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Water activity in polyols: new UNIFAC parameterization

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Water activity in polyol/water systems: new UNIFAC parameterization

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Abstract

Water activities of a series of polyol/water systems were measured with an AguaLab dew point water activity meter at 298 K. The investigated polyols with carbon numbers from n=2-7 are all in liquid state at room temperature and miscible at any molar ratio with water. In aqueous solutions with the same mass concentration, the diols with 5 lower molecular weight lead to lower water activities than those with higher molecular weights. For diols with four or more carbon atoms, the hygroscopicity shows considerable differences between isomers: The 1,2-isomers - consisting of a hydrophilic and a hydrophobic part - bind less strongly to water than isomers with a more balanced distribution of the hydroxyl groups. The experimental water activities were compared 10 with the predictions of the group contribution method UNIFAC: the model predictions overestimate the water activity of water/polyol systems of substances with two or more hydroxyl groups and can not describe the decreased binding to water of isomers with hydrophobic tails. To account for the differences between isomers, a modified UNIFAC parameterization was developed, that allows to discriminate between three types of 15 alkyl groups depending on their position in the molecule. These new group-interaction parameters were calculated using water activities of alcohol/water mixtures. This leads

to a distinctly improved agreement of model predictions with experimental results while largely keeping the simplicity of the functional group approach.

20 1. Introduction

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Organic species are emitted into the atmosphere by a variety of natural and anthropogenic sources. They account for up to 50% of the total fine aerosol mass at continental mid-latitudes (Saxena and Hildemann, 1996) and for up to 90% in tropical forested areas (Yamasoe et al., 2000; Roberts et al., 2002). The amount of water absorbed by aerosol particles can be significantly altered by the presence of organics (Saxena et al., 1995; Saxena and Hildemann, 1997). Conversely, gas/particle partition-

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ing of semivolatile species is influenced by the presence of water in particles (Griffin et al., 2003). To tackle these interdependences, thermodynamic models are required that describe gas/particle partitioning and water activity (Bowman and Melton, 2004). The wide variety of organic species present in the ambient aerosol can only be han-

- ⁵ dled by models that parameterize functional groups rather than individual compounds. The most common group contribution method for organic substances is UNIFAC (Fredenslund et al., 1975, 1977). Although it has been shown that the performance of UNIFAC is not satisfactory for predicting water activities of mixtures containing multifunctional organic species (Saxena and Hildemann, 1997; Peng et al., 2001; Ming and
- Russell, 2002), it is commonly used to describe gas/particle partitioning of semivolatile organic species and water (Pun et al., 2002; Cai and Griffin, 2003; Griffin et al., 2003; Erdakos and Pankow, 2004). Ming and Russell (2002, 2004) have therefore developed an improved parameterization to model the influence of organic compounds in fogs.

This study investigates more closely how the performance of UNIFAC depends on the number and the position of functional groups. This is done for a large variety of alcohols including monofunctional as well as polyfunctional ones. Polyfunctional alcohols – so called polyols – together with polyethers have been identified by HNMR as a main class of the water-soluble organic fraction of atmospheric aerosols (Decesari et al., 2000, 2001). Various individual polyols and carbohydrates have been observed

- in biomass burning samples (Graham et al., 2002; Gao et al., 2003; Claeys et al., 2004; Simoneit et al., 2004). The hydroxyl group can therefore be considered as one of the most important functional groups of organic aerosol constituents. A high variety of mono- and polyfunctional alcohols are commercially available covering a large number of chain lengths and isomers. The focus of this study is on polyols that are
- present as liquids at room temperature and miscible at any molar ratio with water. This allows the measurement of water activities of polyol/water bulk samples over the whole composition range. The experimental data is compared with UNIFAC predictions and used together with vapour-liquid equilibrium data of alcohols at the boiling temperature to develop a new improved UNIFAC parameterization.

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2. Experimental methods

The water activities, a_w, were measured using an AquaLab water activity meter (Model 3TE, Decagon devices, USA). For all measurements, the volatile sample block available as an accessory to the instrument was used. With this sample block, the water
activity in the presence of other semivolatile components can be determined. Experimental errors for the volatile sample block are ±0.015 a_w. To correct for instrument drifts and offset, the performance of the sample block was frequently controlled and readjusted with reference samples. All measurements were performed at 298 K. The substances were purchased from Sigma-Aldrich in the best available purity. For glycerol, ethanediol, 1,2-, and 1,3-propanediol as well as 1,3-, 1,4-, and, 2,3-butanediols the purity was above 99%, for 1,4-pentanediol ≥97.0%, for 1,2-butanediol and 2,4-pentanediol ≥98.0%, for 1,2- and 2,5-hexanediol ≥97.0%, for 1,2- and 1,5-pentanediol 96%, and for 1,2,4-butanetriol and 1,7-heptanediol ≥95%. The substances were used without further purification. The water/polyol mixtures were prepared by mass percent with MiliO water using an accessing a peneltical belonce.

¹⁵ with MilliQ water using an analytical balance.

3. UNIFAC group contribution method

The UNIFAC group contribution method (Fredenslund et al., 1975, 1977) is a broadly used tool for the prediction of liquid-phase activity coefficients parameterized for a wide range of structural groups (Hansen et al., 1991). In the UNIFAC model, the activity coefficients of a molecular component $i(\gamma_i)$ in a multicomponent mixture are expressed as the sum of two contributions: a combinatorial part (*C*), accounting for size and shape of the molecule and a residual part (*R*), a result of inter-molecular interactions

 $\ln \gamma_i = \ln \gamma_i^C + \ln \gamma_i^R.$

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The combinatorial part uses the pure component properties such as volumes and surface areas to account for the excess entropic part of the activity coefficients

$$\ln \gamma_i^C = \ln \frac{\Phi_i}{x_i} + \frac{z}{2} q_i \ln \frac{\Theta_i}{\Phi_i} + \lambda_i - \frac{\Phi_i}{x_i} \sum_j x_j \lambda_j$$
(1)

where

$$\Phi_i = \frac{r_i x_i}{\sum_j r_j x_j}; \qquad \Theta_i = \frac{q_i x_i}{\sum_j q_j x_j}$$
(2)

and

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$$\lambda_i = \frac{z}{2}(r_i - q_i) - (r_i - 1)$$
(3)

with

$$r_i = \sum_k v_k^{(i)} R_k; \qquad q_i = \sum_k v_k^{(i)} Q_k.$$
 (4)

In these equations, x_i is the mole fraction of component *i*, $v_k^{(i)}$ is the number of groups 10 of type k in molecule i, and z is the lattice coordination number, a constant set equal to ten (Fredenslund, 1975). The group volume and surface area parameters R_k and Q_k are based on the work of Bondi (1968).

The residual part of the activity coefficient is given by

$$\ln \gamma_i^R = \sum_k v_k^{(i)} \left[\ln \Gamma_k - \ln \Gamma_k^{(i)} \right],$$
(5)

where Γ_k is the group residual activity coefficient in the mixture and $\Gamma_k^{(i)}$ the one in a reference liquid containing only molecules of type *i*. The residual activity coefficients are calculated as

$$\ln \Gamma_{k} = Q_{k} \left[1 - \ln \left(\sum_{m} \Theta_{m} \Psi_{mk} \right) - \sum_{m} \left(\Theta_{m} \Psi_{km} / \sum_{n} \Theta_{n} \Psi_{nm} \right) \right]$$
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with

$$\Theta_m = \frac{Q_m X_m}{\sum_n Q_n X_n}; \qquad \Psi_{mn} = \exp\left[\frac{-a_{mn}}{T}\right]$$

In these expressions, Θ_m is the surface area fraction of group *m*, X_m the mole fraction of group *m* in the mixture, and Ψ_{mn} the interaction parameter between the groups *m* and ⁵ *n*. Ψ_{mn} is temperature dependent and a function of the group interaction parameter a_{mn} .

For this study, literature values for group volume and surface area parameters R_k and Q_k as well as for the group interaction parameters a_{mn} are taken from Hansen et al. (1991). The values used for the investigated alcohols are given in Tables 1 and 2.

10 4. Results and discussion

4.1. Water activity measurements

Table 3 lists the measured water activities for the investigated polyol/water systems. Only a part of the polyols could be purchased in an anhydrous quality. The water activity a_w of the different polyols before the addition of water varied between 0 and 0.125. Based on these values, the water content was estimated to be between 0 and 1.5 wt% for the different liquids. The compositions of the water/polyol mixtures were therefore corrected to include the water which was present in the compounds as purchased. Such a correction was necessary for glycerol, 1,2-butanediol, all pentanediols,

²⁰ For some of the investigated polyol/water mixtures, water activity data is already available in the literature. Figure 1 shows a comparison of the water activity from this study and literature data for glycerol and 2,3-butanediol. For glycerol/water, water activity has been determined by vapour-liquid equilibrium measurements under isobaric and isothermal conditions at the boiling temperature (Gmehling and Onken, 1977, 2003;

the hexanediols, the heptanediol, as well as 1,2,4-butanetriol.

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(7)

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Gmehling et al., 1981, 1988) as well as by the isopiestic method (Scatchard et al., 1938) and with an AquaLab water activity meter similar to the one used in the present study (Ninni et al., 2000). The quite diverse quality of the different datasets necessitates a careful data selection. Measurement series with large scatter were considered

- as less significant. Figure 1a shows the datasets that were judged to be most reliable for the glycerol/water system. The presented data span a temperature range from 25°–290°C. For 2,3-butanediol/water, the literature data is shown in Fig. 1b. All datasets are based on vapour-liquid equilibria under isobaric conditions at the boiling temperature at 60°–180°C except the one from Borghesani et al. (1989), which was measured with a vapour pressure osmometer at 25°C. It can be seen that there is quite a large scatter
- in some datasets and considerable deviations between datasets.

The data from this study show very good agreement with literature values in the case of glycerol/water, see Fig. 1. For 2,3-butanediol/water they exhibit lower water activities for a given mass concentration than most other datasets, but are in good agreement

- ¹⁵ with Borghesani et al. (1989), whose measurements are however restricted to dilute solutions. In general, water activity measurements by the isopiestic method or with the AquaLab water activity meter are quite accurate (Ninni et al., 2000). In comparison, vapour-liquid equilibrium data at the boiling temperature tend to have larger errors especially when one component exhibits a low but not negligible concentration in the
- ²⁰ gas phase, as it is the case for most polyols in this study. Based on the good agreement for glycerol/water with literature values, and because there is no obvious reason why the accuracy should be worse for other polyols, we assume that the data from this study are accurate within $\pm 0.015 a_w$ as specified for the AquaLab water activity meter with the volatile sample block.
- Figure 2 shows a comparison of the water activities for the investigated diol/water systems as measured in this work. In aqueous solutions with the same mass concentration, the diols with lower molecular weight lead to lower water activities than those with higher molecular weights. For the diols with four or more carbon atoms, there are also considerable differences between isomers. The 1,2-isomers consisting

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of molecules with a hydrophilic and a hydrophobic part bind less strongly to water than the other isomers with a more balanced distribution of the hydroxyl groups.

4.2. UNIFAC predictions

Figures 3 and 4 show the experimental data and UNIFAC predictions for the investigated butanol and pentanediol/water mixtures. For butanol/water, experimental data from vapour-liquid equilibria at the boiling temperature at 25°-35°C and under isobaric conditions at 700–760 mmHg corresponding to a temperature range from 80 to 120°C is shown. There is no temperature dependence sufficiently strong to exceed the scatter between various measurements of the inspected alcohol/water vapour-liquid equilibria. For pentanediol/water mixtures, no literature data is available except for 1,5-pentanediol in the more dilute region (Borghesani et al., 1989). The measurements from this study are in good agreement with this dataset.

Figure 3 shows UNIFAC predictions at 25° and 100°C for the butanol/water systems. The parameters for the calculations are listed in Tables 1 and 2. The UNIFAC parameterization is in good agreement with the experimental data for all butanols but tert-butanol, for which the water activity is overestimated (Fig. 3d). UNIFAC predicts a moderate temperature dependence with stronger water activity depression at higher temperatures.

For the 1,2-pentanediol/water mixture, the UNIFAC prediction at 25°C is in fair agree-²⁰ment with the measurements, whereas for all other pentanediols the water activity at given concentration is clearly overestimated. Figure 4 also shows the water activities for the UNIFAC parameterization by Ming and Russell (2002) with parameters specifically developed for monosaccharides as listed in Table 2. This parameterization is in better agreement with the experimental data as the standard UNIFAC, but it can not account for differences between isomers.

UNIFAC predictions are also shown in Fig. 1 for glycerol and 2,3-butanediol/water mixtures. The Ming and Russell parameterization leads to an excellent agreement for glycerol, but overall the UNIFAC parameterizations are limited by the fact that they do

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not account for differences between isomers (Sandler, 1994).

4.3. New UNIFAC parameterization

In UNIFAC, alkyl groups, whether connected to another alkyl group or to a hydroxyl group, are treated the same way. Such a simple parameterization requires only a low
 number of structural groups, however, can not represent interactions between adjacent atoms in a molecule. UNIFAC does for example not account for proximity effects that occur when two or more strongly polar groups are located on the same or adjacent carbon atoms (Sandler, 1994). One way to overcome this limitation is to define parameters that are specific for certain compound classes. Such parameterizations have
 been proposed by Peng et al. (2001) for dicarboxylic and hydroxyl-carboxylic acids and by Ninni et al. (1999) for poly(ethylene glycol). Ming and Russell (2002) proposed modified parameterizations for long-chain monofunctional compounds, monosaccharides, hydroxyl-acids and diacids. For polyols (Ninni et al., 2000) and sugars (Peres and Macedo, 1997) modified parameterizations within the UNIFAC-Larsen model (Larsen et al., 1987) have been described.

In this work, a different approach is used: the same group interaction parameters are applied to monofunctional alcohols as well as polyols, however, to improve the accuracy, more specific structural groups are defined. The aim of this new UNIFAC parameterization is therefore twofold: (i) to refine the definition of alkyl groups by accounting for their positions in the molecules; (ii) to adjust group interaction parameters

- ²⁰ counting for their positions in the molecules; (ii) to adjust group interaction parameters needed to describe polyols. To reach these two goals, a large dataset is required that includes as many isomers as possible. Besides the water activity data acquired in this study for polyol/water, vapour-liquid equilibria of alcohol/water systems from the liter-ature were also considered. The literature data used for the parameterization covers the temperature range from 20° to 140°C and is listed in Table 4. A fugacity correction
- (Zemp and Francesconi, 1992; Tsonopoulos, 1974) was applied to smaller alcohols.

The UNIFAC parameterization of alcohol/water systems shows two main weaknesses: it underestimates the water uptake of substances with two or more hydroxyl



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groups and it can not account for the reduced binding to water of isomers with hydrophobic tails. A new definition of alkyl groups should therefore improve these two shortcomings. Moreover, it should be unambiguous and easy to apply. Different sets of alkyl group definitions were tested by fitting UNIFAC water activities to the experi-⁵ mental data. In order to achieve this, water-alkyl and water-hydroxyl group interaction parameters were varied while keeping other parameters constant. The UNIFAC volume (R_k) and surface area (Q_k) parameters for the alkyl, hydroxyl and water groups were maintained at the literature values (Hansen et al., 1991) listed in Table 1.

This optimization process led to the definition of the following three types of alkyl ¹⁰ groups:

- CH_n (n=0, 1, 2) with a hydroxyl group: accounts for the induced polarity of alkyl groups directly connected to the electronegative hydroxyl group. This type of alkyl group shows an enhanced interaction with water compared to the other cases.
- CH_n (n=0, 1, 2, 3) in hydrophobic tails: accounts for the non-polar nature of alkyl chains that easily agglomerate and form micelles in water. This type of alkyl group shows decreased interactions with water compared to the other cases. The minimal size of a hydrophobic tail is -CH₂-CH₃. As only exception, for ethanol it is -CH₃.

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- CH_n (n=0, 1, 2, 3) in alcohols: constitutes the general type of alkyl group which applies when the special conditions for the other two types are not fulfilled.

The new group interaction parameters are listed in Table 2. To exemplify the use of the new definitions several examples are given in Fig. 5. Figures 1, 3 and 4 reveal that the new parameterization is able to account for the observed differences between isomers. The agreement with the experimental water activities for the monofunctional alcohol/water as well as the polyol/water systems is distinctly improved. For some mixing ratios, the calculated water activity reaches values above 1 in the butanol/water mixtures (Fig. 3), because the UNIFAC prediction was carried out with the premise that

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the two components are present in the same phase. If the calculations are repeated allowing for the occurrence of phase separations, the new UNIFAC parameterization correctly predicts miscibility gaps for all butanols but tert-butanol, while the parameterization by Hansen et al. (1991) also predicts a phase separation for tert-butanol, which is in disagreement with experimental observations (e.g. Fischer and Gmehling, 1994).

To determine the ability of aerosol particles to serve as cloud condensation nuclei (CCN), an accurate description of their hygroscopicity at high relative humidities is indispensable. The UNIFAC parameterization by Hansen et al. (1991) strongly underestimates the water content of most investigated polyols for high water activities. For the

- pentanediols in Fig. 4, it predicts a mass increase of about 50% at a relative humidity of 95% due to water uptake, whereas the experimental data indicates a triplication of the mass for all pentanediols but the 1,2-isomer, qualifying them as more potent CCN constituents than assumed based on UNIFAC predictions. The new parameterization presented in this study describes the water uptake of polyols satisfactory, enabling the upper of UNIFAC prediction the aritical superacturation for the mass of UNIFAC prediction.
- ¹⁵ use of UNIFAC-predicted water activities to calculate the critical supersaturation for CCN activation based on Köhler theory (Pruppacher and Klett, 1997).

5. Conclusion and outlook

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The analysis of an extensive dataset of vapour-liquid equilibrium and water activity data for alcohol/water mixtures showed that the ability of UNIFAC to describe the non-ideality

- of these systems is limited. The main reason for this limitation is due to the fact that the position of the hydroxyl groups in the molecule influences the interactions with water. The definition of structural groups within UNIFAC can not account for such differences. Therefore we developed a new UNIFAC parameterization that discriminates between three types of alkyl groups based on their position in the molecule. This modified the types of alkyl groups based on their position in the molecule.
- ²⁵ UNIFAC model leads to a distinctly improved agreement with experimental data without losing much of the simplicity of the group contribution approach. In future work, this parameterization shall be extended to mixtures containing further functional groups

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such as carboxylic acid, ether and carbonyl groups with the aim to establish a UNIFAC version that is optimized to describe organic substances of atmospheric relevance and can be implemented in or combined with inorganic thermodynamic models.

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Table 1. UNIFAC volume (R_k) and surface area (Q_k) parameters for subgroup k (from Hansen et al., 1991).

Main group	Sub group <i>k</i>	R_k	Q_k
CH _n (n=0,1,2,3)	CH ₃	0.9011	0.848
	CH_2	0.6744	0.540
	CH	0.4469	0.228
	С	0.2195	0
OH	OH	1.0000	1.200
H ₂ O	H ₂ O	0.9200	1.400

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UNIFAC (Hansen et al., 1991)				
	CH _n	OH	H ₂ O	
CH _n	0.0	986.5	1318	
OH	156.4	0.0	353.5	
H_2O	300.0	-229.1	0.0	
UNIF. paran	UNIFAC (Ming and Russell, 2002,			
paran	CH _n	OH	H ₂ O	
CH _n	0.0	986.5	1318	
CH _n OH	0.0 156.4	986.5 0.0	1318 189.7	

UNIFAC (this study)								
	CH _n with -OH group	CH_n in hydrophobic tails	CH_n in alcohols	OH	H ₂ O			
CH _n with -OH group	0.0	0.0	0.0	986.5	2314			
CH_n in hydrophobic tails	0.0	0.0	0.0	986.5	1325			
CH_n in alcohols	0.0	0.0	0.0	986.5	1890			
OH	156.4	156.4	156.4	0.0	276.4			
H ₂ O	-89.71	362.1	162.3	-153.0	0.0			

Table 2. UNIFAC group interaction parameters a_{mn} between groups *m* and *n*.

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Table 3. Water activities of the investigated polyol/water mixtures at 25° C. Concentrations are given in mole fraction of water $x(H_2O)$. A star (*) marks values corrected for water content of substance as purchased.

		I±						
Ethan	ediol	Glyce	erol*	1,2-Prop	anediol	1,3-Prop	anediol	
$x(H_2O)$	aw	$x(H_2O)$	a _w	$x(H_2O)$	a _w	$x(H_2O)$	a _w	
0.0557	0.048	0.1078	0.113	0.0796	0.083	0.0755	0.072	
0.1104	0.097	0.1977	0.167	0.1699	0.174	0.1687	0.16	
0.1789	0.152	0.3189	0.255	0.2753	0.285	0.2795	0.27	
0.2997	0.258	0.4144	0.346	0.3635	0.38	0.3661	0.362	
0.3956	0.354	0.5171	0.448	0.4330	0.436	0.4403	0.441	
0.4680	0.427	0.6053	0.545	0.5248	0.54	0.5318	0.537	
0.5530	0.511	0.6880	0.636	0.5938	0.604	0.6015	0.608	
0.6300	0.593	0.7615	0.726	0.6750	0.678	0.6805	0.687	
0.6991	0.665	0.8094	0.783	0.7399	0.742	0.7439	0.753	
0.7580	0.737	0.8500	0.837	0.7953	0.79	0.7997	0.808	
0.8078	0.79	0.8853	0.88	0.8547	0.847	0.8535	0.862	
0.8600	0.845	0.9204	0.929	0.9078	0.91	0.9044	0.914	
0.9140	0.915	0.9479	0.956	0.9572	0.961	0.9559	0.964	
0.9610	0.969	0.9678	0.975					
		0.9864	0.988					

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Table 3. Continued.

1,2-Butanediol*		1,3-Buta	anediol	1,4-Buta	anediol	2,3-Butanediol		
x(H ₂ O)	a _w							
0.1078	0.159	0.0838	0.099	0.1153	0.134	0.0864	0.113	
0.1943	0.269	0.1523	0.176	0.2021	0.226	0.1574	0.2	
0.2746	0.379	0.2497	0.29	0.2877	0.327	0.2619	0.332	
0.3574	0.479	0.3357	0.394	0.3663	0.418	0.3454	0.425	
0.4475	0.582	0.4062	0.471	0.4271	0.485	0.4145	0.496	
0.5387	0.67	0.4738	0.54	0.4953	0.559	0.4847	0.563	
0.6192	0.742	0.5482	0.609	0.5633	0.626	0.5588	0.627	
0.6853	0.791	0.6171	0.671	0.6311	0.69	0.6284	0.685	
0.7416	0.832	0.6948	0.742	0.6915	0.746	0.6998	0.744	
0.8096	0.871	0.7669	0.802	0.7529	0.798	0.7682	0.797	
0.8770	0.915	0.8325	0.854	0.8138	0.846	0.8327	0.843	
0.9287	0.942	0.9098	0.926	0.8627	0.887	0.9096	0.916	
0.9619	0.976	0.9661	0.976	0.9062	0.924	0.9670	0.971	
0.9843	0.992			0.9379	0.953			
				0.9627	0.97			
				0.9802	0.986			

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Table 3. Continued.

1,2-Pentanediol*		1,4-Pent	anediol*	1,5-Pent	1,5-Pentanediol*		anediol*	
	x(H ₂ O)	a _w						
	0.1277	0.199	0.1532	0.204	0.0951	0.137	0.1349	0.189
	0.2118	0.342	0.2330	0.315	0.1819	0.245	0.2289	0.32
	0.2871	0.452	0.3055	0.408	0.2585	0.359	0.3009	0.422
	0.3548	0.544	0.3697	0.485	0.3250	0.444	0.3709	0.509
	0.4095	0.61	0.4253	0.545	0.3850	0.516	0.4343	0.582
	0.4753	0.687	0.4935	0.616	0.4538	0.595	0.4997	0.649
	0.5419	0.762	0.5590	0.675	0.5329	0.676	0.5550	0.699
	0.6027	0.817	0.6396	0.75	0.6130	0.754	0.6226	0.755
	0.6778	0.866	0.7062	0.801	0.6792	0.806	0.6996	0.806
	0.7463	0.907	0.7759	0.843	0.7489	0.85	0.7825	0.853
	0.8288	0.933	0.8438	0.886	0.8203	0.891	0.8576	0.895
	0.9188	0.966	0.9116	0.93	0.8957	0.933	0.9110	0.928
	0.9638	0.976	0.9596	0.972	0.9535	0.967	0.9499	0.96
	0.9852	0.989	0.9813	0.986	0.9784	0.982	0.9817	0.987

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Table 3. Continued.

1,2-Hexanediol*		2,5-Hexanediol*		1,7-Hept	tanediol*	1,2,4-Butanetriol*		
х(H ₂ O)	a _w							
0.1553	0.281	0.1570	0.214	0.1524	0.256	0.1510	0.119	
0.2548	0.442	0.2491	0.346	0.2399	0.419	0.2496	0.177	
0.3366	0.568	0.3352	0.463	0.3259	0.552	0.3818	0.297	
0.4047	0.659	0.4136	0.558	0.4001	0.641	0.4732	0.386	
0.4636	0.734	0.4790	0.625	0.4633	0.733	0.5469	0.465	
0.5195	0.797	0.5634	0.716	0.5215	0.795	0.6237	0.551	
0.5688	0.85	0.6341	0.774	0.5885	0.861	0.6952	0.637	
0.6211	0.891	0.7184	0.834	0.6398	0.897	0.7520	0.705	
0.6866	0.938	0.8018	0.878	0.7046	0.937	0.8029	0.772	
0.7489	0.965	0.8785	0.92	0.7823	0.973	0.8491	0.828	
0.8249	0.982	0.9317	0.951	0.8830	0.987	0.8906	0.878	
0.9089	0.988	0.9675	0.98			0.9242	0.921	
						0.9509	0.951	
						0.9731	0.975	
						0.9860	0.989	

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Substance	Number of data points	Temperature (°C)	Pressure (mmHg)	Fugacity correction	References	-		
Ethanol	15	25.13	24-60	yes	Gmehling and Onken (2003), p. 223		Title	Page
2-propanol	12 10	25 80–94 93–101	20–35 760 760	yes yes	Gmehling and Onken (1977), p. 406 Gmehling and Onken (1977), p. 317 Gmehling and Onken (1977), p. 406		Abstract	Introduction
1 buturior	3	103–113 95–99	700 759.81	yes	Gmehling and Onken (1077), p. 400 Gmehling et al. (1981), p. 334 Gmehling and Onken (2003), p. 575		Conclusions	References
2-butanol	8 8	25 87–95	28–33 759.81	yes ves	Gmehling and Onken (2003), p. 586 Gmehling and Onken (2003), p. 587		Tables	Figures
Isobutanol	14	90–107	760	yes	Gmehling and Onken (1977), p. 439			
tert-butanol	2 13	25 79–84	27–37 760	yes ves	Gmehling et al. (1981), p. 343 Gmehling and Onken (2003), p. 618			►1
1-pentanol 3-met-1-butanol	19 6	95–138 95–109	760 760	no no	Gmehling et al. (1988), p. 309 Gmehling et al. (1988), p. 308		•	•
1-hexanol Cyclohexanol	6 13	21 90	6–19 104–560	no no	Gmehling et al. (1981), p. 422 Gmehling and Onken (1977), p. 514		Back	Close

Table 4. Literature data used for the determination of the UNIFAC group interaction parameters.



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b

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0.4

mass fraction of 2.3-butanediol

06

0.8

Fig. 1. Water activity of (a) glycerol/water and (b) 2,3-butanediol/water mixtures. Squares: vapour-liquid equilibrium measurements at constant pressures from 750-760 mmHg (100-290°C), filled circles: vapour liquid equilibrium measurements at constant pressures from 200-600 mmHg (60-180°C), triangle: vapour-liquid equilibrium data at constant temperature at 25°C, all data taken from Gmehling and Onken (1977, 2003) and Gmehling et al. (1988), different colors indicate different measurement series; diamonds: water activity measurements at 25°C (yellow: Ninni et al., 2000, only glycerol; purple: Scatchard et al., 1938, only glycerol; orange: Borghesani et al., 1989, only 2,3-butanediol; green: this study). Lines: UNIFAC predictions at 25°C (black: Hansen et al., 1991; blue: Ming and Russell, 2002; red: using the new parameters from this study).

а

0.8

0.6

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vater activity a

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mass fraction of glycerol

06

0.8

0.8

0.6

0.4

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0

0

≥

water activity a

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Fig. 2. Measured water activities of the investigated diols at 25°C.



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Fig. 3. Water activity of butanol/water mixtures: **(a)** n-butanol, **(b)** 2-butanol, **(c)** isobutanol, **(d)** tert-butanol. Squares: vapour-liquid equilibrium measurements at constant pressures from 700–767 mmHg (80–120°C); triangles: vapour-liquid equilibrium measurements at constant temperatures from 25–35°C, all data taken from Gmehling and Onken (1977, 2003) and Gmehling et al. (1981, 1988), different colors indicate different measurement series. Solid lines: UNIFAC predictions at 25°C, dotted lines: UNIFAC predictions at 100°C (black: Hansen et al., 1991; red: using the new parameters from this study).

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Fig. 4. Water activity of pentanediol/water mixtures at 25°C: **(a)** 1,2-pentanediol, **(b)** 1,4-pentanediol, **(c)** 1,5-pentanediol, **(d)** 2,4-pentanediol. Diamonds: water activity measurements at 25°C (orange: Borghesani et al., 1989; green: this study). Lines: UNIFAC predictions (black: Hansen et al., 1991; blue: Ming and Russell, 2002; red: using the new parameters from this study).

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Fig. 5. Examples to the use of the structural groups of alcohols and polyols.