



A Lipid-Based Depot Formulation with a Novel Non-lamellar Liquid Crystal Forming Lipid

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ABSTRACT

Purpose Non-lamellar liquid crystal (NLLC)-forming lipids have gained attention as a novel component because of their ability to self-assemble upon contact with body fluids. In this study, a novel NLLC-forming lipid, mono-O-(5, 9, 13-trimethyl-4-tetradecenyl) glycerol ester (C17MGE), and a model drug with a middle molecule weight, leuprolide acetate (LA), were used to confirm the usefulness of C17MGE as an excipient for depot formulations with sustained release properties.

Methods A self-constructed depot formulation was prepared by mixing C17MGE and different types of phospholipids. The constructed NLLC structure was evaluated using small angle X-ray analysis and cryo-transmission electron microscopy. In vitro release and blood concentration profiles of LA were investigated.

Results The NLLC structure was confirmed by small angle X-ray analysis. LA release was able to be modified by adding different ratios of various phospholipids to C17MGE. Formulations containing 1, 2-dioleoyl-sn-glycero-3-phosphoglycerol sodium salt with a mixing ratio of 12% or 24% (M_{DOPG12} or M_{DOPG24} , respectively) exhibited sustained release profiles of LA. In addition, the blood concentration of LA was detected over 21 days or more after administration of M_{DOPG12} , and the absolute bioavailability was calculated to be about 100%.

Conclusions A depot formulation using C17MGE was useful to achieve sustained release of LA.

KEY WORDS Lipid-based depot formulation · non-lamellar liquid crystal · self-assembled structure · sustained release

INTRODUCTION

In recent years, the working-age population in developed countries has been declining due to the super-aging of the population (1). Therefore, ensuring working hours for working generations is an important issue. In addition, amidst the novel coronavirus (COVID-19) pandemic situation, many patients hesitate to visit hospital and see their doctor. The development of excipients that enable sustained-release delivery of drugs may improve the quality of life of patients and reduces the risk of recurrence or deterioration of symptoms. In addition, such delivery systems may be helpful to reduce the burden on healthcare professionals. Nowadays, middle-sized molecules (M.W. ca 1–5 kDa) including peptides and nucleic acids are expected as the next target therapeutic drugs because of their high affinity and selectivity to disease sites. Thus, it would be useful to develop new excipients or vesicles that enable sustained delivery for various kinds of therapeutic drugs.

Many reports have been published in the last 5 years showing the usefulness of non-lamellar liquid crystals (NLLCs) as pharmaceutical excipients to obtain sustained drug release (2). NLLCs increases the solubility of poorly soluble drugs and improve the stability of the higher-order structure of proteins (3–5). Ki et al. (6) reported an injectable NLLC system composed of sorbitan monooleate (known as Span 80), phosphatidylcholine, and tocopherol acetate. The NLLC formulation showed sustained release of leuprolide acetate (LA) over 1 month and exhibited a similar bioavailability to the commercial subcutaneous (*s.c.*) depot formulation. In particular, lipids that form an NLLC structure upon contact with body fluids would be useful as injectable excipients. A lipid-based structure is an excellent alternative to polylactic acid-glycolic

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acid copolymer (PLGA) depot formulations. The stability of peptides and proteins entrapped in the depot formulation is a critical point for long-term drug delivery (7). In the PLGA formulation, the acidic products resulting from the PLGA degradation may affect the drug stability by lowering the pH in the formulation (8). Therefore, injectable lipid-based depot formulations may be a promising platform and enable desired drug release and adequate shelf-life longevity.

As mentioned above, NLLC-forming lipids may be attractive pharmaceutical excipient for depot formulations. However, only a limited number of NLLC-forming lipids have been developed, such as glycerol monooleate (GMO), glycerol dioleate, and phytantriol (PHY) (9,10). In the present study, we focused on a newly developed NLLC-forming lipid, mono-O-(5, 9, 13-trimethyl-4-tetradecenyl) glycerol ester (C17MGE), which exists as a liquid state at room temperature and has a low viscosity (0.47 Pa·s at a shear rate of 10 s^{-1}) compared with GMO in a room temperature (11,12).

In our previous report, a newly prepared depot formulation with C17MGE successfully showed an NLLC structure in the body after *s.c.* injection into rats, and the release rate of a hydrophilic middle molecular weight compound, fluorescein isothiocyanate-dextran (M.W.; 4000, FD-4), was controlled by alternation of the interplanar spaces and NLLC structure by adding several types of phospholipids (13). This smart phase transition with C17MGE suggested that it may be a promising drug depot base with sustained release properties of entrapped drugs.

In the present study, LA, a linear nonapeptide with a positive charge at neutral pH (pK_a approximately 6.0: the imidazolyl nitrogen of histidine) (14), was selected as a model middle molecular weight drug. LA is unstable in the skin tissue (15). Therefore, a sustained-release formulation with a high bioavailability of LA is necessary to show the usefulness of C17MGE-based depot formulations. A self-assembled structure formed by C17MGE was evaluated by adding several types of phospholipids, and LA release from the constructed structure and blood-concentration profile were investigated to confirm the usefulness of C17MGE as the excipient for a long-acting depot formulation.

MATERIALS AND METHODS

Materials

LA was purchased from Shin Nippon Yakugyo Co Ltd. (Tokyo, Japan). C17MGE was provided by Farnex Inc. (Yokohama, Japan). Table I shows the chemical structures of LA and C17MGE. Among phospholipids, 1, 2-distearoyl-sn-glycero-3-phosphocholine (DSPC) was selected as a saturated example and 1, 2-dioleoyl-sn-glycero-3-phosphoglycerol, sodium salt (DOPG), 1, 2-dioleoyl-sn-glycero-3-

phosphoethanolamine (DOPE), and phosphocholine (DOPC) were selected as unsaturated phospholipids. These phospholipids were purchased from NOF Corporation (Tokyo, Japan). Table II shows the chemical structure of these phospholipids. This table also shows the abbreviations used for the phospholipids. Polyoxyethylene sorbitan monooleate (Tween 80) was purchased from Tokyo Chemical Industries, Ltd. (Tokyo, Japan). All other reagents were used without further purification.

Preparation of Formulations

Table III shows the composition of the prepared depot formulations. C17 MGE, ethanol, and Tween 80 were weighed and added to a 9 mL vial (laboratory screw tube bottle No. 3, AS ONE K.K., Osaka, Japan) containing a PTFE stirrer (Double Cross-Type D-10 F-4037-01, CFC Corporation, Tokyo, Japan). The weighed compounds were heated at 60°C on a hot plate (Hotplate Stirrer RSH-1DN, AS ONE K.K.) while stirring at 500 rpm for 5 min. Then, weighed saturated or unsaturated phospholipid was added and further mixed at 60°C while stirred at 700 rpm for 60 min. The prepared formulations shown in Table II were abbreviated to $M_{\alpha\beta}$; where α is the type of phospholipid (DSPC/DOPC/DOPE/DOPG) and β is the phospholipid content (i.e., $M_{\text{DSPC}12}$ is a formulation containing 76% C17MGE and 12% DSPC in the formulation). In addition, C17 MGE alone (abbreviated as M_{only}) and C17 MGE with ethanol and Tween 80 (abbreviated as M) were prepared for comparison. Finally, LA was mixed with the prepared formulations using a Pellet Pestle (Thermo Fisher Scientific K.K., Tokyo, Japan) for 2 min. LA concentration in the formulation was 37.5 mg/mL.

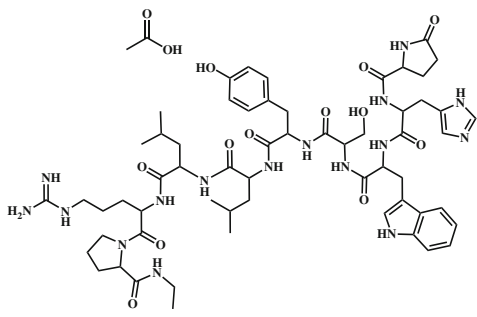
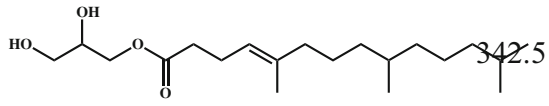
Small Angle X-Ray Analysis

The prepared formulations were analyzed using a small angle X-ray diffraction instrument (Nano-Viewer, Rigaku Co., Ltd., Akishima, Tokyo, Japan) (CuK α radiation, $\lambda = 1.5418 \text{ \AA}$) operated at 30 kV and 40 mA. The camera focal length was set to 700 mm. Formulations immersed into 3 mL of phosphate-buffered saline (PBS) for 6 h were used as samples, and measurements were performed by Kanazawa University (Kanazawa, Ishikawa, Japan). The measurement conditions were as described previously (16). Crystalline interplanar spacing, d , was determined in accordance with the Bragg equation.

In Vitro Release Test from the Formulation

The prepared formulation containing LA (100 μL) was added to dialysis tubing (Pur-A-Lyzer™ Mini 12,000 dialysis kit 25, molecular cut-off 12,000, Sigma Aldrich, St. Louis, MO, U.S.A.), then the dialysis tube was placed into a 25 mL

Table I. Structures of LA and MGE

Drug	Structure	M.W.
Leuprolide acetate		1269.473
Mono-O-(5,9,13-trimethyl-4-tetradecenyl) glycerol ester (C17MGE)		342.5

centrifuge tube (centrifuge tube Mini MINI-2362-025, AGC Technoglass Co., Ltd., Shizuoka, Japan) containing 20 mL of PBS with 0.02% Tween 80. LA release experiments were conducted using the dialysis method.

During the release experiment, the centrifuge tube, covered with aluminum foil to prevent LA degradation, was set in a water bath (AS ONE Corporation, Osaka, Japan) at $37 \pm 0.02^\circ\text{C}$. The outer solution was periodically sampled over 7 days. The same amount of fresh PBS containing 0.02%

Table II. Structure of Phospholipids

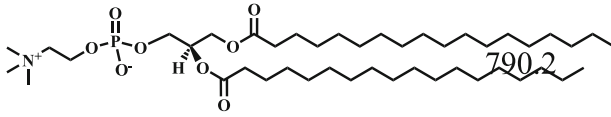
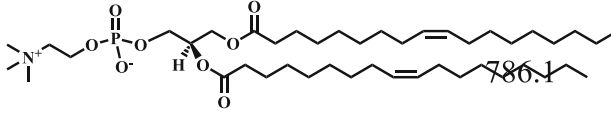
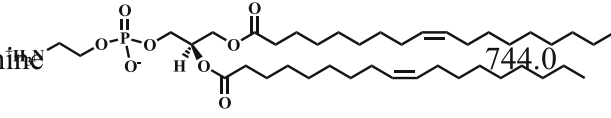
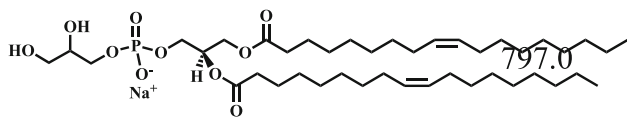
Drug	Structures	M.W.	*XLogP3
1, 2-Distearoyl- <i>sn</i> -glycero-3-phosphocholine (DSPC)		790.2	13.2
1, 2-Dioleoyl- <i>sn</i> -glycero-3-phosphocholine (DOPC)		786.1	12.8
1, 2-Dioleoyl- <i>sn</i> -glycero-3-phosphoethanolamine (DOPE)		744.0	10.6
1, 2-Dioleoyl- <i>sn</i> -glycero-3-phosphoglycerol, sodium salt (DOPG)		797.0	12.5

Table III Composition of Prepared Depot Formulations

		Mαβ									
		M _{DSPC} formulations				M _{DOPC} formulations		M _{DOPE} formulations		M _{DOPG} formulations	
α	β	DSPC				DOPC		DOPE		DOPG	
Formulation name	M _{only}	M	M _{DSPC12}	M _{DSPC24}	M _{DOPC12}	M _{DOPC24}	M _{DOPE12}	M _{DOPE24}	M _{DOPG12}	M _{DOPG24}	
C17MGE	100	88	76	64	76	64	76	64	76	64	
DSPC		–	12	24	–	–	–	–	–	–	
DOPC		–	–	–	12	24	–	–	–	–	
DOPE		–	–	–	–	–	12	24	–	–	
DOPG		–	–	–	–	–	–	–	12	24	
Ethanol							10				
Tween 80							2				

unit: percentage

Tween 80 was added to the outer compartment to maintain a constant volume. The outer solution was agitated using a pipet for 10 s prior to each sampling. The obtained samples were stored at -80°C until measurement. The cumulative percentage of LA released was calculated using the amount of LA loaded in the formulation.

In Vivo Experiments

Male Wistar rats (body weight 200 ± 20 g, 8 weeks old) were purchased from Sankyo Lab Services, Inc. (Tokyo, Japan). Rats were housed in a room maintained at $25 \pm 2^{\circ}\text{C}$ with a 12-h-dark/light cycle (on, off time: 9:00, 21:00). In addition, drinking water and feed (MF, Oriental Yeast Co., Ltd., Tokyo, Japan) were supplied ad libitum. The experiments were conducted in accordance with the experimental animal regulations of Josai University after obtaining consent (JU19009) from the Ethical Commission of Josai University.

Rats were cannulated into the jugular vein. After resting for 1 day post-surgery, the prepared formulation was administered subcutaneously via a 23G needle into the dorsal region. The administration site was shaved before administration. Blood sampling (100 μL) was conducted periodically until 28 days after administration from the cannulation tube. The same amount of saline was injected into the rats through the cannulation tube after every sampling. The blood samples were centrifuged (21,500 \times g, 5 min, 4°C) to obtain plasma. The obtained plasma was stored at -80°C until measurement. The lower limit in the quantitative detection of LA was 1 ng/mL in the present conditions. The area under the curve over 21 days after administration ($AUC_{21\text{days}}$) was calculated using the trapezoidal method. The extent of bioavailability (F) was calculated as the ratio of $AUC_{21\text{days}}$ after *s.c.* and intravenous (*i.v.*) administrations. LA solution dissolved in saline at a concentration of 4 mg/mL was administered through

the jugular vein at a dose of 5 mg/kg, whereas the prepared formulation was administered *s.c.* at a dose of 18.75 mg/kg.

LA Determination

Samples obtained in the *in vitro* release experiment were mixed with acetonitrile at a 1:1 (*v/v*) ratio and vortexed for 5 min and then used as the measured sample. Plasma obtained in the *in vivo* experiment was vortexed with acetonitrile at a ratio of 1:1 (*v/v*) for 5 min and then centrifuged (21,500 \times g, 5 min, 4°C) to use the upper layer as a measured sample.

The liquid chromatography–tandem mass spectrometry (LC-MS/MS) system consisted of a system controller (CBM-20A; Shimadzu Corporation, Kyoto, Japan), pump (LC-20 AD; Shimadzu Corporation), auto-sampler (SIL-20 AC; Shimadzu Corporation), column oven (CTO-20 AC; Shimadzu Corporation), detector (3200 QTRAP; AB Sciex, Tokyo, Japan), and analysis software (Analyst® version 1.4.2; Shimadzu Corporation). The column and the guard column were Shodex® ODP2 HP-2B 2.0 mm \times 50 mm and ODP2 HPG-2A 2.0 mm \times 10 mm, respectively (Showa Denko, Tokyo, Japan). The column temperature was adjusted to 40°C . An internal standard of betamethasone valerate was used for the LA assay. A mixed solution (A:B at a 70:30 ratio) was used for the mobile phase, where A was 0.1% formic acid in water, and B was acetonitrile. The flow rate was 0.2 mL/min, and the injection volume was set to 10 μL . Electrospray ionization was used for LA ionization. The measured molecular weight of LA was set to *m/z* 605.30 for the precursor ion and *m/z* 249.00 for the product ion. The ion spray voltage was 5000 V, the nebulizer gas pressure was 80 psi, the drying gas flow rate was 10 L/min, and the drying gas temperature was 600°C . The lower limit of quantification of this assay was 1.0 ng/mL.

Observation with a Cryo-Transmission Electron Microscope

The formulation immersed into 3 mL of PBS was observed using a cryo-transmission electron microscope (Cryo-TEM) (JEM-3100FEF, JEOL Ltd., Akishima, Tokyo, Japan). For imaging, 1 μL of a 20-fold dilution of the formulation was dropped onto a hydrophilized copper grid (200 mesh, JEOL Corporation) and blotted. The samples were rapidly frozen using ethane as a freezing solvent using a rapid freezing system (EM-CPC, Leica Microsystems Japan, Tokyo, Japan) for observation using the Cryo-TEM at 5–10 μm defocus.

Statistical Analysis

All experiments, except for SAXS analysis, were conducted with more than three replicates per formulation. JMP® Pro software (ver. 14.0.0, SAS Institute Inc., Cary, NC, U.S.A.) was used for statistical analyses. Experimental data were tested for statistical significance ($p < 0.05$) using one-way ANOVA and Tukey's honestly significant difference post hoc analysis. All data were expressed as mean with standard deviation.

RESULTS

Small Angle X-Ray Analysis of the Prepared Formulation

Figure 1 shows the small angle X-ray diffraction peak for each formulation containing LA, and Table IV shows the crystal structure obtained by analyzing the diffraction peak and the inter-pore distances (d) calculated from Bragg's equation. Figure 1a–d shows the results for M_{DSPC} , M_{DOPC} , M_{DOPE} , and M_{DOPG} formulations, respectively. Formulations M_{only} , M_{DOPE12} , and M_{DOPE24} exhibited reverse-hexagonal (H_2) (1: $\sqrt{3}$: $\sqrt{4}$) and cubic structures with Pn3m space groups ($\sqrt{2}$: $\sqrt{3}$: $\sqrt{4}$), whereas M, M_{DSPC12} , M_{DSPC24} , M_{DOPC12} , M_{DOPC24} , M_{DOPG12} , and M_{DOPG24} displayed only cubic structures with Pn3m space groups ($\sqrt{2}$: $\sqrt{3}$: $\sqrt{4}$: $\sqrt{6}$: $\sqrt{8}$: $\sqrt{9}$). Shifted diffraction peaks toward lower diffraction angles were observed in the M_{DSPC} , M_{DOPC} , M_{DOPG} , and M_{DOPE} formulations compared with M. Effect of the addition of LA in the formulation was also investigated (Table IV, SAXS data not shown). The presence of LA induced structural transformation, except for M_{DOPG} formulations.

Figure 2 shows representative cryo-TEM images of NLLC constructed formulations. Cryo-TEM images confirmed the characteristic multilayer pattern of the H_2 structure in M_{only} (a) and the presence of the ordered pattern of cubic phase in M (b) and M_{DOPG24} (c).

In Vitro Release Experiments from the Prepared Formulations

Figure 3 shows the release profile of LA from the prepared depot formulations against the square root of time. Linear lines were confirmed with each LA release. When DSPC and DOPE were included in the formulation (M_{DSPC} and M_{DOPE} , respectively), LA release was higher than from M. Higher release ratios (>40% after 7 days) were obtained from M_{sc} and M_{oe} formulations despite the phospholipid content. On the other hand, DOPG-containing formulations (M_{DOPG12} and M_{DOPG24}) showed lower LA release over 7 days, and only about 20% of the cumulative amount of released LA was observed from the M_{DOPG} formulations.

LA Profile in Blood after *s.c.* Injection of the Prepared Formulations

Figure 4 shows the changes in LA plasma concentration after *s.c.* administration of the depot preparations to a dorsal site in rats. In this experiment, the M_{DOPG} formulations were only administered in phospholipid-containing formulations because they showed lower LA release compared with the other formulations. LA solution and M were also administered to investigate the effect of C17MGE and DOPG in the formulation on the blood concentration profile of LA.

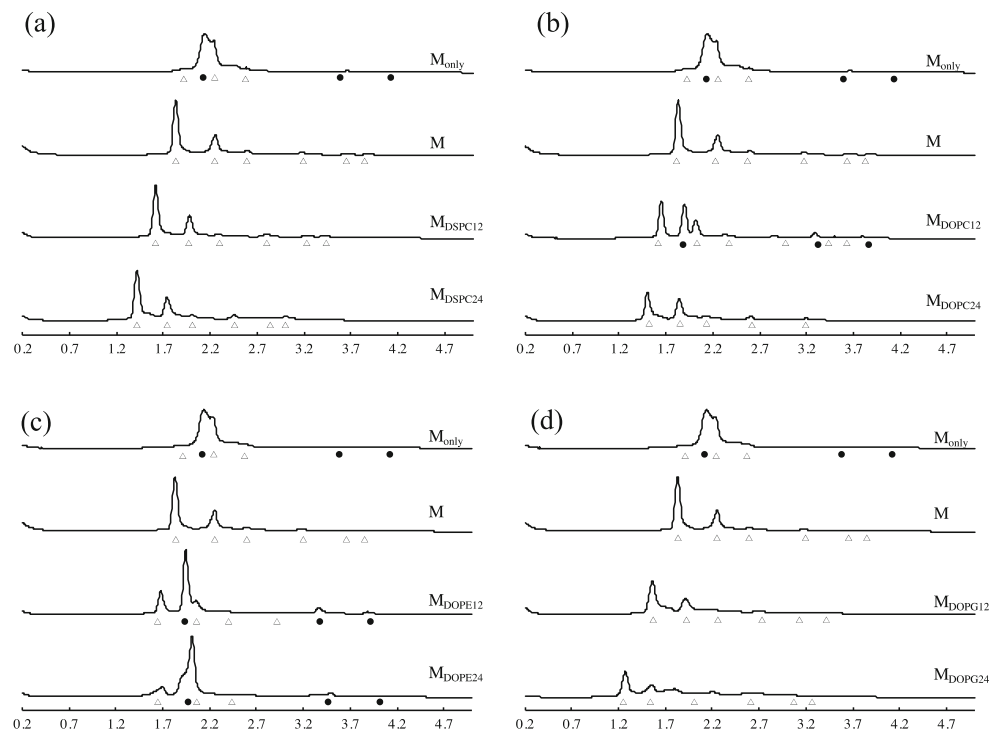
LA was rapidly eliminated from the blood, and no detection was confirmed about 6 h after the *s.c.* administration of LA solution. On the other hand, LA was detected in blood over 21 days after administration of M and Mog formulations. In particular, the M_{DOPG12} formulation provided a significantly higher LA concentration in blood compared with M at 14 days after administration.

Table V summarizes the pharmacokinetic parameters ($AUC_{21\text{days}}$, F) for each formulation. AUC was calculated using the trapezoidal rule over 21 days after administration. The F value obtained with the *s.c.* administration of LA solution was 4.31%, whereas those of the formulations containing MGE were significantly higher ($p < 0.01$) than that obtained using the LA solution. In particular, the M_{DOPG12} formulation exhibited the highest F value (89.1%) among the MGE contained formulations.

DISCUSSION

Lipid-based depot formulations have been actively studied in the last 5 years. In particular, NLLC-forming lipids have gained attention as novel excipients for depot formulations because they conveniently construct a depot upon contact with body fluids. The usefulness of a novel lipid, C17MGE, has been investigated, and it was found that formulations composed of C17MGE can take advantage of transdermal

Fig. 1 Structural analysis of constructed gel by small angle X-ray diffraction. The constructed gel was obtained by dropping the depot formulations of M_{DSPC} (a), M_{DOPC} (b), M_{DOPE} (c) and M_{DOPG} (d) into PBS. Symbols: ●; peak derived from reverse hexagonal structure, Δ; peak derived from cubic structure with Pn3m space group.



(11,12), nose-to-brain (13), oral mucosal (17), and oral (18) delivery routes compared with a conventional, commercial products. In addition, a controlled drug release system based on the NLLC structure was found to be available with

Table IV Constructed Structure and its Interplanar Spacing (d , nm) of Prepared Formulations with or Without LA

Formulation	Liquid crystal structure	d (nm)
M _{only}	$H_2 + Pn3m$ (H_2)	4.10 (4.36)
M	$Pn3m$ (H_2)	4.82 (4.36)
M _{DSPC12}	$Pn3m$ ($H_2 + Pn3m$)	5.46 (4.87)
M _{DSPC24}	$Pn3m$ (H_2)	6.22 (5.49)
M _{DOPC12}	$Pn3m$ (H_2)	5.35 (4.65)
M _{DOPC24}	$Pn3m$ (H_2)	5.87 (4.95)
M _{DOPE12}	$H_2 + Pn3m$ (H_2)	4.30 (4.55)
M _{DOPE24}	$H_2 + Pn3m$ (H_2)	4.37 (4.55)
M _{DOPG12}	$Pn3m$ ($Pn3m$)	5.65 (5.82)
M _{DOPG24}	$Pn3m$ ($Pn3m$)	6.91 (7.31)

The parenthesis shows liquid crystal structure and its d value of the formulation without LA that calculated from SAXS observation results

C17MGE-based formulations (16). Therefore, we investigated the development of C17MGE-based depot formulation that accomplished a sustained release and improved bioavailability of LA, a model middle molecular weight drug, in the present study.

Many reports have been published on controlled drug release with NLLCs. Fong et al. reported temperature-dependent drug release by changing the NLLC structure from a cubic structure to a reverse-hexagonal structure, and vice versa (19). Báez-Santos et al. (20) showed that drug release was increased by alternation of the inter-pore distances in the structure caused by increased water content in NLLC formulations composed of phosphatidylcholine, Span 80, and tocopherol acetate. Tocopherol acetate was added to the formulations as a phase transition modifier. Various other phase transition modifiers have been reported, such as tetradecane, limonene, and triolein (10). However, few reports have studied the influence of phospholipids on the phase behavior and entrapped drug release from NLLC structures. In the present study, different phospholipids were used as phase transition modifiers for C17MGE-based NLLC structures.

Small angle X-ray analysis of the prepared formulations showed that their structure was affected by the type and content of phospholipid in the formulation (Fig. 2). In general, the relationship between the structure of amphiphiles, including phospholipids, and the shape of the self-assembled structures can be considered with the critical packing parameter (CPP) (10). DSPC and DOPC, which were used in this study, are cylindrical in shape with $CPP \sim 1$ and are known to form

Fig. 2 Representative cryo-TEM of M_{only} (a), M (b), and M_{DOPG24} (c) after immersion in PBS for 24 h. Arrows show the presence of a multilayer pattern in (a) and the presence of an ordered pattern (b) and (c). Black bar: 50 nm.

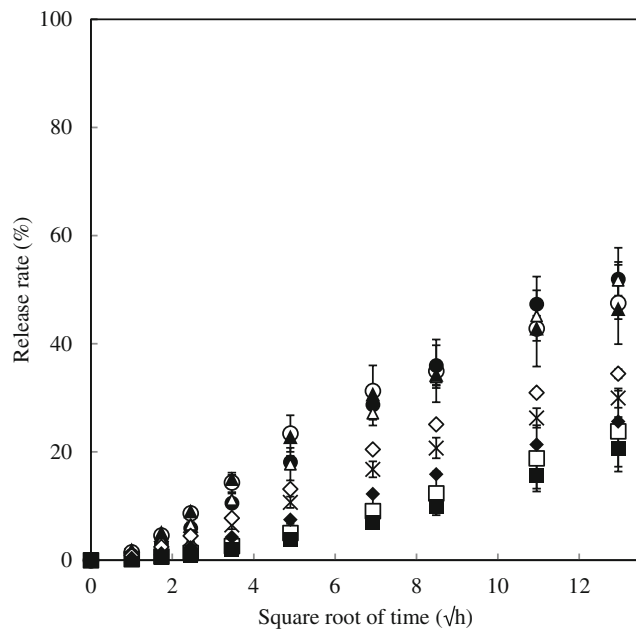
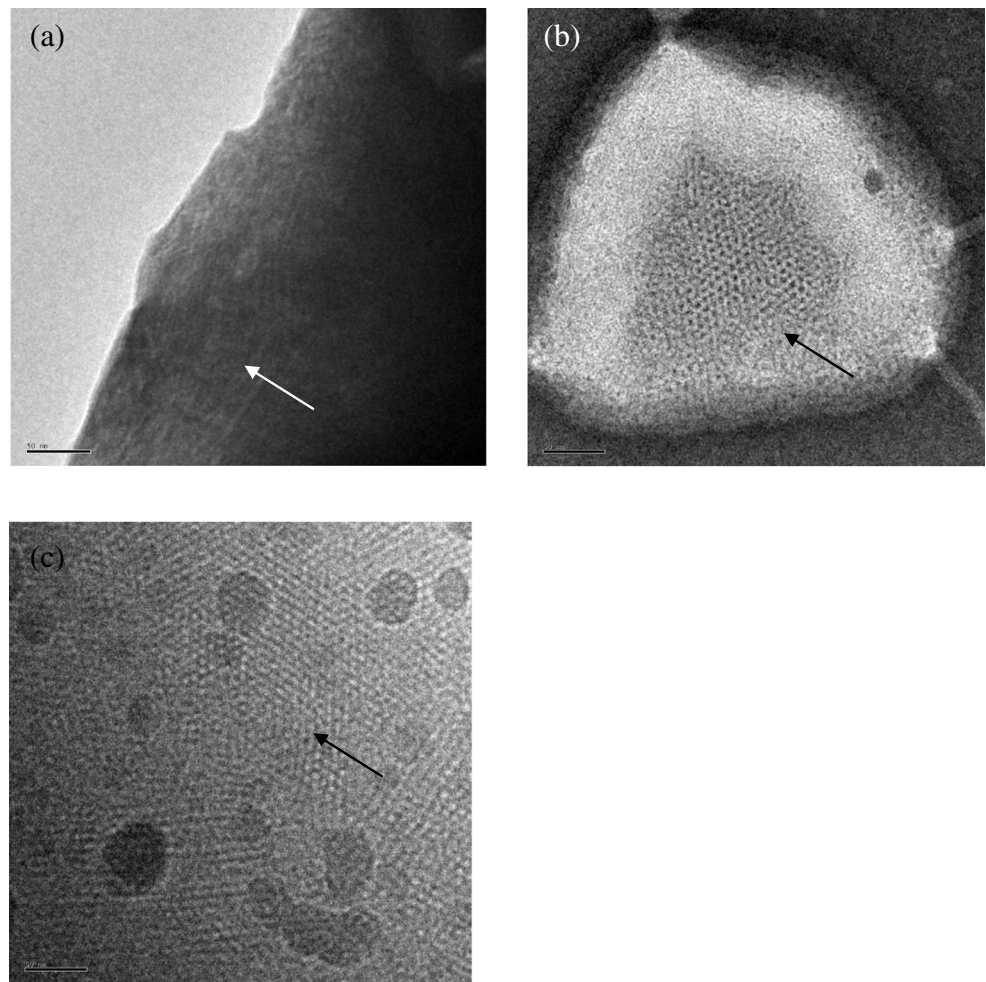
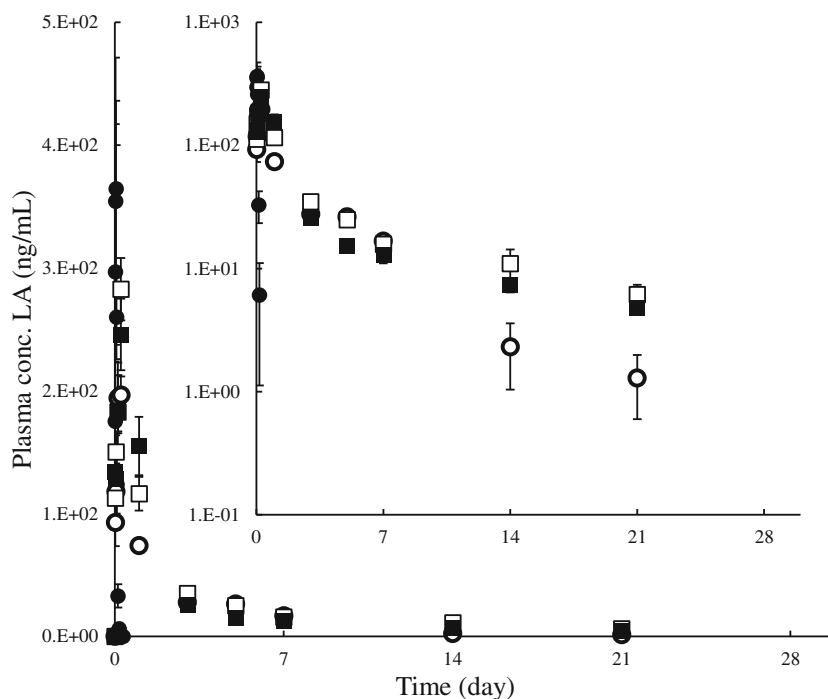


Fig. 3 LA release profile from formulations over 7 days. Symbols: M (x), M_{DSPC12} (o), M_{DSPC24} (●), M_{DOPC12} (◊), M_{DOPC24} (◆), M_{DOPE12} (Δ), M_{DOPE24} (▲), M_{DOPG12} (□), M_{DOPG24} (■). Each value shows the mean \pm S.D. ($n = 3-5$).

bilayer vesicles. DOPE and DOPG have $CPP > 1$ due to their relatively small headgroups compared with the other phospholipids. The addition of phospholipids with $CPP \sim 1$ and $CPP > 1$ in the formulation was thought to provide a phase transition toward lamellar phases and H_2 phases, respectively, by decreasing and increasing membrane curvature, respectively. However, peak shifts toward the lower angle and increased interplanar spacing were observed by increasing the phospholipid amount without a change in the NLLC structure, except for M_{DOPE} formulations (Fig. 2). Yaghmur et al. (21) reported that diglycerol monooleate ($CPP \sim 1$) interacted with the polar head group of monolinolein to increase water channel dimension in a monolinolein-based phase structure. Therefore, the addition of phospholipids with $CPP \sim 1$ may contribute to the induction of increased water channel dimensions. Another factor that decreases membrane curvature is $CPP > 1$ with a charged headgroup such as DOPG. A high phospholipid content is thought to contribute to the stabilization and phase transition of the formed NLLC structure. In the present study, phase transition was only observed in the case of increasing negative curvature by the addition of DOPE.

Fig. 4 Plasma LA concentration–time profile over 21 days after subcutaneous injection of LA sol. (●), M (○), M_{DOPG12} (□), M_{DOPG24} (■). Each point shows the mean ± S.E. (n = 3–5).



In the present study, no SAXS observation, but a polarizing microscopic observation was performed with collected formulation after 6–12 h from the administration site. Specific polarization images derived from NLLC structure were confirmed (data not shown). Therefore, construction of NLLC structure was thought to be obtained at administration site by body fluid after *s.c.* injection.

Release rates of entrapped drugs are faster in the order; lamellar phase, cubic phases, hexagonal phases, and micellar cubic (22,23). However, M_{DOPG} formulations with an H_2 structure showed higher LA release compared with M and M_{DOPC} formulations with Pn3m space groups. In our previous study, the release ratio of fluorescein isothiocyanate dextran (M.W., 4000, FD-4), a hydrophilic drug, from a constructed NLLC structure was increased with an increase in interplanar spacing, despite the difference in the constructed

structure (18). On the other hand, no relationship was found between LA released from the formulations and the value of interplanar spacing in the present study (data not shown). Hydrophilic drugs are located in the water channels or close to the polar head, while hydrophobic drugs and amphiphilic drugs are loaded in the lipid layer and at the lipid layer interface, respectively (22). When effect of the addition of LA on the constructed structure was evaluated with SAXS, the structure was affected by the addition of LA, except for M_{DOPG} formulation (Table IV). This result suggested that LA interacted with the MEG or/and phospholipid, which in turn caused the phase transition. The interaction would be altered by a hydrophilic group in the MGE or /and phospholipid and filled void volume in the structure. Therefore, these interactions might be a reason for higher LA release from M_{DOPG} formulation compared with M and M_{DOPC} formulations in the present study.

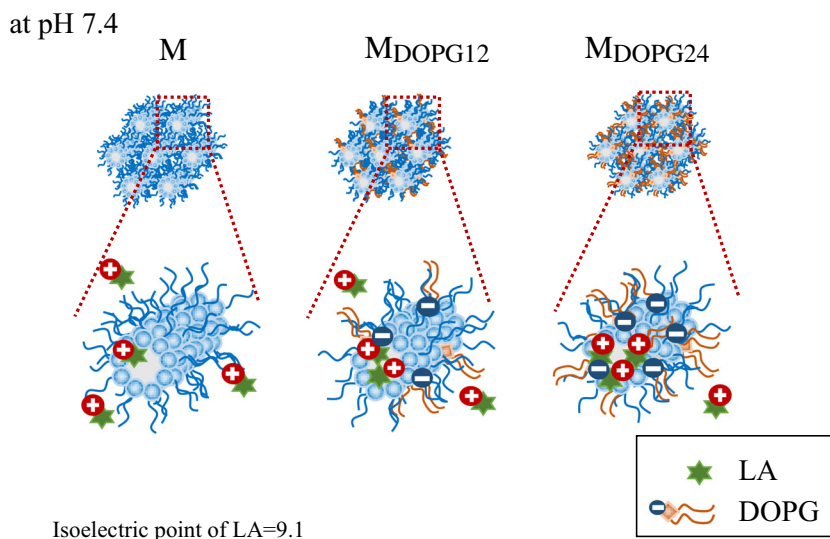
M_{DOPG} formulations showed sustained release compared with the other formulations. Figure 5 shows a schematic representation of electrostatic interaction between LA and DOPG at pH 7.4. LA has a positive charge at pH 7.4 in the solvent environment (isoelectric point of LA = 9.1), and the hydrophilic group in DOPG shows a negative charge, as shown in Fig. 5. Many reports have observed controlled drug release using electrostatic interactions (24,25). Lim et al. reported that the sustained release of positively charged entecavir was observed after injection by interacting with anionic phospholipid of 1, 2-dipalmitoyl-*sn*-glycero-3-phosphatidic acid in acidic conditions (pH 2.5–4.5) (26). M_{DOPG12} showed a sustained release of LA, suggested that 12% of DOPG in the

Table V Pharmacokinetic Parameters of Leuprolide Acetate After S.C. Co-Administration in Rats

Route	Dose (mg/kg)	Formulations	$AUC_{21\text{days}}$		F
			$(\mu\text{g}/\text{min}/\text{mL})$		(%)
<i>i.v.</i>	5	–	228.86 ± 27.53	–	± –
<i>s.c.</i>	18.75	Solution	36.48 ± 4.57	4.31 ± 0.54	
		M	540.55 ± 41.75 *	63.84 ± 4.93	
		M _{DOPG12}	754.06 ± 44.47 *	89.05 ± 5.25	
		M _{DOPG24}	660.71 ± 69.31 *	78.03 ± 8.19	
F = [(AUC _{s.c.} / Dose _{s.c.}) / (AUC _{i.v.} / Dose _{i.v.})] * 100					± S.E.

* $p < 0.01$ Compared with *s.c.* administration of LA solution

Fig. 5 Schematic representation of electrostatic interactions between LA and DOPG at pH 7.4 for M, M_{DOPG12} and M_{DOPG24} .



formulation was enough to control the release, and electrostatic interactions between the negatively charged hydrophilic groups of DOPG and positively charged LA may be a reason for the sustained release.

The M and M_{DOPG} formulations exhibited sustained LA blood levels after *s.c.* injection. The plasma concentration of LA 21 days after *s.c.* injection was in the following order; $M_{DOPG12} > M_{DOPG24} > M$, whereas the amount of LA released over 7 days was in the following order; $M > M_{DOPG12} > M_{DOPG24}$. Unlike the *in vitro* condition, the water absorption rate into the administered formulation might be low due to body fluid restrictions in *in vivo* conditions. Relatively prompt NLLC construction changes from the administered formulation occurred in *in vitro* conditions, which was due to the plentiful existence of water compared with *in vivo* condition. On the other hand, the construction change in the NLLC structure may occur gradually from the peripheral part of the administered formulation even in *in vivo* conditions, and it may take time to construct the NLLC structure in the deeper part of the formulation. This might be a reason for the discrepancy in the observed difference between the release rate observed in the *in vitro* and *in vivo* profiles. When LA release rates of M, M_{DOPG12} and M_{DOPG24} were calculated from *in vitro* release profiles with four successive points from the last (48 h, 72 h, 128 h, and 168 h), the release rate was following order; $M (0.46 \text{ h}^{-1}) > M_{ogDOPG24} (0.44 \text{ h}^{-1}) > M_{ogDOPG12} (0.40 \text{ h}^{-1})$. M_{og12} showed the highest blood concentration of LA, and Met displayed the lowest one among the formulations. Therefore, the LA release rate after sufficient time passed to construct the NLLC structure in *in vitro* experiment was considered to be reflected in the *in vivo* results. A robust correlation of both the rate of drug release obtained from *in vitro* release experiments and the real rate of drug release into the body is still difficult to determine

for depot formulations (8). Therefore, the establishment of an *in vivo-in vitro* correlation (IVIVC) might help in developing depot formulations.

The calculated F value obtained from M_{DOPG12} was about 90%, and the blood concentration of LA was still detectable 21 days after administration. AUC_{∞} was calculated by summation of $AUC_{21\text{days}} + \text{Conc.}_{21\text{days}}/k_{el}$, where $\text{Conc.}_{21\text{days}}$ represents the plasma concentration at 21 days after administration and k_{el} is the elimination rate constant. The estimated F with calculated AUC_{∞} reached almost 100%. The same calculation was applied to the M formulation; the estimated F was about 66%. On the other hand, the F value obtained after *s.c.* injection of LA was about 48%. Instability of LA in the skin tissue was reported, with over 90% of LA degraded within 120 min (15). LA was predominantly metabolized by hydrolysis (27). Several kinds of enzymes are present in the skin, such as peptidases (28). Therefore, the lower F value obtained after *s.c.* injection of LA solution may be ascribed to hydrolysis by such enzymes. On the other hand, an NLLC structure has the potential to protect against enzymatic degradation (29,30). Thus, the stability of LA in skin may be also contributed to by higher F values. The stability of peptides and proteins entrapped in the formulation is one of the most critical considerations for sustained drug delivery. It has been reported that protein instability is attributed to local pH acidification caused by poly(lactic-co-glycolic acid) degradation (31). Therefore, lipid-based depot formulations using NLLC might be a promising alternative to PLGA formulations. Throughout *in vivo* experiment over 21 days, no severe toxicities were detected (no weight loss, no respiratory failure and no inflammation at the administrated site). Further safety evaluation experiments should be done to show the usefulness of MGE-based depot formulation.

CONCLUSION

The controlled release of LA from a C17MGE depot was possible by adding various phospholipids. In particular, M_{DOPG12} showed a highly sustained LA release and improved absolute bioavailability. Recently, hollow microneedles have been developed to achieve direct intradermal delivery of drugs. However, the relative bioavailability of LA from dissolving microneedles was low due to the instability of LA in the skin (25). Therefore, a combination of a microneedle device and NLLC formulation composed of C17MGE may improve instability and accomplish controlled release of LA. The NLLC formulation used in this study might be used as a depot base with controlled drug release, although further investigations are needed.

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