]	3H stretch [v(CH)] [99 K(C—H) _x]	CH stretch [v(CH)][98 K(C—H),]	CH stretch [v(CH)][99 K(C—H),]	7H stretch [v(CH)][99 K(C—H),]	"H stretch [v(CH)][99 K(C—H),]	H stretch (methyl), asymmetric [v(CH)][99	TH stretch (methyl), asymmetric [v(CH)][99	"H stretch (methyl), symmetric [v(CH)][97	1H bend [v(NH)][21 H(H—N—C ₂) + 20 H(N stretch [v(C=N)][53 K(C=N) + 16 K(
	NH suetch [v(NH)][99K(N—H)]	OH stretch [v(OH)][100 K(O—H)]	OH stretch [v(OH)][100 K(O—H)]	CH stretch [v _{CH}][98 K _{(C—H)x}]	CH stretch [v(CH)][99 K(C—H) _x]	CH stretch [v(CH)][99 K(C—H) _x]	CH stretch [v(CH)] [99 K(C—H) _x]	CH stretch [v(CH)][98 K(C—H),]	CH stretch [v(CH)][99 K(C—H) ₁]	CH stretch [v(CH)][99 K(C—H) _t]	CH stretch [v(CH)][99 K(C—H),]	CH stretch (methyl), asymmetric [v(CH)][99 K(Cr-H)]	CH stretch (methyl), asymmetric [v(CH)][99 K(C,—H)]	CH stretch (methyl), symmetric [v(CH)][97 K(C ₁ —H)]	NH bend [v(NH)][21 H(H—N—C _i) + 20 H(C _{ox} —N—H)]	CN stretch $[v(C=N)][53 K(C=N) + 16 K(C=C)_x]$
	3367 NH stretch [v(NH)][99K(N—H)]	3307 OH stretch [v(OH)][100 K(O—H)]	3307 OH stretch [v(OH)][100 K(O—H)]	3066 CH stretch [v _{CH}][98 K _{(C-H)_x]}	3060 CH stretch [v(CH)][99 K(C—H) _x]	3060 CH stretch [v(CH)][99 K(C—H) _x]	3060 CH stretch [v(CH)] [99 K(C—H) _x]	3043 CH stretch [v(CH)][98 K(C—H) _l]	3037 CH stretch [v(CH)][99 K(C—H) ₁]	3037 CH stretch [v(CH)][99 K(C—H) ₁]	3037 CH stretch [v(CH)][99 K(C—H),]				1777 NH bend [v(NH)][21 H(H—N—C ₀) + 20 H(
												2916 CH stretch (methyl), asymmetric [v(CH)][99	2915 CH stretch (methyl), asymmetric [v(CH)][99	2868 CH stretch (methyl), symmetric [v(CH)][97		1719
		3307	3307													1719
	3367	3307	3307	3066	3060	3060	3060	3043	3037	3037	3037	2916	2915	2868	7771	1719
	3367	3307	3307	3066	3060	3060	3060	3043	3037	3037	3037	2916	2915	2868	7771	1719
	3367	3307	3307	3066	3060	3060	3060	3043	3037	3037	3037	2916	2915	2868	7771	1713 w · · 1719
Calculated er Wavenumber (IR-Raman)	3367	3307	3307	3066	3060	3060	3060	3043	3037	3037	3037	2916	2915	2868	7771	1719

Species	Observed Calculated Species Wavenumber (IR) (IR-Raman)	Calculated Wavenumber (IR-Raman)	Assignments (PED, %)
A'	1684 m	1691	CN + CC stretch [v(CC), v(CN)] + [39 K(C_C), + 21 H(C_C), the stretch [v(CC), v(CN)]
A,	1675 m	6991	$CN + CC \text{ stretch } [v(CC) \ v(CN)][24 \ K(C-C) + 30 \ K(C) + 30 \ K(C-C) + 30 \ K(C) + 30 \ K(C)$
¥	1639 s	1645	CC stretch [v(CC)] $35 \text{ K(C-C)}_1 + 33 \text{ H(C-C-H)}_1$
*	1611.s	1610	CC stretch [v(CC)] [49 K(C—C) _x + 17 K(C—C) _t + 14 H(C—C—H) _x]
¥	1592 s	1600	CC stretch [v(CC)][77 K(C—C), + 17 H(C—C—H).1
À	1592 s	1595	CC stretch [v/CC)][54 K(C—C) _x + 21 K(C—C) _t +21 H(C—C—H) _x]
A,	1576 w	1571	CC stretch [v(CC)][41 K(C—C) _x + 34 H(C—C—H) _v .1
A'	1564 w	1564	CC stretch [v(CC)][58 K(C—C) _x + 11 K(C—C) _t + 11 H(C—C—H) ₋₁
A'	1564 w	1557	CC stretch [v(CC)][51 K(C—C) _t + 18 K(C—C) _t + 15 H(C—C—H).1
Α,	1517 s	1517 (CC stretch [v(CC)][29 K(C—C) _t + 15 K(C—C) _r + 14 H(C—C—H).1
A'	1491 s	1478 (CC stretch [v(CC)][46 K(C—C) _x + 10 K(C—C) _t]
Α",	1452 m	1452 (CH bend (methyl) [&(CH)][93 H(H—C—H) + 7 H(C—C—H)]
,ν	1429 m	1450	CH bend (methyl), asymmetric [8(CH)][89 H(H—C,—H) + 5 H(C—C,—H)
Α,	1393 s	1401 C	CC stretch [v(CC)][15 K(C—C) _x + 15 K(C—C) ₁ + 14 K(C ₂ —C) ₁ + 5 K(C ₂ N) ₁ ≤ W(C ₂ C ₁ N) ₂ ≤ W(C ₂ N) ₂ ≤ W(C ₂ C ₁ N) ₂ ≤ W(C ₂
Α'	1393 s	1400	CC stretch [v(CC)][30 K(C—C) _x + 14 K(C _{ox} —C _t)]
Α'	T393 s	1395 C	CH bend (methyl), symmetric [&CH)1[52 H(C_C_H) ± 43 H/H C 103

	Observed	Calculated	
Species	Species Wavenumber Wavenumber (IR) (IR-Raman)	Wavenumber (IR-Raman)	Assignments (PED, %)
Ä	1013 m	1019	NO stretch [v(NO)][55 K(N—O) + 16 K(C _{ox} —C _{ox})]
Α",	994 m	8	CH o.p. bend. $[\gamma(C)][72 P(C)_1 + 15 T(C-C)_1]$
À	972 m	972	CH bend (methyl) [δ(CH)][83 H(C—C—H _D]]
Α".	957 s	955	CH o.p. bend. [y(C)][51 P(C), + 40 H(C—C,—H,) + 14 T(C—C),]
¥	904 m	016	NO stretch [v(NO)][36 K(N—O) + 12 K(C—C),]
Α,,	890 m	890	CH o.p. bend. $[Y(C)[81 P(C)_x + 10 T(C-C)_x]$
×	851 w	850	CN stretch [v(C _{ox} N)][48 K(C _{ox} —N) + 32 F(C _{ox} —N/C _{ox} —N—C _i) + 15 H(C _{ox} —N—C _i)]
¥	830 s	824	NO stretch [v(NO)][23 K(N—O) + 22 K(C—CH ₃) + 36 K(C—C),]
Α",	818 s	814 (CH o.p. bend. $[\gamma(C)][87 P(C)_x + 10 T(C-C)_x]$
Α".	799 m	795 (CN torsion $[\tau(CN)][49 \text{ T(NC}_i) + 30 \text{ P(C)}_i + 18 \text{ P(N)}]$
Α".	761 m	783 (CC o.p. bend. $[\gamma(C)][29 P(C_{ox}2) + 22P(C_{ox}1) + 26 P(C)_x]$
Α"	731 m	739 (CH o.p. bend. [y(C)][80 P(C) _t + 18 T(C—C) _t]
Α,	721 m	734 (CCI stretch [v(CCI)][25 K(C—CI) + 36 K(C—C) _x + 16 H(C—C—C) _x]
, V	704 w	720. N	NO stretch [v(NO)][17 K(N—O) + 12 K(C—CH ₃) + 13 K(N—C ₁) + 13 H(C—C—C),]
Α",	.681 m	692 (CC o.p. bend. $[\gamma(C)][26 P(C_{ox}2) + 22 P(C)_x + 12 T(C_{ox}=N) + 12 T(C_{ox})]$
Α".	w 199	663 C	CH o.p. bend. $[\gamma(C)][40 P(C)_t + 40 T(C-C)_d]$
Α′	630 w	621 C	CCI stretch [v(CCI)][24 K(C—CI) + 13 H(C _{ox} =N—O) + 13 H(C—C—C), 1
Α"	576 m	589 C	CH o.p. bend. $[Y(C)][28 P(C)_x + 25 T(C-C)_x + 18 T(C_{ox}=N) + 2014 P(C_{ox}1)]$