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MesoDyn prediction of a pharmaceutical microemulsion self-assembly consistent with experimental measurements

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In this work, for the first time a coarse-grained model in a mesoscopic dynamic (MesoDyn) simulation for a pharmaceutical nonionic surfactant Brij97 was presented. By using this coarse-grained model, the morphology transition from oil in water (O/W) to a bicontinuous microemulsion formed in the Brij97/isopropanol/isoamyl acetate/H₂O quaternary system was predicted. Furthermore, the effect of emulsifier/isoamyl acetate mass ratio, Brij97/alcohol mass ratio and alcohol structure on the droplets size and kinetic formation process of the O/W microemulsion were studied. These MesoDyn predictions were consistent with experimental measurements. This makes it possible to further predict the formation mechanism of complex drug-loaded aggregates of Brij97 by using this coarse-grained model.

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1 Introduction

Recently, pharmaceutical surfactant self-assemblies have aroused extensive interests due to their excellent abilities of sustained and controlled release for bioactive drugs.¹ Polyoxyethylene oleyl ether Brij97, as a typical non-ionic surfactant, has its advantages in drug delivery on account of the special ability to enhance the permeation of a drug.² In our previous studies, Brij97 could form various aggregates including micellar,³ microemulsion,⁴ lamellar,⁵ hexagonal^{6,7} and cubic^{8,9} liquid crystals. Although the formation conditions of various Brij97-based aggregates have been shown clearly, the detailed information of the self-assembled process of Brij97 in the oil-water interface at the nanometer scale¹⁰ is difficult to obtain from experiments.

Aiming to address the limitation of the experiments, computer simulations have been proven to be valuable tools for providing additional information at molecular¹¹ or mesoscopic scale.¹² In particular, mesoscopic dynamics (MesoDyn) simulation have become a powerful analytical approach for exploring the phase behaviour and dynamic evolution process of polymer or surfactant in solvents at real conditions (pressure and temperature). Li, Y. *et al.*¹³ used MesoDyn method to simulate the aggregation behaviour of pluronic copolymer L64 solutions

in the presence of SDS. The kinetic process of micelles formation underwent transitions from three to two stages with the increase of L64 concentration, which was hard to be observed from the experiments. The necessity of the shear and the weak charge to the formation of hexagonal templates in the P123 copolymer were confirmed by Yuan, S. *et al.* through MesoDyn simulations.¹²

Microemulsions, usually consisting of surfactant, cosurfactant, oil and water, are defined as optically isotropic transparent and thermodynamically stable systems. ¹⁴ Microemulsions have been extensive applied in various fields, such as separation, ¹⁵ chemical reactions, ¹⁶ nanomaterial preparation, ¹⁷ and drug delivery. ¹⁸ Microemulsions application are largely dependent on their different structures, namely, oil-in-water (O/ W), bicontinuous (B.C.), and water-in-oil (W/O) structures. However, as far as we know, the effect of components on the aggregation process of microemulsion from the perspective of phase behaviour and nanostructure ¹⁹ was rarely reported.

In present work, the coarse-grained model of Brij97 in MesoDyn simulation was presented for the first time. The morphologies of Brij97/isopropanol/isoamyl acetate/ H_2O were predicted by using the Brij97 model. The effect of water content, emulsifier/isoamyl acetate mass ratio, Brij97/isopropanol mass ratio and alcohol structure on the morphology and size of microemulsion were further studied by a combination of MesoDyn prediction and experimental measurements.

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2 Materials and methods

2.1 Materials

Polyoxyethylene-10-oleyl ether (Brij97, purity ≥ 99%) was purchased from Sigma Chemical Company. Isopropanol, ethanol, 1,2-propanediol and isoamyl acetate were provided by

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Sinopharm Chemical Reagent Co., LTD (Shanghai, China). Water was deionized and then doubly distilled. All chemicals were used without further purification.

2.2 Simulation method and parameters

Before employing the MesoDyn simulation, the components of our researched system were represented by coarse-grained models. In the present MesoDyn simulation, the coarse-grained model of Brij97 as E₃C₂ was designed and proposed for the first time, where the E beads represented the hydrophilic EO chain and the C beads represented hydrophobic alkyl chain. The ratio of beads E to C was dependent on a simple relationship for atomistic and Gaussian chains^{12,20} with a few modifications. It is believed that this modifications at the mesoscale level does not result in a large deviation.²¹ The molecular structures and coarse-grained models of all components are shown in Fig. 1. As shown, the bead W represented H₂O, the bead A represented alcohol, and the bead O represented isoamyl acetate in the MesoDyn simulation.

The interaction among various beads were represented by interaction parameters. The solvent–polymer interaction energy λ_{ij} are related to the Flory–Huggins interaction parameter χ_{ij} through the following equation:²²

$$\lambda_{ij} = \chi_{ij}RT \tag{1}$$

The Flory–Huggins interaction parameter χ_{ij} can be obtained by the expression:²¹

$$\chi_{ij} = \frac{V_{\rm S}(\delta_i - \delta_j)^2}{RT} + \chi_{\rm S} \tag{2}$$

where δ_i and δ_j are solubility parameters of components i and j, respectively. $V_{\rm S}$ is the molar volume of solvent. $\chi_{\rm S}$ is the entropy contribution to the mixing energy, which is neglected in this study because it is a relatively small correction in comparison to other terms.²¹

The Flory–Huggins interaction parameter between the hydrophobic tail of Brij97 and isoamyl acetate χ_{C-O} was estimated to be 0.1 calculated from the eqn (2), consistent with similar dissolve mutually theory. However, the values of χ_{E-W} ,

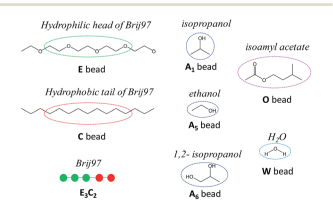


Fig. 1 The coarse-grained models of Brij97, alcohols, isoamyl acetate and H_2O in the MesoDyn simulation. The molecular structure of Brij97 is divided into two types of beads: C and E.

 $\chi_{\text{A-W}}$ and $\chi_{\text{E-A}}$ calculated from the eqn (2) were incorrect. The possible explanation to the trouble is that the hydrogen bonding between beads and solvent was neglected.²³ Hence, the effective Flory–Huggins parameters were defined in Baulin and Halperin's work,²⁴ $\bar{\chi}=\chi_{\text{eff}}+f(\Phi)$. In general, the miscibility between polymer and solvent are dependent on the interaction parameter. Consequently, $\chi_{\text{E-W}}$ used in present study was defined as 0.3 on account of the hydrophilic properties of surfactant EO headgroup, and $\chi_{\text{C-W}}$ was estimated to be 5.0 due to the hydrophobic properties of surfactant alkyl chain tail. The effective Flory–Huggins interaction parameters between various beads used in this work were listed in Table 1.

In MesoDyn simulation, the dimensionless parameters were chosen as follows: the size of cubic grid $32 \times 32 \times 32$ nm, the bond length 1.1543, the bead diffusion coefficient 10^{-7} cm² s⁻¹, the noise parameter 100.0, the compressibility parameter 10 kT, the simulated temperature 298 K, the time step 0.5 ns. For each system, the total number steps of 20 000 were carried out to reach a kinetic equilibrium. All simulations were carried out with MesoDyn module in a commercial software Materials Studio 4.4 from Accelrys, Inc,²⁵ successful used to studied several surfactant²² or polymer²⁶⁻²⁸ solution systems.

2.3 Construction of phase diagram

The pseudo-ternary phase diagram was constructed by the previous method described by Wang $et~al.^4$ Firstly, the surfactant (Brij97) and co-surfactant (isopropanol) were premixed at the weight ratio of $1:1~(K_{\rm m})$ as the emulsifier (EM). Secondly, the mixtures of the EM and isoamyl acetate with the weight ratios (K) varying from 0:10 to 10:0 were stirred to be homogenized at about 50 °C. Finally, individual drops of water phase was added to each mixture under proper magnetic stirring at 25 °C. The microemulsion region was identified by a clear and transparent appearance of the solution on visual observation. Samples near the phase boundary line were equilibrated at 25 °C in a thermostatic water bath for 24 hours.

The microstructural transition from W/O, B.C. to O/W in the microemulsion region was characterized by electric conductivity measurement using a model DDSJ-308A digital conductivity meter equipped with a DJS-1C platinum bright electrode (Shanghai INESA Scientific Instrument Co., Ltd., Shanghai, China).

 $\begin{tabular}{ll} \begin{tabular}{ll} \be$

Bead	E	C	Α	O	W
E	0	1.6	0.8	4.0	0.3
\mathbf{C}	1.6	0	1.0	0.1	5.0
Α	0.8	1.0	0	0.6	1.2
O	4.0	0.1	0.6	0	5.0
W	0.3	5.0	1.2	5.0	0

^a E, C, A, O and W represent hydrophilic group of Brij97, hydrophobic group of Brij97, alcohol, isoamyl acetate, and H₂O, respectively.

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2.4 Diameter distributions of microemulsion formulations

The diameter distributions of different microemulsion formulations were determined using dynamic light scattering (DLS) with a Zetasizer Nano ZS90 particle size analyzer (Malvern Instruments Ltd., United Kingdom). The data of size distribution was collected at the 90° scattering angle and the temperature was kept at 25 \pm 0.1 °C.

3 Results and discussion

3.1 The morphologies of Brij
97/isopropanol/isoamyl acetate/ $\rm H_2O$

As we all know, the hydrophilic EO chain of Brij97 tends to contact with water, and the hydrophobic alkyl chain and oil phase should come to contact each other. Hence, the morphology of Brij97 self-assemblies could be simulated by the isosurface of E and C beads, consistent with mostly previous reports. Here for convenience, the content of each component in system was the weight percentage, consistent with the experiments. While the concentrations used in the MesoDyn simulation were transformed to the corresponding volume fraction. ²²

Fig. 2(a) showed the dependence of simulated morphology of Brij97 assemblies on water content. The components of sample S_1 and S_2 were listed in Table 2. For S_1 with 80 wt% H_2O , the E beads isosurface formed several spherical droplets. In order to

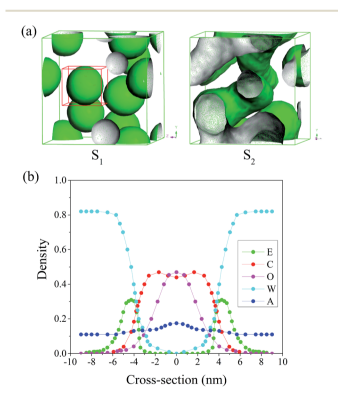


Fig. 2 (a) The beads E isosurface representations for microemulsion formed by Brij97/isopropanol/isoamyl acetate/ H_2O system with different mass ratio of water: S_1 (80%) and S_2 (60%). (b) The radial density distributions of all compositions in the spherical droplet marked by a red cubic grid in S_1 .

inspect the distribution of all compositions in the spherical droplets, a representative droplet marked by red cubic grid was segregated and analysed the radial density distribution. As shown in Fig. 2(b), the density distribution of O, C and W beads presented a single peak, broad peak and valley respectively, indicating the isoamyl acetate molecules and hydrophobic tails of Brij97 composed the core of spherical droplets without free water molecules. For hydrophilic beads E of Brij97, the density distribution presented two low peaks far from the centre of sphere, indicating a hydrophilic shell. This is a typical morphology of O/W microemulsion. While for S2 with 60 wt% H₂O, the E beads isosurface formed several cylindrical channels. It is similar to a classical bicontinuous structure with water tank and oil pipe. The simulation results presented that the microemulsion underwent a transition from B.C. to O/W with water content increasing from 60 to 80 wt%. In order to further confirm the prediction obtained by MesoDyn simulation, the phase diagram of Brij97/isopropanol/isoamyl acetate/ H_2O ($K_m = 1:1$) system at 25 °C was constructed.

3.2 Phase behaviour of Brij97/isopropanol/isoamyl acetate/water system

The pseudo-ternary phase diagram of Brij97/isopropanol/isoamyl acetate/ H_2O ($K_m=1:1$) system at 25 °C was constructed and shown in Fig. 3. The phase boundary was easy to determine by visual observation. As shown, a transparent one-phase microemulsion region is marked " 1Φ " in Fig. 3. The rest of the phase region marked " 2Φ " is a turbid two-phase region which is a microemulsion in equilibrium with an excess water or oil phase. The similar phase behaviour was observed in Brij97/isopropanol/ethyl acetate/ H_2O ($K_m=1:1$) system at 25 °C, whereas the microemulsion area was enlarged by replacing isoamyl acetate with ethyl acetate. This indicated that the microemulsion area was closely related to the chain length of oil molecule, consistent with the previous result. In the microemulsion region marked " 1Φ ".

In order to confirm the prediction of microstructural transition obtained from MesoDyn simulation, further experimental measurement was necessary. It is well known, by measuring the variations of electrical conductivity (κ) of microemulsion as a function of the water content, the microemulsion region was divided into three parts: W/O, B.C. and O/ W (see in Fig. 3). A representative electric conductivity curve was shown in the insert of Fig. 3. It was obviously found that three different types of microstructure: W/O (below 54.9 wt%), bicontinuous (B.C., 54.9-71.7 wt%) and O/W (above 71.7 wt%) occurred in the present system along the water dilution line a (marked in Fig. 3). Comparing with the experimental phase diagram and electrical conductivity results, simulation results can reproduce corresponding microemulsion structure in certain water content, which indicates that the selection of coarse-grained models and interaction parameters is effective and reasonable.13

In order to further explore the aggregation behaviour of Brij97-based microemulsion, several samples were prepared

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Table 2	The components.	size and po	olvdispersity	/ index (PDI) of al	l microemulsions tested ^a

Sample	K	$K_{ m m}$	Alcohol	Water (%)	Size (nm)	PDI	Туре
S_1	9:1	1:1	Isopropanol	80	14.81	0.150	O/W
S_2	9:1	1:1	Isopropanol	60	_	_	B.C.
S_3	8:2	1:1	Isopropanol	80	17.33	0.255	O/W
S_4	9:1	3:1	Isopropanol	80	12.30	0.307	O/W
S_5	9:1	1:1	Ethanol	80	15.52	0.189	O/W
S_6	9:1	1:1	1,2-Propanediol	80	16.02	0.216	O/W

^a K and K_m represent the ratio of emulsifier/isoamyl acetate and Brij97/alcohol, respectively. The values of size and PDI were obtained by DLS measurement.

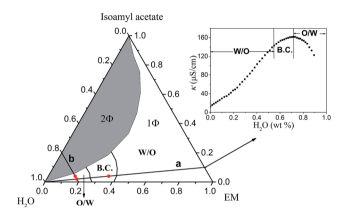


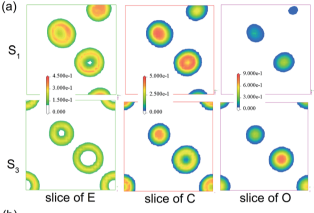
Fig. 3 Pseudo-ternary phase diagram of EM/isoamyl acetate/H₂O system at 25 °C. The insert is a representative curve of electrical conductivity (k) vs. H2O (wt%) along dilution line a

and investigated by using a combination of MesoDyn prediction and experiment methods in the following research. The components and measurement results of all samples tested were listed in Table 2.

Effect of emulsifier/isoamyl acetate mass ratio (K)

To obtain a more clear understanding on the aggregation behaviour of Brij97-based microemulsion, sample S3 with a certain water content of 80 wt% was simulated by changing the mass ratio of emulsifier/isoamyl acetate (K) from 9:1 to 8:2. The MesoDyn simulation could provide the density distribution of different beads in microemulsion droplets, which help us to explore the aggregation behaviour. In the cubic grid, the density slices of beads E, C and O along the x-z plane were created and divided as depicted in Fig. 4. As shown, high density was represented by red colour, and the blue colour represents low density. As shown in the Fig. 4, the beads E was distributed in the outer layer of the droplets in contact with the water phase. While the beads C was distributed in the inner layer of the droplets in contact with the oil phase. These were typical O/W microemulsion structure. Comparing with the density slices of S₁, it was easy to find that the density of beads O of S2 dispersed over a wider area. This induced beads E and C to disperse loosely.

To confirm the MesoDyn prediction, the droplet size distribution of microemulsion S1 and S2 were measured by



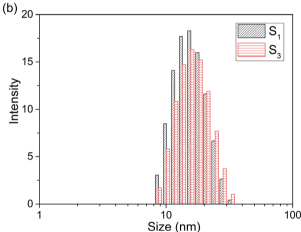


Fig. 4 (a) The density slice of beads E, C and O for microemulsion S₁ and S_3 . (b) The droplet size distribution of S_1 and S_3 .

dynamic light scattering. As shown in Fig. 4 and Table 2, the DLS results showed that the value of both average size and polydispersity index (PDI) increased with the K value decreased. This is consistent with the MesoDyn prediction that beads E, C and O dispersed over a wider area. Moreover, it was obviously found that the droplet size presented in simulated cubic grid (32 \times 32 \times 32 nm) was slightly smaller than the DLS result. The similar result was found that the size of microemulsion droplets in electron microscopy (TEM) images was also smaller than the DLS results.31 This was due to the size obtained by DLS was hydrodynamic diameter rather than real diameter.

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Fig. 5(a) showed the variation of the order parameters of beads W and O as a function of time steps for microemulsion S_1 and S_3 , respectively. As we all known, the formation process of microemulsion is a self-assembling evolution from disorder to order. Hence, the order parameter of beads in system increased along with the aggregation process of microemulsion. It was clear seen from Fig. 5(a), the formation process of the O/W microemulsion can be divided into three stages. In the first stage, all of the order parameter of beads are close to zero, indicating that each component in the system is highly dispersed. In the second stage, the order parameter of each component increased dramatically, which indicated that the ordered aggregates began to form. The third stage is the formation and stabilization of microemulsion. In this stage, the value of order parameter is not constant but waved up and down

Comparing with the order parameters curves of bead E, C and A (not presented), the changes of beads W and O were obvious. As shown in Fig. 5(a), the time of first stage for sample S_3 was shorter than that of sample S_1 . This indicated that the formation rate of microemulsion accelerated with the K value decreasing. The result may be due to the enhancement of hydrophobic interaction. Besides, the order parameter of isoamyl acetate and water increased significantly. The former may be result from the fact that more isoamyl acetate molecules embed into the hydrophobic chains of surfactant to arrange orderly. The latter may be due to the result that larger droplets permitted more water molecules to form hydrogen bond with EO chains of surfactant.

in a small range. It is not surprising because the microemulsion

is a dynamic equilibrium system.

Apart from the size and formation rate of microemulsion, the practical application of microemulsion is also largely dependent on the thermodynamically stability. As we all know, the free energy change of system can reflect the stability. The MesoDyn simulation can provide the free energy change in the aggregation process of microemulsion, while it is hard to obtain from experiment. Fig. 5(b) showed the free energy density plots with time step for microemulsions S_1 and S_3 . As shown, the free

energy density decreased asymptotically to approach a stable value during the simulations when the system reaches dynamic equilibrium. This indicated that the aggregation process of microemulsion is spontaneous. It is worth mentioning that the comparison of the free energy density values between S_1 and S_3 is totally meaningless because it is not routinely calculated for real systems. While in this case, the value of free energy change ($|\Delta G|$) before and after the aggregation could reflect the stability of microemulsion to some extent. As shown in Fig. 5(b), the $|\Delta G|$ value of S_3 was higher than that of S_1 , indicating that the stability of microemulsion increased with the K value decreasing.

3.4 Effect of Brij97/isopropanol mass ratio (K_m)

In general, the formation and stability of microemulsion are closely related to the ratio of surfactant to co-surfactant $(K_{\rm m})$.³⁰ It will be meaningful to predict the effect of $K_{\rm m}$ value on the morphology and aggregation behaviour of microemulsion by MesoDyn simulation. The beads E isosurface representations of sample S_4 was shown in Fig. 6(a). Comparing with the morphology of microemulsion S_1 shown in Fig. 2(a), it is clearly seen that the droplets size of S_4 is smaller than that of S_1 . The

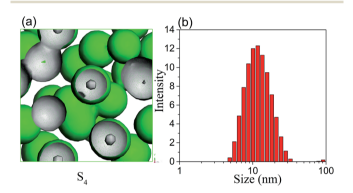


Fig. 6 (a) The beads E isosurface representations for microemulsion S_4 ; (b) droplet size distribution of microemulsion S_4 .

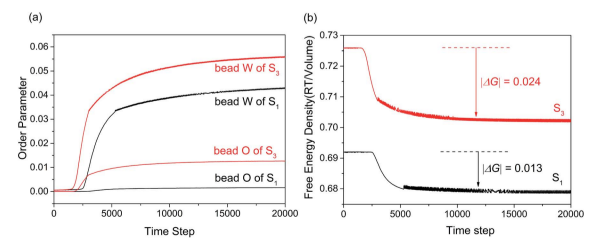


Fig. 5 (a) Order parameter plot of beads W and O with increasing time steps for microemulsion S_1 and S_3 . (b) Free energy density plot with time step for microemulsions S_1 and S_3 .

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DLS resulted showed that the hydrodynamic diameter was also decreased from 14.81 nm to 12.30 nm when the $K_{\rm m}$ value increased from 1:1 to 3:1. It also can be seen from Fig. 6(a), several droplets contacted with each other in the simulated morphology of microemulsion S4, resulting in the nonuniformity of microemulsion. This is consistent with the experimental results that the PDI value of microemulsion S4 is higher than that of S_1 .

Fig. 7(a) showed the variation of the order parameters of beads W, E and C as a function of time steps for microemulsion S₁ and S₄, respectively. From the figure we can see, the formation process of the microemulsion with different $K_{\rm m}$ can also be divided into three stages. While for S4, the first stage of microemulsion formation was remarkably shortened. This is because that the hydrophobic interaction between various components increased with alcohol content decreasing. Besides, the order parameter of surfactant and water increased significantly. Comparing with the hydrophilic headgroup E, the order parameter of hydrophobic tail C increased more significantly. This indicated that isopropanol could weaken the hydrophobic interaction of surfactant by mainly inserting into the hydrophobic tails of surfactant. Fig. 7(b) showed the free energy density plots with time step for microemulsions S1 and S_4 . As shown, the $|\Delta G|$ value of S_4 was higher than that of S_1 , indicating that the stability of microemulsion increased when the $K_{\rm m}$ value increased from 1:1 to 3:1.

3.5 Effect of alcohol structure

Alcohol as a co-surfactant have been widely applied in microemulsion. Generally, alcohol can contribute to the formation of micelle by the means of changing the hydrophile-lipophile balance of surfactant and adjusting the polarity of water and oil phase. Usually, the chain length and polarity of alcohol are responsible for the hydrophile-lipophile ability and surface activity. Here, MesoDyn simulation was used to predict the morphologies and distribution of microemulsion formed by different alcohols.

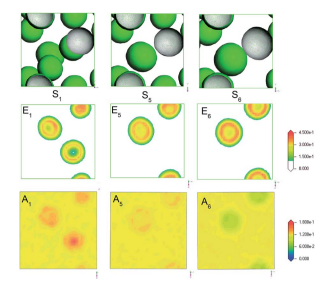


Fig. 8 The E beads isosurface representations (S_1 , S_5 , S_6), density slice of E (E₁, E₅, E₆) and A (A₁, A₅, A₆) for microemulsion with different alcohols: isopropanol, ethanol and 1,2-propanediol.

Fig. 8 S₁, S₅ and S₆ showed the morphologies of microemulsion formed by isopropanol, ethanol and 1,2-propanediol, respectively. All microemulsion were typical O/W microstructure. It was obviously found that the droplets size increased by the order of S₁, S₅ and S₆. The MesoDyn predictions were consistent with the DLS results listed in Table 2. This indicated that the droplets size were dependent on the polarity of cosurfactant alcohol. In order to understand clearly the mechanism of alcohol as a co-surfactant, the density slices of beads E and C of microemulsion with different alcohols were obtained. As we can see from A₁, A₅, and A₆ in Fig. 8, the distribution of alcohol molecules in microemulsion were obviously different. For microemulsion S₁, isopropanol mainly dispersed in oil to improve the polarity of oil. Ethanol mainly dispersed into surfactant molecules to adjust the hydrophile-lipophile balance for microemulsion S₅. For microemulsion S₆, 1,2-propanediol mainly dispersed in water phase to reduce the polarity. The

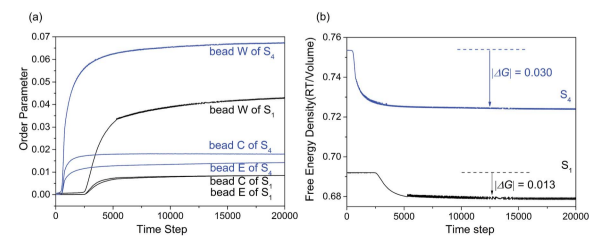


Fig. 7 (a) Order parameter plot of beads W, C and E with increasing time steps for microemulsion S_1 and S_4 . (b) Free energy density plot with time step for microemulsions S₁ and S₄.

different distribution of alcohol molecules may be the reason

why the droplets size of microemulsion changes.

4 Conclusions

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In present work, the phase behaviour and mesoscopic structure of Brij97/isopropanol/isoamyl acetate/water were studied by using a combination of experiment and MesoDyn simulation methods. The structural transition from B.C. to O/W with water content increasing was confirmed by electric conductivity measurement and MesoDyn simulation. At a certain water content of 80 wt%, the size of microemulsion droplets increased with the value of K or $K_{\rm m}$ decreasing. The MesoDyn simulation results showed that the swelling of microemulsion droplets was due to the broad dispersion of isoamyl acetate molecules in the hydrophobic chains of surfactant. Besides, the incensement of alcohol content could induce the formation of large droplets. The effect of cosurfactant on the formation and stability of microemulsion was depended on the hydrophile–lipophile ability of alcohol.

Acknowledgements

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