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Phase Diagrams of the Aqueous Two-Phase Systems of Poly(ethylene glycol)/Sodium Polyacrylate/Salts

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Abstract: Aqueous two-phase systems consisting of polyethylene glycol (PEG), sodium polyacrylate (NaPAA), and a salt have been studied. The effects of the polymer size, salt type (NaCl, Na₂SO₄, sodium adipate and sodium azelate) and salt concentrations on the position of the binodal curve were investigated. The investigated PEG molecules had a molar mass of 2,000 to 8,000 g/mol, while that of NaPAA was 8,000 g/mol. Experimental phase diagrams, and tie lines and calculated phase diagrams, based on Flory-Huggins theory of polymer solutions are presented. Due to strong enthalpic and entropic balancing forces, the hydrophobicity of the added salt has a strong influence on the position of the binodal, which could be reproduced by model calculations.

Keywords: poly(ethylene glycol); sodium polyacrylate; ATPS; Flory-Huggins theory; phase diagram; hydrophobic ions
1. Introduction

Aqueous two phase systems (ATPS) based on water-soluble polymers are highly useful tools for extraction and purification of biomolecules [1-8]. An important aspect of the ATPS investigations concerns the search of systems containing reduced amount of phase forming compounds. Such ATPS have potential application in industry due to the low cost of purification/extraction of biomolecules. ATPS containing polymers that are easily recycled are interesting mostly for environmental reasons. The ATPS here investigated are based on polyethylene glycol (PEG) and sodium polyacrylate (NaPAA), which together with a salt separate into a PEG-enriched top-phase and a NaPAA enriched bottom phase. The PEG from the top-phase can be recycled by adding strongly hydrophilic salts as described by Albertsson [1], and the NaPAA can form a concentrated complex at low pH (<4) with PEG [9]. These properties are thus of remarkable, industrial importance. These polymers are environmentally friendly and are relatively low-cost chemicals. In addition, when mixed together they phase separate at quite low polymer concentrations and at rather mild salt concentrations. In previous works PEG-PAA two-phase systems have been used for the partitioning of the proteins GFP (green fluorescence protein) [6], hemoglobin, lysozyme and glucose-6-phosphate dehydrogenase [5]. Saravanan et al. have studied the effects of pH and salt on the partitioning of myoglobin and ovalbumin in PEG-PAA systems [7]. In these earlier studies on the applicability of the systems for bioseparations it was found that the systems are relatively inert (giving low protein denaturation), have a relatively low viscosity and can separate by gravity within a few minutes [5,6].

The aim of this work was to investigate the physico-chemical properties of the segregating two-polymer ATPS, formed by PEG, NaPAA and different salts: NaCl, Na₂SO₄, NaOOC-(CH₂)₄-COONa (di-sodium adipate), and NaOOC-(CH₂)₇-COONa (di-sodium azelate). Furthermore, the underlying basic thermodynamic driving forces of ion mixing entropy and ion hydrophilicity was investigated in the framework of Flory-Huggins theory [10].

The most studied aqueous two-phase systems are based upon the segregative behavior of two non-charged polymers, for instance PEG and dextran. In such systems, the salt type and concentration have very little influence on the position of the binodal. In contrast, the salt concentration has a very strong influence on two-polymer ATPS, where one or both of the polymers are charged, as shown by Albertsson for the PEG-dextran-sulphate [1], and Bergfeldt et al. [11] for the sodium polystyrene sulphonate-sodium polyacrylate system. The use of charged ampholytic polymers based on acrylate monomers for two-phase extraction of proteins have also been studied by Hughes and Lowe [12]. However, the systems they used contained high amount of relatively complex polymers. Persson et al. have also studied the use of cationic polymers for protein partitioning [13]. Waziri et al. showed that PEG can form a two-phase system with relatively low polymer content with an almost neutral ampholyte polydiallylamino-ethanolate-dimethyl-sulfoxide (PAEDS) [14]. For these cited examples the polymers are still non-commercial and have a complex composition which is a strong drawback for developing a reliable bioseparation process. Recently, the effect of different inorganic ions on the PEG-polyacrylate systems has been investigated [15]. The tendency for a charged and a non-charged polymer to segregate, without salt addition, into two-phases is strongly suppressed by the large difference in small ion concentration between the phases. Therefore, in systems without salt addition, two-phase formation occurs only at very high polymer concentration (e.g., 10 wt% of each polymer)
when the mixing enthalpy becomes strong enough to favor separation, as observed previously [16,17]. However, by addition of salt, phase separation can occur at significantly lower polymer concentrations (e.g., 3–5 wt% of each polymer) [15]. Apart from the salt concentration, the ion type is also important. Gupta et al. [15] found that the binodal curve moves to the water-rich corner in a phase diagram when the salt ion was exchanged from nitrate to sulfate and/or to phosphate for a fixed molarity of the salt. Since systems with low polymer content is desired because of lower cost, viscosity and biocompatibility, there is a natural interest to find out salts that can move the binodal curve closer to the water rich region in a phase diagram. For instance, the charge, hydrophilicity and the concentrations of the ions influence strongly the efficiency of the separation and the shape of the binodal, as shown below. In this work the effects of such salts on the two-phase system is explained by using the Flory-Huggins model for polymer solutions. The presented work here shows novel results in several aspects: the effects of ion concentration and ion partition preference have been separated, namely hydrophobic anions will favor phase separation more than hydrophilic anions at the same concentration. We have also shown that a simple mean field model can explain the changes in terms of balancing the partitioning of the counter ions (in the current work the Na⁺ ions). Finally, the systems presented here are also new with respect to the low polymer molecular weight and low polymer concentration needed for phase separation.

2. Model and Calculation Procedure

The Flory-Huggins model [10] has been extensively used on reproducing both fundamental partitioning effects such as entropic repulsion [18] and more specific effects such as thermoseparation [19] and the electrochemical driving force in aqueous two-phase systems [18,20,21]. The model is based on mean-field statistical mechanical treatment of the combinatorial entropy upon mixing a polymer with a solvent in a lattice system, where the size of the lattice site unit is the same as the smallest component (usually the solvent). All other component sizes and polymerization degree are counted in a number of lattice sites units. The enthalpy of mixing is obtained by simply regarding first neighbor interactions, and the nearest lattice neighbors are always represented by the mean-field composition. In this model no reactions are taking place, thus in order to obtain the phase diagrams it is sufficient to minimize the free energy of mixing. In the search for the minimum global mixing free energy, random amounts of components are moved between the phases. The chemical potentials of the non-charged components are kept equal between the phases. For ionic species, a special treatment is performed. First of all we divide the ions interaction in two parts. One is the coulombic long-range interaction, which due to its strength causes the phases to become electroneutral at equilibrium. The other interaction is the sum of all other interaction types (e.g., polar interactions) and effects apart from the direct coulombic interaction. In the calculations, the coulombic interaction has simply been replaced by setting an electroneutrality condition on each phase. Therefore, by forcing electroneutrality in a model calculation, the fundamental effect of long-range coulombic force is implicitly accounted for in the calculations [21,22]. The polar or hydrophilic character of the ion has been collected into one nearest neighbor interaction. Thus a mixing enthalpy of the system is calculated where the coulombic interaction is omitted, and its free energy is shifted from a real system by a constant term. However, the fundamental effect of coulombic interaction is retained since the system is forced into
The electroneutrality of the ion partitioning will cause the formation of an electrochemical potential or driving force of the system, if the ions have different preferences of the phases. The partitioning of an ion can be better understood by introducing two concepts: “real ion partition” and “intrinsic ion partition”, where the former is simply calculated as the ratio of the ion concentration in the phases and the latter is calculated through the difference in non-ideal chemical potential of the ion in the two-phases, which in turn is obtained by calculating the non-ideal chemical potential of the ion as a non-charged component. The difference in real and intrinsic partitioning corresponds to a free energy cost (per charge), which must attain the same value for all charged species for satisfying a global free energy minimum. Having these constraints, it is possible to treat ion partitioning in the framework of Flory-Huggins theory.

2.1. The Flory-Huggins Model

The entropy of mixing water (1), PEG (2), polyacrylate (3), small cations (4) and anions (5) is given as follows:

\[ \Delta S_{\text{mix}} = -R n_{\text{tot}} \left[ \sum_{i=1}^{5} \phi_i \ln \phi_i \right] \]  

(1)

In the equation \( n_{\text{tot}} \) is the total number of mole of lattice sites, and \( M_i \) and \( \phi_i \) are the molecular size (expressed in units of the water molecules) and volume fraction of the different components, respectively. The volume fraction of a component equals the number of moles of lattice sites of the component divided by the total lattice volume of the phase:

\[ \phi_i = \frac{M_i n_i}{\sum_{j=1}^{5} M_j n_j} = \frac{M_i n_i}{V_{\text{phase}}} \]  

(2)

For simplicity, the lattice size of the water molecule and salt ions are set equal to one, thus:

\[ M_i = 1 \ (i = 1, 4, 5) \]  

(3)

The enthalpy is given by introducing the pairwise nearest neighbor interaction parameter \( w_{ij} \) between component (i) and (j), with the unit J/mol (no interactions of diagonal type, \( w_{ii} = 0 \)). The interaction and size parameters are given in Table 1.

Since the Flory-Huggins theory is a mean field theory the enthalpy is dependent on the average concentration of the components. Thus, we have:

\[ \Delta H_{\text{mix}} = n_{\text{tot}} \left[ \sum_{i=1}^{4} \sum_{j=1}^{5} w_{ij} \phi_i \phi_j \right] \]  

(4)

\[ \Delta G_{\text{mix}} = \Delta G_{\text{mix}} - T \Delta S_{\text{mix}} \]  

(5)

The global mixing free energy of the two phase system is compared with a corresponding homogenous 1-phase system.

For the uncharged species, we add the equilibrium condition:

\[ \mu_{\text{water}}^{\text{top}} = \mu_{\text{water}}^{\text{bottom}} \]  

(6)

\[ \mu_{\text{PEG}}^{\text{top}} = \mu_{\text{PEG}}^{\text{bottom}} \]  

(7)
The chemical potential exemplified for the top-phase for component (i) is:

\[ \mu_{i}^{\text{top}} = \mu_{i} + h_{i}^{\text{top}} - T \xi_{i}^{\text{top}} \]  

(8)

where \( \mu_{i}^{\text{op}} \), \( h_{i}^{\text{op}} \) and \( \xi_{i}^{\text{op}} \) are the standard chemical potential (set to a constant in the calculation), the partial molar enthalpy and the partial molar entropy, respectively. The two latter are obtained in the framework of Flory-Huggins theory [18] as:

\[ \frac{\partial \Delta S_{\text{mix}}}{\partial n_{i}} = s_{i} = -R \left[ \ln \phi_{i} + 1 - M_{i} \sum_{j=1}^{\phi_{j}} \frac{\phi_{j}}{M_{j}} \right] \]  

(9)

\[ \frac{\partial \Delta H_{\text{mix}}}{\partial n_{i}} = h_{i} = M_{i} \sum_{j=1}^{\phi_{j}} \phi_{j} w_{ij} - M_{i} \sum_{j \neq k}^{\phi_{j}} \phi_{k} w_{jk} \]  

(10)

Using the general form of Equations 6–8 and inserting expressions of Equations 9 and 10, it is possible to obtain the intrinsic partitioning of a component (charged or non-charged):

\[ 0 = h_{i}^{\text{top}} - h_{i}^{\text{bottom}} - T \xi_{i}^{\text{top}} + T \xi_{i}^{\text{bottom}} \]  

(11)

and

\[ \ln \frac{\phi_{i}^{\text{top}}}{\phi_{i}^{\text{bottom}}} = \ln K_{i,\text{intrinsic}} = M_{i} \left[ \sum_{j=1}^{\phi_{j}^{\text{top}}} \frac{\phi_{j}}{M_{j}} - \sum_{j=1}^{\phi_{j}^{\text{bottom}}} \frac{\phi_{j}}{M_{j}} \right] - \frac{h_{i}^{\text{top}} - h_{i}^{\text{bottom}}}{RT} \]  

(12)

**Table 1.** Parameters for calculating the phase diagrams of PEG/NaPAA/salt systems. Components: 1 water, 2 PEG, 3 polyacrylate, 4 small cation, 5 small anion (−1 or −2). Temperature 25 °C.

<table>
<thead>
<tr>
<th>Component (charge)</th>
<th>Interaction parameters (J/mol)</th>
<th>Polymerization degree (M_{i}) (corresponding polymer)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water</td>
<td>( w_{12} = w_{13} = w_{14} = w_{15} = 0 )</td>
<td>1</td>
</tr>
<tr>
<td>PEG</td>
<td>( w_{12} = w_{24} = w_{25} = 0 ), ( w_{23} = 2,400 )</td>
<td>202 (PEG 4000)</td>
</tr>
<tr>
<td>Polyacrylate (−85)</td>
<td>( w_{12} = w_{24} = w_{25} = 0 ), ( w_{23} = 2,400 )</td>
<td>202 (PEG 4000)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>404 (PEG 8000)</td>
</tr>
<tr>
<td>Small cation (+1)</td>
<td>( w_{14} = w_{23} = w_{34} = w_{45} = 0 )</td>
<td>335 (NaPAA 8000)</td>
</tr>
<tr>
<td>Small anion (−1)</td>
<td>( w_{15} = w_{35} = w_{35} = 0 ) ( w_{25} = 0 ) for (−1) anion</td>
<td>1</td>
</tr>
<tr>
<td>Small anion (−1) or (−2)</td>
<td>( w_{25} = 0 ) for (−2) anion or ( w_{25} = 27,000 ) (*)</td>
<td>1</td>
</tr>
</tbody>
</table>

(*) value for the “SO_{4}^{2−}” anion in Figure 6 and binodal B in Figure 7.

2.1. Calculation Methodology

The starting up of the calculation of the phase diagrams is performed by defining the total composition of the system. Then an attempt to construct a two-phase system is made by starting up with an unsymmetrical composition of the two-phases. In a program, random amounts of the
components move between the phases until the global minimum free energy is found. The movements are performed under the electroneutrality condition and the condition of chemical equilibrium in Equations 6 and 7 and the electrochemical potential described below.

2.2. Electrochemical Potential

If the ions would partition freely according to their non-ideal chemical potential (i.e., the intrinsic partitioning), then at the lowest free energy, the system, would not be electroneutral. However, since electroneutrality is forced on the system, every ion free energy is raised by a term scaled to the ion charge. This is the electrochemical driving force and is obtained as follows. Firstly, a molar transfer free energy of a charge (from bottom to top-phase) can be calculated from partition coefficient of an ion: \( K_{i,\text{real}} \) (ratio of top and bottom ion concentration).

\[
\Delta G_{T-B,i} = -RT \ln K_i
\]

(13)

The difference in this molar free energy obtained for the real ion partitioning and the “intrinsic” partitioning divided by the charge of the component, is a transfer free energy per charge in the system. This is an electro-chemical driving force that scales with the charge of each component.

\[
\frac{\Delta G_{\text{electro-chemical}}}{RT} = -\frac{(\ln K_{i,\text{real}} - \ln K_{i,\text{intrinsic}})}{z_i}
\]

(14)

or

\[
\ln K_{i,\text{real}} = \ln K_{i,\text{intrinsic}} - \frac{z_i \times \Delta G_{\text{electro-chemical}}}{RT}
\]

(15)

The minimum free energy in Equation 5 is searched under the condition that Equation 14 gives the same value for all charged components and that Equations 6 and 7 are satisfied.

3. Materials and Methods

3.1. Materials

PEG 2000 (molecular weight range 1,900–2,200 g/mol) and PEG 4000 (3,500–4,500 g/mol) were obtained from Merck (Hohenbrunn, Germany), and PEG 8000 (7,000–9,000 g/mol) from Sigma-Aldrich. NaPAA 8000 as 45 wt% pH ca. 7.4 was obtained from Sigma-Aldrich (no data on polydispersity). Adipic acid was obtained from KEBO AB, Stockholm, Sweden. Azelaic acid was obtained from Eastman Kodak Co., Rochester, N.Y. All other chemicals were of pro analysis grade. The di-sodium salts of adipic and azelaic acids were obtained by titrating a water-slurry of the acids with a 50% NaOH solution until pH was between 7.5 and 8.4. The final clear stock solution concentrations of di-sodium adipate and di-sodium azelate were 1.003 and 1.001 mol/kg, respectively.

3.2. Two-Phase Systems and Binodals

For each phase diagram three stock solutions of PEG, NaPAA and salt were prepared. Each solution had the same salt concentration, for instance 3 wt% NaCl or Na₂SO₄. The stock-solutions were mixed in graded 15 mL glass tubes and then centrifuged with a table centrifuge (2,000 rpm, 1 minute). Then
the tube was inspected for formation of two-phases. By titrating with the three stock solutions it is possible to obtain a binodal curve. Generally, the solutions become turbid in the two-phase side of the binodal; however, close to the binodal, the lack of turbidity upon mixing was not a good indication of a one-phase system. This explains the necessity of centrifugation of the systems. For the tie-line determination, corresponding 10 g two-phase systems were made (in triplicate) and were then put in a water-bath for equilibration. The formation of two-phase systems was generally accomplished within 2 min. After the phase separation, the volumes were read and the sample was carefully taken from top-phase using Pasteur-pipettes. Samples from bottom phase were taken after the remaining top-phase had been removed.

3.3. Determination of PEG Concentration

A sample of 1–2 g pure phase (top or bottom) was added to a 15 mL glass beaker. The beaker with the solution was then placed in an oven at 110–150 °C overnight to evaporate the water. About 6 g of warm acetone of high purity (less than 1% water) was added to the dried residue to dissolve the PEG molecules. The acetone-PEG solution was carefully removed from the beaker with a Pasteur-pipette and placed in another glass beaker. Three extractions were made for each residue. The PEG content was then determined gravimetrically by evaporating the acetone.

3.4. Determination of NaCl Concentration

NaCl was determined by the Mohr titration method. Three different aqueous solutions were prepared: Solution 1: 51.97 mmol/kg AgNO₃, solution 2: 5 wt% K₂Cr₂O₇, and solution 3: 0.74 wt% NaHCO₃. One-gram sample from the two-phase systems was mixed in a beaker with 4–5 mL solution 3. Then 5 drops of solution 2 were added to the mixture. Finally, the mixture was titrated with solution 1 until formation of the red precipitate Ag₂Cr₂O₇.

3.5. Determination of Na₂SO₄ Concentration

Na₂SO₄ was determined by precipitation, which was performed by mixing the salt solution with an excessive Ba(NO₃)₂ solution so that all SO₄²⁻ is precipitated as BaSO₄. Since poly(acrylate) is also precipitated by Ba²⁺ (or other multivalent ions), a special procedure was developed. One-gram sample was mixed in a 15 mL plastic tube with 4 mL of an organic solution consisted of 50 wt% acetone and 50 wt% methanol. The mixture became turbid due to a two-phase formation. The top-phase containing the salt was removed and placed in a glass tube. For each sample two organic phase extractions were performed and the organic phases were pooled together (ca. 8 g). Water was added to make the total volume 10 mL. Then 3 mL of a 10 wt% Ba(NO₃)₂ stock solution was added to the glass tube. A white precipitate was immediately formed and centrifuged at 1,500g for 15 minutes in a table centrifuge. The supernatant was discarded and the pellet washed several times from excessive salts by re-suspending it in water and centrifuged again. The glass tube with the precipitate was then dried in an oven overnight. The BaSO₄ was then determined gravimetrically to obtain the original content of Na₂SO₄.
3.6. Determination of NaPAA Concentration

The NaPAA concentration was determined through refractive index measurement of the sample. A sample of the phase was diluted so that the total concentration in the diluted solution of PEG, NaPAA and salt was less than 10 wt%. The refractive index was measured, which gives the total contribution from all components. Previously, linear standard curves were determined separately for aqueous solutions of PEG, salt and NaPAA. The refractive index of the diluted sample was then subtracted by the individual contributions of PEG and salt, and the remaining refractive index gives the concentration of NaPAA through a linear standard curve of refractive index of NaPAA in water. The refractive index was determined with an Abbe type refractometer (OPTECH, optical technology).

4. Results

4.1. Effect of NaCl Concentration

In Figure 1 three binodals of the system composed by PEG 8000 and polyacrylate 8000 with varying amount of NaCl are shown. By increasing salt concentration (1.05%, 3.7% and 6%) the binodal moves closer to the water-rich region, and the changes are larger in the PEG-rich part of the phase diagram. There seems to be a saturation effect of the added salt so that a further increase of the salt concentration does not increase the two-phase region significantly. The straight tie-line shown belongs to the binodal C, and shows a discrepancy between the top-phase composition (PEG-rich phase) and the binodal (for a ternary system the tie-line should connect to the binodal). One explanation can be that the system is not ternary and the mixing point of the tie-line creates a top-phase different from the mixing points of the binodal at the PEG-rich region. The partition coefficient, K, for NaCl was at the mixing point of the tie-line equal to 1.14, i.e., a preference for the PEG-rich phase.

![Figure 1. Phase diagram of the PEG 8000/NaPAA 8000/NaCl system. Binodals A, B and C refer to the systems with 1.05, 3.7 and 6.0 wt% NaCl, respectively. The straight tie-line belongs to binodal C. All binodals determined at 22 °C.](image)
4.2. Effect of Na$_2$SO$_4$ Salt Concentration

In Figure 2 the binodals for systems with different PEG size and Na$_2$SO$_4$ content are shown. The effect of increasing Na$_2$SO$_4$ from 3 to 6 wt% for the PEG 4000/NaPAA 8000 mixture can be seen by comparing binodals B and D. The movement of the binodal to the water region is significant and the changes are more pronounced in the NaPAA-rich region. A straight tie-line for the system with PEG 4000 and 3 wt% Na$_2$SO$_4$ (binodal B) is also shown, and as above, in Figure 1, the tie-line is not connected to the binodal curve as normally found in ternary aqueous two-phase systems. The partition coefficient $K$ for Na$_2$SO$_4$ in the mixing point for the tie-line was 1.0, which is a more even partitioning between the phases than for NaCl. This may be understood by the relatively stronger hydrophilicity of the SO$_4^{2-}$ ion compared to the Cl$^-$ in terms of solvation free energy [23].

**Figure 2.** Phase diagrams of the PEG/NaPAA 8000/Na$_2$SO$_4$ systems. The size of PEG and Na$_2$SO$_4$ concentration has been varied. Binodal A: PEG 2000 and 6 wt% Na$_2$SO$_4$, binodal B: PEG 4000 and 3 wt% Na$_2$SO$_4$, binodal C: PEG 8000 and 3 wt% Na$_2$SO$_4$, binodal D: PEG 4000 and 6 wt% Na$_2$SO$_4$. The straight tie-line belongs to binodal B. All binodals determined at 22 °C.

4.3. Effect of Polymer Molecular Weight

The effect of PEG size on the position of the binodal can be seen in Figure 2, where binodal A (PEG 2000, 6 wt% Na$_2$SO$_4$) should be compared with binodal D (PEG 4000, 6 wt% Na$_2$SO$_4$) and binodal B (PEG 4000, 3 wt% Na$_2$SO$_4$) with binodal C (PEG 8000, 3% Na$_2$SO$_4$). In Figure 3 the binodals for systems containing the hydrophobic divalent salt di-sodium azelate are shown. In both of these examples the binodals move as expected to the water rich region as the molecular weight of PEG increases. The changes are more pronounced at the NaPAA rich region.
Figure 3. Phase diagrams of the PEG/NaPAA 8000/di-sodium azelate (0.211 mol/kg) systems. Binodals A and B refer to PEG 4000 and PEG 8000, respectively. All binodals determined at 22 °C.

4.4. Effect of Salt Type on the Binodal

In Figure 4, the effect of increasing the anion hydrophobicity of the added salt on the binodal curve is shown. The added salts are Na₂SO₄, di-sodium adipate and di-sodium azelate. Interestingly, the binodal changes are significant only at the PEG-rich region. At the NaPAA rich region there is a very weak tendency for an increasing miscibility of PEG and NaPAA.

Figure 4. Phase diagrams of the PEG 8000/NaPAA 8000/divalent salt systems. The concentration of the divalent salt is 0.211 mol/kg. Binodal A Na₂SO₄. Binodal B: di-sodium adipate. Binodal C: di-sodium azelate. All binodals determined at 22 °C.

5. Discussion

5.1. Effect of Salt Concentration

As mentioned in the Introduction, the salt decreases the entropy penalty of compartmentalization of the polyelectrolytes. The more salt ions are added to the system, the stronger is the effect. A saturation
effect upon salt addition in terms of decreased amount of the one-phase region may occur when the salt concentration is high enough that the polyelectrolyte effectively behaves as an uncharged polymer. The effect of increasing the two-phase region upon increased salt addition is modeled as in Figure 5 and all parts of the binodal curve are affected. The saturation effect of the salt observed in the experimental system is however not seen in the model. This is due to the fact that the modeled ions in that case are athermic, which means that the only non-zero enthalpy comes from endothermic repulsion between the polymers. In the experimental case, however, the attractive ion-water solvation interaction may possibly partly counteract two-phase formation by driving the polymers to associate. The relative influence of the polyelectrolyte counter ions (Na\(^+\)) becomes less important if more of the smaller ions are added to the system (e.g., NaCl), since the latter will even out the ion concentration difference between the phases and thus entropically facilitate compartmentalization of the polyelectrolyte. This effect is stronger for divalent than monovalent ions. This allows a separation at lower polymer concentration, which also means that a smaller enthalpic repulsion between the polymers is enough to overcome the effect of the mixing entropy.

Figure 5. Binodal curves for calculated phase diagrams, showing the entropic effect of salt addition. Filled circles, triangles and diamonds: systems containing salt of the Na\(_2\)SO\(_4\) type, but having athermic interaction; empty circles, triangles and diamonds: salt of the NaCl type (athermic). Filled and unfilled squares define the binodal of the experimental system with 1.05 wt% NaCl and polymer concentrations in wt%. Apart from the experimental system, all concentrations are given in volume percent. Concentration of salts (in wt% or volume %): diamonds (1.05), circles (3), triangles (6). Temperature 25 °C. Interaction parameters: for PEG-polyacrylate, \(w_{23}\) is 2,400 J/mol. All small cation and anion interactions were set equal to zero. Flory-Huggins polymerization degree: for PEG is 404, for polyacrylate 335, the value for water and all ions was 1. The polyacrylate polymer has a charge of \(-85\). The concentration of the polyelectrolyte is given as a sum of the polyelectrolyte plus sodium ions for electroneutral balance.

5.2. Effect of Polymer Size

The Flory-Huggins theory [18] predicts a linear dependence on the partition driving force (both enthalpic and entropic contributions) accordingly to the relationship
\[ \ln K \propto M \] (16)

where \( M \) is the polymerization degree of the polymer. The changes in binodal position are shown in Figure 6, where the modeled system contains a “\( \text{Na}_2\text{SO}_4 \)” type salt. The changes in the calculated binodals follow the same tendency as in the experiment (a stronger change at the NaPAA rich region). The position of the calculated binodals has been fitted to approximate the corresponding binodals of the experimental systems (Binodals B and C in Figure 2). The discrepancy between the modeled and experimental is due to a single-parameter fit (PEG-\( \text{SO}_4^{2-} \) interaction) in order to approximate both shape and position of the binodal for different salt concentrations.

**Figure 6.** Binodal curves calculated for the PEG/NaPAA 8000/\( \text{Na}_2\text{SO}_4 \) (3 wt%). Effect of PEG size. Symbols refer to systems with different PEG polymerization degree given in brackets: diamonds (101) PEG 2000, squares (202) PEG 4000, and triangle (404) PEG 8000. The PEG-polyacrylate interaction \( w_{23} \) equals 2,400 J/mol. The PEG-\( \text{SO}_4^{2-} \) interaction \( w_{25} = 27,000 \) J/mol. All other interactions of the small ions were set equal to zero. Temperature 25 °C. The polyacrylate polymer has a charge of \(-85\) and a degree of polymerization of 335. The concentration of the polyelectrolyte is given as a sum of the polyelectrolyte plus sodium ions for electroneutral balance.

5.3. **Effect of Salt Type**

As pointed out in the Introduction, Gupta et al. [15] demonstrated the influence of the salt type on the binodal position of the PEG-polyacrylate system. They explained the effect of ions in terms of water structure breaking or structure making. We present here a more simple and fundamental effect, which is supported by the Flory-Huggins theory. The experimental effect shown in Figure 4, where an increase of the anion hydrophobicity of the added salt moved the binodal to the water-rich corner (increasing the two-phase region) has been reproduced with the model in Figure 7 below. A modeled “hydrophilic salt” of the \( \text{Na}_2\text{SO}_4 \) type gives a binodal far from the water rich region. In order to avoid multi-parameter changes, the salt is made “hydrophilic” simply by letting the \( \text{SO}_4^{2-} \) anions have a strong repulsion to PEG. The comparing “hydrophobic” anion is then modeled by setting this interaction equal to zero (athermic interaction). This causes strong changes in the calculated binodal, an effect that was also found in the experimental case (Figure 4) where the hydrophobic ion is azelate.
The explanation of this effect (displayed in Figure 7) can be understood from the mean field model calculation by considering the Na\(^+\)-ion partitioning. There is a tendency for the small hydrophilic ion to even out the concentration between the phases. As counter ion the strongly hydrophilic SO\(_4^{2-}\), which has a strong repulsion to PEG, cannot distribute itself to the PEG-phase as strong as the intrinsic partitioning preference of Na\(^+\), so a significant amount of polyacrylate takes the counter ion role. When a more hydrophobic divalent anion is used instead, then by lacking a strong repulsion to PEG, it will energetically favor the transfer of Na\(^+\)-ions to the PEG-phase, which in turns allows a polyacrylate to contribute less to this partitioning process, which allows a larger two-phase region. This effect is interesting from a technical point of view since a formation of two-phase systems can be achieved with less amount of polymer. The use of di-sodium azelate as a two-phase inducing salt may be questioned if the two-phase system is intended to be used in bioseparations process, since the alkane dicarboxylates have surfactant-like properties and may denature proteins. However, this type of surfactant has relatively high cmc (>100 mM) and poor hydrophobic solubilization power [24,25].

**Figure 7.** Binodal curve for calculated and experimental PEG 4000/NaPAA 8000/ Na\(_2\)SO\(_4\) 6 wt% system. Square symbols define the experimental system with polymer concentrations in wt% (binodal A). Binodals with circles and diamonds are calculated systems (given in volume %) with PEG-polyacrylate interaction \(w_{23} = 2,400\) J/mol. Line B shows system with PEG-SO\(_4^{2-}\) interaction \(w_{25} = 27,000\) J/mol. Line C shows the system with all salt interactions equal to zero (athermic). Temperature 25 °C. Flory-Huggins polymerization degree is for PEG 202, for polyacrylate 335, and the value for water and all ions is 1. The polyacrylate polymer has a charge of –85. The concentration of the polyelectrolyte is given as a sum of the polyelectrolyte plus sodium ions for electroneutral balance.

![Figure 7](image-url)

6. Conclusions

The experimental and calculated phase diagrams of several PEG/NaPAA/salt systems have been determined. The increase in salt concentration or PEG molecular size always leads to an increase in two-phase region as expected for polyelectrolyte-containing systems. The increase in molecular size of PEG moves the binodal more into the PAA rich region. An interesting effect of hydrophobic divalent
anions was observed when these anions cause a stronger increase in the two-phase region at the PEG-rich region than a hydrophilic anion. The Flory-Huggins theory can be used to reproduce the experimental effects.

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References


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