Analysis of the Thermodynamic Properties of a Binary Mixture of Pharmaceutical Residual Solvents

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Abstract

Deep examination on thermodynamic characteristics (Enthalpy, Entropy, Free energy) across the complete composition range from 298.15 K to 323.15 K at atmospheric pressure was used to study the molecular interaction of pharmaceutical residual solvents mixture (2-Butanol with m-Xylene). The solvation process, or the removal of molecules from their natural environment, is used to make sense of the data and provide context for the results. Then, the Redlich-Kister polynomial equation was used to fit the computed excess thermodynamic characteristics. The molecular interactions between 2-Butanol and m-Xylene are addressed in light of the computed excess characteristics.

Keywords: Residual solvent; Thermodynamic property; Binary mixture; 2-Butanol; m-Xylene

1. Introduction

In terms of output, the pharmaceutical sector is a major consumer of organic solvents[1-3].Properties of binary liquid mixtures are of interest to pharmacists, medical scientists, condensed-matter theorists, experimental chemists, physicists, etc. since organic solvents are present throughout the pharmaceutical manufacturing process. Understanding the intermolecular forces responsible for the many phenomena seen in experiments is crucial for both fundamental and applied research. The physical, chemical, and transport characteristics of fluids are of interest since they are the most common form in which materials are handled in chemical process industries. Understanding the intermolecular interaction between the component molecules and processing product formation are both greatly aided by thermodynamic analysis of liquid mixtures consisting of polar and nonpolar components for a variety of industrial and technical applications. Testing for residual solvents has evolved into an integral aspect of drug quality assurance [4, 5].

Organic volatile molecules utilized or created in the manufacture of drug ingredients or excipients, or in the production of drug products, are referred to as residual solvents in the pharmaceutical industry. In terms of pharmacological significance, 2-butanol (or 2-Bu-OH) and xylene stand out among the residual solvents[6,7].2- Bu-OHis a polar chemical molecule that often appears as a racemic mixture of the two stereoisomers, (R)-()-2- Bu-OH and $(S)-(+)$ -2-Bu-OH. This secondary alcohol is flammable, odorless, and colorless; it dissolves in water and is totally miscible with polar organic solvents like ethers and other alcohols [8, 9]. As a solvent, in the production of chemicals and pharmaceuticals, and in the field of medicine and dentistry [10, 11], xylene has several applications. The most prevalent form of xylene, known as m- xylene (or m-Xln), is a benzene derivative with two methyl groups substituted for the carbon atoms.where "m" stands for "formeta," designating positions 1 and 3 for the two methyl substituents on the aromatic ring.

Professor & Pricipal¹, Associate Professor^{2,3} Dept. of Pharmaceutics, Pharmacology Mother Theresa Institute of Pharmaceutical Education and Research, Kurnool, Andhra Pradesh Due to their widespread use, studies analyzing the nature of molecular interactions and exploring the physicochemical characteristics of such systems [12, 13, 14] have acquired a lot of traction in recent years. A thorough literature search revealed the lack of any comprehensive research focusing specifically on the thermodynamic parameters of the binary combination of 2-Bu-OH and m-Xln. In light of this relevance, it is of interest to research the thermodynamic characteristics in order to understand the interaction behavior in their binary mixes of variousproportionswiththevariationoftemperature.

Thermodynamic characteristics and their excess thermodynamic properties have been computed from experimental measurements of density and viscosity. 298.15K to323.15 K. The Redlich-Kister equation [15] was used to get the values for the excess characteristics, and the resulting enthalpy, entropy, and free energy of activation were then shown graphically.

2. Experimental details

2-Bu-OH and m-Xln used in this research study were purchased from Sigma Aldrich Chemicals Company. According to the manufacturer, the purities of these compounds were >99%. The water used in all experimental work was double distilled in quality. The binary mixtures of 2-Bu-OH and m-Xln were prepared by using an analytical electrical balancewith a precision of \pm o.1 µg and later were converted to different composition of the mixture using dilution method. Special care was taken to prevent evaporation and the introduction of moisture into the experimental samples.

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Densityofallbinarymixturesincludingpuresolvents was measured using an oscillation densimeter (Anton Paar DSA 5000). To measure the viscosity Stabinger viscometer (svm-3000-stabinger-viscometer) was used. The temperature was previously set up by 298.15-323.15 K. The measurement was accomplishedbythedescendingofthetemperaturein theviscometer.Inbothmachineryprocessesthe temperaturewasautomaticallycontrolledwithan uncertainty of ±0.01 K.

Theplotof

$$
\ln\!\!\!\!\frac{(\eta V_m)}{hN)}
$$

vs.**1/T**ofeq.(3)givesa

3. Theoretical aspects

Calculation of different thermodynamic parameters for viscous flow:

Fluidity of liquid follows some mechanisms as liquid is considered as a combination of layers and so it flows asarateprocess.To treattheviscosityasarate process it is assumed that the motion of liquid layers involves the passage of a molecule from one equilibrium position to another in the same layer. To do so, it is necessary that a suitable "hole" or site shouldbeavailable,andtheproductionofsucha

"hole"requirestheexpenditureofenergy,sincework must be done in pushing back other molecules. The jump of the moving molecules from one equilibrium position to the next may thus be regarded as equivalent to the passage of the system over a potential barrier. Eyring and his co-workers using absolute reaction rate theory and partition functions, correlated viscosity [16] as follows:

straight line with slope, $\Delta H^{#}/R$ and intercept, $-\Delta S^{#}/R$ assuming that $\Delta H^{#}$ and $\Delta S^{#}$ be almost independent of temperature. Therefore, $\Delta H^{\#}$ and $\Delta S^{\#}$ can easily be calculated from the slope and intercept of eq. (3) as,

∆H#=SlopeR....................................... (4)

∆S#=InterceptR(5)

The free energy of activation, ΔG^* , for viscous flow has been calculated by using the simple thermodynamic relation,

∆G#=∆H#−T∆S#....................................(6)

Calculationofdifferentexcessthermodynamic parameters for viscous flow

The excess free energy of activation ($\Delta G^{#E}$), excess enthalpy of activation ($\Delta H^{#E}$) and excess entropy of activation $(\Delta S^{#E})$ for viscous flow were calculated by using the following relations,

id id id 1 2 ∆G#E=∆G#−∆G#..................................................................(7) or,∆G#E=∆G#−(X1∆G#+X2∆G#)(8) $\eta = \zeta^{\text{hN}}$ Vm).Exp(∆G#)/RT(1) ∆H#E=∆H#−∆H#................................................................. \cdots (9) or,∆H#E=∆H#−(X∆H#+X∆H#)........................(10) Where, ΔG^{\dagger} isthefreeenergyofactivationpermole forviscousflow,hthePlanck"sconstant(=6.6262x $And\Delta S^{\#E}=\Delta S^{\#}-\Delta S^{\#}$ (11) -34 23 or, $\Delta S^{#E} = \Delta S^{#} - (X_1 \Delta S^{#} + X_2 \Delta S^{#})$ (12) 2

10 J.sec),NtheAvogadronumber(=6.023x10 1

mol⁻¹), R the molar gas constant (= 8.3145 JK⁻¹mol⁻¹) T the absolute temperature in Kelvin scale and η is the observed

viscosity in mPa.s. According to the definition of $\Delta G^{\#}$ eq. (1) reduces to

 $\eta = (hN/V_m) . exp(-T\Delta S^*) / RT$(2)

Where, the subscripts 1 and 2 represent the pure components of the mixture.

The experimentally obtained values of excess properties for a systemwere fitted by the least square method using

Redlich-Kister Eq. (13) of the form:

or,
$$
\ln(\sqrt[n]{\ln m}) = -
$$

\n $\ln^{\Delta S^{\#}} + \Delta H^{\#}/RT$(3)
\n $(Y)^{E} = x_1 x_2 \sum_{i \ge 0} (1 - 2x_1) B_{i}$ (13)

Where, ΔH^* is the enthalpy of activation per moleand ΔS^* the entropy of activation per mole for viscous flow.

4. Resultsanddiscussion

Density and Viscosity of the binary mixture The density and viscosity values of binary mixture as afunctionofthemolefractionof2-Bu-OH(x₁)respectivelyattemperatures(298.15to323.15)Kcan

be known by Figure (1-2). The plotting displays the trend of changing these properties. The changes ofthe properties are evaluated for the changes of each proportion of the mixture component along with the variation of temperatures.

*x***1 (molefractionof 2-Bu-OH)**

Figure1:Densitytrendanalysisofthebinarymixture

Figure2:Viscositytrendanalysisofthebinarymixture

FromFigure2 itcan be observed that viscosityof the binary mixture of 2-Bu-OH + m-Xln increases invery disciplined way with the increasing proportionof 2-Bu-OH but decreases almost linearly with temperature rises. Here the nonpolar m-Xln is low viscous liquid and when it interacts with the polarone it breaks the intermolecular network of the 2-Bu- OH which was built due to a lot of H-bond as well as polar-polar interaction. Thus, there was an induction of polarity occurred in m-Xln and become induced dipole and attached to the polar 2-Bu-OH. Thus, the liquid mixture becomes more viscous. From Figure 2 it can be seen that the rising of viscosity first increases slowly but when the polar proportion is higher it rises sharply upward

Thermodynamicpropertiesmeasurement

The three types of very important thermodynamic propertiesviz.entropy,enthalpyandfreeenergy

were very intensively studied for the binary mixture of polar + nonpolar compound. There excess values were also calculated and fitted in Redlich-kisterfitting equation. Figure 3 shows the plot of enthalpy of activation ΔH^* for the transport process vs. mole fraction of alcohol for the system over the whole rangeofcomposition(0to1).Itisnoticedthatforthe binary system ΔH^* value (Table 1) continues to increase on addition of alcohol and eventually reach the value of pure 2-Bu-OH because H-bondedliquids, like 2-Bu-OH always require higher enthalpy for activation than that of the other low polar or nonpolar liquids. The excess enthalpies of the binary mixturesofaromatichydrocarbonswith2-Bu-OHare graphically represented by the Figure 4. The excess enthalpy of activation for viscous flow (ΔH^{HE}) show the negative values extended over a considerable region of concentrations.

Table1:ExperimentaldataforEnthalpyofactivationofthebinarymixture

Enthalpy, Excess Enthalpy and Fitting value of Excess enthalpy of activation for different molar ratios are listed in the table.

Figure3:Enthalpycurvefordifferentmolefractions

Figure4:Excessenthalpycurvefordifferentmolefractions

Figure 5 is for the plots of entropy of activation ΔS^* for the viscous flow vs. mole fraction of alcohol for this binary system. The corresponding data are listed in Table 2. The typical nature of the small and negative $(\Delta S^{\#})$ value of alcohol indicates that during the viscous flow of initial H-bonded order breaks down to form the activated complex of little ordered structure. On the other hand, negative $(\Delta S^{\#})$ value for the aromatic hydrocarbons revealed that during the flowprocesstheactivatedcomplex formed,aremuch more ordered.

Figure 5 also reveals that with increase of concentrationofalcohol,thenegativevaluedecreases for the binary system of 2-Bu-OH + m-Xln respectively. It indicates that during the flow process at the certain composition of mixtures, the molecular orderoftheactivatedandinactivatedstateforeachof the mixtures is same. The excess entropy ΔS^{*E} of activation for the system are shown (Figure 6) as a function of mole fraction of 2-Bu-OH. The plots of ΔS^{*E} vs. mole fraction of 2-Bu-OH Figure 6 for the binary mixture show an almost similar trend at those of the excess enthalpy ($\Delta H^{#E}$). The curve has been found to be concave in nature. The negative excess entropy signifies that the species formed in the

Table2:ExperimentaldataforEntropyofactivationofthebinarymixture

Entropy, Excess entropy and Fitting value of excess entropy of activation for different molar ratios are listed in the table.

Figure5:Entropycurvefordifferentmolefractions

Figure6:Excessentropycurvefordifferentmolefractions

Table (3-8) shows the variation of free energy of activation ΔG^* fortheviscousflow of the systemasa function of 2-Bu-OH under the whole range of composition at temperature 298.15 K to 323.15 Kandtrendofchangingthisthermodynamicproperty

negative, but with the rise of temperature, the values are less negative, i.e., $(\delta \Delta G^{*E}/\Delta T)_{p}$ is positive. The general natureof thecurvesdoesnotvirtuallychange with the variation of temperature. The negativeexcessfreeenergiesindicateaccordingtotheEyring

aregivenbytheFigure7.The∆G[#]valuesincrease very slowly intheinitialstagewhichisfollowedbya

 $\ln(\eta V_m)$ ΔH^{\neq}

\overline{a} ΔS^2

thattheviscousflow

 \models $____\$ $__\$

relatively greater rise with increasing concentrationof 2-Bu-OH. The curves for ΔG^* for the system are found to be smooth and similar. But a crossover of curves between temperature 298.15 K and 323.15 Kis noticed in the system at higher mole fraction of 2- Bu-OH. It is also revealed from the Figure 7 that the nonpolar (m-Xln)richregiontheΔG[#]valuesincrease slightlywiththeriseof temperaturebutatthealcohol rich region it is vice versa.

Figure 8 shows the variation of excess free energy ΔG^{*E} of activation at 298.15 to 323.15 K over the whole composition range for this binary system (Table3-8).Ineachcase,the Δ G^{#E}valuesare

 (hN) RT R

of the solutions of the aromatic hydrocarbons in 2- Bu-OHisenhanced,causingtheviscositytodecrease from the values expected ideally.

The free energy of activation is often regarded to be an energy barrier that a molecule must surmount in order to make a hole which is a necessaryrequirement for a molecule to flow through. The excess values throughout the whole concentration range are negative with minima falling around 0.6 mole fraction of 2-Bu-OH. The negative excess free energy indicates the reduction of energy barrierheightandhenceincreasesofviscousflow.

Table3:ExperimentaldataforFreeenergyofactivationofthebinarymixtureat298.15K

Free energy, Excess free energy, and Fitting value of excess free energy for different molar ratios at 298.15 K arelisted in the table.

Table4:ExperimentaldataforFreeenergyofactivationofthebinarymixtureat303.15K

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Free energy, Excess free energy, and Fitting value of excess free energy for different molar ratios at303.15 K arelisted in the table.

Table5:ExperimentaldataofFreeenergyofactivationofthebinarymixtureat308.15K

| | | Free Energy | FreeEnergy | Excess | Fitting |
|-----------------------|---------------------|-----------------------------------|---------------------------------------------------|------------------------------------------------------------|---------------------------------------------------|
| Molefractionof | | (Expt.) $\Delta G^{\text{\#}}$ | (Theor.) $\Delta \textbf{G}^{\text{\#}}$ id | Free Energy $\Delta G^{\#E}$ kJmol ⁻¹ | Value $\Delta \mathbf{G}^{\text{\#E*}}$ |
| $2-Bu-OH(x_1)$ | $m\text{-}Xln(x_2)$ | $kJmol-1$ | $kJmol-1$ | | $kJmol-1$ |
| 0.0000 | 1.0000 | 24.8151 | 24.8151 | 0.0000 | 0.0000 |

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Free energy, Excess free energy, and Fitting value of excess free energy for different molar ratios at 308.15 K arelisted in the table.

Table6:ExperimentaldataofFreeenergyofactivationofthebinarymixtureat313.15K

Free energy, Excess free energy, and Fitting value of excess free energy for different molar ratios at 313.15 K arelisted in the table.

Table7:ExperimentaldataforFreeenergyofactivationofthebinarymixtureat318.15K

| Molefractionof | | Free Energy (Expt.) $\Delta G^{\text{\#}}$ | FreeEnergy (Theor.) $\Delta G^{\#}$. | Excess Free Energy ΔG^{H} kJmol ⁻¹ | Fitting Value $\Delta \mathbf{G}^{\text{\#E*}}$ |
|-----------------------|---------------------|------------------------------------------------------------|----------------------------------------------------|---------------------------------------------------------------------------|---------------------------------------------------------------------|
| 2-Bu-OH (x_1) | $m\text{-}Xln(x_2)$ | $kJmol-1$ | id kJ mol $^{-1}$ | | $kJmol-1$ |
| 0.0000 | 1.0000 | 25.3527 | 25.3527 | 0.0000 | 0.0000 |
| 0.0500 | 0.9500 | 25.3106 | 25.4744 | -0.1638 | -0.1545 |
| 0.1007 | 0.8994 | 25.3383 | 25.5977 | -0.2594 | -0.2818 |
| 0.1499 | 0.8501 | 25.3149 | 25.7175 | -0.4026 | -0.3929 |
| 0.1998 | 0.8002 | 25.3255 | 25.8389 | -0.5134 | -0.4990 |
| 0.2496 | 0.7504 | 25.3546 | 25.9601 | -0.6056 | -0.5989 |
| 0.2998 | 0.7002 | 25.4160 | 26.0822 | -0.6661 | -0.6913 |

Free energy, Excess free energy, and Fitting value of excess free energy for different molar ratios at 318.15 K arelisted in the table.

| | | Free Energy | FreeEnergy | Excess | Fitting |
|-----------------------|------------------------------------------------|---------------------------------|-------------------------------------------------|---------------------------------|-----------------------------------|
| Molefractionof | | (Expt.) | (Theor.) | Free Energy | Value |
| | $\mathbf{m}\text{-}\mathbf{X}\mathbf{ln}(x_2)$ | $\Delta \textbf{G}^{\text{\#}}$ | $\Delta G^{\text{\#}}_{\text{\textnormal{id}}}$ | $\Delta G^{H\!E}$ kJmol $^{-1}$ | $\Delta \mathbf{G}^{\text{\#E*}}$ |
| 2-Bu-OH (x_1) | | $kJmol-1$ | $kJmol-1$ | | $kJmol-1$ |
| 0.0000 | 1.0000 | 25.6216 | 25.6216 | 0.0000 | 0.0000 |
| 0.0500 | 0.9500 | 25.5694 | 25.7310 | -0.1616 | -0.1498 |
| 0.1007 | 0.8994 | 25.5985 | 25.8418 | -0.2433 | -0.2705 |
| 0.1499 | 0.8501 | 25.5634 | 25.9496 | -0.3862 | -0.3745 |
| 0.1998 | 0.8002 | 25.5686 | 26.0588 | -0.4902 | -0.4734 |
| 0.2496 | 0.7504 | 25.5927 | 26.1678 | -0.5751 | -0.5665 |
| 0.2998 | 0.7002 | 25.6566 | 26.2776 | -0.6210 | -0.6526 |
| 0.3498 | 0.6502 | 25.6557 | 26.3871 | -0.7314 | -0.7284 |
| 0.3996 | 0.6004 | 25.7032 | 26.4960 | -0.7928 | -0.7917 |
| 0.4498 | 0.5502 | 25.7622 | 26.6059 | -0.8438 | -0.8419 |
| 0.4993 | 0.5008 | 25.8245 | 26.7142 | -0.8897 | -0.8777 |

Table8:ExperimentaldataforFreeenergyofactivationofthebinarymixtureat323.15K

Free energy, Excess free energy, and Fitting value of excess free energy for different molar ratios at323.15 K arelisted in the table.

Figure7:Freeenergycurvesfordifferentmolefractionsatdifferenttemperatures

Figure8:Excessfreeenergycurvesfordifferentmolefractionsatdifferenttemperatures

5. Conclusion

Studiesonthermodynamicpropertiesalongwiththeir

respective excess values in liquids and liquidmixtures is essential to understand the molecular interactions between unlike molecules, to develop new theoretical models [17] as well as engineering applications in pharmaceutical process industries. An understanding of the thermodynamic properties of 2- Bu-OH and m-Xln and their binary systemhave been measured at temperatures (298.15 to 323.15) K andin atmospheric pressure. Excess thermodynamic properties of binary mixture solvents were calculated and fitted with the Redlich–Kister equation. Thermodynamic properties derived from viscosityand its related parameters are also important in designing industrial equipments with better precision [18]. In these mixtures, the predominance of physical effect may arise due to dipole-induced dipole interaction resulting in disruption in the favorable orientation due to breaking of self-network built by H-bond among the polar molecules and rising dispersion force of nonpolar liquid. The observed excess thermodynamic values in all the mixture

indicatethesignificantinteractionbetweentheunlike molecules.

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