Ultraviolet studies of hydrogen-bonding in isomeric toluidines

T. GANGULY AND S. B. BANERJEE

Optics Department,

Induan Association for the Cultivation of Science Jadavpur, Calcutta-700032

Hydrogen bond in toluidines in binary and ternary mixtures in different solvents has been studied by the methods of ultraviolet spectroscopy. The association equilibrium constants K and the hydrogen bonding energies (- ΔH) have been measured. It is observed that values of - ΔH are in the order para-toluidine> meta-toluidine> ortho-toluidine. Part played by steric hindrance in the case of the ortho isomer has been pointed out. Effect of intermolecular hydrogen bond on the frequency and intensity of absorption bands due to electronic transitions in the toluidines is also discussed. It has been concluded that on N--H O bond formation, binding energy of the sp^2 hybridized lone pair on nitrogen in toluidine molecules decreases so that their interaction with π -system increases leading to a large migration moment.

1 IN FRODUCTION

Hydrogen bond is generally studied by methods of infrared spectroscopy. However, ultraviolet spectroscopy is also known to furnish information of intermolecular association (Nagakura *et al.* 1952) and an estimate of the energy of hydrogen bond formation (Nagakura, 1954) can be made. Most of the earlier works in this field were concerned with frequency changes (Pimentel 1957) while only a few authors (Baba *et al* 1961) dealt with change in intensity. Intermolecular hydrogen bonding between toluidines and proton acceptor solvents was shown to exist in earlier infrared studies (Medhi *et al* 1962). In a programme of work undertaken recently, the electronic spectra of ortho-, meta- and para-toluidines were investigated in *neutral* and *proton-acceptor* solvents with the object of analysing the changes in frequency and specially, the intensity of the bands.

2. EXPERIMENTAL

Spectroscopically pure cyclohexane was supplied by B.D.H. Chemically pure dioxane was purified by distillation under reduced pressure. Ortho- and metatoluidine were subjected to fractional distillation and the collected fractions were subjected to distillation under reduced pressure. This operation was repeated four times before the samples were used. Paratoluidine was recrystallized from ethyl alcohol: This operation was repeated five to six times till white coloured crystals of paratoluidine were obtained. Melting point of the compound was then determined. The sample was kept in a vacuum desiccator.

Ultraviolet absorption spectra were measured with a Hilger UV spectrophotometer at different temperatures and 1 cm absorption cells were used For every wavelength, the slit width was maintained at a definite value. For this purpose a wavelength-slit calibration chart was prepared beforehand. Silica-get capsules (used in the spectrophotometer) were changed twice in a day to keep the instrument free from moisture. *Procedure*

Taking the three component systems viz, isomeric toluidines, dioxane and cyclohexane, the last compound being the inert solvent, a set of absorption spectra was measured. In actual practice, dioxane concentration was increased in steps from 0 to 0.5 mole/litre. In every measurement the mixture of cyclohexane and dioxane, in which concentration of dioxane was identical with that in the sample solution (cyclohexane-dioxane-toluidine), was taken as reference. The measurements were made at the three different temperatures $(293^{\circ}K, 303^{\circ}K)$ and $313^{\circ}K$).

The association equilibrium constant (K) was calculated by the equation (Baba and Suzuki, 1961)

where $\varepsilon_f = \text{molar extinction coefficient of the non-hydrogen bonded molecule,}$ $<math>\varepsilon = \text{molar extinction coefficient for a solution in which dioxane concentration}$ is C and $\varepsilon_b = \text{moler extinction coefficient for the hydrogen-bonded molecule.}$

All these extinction coefficients refer to a given frequency. Graphs were plotted taking $1/(\epsilon \cdot \epsilon_f)$ as ordinate and 1/C as abscissa. In every case, an almost linear graph was obtained (fig 1). On extrapolating the straight line to the point where $(1/\epsilon \cdot \epsilon_f) = 0$, the equilibrium constant was obtained as, K = -1/C.

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The free enery changes of the systems of isomeric toluidines were calculated from the familiar equation,



$$\Delta F = -RT\ln K \tag{2}$$

Fig. 1. Plots of $\frac{1}{\varepsilon - \varepsilon_1}$ vs $\frac{1}{C}$ for the determination of K, the association equilibrium $\varepsilon - \varepsilon_1$ constant.

Now, molar extinction coefficients ε_b for the hydrogen bonded species were calculated for every wavelength from the equation derived from the eq. (1),

$$\varepsilon_{\rm b} = \varepsilon + \frac{\varepsilon - \varepsilon_{\rm c}}{KC} \tag{3}$$

By plotting ε_b against ν (frequency in cm⁻¹), the hydrogen bonded curve was obtained (Baba and Suzuki 1961; Suzuki and Baba 1963).

The energy of hydrogen bond formation (ΔH) is related to K and T (temperature in K) by,

$$\Delta H = -\frac{Rd\ln K}{d(1/T)} \tag{4}$$

A plot of $R \ln K$ against 1/T yielded a straight line whose slope gave ΔH (fig. 2). In calculating ΔH and the association equilibrium constant K, method of least squares was used.



Fig. 2. Plots of R ln K vs 1/T for the determination of ΔH , the hydrogen bonding energy

The oscillator strengths (f) for the transitions of both the free (f_f) and the bonded (f_b) molecules were calculated from the well-known relation,

$$f = 4.32 \times 10^{-9} \int \varepsilon d\nu \qquad (5)$$

3. **RESULTS AND DISCUSSIONS**

In cyclohexane solution the isomeric molecules of toluidines are found to yield two systems of absorption bands in the regions 2620 Å to 3090 Å and 2200 Å to 2550 Å. Both are due to π - electronic transitions and are designated as Transition I and Transition II respectively. The wave numbers of bands of the two systems, the oscillator strengths (f) and the frequency shifts due to hydrogen bond formation are given in the Tables 2 and 3. It is seen that with addition of proton-acceptor dioxane to the toluidine-cyclohexane mixture, there is a shift in the positions of the absorption maxima and the integrated absorption values $\int \varepsilon d\nu$ increase. These observations are similar to those reported by Baba and Suzuki (1961) in the case of phenols and naphthols and indicate formation of intermolecular hydrogen bond between toluidine and dioxane molecules.

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In order to find out the effect of gradual increase in the concentration of the proton acceptor molecules on the formation of such bond, the value of the integrated absorption $\int \epsilon dv$ has been plotted against dioxane concentration (fig. 3(a) and 3(b)) from which it can be clearly seen that in all cases and for both the electronic transitions, the value of the integrated absorption first increases and then tends to a constant value above a certain concentration (~0.5 mole/litre) thereby indicating that bond formation is almost complete at such concentration with formation of 1:1 complex between toluidine and dioxane molecules.



Fig. 3.(a) Plots of the integrated absorption $\int \varepsilon dv$ vs dioxane concentration C at the temperature of 303°K for the transition I. $\Box = -$ ortho — toluidine $\Delta = -$ meta — toluidine

0 — para — toluidine



Fig. 3.(b) Plots of the integrated absorption $\int e^{d_v} dv$ vs dioxane concentration C at the temperature of 303°K for the transition II.

For each electronic 'ransition of the individual toluidines, the association equilibrium constant K (in litre/mole) was calculated for various frequencies at the three different temperatures 293° K, 303° K and 313° K.

For a particular toluidine system, K-values determined from the two electronic transitions agreed fairly well with each other as shown in the Table 1. From the linear graph obtained from a plot of $\frac{1}{(\epsilon - \epsilon_f)}$ vs $\frac{1}{C}$, it may be inferred that only one molecule of toluidine is associated with one dioxane molecule in the hydrogen bond formation.

Table	1.	Data	on	equil	ibri	iunı	constan	ts	and	thern	modynamic	parame	ters	for
the	hyc	Irogen	boi	nding	of	the	systems	of	isor	neric	toluidines,	dioxane	and	
	•	-					cycloho	exa	ne					

Systems	Temperature in °K	Associati consta htre	on equilibrium ant K in e/mole	- <u>A</u> fm kcals/mole		∆//in keals/mole
		Transitio 1 (2620 to 3030 Â)	n Transition H (2200Å to 2530Å)	Mean K		
	293	5 89	5 80	5 85	1.034	
ortho-toluidine	303	4 64	4 60	4 62	0.925	4 92
	313	3 058	3 00	3 ()34	0.698	
	293	24 00	24 00	24 00	1 853	
meta-toluidine	303	19.92	20.67	20-30	1 815	\$ 7?
	312	13.00	12 30	12.65	1 558	
	293	25 00	25 00	25.00	1 876	
nara-toluidine	303	18 75	19 40	19 075	1 767	6 17
para-torandine	313	12 07	11.80	11 935	1 551	

From the measured values of ΔH , the hydrogen bonding energy it is found that ΔH is in the order,

p-toluidine > m-toluidine > o-toluidine.

Possible interaction between two reacting groups like NO₂ and OH in orthopositions has been discussed by Sidgwick and Callow (1924). In the case of o-toluidine, one of the groups is CH_3 which is known to be practically inactive

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System	Transition	f* _f	f _b **	f _b /f _f	∆f/f _í	$\Delta f = f_b - f_f$
	I					
ortho-	(2620-3030 Å)	0.03944	0.04648	1.178	0.1785	0.00704
totulaine	п					
	(2200-2530 Å)	0.21340	0.29074	1.360	0.3624	0.07734
	ľ					
meta-	(2640-3040 Å)	0.03226	0.03574	1.108	0.1079	0 90348
toluidine	п					
	(2250-2550 Å)	0.19310	0 24192	1.25	0.2528	0.04882
	I					
para-	(2730-3090 Å)	0.02516	0.02915	1.158	0 1 52 6	0.00399
toluidine	"П					
	(2200-2520 Å)	0.23317	0.25542	1.095	0.0954	0.02225

Table 2. Intensity change due to hydrogen bond formation

* --free, -bonded.

System	Transition	Assignment	٧ı	ν _b	$\delta v = v_b - v_f$	Mean Sv
ortho-			33,888	33,773	(-115)	(-118)
	1	'A' ← 'A'	34,712	34,591	(-121)	
toluidine			42,359	42,181	(-178)	
	II	'A' ← 'A'	44,039	43,846	(-193)	(-185.5)
	I	'A' ← 'A'	34,954	34.772	(-182)	
meta-			34,525	34.375	(-150)	(-166)
	11	'A' ← 'A'	48,181	42,003	(-179)	
toluidine			41.828	41,653	(175)	(-177)
	I	${}^{1}B_{2} \leftarrow {}^{1}A_{1}$	34,425	34,225		
para-			34,354	34,178	· (-176)	(188)
	п	${}^{1}B_{2} \leftarrow {}^{1}A_{1}$	41,828	41,653	(-165)	
toluidine			42,540	42,359	(-181)	(-172)
			41,309	» 41,139	(-170)	

Proton-donor concentration	Proton- acceptor (dioxane) concentration in moles/litre	∫` _E d _V
ortho-toluidine of conc.	0	913 × 106
6.0×10^{-4}	0.05	10.13×10^{6}
moles/litre	0.08	9.92 × 10 ⁶
(TransitionI)	0.1	969×10^{6}
	0 2	10.41×10^{6}
	0.5	10.35 × 108
	Bonded (Calculated)	10.76×10^{8}
ortho-toluidine of conc.	0	4.94×10^{7}
3.1×10^{-5}	0.08	5.44×10^{7}
moles/litre	0.1	6.16×10^{7}
(Transition-II)	0 2	6.59×10^{-1}
	0.5	6.25 × 107
	Bonded (Calculated)	673 × 10°
meta-toluidine of conc.	0	7.47×10^{6}
$3,35 \times 10^{-4}$	0.06	7.93×10^{6}
moles/litre	0.08	7.83×10^{6}
(Transition-1)	0 2	8.03×10^{6}
	0.5	8.17×10^{6}
	Bonded (Calculated)	8 27 × 104
meta-toluidine of conc.	0	4.47 × 107
3.35×10^{-5}	0.08	5.02×10^7
moles/litre	0.1	4.89×10^{7}
(Transition-II)	0.2	5.31×10^7
	0 5	5 40 🔀 107
	Bonded (Calculated)	5.60×10^{7}
para-toluidine of conc.	0	5.825×10^{8}
3.13×10^{-4}	0.08	$5.96 imes 10^6$
moles/litre	0.1	6.21×10^{6}
(Transition-I)	0.2	6.37×10^{6}
	0.5	6.63×10^{6}
	Bonded Calculated)	6.75×10^{6}
para-toluidine of conc.	θ	5.40 × 107
3.13 × 10 ⁻⁵	Ö .1	5.39×10^{7}
moles/litre	0.2	5.51 × 107
(Transition-II)	0.5	5.78×10^7
	Bonded (Calculated)	5.91×10^7

Table 4. Change of intensity with proton-acceptor (dioxane) concentration Temperature : 303°K

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and the C-H compounds are not usually listed as H-bonding acids. (Pimentel and McClellan 1959). Therefore any intra-molecular hydrogen bond, worthy of consideration, between the two groups CH_3 and NH_2 may be ruled out. The smallest value of ΔH in the case of o-toluidine should, therefore, be attributed to steric effect (Zanker and Wi'twer 1959) of the methyl group in this molecule on the amino group in hindering intermolecular hydrogen bond tormation with dioxane molecules. This conclusion is in agreement with that arrived at from infrared studies (Medhi *et al* 1962). From this consideration, it would further be expected that the change in ΔH will be greater in going from the ortho-to 'the meta-isomer than from the meta- to 'the paraisomer. The relative values of ΔH are in accord with this expectation.

The shift in frequency on hydrogen bond formation $\delta v = v_b - v_f$ has been measured, v_b referring to frequency of absorption maxima for hydrogenbonded system and v_f to that for free molecules. It has been noticed that the band maxima shift towards longer wavelengths. The frequency shifts differ in magnitude in the case of the two electronic transitions I and II but for a particular electronic transition all the vibrational components show more or less uniform shift (Table 3) Another significant observation is (Table 2) that for both electronic transitions the oscillator strength (l_b) for the bonded unolecules is always greater than that (f_f) for the free molecules.

From these results it may be reasonably concluded that formation of internolecular N-H . . . O bond introduces larger interaction between lone pair electrons on N-atom and the ring π -electrons resulting in an increase in the migration moment. Following Baba and Suzuki (1961), it may be argued that under the influence of the electron donor oxygen atom, the σ -electrons associated with the N-H bond will be pushed towards the nitrogen atom and the resulting increase in the electron density around the nitrogen atom will lead to a decrease in the binding energy of the sp^2 hybridized lone pair of nitrogen As a result, their interaction with the ring π -system will increase, accounting for the enhanced intensity of the spectra of the hydrogen bonded species. Investigations with several similar systems are in progress.

REFERENCES

Baba H. and Suzuki S. 1961 J. Chem. Phys. 35, 1118.

Medhi K. C., Banerjee S. B. and Kastha G. S. 1962 Indian J. Phys. 36, 457.

Nagakura S. and Baba H. 1952 J. Am. Chem. Soc. 74, 5693.

Nagakura S. 1954 J. Am. Chem. Soc. 76, 3070.

Pimentel G. C. 1957 J. Am. Chem. Soc. 79, 3323.

Pimentel G. C. and Mcclellan A. L. 1959 The Hydrogen Bond (W H. Freeman and Company San Francisco and London).

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Sidgwick N. V. and Callow K. 1924 J. Chem. Soc. 35, 527.

Suzuki S. and Baba H. 1963 J. Chem. Phys. 38, 349.

Zanker V. and Wittwer. A. 1959 Hydrogen Bonding (Pergamon Press).