	Original Research Article
Ultraso	onic Characterization of aqueous Polyvinyl Pyrrolidone (PVP)
ABSTRACT	
	c parameters such as adiabatic compressibility (β_a), relaxation (z), classical absorption ($\frac{\pi}{f^2}$), free length (L _f), isothermal
	expansivity (a) and Moelwyn-Hughes parameter (C1) of PVP the measurement of ultrasonic velocity (u) , density(p) and
Methodology: All these mea concentration of the range 0.0 ultrasonic velocity (u) was obse	surements were made by Pulse Echo technique at different 05 to 0.3wt.% and at 288K, 293K, 298K, 303K and 308K. The erved to vary linearly with concentration of PVP solution. ingle phase formation and the compatibility of aqueous PVP.
The compatibility is due to the and free length decreases line	linear chain structure of the polymer. Adiabatic compressibility early with increase in concentration of aqueous PVP solution eractions, association and non-dominance of PVP in water,
acoustic impedance (z) of aquinteraction between the polym	change in the variation at any concentration. The increase in ueous PVP solution indicates that there is a strong molecular her and the solvent molecules and the structural arrangement is in the structure of solvent or solution is a result of hydrogen

bond formation and structure making property of PVP in solution. **Conclusion:** Due to linear nature of ultrasonic parameters like ultrasonic velocity, adiabatic compressibility, etc. in PVP solution indicates that this PVP water soluble polymer can be

10

11 Keywords: Pulse echo, Polyvinyl Pyrrolidone (PVP), ultrasonic velocity, adiabatic 12 compressibility, other thermo acoustical parameters, etc.

13 14

1. INTRODUCTN

used as tablet binder in pharmaceutical industries.

15 Thermo-acoustic properties of water soluble polymer are of considerable scientific and technological interest ^[1]. These are important for understanding solute-solvent and solute-16 17 solute interactions. Knowledge of volumetric properties of polymer solutions has been 18 proven to be a very useful tool in evaluating the structural interactions occurring in the 19 solution. Many research groups have focused their attention to obtain information on the 20 thermodynamics of aqueous polyethylene glycol (PEG), which is often used in biotechnology. In this respect, thermodynamic properties such as vapors + liquid equilibria, density and viscosity have been obtained ^[1]. However, similar information on the 21 22 thermodynamics of aqueous solution of PVP, which is also a very important system in 23 respect to its use in biotechnology, in scarce. In the present study an attempt has been 24 made to obtain information on hydration² of PVP by measuring various thermo acoustic 25 26 properties such as ultrasonic velocity (u), density (p), viscosity (n), adiabatic compressibility (β_a) , relaxation time(τ), acoustic impedance (z), classical absorption ($\frac{\infty}{f_z}$), free length (L_f), 27

volume expansivity (α) and parameter (C₁) of aqueous PVP solution at different temperature, 28 29 concentration & frequencies (1MHz, 2MHz & 5MHz). 30

2. MATERIALS AND METHODS 31

32 Ultrasonic velocity, density and viscosity values are estimated in the aqueous PVP solution. The solution was prepared by adding a known weight (1gm) of PVP to a fixed volume (100 33 ml) of distilled water and then stirring until clear solution were obtain. The concentration 34 range chosen in the solution are 0, 0.05, 0.1, 0.15, 0.2, 0.25, & 0.3 wt%. Ultrasonic velocity 35 measurements are carried out using ultrasonic pulse echo technique using MHF-400 pulser 36 37 receiver and Tektronix- 2 Channel Digital Storage Oscilloscope from Roop Telsonic 38 Ultrasonix Ltd. Mumbai- India at different frequencies of 1 MHz, 2MHz & 5MHz and at 39 temperature range 288K-308K with an accuracy of ±0.1m/sec. Density has been measured 40 by pycnometer method. The viscosity of liquid was measured by Oswald's viscometer. Temperature is maintained at a fixed range by Plasto Crafts Thermostat and other related 41 parameters calculated by standard formulae given below [6-8]: 42

 $\beta_a = \frac{1}{u^2 \rho}$

 $\tau = \frac{4}{3}\eta \cdot \beta_{\alpha}$

 $\alpha = -\frac{1}{\rho} \left(\frac{\partial \rho}{\partial T} \right)$ Where, *T*- temperature

2.1 Formulae 43

44	1.	Density	$ \rho = \frac{M_l}{M_W} \rho_W $
45	2.	Viscosity	$\eta = \frac{\rho_i t_i}{\rho_w t_w} \eta_w$

- 46 3. Adiabatic compressibility
- 47 4. **Relaxation Time**
- $Z = \rho . u$ 48 Acoustic Impedance 5.

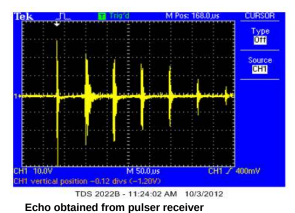
49 6. Classical Absorption
$$(\alpha/f^2)class = \frac{8\pi^2 \eta}{3\rho u^5}$$

50 7. Free Length $L_f = K_j \beta_a^{1/2}$ Where, K_j - Jacobson's constant

- 50 Free Length 7.
- 51 8. Volume Expansivity
- Moelwyn-Hughes parameter $C_1 = [\frac{13}{3} + (\alpha T)^{-1} + \frac{4}{3}\alpha T]$ 52 9.
- 53 54

55

Echo Pattern



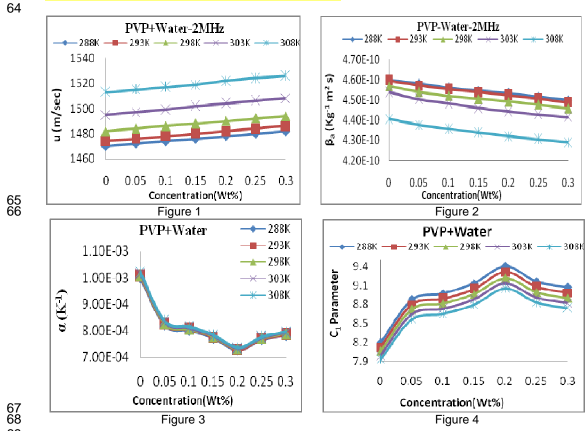
<u>56</u> 58

59

60

61 3. RESULTS AND DISCUSSION

- 62 Graphical and tabular representation of various thermo-acoustic parameters such as (u), (ρ),
- 63 $(\eta),(\beta_a),(\tau),(z),(\frac{\pi}{z_0}),(L_f),(\alpha)$ and (C_1) shown below:



69

70 Table for thermo-acoustic parameters

Parameters/C	0.00	0.05	0.1	0.15	0.2	0.25	0.3
▼onc. (wt%)	◆						
Density(p)							
Kg.m ⁻³	996.798	999.935	1001.86	1002.26	1002.77	1003.48	1004.99
Viscosity(η)							
Pas Sec	8.93E-04	9.02E-04	9.04E-04	9.11E-04	9.12E-04	9.19E-04	9.34E-04
Acoustic							
Impedance (Z)							
<mark>Kg.m⁻²sec</mark>	1.48E+06	1.48E+06	1.49E+06	1.49E+06	1.49E+06	1.50E+06	1.50E+06
Relaxation							
Time(т) <mark>sec</mark>	5.44E-13	5.46E-13	5.45E-13	5.47E-13	5.46E-13	5.49E-13	5.55E-13
Free Length							
(L _f) <mark>m</mark>	4.18E-11	4.17E-11	4.16E-11	4.15E-11	4.14E-11	4.13E-11	4.12E-11
$(\alpha/f^2)_{class}$							
Sec ² m ⁻¹	7.24E-15	7.26E-15	7.23E-15	7.25E-15	7.23E-15	7.25E-15	7.33E-15
Volume							
Expansivity(α)							
K ⁻¹	1.01E-03	8.28E-04	8.07E-04	7.78E-04	7.31E-04	7.71E-04	7.89E-04
Parameter							
(C ₁)	8.06	8.72	8.81	8.96	9.21	8.99	8.9

72

Figure 1 shows variation of ultrasonic velocity with concentration (wt.%) at 288K to 308K. It is observed that the ultrasonic velocity varies linearly with increase in concentration of PVP in solution. It shows molecular association in PVP solution. It also gives the evidence to enhance the miscibility ^[1] among the molecules presence in the PVP solution. The hydration state and its dynamic behavior are also explained in the light of formation of inter- or intramolecular hydrogen bonds between adjacent OH groups ^[2-3].

Figure 2 shows variation of adiabatic compressibility with concentration (wt.%) at 288K-308K and at 2MHz. It gives exactly opposite behavior that of ultrasonic velocity, it shows molecular association in PVP solution ^[4]. This behavior shows strong association at higher concentration of PVP in water.

Table shows Acoustic impedance (Z) increases with increase in concentration (wt.%) of PVP solution. The increase in acoustic impedance (Z) indicates that there is a strong molecular interaction between the polymer and the solvent molecules and the structural arrangement may be affected ^[5]. This change in the structure of solvent or solution is a result of hydrogen bond formation and structure making property of PVP in solution.

88 Variation of density and viscosity with concentration (wt.%) is as shown in table. It 89 increases with increase in concentration of polyvinyl alcohol solution. Increase in density is 90 due to the fact that the number of PVP chains added to the solution increases with 91 increasing PVP concentration (wt.%). Polymers are characterized by the large molecular 92 weight compared to the solvents. This also contributes to increase in the density of the 93 solution. The value of density and viscosity increases with increase in concentration of PVP 94 solution. This change in the structure of solvent or solution is a result of hydrogen bond 95 formation and structure making property of PVP in solution ^[6].

Linear variation of free length with concentration (wt.%) in PVP solution shown in above table. It decreases linearly with increase in concentration of PVP in solution indicates that the strong interactions, association and non-dominance of PVP in solvent, because there is no abrupt change in the variation at any concentration (wt%).

Table shows relaxation time and classical absorption increases with increase in concentration (wt.%) of <u>PVP</u> in solution. It indicates that the solution is highly ordered due to outstanding hydration and such solution generally absorbs more ultrasonic energy ^[6-7].

The variation of density with temperature gives volume expansivity. Figure 3 and 4 shows variation of volume expansivity and Moelwyn-Hughes parameter. It exhibits exactly opposite behavior shows strong association between PVP and water molecules due to the formation of inter or intra-molecular hydrogen bonds between adjacent OH groups^[7]. It means that hydrogen-bond attractions can occur between molecules (intermolecular) or within different parts of single molecule (intramolecular)⁹.

109 4. CONCLUSIONS

110	I.	The linear variation of ultrasonic velocity with concentration gives the evidence to
111		enhance the compatibility among the molecules presence in the PVP solution.

112 II. The linear decrease in adiabatic compressibility and free length with increase in concentration of PVP solution indicates the strong interactions, association and non-dominance of PVP in water.

115 III. The hydration state and its dynamic behavior are also explained in the light of
 116 formation of inter or intra-molecular hydrogen bonds between adjacent OH groups.

- 117 IV. The linearity in all the parameters indicates there is a single phase formation. It
 118 predicts strong polymer-solvent interaction and strong association in the PVP
 119 solution.
- V. This water soluble polymer <u>PVP</u> is used in pharmaceutical industries for tablet
 binder.
- 122
- 123
- 124

125 **REFERENCE**

- 126 [1] Rahmat Sadeghi, Mohammed Taghi Zafarani- Moattar; J. Chem. Thermodynamics 36 (2004)
- 128 [2] Mohammed Taghi Zafarani-Moattar, Zh. Khoshsima; J. Chem. Thermodynamics 40 (2008) 1569-1574
- 130 [3] J. Jayaraju, J. Keshavayya, S.K. Rai, K.C. BASAVARAJU; J. of Macromolelular 131 Science[®], Part A: Pure and Applied Chemistry (2008) 45, 271-275.
- 132 [4] J. Jayaraju, J. Keshavayya, S.K. Rai, Shetty Ravindra Rama; J. of Macromolelular 133 Science, Part B, 47:2, (2008) 296-304.
- 134 [5] A. AL-KATHANI AHMED, J. JAYARAJU, B.S. SHERIGARA, H.S. NAIK BHOJYA; Journal
- 135 of Macromolecular Science, Part A: Pure and applied chemistry (2008) 45, 1055-1063
- [6] Priyanka Tabhane, Omprakash P. Chimankar, Chandragupt M. Dudhe and Vilas A.
 Tabhane; Pelagia Research Library; Der Chemica Sinica, 2012, 3(4): 944-947
- 138 [7] Priyanka Tabhane, O.P. Chimankar and V.A. Tabhane; J. of chemical and 139 Pharmaceutical Research, 2012, 4(6); 3051-3056
- 140 [8] Priyanka Tabhane, O.P. Chimankar and V.A. Tabhane; J. of Modern Physics Letters B, 2012, vol. 26, No. 27.
- 142 [9] IUPAC, Compendium of Chemical Terminology, revised version of 2nd ed. (the "Gold 143 Book") (2006).