Preparation of Hollow Fiber Immobilized Liposome Membrane

Hiroyuki Sugaya¹, ², Hiroshi Umakoshi¹, Yuji Tohtake¹, Ena Oyama¹, Toshinori Shimanouchi¹, and Ryoichi Kuboi¹)*

¹) Division of Chemical Engineering, Graduate School of Engineering Science, Osaka University, 1-3 Machikaneyama-cho, Toyonaka, Osaka 560-8531, Japan
²) Specialty Materials Research Laboratories, Toray Industries, Inc., Otsu, Shiga, Japan

Immobilized liposome membrane (ILM) was prepared in hollow fiber module (HF–ILM), where the liposome was loaded into the hollow fiber membrane module and was physically immobilized by forming the hydrogel in the cavity of the fiber membrane. Basic nature of the hydrogels prepared with xanthan gum (XG) and polyethyleneimine (PEI) via amino coupling method was first investigated in relation to the water contents of the gel matrix, so that the optimal condition for the liposome immobilization was obtained. The HF–ILM was prepared at the optimal condition, resulting in the high contents of immobilized liposome with high stability.

Key words: membrane stress biotechnology / hollow fiber membrane / liposome

1. Introduction

Liposome, a closed phospholipid bilayer membrane, has a nano-order interface (~5 nm) harboring hydration layer and low–k (hydrophobic) layer on its surface. It has been reported that the liposome could recognize the molecule through the combined interactions, such as electrostatic and hydrophobic interaction and hydrogen bonds stability ¹–³). Some new aspects of the liposome membrane itself, which could be induced under stress condition, have recently been reported ⁴): (a) molecular chaperone–like function to assist the protein refolding ⁵–⁷), (b) protein translocation across the membrane ⁸), (c) function as a mediator/initiator of membrane fusion ⁹–¹¹), (d) LIPOzyme functions (Liposome + Enzyme) ¹²–¹⁹). The use of the liposome (or LIPOzyme) could let us design and develop the liposome-based separation/reactor system.

The immobilization of such a liposome is an important technique for the above-mentioned purposes. It has recently been reported that the liposome can be utilized as a molecular recognition element in several analytical methods, such as (i) immobilized liposome chromatography (ILC) / immobilized liposome membrane (ILM) ¹, ³, ²⁰) and (ii) immobilized liposome sensor (ILS) ²¹–²³). The above analytical methods enable us to obtain the independent parameters of the liposome molecule interaction, such as charge, hydrophobicity, and hydrogen bond stability. At the same time, the above immobilized liposome techniques can also be applied for the stress–mediated mutual separation of proteins ²⁰) and for the protein refolding column ¹). In these immobilized liposome matrixes, the liposome was immobilized via (a) physical entrapment method ²⁴), (b) antigen–antibody method ²⁵), (c) hydrophobic ligand method ²⁶), (d) covalent binding method ²⁷) and so on. Although these immobilization methods can be utilized for the bioseparation uses, there are some problems such as the capacity of immobilized liposome, the stability of the immobilized liposome, difficulty or non-scalability of the column operation and so on. It is important to develop a new immobilized liposome matrix to overcome the above problems.

Hollow fiber membrane module is widely applied for the bioremediation, water purification, biomedical uses and could be a powerful tool to solve the above problems ²⁸). Although a possible use of the planer membrane as a support of the immobilized liposome has been previously reported, the stability of the liposome is not so high to apply it for the practical uses.

* Corresponding Author
Tel & Fax : +81-(0)6-6850-6286
E-mail : MSB@cheng.es.osaka-u.ac.jp
Membrane module, consisted of hollow fiber membranes, has been produced in some membrane companies and utilized in the many research areas. Considering the morphology of the membrane itself, more effective method for the liposome immobilization could be developed. Polysulfone (PS) hollow fiber membrane, produced in Toray, is a porous membrane and the pore size is varied in a gradual manner (wide from the outside, narrow to the inside of a fiber). It is expected that the liposome could be easily immobilized through its loading into such a porous space and the following immobilization via polymer gel formation as schematically shown in Fig. 1.

In this study, a method for the liposome immobilization was newly established by using the morphological characteristics of the hollow fiber membrane (HF–M) module produced by Toray. The liposomes were first loaded into the porous space of the membrane module and the hydrophilic gel matrix was formed inside a porous space. The basic characteristics of the hollow fiber type immobilized liposome membrane (HF–ILM) were finally investigated, focusing on the immobilized amounts of liposome, the stability of the immobilized liposome and the characteristics of the membrane module itself.

2. Experimental

2.1 Materials

1-Palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine (POPC) was purchased from NOF Co. Ltd. (Nagoya, Japan). Polyacrylate (PA), hyalurolic acid (HA), and xanthan gum (XG), used for the hydrophobic COOH-harboring polymer, were purchased from the Wako-Pure Industries (Osaka, Japan). Carboxymethyl–cellulose (CMC, product ID: 039–01335) was also purchased from Wako. Polyethyleneimine (PEI), used for the hydrophobic NH2–harboring polymer, were purchased from the Sigma–Aldrich (USA). Rhodamine–phosphatidylethanolamine (Rh–PE), used for the modification of liposome membrane, was purchased from Molecular Probes (MO, USA). Water soluble carbodiimide hydrochloride (WSC, 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide hydrochloride), calcein, and HOBt (1-Hydroxy-1H-benzotriazole, monohydrate) were purchased from Dojindo Chemicals (Tokyo, Japan). Other reagents of analytical grade were purchased from Wako.

2.2 Liposome Preparation

The POPC liposome including 1 mol% Rh–PE and 1 mol% biotinyl–phosphatidyl ethanolamine was prepared by using the following procedure according to the previous work. For the preliminary experiment to test the clearance of the module, the POPC liposome including 30 mol% cholesterol was also prepared. The phospholipid was dissolved in chloroform/methanol. After the solvent was evaporated, the resulting thin film was dried for at least 2 h under a vacuum. The lipid film was hydrated by pure water or 100 mM calcein solution to form the multilamellar vesicles. The solution of the multilamellar vesicle was frozen in dry ice–ethanol (–80 °C) and incubated in the water bath above the phase-transition temperature. The above freezing-thawing treatment was repeated five times to prepare multi-lamellar vesicles (MLVs) with larger size. In order to prepare the single unilamellar vesicle with constant size, the MLVs were ultrasonicated to adjust the liposome size as 30 nm (for the preliminary experiment to confirm the clearance property of the module against the liposome solution) or were treated with the extrusion device equipped with stacked polycarbonate filters of 100-nm pore size (for the immobilization experiment in membrane module).
2.3 Gel Preparation

The hydrogel of anionic polymer and cationic PEI was prepared by using the amino coupling method. After the anionic polymer solution (0.25 wt%), such as xanthan gum (XG), polyacrylate (PA), and hyaluronic acid (HA), was activated with WSC (11 mg/ml) and 11 mg/ml HOBt, the PEI solution at different concentrations (0.1 ~ 1 wt%) was mixed with the above anionic polymer solution at room temperature. The water contents of obtained hydrogel were analyzed by calculating the weight of the hydrogel before and after the drying. In the case of the hydrogel formation on the membrane of the hollow fiber module, the POPC liposome was first loaded into the hollow fiber membrane. After the clearance of the liposome solution was confirmed, anionic polymer (XG (0.5 wt%)) activated by WSC/HOBt was then loaded into the membrane module for 20 min and, secondly, the PEI solution was applied to the membrane module for 30 min.

2.4 Membrane Module Operation

Hollow fibers were obtained from a commercial polysulfone (PS) dialyzer (Toraysulfone TS-U, Toray, Japan). This commercial dialyzer was broken up to retrieve the hollow fibers. A mini module, used for the housing of the immobilized liposome membrane, was prepared by using the retrieved hollow fibers (100 fiber membranes). The membrane module was connected to the silicone tube with the inner diameter of 1 mm with the length of 130 cm. The total volume of the module and the lines were determined by the preliminary experiment and was found to 3 ml. The peristaltic pump was equipped in the flow line of silicone tube. A manometer was set at the side of the filtrate. Before the sample loading, the water solution was applied to avoid the possible contamination by the impurities.

2.5 Measurements

The concentration of the liposome was monitored from the optical density of the solution at 405 nm (OD405). The amounts of leaked calcein loaded in the liposome were determined from the absorbance at 415 nm. The total amount of calcein in the liposome immobilized in membrane module was determined from the calcein leakage after washing module by 1 wt% Triton X–100 solution to disrupt the liposome structure.

3. Results and Discussion

3.1 Polyelectrolyte-Conjugating Polymer Complex to Form Hydrogel

The anionic polymers harboring COOH group can be conjugated with polycationic polymer harboring amine group via amino–coupling method. There are some anionic polymers which are known as a biocompatible polymers in respect to the hydrophilicity and also the high water contents in the formed gel. The preparation and characterization of the conjugated polyion complex were carried out as the preliminary experiment to prepare the immobilized liposome membrane module.

3.1.1 Effect of Polymer Type on Hydrogel Preparation

By using several kinds of polymers, such as polyacrylate (PA), carboxymethyl-cellulose (CMC), hyaluronic acid (HA), and also xanthan gum (XG), the hydrogel was formed by using polyethylene imine (PEI) as an amino–donating polymer through the amino coupling method using WSC. Fig. 2 shows the water contents of the gel prepared by using different kinds of anionic polymers and PEI at its same concentration. Among the possible polymers, the maximal values was obtained in the case of the XG and the value was reduced in the following order: XG > HA > CMC > PA. From the observation of the prepared of hydrogel, the above water contents could also be related to the stiffness of the hydrogel. For example, although the PAA/PEI gel showed solid–like nature, the HA/PEI
gel and XG/PEI gel were found to be soft. However, the HA/PEI gel became hard after the completion of the amino–coupling reaction for more than four hours.

Such tendency could be caused by the density of the carboxyl-group on the surface of single polymer. PAA has a COOH group per single monomer unit, implying that the multi–conjugation points could be formed on the single polymer surface. The COOH density of CM-Cellulose is also high because all the OH–groups on CMC are substituted by carboxy methyl group. On the contrary, both HA and XG are known to be utilized as hydrophobic polymer and have a lower COOH density per monomer unit of the polymers: one and half COOH groups per two glucan molecules (two glucosaminoglucon) in the case of HA and one COOH group per branched five glucan structure. The theoretical density of the COOH group in the single unit of the polymer was also plotted in Fig. 2. The COOH density was well corresponding with the water contents of the gel consisting of the polymers and the PEI.

It was thus found that the XG/PEI gel was optimal as a soft–polymer interface to modify the PS membrane module. The XG/PEI gel was used in the following experiment.

### 3.1.2 Effect of Operational Condition for XG/PEI Gel Preparation

The effect of the preparation condition on the XG/PEI gel formation was investigated. The obtained results were shown in Fig. 3. As shown in Fig. 3(a), the effect of the PEI concentration on the water contents of the hydrogel was first investigated. At the lower concentration region of PEI, the hydrogel was not formed at all. Above the critial PEI concentration (0.25 mg/ml), the XG/PEI gel was formed. It has been reported that the XG and PEI solution could form the polion complex in the aqueous solution because of the formation of hydrophobic and less soluble precipita-

---

**Fig. 3** Effect of operational condition on the water contents of XG/PEI Gel.

---

Fig. 3(a) plot the relationship between the PEI concentration and the water contents of the hydrogel.

Fig. 3(a) shows the relationship between the molecular weight of PEI and the water contents of the hydrogel. The water content with lower molecular weight of PEI was found to be much higher than that with higher molecular weight. As discussed in the previous section, the hydrogel could show solid–like nature through the increase of the cross–linking points between the polymers. The above results on the effect of molecular weight of PEI imply that the concentration of the poly-

The addition of liposome itself, to be immobilized on the hollow fiber module, on the water contents of the hydrogel was investigated. The addition of the liposome did not affect the water content of the XG/PEI gel (data not shown).

It was thus found that the 2 mg/ml XG and 2.5 mg/ml PEI could be effective for the appoprate gel formation in water solution considering the water contents of the prepared gel.
3.2 Preparation of Immobilized Liposome Membrane Module

The immobilized–liposome membrane module, schematically shown in Fig. 1, was prepared based on the above pre-investigation on the hydrogel formation. The operational scheme for the liposome immobilization was shown in Fig. 4. The dialysate side (outside of the fiber membrane) of the membrane module was connected to the reservoir including samples such as liposome and polymers, where this flow line was equipped with peristaltic pump and manometer (Fig. 4(a)). This is because the wide cavity of the channel existed in this dialysate side. The filtrate was recovered from the blood side (inside of the fiber membrane) through the flow line equipped with peristaltic pump (Fig. 4(a)).

The POPC liposome with 100 nm was first loaded the PS membrane module from the dialysate side (from housing side) with circulation mode (Fig. 4(b) Step I), confirming that all the liposome (more than 99%) was cleared in 40 min at 1 ml/min from the reservoir based on the optical density (a typical example was shown in Fig. 5). The aqueous solution of XG (1 mg) activated by WSC was then applied to the module for 20 min to fill the empty space between the liposomes on the PS membrane surface. Almost all XG molecules were entrapped on the PS membrane because of its large molecular weight (average molecular weight: 2000 kDa) (Fig. 4(b) Step II). At the final stage, the aqueous solution of PEI was loaded to the module in order to form the XG/PEI gel on the surface of PS membrane (Fig. 4(b) Step III). After this operation, the pressure drop between the PS membrane was increased from 2.0 kPa to 4.7 kPa, showing that the gel was formed inside a channel of the PS membrane. Although the resistance of the water filtration was increased after the gel formation, the value was found.
to be high enough to be utilized for the reactor and/or separator.

The photo image of the immobilized liposome membrane module was shown in Fig. 6. The POPC liposome was herewith colored with fluorescence probes, such as 100 mM calcein (entrapped inside inner aqueous phase, Orange Color) and Rhodamine–PE (modified with membrane itself at 2 mol%, Red Color) in order to estimate the immobilized amounts of liposome. In contrast to the normal membrane module (Fig. 6, Control), the fluorescence–labelled liposome was observed in the hollow fiber module (Fig. 6, Module (A)), clearly showing that the liposome was entrapped on the PS membrane surface. In addition, the back pressure was applied from the inner side of the membrane fiber to outer side for 20 min. A significant change in the fluorescence was not observed although the liposome loaded without the formation of XG/PEI gel was leaked from the membrane module under such a back pressure condition (data not shown). The result on the back pressure experiment shows that the liposome was entrapped inside the gel matrix of XG/PEI. As a negative control experiment, the gel formation was performed without sequential addition of XG (Step II) and PEI (Step III), where the XG/PEI mixture was applied to the module after the liposome loading (Step I). Although the color of calcein and Rhodamine–PE was observed similarly, the fluorescence color was not deep (Fig. 6, Module (B)).

3.3 Characterization of Immobilized Liposome Membrane Module

The obtained membrane module immobilizing the liposome membrane (HF–ILM) was further characterized, focusing on the (i) immobilized amount of the liposome, (ii) stability of the immobilized liposome and (iii) filtration constant of the membrane module itself.

The calcein was used as a measure of liposome immobilization because the calcein can be easily released by the addition of an excess amount detergent. The amount of calcein leakage is proportional to the concentration of liposome entrapped because the calcein concentration inside the liposome is constant at 100 mM. The Triton X–100 solution (1 wt%) was applied to the membrane module for the solubilization of the liposome membrane and the total amounts of calcein leakage were measured as shown in Table 1. The calcein leakage of sequential immobilization (Fig. 6, Module (A)) was found to be 10 times higher than that of the non–sequential immobilization (Fig. 6, Module (B)). This could be because of the disruption of the liposome during the XG/PEI gel formation. In the case of the sequential loading of the polymer, the XG polymer was first packed inside a channel and the PEI cross–links the packed XG gel, resulting that the liposome was not disrupted during this operation.

The result on the immobilized liposome, determined from the calcein leakage, was compared with previous data on the immobilized liposome chromatography (Table 1). The amount of immobilized lipid shows 820 μmol/ml–PS–membrane and is 27 ~ 40 times larger than that in the case of immobilized liposome hydrogel 32, 33). The difference of the present membrane module and previous immobilized liposome gel matrix is caused by the difference of the immobilization method. In the case of the immobilization of liposome on the gel matrix, the surface area of the gel surface to be immobilized was restricted 32). However, the present method enables us to immobilize the liposome because the liposome was first loaded into hollow fiber module with highly porous membrane. In addition, various kinds of liposome could also be immobilized by using this method because the loaded liposome was physically immobilized through the creation of gel matrix in between the liposomes.

The stability of the liposome immobilized on the PS membrane surface is also important factor for the practical use 32, 33). The time course of the calcein leakage was monitored after the above HF–ILMs were pre-
pared as shown in Fig. 7. The calcein leakage was not observed in the case of module although the calcein was leaked in the case of module B. The calcein leakage from the immobilized liposome in gel matrix was previously reported, where the calcein leakage was not observed at least for 3 weeks 32). In the case of the lipid membrane in the extremely porous membrane (cellulose acetate membrane), the life time of the calcein leakage was not so high (5 days) owing to the high shear stress of the liquid flow across the filter membrane (data not shown). The immobilization of the liposome on the SAM–supported solid surface was also performed and the immobilized liposome was found to be ruptured in one week. The lifetime of the present HF–ILM was found to be at least two weeks without significant leakage of the calcein. The above results show that the stability of the immobilized liposome is applicable for the analytical use and, also, preparation use.

The filtration constant was also shown in Table 1. The value was slightly reduced in the case of immobilized liposome PS module. It has been reported that 230 ml/(hr m² kPa) in UFRS value is applicable for the dialysis use. The obtained value was lower than the

<table>
<thead>
<tr>
<th>Matrix Type</th>
<th>Immobilization Method</th>
<th>Amount of Immobilized Lipid [μmol/lipid/ml-matrix]</th>
<th>Stability (Calcein Leakage [%])</th>
<th>UFRS [ml/hr m² kPa]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control Module</td>
<td>PS Membrane</td>
<td>None</td>
<td>0</td>
<td>--</td>
</tr>
<tr>
<td>ILM-Module-(A)</td>
<td>PS Membrane</td>
<td>XG/PEI(Sequential)</td>
<td>820.0 (*1)</td>
<td>0.89 (*5)</td>
</tr>
<tr>
<td>ILM-Module-(B)</td>
<td>PS Membrane</td>
<td>XG/PEI(Preforming-Gcl)</td>
<td>49.1 (*1)</td>
<td>72.0 (*5)</td>
</tr>
<tr>
<td>ILC</td>
<td>Sepharose</td>
<td>Avidin-Biotin</td>
<td>37.6 (*2)</td>
<td>0.53 (*6)</td>
</tr>
<tr>
<td>ILC</td>
<td>Sephasclry</td>
<td>Avidin-Biotin</td>
<td>41.5 (*2)</td>
<td>0.77 (*6)</td>
</tr>
<tr>
<td>ILC</td>
<td>Sephadex</td>
<td>Avidin-Biotin</td>
<td>30.2 (*2)</td>
<td>1.08 (*6)</td>
</tr>
<tr>
<td>ILC</td>
<td>TSK</td>
<td>Avidin-Biotin</td>
<td>30.3 (*2)</td>
<td>0.74 (*6)</td>
</tr>
<tr>
<td>ILM</td>
<td>TSK</td>
<td>Covalent Binding</td>
<td>34.2 (*3)</td>
<td>--</td>
</tr>
<tr>
<td>ILM</td>
<td>Cellulose</td>
<td>Covalent Binding</td>
<td>14.5 (*4)</td>
<td>81.2 (*7)</td>
</tr>
</tbody>
</table>

*1 The amount of the immobilized lipid was calculated from the calcein leakage after the wash of the non–entrapped calcein outside the liposome by buffer solution and the solubilization of immobilized liposome by using Triton X–100 solution. Triton X–100 solution more than its critical micelle concentration is known to disrupt the liposome structure, resulting in the leakage of the calcein from the immobilized liposomes. The total amount of calcein leaked from the module is equivalent to the total calcein entrapped inside the liposome immobilized in the module. The number of liposomes can be estimated from the total amounts of leaked calcein because the calcein concentration inside a liposome is set at 100 mM. The number of lipid molecules of one liposome (size: 100 nm) can be calculated as 79725 by assuming 0.72 nm² as surface area of single lipid and 4.5 nm as the bilayer thickness.

*4 Data from Unpublished Work
*5 The value was calculated as the accumulated amounts of calcein for several days at 25 °C against its total amounts
*7 The value was recalculated based on data from unpublished work
maximal level of the filtration resistance, showing that this immobilized liposome module can be applied for the practical usage.

It was thus found that the prepared HF–ILM can be used in relation to the (i) immobilized amounts of liposome, (ii) its stability, and (iii) filtration nature.

4. Conclusion

The immobilized liposome membrane module was newly developed by employing the sequential immobilization technique of liposome and polymers for gel matrix. The XG/PEI gel was employed as a gel for the liposome immobilization. The immobilized liposome membrane module, prepared at optimal condition, was found to exceed in the following aspects: (i) high immobilization, (ii) high stability, (iii) high efficiency. Considering the immobilization principle, this method could be applied for the immobilization of various kinds of liposome. It has previously reported that the liposome has a molecular chaperone–like function and the immobilized liposome gel column was applied to the protein refolding column. Recently, the liposome could induce the enzyme–like function (LIPOzyme). The previous findings show that the liposome could be utilized as a catalytic material. The combination of the liposome and/or LIPOzyme with the present immobilized liposome membrane module could enable us to develop the liposome-based bioreactor. In addition, the liposome immobilization could change the biocompatibility of the membrane surface itself. The obtained membrane module could contribute to the possible applications shown above. The possible applications are now under investigation.

Acknowledgement

The fundamental concept of this study was supported by the Research Group of “Membrane Stress Biotechnology” and “Engineering Science of Liposome”. It is partly supported by a Grant–in–Aid for Scientific Research (No. 15206089, 16686046, 16760635, 17656268, 19656203, 19656220, and 20360350, 20760539, 21246121) from the Ministry of Education, Science, Sports, and Culture of Japan, a grant from the 21st Century COE program “Creation of Integrated EcoChemistry” and the Global COE program “Bio-Environmental Chemistry” of the Japan Society for the Promotion of Science (JSPS). The authors are grateful to the Research Center for Solar Energy Chemistry of Osaka University and the Gas hydrate Analyzing System of Osaka University.

Literature Cited

15) Tuan LQ, Umakoshi H, Shimanouchi T, Kuboi R :

(Received 26 December 2008; Accepted 12 June 2009)