

Cooperativity during Melting and Molecular Exchange in Micelles with Crystalline Cores

Nico König,^{1,2,§} Lutz Willner,^{1,*} Vitaliy Pipich,^{3,¶} Thomas Zinn,^{2,†} and Reidar Lund^{2,‡}

¹Jülich Centre for Neutron Science (JCNS) and Institute for Complex Systems (ICS),
Forschungszentrum Jülich GmbH, 52425 Jülich, Germany

²Department of Chemistry, University of Oslo, P.O. Box 1033, Blindern, 0315 Oslo, Norway

³Jülich Centre for Neutron Science (JCNS) at Heinz Maier-Leibnitz Zentrum (MLZ),
Forschungszentrum Jülich GmbH, 85747 Garching, Germany

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Molecular exchange processes are important equilibration and transport mechanisms in both synthetic and biological self-assembled systems such as micelles, vesicles, and membranes. Still, these processes are not entirely understood, in particular the effect of crystallinity and the interplay between cooperative melting processes and chain exchange. Here we focus on a set of simple polymer micelles formed by binary mixtures of poly(ethylene oxide)-mono-*n*-alkyl-ethers (C_n -PEO5) which allows the melting point to be tuned over a wide range. We show that the melting transition is cooperative in the confined 4–5 nm micellar core, whereas the exchange process is widely decoupled and unimeric in nature. As confirmed by differential scanning calorimetry, the total activation energy for ejecting a molecule out of the micellar core below the melting point is the sum of the enthalpy of fusion and the corresponding activation energy in the melt state. This suggests that a “local, single-chain melting process” preludes the molecular diffusion out of the micelle during chain exchange.

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Partially crystalline, self-assembling systems with multiple components are omnipresent in nature with living cells as a prominent example. Their structural integrity is maintained by a membrane consisting of various lipids that can be in a liquid, gel, or crystalline state. For other self-assembling systems like block copolymers with crystallizable blocks, crystallization may even be the driving force behind self-assembly and can yield a whole range of remarkable nanostructural morphologies [1]. In the resulting micellar systems with crystalline cores of typically a few nanometers in diameter, interesting questions relate to the role of confinement. For example, it was recently discovered that the melting points of *n*-alkanes under soft nanoscopic confinement in micellar cores exhibit a simple Gibbs-Thomson behavior [2], typical for hard-confined liquids. In this context, it is also interesting how the dynamics of self-assembling systems, like molecular exchange kinetics, are affected by confinement and crystallinity, as these processes control the equilibration and sometimes even the resulting morphology.

So far, the existing theories for micelle kinetics have been devoted to amorphous micelles. For instance, the general theory developed by Halperin and Alexander [3] considers starlike micelles with molten cores where the chain expulsion is the rate-determining step. The expulsion is pictured as a process in which the chain diffuses over a free energy barrier with an activation energy E_a , which stems from the additional solvophobic surface created upon chain expulsion. This is a first-order kinetic

process where the fraction of exchanged chains is described by an exponential relaxation function $R(t) = \exp(-t/\tau)$. Here the characteristic timescale is given by $\tau = \tau_0 \exp[E_a/(k_B T)]$, where k_B is Boltzmann’s constant, T the absolute temperature, and τ_0 a prefactor that is associated with the attempt time, i.e., the time between two consecutive attempts of a molecule to leave the micellar core. In a previous study of C_n -PEO5 micelles [4], we could show that in molten cores the activation energy E_a scales linearly with the length of the solvophobic block n because the relatively short alkyl block does not fully collapse into a spherical shape—in contrast to the $E_a \propto n^{2/3}$ scaling of fully collapsed solvophobic blocks predicted by Halperin and Alexander [3]. Since $R(t)$ is an exponential function of τ and the exchange time τ in turn depends exponentially on $E_a \propto n$, the relaxation function shows an extremely strong sensitivity to the solvophobic block length. For block copolymers with even a narrow distribution of n (dispersity M_w/M_n close to unity), there is thus a relatively broad distribution of exchange rates which results in a close-to-logarithmic decay instead of a single exponential [5–7]. This renders an exact determination of τ and E_a impossible for block copolymers with finite dispersity. The effect is even more drastic in binary block copolymer mixtures of different molecular weight, so only qualitative conclusions can be drawn [8,9]. In this work, our goal is twofold: (i) understand the role of crystallinity and possible effects of cooperativity quantitatively and (ii) understand the mutual

influence on the dynamics in systems with multiple components.

Until now, it has been found that in micellar systems crystallinity yields an additional energy barrier upon chain expulsion [4,10], but the interplay between the cooperative melting processes and chain exchange is still unknown. An interesting approach to shed light on these processes is coassembling different amphiphiles which allows the melting point to be tuned. Here we use binary mixtures of amphiphilic, micelle-forming copolymers as a model system, namely, two poly(ethylene oxide)-mono- n -alkyl-ethers (C_n -PEO5 with different alkyl lengths $n = 22, 28$ and a PEO molecular weight of 5 kg/mol) in aqueous solution. The polymers are prepared by well-established living anionic polymerization techniques leading to materials with desired PEO molar mass and narrow molar mass distributions of $M_w/M_n < 1.04$ [11]. The n -alkanes, on the other hand, are monodisperse, $M_w/M_n = 1$, which assures a single expulsion time, and do not exhibit the strong broadening inherent in polydisperse systems. Calorimetry and densimetry data show that the hydrophobic blocks mix within the micellar cores, and the melting point can be tuned continuously from 29 to 57°C. In contrast, neutron scattering measurements reveal that the molecular exchange kinetics is governed by two unique expulsion rates corresponding to each component, both in the molten and crystalline state. That means, while the individual diffusion processes are decoupled, the two components are still subject to the same cooperative melting process which, below the melting point, imposes an additional activation energy term equal to the enthalpy of fusion ΔH_{fus} .

The micelles were structurally characterized by small-angle neutron and x-ray scattering (SANS, SAXS). Figure 1(a) shows representative SANS data of neat C_{28} - and C_{22} -PEO5 solutions and of a solution of a premixed 1:1 blend. A fit analysis of the scattering data using a

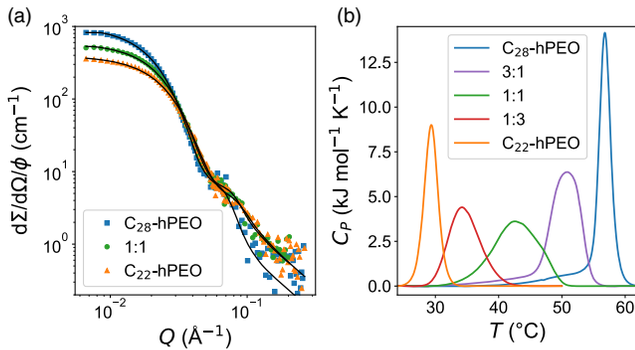


FIG. 1. (a) SANS data of pure C_{28} - and C_{22} - h PEO5 micellar solutions and of a premixed 1:1 blend (volume fraction $\phi = 0.25$ vol%). Solid lines represent fits according to a quantitative core-shell model reported in Ref. [11] and outlined in the Supplemental Material [12]. The key fit parameters are shown in Table I. (b) nanoDSC traces of the same samples and two additional mixtures measured at a scan speed of 2 K/min.

previously reported model [11] shows that the micelles exhibit a typical core-shell structure consisting of a fully segregated, solvent-free core of constant density and an extended starlike PEO corona. The structural key parameters are given in Table I. Complementary SAXS data are shown in Fig. S1 in the Supplementary Material [12], together with the resulting fit parameters and a brief outline of the fit model. Evidently, the size of the 1:1 mixed micelles is in between those of the neat ones, indicating that both species mix on a molecular level. This is further supported by differential scanning calorimetry (nanoDSC) shown in Fig. 1(b). For mixed micelles, only a single exothermic peak is observed which is attributed to a cooperative melting process in the micellar core. The individual melting transitions widely disappear and therefore strongly suggest mixing of C_{28} and C_{22} blocks, even though the broadening of the transition indicates some local heterogeneities and a distribution of mobility caused by the small number of molecules present in the core and their tethering to the core-corona interface. The melting points and fusion enthalpies are also given in Table I. Noteworthy, the observed melting transitions are in quantitative agreement with density measurements performed on the same samples (see Fig. S2 in the Supplemental Material [12]). It is interesting that we observe miscibility of the C_{28} and C_{22} blocks ($\Delta n = 6$) in the core, even though bulk alkanes with $\Delta n > 4$ are not miscible according to Kravchenko's rules [13,14]. This is consistent with studies reviewed by Su *et al.* [15] which show that micrometer confinement enhances the miscibility of n -alkanes. The enhanced miscibility is attributed to a suppressed lamellar ordering and longitudinal diffusion under confinement, which are processes involved in the phase separation of length-mismatching n -alkanes in bulk.

The kinetic experiments were performed by time-resolved SANS using a kinetic zero-average-contrast experiment [16]. Briefly, the method works by mixing two populations of micelles, one with a deuterated and the other with a protonated PEO block, both in a zero average contrast H_2O/D_2O mixture ($\phi_{D_2O} = 66.7$ vol%), which matches exactly the average scattering length density of

TABLE I. Structural and calorimetric data of the investigated micelles obtained from fitting SANS curves and analyzing DSC traces, respectively: N_{agg} , aggregation number; R_m , micellar radius; R_c , core radius; T_m , melting point; ΔH_{fus} , enthalpy of fusion.

	N_{agg}	R_m	R_c	T_m	ΔH_{fus}
		(Å)	(Å)	(°C)	(kJ/mol)
C_{28} - h PEO5	130 ± 5	135 ± 5	28 ± 1	57 ± 1	40 ± 3
3:1	108 ± 5	127 ± 4	26 ± 1	51 ± 1	39 ± 3
1:1	84 ± 4	121 ± 4	24 ± 1	43 ± 1	33 ± 2
1:3	72 ± 3	116 ± 3	22 ± 1	34 ± 1	27 ± 1
C_{22} - h PEO5	57 ± 2	107 ± 3	20 ± 1	29 ± 1	25 ± 1

the two PEO blocks. We note that the deuterated polymers have almost the same PEO length as their proteated counterparts and that the deuteration has no significant effect on the micellar structure (as shown in the Supplemental Material of Ref. [4]). Upon mixing equal volumes of the oppositely labeled micellar solutions, initially at $t = 0$, the two micelle populations scatter with maximum contrast. But as soon as molecules are exchanging, the contrast decreases to a minimum at $t = \infty$, where the h - and d PEO chains are fully randomized and matched out by the solvent on larger length scales. Rapid mixing in the millisecond range was accomplished with a stopped-flow apparatus equipped with a separately temperature-controlled neutron probe head which ensures a high thermal stability in the range between 5 and 55 °C. From the integrated detector intensity $I(t) = \int I(Q, t) dQ$ the relaxation function $R(t) = \{[I(t) - I(\infty)]/[I(0) - I(\infty)]\}^{1/2}$ can be calculated [7], with the initial intensity immediately after mixing $I(0)$ and the intensity from randomly mixed micelles $I(\infty)$. This procedure allows for a much better time resolution (down to 25 ms) compared to tracking the full scattering pattern $I(Q, t)$.

Example curves of C_{22} -PEO5, C_{28} -PEO5, and the 1:1 mixture at various temperatures are shown in Fig. 2. The neat micelles exhibit an exponential relaxation function, as predicted by the theory of Halperin and Alexander [3]. Surprisingly, the mixture shows two distinct exchange processes which can be modeled simply by a sum of two exponentials without any broadening. We intuitively attribute the faster process to C_{22} - and the slower process to C_{28} -PEO5 molecules exchanging. That suggests that even though the different alkyl blocks mix inside the micellar core and crystallize cooperatively, the exchange processes are still decoupled, in agreement with previous reports on amorphous micelles [8,9]. This is shown more quantitatively in Fig. 3 which displays an Arrhenius plot for the

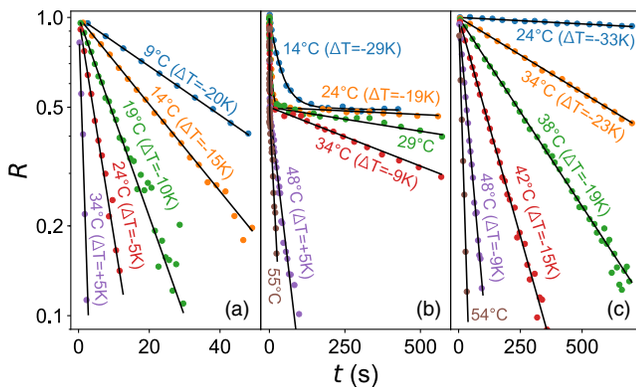


FIG. 2. Experimental relaxation functions and exponential fits at various temperatures. (a) C_{22} -PEO5, (b) $C_{28/22}$ -PEO5 1:1, (c) C_{28} -PEO5. The temperature difference $\Delta T = T_{\text{exp}} - T_m$ to the melting point is given in brackets. Some curves have been left out for reasons of clarity.

exchange times in the 1:1 mixture compared to those in the neat C_n -PEO5 micelles. For an accurate comparison, a contour plot of the corresponding DSC traces, which shows the melting transition, has been included.

Evidently, two distinct regimes with different slopes emerge which reflect the molten and crystalline state as determined by nanoDSC. Moreover, the difference in slope in the Arrhenius plot, and hence, activation energy, exactly matches the enthalpy of fusion as determined from the nanoDSC experiments. For example, for neat C_{22} -PEO5 we obtain $E_a = 105$ kJ/mol below melting and $E_a = 80$ kJ/mol above melting, which corresponds very nicely with $\Delta H_{\text{fus}} = 25$ kJ/mol. Unfortunately, we could not compose a similar plot for neat C_{28} -PEO5 because temperature stability in the stopped-flow setup was not achievable above the melting point $T_m = 57$ °C. Thus, the respective plot only exhibits a uniform slope below T_m corresponding to the activation energy of C_{28} -PEO5 in the crystalline regime (170 kJ/mol). However, via reducing the melting point by mixing C_{28} - and C_{22} -PEO5, we were able to obtain similar results for C_{28} -PEO. Analogous to neat C_{22} -PEO5, the reduced melting point in the 1:1 mixture allows for an Arrhenius plot which is separated into the molten and crystalline regimes. Likewise, the activation energies of both regimes differ by the enthalpy of fusion. Additionally,

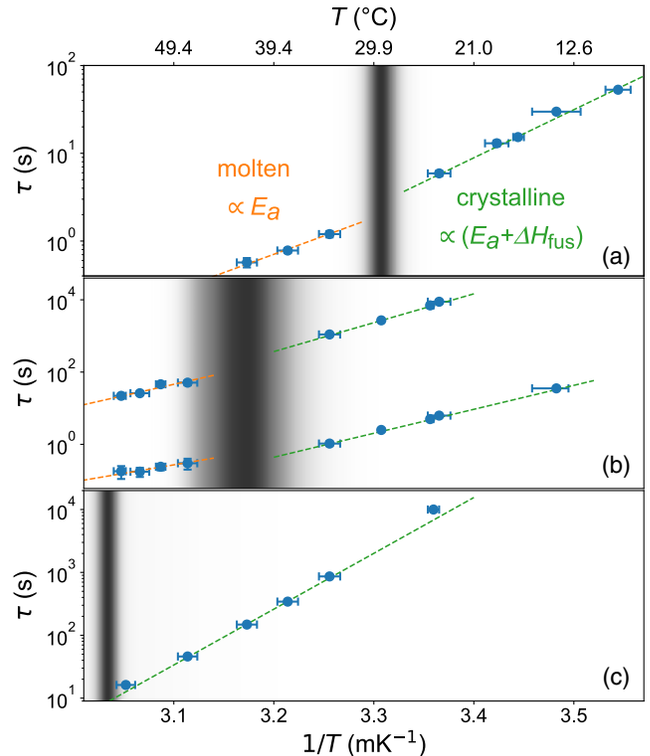


FIG. 3. Arrhenius plot of the temperature-dependent characteristic exchange times of (a) C_{22} -PEO5, (b) the $C_{28/22}$ -PEO5 1:1 mixture, and (c) C_{28} -PEO5. The gray shade is a contour plot of the respective nanoDSC melting curve and thus separates the molten and crystalline regimes.

we performed a similar analysis of 1:3 and 3:1 mixtures (see Figs. S3 and S4 in the Supplemental Material [12]).

Based on these findings, activation energies obtained below T_m have been corrected for the enthalpy of fusion, as measured by nanoDSC, via $E_a \rightarrow E_a - \Delta H_{\text{fus}}$. The thus obtained individual net activation energies E_a of C₂₂- and C₂₈-PEO5 in the different compositions are shown in Fig. 4(a). The plot clearly reveals that within experimental uncertainty, E_a remains virtually unchanged for both alkane lengths in mixed micelles. Apparently, the individual expulsion of the alkane block is almost independent of the core composition, thus, confirming the “independent chain hypothesis” [9]. This is straightforwardly explained by the fact that—in the molten regime—the core blocks only “feel” an alkane environment, and the specific length of the neighboring chains does not significantly influence the energy barrier imposed by the additional hydrophobic surface upon chain expulsion. Nevertheless, when inspecting the C₂₂-PEO5 activation energies more closely, there is a trend of increasing E_a with increasing C₂₈-PEO5 content. We attribute this to the influence of core-corona-interface area per chain as described by Zhao *et al.* [17]. As the average alkyl chain length in the mixture increases, so does the aggregation number and core radius (compare Table I), but the interface area per chain decreases $\propto 1/R_c$. The “orifice” through which the alkyl block leaves the core thus becomes smaller and the corona denser, both effects hindering chain expulsion and increasing the apparent activation energy.

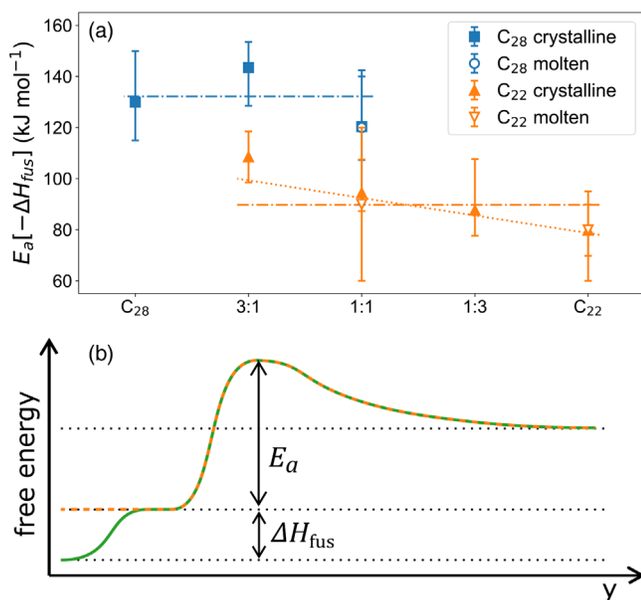


FIG. 4. (a) Activation energies of the chain expulsion step of C₂₈- and C₂₂-PEO5 in various mixtures. Values obtained from the crystalline regime were corrected for the enthalpy of fusion determined from nanoDSC. The (dashed-)dotted lines are guides to the eye. (b) Tentative free energy landscape of the expulsion step where y is the reaction coordinate.

We should note that the experimental uncertainty of the C₂₈-PEO5 activation energies is generally larger than those of C₂₂-PEO5, and hence prevent similar deductions for C₂₈-PEO5. This is basically due to the necessary long collection times (several hours) to get the full relaxation profile and to determine accurately the final intensity, $I(\infty)$.

Inspecting the corresponding attempt times τ_0 yields a similar picture (see Fig. S5 in the Supplemental Material [12]): When correcting for the entropy of fusion $\Delta S_{\text{fus}} \approx \Delta H_{\text{fus}}/T_m$ [$\tau_0 \rightarrow \tau_0 \exp(\Delta S_{\text{fus}}/R)$, where R is the universal gas constant] [4], the attempt times in the molten and crystalline regimes agree [18]. We thus propose that the cocrystallization gives an additional, cooperative energy barrier to the otherwise decoupled expulsion processes in mixed micelles as is depicted in Fig. 4(b).

To summarize, employing this well-defined model system, we show that cooperative solid-liquid phase transitions in nanocrystalline self-assembled micellar systems not necessarily imply cooperativity in intermicellar exchange processes. That implicates that the segmental diffusion (i.e., local conformational transitions and segmental motion) is essentially decoupled from the center-of-mass diffusion (i.e., molecular exchange). However, it should be noted that this does not translate to an effective friction coefficient but rather a modification of the activation energy of the expulsion step. Even though the expulsion process includes a partial premediating melting process as the chain escapes the crystalline core [19], we still find two unique release processes corresponding to each component in the binary mixtures. Furthermore, the data show that the activation energy of the expulsion step governing the diffusion process out of the core undergoes an abrupt reduction at the melting point that exactly amounts to the enthalpy of fusion. Very interestingly, the transition temperature can be accurately tuned by coassembling polymers with different n -alkyl length. Above the melting transition, this occurs without affecting the activation energies of the individual compounds, which only depend on the hydrophobic surface created upon chain expulsion. However, below the melting transition, an additional term ΔH_{fus} is added to the total activation energy, a quantity which depends on the mixing ratio and the nature of the n -alkane. Hence, cocrystallization, although less effective than varying the core block length [5,7], corona block length [20,21], and molecular architecture [22–24], represents a handy tool to fine-tune the temperature response of the timescale on which the molecular exchange takes place. This is important insight for the accurate design of multicomponent self-assembling systems, although in the case of mixing chemically dissimilar components, additional modification of the dynamics may be expected, e.g., a change in the local segmental motion. We also shed light on the thermodynamics and kinetics of low-molecular-weight molecules in nanoconfined systems. It has been reported that confinement may induce demixing of small molecules like, for

instance, water and methanol [25] or colloidal mixtures [26]. In our case, however, the data demonstrate that the spherical nanoconfinement greatly enhances the miscibility of crystalline long n -alkanes which are incompatible on a macroscopic scale—in agreement with previous studies on microconfinement [15]. Moreover, we show that the dynamics in crystalline self-assembled materials can be rather straightforwardly predicted from their amorphous state based on quantities easily measurable using standard techniques such as DSC. Similar ideas and experimental approaches can be applied within biophysics to understand the properties of membranes that consist of a range of lipids with similar chemical compositions but varying melting points.

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*l.willner@fz-juelich.de

ORCID ID:0000-0001-6482-7937

†Present address: ESRF—The European Synchrotron, 38043 Grenoble Cedex 9, France.

‡reidar.lund@kjemi.uio.no

ORCID ID:0000-0001-8017-6396

§ORCID ID:0000-0003-3319-5708

¶ORCID ID:0000-0002-3930-3602

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