Rajshahi-6205

Bangladesh.

RUCL Institutional Repository

http://rulrepository.ru.ac.bd

Department of Chemistry

MPhil Thesis

1980

The Formation Constant of Dimer of Acetic Acid in Different Solvents.

Islam, M. Saidul

University of Rajshahi

http://rulrepository.ru.ac.bd/handle/123456789/861

Copyright to the University of Rajshahi. All rights reserved. Downloaded from RUCL Institutional Repository.

The FORMATION CONSTANT OF DIMER OF ACETIC ACID IN DIFFERENT SOLVENTS

By

M. Saidul Islam

A Thesis

Submitted to Rajshahi University for the Degree of Master of Philosophy

December, 1980.

Department of Chemistry University of Rajahahi Rajahahi, Bangladesh

ACKNOWLEDG EMENTS

It is impossible to acknowledge adequately the assistance and guidance I received from my Supervisor Nr. Zillur Rahim, Associate Professor, Department of Chemistry, University of Rajshahi in the preparation of this Thesis. But for his help and far-sighted guidance the producing of the thesis would have been an impossible task. I owe to him the most and consider it a rare fortune to work under him.

A note of thanks is due to Dr. M. Bazlur Rahman, Chairman, Department of Chemistry, Professor M. A. Sattar and Professor Alauddin Ahmad, ex-Chairman, Department of Chemistry for giving encouragement and providing laboratory facilities.

Acknowledgements are expressed to Dr. Nuruddin Ahmed, Associate Professor, Department of Chemistry for his helpful suggestions and comments on the final draft; and to Dr. M. Yusuff Ali, Mr. Sayenuddin Ahmed and Mr. Mohsin Ali for their occasional help.

Gratitude is expressed to my friends Messrs. M. A. Bari, Assistant Curator, Varendra Research Museum and Saiful Islam, Research Fellow, IBS, University of Rajshahi for their constant help. Messrs. Ferdous Hossain, Aminul Islam, Dnam Elahi, Zakir Hossain and M. Saber are acknowledged for their different sorts of help.

Special thanks are extended to Mr. Momtazuddin, Draftsman,
Department of Geography and Mr. Kibriya Moonzer for their painstaking
work in preparing graphs and figures. Mr. Hossain Ali is also acknowledged
for typing the thesis.

I am thankful to the Rajshahi University authority and University Grants Commission for awarding fellowship for the completion of the thesis.

Lastly but not the least, thanks are due to my wife, Hamida Islam for her help and constant encouragement throughout the M. Phil.programme.

The many transformation of the state of the transformation of the state of the stat

There is a first the state of the company of the state of

కాం. 🕽 లాండు కాండుకుండా జాలు కుండిని అయిని అందిని అక్కి అయిని అందిని మంటుకుండిని చేసినికి ఈ అయినుకుండిని ఈ అంది.

M. Saidul Islam

The color relies and solve private were extracted by securior extract the colorina by securior explain the colorina good distincts and extract the colorina colors of the color extract the color of the color extracts and the color of the color extracts and the color of the color

The strailer of and thereasize then in Ifer a demons, who was the company of a company, the action of a big interfering by the common god or the reservables there been demonstrated by the action of the company of a company of

ABSTRACT

For the determination of Formation Constant of acetic acid dimer (linear and cyclic), a theoretical frame work has been constructed where it is shown that

 $PS = \frac{K}{D} \times \frac{(P_2 - 2PS)^2}{V} + P_1; \text{ where } P_1, P_2, D, PS, V \text{ and K are that of at measured concentrations motor property of monomer, dimer, } D = P_2 - 2P_1, \text{ apparent motor property of volume which contain one mole of species and equilibrium constant respectively.}$

The molar volume and molar polarization were obtained by measurement of specific volume and dielectric constant in solutions of varying for the concentrations for which PS and V were calculated. Graphical methods were employed on the basis of the above relation to obtain P₁, P₂, KD and finally K.

The results of such investigation in five solvents, such as, CCl₄, benzene, chlorobenzene, high boiling hydrocarbon and cyclohexane have been found to be satisfactory on careful considerations of various effects of solvents and also of their peculiar structures.

斯特克雷斯 医舒适伯斯氏核水体色谱 横尾 横点的八字电

D (P2	- 2P ₁) : Change of molar property.	0 = 7 &	W.
K ₂ or K	: Formation constant of dimen	, 2 2 3	18
W	: Weight fraction		
v _f	: Volume of the float		
W _f	: Weight of the float	0.650	
△ ♥	Change of specific volume (i.e. observed - calculate	ed)	3
△ €	: Change of dielectric consta	int ed)	4
wt. fr.	or wt.fraction : Weight fraction	57774	21
	发生物的学术 在下 元素代数中各分数 是在30元	2	2.5
	Classic Libra of Relocates Timesia April 1888 Excline	D L 1 4 0	
	Country or t of Sylveges trad Parkettes	4 4 9 5	14
	Cott a rectorion and Equilibrium Conntent The Address of Solf-Associated Therefore	****	17
	and the section	***	49
	STATE MARKET		\$4
	And the Mark	5149	\$1
ä	Three Colons, Silveras, PM	6115	23
	Tollies Discould on Thirtigas transiting	€ + + ×	28
	historic Compos on Agircam Pacific	4 × × 9	1.19
	Throw to the Transferra	3 4 + 2	40
	Linger Transferen	新 引 水 准	41

		PAGE
		April production of
CHAPTER		
Contract to the contract of th		4.55
3 EXPERIMENTAL	••••	43
Description of the Magnetic Float	Ø 0 V 6	
Densimeter and the content		43
		10
Thermostatic Air-Bath	••••	47
Magnetic Faqat	• • • •	48
Solution Container	***	49
Micrometer Syringe, MS		49
Manual Manual Manual	7 4 - 5	20
Temperature Monitoring Device	•••	50
System for the Measurement of Current	2 5 A 0	Sept. 3 2 + 4
Through Solenoid	••••	52
	0000	7/4
Calibration and Calculation of the		
Densiaeter	0.7 . 4	53
A Detail Description of the Working of	4	48.0%
Magnetic Faoat Densimeter	•••	56
Dipole meter's headershad god of strik	48P	59
and the stranger	8 4 5 2	p=15-3
Types of Cell used	••••	59
Measuring Range for DFL-1 and DFL-2	少安 张 张	
type cells	••••	60
Calibration of the cell DFL-1 and E	FL-2	61
Measurement stras ef Total and	••••	61
Working Procedure		62
Constant Temporature Bath used in	***	100
Dielectric Measurement Carlot sans	••••	64
Reagents and Their Purification	••••	66
Acetic Acid. Methanel	••••	66
Purification of Solvents	••••	66

	Si u	•		
CHAP!	r er			
			7 · · · ·	
4	RESULTS		••••	69
	Syst	em: Acetic Acid in CCl4	••••	69
		Results of Specific Volume Measurement and obtaining Refined	••••	69
	- 1,0 2	values.	2 G # 3	144
	ي درود	Results of the Dielectric Measurement	• • • •	71
		Calculation of S and V and the plot	Sint	3.7
		of VS against V	••••	71
£1	20100	Calculation of P, S and PS for		
8		polarization "	• • • •	72
		A Comparative Study of VS and PS	90 S S S	
	1	Curves	• • • •	74
		The plot of V. PS versus V and Calculation of KD and K	K. S.	77
	Syst	em: Acetic Acid in Benzene	••••	80
		Results of Specific Volume and	0.112	510
	N N = 7	Dielectric Measurement and Obtaining		
	* 1	Refind Values	••••	80
	a se		10 v + V	140 8
		Plot of VS and PS Against V	••••	82
	APPLY TER	-116	8 - ' 9	生生国
	i i . Še, p.j.	\triangle (V.VS) and \triangle (V. PS) Plots.	••••	85
	Syst	em: Acetic Acid in C ₆ H ₅ Cl	••••	87
	Solv	ent Behaviour of HBH and	4000	145
	Cycl	ohexane to Acetic Acid	••••	93
	Syst	em: Acetic Acid in RBH	••••	95
	Syst	em: Acetic Acid in Cyclohexane	••••	101

PAGE

	,		•	9		
CHAPT	ER)			_		
5	DISCUSSION				••••	109
	Molar Vol	ume		*	••••	109
	Monon	er			••••	109
	Dimer	•		~	****	111
	Chloroben	zene- Co	ap 1	ex Formation	••••	114
	Molar Pol	arizatio	n		••••	117
	Equilibri	um Const	ant		••••	120
6	SUMM ARY			Tile I	••••	123
,		Suss		in an e		
	APPENDIX-A. 1	Tables		AcOH - CCl ₄ system	••••	127
	APPENDIX-A.2	Tables	for	AcOH- Benzene system	••••	132
	APPENDIX-A.3	Tables	for	ACOH- C6H5Cl system		136
	APPENDIX- A. 4	Tables	for	ACOH-HBH System	••••	140
	APP MNDIX-A.5	Tables System	for	AcOH - Cyclohexane	••••	144
	APPENDIX-B.				••••	148
				on of the Solute		
	Solvent B	_	41100	, was mores or a	••••	148
	BIBLIOGR APHY					4 67

PAGE

8

or Areales them of Essalisation

Chemical Bond and Dissociation Energy

Much of science of chemistry is concerned with making and breaking of chemical bonds. Pauling has defined chemical bond as follows: "There is a chemical bond between two atoms or groups of atom in case that the forces acting between them as such as to lead to the formation of aggregate with sufficient stability to make it convenient for the chemist to consider it as an independent molecular species".

The word 'stability' means the 'strength of chemical bond' which is given by bond energies expressed in Kcal/mole. At room temperature, the average translational energy in the direction of one coordinate is % RT or .3 Kcal/mole. A simplified expression for the fractional number of molecules possessing translation energy E or greater in two degrees of freedom is $\frac{n_E}{n_0} = \frac{E}{e^{-RT}}$. If two molecules collide, then it is reasonable to assume that the combined translational energy along the line of collision represent translational energy available for chemical reaction (breaking of a chemical bond). Ordinary covalent bonds have enthalpy between 200 — 90 Kcal/mole, whereas the forces that come into operation, which have been recognized as inter-molecular forces, involve energies from less much than 1 to 15 Kcal/mole. We can now look at the stability of "chemical bonds" by computing n_E/n_0 at 300° K for different values of E indicates that at room temperature $(300^{\circ}$ K) only 1% by molecular collision involve 3 Kcal/mole and a very negligible number of

collision, 10-42% have energy equal to or greater than 60 Kcal/mole. Thus it is clear that bond energy involving more than 20 Kcal/mole should have very little measurable dissociation of bond at ordinary temperature.

Inter-Molecular Forces

Six types of inter-molecular forces are generally recognized:

- (i) Repulsive forces, due to repulsion effect between closed electron shells which is a very short range interaction;
- (ii) Dispersion forces, which are always attractive for molecule or atom in their ground state and arise even between neutral, non-polar molecules or atoms and are due to average interaction of an instantaneous moment of one molecule brought about by charge density fluctuation and the electric moment induced on the other molecule or atom; (iii) Polarization forces, which arise from the polarization of atom or molecule by an approaching atom or molecule forcing redistribution of charges with concomitant effects on it s permanent electric moment, due to the effect of charges or permanent electric moments of the approaching atom or molecule;
- (iv) Orientation forces, due to mutual coloumbic attraction or repulsion of the net charges or electric moments carried by two interacting atoms or molecules, such as interaction between two ions, ion-dipole and dipole-dipole interaction. In a high dielectric medium at a distance

- of 5 Å E can be nearly 5 Kcal/mole for two unit of charges of opposite signs;
- (v) Mulliken charge transfer forces, leading to transfer of an electron from an electron donor molecule to an electron acceptor molecule forming a well defined complex, known as charge transfer complex, with concomitant appearance of a characteristic intense band in the visible or UV spectrum of the region². The thermodynamic parameters for charge transfer complexes are of the same magnitude as for hydrogen-bonded complex and
- (vi) Hydrogen bonding, due to involvement of specific H-atom of the proton donor group with a localized site of high electron density in the same or another molecule.

ing private of the water solveniles. These that

Repulsive, dispersive, polarization and, most of the time, orientation forces are very weak forces and hence are not "stable" enough leading to the formation of an aggregate with sufficient stability to be considered as independent molecular species. On the other hand charge transfer and hydrogen-bonding have bond energies from 30 Kcal/mole down to 4 Kcal/mole or less. The important features of these two types of complexes is that, at ordinary temperature, only a fraction of the molecule are associated and at equilibrium, due to kinetic energy of motion of the molecules, a certain number of new molecules are continually broken while the same number of new complexes are continually formed.

Hydrogen-Bond

Hydrogen-bonds are almost as frequent and almost as important in nature as ordinary chemical bonds and they will probably remain a target for intensive search for many generations to come. Hydrogen-bond formation is a special kind of dipole-dipole interaction. When hydrogen occurs linked to an electronegative element or group, the polarized H-atom permits a close approach to another negative atom having a lone pair, giving rise to a specific bond formation, represented as A — H B and termed as H-bond.

The concept of H-bond was first introduced by M. L. Huggins and then Latimer and Rodebush in 1920 during their study about the investigation of the associative nature of the water molecules. Since that time, there has been considerable discussion of a suitable definition of the H-bond 5.

A competent description of the hydrogen-bond has to be able to take into account a number of important properties of H-bonded aggregate for which extensive experimental evidence are available:

more and the contraction of a first and are appropriate to the same substitution of

(i) Molecules forming H-bonds come much closer than the sum of the vander waals radii of the nearest atoms would allow. In Figure 1.1 this fact is demonstrated with the aid of two typical H-bonded complexes - (NF)₂ and (N₂0)₂.

respective the Legion filities engineer in continuous and form publish, they

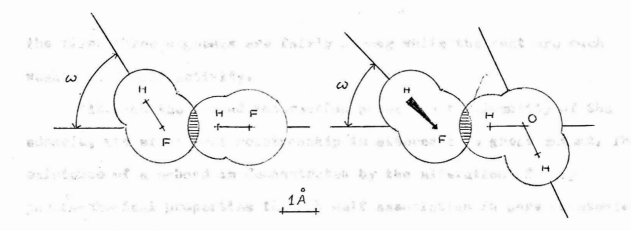


Fig. 1.1. Equilibrium geometry of (HF)₂ and (H₂O)₂. The circles around the nuclei represent the vander waals radii⁶;

(ii) R-bonding increases the polarity of the molecules concerned. The dipole moment of an H-bonded complex is larger than vectorial addition of the monomer moments would suggest:

for concluse formation between dearer and permeter enlanded to the

- (iii) As a consequence of H-bonding the lengths of the HX-bonds concorned are somewhat increased, IR stretching frequencies are shifted to smaller wave numbers and larger integral IR intensities of the corresponding absorption bands are found and
- (iv) Electron densities at the protons involved in H-bonds are decreased and consequently NMR signals are shifted to lower magnetic fields.

The most common H-bond donor segments in the acidic molecule, in order of strength, are F-H, O-H, N-H, Cl-H, S-H, P-H and in certain situations C-H (as in chloroform, where C is attached to many negative atoms or in acetylene, where carbon atoms are linked with triple bond rendering the C-atom highly negative in character) can form H-bond. Only

the first three segments are fairly strong while the rest are much weaker in N-bond activity.

Although the H-bond interaction preserves the identity of the adducts, the structural relationship is altered to a great extent. The existence of a H-bond is demonstrated by the alteration of many physio-chemical properties through self association in pure substances or by complex formation between donor and acceptor molecules in the mixtures. The following are the causes of such alterations:

- (i) Molecular aggregates of two or more come into existence, thereby altering the boiling point, melting point, dielectric constant and the colligative properties;
- (ii) As the N-bond interaction is attended with shortening of X ... Y distance, alteration of electronic environment at this zone, the density, refractive index and proton magnetic resonance are also altered and (iii) As this combination will also alter in many way the vibrational spectra of the uncombined molecules and a face new vibrations should also appear.

If these expectations are fulfilled in any particular situation, then it is assumed that H-bond exists. The effects of the H-bond on self-association or association with a donor molecule on various physico-chemical properties described giving the reasons of such changes in the Table 1.1.

Behaviour of H-bonded

Table 1. 1

Effect of Hydrogen Bonding on Some Physico-Chemical properties of Liquids in Liquid Mixture.

I have the country of the same than the same the description of the same of

	Property	compounds relative to non-bonded state	Causes of the behaviour
•	Molecular weight (by cryoscopy)	Higher	Number of molecule decreases due to H-bond intermolecularly.
•	Density	Higher	Contraction of volume.
•	Refractive Index	Higher	V_1) V
•	Dielectric Constant	Mostly higher	Vector addition of moments of
•			Resistance to breakage of H-bond.
•	Thermal conductivity	Higher	Protons high vibration freedom facilitate transfer of internal kinetic energy in the bulk.
•	Electrical conductivit	y Higher	Proton tunneling and rotation of molecule help electrical charge transfer.
٠	Acoustic conductivity	Higher	Extensive molecular arrange-
	the Valle of gard	the the matter that probable	w may the restrated to Coreta
•	Viscosity	Much higher	Longer & complex molecular size.
0.	Surface tension	Higher makent I be	Molecules held together by H-bond.
1.	Vapour pressure	lead Lower a have been	Molecules held together by
2.	Melting and boiling	Higher	Resistance to break H-bond holding together many molecules.
3•		Shift to lower field generally	Rapid exchange of H-atom between two electronic environment.

Pimentel and McClellan have thoroughly described the uses of these physico-chemical properties in their book. The importance of these properties for investigation of H-bond interaction, in order of frequency of their use, are:

- (i) Spectroscopic methods, e.g. U. V., I. R., Raman spectra, and Nuclear magnetic resonance, and
- (ii) Nonspectroscopic methods, among these methods, the dielectric properties have been largely preferred over others, mainly due to the relative case of their quantitative interpretation.

A brief discussion on the spectroscopic manifestations will convince one that M-bonding has all the attributes of a regular chemical bond with some interesting differences.

Spectroscopic Manifestations of Hydrogen Bond

Infrared and Raman spectra reveal the characteristic frequencies of molecular vibrations. Such frequencies are fixed by the masses of the vibrating atoms, the molecular geometry and the restraining forces holding the atoms in their equilibrium position in the molecule. Since the restraining force (force constant) is simply related to chemical bond order, vibrational spectra have been an excellent source of information concorning the H-bond. The different vibrational mode in the H-bonded substances are shown below:

1.
$$\gamma_S$$
 R - A - H \longrightarrow ... B \longrightarrow A - H stretch(anti-symmetric) (3500-2500 cm⁻¹, 3-4 μ)

2.
$$\mathcal{V}_{B}$$
 R - A - H ... B R - A - H bond (in plane)
(1700 - 1000 cm⁻¹, 6-10 \mathcal{M})
3. \mathcal{V}_{A} R - A - H ... B R - A - H torsion (out of plane)
(900 - 300 cm⁻¹ 11-30 \mathcal{M})
4. \mathcal{V}_{A} R - A - H ... B stretch
(250 - 50 cm⁻¹, 40-200 \mathcal{M})
5. \mathcal{V}_{B} R - A - H ... B bend (\mathcal{L}_{A} 50 cm⁻¹, 200 \mathcal{M}).

The most pronounced changes that occur are (i) A-H stretching frequency () is shifted to lower frequency, (ii) the breadth of this frequency increases markedly and (iii) also the intensity is unusually large. This anti-symmetric stretching frequency, occurs near 3500 cm⁻¹ (3 M) and these changes have properly assumed the importance of qualitative criteria and quantitative indices of the H-bond strength, bond energy and other bond properties. This stretching frequency shows spectacular changes with change of temperature, concentration and to a lesser extent, change of pressure. The increase of temperature reduces the intensity and so does the reduction of concentration, the increase of pressure, on the other hand, has the effect like the increase of concentration. The extreme sensitivity of by to these changes clearly indicates that H-bonding systems involve the non-bonded species and the bonded species in rapid equilibrium. The intensity of IR stretching frequency (\times \times_3) is much used for the measurement of equilibrium constant,

calculation of the thermodynamic quantities $\triangle \mathbb{N}$, $\triangle S$ and $\triangle F$.

K of the H-bonding in self-association or of association with another base molecule. The measurement of K at different temperatures permits

The shift of stretching frequency ($\triangle \mathcal{V}_{S}$), brought about by H-bond formation, has been found to be related to various physicochemical properties. The first and most important relationship involving $\triangle \mathcal{V}_{S}$ was proposed by Badger and Bauer 10 in 1937. Since then, through the works of many other authors 11-18, there has been a general acceptance of the proposal that $\triangle \mathcal{V}_{S}/\mathcal{V}_{S}$ provides an index of $\triangle \mathcal{H}$ of H-bond information. Also, various other physical properties have been found to be directly related to $\triangle \mathcal{V}_{S}$; such as acid strength 19, base strength 20, reactivity or rate constant 21, Hammett sigma function 22 and Henry's law constant 23.

H-bonding is the proton magnetic resonance. H-bonding causes a very large shift to low field of the X-H proton resonance. The position of the proton signal for X-H involved in H-bonding is both temperature and concentration dependent. The H-bonding being essentially dynamic in nature, the formation and breaking of the bond occurs rapidly. Because of the relatively long "observation" time of NMR measurements, only once signal is observed, corresponding to the average environment of the preton distributed between free and complexed state. The frequency of the

observed peak is thus the weighed mean of the frequencies of all the components present at the equilibrium 24.

The measurement of the association constant at different temperatures loads to the thermodynamic quantities, \triangle and \triangle s. Several workers $^{25-27}$ have obtained these quantities from the measurement of chemical shifts. Many workers claim that the change in chemical shift on 15-bonding, \triangle , is a measure of the strength of the bond. The existence of roughly linear relations between \triangle and \triangle \bigcirc_{S}^{29} , \triangle and pka or between \triangle and \triangle and \triangle indicate in a semiquantitative way the linear relationship among \triangle , \triangle , \triangle h and b and distance of X Y.

Theory of Hydrogen-Bond

A successful theory of the H-bend has to be able to explain all the proporties for which extensive experimental evidences are available. The recent development in theory made it possible to define certain contributions to H-bend energies. Estimate on the relative importance of forces of different origin (coulomb, polarization, dispersion forces etc.) became accessible to various quantum mechanical approach and give a fairly correct description of isolated dimer or in some cases, isolated trimer but fails to deal with larger clusters.

Sometime ago it had already been suggested by Coulson and Danielson and independently by Tsubomura 33 that the main problem in

the theory of hydrogen-bond is brought about by a complicated super position of the five men contributions which are of similar magnitude:

Translate

ecopy teles.

habita a strake, exiets.

- (i) Electrostatic or coulomb energy,
- (ii) Exchange and repulsion energy,
- (iii) Polarization energy,
- (iv) Charge transfer energy or covalent contribution and

wallers in with one or work

(v) Dispersion energy.

The efforts of recent calculations meet exactly these ideas.

Various methods of application of perturbation theory and M. O. methods to theoretical investigation on H-bond have been described in details in "The Hydrogen Bond", vol.-1(by Schuster, Zundel and Sandorfy)North-Holland publishing company.

All reliable calculations agree in the fact that considerable overlap between the molecule forming the H-bond occurs, and the whole phenomena can be described neither by the electrostatic theory nor by weak covalent bonding alone. A nice example demonstrating the superposition of different contribution to the H-bond energy was presented by Kollman 34, who was able to explain the stereo-chemistry of H-bonded structures by superposition of coulomb and charge transfer energy.

Classification of Molecules vis-a-vis Hydrogen-Bonding

with \$4.5 har Sala and the last and the second and the salation of the salatio

All molecules can be conveniently classified into four types with respect to their ability to participate in the hydrogen-bonding(Table-1.2).

colone is a typical stanting

money is a affect to to reid Table 1. 2

Term & place two II melecular form delimited continuous from

Classification of Molecules according to Pimentel and McClellan 35.

×	Туре	Description	Examples
		molecules with one or more donor groups (acids) and no acceptor groups	haloforms, highly halo- genated compounds and acetylenes.
	II Later	molecules with one or more acceptor groups (bases) and no donor groups	nitriles and isonitriles.
	III	molecules with both donor and acceptor groups.	alcohols, water, phenols, inorganic and carboxylic acids, primary and secon-
	Cition and 3	principles of the donor and the acre	dary amines.
	*	molecules with neither donor nor acceptor groups.	saturated hydrocarbonés, carbon tetrachloride and carbon disulfide.

Hydrogen bonding molecules are divided into types I through III, while molecules incapable of H-bonding form type-IV. The latter includes compounds which are used as the so called inert solvent in studies of H-bonded molecules.

Various N-bonded systems occur when one or more of the three types of molecules are present. The bulk of studies dealing with N-bonded systems have been performed in the liquid phase, and unless otherwise specified we will restrict ourselves to solutions. It is instructive to consider some of the types of N-bonded systems (N-bonded equilibria) formed in solutions of various types of molecules.

Type I plus type II molecules form N-bonded complexes, frequently in a simple 1:1 ratio. The strength of N-bonding depends primarily on the relative acidity of I and the basicity of II. The system chloroform acetone is a typical example:

$$\text{Cl}_3\text{C} - \text{H} + (\text{CH}_3)_2 \text{C} = 0 \Longrightarrow \text{Cl}_3\text{C} - \text{H} \dots 0 = \text{C}(\text{CH}_3)_2$$

Type III molecules can self-associate by H-bonding with themselves.

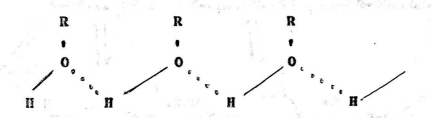
Two types of H-bonded complexes may be formed:

(ii) intromolecular, involving two or more separate molecules; and
(ii) intromolecular, involving donor and acceptor sites within the
same molecule. The strength of H-bonding depends on the relative acidities and basicities of the donor and the acceptor sites and in case
of intramolecular H-bonds, on the spatial arrangement of donor and
acceptor sites. Self association through intermolecular H-bonds can
form a large variety of open (linear) and cyclic (closed) polymers.
The exact nature of the H-bonded molecular species present in water,
liquid alcohols, and phenols, and in their solutions in inert solvents,
remains unclear inspite of a great deal of effort over the last 20 years.

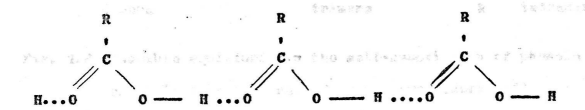
Consequence of Hydrogen Bond Formation

A direct consequence of H-bond formation is the "self-association" of a compound to give dimer, trimer, or higher polymers. Hydrogen bond formation causes water to attach itself to foreign molecules, so that their solubility in water is increased. Multiple bounded arrangements

occur in liquid water and in liquid HF and they are common in carboxylic acids, alcohols, phenols, amides, proteins, polypeptides and polyhydroxy organic and inorganic materials. How these alcohols as well as phenols and carboxylic acids form these polymers, are shown below:



(In case of alcohol and phenols)



(In case of carboxylic acids)

Self Association and Equilibrium Constant for Different Self-Associated Species

Many N-bonded molecules contain both donor and acceptor groups and such molecules can self associate through N-bonds in two or more different ways into a number of complex species (polymers, multimers, n-mers) which may co-exists with one another in a series of complicated

equilibria. The most complicated cases of self-association are the ones where the cyclic and linear H-bonded species co-exist. The best known examples are water, alcohols and phenols. Their self-association can be generalized schematically as in Fig. 1.2.

Fig. 1.2 Possible equilibria in the self-association of phenols and alcohols (The terminal OH groups are underlined).

న కేర్ ముక్రంలో అకోర్ అక్కి కారుకోంది. ఈ కోట్లాయుచ్చేంద్రీయ భారాలకేంద్రంలో ఉన్నాయి. కోర్య

It is obvious that analysis of such systems requires the determination of many equilibrium constants. Rigorous treatments using IR absorption measurements are possible in principle 36, but difficult to carry out in practice.

The usual approach in studies of self-association using IR and PMR techniques has to develop a model which assumes that only a few H-bonded species are present (usually two or three) and then to carry out computations from the experimental data to ascertain whether con-

-sistent values of the association equilibrium constants are obtained. In a typical example, ethanol was assumed to exist in CCl₄ solution as monomer, open dimer and trimer, and cyclic tetramer³⁷. This interpretation yielded $K_{\text{dimer}} = 0.95$ litre/mole, $K_{\text{trimer}} = 95$ litre²/mole² and $K_{\text{tetramer}} = 650$ litre³/mole³.

In the same study the dielectric polarization of ethanol was measured and found to increase upto a mole fraction of 0.001 and then to decrease with further increase in ethanol concentration. This evidence was judged to indicate increasing dipole moments in the order monomer < dimer < trimer, due to open or linear form, and decreased dipole moment in the tetramer, due to the latter's cyclic form.

Acetic Acid:

It is now clear that a close study of even monomer-dimer equilibria is infested with various formidable complications. The present work was undertaken to study only the monomer-dimer equilibrium of acetic acid in various types of solvents.

negman³⁸ postulated the existence of monomer-dimer equilibrium in order to explain the anomalous freezing point depressions of carbo-xylic acids in inert solvents. Since this early work acetic acid has been extensively studied³⁹ and the presence of a monomer-dimer equilibrium has been corroborated.

Using electron diffraction Karle and Brockway⁴⁰ concluded that in the gas phase acetic acid exists as a cyclic dimer. Based on a model assuming only a simple monomer-dimer equilibrium, several attempts to measure this hydrogen bonding equilibrium constant have resulted in a considerable range of values⁴¹⁻⁴⁵.

acid in Cta,. This work was estanded by Gorves and others. " . Fee ofly

In the infrared spectroscopy, the extreme sensitivity of the IR stretching frequency (\mathcal{D}_{ς}) indicates that H-bonding systems involve the non-bonded species in rapid equilibrium. The intensity of stretching frequency (\mathcal{D}_{ς}) is much more used for the measurement of equilibrium constant K of the H-bonding in self-association. It is disturbing to note that the magnitude of the equilibrium constant (dissociation of the associated species) obtained from infrared intensity data decreases rapidly with increase of acetic acid concentration $^{42-44}$.

Nowever, the dimerization of acetic acid has been studied extensively and the dimerization constant can be determined by various methods. They include cryoscopy, ebullioscopy at atmospheric and reduced pressure, Isopiestic measurements, absorption spectroscopy (visible, UV and IR) and altrasonic absorption. Useful summaries of such works upto about 1953 were included in the papers of Lefevre and Vine and Allen and Caldine 4. Almost all of the recent investigations are limited to IR 47-49, NMR 50, dielectric polarization 51,52 and ultrasonic studies 53.

It is well known that the concentration dependence of the NMR chemical shift can be used to study hydrogen-bonded systems 54. Huggins,

crack a think a condact,

Pimental and Shoolery 55 first studied the NMR chemical shift of acetic acid in CCl_4 . This work was extended by Reeves and others $^{56-58}$. Recently the NMR chemical shift of the acid proton of acetic acid in CCl, has been investigated by Goldman and Emerson 50. It was observed that concentration dependence of the acid proton-chemical-shift can not be accounted for quantitatively by a simple - monomer-cyclic dimer equilibrium. One must conclude that other hydrogen bonded species are present in solution. The observation of long chain polymers in crystalline acetic acid along with cyclic dimers in the gas phase lead one to postulate that acetic acid in the liquid state can be described as an equilibrium mixture of the gas and solid states. Recently infrared and Raman evidence indicating the presence of chain polymers has been reported . Goldman and Emerson explained the observed chemical shift quantitatively by assuming that acetic acid is present as an equilibrium mixture of monomers, cyclic dimers, linear dimers and linear polymers.

The structure of acetic acid dimer, trimer and tetramers as represented by Ritter and Simons is given in the following:

$$CH_3 - C$$
 $O - H - O$
 $C - CH_3$
 $CH_3 - C$
 $CH_3 - C - O - H - O = C - CH_3$
 $CH_3 - C - O - H - O = C - CH_3$
 $CH_3 - C - O - H - O = C - CH_3$
 $CH_3 - C - O - H - O = C - CH_3$
 $CH_3 - C - O - H - O = C - CH_3$

tetramer

As we now see, even monomer-dimer equilibrium of acetic acid in solvents is not without problems. Let us look at the following equilibriums occurring simultaneously,

$$\begin{array}{c} A+A & \stackrel{K_1}{\longrightarrow} A_2 & \text{(open chain or linear)} & \text{and} \\ \\ K_3 & \\ A+A & \stackrel{K_2}{\longrightarrow} A_2^* & \text{(cyclic chain)} \end{array}$$

walling to this to be landwared to lead at the tests bullet. It is

where.

where
$$K_1 = \begin{bmatrix} A_2 \end{bmatrix}$$
 is (quantities) observe very wide respectively.

and

$$\mathbf{K_2} = \frac{\begin{bmatrix} \mathbf{A_2} \end{bmatrix}}{\begin{bmatrix} \mathbf{A} \end{bmatrix} \begin{bmatrix} \mathbf{A} \end{bmatrix}}$$

e all world of the interestina.

$$K_3 = \frac{K_1}{K_2} = \frac{\left[A_2\right]}{\left[A_2\right]} = \frac{5,000-25,330}{\left[A_2\right]}$$

Therefore, it appears that in a simultaneous formation of linear and cyclic dimer, the ratio of concentration of two forms remain in a constant proportion and in such a case, this will appear one interaction only as $A + A \rightleftharpoons (A_2 + A_2^*)$; where a different equilibrium

constant, K, appears. The large to la sure represents the large t value.

Thus,
$$K_4 = \frac{\left(A_2 + A_2^{\bullet}\right)}{\left[A\right]} = K_1 + K_2^{\bullet}$$

These pictures are applicable to ideal solutions only. The solvent effect frequently complicates physico-chemical studies in solution; of which we get ample evidence in the present work. For this reason we must take into account the factor which may distort the equilibria of the interactions.

the different three without frame work and distart want or without

on when there the perfect wats and kenterow by the a 47.7 therefore a cra-

Finally it will be of interest to look at the table below. It is seen that the value of K (association) cover a very wide range of values from 27.7 to 5550 litre/mole.

stockels take these salv Table 1. 3 a laterestian between estreet and

Solvent	Temp. (°C)	K _{Association} (litre/mole)	Method **	References
CCl	25	5,550-3200	IR	42
gargo a seas	25	1,000-2,650	r ir s	43 3 4 1
	31	63.57 (linear)	NMR	50
	31	181 (cyclic)	NMR	50
sevel singl	25	1800	IR	47
Edult : SE Fie	Intermed forem	n which ecutivos the interval	aris oles	34.463 AM
Benzene	30	370	D. C	84
ben zene	25 25 25 25 25 25 25 25 25 25 25 25 25 2	116.3(polar)	D.C	85
which was a	140 25 Vich i	45.45(non-polar)	D.C	85
The extitions	20 wivest	27.70	IR	86
production of the second	28	500	IR	87

The lower values seem to be more reasonable. The lowest value, as seen here for acetic acid and benzene system, 27.7 litre/mole compares well with the results of this work. The reasons for such divergence of results are due to experiments at different concentration ranges, use of different theoretical frame work and disregard of other interactions exhibited by different solvent effects. Unfortunately most of these reported works do not give the experimental data. For this reason their comparison in the same theoretical frame work could not be possible.

Solvent Effects

The closest approximation to a truely ideal system is the vapour phase. In liquid state no completely neutral solvent can exist since dissolving take place only through some interaction between solvent and solute molecules. In his original development of the method for the determining dipole moment through measurement of dilute solution, Debye gives due consideration to the question of whether the values so obtained would be independent of the nature of the solvent and identical with the moments derived from the measurements on the vapour. In 1929 he suggested that, as the internal forces which control the interatomic distances in a molecule are of electrical origin, one might expect small difference, which would increase with increasing dielectric constant of a solvent. The existence of a "solvent effect" was not given full credence until attention was drawn to the matter by Muller 1935. The first attempt

of goodpans in the ention terment the governint

to evaluate solvent effect theoretically was by Weigle on the distinguished three possible factors which may contribute to the effects which are observed. These are,

in a color color des la fan auto li

e 14 a

- (a) in extra contribution to the (apparent) dipole moment, acting in the same direction as the latter, through the orientation of solvent molecules by the field around the dipole;
- (b) The change in the mean polarizability of the molecules under the influence of an external field due to the orientation produced by the dipole and
- (c) The induced moment produced in the medium through the asymmetric geometrical form of the dipolar molecule.

Since 1933 there have been innumerable attempts to derive equations relating to the apparent dipole moment of the substance in solution with the dielectric constant of the medium, with the eventual aim of permitting the calculation of the true moment of the isolated molecule from measurement on solution. The theories and emperical equation so far attempted in pure liquid state interms of their distortion polarization and dipole moment have not been completely successful. It has been generally realized that Debye theory, developed for gases and vapour and latter extended to dilute solution, where the polar molecules are

relatively far apart, is inadequate to cover the strong interactions which must occur between polar molecules in the pure liquids, and therefore the treatment of the problem suggested involved drastic modifications. None of the theories have been thought satisfactory; nevertheless they have made important contributions to the understanding of the subject. A full discussion on the subject have been excellently reviewed by Smith. The important types of interaction that take place between solvent and solute may be classified as:

- (i) Dipole dipole interaction where both solute and solvent are polar;
- (ii) Solute permanent dipole solvent induced dipole interaction where the former is polar and the latter is not;

of the fire of sounding bare trees solutioning discussioning

- (iii) Solvent permanent dipole solute induced dipole interaction for the reverse case;
- (iv) Dispersion interaction between the solute transition dipole and the dipole induced in the solvent;
- (v) Specific interactions such as hydrogen bonding, charge trans-

on 1 - 10 60

(vi) Solvent - cage compression effects on the solute where the solute occupies a volume larger than the site in the solvent quasi - lattice.

(i-iv) may be classified as general electrostatic solvent effects and much of the theoretical work on solvent shifts involves these interactions. However, where present, (v) and (vi) are likely to dominate the solvent - solute interaction and are generally treated as strong interactions leading to complex formation. But the other effects (i) - (iv) can not be avoided by choice. The effects of solvents on charge transfer complex have been extensively discussed in "Molecular Association "Vol. I, by K. M. C. Davis, (Page 150-204), Academic Press, 1975. The H-bond complexes are not immune from such effects. We will have more on these points in our discussion on the results of the present investigation.

About this Work

In the present work an investigation on only monomer-dimer association equilibrium in five solvents of diverse nature by measurement of specific volume and dielectric constant have been undertaken. The five

于电影的 化克尔克斯 人名马克斯巴拉通的大名。 蒙然 图象

religious to a la come recognition has builded a alternative. L'union hacament rans

The structure of chiesenseastern it caves may

solvents are: we are increasing by way, (i) paditionly painting

- (i) a Carbon tetrachloride, and the for intermediate. We will find
- (ii) Benzene, a sy there willed by our present largetter days.
- (iii) Chlorobenzene,
- (iv) Cyclohexane and
- (v) High boiling hydrocarbon (HBH) (150°C 200°C).

Carbon tetrachloride: Although it is ordinarily thought that this solvent is inert and have very little effect on solute molecules, it is to be noted that sharpe and walker have published several papers concerning interactions of carbontetrachloride with alkyl and heterocyclic amines, ethers and sulphides 64-66. In general the interaction of poly halogenated alkanes with a variety of lone-pair donors are no longer in doubt 67.

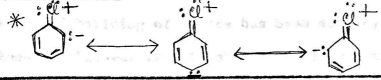
I therefore in the compact water there are a compact

(1) 1 (1) (1) 1 (

Benzene: This solvent is widely known to be a T-electron donor 68 which readily form a weak H-bond, O-H-OAC with highly polarized H-atom such as alcohol, carboxylic acids etc.

安全一支法主义的新闻 水管 化中央水色系统 计复杂符 电影光点中态设置 电双角电视多种共享多效 通报法 數數 医皮肤红色

Chlorobenzene: The structure of chlorobenzene is given as,



^{*} Taken from "Physical Chemistry" by W. J. Moore, page-706, Prentice-Hall, Inc., Englewood Cliffs, New Jersey, 1972.

[&]quot;Electric Dipole Moments" by Smith, Page-200, London Butterworths Scientific Publications, 1955.

Thus we have two most important sites, (i) positively polarized C1-atom and (ii) donor ring orbital for interactions. We will find profound effects of these sites in our present investigation.

Cyclohexane and IMH: Although, in chemical sense, these two solvents should be 'inert' in comparison to all other kinds. We have here a new kind of effect which will be called "solute - hole interaction".

In this work it is proposed that, at first, a very detailed picture of variations of a measured property should be build up (in our case specific volume and dielectric constant), by very accurate and careful experiment approaching from very high dilution to more concentrated solution: so that it is visibly evident that some kind or other interactions are most probably taking place within the range. The importance of finding the ranges of various interactions is that the variations of property with changing concentration can be examined analytically so that it become pertinent to subject them to particular calculation of important parameters. The measurement specific volume has been done by "Magnetic Float Densimeter" and dielectric constant by "Dipole Meter" using a very large cell; the examination of error of data and building of curves has been done by a powerful graphical method known as "Linear Transform" and the examination of these Variations of property have been done by a new formulation of variation of apparent molar properties with the change of concentration.

more than bendering

relief of the sea herman amile

An the surrower of Temples, therefore, the distance because A set D decord to be received to the fellowing distance.



THEORETICAL DISCUSSION

THEORETICAL DISCUSSION

THEORETICAL DISCUSSION

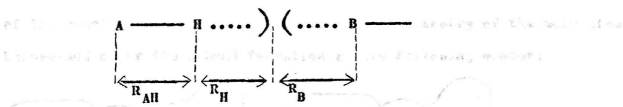
TO A CONTROL OF THE PRODUCT OF THE PROPERTY OF THE PRO

The mindy word on the party-landing of vincer week's rolls of a not a new the rolls of a not a new the rolls of a not a new a new a new and twitten and a new a new a new and a new a new and a new

THEORETICAL DISCUSSION

Volume Changes on Hydrogen-Bonding

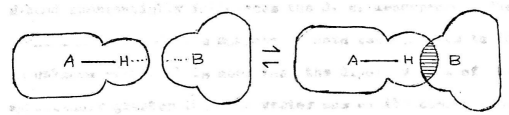
In the absence of H-bonding, therefore, the distance between A and B should be as represented in the following diagram:



where R_{AH} is the covalent bond distance of A —— H, R_{H} is the vander waal's radius of the H-atom and R_{B} is the vandar waal's radius of the B-atom. Therefore, the closest distance between A and B in absence of any interaction should be equal to ($R_{AH} + R_{H} + R_{B}$). Therefore, the observed distance between A and B when subtracted from this calculated distance, gives the contraction of A B distance on H-bond formation, denoted by S(A....B), and similarly the H....B distance is less than $R_{H} + R_{B}$ by an amount S(H....B). It is to be noted that S(A....B) is not exactly equal to S(H....B) and which indicates that the H-bonding is never without an effect on the length of the original covalent A-H bond. This effect seems to be the expansion of this bond under some sort of an attractive force on the H-atom.

The study made on the over-lapping of vander waal's radii of H and B and the position of H-atom with relation to A and B by Barman and Rahim is worth-noting 6, where it is indicated that H-bond formation not

only shortens the H....B distance but also leaves an overall shortening of A....B distance of comparable in magnitude and which must cause
an overall volume contraction. Thus, there is an excellent possibility
of detecting the H-bond formation by measuring the volume contraction
of the reacting species. We may represent the geometry of the molecules
before and after the H-bond formation in the following manner:



Overlapping zone, indicated by the shaded portion, represents the volume contraction consequent to the shortening of H.... B distance. Therefore, the important postulates are:

marks for fine according at fortax to Levister Belsent stop for

- (i) There is a net definite volume contraction for the formation of particular molecular complex, and
- (ii) The shorter is the H-bond interaction the greater is the volume contraction.

Dielectric Changes on Hydrogen-Eonding

The formation of a H-bond A-H....B leads to an increased polarity of the bond A-H and hence, to a large d. c. and greater dipole moment.

The d. c. (permittivity) is proportional to M. Thus, the two dipole A-H.

each of moment $\mathcal M$, associate to form a structure - $\mathcal M$ -H... A-H of moment 2 $\mathcal M$, the increase of permittivity is proportional to an increase of 2 $\mathcal M$ to $(2 \mathcal M)^2$. This has been viewed for basic reason for the abnormally large permittivity of water.

Mutual polarization of the donor and acceptor molecule involving II-bond substantially influences the d. c. measurement. These polarizations change the dipole moments of both the partners in the complex by an unknown amount it is seen that the dipole moment of the complex is appreciably greater than the vector sum of the components. The use of permittivity measurements in the study of intermolecular H-bonding in solutions is exemplified by the isomers of octanol by Dannhauser and his collaborators ⁵⁹. It is possible to distinguish linear and cyclic self-association of various alcohols.

When the complexes studied dissociate strongly we have specific design of experiment and specific design of mathematical processing for the different physical methods. There are general approaches for studying weak intermolecular reaction some of which are graphical interpolation method which are widely used and can be termed "Linear-ization of relation with insufficient number".

Theoretical Treatment

present work utilizes a new theoretical frame work developed

It use the maker gramming apporting property was

erry, thought of those to a directation of the

by Rahim⁷¹. The ingeneous way it has been developed is presented in the following:

1. The molar property of a pure substance is calculated by multiplying the specific property by its molar volume and the molar property
of a solution is calculated by multiplying specific property of the
mixture by its molar volume, which can be expressed as:

 V_m here is the volume in which $\sum_{i=1}^n$, where x_i , the mole fraction of ith species and thus molar volume is calculated by $V_m = \sqrt{2}x_i$ M_i in which $\sqrt{2}$ is the specific volume of the solution $\sqrt{2}$.

ty w. . the sold fraction of minimum

There are various molar properties which are found to be structurally additive (vectorically or otherwise, such as, molar volume, molar refraction and molar polarization). These properties seem to follow simple additivity rule in mixtures where no appreciable interesaction between species is expected; the additivity rule can be expressed as,

where, P_i , p_i and \mathcal{L}_i are the molar property, specific property and mole fraction of ith species. Whenever there is a deviation of the observed molar property P_m from P_i , an interaction between species are

taken into consideration. This rule holds fairly well so that this deviation of molar properties are widely used in the studies of various molecular interaction in solution 73-80.

Suppose we have \mathbf{x}_A mole of a substance A, which remain in solution in various polymeric forms, such as, monomer, dimer, trimer etc., designated as \mathbf{A}_1 , \mathbf{A}_2 , \mathbf{A}_3 , in a solvent B, then the molar property would be $\mathbf{P} = \{\mathbf{p}_i\mathbf{x}_i + \mathbf{P}_B | \mathbf{x}_B(2.3) \}$ where \mathbf{P}_B and \mathbf{x}_B denote molar property and mole fraction of the solvent B. Let us divide both sides of equation (2.3) by \mathbf{x}_A , the mole fraction of substance A and we have the equation (2.4)

$$\frac{P}{X_A} = \sum p_i \frac{X_i}{X_A} + P_B, \frac{X_B}{X_A} \qquad 2.4.$$

The equation (2.4) can be expressed as.

$$\overline{P} = \sqrt{p_i X_i + s} \qquad 2.5$$

where, $\overline{P} = P/x_A$ and $S = \frac{P_B x_B}{x_A}$ and X_i is the mole fraction of ith species which is equal to x_i/x_A . Please note that the mole fraction, X_i is now calculated only in terms of species A alone and solvent species is kept apart being calculated on the basis on its actual presence per mole of substance A.

Suppose one mole of A is dissolved in X_8 mole of solvent B in which only monomer and dimer exist. In that case, the equation (2.6) becomes $PS = P_1X_1 + P_2X_2 + \cdots + P_3X_4 + P_3X_5 + \cdots + P_3X_4 + P_3X_5 + \cdots + P_3X_5 + \cdots$

Remembering that two molecules of monomer join to form one molecule of dimer and that the total monomer, either as dimer or monomer, must be equal to one mole,

is to be a critical on to love a

We have.

$$X_1 + 2X_2 = 1$$

$$X_1 = 1 - 2X_2 \qquad ... \qquad 2.8$$

Therefore, $PS = P_1(1-2x_2) + P_2x_2$

Or,
$$PS = P_1 + X_2(P_2 - 2P_1)$$
.

where $D = P_2 - 2P_1$, which is the change of molar property due to formation of 1 mole of dimer from two moles of monomer. Therefore, we write, from equation (2.9)

$$x_1 = 1 - \frac{2(P_S - P_A)}{D} = \frac{D - 2P_S + 2P_A}{D}$$

Now, substituting $D = P_2 - 2P_1$ in the relation (2.8), which is

$$X_1 = 1 - 2X_2$$
, we have
$$X_1 = 1 - 2(PS - P_1)/D$$

$$Or, X_1 = \frac{D - 2(PS - P_1)}{D}$$
2.11.

2. If in a solution species A are in different polymeric forms A_1 , A_2 , A_3 etc. which remain in equilibrium as follows:

which is expressed in mathematical form,

$$K_{2} = \frac{\left[A_{2}\right]}{\left[A_{1}\right]^{2}}$$

$$K_{3} = \left[A_{3}\right] \left[A_{1}\right]$$

$$K_{4} = \left[A_{4}\right] \left[A_{3}\right] \left[A_{1}\right]$$

$$Or, K_{5} = \left[A_{3}\right] \left[A_{1}\right]^{3}$$

$$K_{6} = \left[A_{4}\right] \left[A_{1}\right]^{4}$$

where, $K_5 = K_2 K_3$ and $K_6 = K_2 K_3 K_4$; $\begin{bmatrix} A_1 \end{bmatrix}$, $\begin{bmatrix} A_2 \end{bmatrix}$, $\begin{bmatrix} A_3 \end{bmatrix}$ refer to concentration in mole/litre of particular species and K with subscripts refer to different equilibrium constants. It is obvious that when solution of A in B is very dilute the higher species tends to decrease.

Suppose, the concentration is so low that only monomer and dimer are in sufficient quantity to have perceptible effect on the observed properties of the mixture, then the equation (2.11) and (2.13) become useful. The equation (2.13) is now written in the following form:

$$K_2 = \frac{X_2 \cdot V}{(X_1)^2}$$
 2.14

because,
$$\begin{bmatrix} A_1 \end{bmatrix} = \frac{X_1}{V}$$
 and $\begin{bmatrix} A_2 \end{bmatrix} = \frac{X_2}{V}$

On substituting the values of X_1 and X_2 from equation (2.10) and (2.11) into the equation (2.14) we write,

$$K_2 = \frac{\left(\frac{P_3 - P_1}{D}\right) \vee \left(\frac{P_2 - 2P_3}{D}\right)^2}{\left(\frac{P_2 - 2P_3}{D}\right)^2}$$

or,
$$K_2 = \frac{p(PS - P_1) \vee (P_2 - 2PS)^2}{(P_2 - 2PS)^2}$$

$$\frac{\left(\frac{P_2 - 2PS}{D}\right)^2}{\frac{B(PS - P_1)}{(P_2 - 2PS)^2}}$$

$$PS = -\sqrt{\frac{D}{4K}}\sqrt{\sqrt{V} + \frac{P_2}{2}}$$

For simplicity we write $K_2 = K$ and then transposing we have,

$$\frac{K}{D} \times \frac{(P_2 - 2PS)^2}{V} = (PS - P_1) \qquad 2.15$$

Again transposing, PS =
$$\frac{K}{D} \times \frac{(P_2 - 2PS)^2}{V} + P_1 + P_1 + P_1 + P_1 + P_2 + P_3 + P_3 + P_4 + P_3 + P_4 + P_5 + P_$$

In this equation observable are only PS and V and there are three unknowns in an involved form, A direct solution of this equation is not possible.

3. When the solution is very dilute the monospecies predominate; this means, PS tend towards P_1 and therefore P_2 —2PS becomes nearly equal to P_2 —2P₁ i.e., equal to D. Thus, by approximation, we arrive to relation from equation (2.16):

The compared
$$PS = \frac{K}{D} \cdot J \frac{D^2}{V} \cdot P_1$$
 and is simultance to equation (2.77) where the product is a raise (2.77) if the raise (2.77) if the product is a raise (2.77) if the raise (2.77) if

and two assignments of the process of
$$V$$
 when the state of the content V are the process of the process of the state of the content V

we can also write,
$$V.P. = P.V + KD$$

Hirds of disconfinentials, (i) each

Here we arrive at two most important relations, equation (2.17) and (2.18). Equation (2.17) shows that PS plotted against 1/V give a straight line, in which a dimer association interaction is taking place, whose slope give KD and intercept give P₁. The plot PS versus 1/V has

utilized to obtain P4. There should be no doubt that the plot of PS versus 1/V should give only one straight line in absence of any association interaction; this should be evident by the following reasoning. Equation (2.5) give the relation,

$$\overline{P} = \sum P_i X_i + S$$

$$or_2 \overline{P} = S = \sum P_i X_i$$

Therefore, $PS = P_4X_4$ when only one kind of species is present in the solution when without doubt X4 = 1 .

min record some gas error of

while at the my epolice Albert wind

This shows that PS remain constant and is also true in equation (2.17) when KD = 0, giving a straight line parallel straight line to axis, 1/V. In practice, the zero (0) slope is never realized because of the presence of some kind or other interaction between solvent and solute and the extrapolated volue of P, thus obtained, indicate the molar property of the monomer species in the environment of the dominant solvent atmosphere.

There are possibilities of two kinds of dimer formation, (i) open chain dimer and (ii) cyclic dimer. These two types of interaction take place in the same solvent, but either as simultaneous or consecutive interaction, at first one of the kind is formed and after it reaching a point the next one begins to be formed. In these cases, we should

have two linear portions in plot, PS versus 1/V (or V.PS versus V plot). In case of simultaneous interaction in which both the kind appear in solution following their own equilibrium conditions. We should have only one linear section related to dimer interaction because they will appear as one interaction. Although the reason for this has already been discussed in the Introduction, yet here it is:

Let
$$K_{(1)} = \frac{\begin{bmatrix} A_2 \end{bmatrix}}{\begin{bmatrix} A \end{bmatrix}^2}$$
 and $K_{(2)} = \frac{\begin{bmatrix} A_2^* \end{bmatrix}}{\begin{bmatrix} A \end{bmatrix}^2}$

where, $K_{(1)}$ and $K_{(2)}$ are for open chain or cyclic dimer equilibrium in simultaneous interaction of the type shown below:

(i)
$$\Lambda + \Lambda \rightleftharpoons - \Lambda_2 - \text{(open chain)}$$

and (ii) $A + A = A_2$ (cyclic)

$$\frac{K_{(1)}}{K_{(2)}} = \frac{\begin{bmatrix} A_2 - \end{bmatrix}}{\begin{bmatrix} A_2 \end{bmatrix}}$$

$$\begin{bmatrix} A_2 - \end{bmatrix} = \frac{K_{(1)}}{K_{(2)}} \cdot \begin{bmatrix} A_2^* \end{bmatrix}$$

Thus substituting this relation in either of the two equilibrium equation

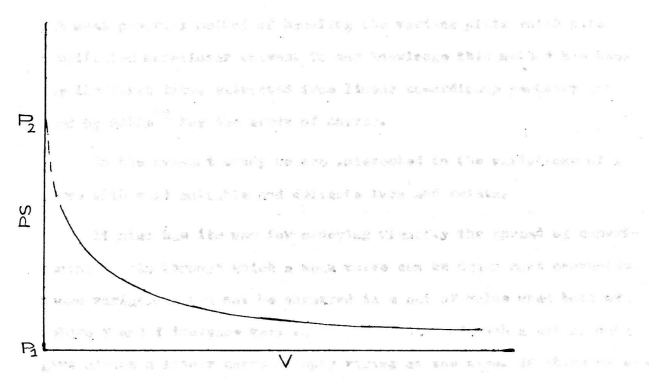
$$K_{1} = \frac{K_{(1)}}{K_{(2)}} \cdot \left[A_{2} \right]^{2} \qquad K_{2} = \left[A_{2} \right]^{2}$$

This means that both the species, open and closed dimer appear in this solution in a constant ratio to each other and they appear to be only one interaction which is simply an interaction of dimer mixture of constant ratio. (Equation (2.18) has its usefulness because the KD appear as intersect which is far more easier to determine. It does not mean the equation (2.17) and its plot can not be used for finding KD, appearing in the slope of the linear portion of the curve; but the determination of any slope value is always, for that matter, subject to some confusion and error.

Thus, finding PS and V from experiments in very dilute solutions and using the relation (2.17) and (2.18) we can find, at least approximately the value of KD and also P_1 . But unless we find the value of D, which is equal to $P_2 = 2P_1$, we can not find K. Thus, the problem now is to find the value of P_2 .

that when V is made bigger and bigger i.e. when the solution is progressively diluted, PS must and being almost equal to P₁ and on the other hand when the volume V is continuously made smaller i.e., the solution is progressively made concentrated then PS approaches slowly towards P₂. We have calculated PS and V by using a suitable computer programme. Using the equation (2.15) for a system in which only one

dimer is formed, whose $P_1 = 55$ c.c., $P_2 = 2\times60$ c.c. and K = 100 l/m. The plot of PS versus V presented in the following Figure:



In this curve it appears that a careful extrapolation to V=0
we can get a reasonable value of P₂ which should help us to find an acceptable value of K and D. It must be noted that in various methods that have been used at different lines using various properties the procedure for finding P₁ and P₂ is always extrapolation. The difficulties of separating D from KD has always been recognized at least in bear interactions. It must be said at this point that the procedure that has been developed here is no worse, if not better than others.

Thus finally is he find KD from that
PS . Vs. & (refer equation 2.17) and already tre
have Pr and Pr as directed about is therefore
the can obtain K brownij that D= 2P1-P2.

Linear Transform , this should and policyst it from the walness of

But before closing this chapter let us discuss an unfamiliar but most powerful method of handling the various plots which give complicated non-linear curves. To our knowledge this method has been for the first time, extracted from linear co-ordinate geometry and used by Rahim 81 for the study of curves.

In the present study we are interested in the variations of a curvo with most suitable and delicate turn and twists.

It also has its use for studying visually the spread of experimental points through which a mean curve can be drawn most conveniently. These variations can not be observed in a set of value when both of values X and Y increase very rapidly. The plot of such a set of data give almost a linear curve steeply rising at the time. If there be some small variations, even a plotting on a very big graph sometimes are not of much use. In these cases "Linear transform" is very helpful.

even in a statement, the transfer of to

softe by linear entranclistions, has been entra-

Suppose we have a curve which is, in reality, an interconnection of g two straight lines at a very wide angle of very nearly of 180° whose equations are(a) $Y=a_1x + b_1$ and (b) $Y=a_2x + b_2$ and whose common point, of course, is at $x = \frac{b_1 - b_2}{a_1 - a_2}$. Now let us transform this equation with reference to another straight line which passing very closely to these straight lines whose equation is $Y=a_3x + b_3$. Let us calculate

 Y_1 at a value x by this equation and subtract it from the values of Y, obtained for either equation (a) or (b) to obtain $\triangle Y=Y-Y_1$, then we have, for straight line of equation (a), $\triangle Y=(a_1-a_3) \times +(b_1-b_3)$ and for the other equation (b), $\triangle Y=(a_2-a_3) \times +(b_1-b_3)$. These two again, as we can see, are straight line equations whose common point, of course, is at $x=(b_1-b_2)/(a_1-a_2)$. Thus we see that $\triangle Y$ versus x give a curve whose intersection point in reference to x remain unchanged and the angle become comparatively acute. Thus it is evident that by suitable choice of straight line equation as a reference, the transformed value of Y, $\triangle Y$ versus X can give a kind of plot which is not only equivalent to using a very large graph but also the increase of sharp and more acute bending of curve give a far greater insight into nature of the curve. These findings will convincingly be clear in our first example of study on acetic acid in CCl_4 system.

This method of studying curves, help to find accurately the slope and determining intercepts by linear extrapolations, has been extensively used in the present work.

The following the same

or on the besidence in the book of this

EXPERIM INTAL

and the contract of the contra

EXPERIMENT AL

Description of the Magnetic Float Densimeter

The magnetic float densimeter used in the present investigation is similar to one developed by Barman and Rahim which is based on the principle to the one developed by Lamb and Lee 83 with one important modification.

A float made of glass bulb in which a permanent magnet is fixed at the lower end. The weight of the float is so adjusted that it just floats when immersed in a liquid (whose density is to be measured) within a glass vessel. A solenoid is attached at the base of this vessel. When a large amount of current is passed through the solenoid the magnetic field so produced drags the float down to the bottom. When the current is decreased slowly a point is reached when the magnetic float just begins to float. At that point the current is measured, termed as the "equilibrium current". Then the weight of the liquid displaced by the completely immersed float must be equal to the sum of the actual weight of the float and a weight which is equivalent to the magnetic attraction between the solenoid field and the magnet. As the magnetic attraction is directly proportional to the applied current, we can write,

where,

w, = the weight of the float

v. = volume of the float

d = density of the experimental liquid

i = the minimum current required to hold the float bottom

k = weight equivalent of the current.

The magnetic float densimeter consists of the following assembly:

- (i) A thermostatic air-bath (Fig. 3.2) in which the densimeter is placed (Fig. 3.1) with the help of a clamp and stand. The different components of the densimeter are,
- (ii) A magnetic float(C) which is immersed in the liquid in the container;
- (iii) Solution container, (A) in which liquid is taken whose density is to be measured. A quickfit lid(E) is to cover the mouth of the container;
- (iv) A micro-meter syringe(MS) by which one of the components is dropped to the container to vary the concentration of the solution in the container, and a syringe(PS) is attached with a suction arrangement for homogeneous mixing of the mixture and
- (v) Contact thermometer(CT) (Fig. 3.2), fans (F₁, F₂), electric bulb
- (L) is for monitoring the temperature of the air-bath.

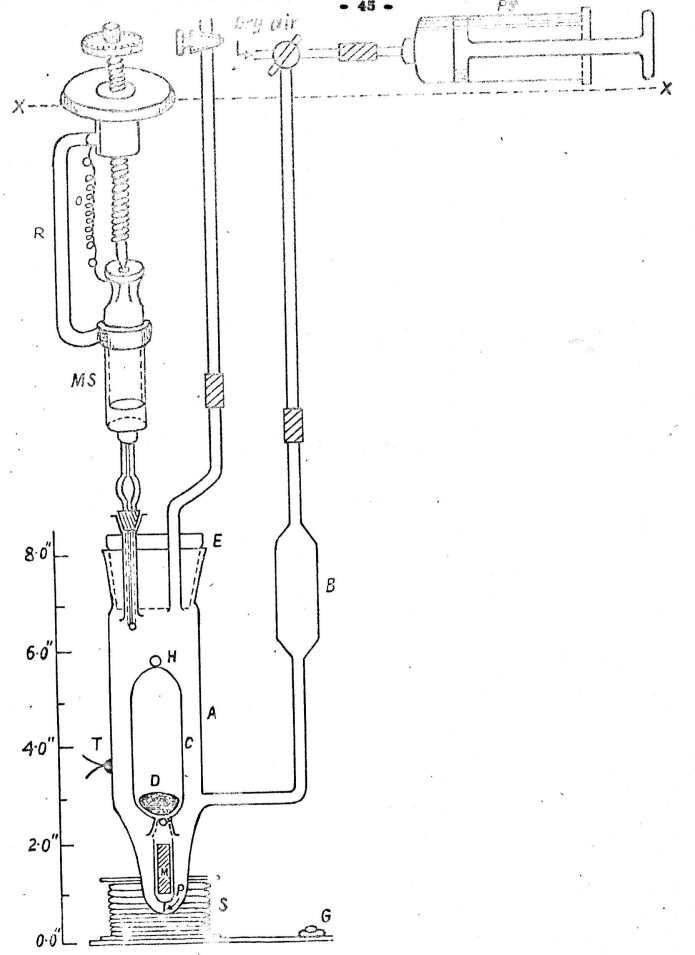
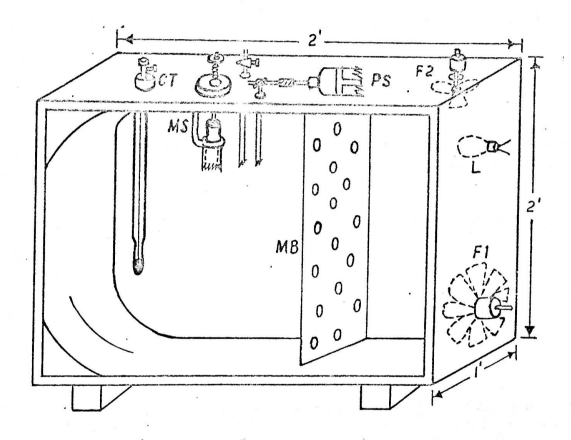


Fig.-31 Schematic of the Numetic float densitemeter. A - solution container, B - Bulb, E - Quicklit lid, C - Magnetic float, D - Mercury, M - Magnet, P - Platinum tip, H - Platinum ring, T - Thermistori, S - Polenoid, I - Spirit level, MS - Micrometer syringe, PS - 20 cc. syringe, R - Brass (2012, I ... I represent of the thermostatic air bath showing the



PS - 20 cc. syringe, FI - and F2 - Fans, L - 60 watt bulb, M8 - A thin metal foil barrier with perforations. (Electric bulb and contact thermometer are connected to an electronic relay).

a thermostatic air-bath: The densimeter assembly is placed inside a thermostatic air-bath. It is a double walled hardboard box of 60x30x60 cm. The external surface of the bath has been heavily insulated with polyurethane foam sheet. The corners of the left wall of the box have been made round. The front wall of the box is the door, which is made of glass and covered with insulating materials on both sides, keepinga portion bare for observing the dessimeter assembly. The thermostatic system consists essentially of two fans, a 60w electric bulb, a contact thermometer, and an electronic relay as shown in (Fig. 3.2). The electrical connections of the control system are shown schematically in (Fig. 3.3).

There is a metallic barrier containing a few holes separating the heater and fans from the densineter assembly for fair mixing of hot and cold air. For the maintenance of a nonfluctuating and uniform temperature around the liquid container and micrometer syringe, these two are placed inside a replaceable metallic enclosure. The metallic enclosure has been provided with a transparent window for viewing the movements of the float within the solution container.

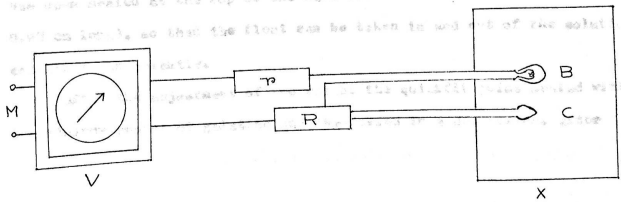


Fig. 3.3. Electronic Connection of the Control System.

drying, it is immersed in dry and pure CCl₄. Before starting work the float is rubbed with a piece of dry cloth and then weighed.

about 100 c.c. capacity (diam. 1.8" and height 7"), is provided with a side tube having a bulb, B. The top of the container is fitted with a quickfit lid, E. This lid consists of two narrow holes: one being pressure outlet and another for dropping one of the components of the mixture from a micrometer syringe, MS. The solution container is fitted with a solenoid, S, at the bottom. The solenoid of length 3.0 cm. consists of 11500 turns of 42 gauge enamelled copper wire wound on a plastic former and its resistance is about 2000 ohms.

There is a bakelite base fixed below the solenoid. A spirit level, G, fixed on the base, is used for getting a definite position of the solenoid relative to those of magnet in the float and earth's magnetic field.

The solution container can be held in position by clamp and stand arrangement (not shown in the figure). Only non-magnetic materials have been used in the construction of the stand. The spirit level below the solenoid can be adjusted by screw legs in the base of the stand.

(iv) Micrometer Syringe, MS: The micrometer syringe is a clinical syringe, MS, of 5 ml. capacity, which is attached to the brass frame R (Fig. 3.1). The piston of the syringe can be moved by means of a micrometer, and a spring G, which are attached with the brass frame. It

has a long glass needle (8.75 cm long) with a small bulb at the same middle. In the same to the heidless should be bridge sixents to be

The micrometer syringe is calibrated in situ by dropping purified kerosene of known density (on turning a certain number of terns)
from the micrometer syringe into a small weighed conical flask and
weighing the conical flask with kerosene.

varied by decopping one of the components from the micrometer syringe.

The resulting solution can be made homogeneous by stirring it with the help of up and down movements of the float and also by transferring some liquid in the side tube, B, with the help of the suction arrangement attached to one of the tube through the lid of solution container (Fig. 3.1). The float can be danced if the current passing through the solenoid is followed by a sudden cut.

mental liquid has been monitored by a bead type thermistor (F22, standard Telephone and Cable Co.) whose resistance is about 125 chas. at 30°C having temperature coefficient approximately 4% per °C. The thermistor has been fixed on the glass surface of the liquid container at T (Fig. 3.1) and it is used as an arm of a wheatstone resistance bridge as shown in (Fig. 3.4). The other three arms of the resistance bridge, each about 125 chas, have been made of high resistance Nichrome wire and these are placed inside the thermostatic air-bath. The e.m.f.

across the bridge is maintained at 1.3 volts. and a sensitive galvanometer is connected to the bridge. Wheatstone bridge circuit containing the thermistor gives a balance in the galvanometer at a definite temperature $(30^{\circ}\text{C} \pm .005^{\circ}\text{C})$. The necessary measurements for density data are made when the spot of the galvanometer remain within - 0.1 cm. to + .01 cm. of the scale.

The part and the property of the second state of the second state

Fig. 3.4 Wheatstone bridge circuit.

than inarthered with seal the equal thrive autrent is edicined from the

A thermister is attached on the glass of the solution container. It is covered thickly with wax so that the short term temperature fluctuation in the air-bath may not have serious effect on it and the temperature of the solution can easily pass through the glass to the thermister for monitoring.

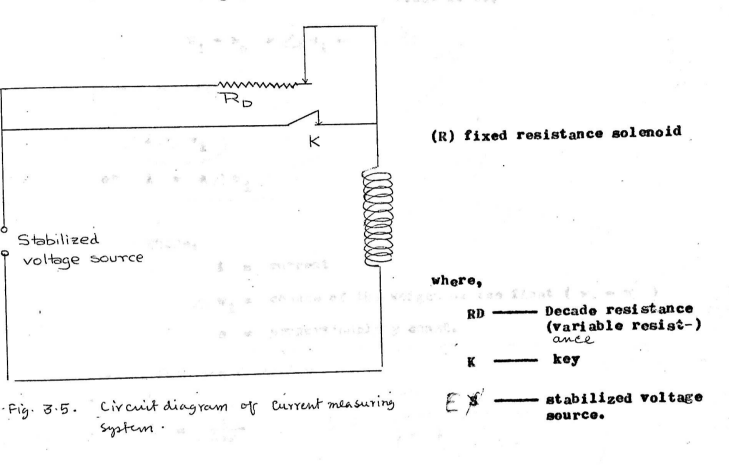
System for the Measurement of Current through Solenoid

In Barman and Rahim's system the equilibrium current is supplied from a battery through a fixed resistance to the solenoid, all connected in a series, in which current is measured with the help of a potentiometer by noting the potential drop across the known fixed resistance. In this system one would require a battery for supplying the current to the solenoid, another battery for the potentiometer and a standard cell to standardize the potentiometer. A lot of time and attention is thus wasted to note the actual potential drop each time. Another source of genuine trouble is the short term fluctuation of voltages of these batteries.

In our modification the current is now supplied from a stabilized voltage source. Thus the use of potentiometer and batteries are altogether despensed with and the equilibrium current is obtained from the reading on decade resistance box, RD (Fig. 3.5).

count the Italian temperature of the late. But work he dispersity present the country

The circuit diagram is shown below:



Calibration and Calculation of the Densimeter

To establish a relation for the calculation of the specific volume, the weight of the float is so adjusted that when it is immersed in a solvent in the solution container the equilibrium current is zero. Suppose that the weight of the float at that condition is \mathbf{w}_0 and the resistance is \mathbf{r}_0 . Again suppose the weight of the float is increased to \mathbf{w}_1 and \mathbf{w}_2 for which the resistances obtained are \mathbf{r}_1 and \mathbf{r}_2 respectively.

Therefore, the equilibrium current passing through the solenoid across the fixed resistance(R) (Fig. 3.5) must be directly proportional

to the change of the weight of the float i. e..

Therefore, we can write,

or
$$i = a \triangle w_1$$

Where,

i = current

 $\triangle w_1$ = change of the weight of the float ($w_1 - w_0$)

a = proportionality const.

Again, from the Ohmas law

ETYP

where,

S = voltage

R = fixed resistance

r = variable resistance.

Therefore, from the above relation,

$$\frac{\$}{R+F_4} = a \triangle w_1 \qquad \dots \qquad 3. 1$$

The Where, a sed the is properly and the Action in a -- The e-

$$\triangle v_2 = v_2 - v_0$$

bna

 r_1 and r_2 are variable resistance for v_1 and v_2 respectively.

Prince & our will risk to passived their the retirement

From equation (3.1) and (3.2)

$$R = \frac{\triangle w_1 r_1 - r_2 \triangle w_2}{\triangle w_2 - \triangle w_1} \quad \text{and} \quad S/a = \triangle w_1 \quad (R+r_1)$$

$$er_{\bullet} \triangle w_2 \quad (R+r_2) \quad .$$

At equilibrium current the equivalent weight .

$$w_1 = \frac{S/a}{R+r}$$

Knowing the value of w, specific volume can be calculated from the following equation:

peak is noted this this election that is not worth, the famous waits

Results and Calculations

Weight of float
 Resistance

$$w_0 = 22.56056$$
 $r_0 = 7713$
 $w_1 = 22.56695$
 $r_1 = 7769$
 $w_2 = 22.57105$
 $r_2 = 7834$

The value of R and S/a is calculated and the value is R =-7935.29 and S/a = (-1.06256)

Volume of the fleat is found out from the relation (3.3) by a solvent of known specific volume and it is found $V_r = 25.9976$ c.c.

Hence, specific volume can easily be calculated by reading the resistance from the resistance box.

Which when being the bear of the control of the colors

A detail Description of the Working of Magnetic Float Densimeter

- (i) A stoppered dropping funnel is filled with the purified and dried solvent and weighed. Then the solvent is poured into the solution container(A) (Fig. 3.1), close the lid(E) and weight the funnel again. The difference of the weight gives the weight of the solvent taken in the container. The volume of the solvent in the container should be such that the float can completely sink in the liquid when pulled down by magnetic force.
- (ii) The adjusted float (described in the magnetic float heading) is then immersed in the liquid of the container by opening the lid(E) (Fig. 3.1) and close it again.
- (iii) By screw adjustment the spirit level is brought into proper position.

- (iv) The container is enclosed in the replaceable metallic enclosure (described earlier in the heading thermostatic air-bath).

 The micrometer syringe(MS) is filled with the mixture of accurately known composition (acetic acid and solvent, usually 6%) and introduced into the solution container.
- (v) The door is closed and the instrument is switched on and the thermostatic air-bath is allowed to attain equilibrium temperature which takes over an hour, which is indicated by the galvanometer of the thermister circuit (Fig. 3.4). During this time the liquid in the container is stirred at intervals of 5 10 mins. by up and down movements of the float and also by transferring some liquid in the side tube B (Fig. 3.1) (as described earlier).
- (vi) Then the resistance, r is read from the decade resistance box. Special care should be taken to take the final reading. At first by trial and error reading is taken which is say 3236. Next time to start with the resistance is fixed at 2000. The float dragdown to the bottom of the container by "hold down current". After sometime the resistance is increased by one thousand, then by one hundred at each step and then by 10 \(\text{\text{till}} \) it reaches 3230. Then by care and caution the

when a large amount of current is passed through the solenoid across the fixed resistance (R), it drags down the float to the bottom of the container. This current is termed as hold down current.

resistance is increased by one thousand, and then by one hundred at each step and then by 10Ω till it reaches 3230. Then by care and caution the resistance is increased by one Ω and allow some time to increase by another Ω till the float just begins to rise up. This is done as the current gradually reaches to the equilibrium current the hold down current becomes zero. The float may begin to rise before equilibrium current is reached due to the agitation or mobility of the liquid.

It should be remembered that temperature control is the most important factor for obtaining a good result. When the reading is noted the care should be taken that the spot of the galvanometer remain within ± 0.1 cm of the scale.

(vii) The concentration of the liquid in the container is now varied by adding a desired amount of mixture from the micrometer syringe followed by homogeneous mixing by the method described earlier. The system is again brought to the equilibrium temperature and the resistance for equilibrium current is recorded. Thus, by repeating this process, data for several concentrations which lie very close to each other within a particular section are obtained. At last the float is removed from the container, it is dried by rubbing with the cloth and kept in dry CCl₄ solvent.

The float can be readjusted and used for other system.

Dipolemeter

The dipolemeter, type DM-O1, represents a precision instrument for measurement of the dielectric constant of a pure liquid or mixture measurable upto the 5th place of decimal with measuring sensitivity (4-8)X10⁻⁵ depending upon the cell used.

ther rolls are provided with a voter facilit

This instrument operates on the superposition method, by which the oscillations of two high frequency ascillators are brought to superposition in a mixing section. After subsequent amplification, the resulting beats are brought to the screen of a cathode ray tube as the indicator. By this method it is thus possible to observe even fractions of a 1HZ(cycle) beat frequency.

and the variable measuring condenser connected in parallel to it are parts of one of the oscillators. By changing the measuring condenser the sum of both the capacitance values is set so that the same frequency results as by the other oscillator (zero beat).

The double geared, precision scale, with 4500 divisions(scale length about 33 ft (10 meters)) permits determination of the measuring condenser position with the required reading accuracy.

Types of Cell used: The sample holding cells used in connection with the apparatus for dielectric constant measurements are cylindrical condensers. These are of DFL types.

All the sample holding cells are provided with a water jacket for maintaining the constant temperature of the experimental solution.

The surface of the electrodes of DFL type cells are gold plated,

The sample volume of the DFL-2 is 4 c.c. ceramic is used for the insulation of the inner cylinder. Teflon is used for the gaskets. The sample volume of the DFL-1 is 20 c.c. Both the types have a threaded cover, which can be unscrewed. These types have ground glass joint inlets. The DFL-1 type has a bottom outlet drain fittings. The DFL-2 has in addition a teflon stopcock at the bottom. The DFL-2 does not have any drain.

Measuring Range for DFL-1 and DFL-2 Type Cells: A specific cell can be used for the measurement of dielectric constant of a certain range.

The measuring range for DFL-1 and DFL-2 type cells are shown below -

	Sample holding cell			Switch position			
				D ₁	$\tilde{\mathfrak{d}}_{2}$	M ₁	
		DFL-2			2.0-4.7	4.2-6.9	1.0-3.7
		DFL-1	elie la	pingina	1.0-2.4	2.0-3.4	1 - 7 - 7 - 8

By using the cell DFL-1, scale reading is obtained in the range switch at D_1 and D_2 . For the cell DFL-2, the reading is obtained at all the three positions of the switch.

军武功公 南北 称:在 由于是是 名称由 医经验性外部检查 医皮 医病医疗学 了巴斯 西斯西里巴克克西托亚

the setura of the content of all a leader bush the

Calibration of the Cell DFL-1 and DFL-2: The calibration curves with scale divisions against dielectric constant for the cell MFL-1, supplied by the company. The curve is a straight line and the equation of the curve has been found to be:

dielectric constant(d.c.) = 0.0018375x scale reading(at M₁ position)

with the help of this calibration curve, cell DFL-1 and DFL-2 have calibrated and verified by the pure liquid of known dielectric constant. The equation for these calibration curves are.

I compare the employ of up and down to return

a new regularia by forfiner different

For DFL-1

d. c. = 0.0008555186X scale reading (at D₄) + 1.59220.

receive on a fill-is of-

and.

For DFL-2

d. c. = 0.0004783X scale reading (at D₁) + 0.72567.

d.c. = 0.0618677x (call recording (a1 p.) + 3 40968

Measurement

The dipolemeter is plugged into the 220 volts A. C. line. The instrument is turned on. The internal thermostat of the low/er housing should also be turned on. (The panel lamp above the switch indicates that the heaters of the built in thermostat are operating). After about 30 minutes, during which the thermostat and its panel lamp should often turn on and off, the instrument is ready for operation.

melotor. I operative as course outd in source to it by a eprice was not de-

Focus and intensity of the cathode ray tube can be adjusted by the two small knobs on the upper panel below the screen. With the range switch at the Korr. position, the measuring oscillator is brought into resonance with the reference oscillator by means of the correction knob. In the Korr. setting of the range switch the position of the measuring condenser has no influence upon the resonance point. By turning the correction knob, the horizontal line on the cathode ray tube broadens vertically and wave forms become evident. Lissajous figures are formed between the beat frequency and the cathode ray tube scanning frequency 50c/s, which at ± 50c/s change over into the well known circle or ellipse.

At exact resonance the line remains still, moving up and down with very small changes. The number of up and down movements per second indicates the beats per second. By further divergence from resonance, these up and down movements again Lecome Lissajous figures. After the zero beat has been adjusted in Korr. position, the instrument is ready for measuring. The correction knob must not be disturbed ence the adjustment has been made as this will destroy the resonance.

Working Procedure

(i) A range of solution from pure solvent to 0.4% by weight have been prepared by the following procedure.

A flat bottom flask of capacity 100 ml. is cleaned, dried and weighed. A quantity of acetic acid is taken in it by a syringe and again

weighed. The difference between the two weights gives the weight of acetic acid. The pure and dry solvent is then added and weighed again. The difference between 3rd weight and first weight gives the total weight of the solvent and Acon. The concentration in weight percentage is then given by an emore all a builty washed with describing and

suppose a 1 f
$$\frac{1}{w_1}$$
 Ty alter to the the a function of the upper right that it is

where.

evis propredite w. = weight of acid taken. The largery the suppose and the

constant to way = Total weight of solvent and AcOH. - when to order to

w_ = weight percentage of the solution.

This selution is termed as stock solution. Solutions of low concentrations are prepared from this stock solution. Next another flask (already cleaned and dried) is weighed. Then the stock solution is poured into it and weighed again. The difference between these two weights gives the weight of stock solution. Then pure and dry solvent is added and weightd. The difference between this last weight and first weight gives the weight of solvent and stock solution. Hence concentration of dilute solution is computed by,

column was a weight of stock solution.

w4 = total weight of solvent and stock solution.

In this way a series of solutions are prepared for measurement. All these operations excepting weighing is done in a moisture free box containing $P_2 0_5$.

(ii) The measuring cell to be used is kept for several hours at 120°C in the drying chamber after being washed with distilled water and acetone and finally allowed to cool in a dessicator. Now this cell is inserted into the co-axial connection at the upper right side of the instrument. This cell is leveled to avoid spillage, with its central axis perpendicular to the table top. After starting the measurement the cell is not removed from, moved or turned in its connector in order to avoid errors due to slight change in electrical contact. The constant temperature of the cell is maintained by circulating water from constant temperature bath (described latter). The reading of the empty cell is taken by operating the range switch at M4 and D4 or D2. as required by the sample holding cell used and by turning the measuring condenser resonance (Zero beat) is again established. Then the cell is filled with the liquid (or mixture) and reading is taken in the same way. In each case three readings are taken, and the average value is taken as the valid figure. After measurement of a sample is completed, the range switch is turned to Korr. position.

Constant Temperature Bath used in Dielectric Measurement

The thermostatic water bath is equipped with a GallenKamp thermostirrer(Cat.No. TM 860) which controls the temperature of the water bath at 30° C $\pm .01^{\circ}$ C. The control of temperature at $30 \pm .01^{\circ}$ C is achieved by maintaining the main voltage to 180 volts by a variac transformer and connecting an extra resistance in series with the thermostat-heater. An extra heater is also used to cut the initial heating period. Although, it has been written in the catalogue that the temperature monitors at 30° C $\pm .01^{\circ}$ C, we have observed that it maintains the temperature at 30° C $\pm .002-3^{\circ}$ C.

ante la tre albertal lebe anather finely being measure of fractional expensional provincial constant to expension the expension developed being particular to the site of an elemental finely of the site of the s

Evilledian of Delresia

Constally, at first the solvents are arised by high and them the theory of sector in the traces of sector in the content of sector in sector.

First at a transment of the Later the release to distill and to treated with Market and Later the Sector to Dispute the State Contract to the Sector to Sector the Sector to Sec

of Constant bolding point in the

Reagents and Their Purification

Acetic Acid

E. Merck acetic acid is dried by shaking with P₂O₅ in cold and allowing to stand for several hours; then the liquid is drained off and distilled. The distilled liquid is frozen in an ice-bath. When half portion of acid is transformed into fine fibrous crystal, the remaining acid is transferred into another flask. This procedure of fractional crystallization is repeated several time and is finally kept in an airtight flask placed in a dessicator containing P₂O₅. A stock can be used for about one and half month in dry season and 15 days in rainy season. Each time acetic acid is needed for a set of experiment it is frozen, the liquid part is separated and the crystals are taken.

fore property of the weak was to place the forest of

Purification of Solvents

Generally, at first the solvents are dried by P205, and then Na-threads are dropped into the liquid to remove the traces of water in the solvent. But this method does not remove sufficiently the traces of water. We adopt the following method.

After the treatment with P₂0₅, the solvent is distilled and is treated with Na-threads. After several hours the liquid is distilled and treated with a little freshly purified acetic acid (.06 gms acid is added in 100 gms of solvent) followed by Na-threads. Na-threads are

added till the reaction is complete. Then it is finally distilled and a centre cut of constant boiling point is taken.

In support of this procedure we have to state the following:

In our earlier work in acetic acid - CCl₄ system it was found that even after drying with Na-threads, the dielectric constant progressively decreased from that of pure solvent upto a point, from which again, dielectric constant increased with increasing concentration of acetic acid. It was thought that addition of acetic acid to CCl₄, dielectric constant should, on no account, be less than that of solvent. To account for this strange behaviour we concluded that there are traces of N₂O in CCl₄ which form addition compound with acetic acid, which most probably, has the following symmetrical structure with very little dipole,

deconoted and apptition. Then it is tranted with acrese and follower

Thus, it was concluded that the last traces of H₂O remain inside interstices of the solvent from which it is difficult to be removed by interaction with sodium. If pure dry acetic acid in trace quantity (.6 gm in 1000 gm of solvent) is added, will react with H₂O to form the addition compound; which is, as a whole a strong acid. This compound will therefore react strongly at once it come in contact with

E TOTA

the love the ten and in the Continue to the children

is public of front to the every money property and this into the following the con-

who would be added to the first dension of species and is the part of the control of the first dension for the part of the first dension for the part of the first dension of the part of the first dension of the first density densit

there we do be sound in tile prove this there are three divinest charters of direction. There is indiction would not be be preserved in the sense to be a contract of the sense the contract of the sense to be a contract of the sense of the sense to be a contract of the sense of th

RESULTS

System: Acetic Acid in Carbontetrachloride.

Results of Specific Volume Measurement and Obtaining Refined Values

The specific volume of solution of acetic acid in CCl₄ was measured by magnetic float densimeter in the range between pure solvent and .002 weight fraction following the procedure described in the previous chapter. The data obtained by such experiment is given in the Appendix-A.1, along with the linear transform of specific volume (referred to as $\triangle v$) by a reference line, $v_{cal} = .634985_{+} .30303 \text{ X}$ w, where w is the weight fraction of acid. The plot of the $\triangle v$ versus w is shown in the Fig. 4.1 curve(a).

changes of direction. These inflections could not be observed without the help of this linear transform plotting. Here we see that the mean deviation no where is more than 2×10^{-6} c.c/gm. There is a curvature from 0.0 to about .0012 weight fraction showing a gradual increase of specific volume and then again there is a sharp decrease of specific volume upto .0015 weight fraction and from that point there is again a sharp increase of specific volume upto the highest measured point. A

sample calculation from the line of the curve shows that the standard deviation is $\pm 1 \times 10^{-6}$ c.c/gm. Thus, we can say that the precision of the measurement are highly satisfactory.

d from the every following to

Results of the Dielectric Measurement

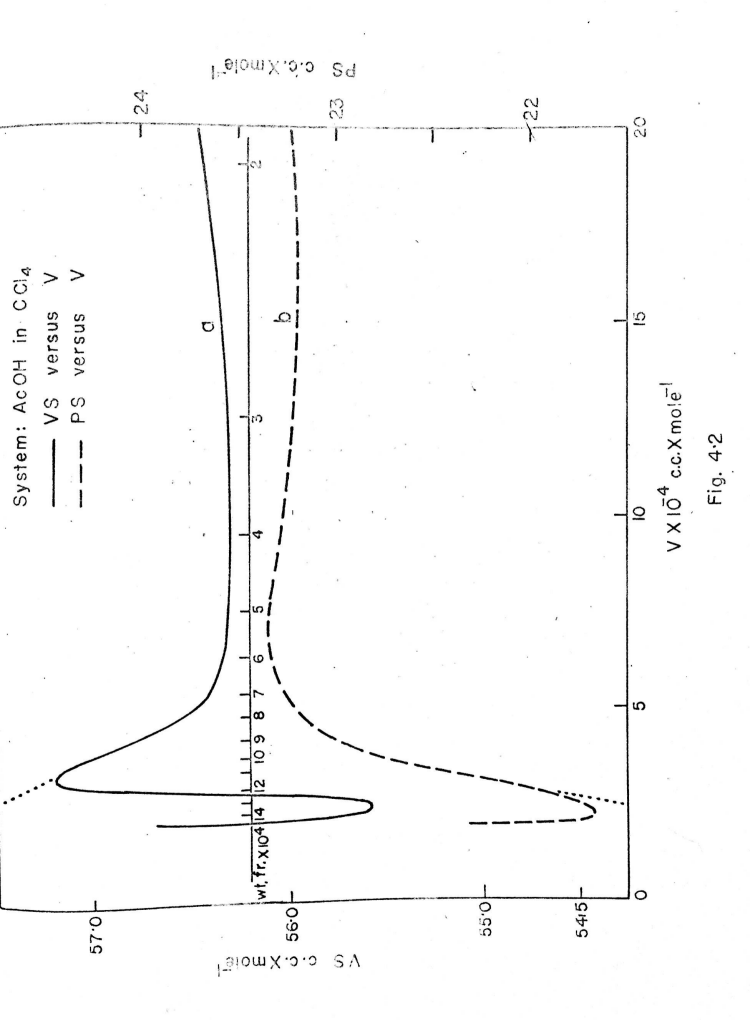
The data of dielectric measurement of solutions of acetic acid in CCl₄ between 0.0 to .0027 weight fraction is recorded in the appendix-A.1, along with the linear transform, \triangle from the straight line of equation, $C_{cal} = 2.251207 + 1.21415$ XW, whose plot is shown in Fig. 4.1 curve(b). The transform plot shows that the mean deviation of the experimental data from the drawn line is 2×10^{-5} c.c/gm, the percentage deviation of course is of the same order of as that of specific volume. This means that dielectric measurement also has a high precision. The curve shows a number of inflections keeping in line with the inflections shown in the specific volume - wt. fraction curve.

Calculation of S and V and the Plot of VS against V

Using the following relations, V and VS is calculated:

$$V = \frac{V}{W} X M$$
; $S = V_S X \left(\frac{1 \leftrightarrow W}{W} \right) X M and VS = V-S$.

Reading out carefully the value of \triangle v from linear transform curve and calculating back to trefined "value" of specific volume as described a



"refined specific volume" already calculated and tobulated at these points previously.

The P, S and PS are calculated using the following relations:

the curve herbeen 5 to 10 note (for after to 1000 Mt. francistic). In

of to red of these of polarisers and tests the forestign of eyelic
$$\mathbf{P} = \frac{\mathbf{b}}{\mathbf{v}} \times \mathbf{M}$$
; there is value of relax polarisation of dimer, i. and i.e.

the
$$x \le x = \frac{1-y}{y}$$
); or how is he obtained by extrapolation of

and PS = P-S;

The plot of PS versus V is shown in Fig. 4.2 curve(b).

Ps-V

ad from thomas someon by remain

A Comparative study of VS and PS Curves

In Fig. 4.2 the curve(a) and (b) show the variation of VS and PS versus V. The straight line at the middle of the graph, with marking from 2 to 14 is given to indicate the weight fraction composition of the mixture at which such values are obtained, to compare with other systems, interpret and compare these curves more intelligently and to speculate about the regions at which different kinds of interaction takes place. Let us start our inspection from the right side of the curves whose V is very high. It is seen that VS and PS both attain nearly a constant value between mark 2 to 6 (i.e. wt. fraction .0002 to .0006). Then there is a rise of VS and on the other hand, a fall of PS upto about .0012 wt. fraction. From this point onward, there is a fall of

VS and rise of PS. Thus, it appears that from .0002 to .0006 wt. fraction PS and VS change very little which indicate that only monomer predominate at this region; and, after .0006 wt. fraction the increase of VS and decrease of polarization indicate the formation of cyclic dimer. To obtain the value of molar polarization of dimer, P₂ and V₂, the molar volume of this dimer has to be obtained by extrapolation of the curve between 6 to 12 mark (i.e. .0006 to .0012 wt. fraction). By these method, we obtain,

$$V_2 = 2x 59.08 \text{ c.c/mole}$$

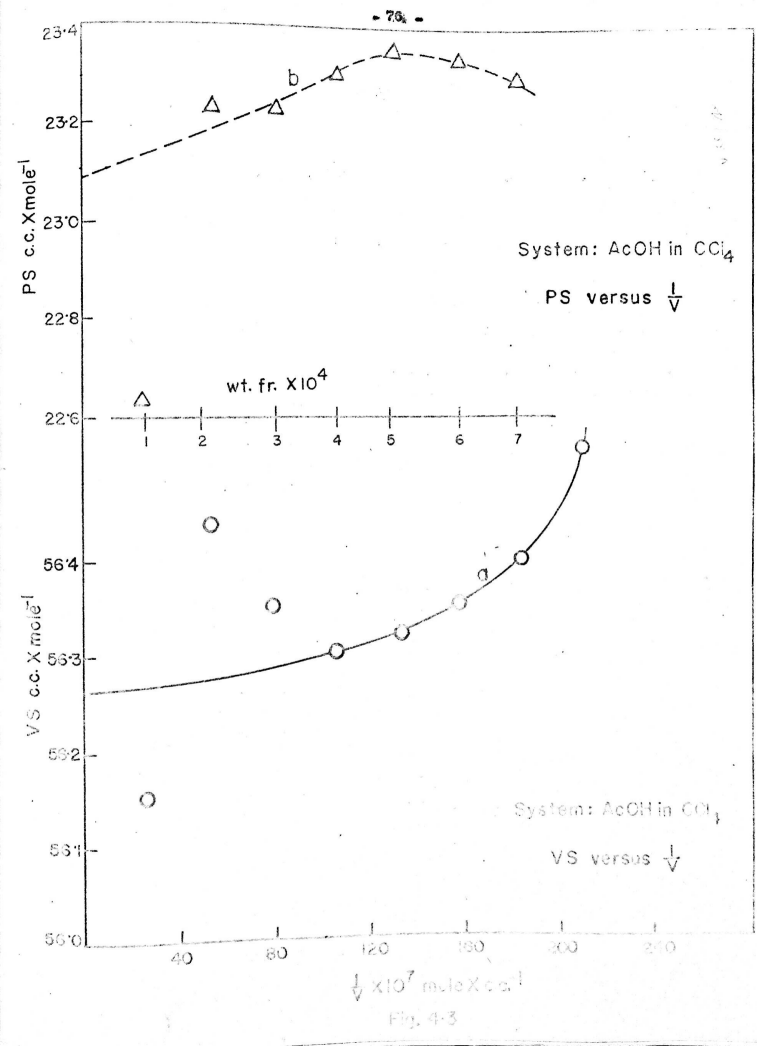
The value of V_1 and P_1 can be obtained from these curves by rough extrapolation. But we have a better method; according to equation (2.17), the extrapolation of the plot (Fig. 4.3) PS (or VS) versus $\frac{1}{V}$, should give P_1 or V_1 as the case may be . By that method, we obtain,

$$P_1 = 23.10 \text{ c.c/mole}$$

$$V_1 = 56.26$$
 c.c/mole

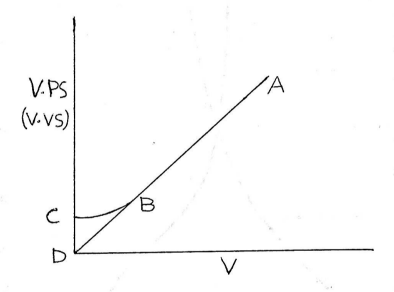
D for molar volume = 5.64 c.c/mole

D for molar pelarization = -7.82 c.c/mole.

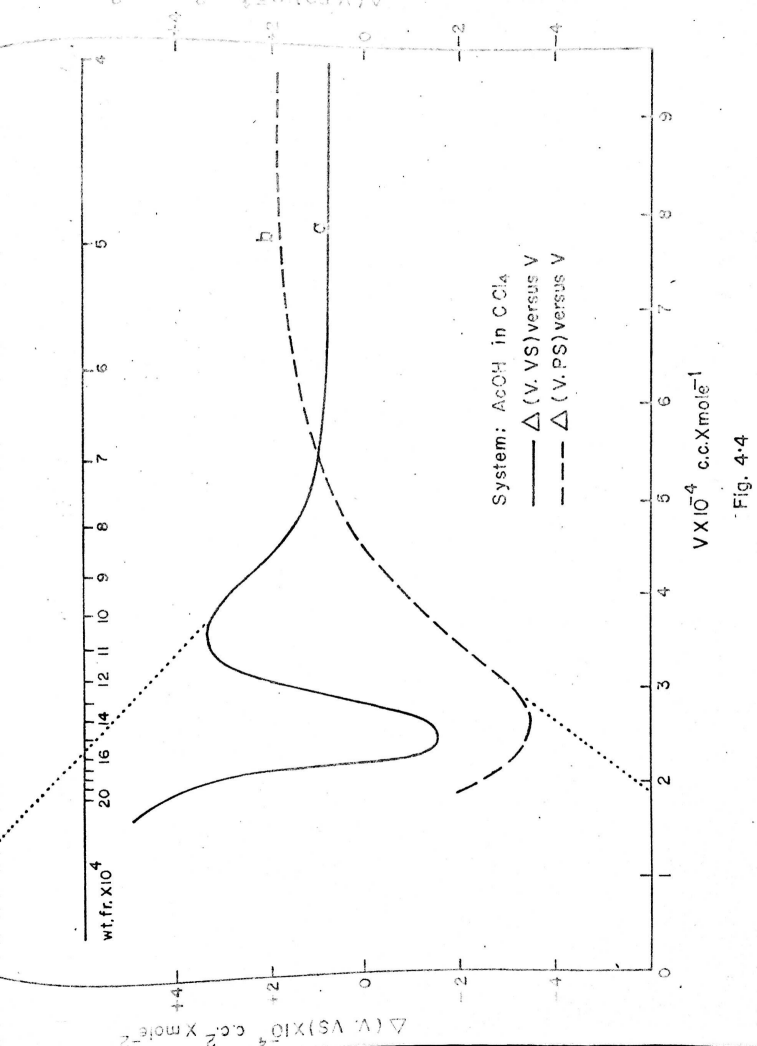


The Plot of V. PS(or V. VS) Versus V and Calculation of KD and K

V. PS (or V.VS) versus V plot give a curve is not satisfactory for convenient estimation of KD. The following figure shows an exaggeration of such a plot.



The part of the curve AB shows that at this region only monomer exist althoughout . As we have already shown, if only monomer existed althoughout then the line should have been ABD. Here CD is the intersection of ABC curve. In ordinary plotting actually it falls very near to zero. Therefore, V. PS or V. VS versus V have been transformed by the straight line ABD, whose slope is equal to Y_1 (or V_1) and intersection is zero. Thus, \triangle (V.PS) and \triangle (V.VS) have been calculated and given in Appendix- A.1. The plots are shown in Fig. 4.4.



It is to be noted that from high dilution to .0006 wt. fraction in both cases shows a flat portion. This part is obviously concerned with existence of monomer and weak interaction of this monomer with solvent. In V.VS then rise upto .0012 wt. fraction and in case of polarization V.PS fall down. This region indicate the equilibrium reaction of dimer formation. The linear extrapolation of these curves leads the following values of KD.

For molar volume, KD = 100600 c.c²/mole²
For molar polarization, KD = - 105000 c.c²/mole²

But we have D for molar volume = 5.64 c.c/mole
and D for molar polarization = -7.82 c.c/mole.

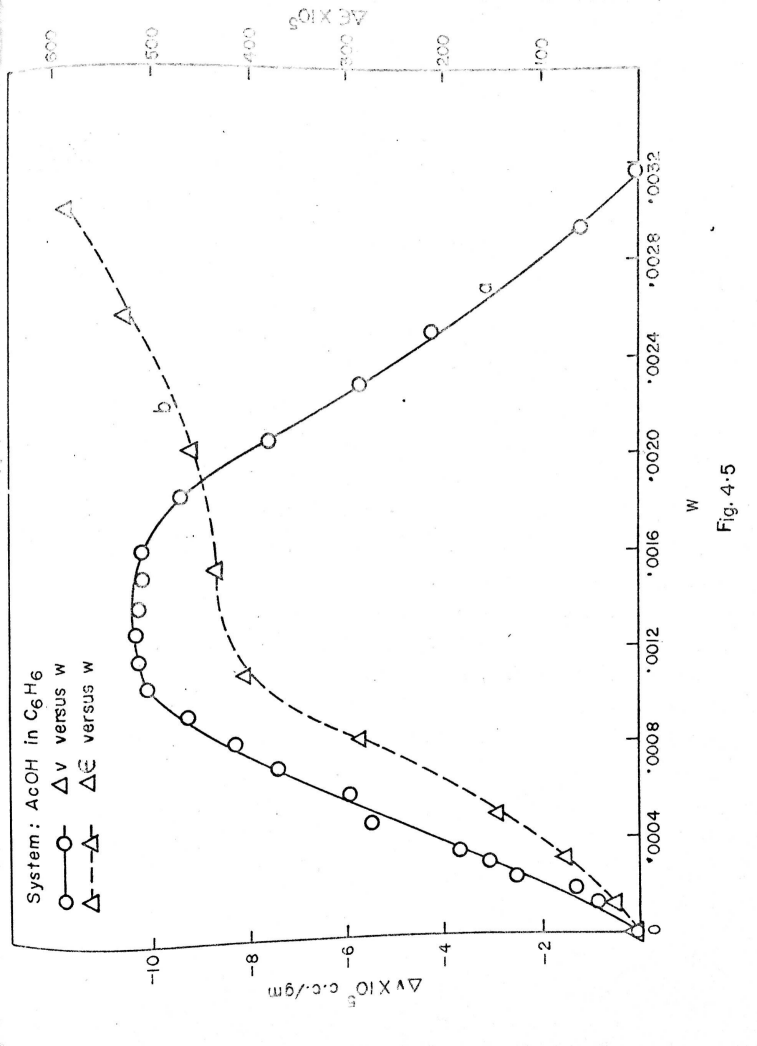
Therefore, K are 17.84 1/mole and 13.43 1/mole for specific volume and polarization respectively. The results are tabulated in the Table 4.1.

. C. I John Mid the pittich on active of weather at the charm as Fig. 4.5

gurray (c) end or transport TABLE 4.1 Price 4.5 carres (a).

Molar property	Monomer (c.c/mole	Dimer (c.c/mole	$(c.c^2/m_0le^2)$	K (1/mole)
Volume	56.26	59.08X2	100600	17.84
Polarization	23.10	19. 19X2	-105000	13.43 Diverse

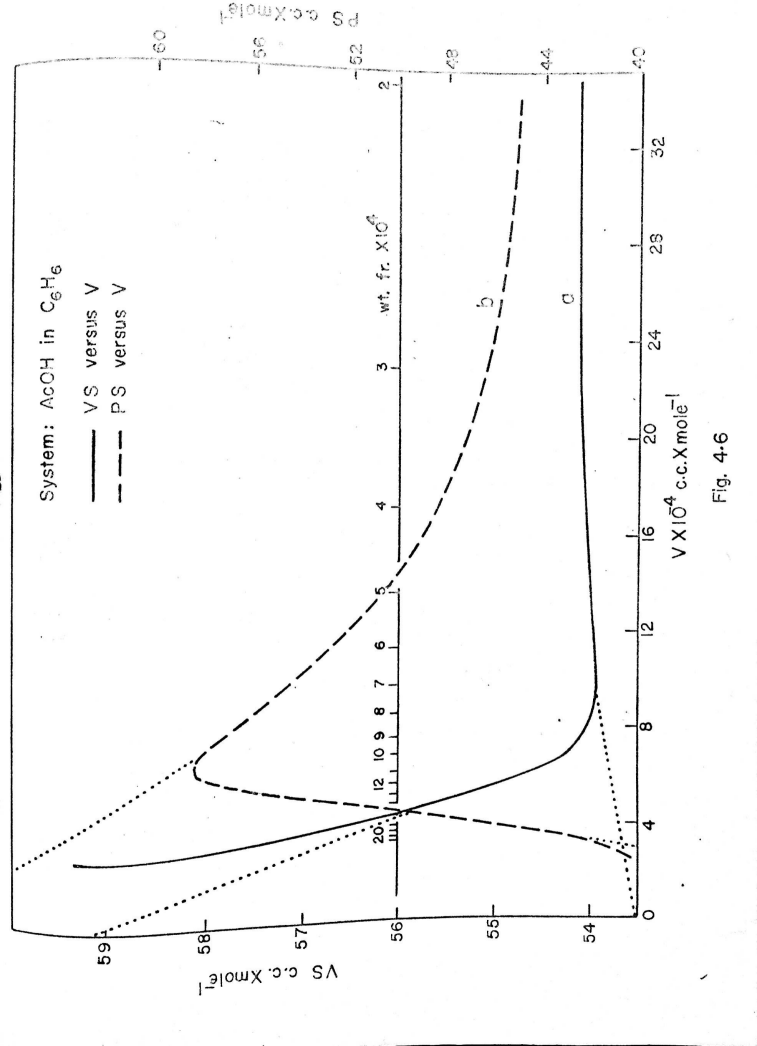
things walked of specific volume and factories constant (as described



in the case of CCl₄) to calculate S, P, V, Vs, Ps etc. at equally spaced chosen points of composition.

Plot of VS and PS Against V

Vs (and PS) versus V are pletted and shown in Fig. 4.6 curve(a) and (b). The curves in the figure show the variation of Vs and PS versus V. (The straight line at the middle of the graph marking from 2 to 20 has been given to indicate the wt. fractions of acid in the solution). From .0002 to .0008 wt. fraction there is an increase of PS and decrease of VS. The increase of PS shows that a species is continuously forming with the increase of concentration has higher polarization than the monomer. supposed to exist abundantly in the dilute solution. In VS we observed that there is a weak but definite tendency of Vs falling down indicating that the species is formed with a little contraction of volume on the combination of two monomers. So, we are led to consider the formation of open chain dimer in this concentration range. Then after the solution has attained the concentration .0009 and goes on increasing, we find that PS decrease and VS increase. This section strongly indicate the formation of cyclic dimer, as we know, the formation of which is always attended the increase of volume and decrease of polarization. Thus, we have calculated the important parameters for



two distinct interaction, in succession:

(a) $A + A \rightleftharpoons A - A$ (open chain dimer)

and

(b) A + A = (A) (closed chain dimer)

The justification of such a supposition will be perfectly clear when we examine \triangle (V.PS) and \triangle (V.VS) variation with the variation of volume.

The molar volume of dimer, open chain and cyclic have been obtained by extrapolation of these curves according to the procedure already laid down. The values obtained by such operations are shown below:

 V_2 (open) = 2x52.53 c.c/mole, P_2 (open) = 2x72.57 c.c/mole

 V_2 (cyclic) = 2x59.13 c.c/mole, P_2 (cyclic) = 2x16.17 c.c/mole. The determinations of V_1 and P_1 was carried out by VS (and PS) versus 1/V plot and extrapolation to 1/V = 0(not shown in figure). From these operations we obtained the value V_1 = 54.18 c.c/mole and P_1 = 40.20 c.c/mole. Thus, the values of P_1 , V_1 , P_2 , V_2 and D obtained are shown in the Table 4.2.

The filter partition of dilette reviews could be rear unit a made

THE STANDARD DIESE OF THE THE TALLS OF TOTAL OF A THE FOREST LINE OF BOOK WHEN

A Law F.

1 - 10 200.

Table 4.2

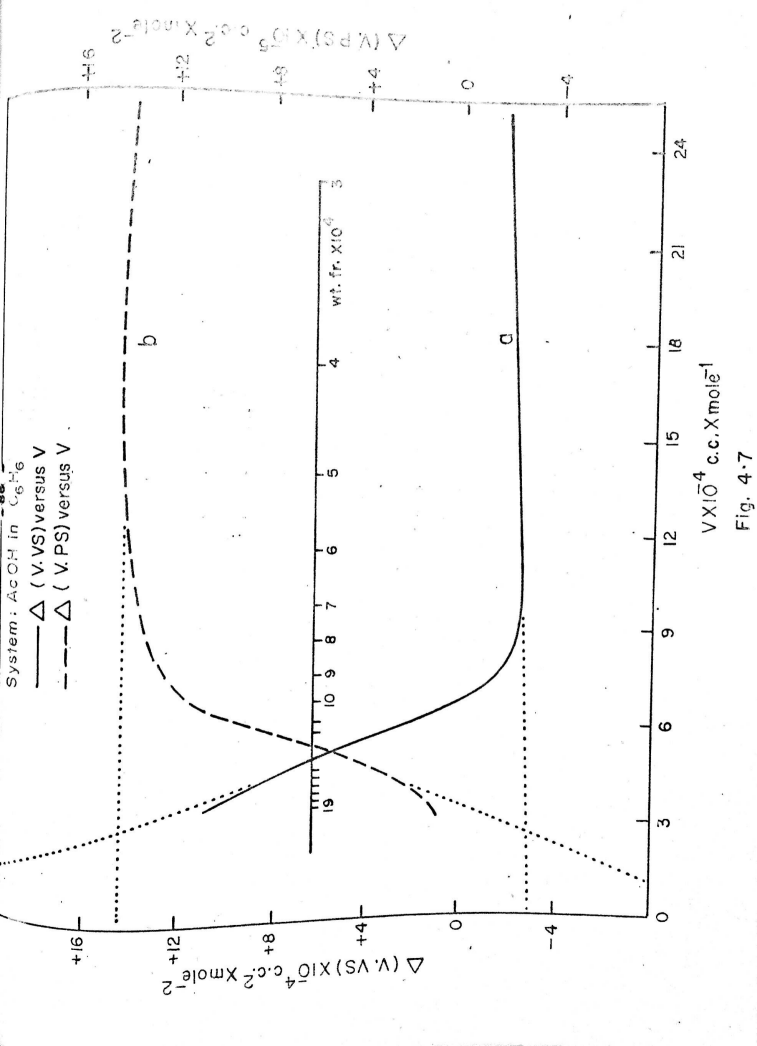
V₁ = 54.18 c.c/mole;

 $P_1 = 40.20 \text{ c.c/mole.}$

r properties	Open chain (c.c./mole)	Closed chain	ton spirit dec
v ₂	2x53.53	2 x 59. 13	W - 1 j
D (volume)	- 1.3	9.9	
P ₂	2x69.60	2x16.17	- 4
D (Polariza- tion)	58.80	- 48.06	

The \triangle (V. VS) and \triangle (V.PS) versus V are shown in Fig. 4.7. In both the curves there are two clear sections: (a) from most dilute to .0008 wt. fraction a straight line nearly, but not exactly, parallel to the V-axis, and (b) another linear section from .0010 to .0017 wt. fraction.

The first portion at dilute region could be very well a weak interaction just, as in the case of CCl4 or the formation of open chain dimer.



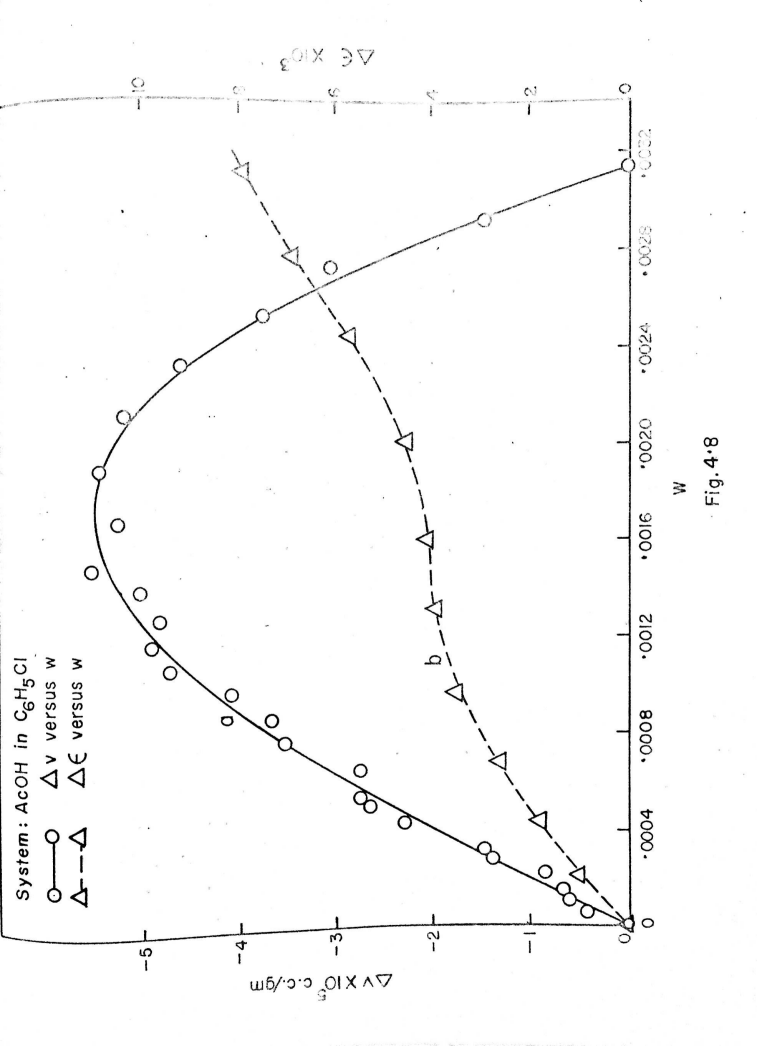
Here the indication by comparison with VS (and PS) versus V plot we are convinced of an interaction and the second portion of interaction line of closed chain dimer. The extrapolation of these relevant line leads to the following results:

Table 4.3

Open chain			Closed chain	
	Vol.	Pol.	Vol.	Pol.
KD(c.c ² /mole ²) - 30000	1440000	260000	- 1120000
K (1/mole)	23.08	24.49	26.26	23.30

System: Acetic Acid in C6H5Cl

The specific volume and dielectric constant of the system has been measured from pure solvent upto .0025 wt. fraction. The data is given in the Appendix A.3, along with the other parameters (S, P, V, \triangle C, \triangle V, VS, PS etc.). The linear transform equation for specific volume is $V_{\text{cal.}} = .912825 + 0.24650 \times v$ and for dielectric constant is \triangle C = Cal = 5.610160. The plot of \triangle V and \triangle C are shown in the Fig. 4.8 curve (a) and (b). The plots show clearly the spread of error of measurements



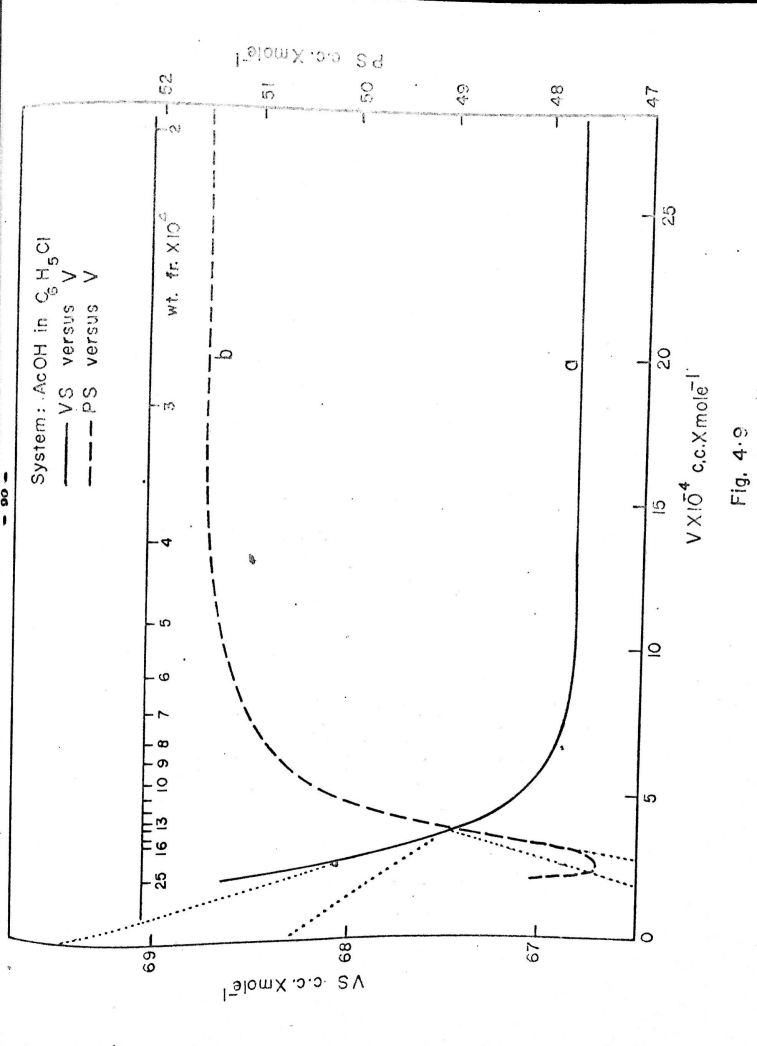
which are as usual within reasonable limits. The variations of the properties due to interaction are also evident. The mean lines are drawn very carefully from which refined and acceptable values have been specified and chosen for calculations of other important parameters.

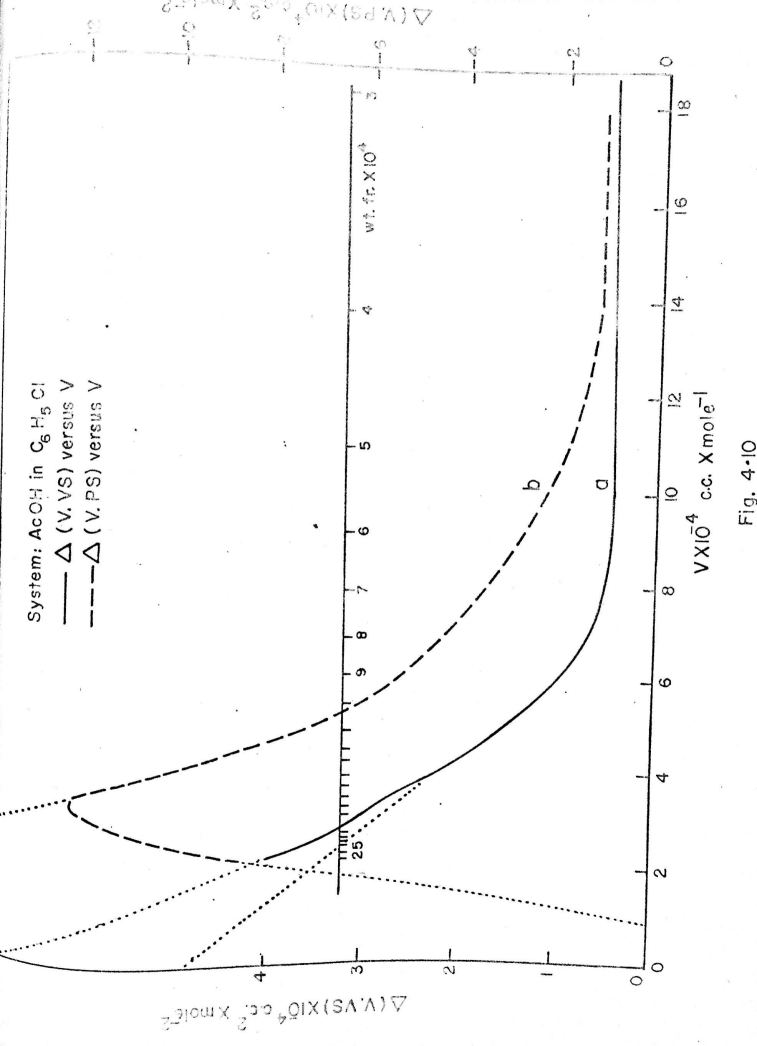
with the refined values at specified points the V, P and S are calculated from which we obtained PS and VS versus V. The plot of VS (and PS) versus V are shown by two curves in the Fig. 4.9 curve (a) and (b).

There are two distinct sections:

(a) from extreme dilution to 0.0005 wt. fraction and (b) from 0.0005 to 0.0019 wt. fraction, revealed in the PS versus V plot. In VS versus V plot the second section can not be clearly marked out and this is because of the fact that there is a continuous increase of volume with the increase of acetic acid and this must be due to formation of such complexes in succession which have greater molar volume than its predecessor. This feature is distinctly different from those two systems we have already studied, which will be discussed later.

In \triangle (V.VS) and \triangle (V.PS) versus V plot there are two distinct sections Fig. 4.10: (i) from dilute to 0.0015 wt.fraction, at which range both polarization and volume of species formed, is attended with increase of volume and polarization; which shows something unusual; (ii) in the next portion, from 0.0015 wt. fraction upward, volume increases and polarization decreases. We will presently find that





this section does not conform to any monomer-dimer equilibrium. All these will be explained at a latter part of this work.

Following the usual procedure the parameters have been estimated which is given in the following Table. V_1 and P_1 were found out by extrapolation of 1/V = 0 from VS or PS versus 1/V plot (not shown in Figure).

Table 4.4

Molar proper- ties	Monomer (c.c/ mole)	open chain	(c.c/	chain	1.	chain	KD cyclic (c.c ² / mole ²)	K open chain (1/mole)	K cyclic (1/mol
Volume	66.81	2x68.3	2x69.47	2.98		48500 NS	68000	16.28	12.78
Polari- zation	51.56	2x45.46	2x42.88	-12.2	-17.36	-230000	92000	18.85	-5.30
zarron		of thes	eri turk	13 B S F F	er post	ef ita ya	inga siki	i in the t	57. 27-46

Here again we see an unexpected thing, the value of V is higher than expected.

整个人的主义的 经证据 自然主义的 医结束心脏 (美国的农民类似 是 MESTIN 知识的 ANDER 1777 (1974) (1974) (1974) (1974) (1974)

There is another section which invites our attention, is from 0.0016 to 0.0024 wt. fraction. Here the PS shows down ward movement with clear linear section and its counterpart in VS also shows upward movement

of clear linear section. Does it indicate another dimer interaction? The extrapolation of PS gave KD is $+92 \times 10^3 \text{ c.c}^2/\text{mole}^2$ and D for polarization is -17.36 c.c/mole which gives K = -5.30 l/mole. As K can not be a negative value, this clearly indicate that this can not be indicative of a dimer interaction. The speciality of association interaction of acetic acid in this solvent has been discussed in the section dealing with interpretation of results.

Solvent Behaviour of high Beiling Hydrocarbon and Cyclohexane to Acetic Acid.

High boiling hydrocarbon (HBH) and cyclohexane are universally accepted as "inert" solvents, at least more inert than CCl₄. Inspite of being so, we have trouble ahead. We have very strong indications of solute solvent interaction of a very unusual kind. This came to our notice in this laboratory a few years back. It was found that in very dilute solution of both cyclohexane and HBH acetic acid molecule (most probably monomer) looses a great part of its volume which is very surprising and quite inesplicable. A work was undertaken to investigate this aspect more thoroughly. The result of such investigation was very interesting. It was found that the apparent molar volume of acetic acid

^{*}A project "Equilibrium Interaction of Solute species entering into the holes of a solvent body" was undertaken by Mr. Zillur Rahim, Associate Professor of Chemistry, in the year 1977, in which the present author was a research Fellow. The Final report was submitted in the year 1980.

molecule, VS attained 40 c.c at very high dilution in HBH and 10 c.c in cyclohexane, which gradually increase with increasing concentration. The variation of the apparent molar volume(VS) of acetic acid in these solvents was explained assuming holes in the solvent. It was proposed that the solvents have a number of "holes" within the solvent body in which the species enter and looses its own volume; but this entry and exit follows a dynamic process which can be expressed by a dynamic equilibrium in the following:

free Solute hole solute in hole

where, where
$$K = \begin{bmatrix} A \\ A \end{bmatrix}$$
; in ferror a second in the s

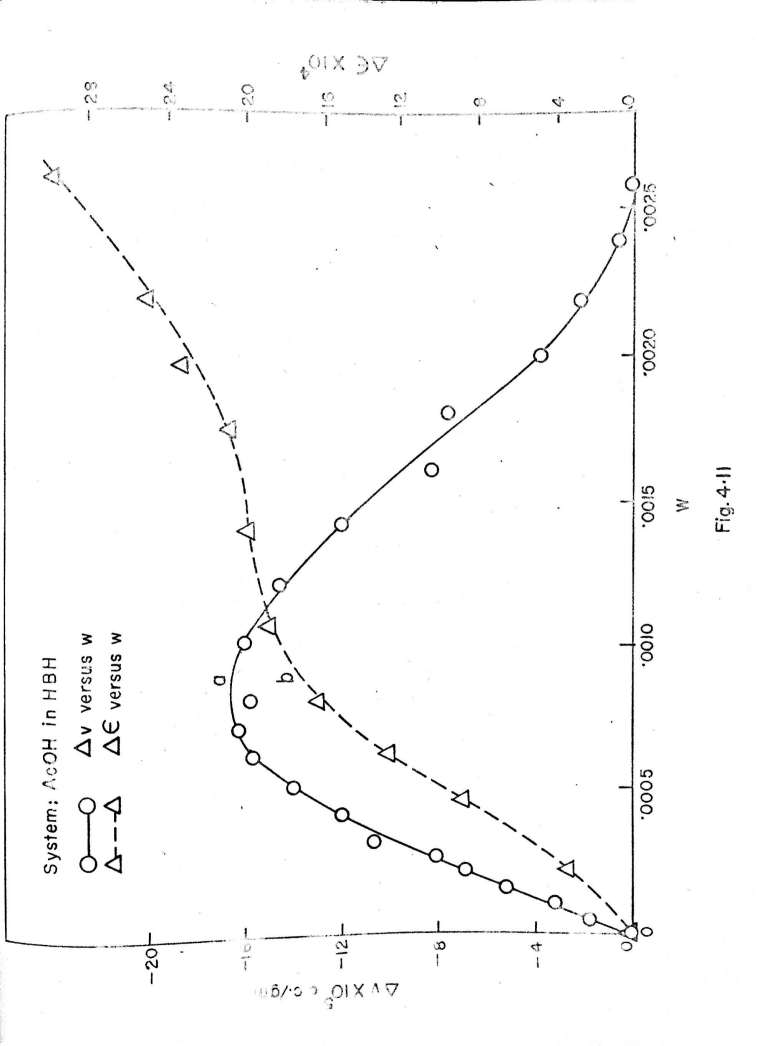
which [A], [A] and [O] indicate the concentrations of A, A and the holes.

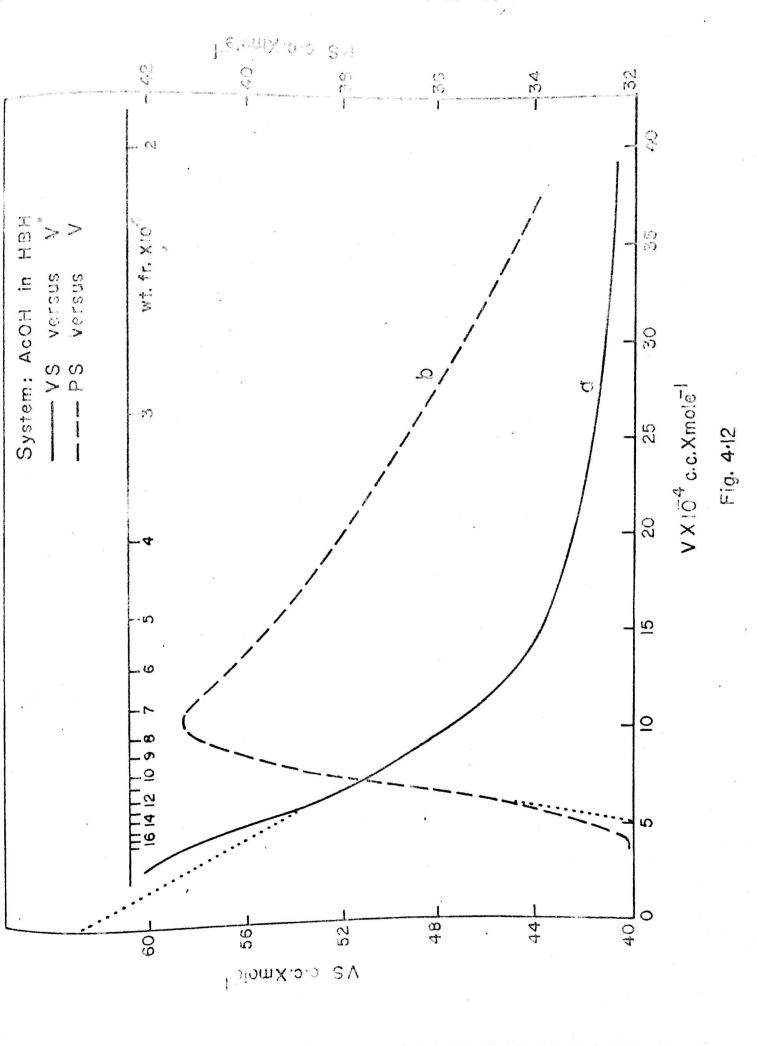
It must be emphasized that the law of mass action is essentially statistical in nature. Hence, the above formulation is not altogether illogical. The important point is whether this formulation can stand the test. The result of such investigation is given in the Appendix-B which amply show that in the dilute part of acetic acid in HBH and cyclohexane behave very differently whose dominant characteristic is an interaction of the type termed as "solute - solvent hole" interaction.

This interaction is followed in case of HBH acetic acid systems upto 0.0006 wt. fraction and in cyclohexane upto 0.0008 wt. fraction. Thus, this part of the solution has been relegated to "solute hole" interaction and when we present the results of our present investigation for determination of monomer-dimer equilibrium we will find its pronounced effects.

System: Acetic Acid in High Boiling Hydrocarbon (150°C - 200°C)

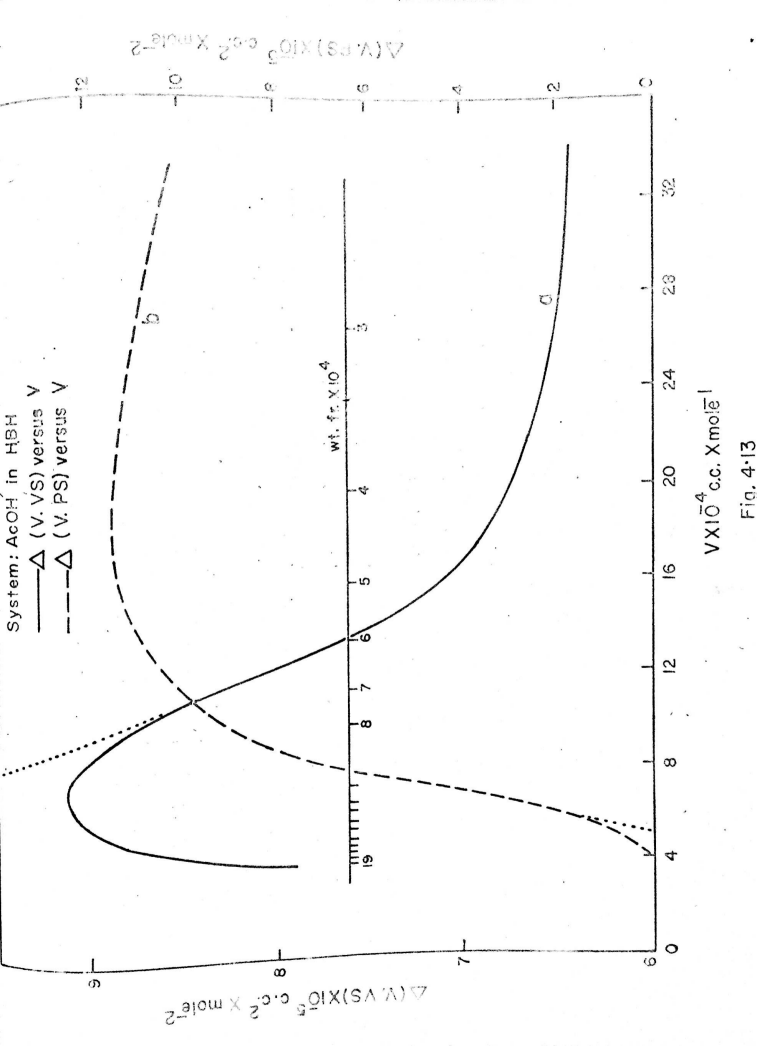
The results of specific volume and dielectric constant along with the calculations of other parameters (V, P, S, VS, Ps etc.) are given in the Appendix A.4. The plot of linear transform of specific volume and dielectric constant (by the reference lines quoted in the Appendix A.4) given in the Fig. 4.11. The errors in the measurement are of the same dimension as in the other cases. From the carefully drawn curvos the relevant values have been read out at properly spaced specific points, from which P, S, VS, PS and other parameters have been calculated. The PS and VS versus V plots are given in the Fig. 4.12. Both the curve show some abnormal features in the most dilute part of the system. The VS, which shows the p apparent molar volume of the species, at very dilute solution is about 40 c.c. which ultimately rise to a value of 60 c.c and PS showing the apparent molar polarization of the species at first rise from the low value 30 c.c to about 58 c.c at





0.0007 wt. fraction and then begins to fall continually to a very low value. It has been assumed on the ground of our previous investigation on "solute - hole interaction" that the section of the curve from most dilute to 0.0008 wt. fraction must be mostly relegated to solvent solute interaction and it is also assumed that after this section the solute in monomeric form are comparitively free to participate in the monomer dimer equilibria whose structure is suspected to be cyclic. V_2 and P_2 for dimer have been obtained by plotting VS and PS versus V on a larger graph. The values obtained are $P_2 = 2x19.99$ c.c and $V_2 = 2x63.40$ c.c. In case of polarization P_1 was obtained by extrapolation to 1/V = 0 in PS versus 1/V plot. But in case of V_1 , although we can get by such a proceeding, the value, 39.0 c.c which seem to be of dubious nature; because V_1 under these circumstances, is an ill defined quantity, varying continuously with the change of concentration under the influence of "solute-hole interaction".

△ (V.VS) and △ (V.PS) against V are plotted and shown in the Fig. 4.13. The effect of "solute hole interaction" is evident at dilute solution upto .0007 wt. fraction. From .0007 to about.0014 wt. fraction we find, in both the curve linear sections clearly indicating the monomer dimer equilibrium. In case of molar volume we have a slight difficulty in ascertaining the slope within this sections; however, the points at .0005 to .0009 wt. fraction have been extended to get the intersection



for KD value. In case of polarization the fall of PS indicate the formation of cyclic dimer. The finding of the intersection for KD is not a problem. We can find V_1 or K for volume property by using the values obtained in polarization study.

Now if we assume that K = 46.68 l/mole (polarization), then deviding KD ($1018627.5 \text{ c.c}^2/\text{mole}^2$ for volume) by 46.68 l/mole (i.e., 46680 c.c/mole), we obtain D = 21.82 c.c/mole. But $2V_1 = V_2 = 21.82 \text{ c.c/mole}$ and therefore, we have finally, $V_1 = 52.49 \text{ c.c/mole}$. This is an excellent finding; because in case of CCl₄ and benzene we get $V_1 = 56.26 \text{ c.c/mole}$ and 54.18 c.c/mole respectively. Thus, we can claim that KD obtained by both the studies agree well with each other as there is a clear linear section between .0008 and .0014 wt. fraction. The result of such investigation is given below:

tard value to highest paint (1860 of fraction) to \$6 c.s. This directly

	Monomer (c.c/mole)	Dimer (c.c/wole)	(c.c ² /mole ²	K (1/mole)
Volume	2 - V pr. 4 (2x63.4	10 18627.4	i — A pivolur v arializa
Polarization	32.2	2X19.99	1140000	46,68
	alam adilys. 1	7.3 6.0 444	tion that pass	\$ 2% alowing rices

as of select of the wheeler, assitt

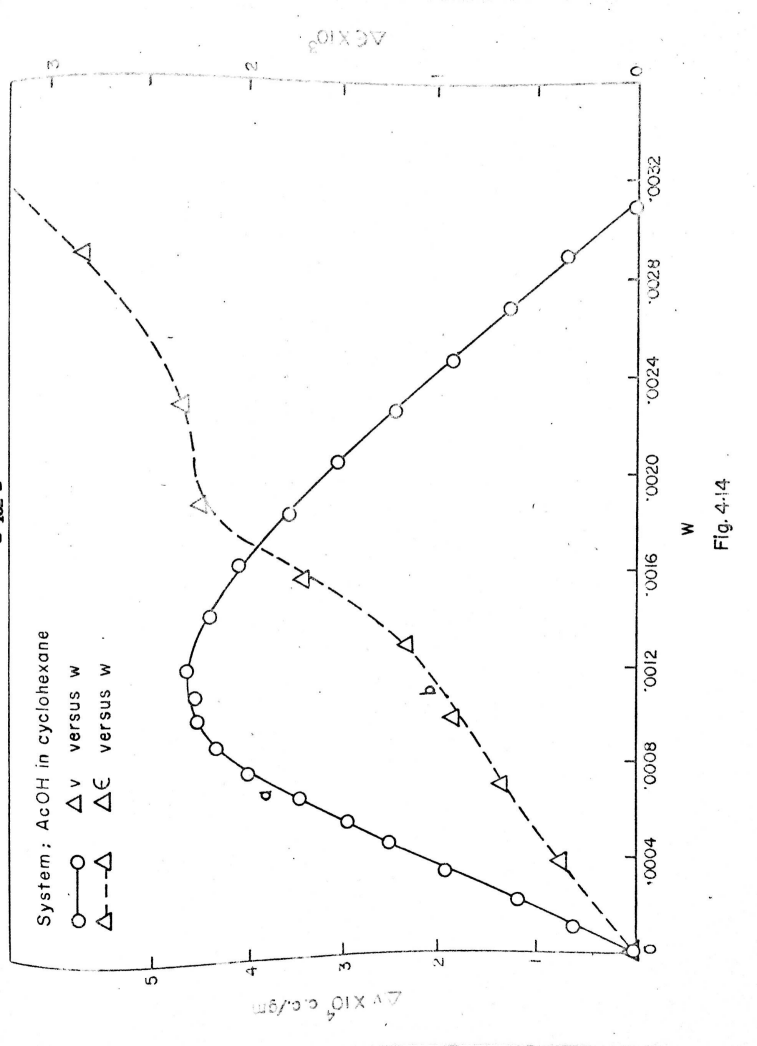
System: Acetic Acid in Cyclohexane

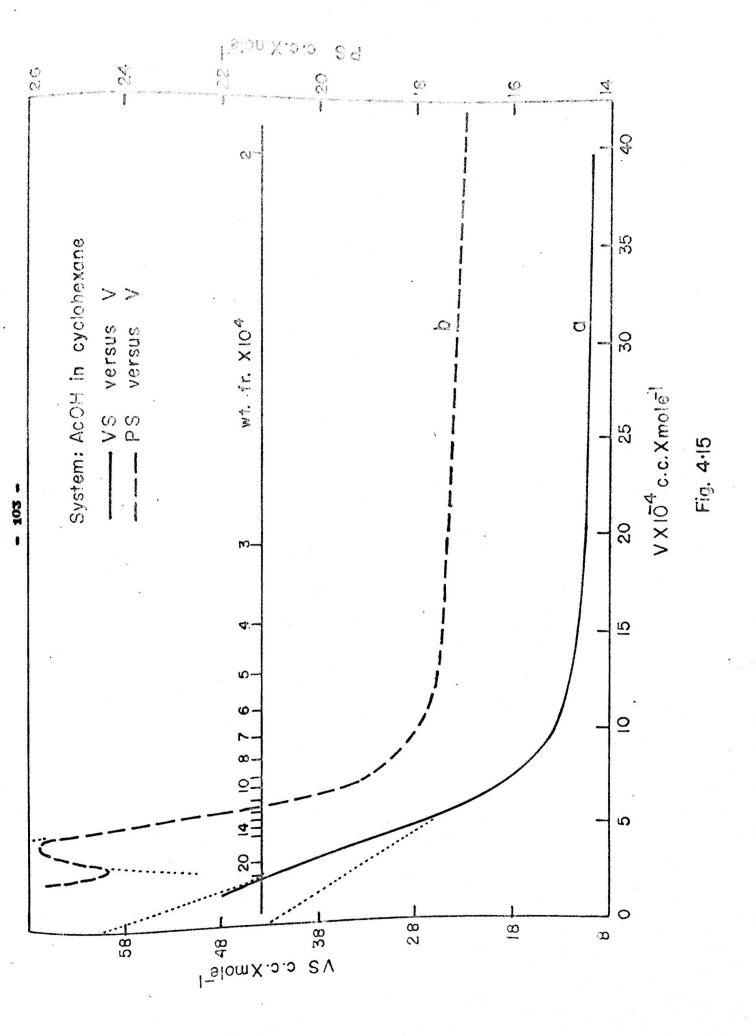
The results of specific volume and dielectric constant measurement and other subsequent calculations have been given in the Appendix-A.5. The plot of linear transformation of specific volume and dielectric constant have been given in the Fig. 4.14. The results are excellent as regards the mean deviation and standard deviation. From the carefully drawn lines, values of specific volume and dielectric constant have been read out at the properly spaced specific points from which Vs. ps. p.s etc. have been calculated Ps and Vs both have been plotted against V and are given in the Fig. 4.15.

We observe that VS, apparent molar volume keeps on a very low value, 8-9 c.c upto .0004 wt. fraction and then upto .0007 wt. fraction it reaches to 13 c.c, from that point there is a continuous rise of this value to highest point (.0030 wt. fraction) to 44 c.c. This clearly indicates that there is a tremedious loss of volume of the species, acetic acid all throughout this range. This shows that a very unusually strong solute-solvent interaction of the type 'solute hole interaction' is taking place.

In PS versus V plot (curve (b)) also, we find a similar variation.

Upto .0007 wt. fraction the solute in the solvent have a very low app
ment molar polarization, 17.5 c.c and from that point it slowly rises





upto .0010 wt. fraction and then there is a swift rise to 26 c.c. at .0018 wt. fraction. From this point upto .0024 wt. fraction there is a fall of this value and then again a rise. These indicate a number of interaction with the variation of concentration. These will be indicated very clearly in the next plots, \triangle (V. VS) and \triangle (V. PS) against V in Fig. 4.16 curve (a) and (b).

In these curves (a & b) in Fig. 4.16 we identify two linear sections in each of the curves:

- (i) From .0008 to .0014 wt. fraction in molar volume curve and from .0011 to .0018 wt. fraction in molar polarization curve;
- (ii) From .0018 to .0024 wt. fraction in both molar polarization and volume curves. The open chain dimer is identified by increase of molar volume and decrease of molar polarization (the lower part of the curve, i.e., .0008 wt. fraction down to more dilute solution the curve indicate the solvent-solute interaction as has already been indicated as 'solute hole interaction'.). The linear portions as indicated in the two curves have been extrapolated to obtain the intercept for the value of KD for the molar volume and molar polarization.

 P_2 and V_2 obtained by extrapolation of the curves, PS and VS against V plot (Fig. 4.15) at proper sections of the curve P_1 was obtained by PS against 1/V plot, extrapolation to 1/V = 0. As we have

already discussed in the case of HRH that v₁ obtained by VS against 1/V plot give a value, 8.2 c.c/mole which is not to be taken very seriously, because with the continuous variation of the apparent molar volume of the monomer varies continuously with the change of concentration at this region. However, the results of such investigation is tabulated below:

Table 4.6

	1	0	imer (c.c/mole	The total of the t		K (l/mol	e)
Molar property	Monomer (c.c/mole)		cyclic	open	cyclic	open	closed
Volume		43x2	60x2	1380000	750000	e tolky m	

We see that we can confidently obtain K values for both types of interactions - open chain dimer and closed chain dimer association interaction by polar method because molar polarization remain uneffected by solute-hole interaction, because the volume effect have been incorporated properly in specific polarization calculation. But in the case rated properly in specific polarization calculation. But in the case of molar volume the effect of 'solute-hole interaction' largely remains.

Thus, the calculation V_1 (although we obtain by usual method to be 8.2 c.c/mole) is not useful for determination of values of K for both these interactions. One way of finding V_1 would be to calculate them by using the values of K from polar data. Thus, we obtain the V_1 values for open and closed chain dimer as follows:

Open chain : $\frac{1580000}{26960}$ = 58.60/2 = 13.7 c.c/mole

Closed chain: $\frac{750000}{25000}$ = 45 c.c/mole.

Thus we find $V_1\approx 14$ c.c. at a stage of interaction when open chain dimer is formed. That means only few of the added solute are out of "holes". Then again $V_1\approx 45$ c.c/mole when cyclic dimer begins to form. This would mean that quite a large number of solute molecules are outside the "hole" at this stage. It is to be noted with some satisfaction that V_2 for cyclic dimer in CCl₄ and C_6H_6 are about 2X59 c.c/mole; where as, in IBH is found to be 2X63 c.c/mole and cyclohexane 2X60 c.c/mole. This close agreement is quite remarkable. Note in the case of chlorobenzene the cyclic dimers have $V_2=2X69.47$ c.c/mole. The reasons will be discussed in the next chapter.

The results of all the five systems are given in the table 4.7.

Table 4.7

Consolidated Results, of the Systems of Acon with Different Solvents.

	Molon	Monogler	Dimer(c.c/mole)	mole)	K (1/mole	10)	1 1
Acon in	ty	(c.c/mole)	Open	Cyclic	Open	Cyclic	1
5	Volume	56,26		118.16	•	17.84	
4	Polarization	23.10		38,38		13.43	
:	Volume	54.18	107.06	118.26	23.08	26.26	
8 8 8 8	Polarization	40.20	139.20	32,34	24.49	23,30	
5	Volume	66.81		136.60		16.28†	
6.37	Polarization	51.56	•	90.92	=	18.85†	
Ē	Volume	52.49*	. 7. 6	126.8	*	:	
	Polarization	32.2	1	39.98	•	46.68	
Cyclohexane	Volume	13.7 45.0	86.0	120.0	*	:	
	Polarization	16.4	73.6	22.4	26.96	25.00	

· Calculated values.

The value could not be calculated since D was not known.

⁺ Most probably a distinct complex of linear dimer-chlorobenzene is formed.

Pit P St St

If the relation of the property and the property of the proper

CHAPTER-5

DISCUSSION

Fotopinga, value

Me. Set

condition a plantage expects exercisely as authors as any evening state expenses

elas, Es ens, dae de existencessarios i forcables pa electe de Mig. Mel.a

come a well have a keeps at according with contra accid

Will and the section of Discussion a descripe a seculi this contracts

Molar Volume

Monomer: If is encouraging to note that the molar volume obtained in CCl₄, benzene and HBH are 56.26 c.c., 54.18 c.c. and 52.49 c.c. respectively, which can be considered very close to each other. The reasons for slight variations among them are not very difficult to find on the basés of various solvents effects noted in the Introduction (page-22).

Carbontetrachloride possibly form a weak charge - transfer complex.

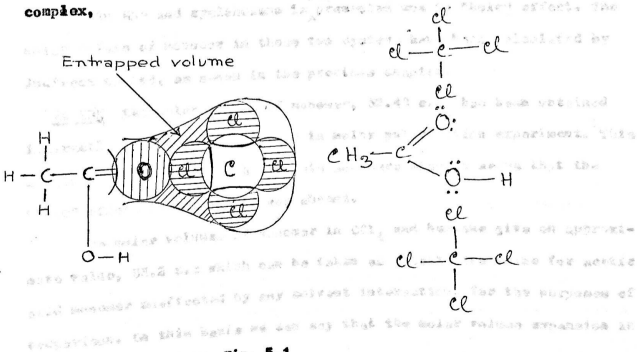


Fig. 5.1

causing a slight overall expansion of volume of acetic acid monomer

due to entrapped volume within adducts despite a small volume contraction, if any, due to oxygen-chlorine interaction as shown in Fig. 5.1.

Benzene forms a well known type of H-bonding with acetic acid as shown,

which causes volume contraction. The said to the districtions The said to

The straight forward determination of apparent molar volume of monomer in HBH and cyclobexane is presented due to "hole" effect. The molar volume of monomer in these two systems have been calculated by indirect method, as shown in the previous chapter.

captain by combine that of measures give for widers the volume constractions

In HBH the molar volume of monomer, 52.49 c.c. has been obtained indirectly from K-value obtained in molar polarization experiment. This should be taken as a rough estimate and here also it seems that the "hole" effect is not altogether absent.

The molar volumes of monomer in CCl₄ and benzene give an approximate value, 55.2 c.c which can be taken as a tentative value for acetic acid monomer uneffected by any solvent interaction, for the purposes of comparison. On this basis we can say that the molar volume expansion in

CCl₄ and volume contraction in benzene is about 1 c.c., while in HBH the volume loss of the monomer species is about 2.5 c.c. - a negligible amount; and in cyclohexane volume loss is about 10 c.c. which is considerably high.

Dimer: Only in benzene and cyclohexane, as already stated,
linear dimer could be identified, whose molar volume are found to be
107.06 c.c. and 86.00 c.c. respectively. The molar volume of dimer
should be double that of monomer plus (or minus) the volume contraction
(or expansion) due to structural changes and bond formation. The molar
volume of monomer in benzene being 54.18 c.c., we have here a volume
contraction due to linear dimer formation of 1.3 c.c. But more careful
calculation should be on the following basis,

Therefore, D = 54.18 + 55.2 - 107.06 = 2.32 c.c, where D is the volume contraction.

In case of cyclohexane, the molar volume of monomer being approximately 45.0 c.c.D for linear dimer formation should be as follows:

D = 2x45.0 - 86.0 = 4 c.c;

which seem to be a little higher than expected. But considering that 45.0 c.c is only an approximate (obtained indirectly), we can say that the method employed here give a value not far off from the mark. Here we find that the "hole" effect on linear dimer is as strong as on the monomer.

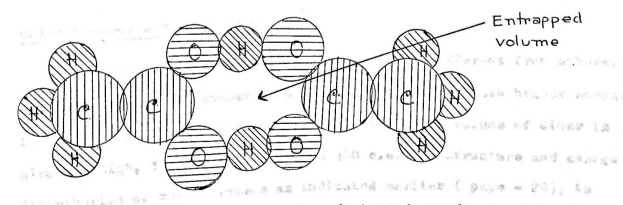
In CCl₄, benzene, cyclohexane and HBH the molar volume of dimer have been found to be 118.16, 118.26, 120.0 and 126.80 c.c respectively. The first two values are remarkably closer to each other; the value in cyclohexane is not far off; and in HBH a slightly higher value is obtained. If we take into account of the fact that in each case, the presence of different amount of linear dimer is likely to be present along with the cyclic form, then these values should be considered quite satisfactory. Thinking in these line we can say that in HBH dimer may consist mostly of cyclic form and in cyclohexane a little more of linear form is present, while in CCl₄ and benzene that is present in higher quantities due to an equilibrium of.

$$cH_{3}-e^{-1}$$
 $cH_{3}-e^{-1}$
 $cH_{3}-e^{-1}$
 $cH_{3}-e^{-1}$
 $cH_{3}-e^{-1}$
 $cH_{3}-e^{-1}$
 $cH_{3}-e^{-1}$
 $cH_{3}-e^{-1}$
 $cH_{3}-e^{-1}$
 $cH_{3}-e^{-1}$
 $cH_{3}-e^{-1}$

As we have indicated that due to interaction of solvents with monomer and linear dimer, the linear form should be present to a greater extent in CCl₄ and benzene.

The cyclic form has the following structure,

This closed structure should have some special characteristics, such as, low dipole moment (consequent to which it should have lower dielectric constant), less solvent effects, such as, weak charge-transfer and other dipole- dipole interactions and large volume expansion due to cyclic structure. A model calculation on volume effect, as shown in the following figure, give a net volume expansion of about 9-10 c.c, taking



into account the volume contraction of about 4 c.c for two H-bonding.

The volume expansion is due to "entrapped" volume inside the ring as shown in figure.

error of he a this tend to have a fine-boar complex mas, may strongly

We find the following values of D, by the following relation,

 $D = 55.2\chi^2 - molar volume of dimer;$

D in CCl_A - 7.76 c.c expansion

Benzene - 7.86 c.c expansion

Cyclohexane - 9.60 c.c expansion

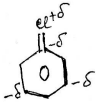
HBH - 15.6 c.c expansion

The value obtained in HEH seem to be a little too high and that may be due to progressively increasing formation of cyclic trimer, which also has smaller dipole, with the increase of concentration of the solution. This is very likely, in non-polar solvent the dissolved species tend to become as much non-polar as possible by induction effect or structural change by self-association or some other changes within the structure.

Chlorobenzene - Complex Formation

The case of chlorobenzene system is quite different from others. The molar volume of monomer is as high as 66.81 c.c (much higher compared to 55.2 c.c in HBH), on the other hand, the molar volume of dimer is also too high, 136.6 c.c compared to 120 c.c. The structure and charge distribution of chlorobenzene as indicated earlier (page - 26), is

like



It is obvious that C=0 group of acetic acid can form a good charge transfer complex with Cl-atom and on the other hand, the H-atom of OH
group of ACOH will tend to form a T-H-bond complex much more strongly
than benzene, forming a complex like,

(chlorobenzene - acetic acid complex)

Thus D for molar volume change, shows an expansion of volume of 11.61 c.c. This is not unreasonable, when we look at entrapped volume within structure of the chlorobenzene - acetic acid complex. We should calculate D for dimer formation on the following basis:

D = 66.81 + 55.2 - 2.3 - 136.60 = 16.89 c.c.

This means that another, 16.89 - 11.61 = 5.28 c.c. expansion is attended on attachment of second monomer to chlorobenzene - acetic acid complex. We must then seek a Structure which can justify this second volume expansion. This can be achieved by the following proposed structure.

or a comparation of according to the different and control of the control of the

In this structure one end of the linear dimer i.e. CO group attaches itself to Cl⁺⁶ atom while the free acidic H-atom attached itself to T electron of the ring.

The consideration of vector addition of moment of this structure indicate that the dimer-complex should have moment or molar polarization less than the double of that of monomer - complex. That is what we have obtained in molar polarization experiment. The monomer complex has molar polarization, 51.56 c.c and dimer complex has 90.92 c.c which shows that 12.2 c.c molar polarization is decreased on formation of dimer complex.

Thus this probable structure satisfy two observable molar properties V_2 and P_2 and D for molar volume and polarization.

sons tore bory carefully on the basis of intersections on have already

Molar Polarization

Molar polarization of monomer obtained in different solvents are widely apart from each other, whereas, we have in three solvents the molar volume, very close to each other, as shown in the following:

are alway	Carbon tetra	Benzene A e. p	нви 👉 с	yclohexane 🐭	C6H5CI
Molar volume	56.26 C.C.	54. 18 C.C.	52.49 €.€	45.0 C.C	66.81 C.C
Molar polarizat:	ion 23.10 C.C	40.20 C.C	32.20 C· C	16.4 C.C	51.56 €.€.

weak charge - transfer complex and dipole - dipole interactions can lead to change of volume from less than 1 c.c to about 5 c.c. This explains the close resemblance of molar volumes of acetic acid monomer in CCl₄, benzene and HBH. The somewhat larger departure of molar volume of acetic acid monomer in cyclohexane and C₆H₅Cl has been shown to be due to "hole" effect and a complex formation respectively. Here in case of molar polarization we are confronted with the problem of solvent effects of various kinds.

88,92,93

Thus a conformity of results of molar effects of various kinds.

polarization of monomer acetic acid, which is a very highly polar molecule, in different solvent is can be expected. But we can discuss each case very carefully on the basis of interactions we have already identified.

i) In Carbon Tetrachloride the formation of a weak complex have been suggested by the Fig. 5.1. The main dipole is along the C - C bond and the induced dipole by CCl₄ molecules on both the oxygen attached carbon are along the right angle to it in opposite directions, as shown below:

cely

$$Cel_4$$
 Cel_4
 $Cel_$

This explains why the monomer acetic acid should have lower moment and consequently molar polarization in CCl₄. Had the adduct joined along the main axis the moment would have increased.

ii) The formation of W-H-bond complex with benzene along the axis of main dipole of acetic acid obviously increase the dipole of the

complex and the highest value obtain in benzene compared to those in CCl₄. IBH and cyclohexane is thus satisfactorily explained. The molar polarization of linear dimer in benzene is also very high for the same reason.

- iii) The value obtained in IBH, it seems to us, can be taken as the "true" value for monomer unaffected by any solvent effect. The reason for such assignment is quite simple. Although there is a "hole" effect which cause a little volume loss the monomer can freely orientate itself under alternating electrical field in a dielectric measurement.
- iv) The "hole" effect in cyclohexame on the monomer, as well as linear and cyclic dimers seem to be quite strong. We are lead to believe on the evidence of these surprisingly low values of molar polarization of monomer and dimers that the species in the "hele" are very strongly prevented from free movement. Hence the contribution of orientation polarization of these species in the "hole" are bound to be less. Thus taking the value 32.20 c.c as normal (in HBH), 16.4 c.c in cyclohexane is certainly very low.

The value for linear dimer is only 73.6 c.c whereas it should have been somewhat less than 139.20 cc which is the value for linear dimer in benzene. Again we find that the even the cyclic dimer in this solvent have a very low value compared to others. All these indicate

that "hole" has restrictive effect on the free movement of the species in such a condition.

- zation than monomer and linear dimer is obvious due to its symmetrical structure. They seem to be less effected by solvent effects. In CCl₄, benzene and IBH the values are 38.38, 32.34 and 39.98 c.c respectively. The small differences between them is not easy to explain. But we advocate that the value obtained in IBH should be taken as a reasonable value because of the possibility that it is less effected by the solvent interactions. The smaller value in cyclohexane is already discussed above.
- lead to the formation of complexes and has already been shown. A careful explanation of this structure shows that they should have higher molar polarization compared to others. It has already been shown in the experimental results that cyclic dimer in chlorobenzene can not exist. The explanation is simple. The complex formation with monomer and linear form is so strong that the cyclic form is broken up as readily as its formed.

Equilibrium Constant

In the present investigation, in each set of experiments we have determined three unknown quantities, molar properties of monomer and

dimer of AcOH and the equilibrium constant related to particular type of association (linear or cyclic). Looking at the Table 4.7 we find tolerably closer values of some of these quantities. Most remarkable conformity or closeness of values are found in the values of equilibrium constants. There are two sets of values:

to B handary with The electrical of homeoner with the color of crisis in the

17.84, 13.43 and 16.28, 18.85 l/mole

(for CCl_A and chlorobenzene)

the nation of policing the benefits welcoming linksburges their not been,

Corporate and

fa fally with the late

(ii) 23.08, 26.26, 24.49, 23.30 and 26.96, 25.0 1/mole (for benzene and cyclohexane).

The values in CCl, and chlorobenzene are much lower than those of benzene and cyclohexane, while the value in HBH is still higher than those and stand apart. It is also interesting to note that the equilibrium constants for linear and cyclic dimer association in benzene and cyclohexane are nearly equal to each other. We have equilibrium constant in HBH for only cyclic form which is nearly the sum of equilibrium constants for linear and cyclic dimers in benzene or cyclehexane.

In two types of association linear and cyclic, we find in the processing of results that they occur in stages and not simultaneously, contrary to expectation according to the scheme of interactions presented earlier (page- 20). The explanation of this effect is not easy. In case allegai tota es contes which can bern allegarosoletisteria. These in the expective

that the west of discussions therein will receive actions within a

of benzene system acetic acid monomer has greater tendency to form a H-bonding with ring W-electrons of benzene and so does the linear dimer. It indicates that in such a system dimer will remain more in open form than cyclic a so long the concentration of acetic acid monomer and linear dimer are not too high to be able to remain engaged in H-bonding with W- electron of benzene. But in case of cyclohexane the explanation will not be easy until the "hole" mistry of this solvent is fully understood.

2/2/

In two sets of values, the higher values, including that of HBH, are associated with hydrocarbon nature of the solvent, i.e. non-polarizability of the atoms, while the lower values of equilibrium constant is associated with chloro-compounds in CCl₄ and chlorobenzene. In these two latter solvents we have indicated that Cl-atom can form charge-transfer complex with C=O group. This complexation virtually remove, or in other words, inactivate the donor property of C=O which consequent lead to formation of lesser quantity of dimers by self association which obviously must lower the equilibrium constant of the association interaction.

In conclusion we are to remark that our equilibrium constant values are the lowest so far obtained and it nearly conform to the value obtained by Simamura. The method developed in this work has a general importance in the fact that this proceedings can be applied to any other chemical species which can have self-associations. Thus it is expected that the method described therein will receive serious attention it deserves.

F1121 63

- A Timerumies on the restrict accessive with the determination of Friedrich Counties of sortic and discor in different contains in the literature and the restrict and the restrict and by the face (i) the har secondard in province a different and by the 1, the war (it was and optic) and bighter prights of the vertices types whenly exist planking county in continion and first different advanced about the periods at a section of the transfer and characters and the periods are formally at the conscious of the confidence of the cold that an exclusive content and the formalisms are exclusived as the arrangement of the content and the content and the transfer and transfer and the transfer and t
- A themselfond from new him being continues for the wildy of the water production for the wildy of the water product of the wild as well to invest the wild from the water to invest the wild from the water to be a continued to the product of the product to the product of the product of the product to the water to be a second to the wind of the water to the water to be a second to the water to the water to be a second to the water to the water to be a second to the water to the water

of a following to the test of applicable and the send to a probable to a to a probabl

S. (1) A description of the soline for the Swinning time of appoint

SUMMARY

s of the transferentian and confere

- A discussion on the problems associated with the determination of Formation Constant of acetic acid dimer in different solvents in the Introduction show that (i) various associated species dimer (linear and cyclic), trimer (linear and cyclic) and higher polymer of the various types should exist simultaneously in equilibrium and (ii) different solvents should have various "solvent effects" including the possibility of formation of distinct complexes. At a suitably chosen dilute region, it is expected that only the first association i. e. dimer formation, can exist with the effective exclusion of others and the investigation can be conducted at this region to study this particular association interaction.
- 2. A theoretical frame work has been constructed for the study of the variation of "apparent molar property", such as molar volume and molar polarization, whereby it is shown that the "apparent molar property", PS is related linearly to volume, V which contain one mole of the added species, as follows:

 $PS = \frac{K}{D} \times \frac{(P_2 - 2PS)^2}{V} + P_1$; where the symbols has the usual

significants according to list of symbols. How this relation has been used, to determine P₁, P₂ and KD graphically and finally D and K, is explained.

3. (i) A description of the method for the determination of specific washed volume of liquid solution with the help of a Magnetic Float Densimeter

is presented along with the details of the construction and manipulation of the device. (ii) The method for the determination of dielectric constant with the help of a Dipole meter has been described with full particulars. (iii) The details of purification and particularly, careful dehydration have been fully described. (iv) Finally, how the variation of concentration of solution have been accomplished is reported.

- benzene, cyclohexane and high beiling hydrocarbon (HBH), the specific volumes and dielectric constants have been measured. The linear plots of these quantities against weight fractions have been carefully scrutinized to obtain finally a minimum error-free values at regularly spaced weight fractions. From these " refined values" the various parameters, P, V, PS, VS etc, have been calculated; from which various plots have been constructed related to various equations to obtain P₁, P₂ and KD. The results of such efforts have been presented in the Table 4.7.
- formation dimer is obtained between values 13.43 to 18.85 litre/mole.

 (ii) In benzene and cyclohexane the value is between 23.08 to 26.96

 (iii) The value in HBH is 46.68 litre/mole.

 The lower value of equilibrium constant in (i) has been explained as due to weak charge-transfer interaction between Cl-atom of solvent with

6.

C=O group of the acid. In case of benzene and cyclohexane two types of dimer, linear and cyclic form, the K values are nearly same in both cases; whereas in MBH only cyclic form is observable, in which case K value is almost equal to the sum of the values for linear and cyclic form in benzene and cyclohexane.

(i) The molar volume of cyclic dimer obtained in different solvents are found to be very close to each other- 118.16 c.c in CCl4, 118.26 c.c in benzene, 120 c.c in cyclohexane and 126.8 c.c in HBH. It is suggested that the closed structure of cyclic dimer form is likely to be least affected by solvents and they should have very similar values. The formation of cyclic form is attended with volume expansion of about 10-14 c.c which is accounted for by considering the entrapped volume in the closed structure. (ii) The linear dimer in benzene have molar volume 107.06 c.c which is attended with volume contraction of 2.32 c.c - this value is reasonable for one H-bond considering the contraction of volume of about 1 c.c for T-H-bond formation with the end H-atom of the linear dimer. The molar volume of this form in cyclohexane is much less than expected and is explained by the loss of volume due to the "hole" effect. The calculation shows that the formation of this type, as expected, is attended with volume contraction of 4 c.c this value seem to be a little higher, although not unexpected. (iii) The range of values obtained for molar volume of monomer (excepting in chlorobenzene where complex formation takes place and in HBH and cyclohexape where the "hole" effect is strong) are quite

close to each other, between 54.18 to 56.26 c.c.

(i) The molar polarization of monomer have been found to be different in different solvents. It has been pointed out that the various effects of solvents including complex formation and "hole" effect, would always have profound effect on the measured molar polarization. (ii) The molar polarization of linear dimer in benzene have a higher value due to linear alignment of two monomers. This is found to be true in case of cyclohexane if the molar polarization of monomer in this solvent is taken into account. (iii) The molar polarization of cyclic form have much less values than that of monomer and linear dimer as can be expected on the symmetrical structure.

It has been pointed out that the molar polarization of monomer, linear dimer and cyclic dimer in cyclohexane are always much less than the corresponding values of these species in other solvent. It has been argued that the species in "hole" of the cyclohexane solvent have a profound restrictive effect on their free movement which consiquently keeps the orientation polarization low.

In conclusion, it is hoped that the method developed in this work will be considered to be applicable to any other monomer-dimer equilibrium system for investigation and evaluation of different parameters.

Frankli werde grid in the

This is the presidental results of specific volume

	The second second second second second	piffermon in ager ciffe rollings of (observable)
7.4	0.471201	4 9
86.77.5974	0, 817723	e i
0,40000	477.153	· 3
1.000000	1 (1.22)	117 Q
0.0411100	PPENDIX	* 1
	D. 1004 A	* 6
. 0.606763	C. 6 55784	a 22
0.009.804	0.631227	× £
0,000.00	6,012.90	
	0.63 22 6	
	47、60万D 14.5	
n. (C. 450)	3.40 PBS 3	
G. (3519)		* 5
\$ 140 FP	0.435/11	
0.17169	V 4 5 5 50 6 3	40 m ⁻² / ¹
0.46840	2.60387	
0. 273330	0.61 585	4.2
0. 105610	G. Sarbe Es	
0.21K-350	0.635066	***

TOTAL

APPENDIX- A. 1

System: Acetic Acid in CCl4

Table 1. Experimental results of specific volume

fraction(w)X10 ²	Observed specific volume(v)c.c./gm.	Difference in specific volume, △v (obs cal.) X10 ⁶ c.c./gm.
0.0	0.635001	+16
0.011489	0.635021	* 1
0.022936	0.635053	- 2
0.034343	0.635088	4.3 - 1
0.045709	0.635125	-C+1
0.057034	0.635156	-8 - 2
0.066765	0.635189	+ 2
0.079564	0.635227	and * 1
0.090769	0.635268	+ 8
0.101934	0.635306	+12
0.113059	0.635344	+16
0. 124140	0.635371	+10
0.135190	0.635390	- 5
0.146200	0.635411	- 17
0.157169	0.635441	-20
0.168100	0.635501	+ 7
0. 178990	0.635565	+38
0.189840	0.635615	+55
0.200660	0.635666	+73

Vcal. = 0.634985 + 0.30303XW.

Table 2. Experimental result of dielectric constant.

(Weight fraction	on)X10 ²	Observed d.c	△€(obscal.)X10
0.0	6.57 s.x.	2.251207	0.0
0.01520	0.003	2.251372	- 20 498
0.03040	6 CO 5 43	2.251516	- 60
0.05167 0.06951	0,858670	2.251749 2.251957	- 85 - 94
0.08780	9.01 1.6	2.252164	-109
0.11073	0.000100	2.252328	-223
0.14176 0.16232	0.61/107	2.252404 2.253478	-524 -699
0.18413	0.635139	2.252596	-847
0.20150	0.435230	2.252756	-898
0.22627	6,575768	2.253121 2.253589	-833 -569
0.26097	o, e ilisio	2.254107	-269
0.27583	0.077553	2.254556	0.0
L3788 -	6,8,31.7	2,207.15	0.907109
	8,610013	= 2.251207 + 1.21	0.407104
	eal.	# 2.25(20) + 1.21	4. W. D.
, VD 15	0.375 334	2,252476	0. 167 (9)
, 1.5° \$60	0.4.551	2,532,638	0. 117 117
	0,5000	S. 752 557	6. 307 274
1,1158	6.200270	\$ 。 它 \$20 古佛等	9.28.203
	6.63607	意。不是有意味。	0,00720
8.00	4.837516	5,057748	5, 537, 53

The set of the foliated parameters from sectional datas

Table 3. Refined values of specific volume and dielectric const. The state of the s

10000	Weight fraction	Specific volu (c.c./gm.)		d. c.	Specific position (c	
0.1893	5 / / / 4,75	The state of the s	20.31		4.24.54.40	
CARCI	0.0	0.634986		2.251207	0.186888	
· ALL TEN	0.0001	0.635015	2 ·	2.251306	0.186907	y 136, (
0,5	0.0002	0.635046		2.251410	0.186928	
0.000	0.0003	0.635076		2.251526	0.186948	
C	0.0004	0.635106	25.30	2.251636	0. 186968	
5 - 18 W	0.0005	0.635136	27.	2.251744	0.186989	244.50
P. 图 4.0	0.0006	0.635167	70.0	2.251855	0.187009	2010 17
a.dugo	0.0007	0.635198	55 21	2.251954	0.187029	14" , E
0,30,30	0.0008	0.635230	5.160	2.252035	0.187047	
0,000	0.0099	0.635265	dig to	2.252108	0.187065	1 St. 16. 4
0.6943	0.0010	0.635300	21.3	2.252162	0.187081	51.79.8
0.7011	0.0011	0.635333	22,00	2.252200	0. 187095	5 704,80
Sicori	0.0012	0.635362	31.75	2.252255	0. 187 109 0. 187 121	3.5 004, 57
0,0010	0.0013	0.635381	23.68	2.252316	0.187121	Tan 1. 18
\$ 16.5°	0.0014	0.635399	23.30	2.252479	0. 187 150	North Hi
0.00 %	0.0015	0.635421	21,73	2.252556	0. 187 167	2000
0.0117	0.0016	0.635512	31,9	2.252547	0. 187 184	23F9 S. 70
6.00.20	0.0017	0.635570	20,01	2.252565	0. 187203	1967
	0.0018		the state of the state of the state of	2.252654	0. 187226	lander progression of the lander of the
A william growth state that I have been strongly to the	0.0019	0.635617		2.252745	0. 187250	
	0.0020	0.635664			0. 10.200	

Table 4. Calculated parameters from refined data.

weight fraction	V(c.c/mole)	Vs(c.c.	/mole)	PS(c.c/mole)) △(V•VS)cc ² /mole ²	² △(v.Ps) c.c ² /mo
u u	and the training		9/4770	(moin(s.e)	Patricial maps parame	1.00 20 3
0.0001	381326.51	56. 15	Service was a service	22.63	-22880.0 -	179223.40
0.0002	190672.56	56.44		23.23	+43855.0 +	24787.40
0.0003	127121.05	56.35	25	23.23	+17796.9 +	16525.70
0.0004	95345.29	56.30	5	23.30	¥ 8581.1 +	19069.10
0.0005	76279.89	56.32	17	23.35	+ 8390.8 +	19070.00
0.0006	63569.63	56.35	34	23.33	+ 8899.7 +	14621.00
0.0007	54490.91	56.40	-	23.30	+ 10353.3 +	10898.20
0.0008	47681.95	56.52	15.	23.15	+ 14781.4 +	2384.10
0.0009	42386.29	56.81	1	23.04	+ 25431.8	2543.17
0.0010	39149 .77	57.05	27	22.83	+ 32045.8 -	10300.44
0.0011	34683.41	57.08		22.54	+ 33643.0 -	19422.74
0.0012	31794.57	56.99	CALLS SEE SEE SEE SEE	22.27	+ 24799.8 -	26389.Z 9
0.0013	29349.71	56.42		21.99	+ 6163 .5 -	32578.18
0.0014	27254.08	55.89		21.90	- 8721.3 -	32704.89
0.0015	25438.02	55.58		21.72		35104.47
0.0016	23849.27	55.62		21.68	- 14071.1 -	33865.96
0.0017	22448.53	56.75		21.68		31876.91
0.0018	21203.32	57.65		21.73		29048.55
0.0019	20088.84	58.10		21.91		23905.72
0.0020	19085.81	58.52		22.09	+ 44088.3	19276.67

Table 5. Parameters VS, .PS and 1/V .

	Weight	fraction X 104	1/VX10 ⁷ (mole/c.c)	VS(c.c/mole) PS(c.c/mole	e)
0			-	· · · · · · · · · · · · · · · · · · ·	•
	Conservation Comments		26.22	56.15 - 10.00 22.63	
	Afficially and the second of the second of the second	2	52.44	56.44 23.23	
٠	0.0	3	78.66	56.35 23.23	
		4	104.88	56.30 23.30	
	8 St. Co.	5	131.09	56.32 23.35	
u .	4, 7, 75	6	157.30	56.35 23.33	
1 *	10 Fa 14574	7	183.51	56.40 23.30	
4	Page 124	8	209.72	56.52 23.15	
	有,是实现 。		5-122231	- 8, 47	
*	7,000		AL L'OCTAGE!	- 5, 42	
					-
	0.701.3		5. 98 3355	9, 33	
	Con a sty or		A 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	~ 10,09	
	0.13002	, &	4-101011	- 19,24	
	0.00		10 F 15 B	- 30 4 1	*
	66 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	,	14.3842*O	一 数十二章	4
	De Charles		8. 3.11 *160	\$17 a \$7"	
Š	e, 10274		1. 13/107	- E0.28	-
			1. 161117, ***	- 9.37	Ų.
*	台。据符》。		1,751301	7.89	
*	ALL THE STATE OF	. *	1. 3.258	- 3.73	
			1. 10 374	- 4. 63	
	1.304		5. 15 10.33		
	4.1 2			4, 8, 99	
	7,1207		1.101035		
	0. \$ 75.5		3. 13.44.23	5.0	

\$ 10 to 10 1990 . (- 0, 15 12) w .

APPENDIX-A.2

System: Acetic Acid in Benzene

Table 1. Experimental results of specific volume

(Weight fraction)X10 ²		Observed specific volume Difference in specific (v) (c.c/gm) volume, \(\sigma v \) (obscal.		
		A CONTRACT	the graph of	(c.c/gm.)
	0.0	1. 151490	3.70% 23 0.0	
	0.01189	1. 151464	0.81	
	0.01782	1. 151450	1.31	
	0.02373	1. 151429	- 2.52	
	0.02964	1. 151415	3.02	٠
	0.03553	1. 151400	- 3.63	
10 A	0.04728	1. 151464	- 5.46	
	0.05899	1. 151341	- 5.99	
	0.07065	1. 151309	-7.42	
	0.08226	1. 151282	8.37	e
	0.09383	1. 151255	- 9.32	
	0.10536	1. 151230	-10.07	
*1.1	0.11684	1. 151211	- 10.24	
	0.12827	1. 151193	- 10.31	. je
	0.13966	1.151176	-10.29	3 4 4
	0.15101	1. 151160	- 10. 17	, , , , , , , , , , , , , , , , , , ,
	0.16231	1. 151142	-10.26	
	0.18478	1. 151117	- 9.37	
	0.20709	1.151101	-7.59	
	0.22922	1. 151086	- 5.75	
	0.25118	1. 151068	-4.23	
	0.27298	1. 151052	-2.53	
	0.29461	1. 151033	- 1. 16	
	0.31608	1. 151012	0.0	
	4107-			

V_{cal}. = 1.151490 + (= 0.1512) w .

Table 2. Experimental results of dielectric constant

Some the trifficant was one of whomethe walling and distance the

is -	(Weight fraction)X1	O ² Transport Accommon B. C	
Cal.	N 48 52540	2.21700	0,3377,47
1.66	0.0	2.295660	6.337.536
0.4.0			0.377312
0.655	0.0110	2.295901	
0.00	0.0302	2.296421	0.000
0.00	0.0490	2.297120	- 0. A 17 to 2
	0.0819	2.298539	9.327653
O. Atlat	1.4 4314	2 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 -	0.307909
Note 15th	0.1095	2.299750	0.247 ec
16. B.F. 6	0.1541	2.300009	4. July 18
	0.2016	2.300232	0.30733
77 12 8 3		F. Land	0.057900
0.40%	0.2582	2.300910	, ASSET O
6-14-7	0.3021	2.301509	
50 g =	0.3496	2.302381	0.090109
A 600 E.S	4. 20.154	2.000	. 4.3523
4.00	表。 是 3 代表 4 7 7	3.30000	. 2 3 2 2 4 7 5
0.000		9 205660	0.219 -1
6,000	AE = E	cal - 2.295660.	
6,12,30	3-422443	2 : 501	6.7/FQ(3)
A Base Control	E. 17 1 10 7	2 - 10 2 6 6	0.00000
(1.00)		2.37字行者	is a second
	表。1775日4	the telephone to	0.305079
	1. 15 (.46	2.777631	0.30010
	中国	是。是上的有一位。	0.34863
1.70	4. 17.1677	0.300773	0.540.40

Table 3. Refined values of specific valuee and dielectric constant.

			(c.c/ga.)
0.0	1. 151490	2.295660	0.347313
0.0001	1. 1514/65	2.295871	0.347345
0.0002	1. 151440	2.296179	0.347395
0.0003	1. 151414	2.296461	0.347440
0.0004	1. 151389	2.296802	0.347496
0.0005	1. 151364	2.297182	0.347560
0.0006	1. 151333	2.297610	0.347630
0.0007	1.151311	2.298041	0.347705
0.0008	1. 151287	2,298480	0.347780
0.0009	1. 151263	2.298983	0.347866
0.0010	1.151241	2.299431	0.347943
0.0011	1. 151223	2.299749	0.347997
0.0012	1. 151207	2.299901	0.348020
0.0013	1. 151191	2.299952	0.348025
0.0014	1. 151176	2.300000	0.348030
0.0015	1. 151161	2.300000	0.348025
0.0016	1. 151147	2.300000	0.348021
0.0017	1. 151134	2.300031	0.348023
0.0018	1.151122	2.300049	0.348023
0.0019	1.151113	2.300101	0.348030
0.0020	1.151105	2.300199	0.348046
0.0021	1. 151098	2.300298	0.348062
0.0022	1. 151092	2.300401	0.348079
0.0023	1. 151086	2.300521	0.348100
0.0024	1.151079	2.300650	0.348122

Table 4. Calculated parameters from refined data.

Weight V(c.c/mole) fraction	VS(c.c/mole) FS(c.c/mole)	△(V.VS) △(V.PS)
THE STREET AND	in the rest appetite waters	(c.c ² /mole ²) (c.c ² /mole
0.0001 691454.73	54.14 40.08	- 27644.0 - 82975
0.0002 345719.82	54.10 45.48	- 27657.0 1825401
0.0003 230474.62	54.09 46.27	- 20742.0 1398981
0.0004 172852.33	54.05 48.40	- 22471.2 1417388
0.0005 138278.80	54.00 50.52	- 24890.3 1427037
115229.29	53.95 52.59	- 26502.9 1427690
98766.05	53.90 54.48	- 27654.5 1410378
0.0008 86418.45	53.88 55.92	- 25925.6 1358498
0.0009 76814.83	54.01 57.69	- 13058.4 1243491
0.0010 69132.01	54.18 58.67	• 0.30 1276868
62846.29	54.55 58.20	+ 23253.1 1131233
57608.30	54.97	+ 45510.4 9246134
0.0013 53176.18	55.34 53.74	+ 61684.2 720005
0.0014 49377.22	55.68 51.59	+ 74066.3 562406.8
17.	55.99 49.36	+ 83413.5 492137.3
	56.26 47.42	+ 89864.0 311932.5
	56.56 45.92	+ 96775.5 232587.6
	56.87	+ 103303.2 166284.1
	57.23 23 43.50	+ 110962.6 120058
36381.22	57.58 . 8 13 42.85	+ 117510.2 91588.9
0.0020 34561.92	57.93	+ 123435.0 68136.5
32915.91	58.29 41.76	+ 129134.2 49014.9
0.0022 31419.58	58.59 41.39	+ 132535.5 35763.4
30053.35	41.09	+ 134788.60 25632.9
28800.95		-1, 43
2.36.7	0.8(1832	秦司 建200

APPENDIX A. 3

System: Acetic Acid in C₆H₅Cl .

Table 1. Experimental results of specific volume

		erved specific		lume Av(ob	specifi scal.) (c.c/gm.
e e e e e e e e e e e e e e e e e e e	E-12-01			4	
0.0		0.912825	1 2 1 3 2 4 3	0.0	
0.0055	0.0070	0,912834	5.611.64	-0.45	
0.0110 9.0156	50, 1150 B	0.912846 0.912857	. S.R. PRO	-0.61 -0.64	
9.0221	型,2000年第	0.912871	0.763730	-0.84	
0.0280 0.0332	新,艺术教	0.912880 0.912892	#. 5 <u>(</u> < ₹ 4 €	-1.40 -1.48	
0.0443 0.0509	0.183	0.912911 0.912924	5. 6 to - 10	-2.31 -2.64	
0.0553	\$ 10 A	0.912934	5.61170	-2.73	
0.0662	\$, 1190 s	0.912961 0.912908	3.5.73.3	-2.71	
0.0880	17 . 5 . 24	0.913005	3, 5 to 10 d	-3.69	
0.0993	8.3717	0.913030	· 6 6 2 120	-4.10	
0.1088		0.913046	· · · · · · · · · · · · · · · · · · ·	-4.71	
	た。24 達	0.913071		-4.90	
0.1197 0.1298 0.1401		0.913097 0.913120	and referen	-4.79 -5.03	
	Ast .	0.913140	. \$ 5 (CE)	-5.52	
0.1502		0.913194		-5.25	4
0.1710		0.913243 0.913298	3	-5.47 -5.20	
0.2130		0.913353		-4.60	
0.2530		0.913411 0.913467		-3.76 -3.06	
0.2729		0.913532		-1.45	
0.2927 0.3131		0.913597		0.0	

. Intia b. nothers whitem of according welcome and bisherric

Table 2. Experimental results of dielectric constant.

the state of the state of the	(Madaba a	. 2		17.00
19 ₁ (3)	(Weight fraction) X 10 ⁻²	D. C. (€)	2 2 2 3 3 3 4 4 5 5 4 4 6
0.2		*		
S. Garage	0.0	*	5,610160	6. 7 TO . TH
	0.0201		5.611201	い。例如では何 の、数ではもでき
Martines	0.0440		5.611962	
0.000	0.0681		5.612769	C. Prograd
	19. A 4 1 1 1 3 D		5.6.17.559	44 Table 64
0.01	0.0985		5.613710	sr. * Const
15. d	0.1321		5.614149	\$10000
Page 1	0.1610		5.614230	D. A. 197
0.050	50 1 1 19		きょちょうアデュ	A . 1 1 . 1 . 1 . 1
0.005	0.2021		5.614701	7 . 29
Fr 1 45	0.2461		4, 4340 5.6 15889	Parties of
5,000	0.2795		5,617109	6, * 100
0.3514	4 2 2 2 2 2 4 4 5 5 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6		6,414000	
0.00	0.3122		5.618120	6-3-5-5
	0.3413		5. 7. 2. 5.618951	
	0,010	*	5.6 - 441	5.50
	0.50		To \$1,577.0	A
4, 5, 7,		cal.	- 5.610160	4.50
	6. R (%) *	Car	5,514283	
CAR WITTE	6,8450		Maria Sala	
0.000	0,813.85		5.0 10 10 10 1	10 m
0. 44			5.3339	
0.000	0.033.55	1	8.819928	
0.088	0:4:0.7.3		E. G 17 (H)	10 m 10 m
6.427	0.7.473			

Table 3. Refined values of specific volume and dielectric constant.

Wei	ght fraction	specific volume (c.c/gm.)		Specific polarization (c.c/gm.)	
84 S 179 I	0.0	0.912825	5.610250	0,552985	
	0.0001	0.912845	5.610701	0.553018	
	0.0002	0.912865	5.611029	0.553046	
	0.0003	0.912884	5.611429	0. 553076	
	0.0004	0.912905	5.611800	0. 553 106	
	0.0005	0.912925	5.612171	0.553136	
	0.0006	0.912942	5.612549	0.553164	
	0.0007	0.912965	5.612870	0.553193	
	0.0008	0.912986	5.613171	0.553220	
	0.0009	0.913007	5.613472	0.553247	
te 多額	0.0010	0.913029	5.613750	0.553274	
A Page 1	0.0011	0.913051	5.613901	0.553294	
	0.0012	0.913073	5.614050	0.553315	
2.07.83	0.0013	0.913095	5.614149	0. 553333	
14/4 38	0.0014	0.913118	5.614200	0.553349	
14 TH 2 2 A	0.0015	0.913142	5.614232	0.553365	
1. 2. 1.	0.0016	0.913166	5.614251	0.553380	
	0.0017	0.913190	5.614301	0.553397	
	0.0013	0.913215	5.614379	0.553416	
1,		0.913239	5.614519	0.553437	
	0.0019	0.913265	5.614650	0.553459	
5.5.2¥	0.0020	0.913290	5.614852	0.553484	
1	0.0021	0.913317	5.615101	0.553512	
4.21.3	0.0022	0.913345	5.615380	0,553542	
1.000	0.0023	0.913373	5.615669	0.553573	
W/5 5	0.0024	0.913402	5.616000	0.553606	

Table 4. Calculated parameters from refined data.

	(c.c ² / ole ²)
0.0001 548163.48 66.89 53.26 43853.0 + 93187	- 4.5 Z
0.0002 274087.71 66.93	
0.0003 182728.97 66.84	
0.0004 137049.87 66.84 51.47	
0.0005 109642.34 66.05 54.70	
0.0006 91370.25 66.86 51.13 4568.5 - 39289	
0.0007 78319.34 66.89 51.06 6265.5 - 39159	
0.0008 68531.02 66.91 50.90 6853.1 45230	
0.0009 60917.84 66.95 50.70 8528.5 52389	1890 18578
0.0010 54827.39 67.07 50.56 14255.1 54827	
0.0011 49844.26 67.13 50.08 15950.1 73769.	
0.0012 45691.71 67.24 49.72 19647.4 - 84072	
0.0013 42177.98 67.31 49.28 21089.0 -96165.8	
0.0014 39166.24 67.39 48.83 22716.5 - 106923	3.9
0.0015 36556.11 67.50 48.42 25223.7 - 114786	5.2
0.0016 34272.26 67.61 48.05 27417.8 - 120295	5.7
0.0017 32257.09 67.71 47.78 29031.4 - 121931	1.8
0.0018 30465.86 67.82 47.59 30770.5 - 120949	.5
0.0019 28863.17 67.91 47.52 31749.5 - 116607	.2
0.0020 27420.77 68.01 47.44 32904.9 - 112973	.6
0.0021 26115.75 68.12 47.48 34211.6 - 106552	.2
0.0022 24929.41 68.26 47.61 36147.7 - 98471.	1
0.0023 23846.24 68.39 47.75 37677.1 -90884.2	
0.0024 22853.36 68.53 47.92 39307.8 - 83186.	
0.0025 21939.91 68.67 48.12 40808.3 - 75473.	3

APP ENDIX-A.4

System: Acetic Acid in High Boiling Hydrocarbon (HBH) (150°-200°C).

Table 1: Experimental results of specific volume.

			lume (V)	c.c/gr	ΔV (obscal.)X10 ³ (c.c/
0.0	\$ 4 0 10 3		1.323345	,	2,080,040	0.0	
0.0050	0.6461		1.323312	3	0.4 1734	1.69	
0.0103	3,0003		1.323280	÷	2.1-1-1-1-1-	3.20	٠
0.0152			1.323245	ć	46	5.20	
0.0207	0.40 120	•	1.323210		January St.	6.90	
0.0260	1. 1. 1.		1.323181	•	2,000,000 -	8.10	
0.0310	4. 1522		1.323139	3	1.827265	10.70	100
0.0413			1.323092	1	2,033011	12.10	
0.0515	(23.3		1.323039	:	# (trig=1)	14.10	, 5 ×
0.0617	3.20	•	1.322990	į.	2 - 44 S. 33 -	15.70	
0.0718			1.322951	*	Bar Grief	16.29	
0.0819	0.50.3		1.322925		2,05,039	15.83	
0.1021			1.322855	•	-	16. 13	
0.1221	0.000		1.322805	1		14.69	
0.1419			1.322768	•		12.01	1 A
0.1615			1.322742	k		8.30	,
0.1811	*	*	1.322685	•		7.69 3.98	
0.2004	∴	e and	1.322660	9		2.26	
0.2197			1.322571	•.		0.51	
0.2388			1.322515	*		0.0	Y
0.2577			•4				

V_{cal} = 1.323345 + (- 0.32207)Xw

RT 12 4年 1874 。

0.4.5

Table 2. Experimental results of dielectric constant.

were to moralfie mainum and dialestria

	(Notable Const.)	2.0,224	n, 3000an
0.00	(Weight fraction)X40	g. D. C.	0,350 13
	1.3.3218	2.000	. 0.3/5-26.4
0-1,17	0.0	2.063891	0.4.037
0, 1	0.0225	2.064240	6.03605
19 4 M	0.0461	2.064789	0.040704
0.143	t, 5: 4: 43	Telis	0.343833
	0.0625	2.065141	0.045702
	0.0810	2.065511	- 6,7837 tg
12-1-3	0.1082	2.065692	· 0.5%3753
Mesting.	4,37, 500	k. 90 % D	4.545008
4.4. 11	0.1422	2.065790	6,3047.6
0-1-54	0.1765	2.065971	Carlot Top
Garage T	0.1989	2.066210	6,57339
5. P. W.	10.72.7.7	g, the sout	11,74 45
4.013	0.2205	2.066391	6.016.31
0.000	0.2603	2.066849	0.3 19
0.10.7	0.3015	2.067890	· 5 356763
0.000	1.222100	7, 12 2003	· 6,500 3
0.1630	\$ 2. 2. 3. 4. 5	温。作を分析する	0.34 - 3*
e mil	4.43	2.18 Sept 3	· 位。《法》等"直
0.000	\$1,750.500	2.674700	· U ARPETA
0,00.2	$\triangle \in = \in_{cal} = 2$.063891	0.3-5-71
Oath Ar		3 在社会分享	O. mileta
	1. My 197	2,45,450	10.73 8.15
0.08.74	4 0 3 7 7	. 8,1007 744	4.01000

Table-13.. Refined values of specific volume and dielectric constant.

	Weight fraction	Sp.volume(c.c/gm)	D. C.	Sp.polarizat (c.c/gm.)	tion
the Proc	0.0	1.323345	2.063891	0.346440	(4.25)
growing and property of the second con-	0.0001	1.323279	2.064040	0.346459	Lus Penle
	0.0002	1.323216	2.064204	0.346482	
0,000	0.0003	1.323155	2.064419	0.346517	. 117 (A) . (
9,070	0.0004	1.323099	2.064640	0.346555	4 9 7 27 4
P. F. 103	0.0005	1.323045	2.064862	0.346594	100000
3.755	0.0006	1.322998	2.065079	0.346635	李/36人下进。
0.12 3	0.0007	1.322959	2.065321	0.346682	t . 7 7 .
0.000	0.0008	1.322923	2.065482	0.346711	
0.0007	0.0009	1.322891	2.065601	0.346732	18,27 18 22
CARL	0.0010	1.322862	2.065660	0.346739	50000000000000000000000000000000000000
0.0076	0.0011	1.322835	4 2.065692	0.346739	P. 200 44
\$ - C 10	0.0012	1.322810	2.065720	0.346739	55307.7
6. 4114	0.0013	1.322789	2.065739	0.346739	35×37×4
6.0013	0.0014	1.322771	2.065780	0.346744	
4,0:10		1.322754	2.065831	0.346751	14127.7
0.00.14	0.0016	1.322737	2.065880	0.346759	B 36 3, 9
0.9.15	0.0017	1.322719	[®] 2.065939	0.346768	53 al. 0
0.60%	0.0018	1.322700	2.066001	0,346778	2017
3. Ch 17	0.0019	1.322681	2.066100	0.346797	\$164. Y
0.000	0.0050	1.322661	2.066202	0.346815	Espera.
0.000	0.0021	1.322639	2.066300	0.346834	4393,4
0.6 7.0	0.0021	1.322615	2.066399	0.346851	5703,5
0,0003	0.0023	1.322592	9.066479	0.346864	46 42.7
D. Geral	0.0020	1.322567	2.066619	0.346891	\$ 37.5
	0.0021	1.322539	2.066740	0.346913	10 3.X
1.400	0.0025	\$5. ind	30.00	春春 名[2] 古。秦	さんない。ま

A CARLA

Table 4. Calculated Parameters from refined data.

21 711 - 4 5

					PS(c.c/mole)	$(c.c^2/mole^2)$	(c.c/mol
\$ 774	100				X	(coc /more)	(6.6/10)
0.0001		794629.03	39.83		32.04	659542.0	- 127140.
0.0002	or or was the	397295.60	40.74		33.40	691294.0	476755.
0.0003		264851.52	41.44	10.00	36.34	646237.0	1096485
0.0004		198630.23	42.53	9. 15.		701164.7	1189795
0.0005	De	158897.70	43.44	1,20	39.37	705505.7	1139296
0.0006	17	132410.04	44.74	1. 30	40.33	760033.6	1076493
0.0007		113490.94	46.43	1, 3 %	41.29	843237.7	103 1632
8000.0	É.	99301.93	48.81	有意と行う	41.19	874850.0	892724.
0.0009	19 44	88266.24	49.19	2. F 30	40.28	899433.0	713191.2
0.0010	1) 2	79437.86	50.46	1.	38.74	910357.9	519523.6
0.0011		72214.75	51.61	1,70	0.012	910628.0	354574.4
0.0012	0.	66195.60	52.68	1,517	35.79	931895.8	237642.2
0.0013	Ĉ.	61102.69	53.80	1 × 2 × 3	34.61	904319.8	147257.5
0.0014	E a	56737.44	54.86	1, 33	33.83	899855.8	92482.0
0.0015	60	52954.25	55.81	S = 1 (p)	33.25	890160.9	55602.0
0.0016		49643.97	56.65	1.200	32.77	876216.1	28297.0
0.0017	O.	46723.09	57.35	1.20	32.40	857368.7	9344.7
.0018		44126.75	57.96	2, 513	32.08	836643.2	5295.2
.0019	Û.	41803.69	58.50	1.2.47	32.09	815171.9	4598.4
0.0020	E.	39712.90	58.93	1. 27	32.08	791478.0	4765.5
		37821.17	59.28		32.07	767013.3	4916.7
0.0021	G.	36101.37	59.54		32.03	741522.1	6137.3
0.0022		34531.15	59.81	1,3%	31.87	718593.2	11395.3
.0023		33091.73	60.00		32.09	694926.4	3640.1
.0024		31767.39	60.11	. O. S	32.17	670609.6	953.0

APPENDIX- A. 5

System: Acetic Acid in Cyclohexane

Table 1. Experimental results of Specific volume.

(Weight fraction)X10 ²	volume, v (c.c/gm	cific volume $\triangle v(\text{obscal.}) \times 10^4$ (c.c/gm)
0.0	1.299916	3.0.270 . 0.0
0.01122	1.299790	- 0.63
0.02241	1.299673	- 1.18
0.03355	1.299536	1.01100-1.92
0.04465	1.299413	a. 4 75 2 - 2.54
0.05571	1.299307	2.98
0.06673	1.299195	- 3.48
0.07770	1.299080	2.94476 - 4.02
0.08864 6.2.37	1.298986	4.35
0.00953	1.298908	- 4.53
0.11039	1.298842	-4.58
0.12119	1.298775	4.65
0.14270	1.298680	2.00 × 2.00 - 4.40
0.16404	1.298589	- 4.12
0.18523	1.298525	- 3.58
0.20625	1.298460	- 3.05
0.22713	1.298401	- 2.48
0.24785	1.298347	- 1.86
0.26842	1.298290	- 1.29
C.28384	1.298235	- 0.70
0.30911	1.298192	0.0

v_{cal} 1.299916 + 0.5577XW

Table - 2. Experimental results of dielectric constant.

		10000 美元 美元 30000	(30, gH)	to Ce	
W (W	eight fra	ction)X10 ²		D. C.	0.7
£		a solected		6-8-56 3	0,173.0
	0.0	1,000		2.511213	A. 30 000 c
	0.0	2 . 3 . 48.73		2.0123	1 4 4 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
	0.0380	5-18-185		2.7 2.0127	01
t	0.0701	1. 《湖》。	. *	2.0130	09 - 0, 327633
Contraction	0.0984	The state of		2.0132	50
3.6		5.75 G.		Dall In ter	C. SEY 25
1111	0.1303	\$1.29 C.S.		2.0134	89
4	0.1572	t.1.20973.		2.0140	52
0.00	0.1884			2.0145	81
6.36		Section .		2.073	9, 707 117
	0.2287	6 1726.		2.0146	80
Balance	0.2889			2.0151	62
6. 7.	0.3199	4, 100		2.0156	
				2.0163	50
	0.3462				THE REPORT OF THE
0.100		1. 186.		A.816373	0,35913
Barrell Co		\$27.45 A		2.61 (4)	0.304136
September 2		2,75,073.		2.015.515	0,078
		1.5.8-42.	2.0	12330.	0, 201 14
0.1114		△E = € cal			
3.4.22		1.0124-2		2,0,273	4. * 47 * 5
P. 1500				3.51120	6,30+360
6.01		4,1,20		\$ 0 1 4 7 7 A	Q. 505 379
A DETERM		t, states		2.0:47:53	
6 ,				W. 01-0/2)	# 32° 12'
3.0-17		1 1 1 1 1 1 1			0. 368 × 9
ALCOY II		4. 7. 7. 7. 8. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1.			
0.0000				2.015)3	6. METER
		E. C.		ances 111 (1970) 전 관광이	when the statement of the statement of

Table-13., Refined values of specific volume and dielectric constant.

美术的人 化二氢化物铁 鎮土電池

	eight fraction	Specific volume	(cc/gm)	D. C.	Specific polarization (c.c/gm.)
0.0703	0.0	1.299916		2.012330	0.327974
#L6-07	0.0001	1.299800	12 31	2.012425	0.327968
	0.0002	1.299689	2 2 1	2.012513	0.327961
ch and a	0.0003	1.299578	24.24	2.012613	0.327958
	0.0004	1.299463		2.012725	0.327956
	0.0005	1.299360	17.00	2.012820	0.327953
Varience	0.0006	1.299256		2.012913	0.327949
Dated T	0.0007	1.299155		2.013013	0.327948
0,1988	0.0008	1.299057	49 g to 13	2.013093	0.327942
Charle Marin	0.0009	1.298972	17.76	2.013175	0.327941
(4,00 m	0.0010	1.298903	\$5.00	2.013255	0.327943
0,4001	0.0011	1.298842	374,37	2.013325	0.327945
0,141,8	0.0012	1.298786	医透水红素	2.013400	0.327949
0.04.13	0.0013	1.298731	10.07	2.013518	0.327963
Dark M	0.0014	1.298685	30 × 15	2.013675	0.327990
2, 11 13	0.0015	1.293644	25013	2.013875	0.328028
0,000	0.0016	1.298603	3.3 . 34	2.014100	0.328072
A.C.J.17	0.0017	1.298573		2.014300	0.328113
0.0019	0.0018	1.298540		2.014463	0.328144
kn (R) +1)	0.0019	1,298511		2.014588	0.328166
3. 7/03	0.0020	1.298482	e de	2.014650	0.328174
O. Olert	0.0021	1.298451	8 7 4 4 5 8 5 7 6 6	2.014675	0.328172
9.0000	0.0022	1.298422	73.34	2.014675	0.328165
0.00	0.0023	1.298395	24.61	2.014675	0.328158
	0.0024	1.298367		2.014700	0.328157
4,3715	0.0025	1.298341	24.58	2.014775	0.328169
1. 3. 5.	0.0026	1.298316		2.014862	0.328184
000	0.0027	1.298288	24.65	2.014950	0.328198
0.0.V9P	The state of the s	1.298261	Sacrate ST 18	2.015063	0.328218
and the second s	0.0028	1.298234		2.015175	0.328238
	0.0029	1.298209		2.015313	0.328265
	0.0030				

Table 4. . Calculated parameter from refined data

Wt. fraction V(c.c/mole) VS(c.c/mole²) PS(c.c/mole) Δ(V.VS) Δ(V.PS)

(c.c/mole²) (c.c/mole²) (c.c/mole²)

0.0001 780529.90 8.41 18.50 163901.3 1639113.0

		Addition of the second		(c.c/mole ²)	(c.c/mole ²
0.0001	780529.90	8.41	18.50	163991.3	1639113.0
0.0002	390231.62	9.91	16.99	667296.1	230236 .7
0.0003	260132.19	10.40	17.29	574892.8	231517.6
0.0004	195081.68	10.06	17.59	362853.0	232147.4
0.0005	156053.13	11.28	17.65	480643.7	195066.4
0.0006	130033.86	12.01	17.59	495429.0	154740.2
0.0007	111448.93	12.89	17.47	522695.5	119250.4
0.0008	97510.46	13.58	17.59	524606.3	116037.4
0.0009	86670.30	15.08	17.76	596291.7	117871.6
0.0010	77999.13	17.23	18.08	704332.2	131038.5
0.0011	70904.96	19.43	18.33	796262.7	136846.6
0.0012	64993.41	21.51	18.64	865062.3	145585.2
0.0013	59991.38	23.32	19.37	907069.6	178174.4
0.0014	55704.31	25.26	20.56	950315.5	231730.0
0.0015	51989.05	27.14	22.02	984672.6	292178.4
	48738.19	28.78	23.52	1003032.0	347015.9
	45370.18	30.63	24.75	1028868.0	383016.0
	43320.74	32.16	25.50	1037964.0	394218.7
	41039.78	33.66	25.89	1044872.0	389467.6
30 M	38986.92	35.01	25.82	1045239.0	367256.8
	37129.52	36.17	25.48	1038512.0	337136.0
	35441.02	37.29	25.02	1030979.0	305501.6
	33699.40	38.35		1022066.0	278314.0
	32486.22	39.30	24.27	1010321.0	255666.5
0.000	31186.15	40.23	24.38	998890.8	248865.4
	29986.11	41.11	24.55	986842.8	244386.8
0.0040	28874.89	41.86	24.66	971928.8	238506.5
0.002	27843.06	42.57	24.93	956966.0	237501.3

V= S+(U-x) Ky + x 1/2

of the Market

APPENDIX-B

135-0) = 2 Equilibrium Interaction of the Solute Species Entering into the Holes of a Solvent Body.

Theory of a Solvent Body.

with the discovery of interestitial compounds and various other observations along with the advancement of theories on the nature of solvents one is lead to suspect the presence of ordered structure of liquids having some unoccupied spaces which is referred to in this paper as "holes". It is possible that very minute quantity of solute species of tolerably small size temporarily occupy some of these "holes". The duration of such occupation may be attended with some distortions of both solvent "holes" and the solute species depending on the polarizability and structural differences of the components. One of the logical treatments that can be conceived on the basis of law of mass action (which is essentially statistical in nature) take the following form:

Species + holes - "species in hole"

MITTER FAKET

where bracket indicate the concentration in terms of number as well as "holes" per unit volume.

Testing of this "model of interaction" can be based on the assumption that species entering into the "holes" lose a part or whole of its

$$V = S + (1-\alpha) V_1 + \alpha V_2$$

$$VS = V_1 + \alpha (V_2 - V_2) = V_1 = \alpha D$$

$$K \neq BS - \alpha = \frac{\alpha}{1-\alpha} = \frac{V_1 - V_5}{V_5 - V_2}$$

$$K\beta - \frac{\lambda B}{V} (V_1 - V_5) = \frac{V_1 - V_5}{V_5 - V_2}$$

$$\beta - \frac{V_1 - V_5}{D \times V} = \frac{V_1 - V_5}{V_5 - V_2} + \frac{\lambda B}{V_5}$$

$$\beta - \frac{V_1 - V_5}{D \times V} = \frac{V_1 - V_5}{V_5 - V_2} + \frac{\lambda B}{V_5}$$

$$\beta - \frac{V_1 - V_5}{D \times V} = \frac{1}{V_5 - V_2} + \frac{\lambda B}{V_5}$$

$$\beta - \frac{V_1 - V_5}{D \times V} = \frac{1}{V_5 - V_2} + \frac{\lambda B}{V_5}$$

$$\beta - \frac{V_1 - V_5}{D \times V} = \frac{1}{V_5 - V_2} + \frac{\lambda B}{V_5}$$

$$\gamma_5 - V_2 + V_5 = \frac{1}{V_5} + \frac{V_1 - V_5}{V_5 - V_2}$$

$$\gamma_5 - V_2 + V_5 = \frac{1}{V_5} + \frac{\lambda B}{V_5} + \frac{\lambda B}{V_$$

12+601 = 1 (18-18) +17

15-14 = OVCH-15) volume, if calculated on the basis of simple additivity rule*. If V is expressed as the volume which contain one mole of $A_{f e}$ of which arphi fraction has entered into the "holes" and each c.c of solvent have only & moles of "holes" (i.e. 10²³x6.06 x \beta holes per c.c.) then

$$K = \frac{\alpha V}{(1-\alpha)(\beta S-\alpha)}$$
; whore S express the actual volume of

the solvent, initially added to make the mixture.

Now, the volume, V contain solvent S and free A, while the rest of A remain inside the "hole" without contributing its volume to the bulk, thus,

where, V₁ is the molar volume of A.

Hence we can write,

$$V = S = VS = (1-\alpha) V_1$$

or,
$$1-\alpha = \frac{v_s}{v_1}$$

or,
$$\angle = \frac{v_1 - v_S}{v_1}$$

the look the close of this plac gives

⁼ w1 + w2 ; where w is specific volume of the mixture, v4 and v2 are that of solute and solvent, and 1 = v1 + v2 .

Now, putting Z = V₁ - VS, we get,

$$\alpha = \frac{z}{v_1}$$
 and $\frac{\alpha}{1-\alpha} = \frac{z}{v_S}$.

Or,
$$\mathbf{K} = \frac{2 \mathbf{V} \mathbf{V}_1}{\mathbf{V} \mathbf{S} (\beta \mathbf{V}_1 \mathbf{S} - \mathbf{Z})}$$

$$\frac{K}{V_1} = \frac{(\beta V_1 S-Z)}{Z} = \frac{V_1 S-Z}{V_2 S} = \frac{V_2 S-Z}{V_2 S} = \frac{V_1 S-Z}{V_2 S} = \frac{V_2 S-Z}{V_2 S} = \frac{V_1 S-Z}{V_2 S} = \frac{V_2 S-Z}{V$$

or,
$$\frac{V}{VS} = \frac{K\beta s}{2} = \frac{K}{V_1}$$
 as Mark the state of the first section of the section o

Putting back Z = V1- VS, finally we have the relation.

the annual tending. The results of A and B and mires in The to Table

Therefore, a plot of $\frac{V}{VS}$ against $\frac{S}{V_1-VS}$ will give a straight line at the lower end of concentrations. The slope of this plot gives K/V and the intersection give K/V_1 . Holar volume of AcOH monomer known to be about 55.2 (average V_1 value in CCl₄ and Benzene), β and K can be found.

Results

In the present work of two systems, (i) acetic acid - Kerosene (HBH) and (ii) acetic acid - cyclohexane, acetic acid as the solute species have investigated for the "solute - hole equilibrium effect."

It must be noted that extremely accurate and precise measurement of density (or specific volume) is needed to investigate the emperical basis of such a construct. The measurement of specific volume have been carried out by magnetic float densimeter, as described in the main bedy of this thesis (page-43). V and S have been calculated by the relations described already (Appendix- A.4 and A.5). Molar volume of acetic acid monomer is about 55.2 c.c(obtained by indirect method shown previously). Now V/VS and S/V₁-VS for both the systems (HBH and cyclohexane) have been calculated and given in Table 7.1 and 7.2 respectively and the plots of V/VS versus S/V₁-VS are shown in Fig. 7.1 and 7.2 respectively. The results of K and B are given in Table 7.3.

These results were obtained by using a much bigger graph which is shown in the Fig. 7.1 and 7.2 in a smaller scale. The intersections are in the negative which rightly conform to the equation (7.1). The slepes and intersection so obtained were used to calculate B and K.

7 5 5 NS

Table 7-1

System: Acetic Acid in HBH

(Wt. fraction)X104	VS	V ₁ -VS
	1000000000	# TF 1.37
1	19950.52	51697.41
		#6 Te. 0 *
2	9751.98	27472.67
		Saul 5" , 359
3	6391.20	19244.92
	2.0	4 2 3 4 4 6
4	4670.36	15673.85
		発力力を支援す
5	3657.87	13508.02
		56.10.45
6	2959.54	12654.43
•	9 64. 4	0.13.60
7	2444.35	12935.52
•		2504,59
8	2077.01	13430.87
6	-7	2100.30

Table 7.2- | - 2

System: Acetic Acid in Cyclohexane

(Wt. fraction) x 10 ⁴	v Vs	v ₁ - vs
ergyeer 1	92809.74	16681.37
2	39377.56	8616.07
and the second s	24988.68	5807.59
4	19391.84	4321.48
\$	13834.50	3552.87
6	10827.13	3010.46
7	8646.15	2633.80
8	7180.45	2342.35
9	5747.37	2159.90
10	4526.94	2053.78

Results

Table 7.3 1 3

Solvent	K/V ₁	K (litre/mole)	β K	β (mole/litre)
HBH (High Boiling hydrocarbon)	1.8X10 ³	220.8	0.4151	0.00418
Cyclohexane	4.0x10 ³	99.36	5.0322	0.0227

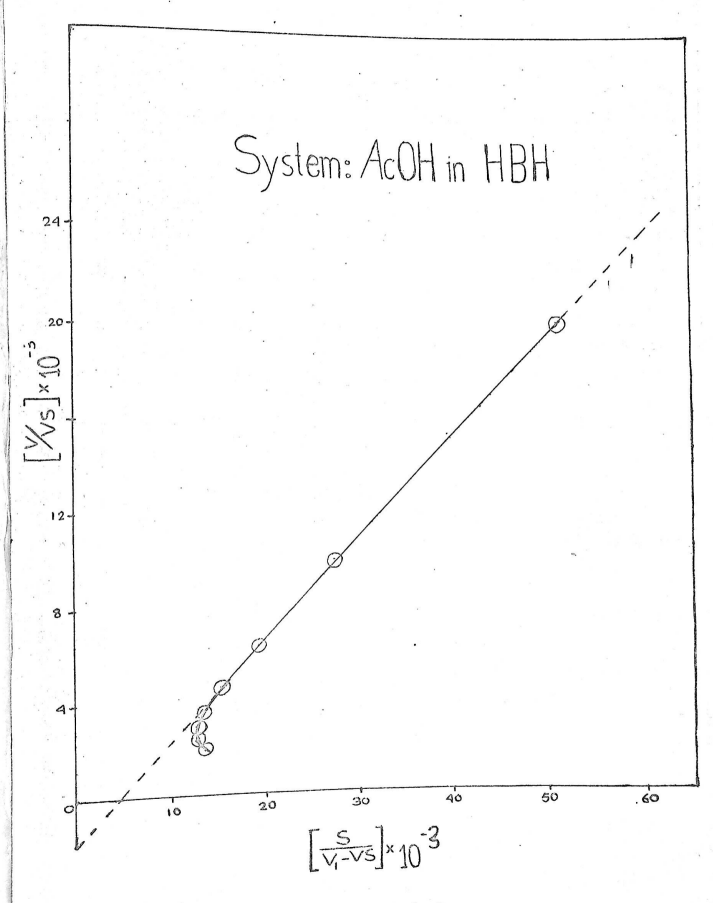


Fig. 7-11.1

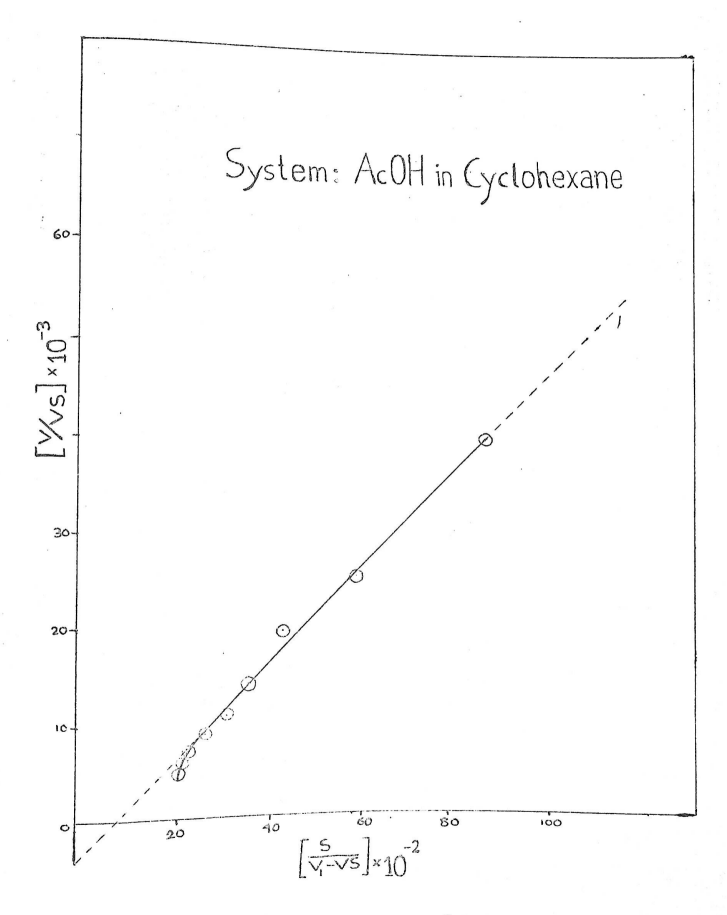


Fig. 7-2 1.2

Discussion

Fig. 7-1 and 7-2 show that the equation (7.1) very closely represent the variation of the parameters from near zero concentration upto about .0006 wt. fraction solution. The equilibrium constants are neither too high nor too low. Similarly the number of "holes" are within reasonable value. A sample calculation for cyclohexane system indicate that only one "hole" of the size of one molecule of acetic acid monomer exist per 450 molecules of cyclohexane. In HeH such calculation is not possible as the molecular weight is not known but the molecular "holes" exist only one - fifth times that of cyclohexane.

References 1. the state of the s

BIBLIOGRAPHY

- Same a financia de ser for sincipio de la Pare de Companya de la Pare de Companya de Compa
- in the contract of the contrac
- and the same was finding to the Property of the State of
- and work to be a second of the second of the
- er i de Charle (1865) i de Charl
- to. Quely mid Chemiora 2 d. Cross Physic, C. D.
- \$8. South on the Transfer I do to Them the to be the control of the total control of the control

BIBLIOGRAPHY . See., M. SAVE (1940).

100	John C. D. St. St. C. 1	A M. Then. Mich. 17, 72 (1203).
1.	Pauling, L	: "The Nature of the Chemical Bond", P-6, Third Edition, Oxford and IBH Publishing
	1 7 % tellings	Co., 1975 (Indian print - New Delhi).
2.	Mulliken and Pearson	: "Donor - Acceptor Complexes", Ann. Rev.
	unifor each Restord a	Phys. Chem. 13, 107 (1962); "Molecular Association". Volume I. Academic Press,
- L #	1 3 2 1 3 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	1975.
3. ,	Huggins, M. L. Markett	: Angew. Chem. internat. Edit., 10, 147 (1971), No. 3.
4.	Latimer and Rodebush	: J. Am. Chem. Soc., 42, 1419 (1920).
5.	Bline and Hadsi	"Hydrogen Bonding", (Pergamon, 1959); London Abstracted in Angew. Chem., 69,
1g	TRANSPORT OF A STATE OF	755 (1957).
6.	Barman and Rahim	: Acta Cryst., A34, 761 (1978).
7.	Schuster, Zundel and Sandorfy	: "The Hydrogen Bond", P-28, Vol. I, North Holland Publishing Company, 1976.
8.	Schuster, Zundel and Sandorfy	: "The Hydrogen Bond", P-81, Vol.1, North Holland Publishing Company, 1976.
9.	Pimentel and Mc Clellan	: "The Hydrogen Bond", P-51, Freeman and Co., 1960.
10.	Badger and Bauer	: J. Chem. Phys., 29, 1193 (1937).
11.	Albert and Badger	: J. Chem. Phys., 29, 1193 (1958).
12.	Badger, G. M	: J. Chem. Phys., 8, 288 (1940).
13.	Herman, R. C	: J. Chem. Phys., 8, 252 (1940).
14.	Fox and Martin	: Proc. Roy. Soc., (London) 162 A, 419 (1937).
15.	Gordy and Stanford	: J. Chem. Phys., 9, 204 (1941).
16.	Searles and Tamres	: J. Am. Chem. Soc., 73, 3704 (1951).

```
17.
     Tamres, M
                          : J. Am. Chem. Soc., 74, 3375 (1952).
     Barrow, G. M. et al. :
18.
                              J. Am. Chem. Soc., 75, 71 (1953).
19.
     Barrow. G. M
                              J. Phys. Chem., 59, 1129 (1955).
20.
     Fermer, Hardie and
     Thomson
                              J. Chem. Phys., 8, 3600 (1959).
21.
     Gordy and Stanford
                              J. Chem. Phys., 8, 170 (1940).
22.
     Josien, M. L
                              Compt. rend., 237, 175 (1953).
23.
     Josien and Sourisseau:
                             Compt. rend., 238, 2525 (1954).
24.
     Eyman and Drago
                              J. Am. Chem. Soc., 88, 1617 (1966).
25.
     Becker. Riddle and
                              J. Mol. Epect. P-211 (1958).
     Shoolery
     Takahashi and Li
                              J. Phys. Chem. 68, 2136 (1964).
26.
     Cresswell and Altred: J. Am. Chem. Soc., 68, 2136 (1964).
27.
                              Opt. Spekt., 16, 430 (1964).
     Bystrov
28.
                              J. Am. Chem. Soc., 58, 1617 (1966).
     Eyaman and Drago
29.
                              Ber. Beuseu. Phys. Chem., 17, 1104 (1967)
30.
     suhr
                              Trans. Farad Soc., 63, 1063 (1967).
     socrates
31.
     Coulson, C. A. and
                              Arkiv Fysik 8, 205 (1955).
32.
     Danielson, U
                              Bull. Chem. Soc., Japan, 27, 445 (1954).
     Tsubomura, M
33.
     Kollman. P. A. and
                              Chem. Rev. 72, 283 (1972).
34.
     Allen. L. C
     Pimentel and
                              "The Hydrogen Bond", P-39, Freemant Co.,
35.
     Mc Clellan
                              1960.
                             J. Chim. Phys., 63, 239 (1966).
     Vinogradov
36.
                             J. Chem. Soc., (B), 80 (1967).
     Ibbitson and Moore
```

37.

me y y

- 38. Beckman, E : J. Phys. Chem., 6, 444 (1890)
- 39. Allen, G. and Caldin, E. F Quart. Rev. 7, 255 (1953) and references therein.

* 19do to may

- 40. Karle, J. and Brockway, L. 0 : J. Am. Chem. Soc., 66, 574 (1944).
- 41. Taylor, M. D : J. Am. Chem. Soc., 73, 315 (1951).
- 42. Harris, J. T. Jr. and
 Hobbs, M. E : J. Am. Chem. Soc., 76, 1419 (1954).
- 43. Barrow, G. M. and Yorger, E. A : J. Am. Chem. Soc., <u>76</u>, 5248 (1954).
- 44. Wenegred, J. and Sprurr, R. A : J. Ap. Chem. Soc., 79, 9844 (1957).
- 45. Fredman, E : J. Chem. Phys., 21, 1784 (1953).
- 46. Le Fevre, R. J. W. : J. Chem. Soc., 1795 (1938).
- 47. Coburn, W. C. Jr. and J. Am. Chem. Soc., 80, 1318 (1958).
 Grunwald, E
- 48. De. Tar, DeLos. : J. Am. Chem. Soc., 92, (5), 1361 (1970).
 Novak, R. W
- 49. Mathows, D. M. and Sheets, R. W : J. Chem. Soc., (4), 2203 (1969).
- 50. Goldman, M. A. and : J. Phys. Chem., 77, (19), 2295 (1973). pmerson, M. T
- 51. Jenkins, J. O. and : J. Chem. Soc., (B), 1538 (1970).

 Smith, J. W
- 52. Liszi, Jamos : Acta Chem. 67(1), 5-14, (1971) (Cf. Chem. Abs. 103756h), 74 (1971).
- 53. Corsaro, Robert, D., : J. Chem. Phys. 54 (9), 4090 (1971). et al.
- 54. Arnold, J. T. and J. Chem. Phys., 19, 1608 (1951).

- 55. Ruggins, Pimentel, G.C. : J. Phys. Chem., 60, 1311 (1956). and Shoolery, J.N.
- 56. Reeves, L. W. and Schneider, W. G : Trans. Farad. Soc., <u>54</u>, 314 (1958).
- 57. Reeves, L. W : Trans. Farad. Soc., 55, 1684 (1959).
- 58. Davis, J. C. Jr. and Pitzer, K. S : J. Phys. Chem., 64, 886 (1960).
- 59. Jones, R. E. and . Acta cryst., 11, 484 (1958).
- 60. Lascombe, J. et al. : J. Chem. Phys., 59, 1233 (1962) 23
- 61. Bellamy, L. et al. : Spectro Chim. Acta., 19, 443 (1963)
- 62. Hauric, M. et al. : C. R. Acad. Sci. Rev. C, 264, 694(1967).
- 63. Ritter, N. L. and
 Simons, J. N : J. Am. Chem. Soc., 67, 757 (1945) and
 references therein.
- 64. Sharpe, A. N. and : J. Chem. Soc., 2974 (1961).
- 65. Sharpe, A. N. and : J. Chem. Soc., 157 (1962). Walker, S
- 66. Sharpe, A. N. and : J. Chem. Soc., 2340 (1964). walker, S
- 67. Kulevsky, N : "Molecular Association", P-144, Volume 1, Academic Press, 1975.
- 68. Vinogradov : "Hydrogen Bonding", P-14, Van Nostrand Reinhold Company, 1971.
- 69. Johari and Dannhauser : J. Phys. Chem., 72, 2373 (1968).
- 70. Cantacuzene : J. Bull. Soc. Chim., Fr., No. 4, 747 (1962); (Cf. Donor Acceptor Bond, E.N. Gur'yanova, et al. John Wiley & sons, New York, Toronto, 1975).

71. Rahim, Z

Private Communication (Associate Professor of Chemistry, Rajshahi University).

72. Glasstone, s

P-531. MacMillan and Co. Ltd., 1969.

Band the new way before the two controls of the control of the con

The form of the second

73. Smyth

"Dielectric Behaviour and Structure", P-401, (Mc. Graw).

74. Smyth

- : J. Am. Chem. Soc., 51, 1734 (1949).
- 75. Baur, M.E., Horsma, D.A. Knobler, C.M. and Perez, P
- J. Phys. Chem., 73, 641 (1969).
- 76. Baur, M.E. Knobler, C.M. Horsma, D.A. and Perez.P
- J. Phys. Chem., 74, 4594 (1970).

Bally thy . was a to be to because

- 77. Voronkov, M.G. and Deich,
 A.Y. Teor. i Eksperim.
 Khim.
- Akad. Nauk Ukr. SSr (Eng. Trans.), 1, 443 (1965).

from the Parterage State State Mala a March (

- 78. Voronkov, M.G. Deich, A.Y. and Akatova, E.V. Khim. Geterotsikl. Soedin.
- Akad. Nauk Lat. SSR (Eng. Trans.), 2, 5 (1966).

79. Kulovesky, N

- "Molecular Association", P-115, Vol. 1, Academic Press, 1975.
- 80. Guryanova, E. N. et al. :
- "Donor-Acceptor Bond", P-59, John Wiley & Sons, New York. Toronto, 1975.

- 81. Rahim, Z
- Private Communication (Associate Professor of Chemistry, Rajshahi University).
- Barman and Rahim
- Rev. Sci. Instrument, 48, 1695 (1977).
- 83. Lamb. A. B. and Lee, R. E
- J. Am. Chem. Soc., 35, 1666 (1913).
- 84. Pohl. Hobbs and Gross
- : J. Chem. Phys., 9, 408 (1941).
- 85. Buckingham, A.D. and Raab, R. E
- Trans. Faraday Soc., 55, 377 (1959).

- 162 -

86.	Nagai and Simamura		Bull. Chem. Soc., Japan, 35, 132(1962).
87.	Satchel, D. P. N. and Wardell, J. L	:	Trans. Faraday Soc., 61, 1199 (1965).
88.	Dobye, P	:	Polar Molecules, Page-54, New York, 1929
89.	Muller, F. H.	:	Phys. Z., 34, 689 (1933).
90.	Weigle, J.	:	Helv. Phys. Acta, 6, 68 (1933).
91.	Smith, J. W.	:	"Electric Dipole Moments", Page-127-167 London Butterworths Scientific Publica- tions, 1955.
92.	Müller, F. H.	:	Trans. Faraday Soc., 30, 729 (1934).
93.	Müller, F. H.	:	Phys. Z., 35, 346 (1934).

N. B.: References 64-66 and 75-79 were cited in "Molecular Association", volume 1, Edited by R. Foster. Academic Press, 1975; References 80 was obtained from "Donor-Acceptor Bond", John Wiley & Sons, New York. Toronto, 1975 and References 88-90 and 92, 93 were obtained from "Electric Dipole Moments", by Smith, London Butter worths Scientific Publications, 1955.