Phase behavior of temperature-responsive polymers with tunable LCST: An equation-of-state approach

Alexei M. Kisselev\textsuperscript{a}, E. Manias\textsuperscript{b,∗}

\textsuperscript{a} Department of Physics, Pennsylvania State University, University Park, PA 16802, USA
\textsuperscript{b} Department of Materials Science and Engineering, 325-D Steidle Bldg, Pennsylvania State University, University Park, PA 16802, USA

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Abstract

The aqueous solution phase behavior of (ethylene oxide)/ethylene copolymers with varying degree of hydrophobicity is explored using an equation of state approach. The general formalism of the lattice-fluid with hydrogen-bonding theory is employed after a minor adjustment to account for multiple types of hydrogen bonds. The theoretical model is shown to be effective in describing the phase behavior of these systems, and the model parameters seem to be transferable between different homologous copolymer series. Despite limitations of the model, the calculated phase diagrams and the dependence of lower critical solution temperature (LCST) on the hydrophilic/hydrophobic content of the polymer show good correspondence with the available experimental data.

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1. Introduction

Aqueous polymer solutions are important from a technological point of view [1], but, at the same time, represent systems whose theoretical description can be challenging [2]. These challenges arise from the fact that systems of molecules interacting with strong specific interactions, such as hydrogen-bonding, deviate remarkably from normal solution behavior. However, there is a strong interest in designing polymer systems – going beyond hydrogels – with a controlled solubility in water, since they could potentially be used in a variety of applications requiring smart/responsive materials, e.g. sensors, actuators, cell patterning, and smart/triggered drug delivery. Temperature-sensitive solubility can be rendered from the existence of the lower critical solution temperature (LCST), above which the polymer becomes insoluble in water. Thus, an analytical model which would describe the LCST-type phase behavior of aqueous polymer solutions in a clear, concise and self-consistent manner and yet having a certain predictive power is very desirable.

Poly(ethylene oxide) (PEO) is historically one of the most investigated water-soluble polymers with an LCST, including considerable theoretical attention in recent years [3–5]. Its LCST has been associated with its ability to hydrogen bond to water, and occurs at temperatures above 100 °C – i.e. above the boiling point of water for ambient pressures – thus limiting its use for applications requiring a temperature-response. However, it has been predicted that the LCST of water-soluble polymer can be altered by controlling the balance of hydrophilic and hydrophobic segments in the polymer chain [6]—e.g. PEO’s LCST can be lowered by proper addition of hydrophobic groups [7]. This approach was tested for various polymers, such as poly(N-isopropyl acrylamide) (PNIPAM) [8], poly(siloxylethylene glycol) [9], random copolymers of ethylene oxide and propylene oxide (EOPO) [10], as well as alternating segmented copolymers of ethylene oxide and ethylene (EO–EE) [11]. A sharp LCST transition, which corresponds to a genuine coil-to-globule transition [12] and which is determined...
by the hydrophilic/hydrophobic balance, dictates a polymer structure where hydrophilic and hydrophobic segments are not lumped together in extended blocks [7,11], since such blockiness facilitates micellar type of aggregates/collapse [13,14]. In this work, we will focus on this type of oligo-ethylene-oxide/oligo-ethylene linear copolymers, which were shown to exhibit LCST phase behavior similar to that of pure poly(ethylene oxide) with a sharp LCST transition tailored at varied temperatures (from 7 to 80 °C) via the copolymer composition [11]. Specifically, focusing on these polymers, we employ a well-established equation of state theoretical framework [2,15] to describe their phase behavior.

From an Equation-of-State theoretical viewpoint, one can treat physical (van der Waals) interactions with the well-established Sanchez–Lacombe lattice-fluid (LF) model [15,16], which is a compressible lattice theory (compressibility effects also give rise to an LCST, in the general case of polymer solutions) and thus can overcome the drawback of incompressible classical Flory–Huggins theory that is inherently unable to account for an LCST [17]. In addition, combining the lattice-fluid approach with the chemical association approach [18], enables the thermodynamic description of hydrogen-bonded polymer solutions. In a model proposed by Panayiotou and Sanchez – known as the lattice-fluid with hydrogen-bonding (LFHB) theory [19]– the physical contributions are treated based on the combinatorial expression for the number of configurations available to ark-mer in the close-packed pure state; ωk is treated as a constant and will be decoupled, i.e., the canonical partition function can be expressed as

\[ Q = Q_{LF}Q_{HB} \] (1)

We define our system to contain r types of molecules, with \( N_k \) molecules of k-th type at temperature T and external pressure P. Also, there are m types of proton-donor groups and n types of proton-acceptor groups, with \( a_{ij}^k \) being the number of donor groups of i-th type in each molecule of k-th type and, equivalently, \( a_{ij}^k \) being the number of acceptor groups of j-th type in each molecule of k-th type.

2. Theoretical model and framework

The specific goals of this study are (a) to apply the LFHB to the LCST phase behavior of (ethylene oxide)/ethylene copolymers, an application which necessitates a minor modification so as LFHB can address multiple types of hydrogen bonds; and (b) to verify the assumption that the phase behavior of these systems is of the same nature as that of PEO, but with an LCST lowered in value by the hydrophobic character of the ethylene units.

2.1. Lattice-fluid

According to the lattice-fluid theory, molecules are arranged on a quasi-lattice of \( N_t \) sites, \( N_0 \) of which are empty. Each molecule of k-th type is divided into \( r_k \) segments of close-packed volume \( v_k^* \) in the pure state and average (mean-field) interaction energy \( \epsilon_k^* \). The total number of lattice sites is then

\[ N_t = rN + N_0 \quad \text{with} \quad r = \sum_{k=1}^{t} r_k x_k, \] (2)

where \( rN \) is the total number of molecular segments in the system and \( x_k \) is the mole fraction of k-th component in the mixture (\( x_k = N_k / \sum_{k=1}^{t} N_k \)). The following combining and mixing rules are assumed

\[ v^* = \sum_{k=1}^{t} \phi_k v_k^* \quad \text{and} \quad \epsilon^* = \frac{1}{2} \left( \sum_{p=1}^{t} \epsilon_p \right) \left( \sum_{k=1}^{t} \sum_{l=1}^{t} \theta_k \theta_l \epsilon_{kl} \right), \] (3)

where \( \phi_k \) are the segment fractions (defined as \( \phi_k = x_k r_k / r \)) and \( \theta_k \) are the surface fractions (defined as \( \theta_k = \phi_k s_k / \sum_{l=1}^{t} \phi_l s_l \)), where \( s_k \) is the average number of contacts per k-th segment, equivalent to a surface to volume ratio of that segment). A Berthelot-type combining rule is adopted for \( \epsilon_{kl} \)

\[ \epsilon_{kl} = 2 \tilde{\epsilon}_{kl} \left( \frac{\epsilon_{kl}^*}{s_k s_l} \right)^{1/2} \] (4)

where \( \tilde{\epsilon}_{kl} \) is a dimensionless parameter, expected to have values close to unity. The total lattice-fluid volume of the system is given by \( V_{LF} = N_k v^* = rNv^* \tilde{v} \), where the reduced volume (\( \tilde{v} = 1 / \tilde{\rho} \)) is defined on the basis of a reduced density (\( \tilde{\rho} = rN / N_t \)). Similarly, the total potential energy of the system, as derived in Refs. [15,19] taking into account only nearest-neighbor interactions and ignoring interactions with empty sites, is given by \( \Delta E_{LF} = rN \tilde{v} \tilde{\epsilon}^* \). On the basis of this definitions the canonical partition function of the physical interactions is given [19] by:

\[ Q_{LF}(T, N_0, \{ N_k \}) = (1 - \tilde{\rho} - N_0 \rho) \prod_{k=1}^{t} \left( \frac{\omega_k}{\phi_k} \right)^{N_k} \exp \left[ - \frac{E_{LF}}{RT} \right] \] (5)

where \( \omega_k \) is the number of configurations available to a \( r_k \)-mer in the close-packed pure state; \( \phi_k \) is treated as a constant and will
cancel out in all applications of interest here (this is not obvious in cases of PEO physisorption on solid surfaces, or in extreme, sub-$R_g$, confinements [22–24]).

2.2. Hydrogen-bonding

The interaction energies due to hydrogen-bonding contribution are in excess of the physical interactions (which are accounted for by the lattice-fluid part of the partition function). For the hydrogen bond between a donor of the $i$-th type and an acceptor of the $j$-th type, we define $E_{ij}$ to be the favorable energy change upon hydrogen bonding, $S_{ij}^0$ the entropy loss associated with the $(i,j)$ bond formation, and $V_{ij}^0$ the respective volume change. In the general case, there are $N_{ij}$ bonds of the $(i,j)$ type, thus the total hydrogen-bonding energy of the system is

$$E_{HB} = \sum_{i=1}^{m} \sum_{j=1}^{n} N_{ij} E_{ij}^0$$  \hspace{1cm} (6)

The number of ways of distributing the $N_{ij}$ bonds among the functional groups of the system is given by the combinatorial expression corrected by the mean-field probability of bond formation, as derived in Ref. [19]:

$$\Omega = \prod_{i=1}^{m} \frac{N_{i}!}{N_{i0}!} \prod_{j=1}^{n} \frac{N_{j}!}{N_{j0}!} \prod_{i,j} P_{ij}^{N_{ij}} N_{ij}!$$  \hspace{1cm} (7)

where $N_{ij}$ is the total number of donor groups of $i$-th type ($N_{i} = \sum_{k=1}^{i} a_{ik}$) and $N_{i0}$ is the total number of acceptor groups of $j$-th type ($N_{j} = \sum_{k=1}^{j} a_{jk}$), leaving $N_{i0}$ number of unbonded donors of $i$-th type and $N_{j0}$ number of unbonded acceptors of $j$-th type ($N_{i0} = N_{i} - \sum_{j=1}^{n} N_{ij}$ and $N_{j0} = N_{j} - \sum_{i=1}^{m} N_{ij}$), and $P_{ij}$ is the mean-field probability of $(i,j)$ bond formation

$$P_{ij} = \frac{\rho}{rN} \exp \left[ \frac{S_{ij}^0}{R} \right]$$  \hspace{1cm} (8)

Following this formalism the canonical partition function for hydrogen bonding can be written as [19]

$$Q_{HB}(T, N_0, \{N_k\}) = \frac{\sum_{\{N_{ij}\}} \rho^{\sum_{ij} N_{ij}} \prod_{i=1}^{m} \prod_{j=1}^{n} \frac{N_{i}!}{N_{i0}!} \frac{N_{j}!}{N_{j0}!} \prod_{j=1}^{n} \frac{1}{N_{ij}!} \exp \left[ -N_{ij}(E_{ij}^0 - TS_{ij}^0) \right]}{rN^{\sum_{i=1}^{m} \sum_{j=1}^{n} N_{ij}}}}$$ \hspace{1cm} (9)

Various hydrogen-bonding fractions are defined by

$$v_{ij} = \frac{N_{ij}}{rN}, \quad v_{ij0} = \frac{N_{i0}}{rN}, \quad v_{ij0} = \frac{N_{j0}}{rN}, \quad v_{ij} = \frac{N_{i}^j}{rN}$$ \hspace{1cm} (10)

2.3. Free energy and equations of state

The Gibbs free energy in a framework of the lattice fluid theory with hydrogen-bonding (LFHB), developed by Panayiotou and Sanchez [19], consists of two terms: lattice-fluid and hydrogen-bonding

$$G = G_{LF} + G_{HB}$$  \hspace{1cm} (11)

which follows immediately from the expression for the Gibbs partition function given by

$$\Psi(T, P, \{N_k\}) = \sum_{N_0=0}^{\infty} Q_{LF}(T, N_0, \{N_k\}) Q_{HB}(T, N_0, \{N_k\})$$ \hspace{1cm} 

$$\times \exp \left[ \frac{-PV}{RT} \right]$$ \hspace{1cm} (12)

with $G = -kT \ln \Psi$ and $V$ is the system volume

$$V = rNv^a + \sum_{i=1}^{m} \sum_{j=1}^{n} N_{ij} V_{ij}^0$$  \hspace{1cm} (13)

While the reduced pressure ($\tilde{P}$) and reduced temperature ($\tilde{T}$) are:

$$\tilde{P} = \frac{P}{\rho e^*} \quad \text{and} \quad \tilde{T} = \frac{T}{e^*}$$  \hspace{1cm} (14)

the lattice-fluid contribution can now be expressed by

$$\frac{G_{LF}}{k_B T} = rN \left\{ -\tilde{P} + \frac{\tilde{P} \tilde{V}}{\tilde{V}} + (\tilde{V} - 1) \ln(1 - \tilde{\rho}) \right\}$$

$$+ \frac{1}{r} \ln \tilde{\rho} + \sum_{k=1}^{i} \frac{\Phi_k}{\omega_k} \ln \frac{\phi_k}{\omega_k}$$ \hspace{1cm} (15)

and the hydrogen-bonding contribution is given by

$$\frac{G_{HB}}{k_B T} = rN \left\{ \sum_{i=1}^{m} \sum_{j=1}^{n} \frac{v_{ij}}{v_{ij0}} \ln \frac{v_{ij0}}{v_{ij}} + \sum_{j=1}^{n} \frac{v_{ij}}{v_{ij0}} \ln \frac{v_{ij0}}{v_{ij}} \right\}$$ \hspace{1cm} (16)

The Gibbs partition function was evaluated in the usual way by using the maximum term approximation [25]. This is equivalent to using the generic term in (12) for the Gibbs free energy (substituting the factorials with the Stirling approximation) and implying the minimization conditions with respect to $\tilde{\rho}$ and $N_{ij}$

$$\frac{\partial G}{\partial \tilde{\rho}} \bigg|_{T,P,\{N_k\},\{N_{ij}\}} = 0 \quad \text{and} \quad \frac{\partial G}{\partial N_{ij}} \bigg|_{T,P,\{N_k\},\{N_{ij}\}} = 0$$ \hspace{1cm} (17)

The first condition of (17) yields the equation of state for the reduced density

$$\tilde{\rho}^2 + \tilde{\rho} + \tilde{T} \left[ \ln(1 - \tilde{\rho}) + \tilde{\rho} \left( 1 - \frac{1}{\tilde{\rho}} \right) \right] = 0$$ \hspace{1cm} (18)

where $1/\tilde{\rho} = 1/r - v_{ij}$ and $v_{ij} = \sum_{i=1}^{m} \sum_{j=1}^{n} v_{ij}$. The second condition in (17) yields a system of equations for the fractions
of hydrogen bonds in the system

\[
\frac{v_{ij}}{v_{0i}v_{0j}} = \tilde{\rho} \exp \left[ -\frac{G_{ij}^0}{RT} \right]
\]
(19)

where \( G_{ij}^0 = E_{ij}^0 + PV_{ij}^0 - TS_{ij}^0 \). The reduced density equation of state (18) and the H-bond fractions equations (19) are used together and serve as a system of equations of state

\[
\left\{ \begin{aligned}
\tilde{\rho}^2 + P + \tilde{T} \left[ \ln(1 - \tilde{\rho}) + \tilde{\rho} \left( 1 - \frac{1}{\tilde{T}} \right) \right] &= 0, \\
\frac{v_{0i}v_{0j}}{\tilde{\rho}} - v_{ij} \exp \left[ -\frac{G_{ij}^0}{RT} \right] &= 0
\end{aligned} \right. \]
(20)

Chemical potential of k-th component is obtained as follows

\[
\mu_k = \left( \frac{\partial G_{LF}}{\partial N_k} \right)_{T,P,N_j,\{N_{ij}\}} + \left( \frac{\partial G_{HB}}{\partial N_k} \right)_{T,P,N_j,\{N_{ij}\}}
\]
(21)

and is given by

\[
\frac{\mu_k}{k_B T} = \ln \phi_k + \left( 1 - \frac{r_k}{r} \right) \phi_k
\]

\[
+ r_k \tilde{\rho} \left( \sum_{i < \ell} \frac{s_k}{s_i} X_{ik} + \sum_{i > j} \frac{\theta_i \theta_j}{s_i} X_{ij} \right)
\]

\[
+ r_k \left( \frac{-\tilde{\rho}_0 + P_k \tilde{\rho}}{T_k} + (\tilde{\rho} - 1) \ln(1 - \tilde{\rho}) + \frac{1}{r_k} \ln \frac{\tilde{\rho}}{\omega_k} \right)
\]

\[
+ r_k \sum_{m=1}^{n} \sum_{j=1}^{m} v_{ij} \frac{d^k \ln \frac{v_{ij}}{v_{0j}} - \sum_{j=1}^{n} a_j \frac{d^k \ln \frac{v_{ij}}{v_{0j}}}{d^k v_{0j}}}.
\]
(22)

where \( X_{ij} \) is the reduced average interaction between k-mer and \( i \)-mer and \( P_k = P/P_i^* = P v_i^*/\epsilon_i^* \) and \( T_k = T/T_i^* = RT/\epsilon_i^* \).

2.4. Binary mixture

Focusing on the phase behavior of a polymer–solvent binary mixture (index 1 denotes solvent, and index 2 polymer) \( r \) becomes \( r = r_1, x_1 + r_2, x_2 \), and the mole fractions (\( x_i \)) and the corresponding segment fractions (\( \phi_i \)) are simply:

\[
x_2 = 1 - x_1 = \frac{N_2}{N_1 + N_2} \quad \text{and} \quad \phi_2 = 1 - \phi_1 = \frac{r_2 x_2}{r}
\]
(23)

and the surface fractions become

\[
\theta_2 = \frac{\phi_2}{\phi_2 + \phi_1(s_1/s_2)} \quad \text{and} \quad \theta_1 = 1 - \theta_2 = \frac{\phi_1}{\phi_1 + \phi_2(s_2/s_1)}
\]
(24)

where \( s_1/s_2 \) is ratio of the surface area per unit characteristic volume for solvent and polymer and can be calculated/approximated by a hard-sphere model or via van der Waals radii. The mixing and combining rules simplify to

\[
v^* = \phi_1 v_i^* + \phi_2 v_j^* \quad \epsilon^* = \phi_1 \epsilon_i^* + \phi_2 \epsilon_j^* - \phi_1 \theta_2 R T x_{12}.
\]
(25)

where \( x_{12} = (\epsilon_i^* + (s_1/s_2) \epsilon_j^*) - 2(s_1/s_2)^{1/2} \epsilon_i^{1/2} \epsilon_j^{1/2} / R T \) with \( \epsilon_i^{1/2} = \xi_{12}^{1/2} / \epsilon_i^{1/2} \epsilon_j^{1/2} \). The dimensionless parameter \( \xi_{12} \) is the only free parameter of the model, and is expected to have values close to unity.

In the binary case of the model, the phase behavior can be determined in terms of two independent variables – composition \( x_2 \) and temperature \( T \) – along with a number of parameters, including external pressure \( P \). External pressure is always kept constant in our considerations, thus it is treated as a parameter rather than a variable. Other parameters of the model include, from the lattice-fluid part: average interaction energies \( \epsilon_1^*, \epsilon_2^* \), close-packed volumes \( v_1^*, v_2^* \), and numbers of segments \( r_1, r_2 \); and, from the hydrogen-bonding part, for each \( (i, j) \) bond type: favorable energy change \( E_{ij}^0 \), entropy loss \( S_{ij}^0 \), and volume change \( V_{ij}^0 \). All these parameters can be obtained from experimental PVT data, from mixing rules, and from hydrogen-bonding interactions data, all of which can be found in literature for a variety of components [25].

The coexistence curve (binodal) – depicting the composition of coexisting phases at different temperatures – is obtained by the typical thermodynamic stipulation that the chemical potential of both component is the same in all coexisting phases, i.e. at a given temperature \( T \) mole fraction points \( x_1^* \) and \( x_2^* \) belong to the binodal curve if the following conditions is satisfied

\[
\mu_1^A - \mu_1^B = 0, \quad \mu_2^A - \mu_2^B = 0
\]
(26)

To find a pair of binodal points one needs to solve (26) and the system of equations of state (20) simultaneously. The corresponding spinodal curve, which separates the region of thermodynamic instability from the region of metastability and at a given temperature \( T \), is determined by

\[
\frac{d\mu_2}{dx_2} = 0
\]
(27)

Consequently, it is required to solve simultaneously (27) with the system of equations of state (20) and the system of first composition derivatives \( d/dx_2 \) of the equations of state. The extremum point of the spinodal curve, corresponding to the critical point of the system, is given by

\[
\frac{d^2 \mu_2}{dx_2^2} = 0
\]
(28)

Finding a critical point is equivalent to the simultaneous solution of (28) with the system of equations of state (20) along with first \( d/dx_2 \) and second \( d^2/dx_2^2 \) composition derivatives of the equations of state, which adds up to a rather large system of nonlinear equations. The system of this size and complexity cannot be solved analytically even in relatively simple cases, thus requiring a numerical approach to the finding of the solution. The explicit form of the equations for binodal, spinodal and critical point are provided in Supplemental Materials, whereas the numerical solution procedure is outlined in Appendix A.
### 3. Results and discussion

#### 3.1. PEO/water model

Probably the most investigated polymer with an LCST in water solution is poly(ethylene oxide) (PEO). It has been established, both experimentally and theoretically, that the phase behavior of PEO in water is determined by the hydrogen bonding balance between water and PEO molecules. Let us first consider an application of LFHB theory to the case of water/PEO mixture. Each OH group acts as both a donor and an acceptor, thus a water molecule has $d_1^1 = 2$ donors and $a_1^1 = 2$ acceptors. Let us denote the number of –O– groups per PEO molecule with $a$, hence the PEO molecule has $a_2^2 = a$ acceptors, where $a$ is the degree of polymerization of PEO. There will be two types of H-bonds in this system (1, 1) and (1, 2), water–water and water–polymer correspondingly. Chemical potential of the polymer $\mu_2$ and of water $\mu_1$ can be obtained from (22), by rewriting for the binary case of two types of hydrogen bonds. There will be three equations of state—one for the reduced density and one for the H-bond fraction of each type $v_11$ and $v_12$ (20).

Lattice-fluid and hydrogen-bonding parameters of the model, shown in Tables 1 and 2, are obtained from the literature [19,26,27]. Tabulated data can be translated into the terms of LFHB model via obvious relationships

$$
\epsilon_k^* = RT_k^*, \quad \psi_k^* = \frac{\epsilon_k^*}{P_k^*}, \quad r_k = \frac{M_{ak}}{\rho_k^* h_k^*}
$$

where $M_{ak}$ is the molecular weight of k-th component. The ratio of surface areas per unit characteristic volume for water and ethylene oxide can be calculated based on hard-sphere model to be $(s_1/s_2)^{EO} = 1.3424$. The dimensionless variable $\xi_{12}$ is treated as a free parameter of the model.

Finding the pair of binodal points at given temperature $T$ involves solving the system of 8 nonlinear equations, finding a concentration and a temperature of a spinodal point is equivalent to solving the system of 7 equations, and the critical point can be found as a solution to the system of 11 equations, solved numerically as described in Appendix A.

At the limit of high molecular weights the phase behavior of polymers becomes almost molecular weight independent, which allows us in the first approximation to disregard the effects of the molecular weight, i.e. keep $M_w$ in all of our calculations in this section constant and equal to $6 \times 10^5$, the approximate molecular weight of $m = 13$, $n = 6$ ester copolymer [11]. Following this approach, the phase diagram of $M_w = 10^5$ PEO [28] was fitted with the calculated PEO binodal using the numerical procedure described in Appendix A, yielding a value for the dimensionless interaction parameter $\xi_{12}^{EO} = 1.0472$, the experimental and the calculated phase diagrams are shown in Fig. 1a. This value of $\xi_{12}^{EO}$ was used in later calculations of the phase diagrams of our “polyester” and “polyamide” ethylene oxide copolymers without any further change.

#### 3.2. P(EO–EE)/water model

The polymers of interest are based on a linear alternating sequence of oligo($m$)-EO and oligo($n$)-EE [11] with

\[\begin{array}{cccc}
(i, j) & E_{ij}^0 \text{(J/mol)} & S_i^0 \text{(J/(mol K))} & V_{ij}^0 \text{(m^3/mol)} \\
\text{OH} \cdots \text{OH} & (1, 1) & -1.52 \times 10^4 & -16.6 & -4.2 \times 10^{-6} \\
\text{OH} \cdots \text{O} & (1, 2) & -1.42 \times 10^4 & -16.0 & -8.5 \times 10^{-7} \\
\text{OH} \cdots \text{C} \equiv \text{O} & (1, 3) & -1.60 \times 10^4 & -15.8 & -8.5 \times 10^{-7} \\
\text{N} \cdots \text{H} \cdots \text{OH} & (2, 1) & -1.25 \times 10^4 & -7.8 & -8.5 \times 10^{-7} \\
\end{array}\]

![Fig. 1. (a) Experimental coexistence curves fitted with calculated binodals (solid line); (b) LCST dependence on hydrophilic/hydrophobic balance $m/n$ for the general case of [$(EO)_m$–$(EE)_n$] copolymer as described in Section 3.2.](image-url)
the aim of tailoring polymer LCST by controlling the hydrophilic/hydrophobic balance \((m/n)\). These copolymers exhibit phase behavior that appears similar to that of PEO, but have an LCST at substantially lower temperatures. One of the goals of this study is to see whether the LFHB model of PEO as presented above can be extended to describe the phase behavior of the \((|EO|_m - |EE|_n)\) copolymer, which would imply that the copolymer phase behavior can be considered similar in nature as the aqueous phase behavior of PEO only temperature-shifted by the hydrophobic EE group contributions. This can be done by introducing the following simple combination rule (the weighted average) for the EO and EE dimensionless parameters

\[
\xi_{12} = \frac{m}{m + n} \xi_{12}^{EO} + \frac{n}{m + n} \xi_{12}^{EE},
\]

\[
s_1 = \frac{m}{m + n} \left( \frac{s_1}{s_2} \right)^{EO} + \frac{n}{m + n} \left( \frac{s_1}{s_2} \right)^{EE},
\]

\[
s_2 = \frac{m}{m + n} \left( \frac{s_1}{s_2} \right)^{EO} + \frac{n}{m + n} \left( \frac{s_1}{s_2} \right)^{EE}.
\]  

(30)

The ratio of surface areas per unit characteristic volume for water and ethylene can be calculated based on the hard-sphere model \((s_1/s_2)_{PEO} = 1.3266\), and \(\xi_{12}^{EE}\) can be obtained from the best fit of the experimental phase diagram of one of the P(EO–EE) copolymers. Replacement of EO segments with EE segments decreases the number of hydrogen bonding sites per polymer molecule, effectively reducing the hydrogen bonding interactions, which is expected to result in lowering of the LCST. The model produces satisfactory results in terms of describing the phase behavior of the series of copolymers as a function of hydrophilic/hydrophobic balance \(m/n\) with no further modifications (Fig. 1b), simply by scaling the number of hydrogen bonding sites per polymer molecule to equal to \(a_2^2 = (a_2^2)_{PEO} m/(m + n)\), where \(a_2^2_{PEO} = M_w^{poly}/M_w^{PEO}\).

The application of the model to P(EO–EE) was tested in the most simple case, which ignores the presence of the linkage groups and accounts for the difference between PEO and P(EO–EE) by using the combination rule for dimensionless parameters and by appropriate rescaling of the number of the polymer hydrogen bonding sites as described above. The dimensionless interaction parameter \(\xi_{12}^{EE} = 1.0860\) was obtained from the fit of the experimental coexistence curve of \(m = 13\), \(n = 6\) ester copolymer [11] with the LFHB binodal. This value of \(\xi_{12}^{EE}\) arguably contains implicit contributions from the ester groups that link the EE and EO sequences. Despite the crudeness of this first approximation, LFHB can capture the trends of the experimental data for the dependence of the calculated LCSTs on the hydrophilic/hydrophobic ratio \(m/n\) for these ester copolymers. The phase behavior of other homologous series of EO/EE copolymers containing amide groups can be described in the same manner, but will obviously require obtaining different value for the \(\xi_{12}^{EE}\) parameter (since now this parameter will contain implicitly contributions from the amide linkage groups). Given that the differences in phase behavior of these homologous polyester and polyamide copolymers should be explained by accounting for the differences between the respective linkage groups, it is more desirable to extend the model to include the linkage groups explicitly by introducing a more rigorous H-bonding counting scheme, and fix the \(\xi_{12}^{EE}\) parameter to be the same across all EE-containing copolymers with different linkage chemistries.

### 3.2.1. Polyester EO–EE copolymers

\[
\left\{ \begin{array}{c}
\text{CH}\_2\left(O\text{CH}_{2}\text{CH}_3\right)_m\text{CH}_2\text{C}^\text{\textdegree}\text{O}\text{O}^\text{\textdegree}\text{C}\_n\text{CH}_2\text{CH}_3\_N^\text{\textdegree}\text{C}\_n
\end{array} \right\}
\]

(31)

The ester oxygen (C=O) in the polyester copolymers (31) introduces an additional type of acceptor to the model. Now there are three types of H-bonds: \((1, 1), (1, 2), (1, 3)\)—water–water, water–polymer(ether oxygen, –O–), water–polymer(ester oxygen, C=O). Each water molecule has \(d_1^1\) = 2 donors and \(a_1^1\) = 2 acceptors of the first type, each P(EO–EE) molecule has \(a_2^2\ = m \cdot N\) acceptors of the second type, equal to the number of EO segments, and \(a_3^2\ = 2 \cdot N\) acceptors of the third type, equal to the number of C=O groups. We can assume that the van der Waals contribution from the C=O groups to the lattice-fluid part of the model will be contained in the contribution from EE segments and thus can be accounted for by the binodal fitting through the combination rule (30). This leaves the lattice-fluid part unchanged, except for the modified value of the dimensionless parameters \(\xi_{12}^{EE}\), but it introduces one additional term to the hydrogen bonding part of the chemical potential and an additional fourth equation to the system of the equations of state. The critical point can be now found as a solution to the system of 14 equations.

Lattice-fluid and hydrogen-bonding parameters of the model from Tables 1 and 2 can be used in the model with the Eqs. (29) and (30). The ratio \((s_1/s_2)_{PEO} = 1.3266\) was calculated based on hard-sphere model, and the dimensionless parameter \(\xi_{12}^{EE}\) can be obtained from fitting the experimental phase diagram of polyester; for example, \(\xi_{12}^{EE} = 1.0537\) is obtained by fitting experimental phase diagram for \(M_w = 6 \times 10^5\), \(m = 13\), \(n = 6\) polyester with the binodal calculated for the EO–EE copolymer with the ester linkage group, the result of the fit is shown in Fig. 1a and appears to be reasonably close to the experimental phase diagram [11]. The dependence of the LCSTs of polyester series on \(m/n\) ratio shown in Fig. 2 was calculated with the aforementioned values of parameters, varying only \(m\) and \(n\). The calculated trend exhibits the same behavior as shown by the experimental data [11]. One can see that the curve levels off at large values of \(m/n\), which corresponds to the polymer phase behavior being dominated by EO segments, asymptotically approaching the LCST value of pure PEO.

### 3.2.2. Polyamide EO–EE copolymers

\[
\left\{ \begin{array}{c}
\text{CH}\_2\left(O\text{CH}_{2}\text{CH}_3\right)_m\text{CH}_2\text{C}^\text{\textdegree}\text{O}\text{O}^\text{\textdegree}\text{C}\_n\text{CH}_2\text{CH}_3\_N^\text{\textdegree}\text{C}\_n
\end{array} \right\}
\]

(32)

Similarly to the esters, when amide linkage groups (32) are introduced in the EO–EE copolymer, the N–H group brings in an additional type of donor, in addition to the ester C=O discussed above. Now, each P(EO–EE) molecule has \(d_2^2 = 2 \cdot N\) donors of the second type, equal to the number of N–H groups, \(a_2^2 = m \cdot N\) acceptors of the second type, equal to the num-
ber of EO segments, and \( a_2^2 = 2 \cdot N \) acceptors of the third type, equal to the number of C=O groups. Polymer–polymer hydrogen bond formation has a relatively small probability and can be ignored in the case of dilute solutions, thus we consider four types of H-bonds: (1, 1), (1, 2), (1, 3), (2, 1)—water–water, water–polymer(–O–), water–polymer(C=O), polymer(N–H)–water. As in the case of the polyester, this modification leaves the lattice-fluid part unchanged, but introduces two additional terms to the hydrogen bonding part of the chemical potential and results in two additional equations to the system of the equations of state. The critical point can now be found as a solution to the system of 17 equations.

In this manner, after the introduction of this additional hydrogen-bonding, the previous LFHB is used with the same values as for the esters and with no further fitting, i.e., with the parameter values from Tables 1 and 2, and the same pair of \( \xi_{12} \) parameters, which were obtained from the binodals of PEO and of the homologous polyester. The LCST versus \( m/n \) curve calculated for the copolymers with the amide linkage group is shown in Fig. 3 and exhibits excellent correlation with the experimental data [11]. This agreement is despite the fact that none of the parameters in the LFHB model were recalculated or adjusted for this system, which, to a certain extent, proves the consistency of the model and the validity of the assumption that the EO–EE copolymer phase behavior can be accounted through a “superposition” of the EO and EE aqueous phase behaviors.

### 3.3. Effect of molecular weight

To further test the validity of the model and of our assumptions, the influence of other parameters of the model was also investigated. In particular, we checked the dependence on polymer molecular weight, since the model has a complex explicit dependence on \( r \), and consequently on the size of the polymer (i.e., the number of segments on the quasi-lattice is dependent upon the polymer size). Focusing on PEO homopolymers, the calculated LCST values as a function of the \( M_w \) are plotted in Fig. 4a, and are compared with experimental values...
Fig. 4. (a) LCST dependence on molecular weight $M_w$ for PEO, and EO-EE polyester and polyamide copolymers. (b) LCST as a function of hydrophilic/hydrophobic balance $m/n$ for polyester series with explicit $M_w$ dependence (the star symbols are calculated LCST accounting for the experimental $M_w$ of each system).

[28,29] showing a decent qualitative agreement for the trend of polymer size on the PEO LCST. Specifically, the phase diagram of $M_w = 10^5$ PEO [28] was fitted with the calculated PEO binodal setting $\xi_{12}^{\text{EO}} = 1.0464$ (the fit of PEO coexisting curve for $M_w = 1.02 \times 10^5$ [29] returned a similar value $\xi_{12}^{\text{EO}} = 1.0467$). In addition, after obtaining the corresponding $\xi_{12}^{\text{EE}} = 1.0598$ as before, from the fit of $M_w = 5.75 \times 10^5$, $m = 13$, $n = 6$ polyester coexistence curve, the critical temperatures of the polyester copolymers were also calculated for the molecular weight values reported for each polymer [11], i.e., $M_w = 3.80 \times 10^5$ for $m = 13$, $n = 3$, $M_w = 2.45 \times 10^5$ for $m = 5$, $n = 3$, and $M_w = 3.00 \times 10^5$ for $m = 4$, $n = 3$.

The calculated LCST values, shown in Fig. 4b, exhibit excellent agreement with the experimental data, compared to the previous case which disregarded the molecular weight dependence (cf. line in Fig. 4b). Along the same lines, we also show in Fig. 4a the molecular weight dependence of the EO–EE copolymers with ester and amide linkage groups. As expected [7], there is a weaker $M_w$ dependence on the LCST for the copolymers compared to PEO, which is of importance for the design and synthesis of these systems, but which has not been systematically verified experimentally.

Finally, the experimental cloud point temperatures reported in Ref. [11] are in fact measured for 1 wt% polymer solutions and do not strictly correspond to the LCST values (although $\phi_p = 1$ wt% lies in the near vicinity of the critical concentration for the reported molecular weights of the copolymers). Thus it makes some sense to compare those experimental data against the binodal temperatures at a weight fraction of polymer equal to 0.01 and for the experimentally reported $M_w$, rather than against against the LCST of a representative $M_w$ (cf. Fig. 2). This comparison is shown in Fig. 5 for the ester copolymers, showing a very good agreement with the experimental data.

4. Concluding remarks

The LFHB model in the form of the equations of state is suitable for describing the thermodynamic properties of polymer solutions over an extended range of external conditions. Unlike many alternative approaches, which ignore the water–water interaction, the present model takes into account both water–polymer and water–water H-bond formation, however the orientation of donor and acceptor sites with respect to each other is not accounted for explicitly, due to the purely statistical nature of the chemical part of the model. The model retains lattice-fluid mean-field character and inherent drawbacks, which makes it impossible to account for difference between alternating, random, and especially highly segmented or block copolymers, or account for end-group effects (all of which are extremely important for the phase behavior of the copolymers in question).

Despite these drawbacks, the LFHB model, as applied here with minor alterations to account for multiple hydrogen bonding types, reproduced the experimentally observed phase behavior
for aqueous solutions of PEO, and EO–EE ester and amide copolymers. In particular, it captured the critical temperature dependence on the hydrophilic/hydrophobic balance for two series of copolymers, with the model parameters obtained from one PEO homopolymer and one polyester copolymer and used without further adjustment to the rest of the esters and to all the amide series. This agreement leads to a few important, albeit rather qualitative, conclusions on the phase behavior of these uncharged copolymers: (a) the phase behavior of such systems is heavily controlled by the hydrogen-bonding, rather than the van der Waals or other interactions; (b) the LCST, and to some extent the binodal, can be predicted on the basis of the PEO aqueous phase behavior after simple weighted-addition of hydrophobic contributions, and thus can be related back to the hydrophilic/hydrophobic balance in the copolymer composition; (c) suggests that the effects of the water network distortion by the introduction of hydrophobic groups, beyond the trivial re-enumeration of hydrogen bonding probabilities, are probably small. On this last remark, one should consider that the parameter values, as obtained from the fitting of binodals of aqueous solutions, do arguably include implicitly the energetics of the water (solvent). Thus a safer wording would be that the effects of the water network distortion for the (ethylene oxide/ethylene) copolymers are similar to the effects in poly(ethylene oxide) aqueous solutions.

Higher fidelity mean-field approaches, including modifications based on a quasi-chemical lattice-fluid framework [30,31] and on hydrogen bonding cooperativity [2,32], as well as molecular based simulations [33], are the focus of current work, so as to obtain more insights into the phase behavior – and ultimately into quantitative design principles – of polymers with tunable temperature response.

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Appendix A. Numerical procedure

Experimental polymer/water phase diagrams were fitted with the present model using a nonlinear least-squares procedure. Naturally, the fitting process involved the finding of numerical solution of the system of equations (26) repeatedly for different values of parameter $\xi_{12}$. To facilitate the numerical procedure and to avoid the finding of the trivial solution, equations (26) were rewritten in the following equivalent form

$$\left(1 - \frac{\mu_1^B}{\mu_1^A}\right) \frac{1}{x_2^A - x_2^B} = 0,$$

$$\left(1 - \frac{\mu_2^B}{\mu_2^A}\right) \frac{1}{x_2^A - x_2^B} = 0 \quad (A.1)$$

The system of binodal equations (A.1) and the equations of state (20) was solved numerically using the trust-region dogleg method [34]. It appears to be more logical to build a binodal curve by solving for $x_2^B$ and $T$ for a series of $x_2^A$ values, instead of solving for $x_2^A$ and $x_2^B$ for every given temperature value. Due to the system complexity, the choice of initial approximation is crucial to the successful finding of the roots and has to be automated in order to be used in the fitting procedure. Fortunately, the system of the binodal equations has a much better tolerance to the initial approximation, which allows for solving for a point on the spinodal curve, which, in turn, can be used as a good initial approximation for the solution of the critical point equations (taking into account the fact that the critical point is the extremum of the spinodal curve). Taking two points closely on opposite sides of the critical point provides a sufficiently good initial approximation for the system of the binodal equations. Naturally, after the first pair of binodal points is found, it is used as the initial approximation for the solution at the next value of $x_2^A$; granted sufficiently small step size, the described algorithm can be fully automated and can successfully construct a binodal which can fit the experimental phase diagram data.

Appendix B. Supplementary data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.fluid.2007.06.018.

References