Design of Controlled-Release Morphine Suppositories Containing Polyglycerol Ester of Fatty Acid

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Controlled-release morphine suppositories were prepared by utilizing polyglycerol ester of fatty acid (PGEF). The addition of PGEF to fatty suppository base Witepsol H15 resulted in a decrease of morphine release rate from suppositories. Among PGEFs examined, decaglycerol heptabehenate (HB750) was the most effective additive for the controlled-release of morphine from fatty suppositories. The apparent viscosity of suppository bases increased with the increased HB750 content, and good correlation was observed between the apparent viscosity of suppository bases at 37°C and the amount of HB750 added in the mixed base. The in vitro release rate of morphine was decreased by the addition of HB750 and the release rate constant (Higuchi’s rate constant) for morphine release was significantly correlated with the HB750 content in the mixed bases as well as the apparent viscosity of mixed bases, indicating that the release of morphine from the mixed bases could be regulated by the HB750 content in the mixed bases. After rectal administration of Witepsol H-15-HB750 mixed suppositories to dogs, plasma concentrations of morphine did not increase rapidly at early time periods, but relatively high levels of morphine in plasma were sustained for longer time periods. Mean residence time of morphine was considerably prolonged without changing relative bioavailability in the case of the mixed base suppositories containing 15—17% HB750, compared with the Witepsol H15 suppository, clearly indicating that the mixed bases containing HB750 are expected to be useful for the design of controlled-release morphine suppositories.

Key words: morphine suppository; fatty suppository; polyglycerol ester of fatty acid; viscosity; release control

Morphine is widely used for the management of severe cancer pain. The oral route is the first choice in administering medication to relieve it, but clinical experience has shown that alternative routes of administration should be employed for patients suffering from nausea and vomiting. Furthermore, as morphine is subjected to the extensive first-pass metabolism after oral administration, the rectal administration of morphine, which can partially avoid the hepatic first-pass metabolism, leads to higher bioavailability, compared with the oral dosing. Therefore, the rectal route is very important for the management of severe pain by morphine. Morphine suppository preparations used clinically are not of the sustained-release type, so more than 3 doses per day are required to maintain the analgesic effect due to the rapid elimination of morphine. Therefore, controlled-release dosage forms are considered to be particularly appropriate for morphine. Sustained-release suppositories of morphine have been investigated, and the release of morphine was controlled by hydrogel, modified suppository base such as light anhydrous silicic acid (Aerosil®) or algicin acid. In another case, a controlled-release morphine tablet was investigated for rectal use. Polyglycerol ester of fatty acid (PGEF) is a solid fat and has excellent characteristics as an additive of controlled-release suppository. PGEFs have a well affinity with the fatty suppository bases, and the matrix bases can be homogeneously prepared by adding PGEFs to suppository bases, which may provide a stable controlled-release system. In the present study, a novel type of controlled-release suppositories has been developed for morphine by using PGEF as an additive of suppository. Physicochemical characteristics of the suppositories and the release properties of morphine from the suppositories were evaluated in an in vitro study. Furthermore, in vivo absorption studies were performed in beagle dogs.

MATERIALS AND METHODS

Materials: Morphine hydrochloride was obtained from Dainippon Pharmaceutical Co., Ltd. (Osaka, Japan). Witepsol H15 was purchased from Mitsuba Trading Co., Ltd. (Tokyo, Japan), and hexaglycerol pentastearate (PS500), decaglycerol decastearate (DAS750) and decaglycerol heptabehenate (HB750) from Sakamoto Yakuhin Kogyo Co., Ltd. (Osaka). Other reagents used were analytical or special reagent grade commercial products.

Preparation of Suppositories: PS500, DAS750 and HB750 were chosen as PGEF in the present study. Suppositories were prepared according to the fusion method. At first, PGEF (PS500, DAS750 or HB750) was mixed well with Witepsol H15 at 60°C. Then, morphine hydrochloride, sieved by a 50-mesh screen, was mixed with melted bases (60 mg/1.6 g suppository) and dispersed homogeneously. The resulting mixture was then poured into plastic molds and allowed to cool at room temperature. Separately, Witepsol H15 suppository was prepared following the same procedure (30 mg of morphine/1.6 g suppository). Prepared suppositories were stored at 4°C until use.

Differential Scanning Calorimetry (DSC) Analysis: DSC thermograms were obtained using a DSC 2920 system.
In order to shear the melted suppository, a Teflon stirring bar 25 rpm and the dissolution medium was maintained at 37 °C. The release profile was described by the following equation:

\[ Q = k_Q t^{1/2} \]

where \( k_Q \) is the Higuchi’s rate constant. The value of \( k_Q \) was obtained as a slope of the linear portion of \( Q \) versus \( t^{1/2} \) plot for each mixed base by the least squares method.

Measurement of Viscosity A PGEF and Witepsol H15 were melted and mixed at the arbitrary ratio at 60 °C, and solidified at room temperature. The apparent viscosity of mixed bases (approximately 1 g) was measured with a cone and plate viscometer (Visconic® model E, Toki Sangyo Co., Ltd., Tokyo) at 37 °C.

In Vitro Release Study The release study was performed by PTLSW-type rotating dialysis cell (Pharma Test, Hamburg, Germany).\(^{10}\) The dissolution medium was 900 ml of phosphate buffer (pH 7.3), and 3 ml of the phosphate buffer was placed in the rotating cell. The cell was rotated at 25 rpm and the dissolution medium was maintained at 37 °C. In order to shear the melted suppository, a Teflon stirring bar \((f = 6 \times 30 \text{ mm})\) was placed in the rotating cell. The drug concentration in the dissolution phase was assayed spectrophotometrically at 240 nm.

The release profile was analyzed based on Higuchi’s model,\(^{11—13}\) where the cumulative amount \( (Q) \) of morphine released versus time was described by the following equation:

\[ Q = k_Q t^{1/2} \]

where \( k_Q \) is the Higuchi’s rate constant. The value of \( k_Q \) was obtained as a slope of the linear portion of \( Q \) versus \( t^{1/2} \) plot for each mixed base by the least squares method.

In Vivo Absorption Study Male Beagle dogs (8—12 kg), maintained at 20—26 °C and 40—70% humidity, were used for all experiments. Prior to the experiment, the fecal content in the rectal canal was reduced by fasting for 24 h. After rectal administration, 3 ml of blood samples were collected from antibrachium cephalic vein using heparinized syringes, and plasma was obtained by centrifugation. For the intravenous administration, 2.5 ml of morphine hydrochloride aqueous solution (1 mg/ml) was injected from antibrachium cephalic vein.

RESULTS AND DISCUSSION

DSC Thermograms and Viscosity of Mixed Bases of Witepsol H15 with PGEF Figure 1 shows the DSC thermogram of Witepsol H15-DAS750 as a typical example of the Witepsol H15-PGEF mixed bases, and endothermic peaks of the solid fat and the mixed base are presented in Table 1. Witepsol H15 showed an endothermic peak at 29.6 °C. When Witepsol H15 was mixed with DAS750, the main endothermic peak derived from Witepsol H15 was hardly shifted (29.8 °C), and the secondary peak (39.9 °C) was dissociated at the higher temperature side of the main endothermic peak. As shown in Table 1, it was confirmed that PS500 and HB750 showed the same properties as observed in the previous study.\(^{10}\) In the case of HB750, briefly, the main endothermic peak derived from Witepsol H15 was slightly shifted to higher temperatures (32.2 °C), and secondary peak (56.2 °C) was clearly separated from the main

Figure 1. DSC Thermograms of Witepsol H15-DAS750 Mixed Base
(a) Witepsol H15, (b) DAS750, (c) Witepsol H15-DAS750 mixed base containing 20% DAS750.
endothermic peak. Table 1 also shows that the temperature of secondary peak, which is derived from each PGEF, is different among PGEFs. As PGEFs are prepared by esterification of polymerized glycerol with various fatty acids, their physicochemical properties are dependent on the length of polymerized glycerol and/or esterified fatty acids. PS500 and DAS750 are stearic acid esters of polyglycerol, while HB750 is a behenic acid ester. Therefore, the much higher temperature of the secondary peak for Witepsol H15-HB750 mixed base comparing with other mixed bases, might be ascribed to behenic acid.

Figure 2 presents the fraction of secondary peak area in DSC thermograms of Witepsol H15-PGEF mixed bases as a function of PGEF content. The fraction increased with the increase of PGEF content in every mixed base examined and the statistically significant correlation ($p<0.05$) was observed with squared correlation coefficients of 0.990 (PS500), 0.980 (DAS750) and 0.992 (HB750), showing that the fraction un-melted at 37 °C increased by adding each PGEF into Witepsol H15.

The apparent viscosity was measured for Witepsol H15-PGEF mixed bases at 37 °C, and the relationships with the content of PGEFs are shown in Fig. 3. The linear and significant relationship was observed for every mixed base examined and the squared correlation coefficients for PS500, DAS750 and HB750 were 0.934, 0.959 and 0.997, respectively. These results indicate that the apparent viscosity of the mixed bases can be regulated quantitatively by adding PGEF into Witepsol H15, which could be ascribed to the amount of un-melted PGEF at 37 °C in each mixed base as shown in Fig. 2. Figure 3 also shows that HB750 makes it possible to prepare more viscous mixed base than PS500 and DAS750.

**Drug Release from Witepsol H15-PGEF Mixed Base Suppositories** Figure 4 shows the release profiles of morphine from suppositories made from Witepsol H15-PGEF mixed bases. The content of PS500, DAS750 or HB750 was 20%. Addition of each PGEF to Witepsol H15 delayed the release of morphine from a suppository. As the addition of HB750 resulted in the slowest release of morphine among PGEFs examined, the effect of HB750 amount contained in the mixed base on the release of morphine was investigated (Fig. 5). The release of morphine from the Witepsol H15-HB750 suppository decreased markedly with the increase in the weight fraction of HB750 in suppositories (Fig. 5).
The analysis of the results of the in vitro release study showed that the release rate of morphine from suppositories could be described by Fickian diffusion model (Higuchi's model), suggesting that the drug release from the suppository is regulated by the drug diffusion through the melted base. Then, the release rate constant of morphine ($k_{rH}$) was calculated, and the relation with the weight fraction of HB750 in the Witepsol H15-HB750 mixed bases (Fig. 6A) or the apparent viscosity of the mixed bases (Fig. 6B) was examined. The result shows that the release of morphine from the suppositories decreases with the increase in weight fraction (Fig. 6A) and the apparent viscosity of the mixed bases (Fig. 6B). This result indicates that the release of morphine from Witepsol H15-HB750 suppositories could also be regulated by changing the apparent viscosity as reported previously. As shown in Fig. 3, as the apparent viscosity of Witepsol H15-HB750 mixed base is significantly correlated with the content of HB750 in the mixed bases, the increased apparent viscosity by an un-melted solid fat would lead to the reduction of drug diffusion within the melted base.

**Absorption of Morphine from Witepsol H15-HB750 Mixed Base Suppositories** Figure 7 presents the plasma concentration of morphine after single rectal administration of Witepsol H15-HB750 suppositories in dogs, and pharmacokinetic parameters are summarized in Table 2. The dose of morphine hydrochloride was 60 mg/dog except for the suppository made from Witepsol H15 alone (30 mg/dog), because the rectal administration of 60 mg/dog by the Witepsol H15 suppository would lead to serious toxicity. But preliminary study showed that the linear pharmacokinetics of morphine after rectal administration of 60 mg/dog or less. After rectal administration of the Witepsol H15 suppository, plasma level of morphine increased rapidly with a $T_{\text{max}}$ of 0.4 h, followed by a fast decrease. On the other hand, plasma levels of morphine after rectal administration of Witepsol H15-HB750 suppositories gradually increased at early time periods and were kept at higher level for longer time periods, comparing with the Witepsol H-15 suppository.

Witepsol H15-HB750 suppositories showed the significant prolongation of MRT with the increase in HB750 contents, although there was no significant difference between 15% and 17% HB750. This prolongation of MRT was significantly correlated with the decrease in Higuchi's rate constant shown in Fig. 6 ($r=0.9772, p<0.01$). The significant delay of $T_{\text{max}}$ was observed, and the decrease in $C_{\text{max}}$ was also significantly correlated with the decrease in Higuchi's rate constant ($r=0.9203, p<0.05$). These results clearly indicate that the absorption of morphine was successfully sustained by mixed suppositories containing HB750. Relative bioavailability (BA) values for mixed base suppositories, compared with the Witepsol H15 suppository, are also summarized in Table 2. A mixed base suppository containing 10% HB750 decreased the relative BA, but the reason might be that the real $C_{\text{max}}$ results are expressed as the mean±S.E. of 4—5 experiments.

### Table 2. Pharmacokinetic Parameters for Morphine after Rectal Administration of Mixed Base Suppositories of Witepsol H15 with HB750 in Dogs

<table>
<thead>
<tr>
<th>Base</th>
<th>Dose (mg/dog)</th>
<th>$T_{\text{max}}$ (h)</th>
<th>$C_{\text{max}}$ (ng/ml)</th>
<th>MRT (h)</th>
<th>$AUC_{0-24h}$ (ng*h/ml)</th>
<th>Relative bioavailability (%)</th>
</tr>
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<tbody>
<tr>
<td>Witepsol H15 alone</td>
<td>30</td>
<td>0.4±0.1b,c-de</td>
<td>39.1±10.1b,c-de</td>
<td>6.6±1.3b,c-de</td>
<td>136.0±32.5b,c,d</td>
<td>100.0±23.9b,c,e</td>
</tr>
<tr>
<td>with 10% HB750</td>
<td>60</td>
<td>2.6±1.3c-de</td>
<td>25.4±5.0e</td>
<td>8.2±1.4c,d</td>
<td>192.1±27.1a,c,d</td>
<td>70.7±10.0c,d</td>
</tr>
<tr>
<td>with 15% HB750</td>
<td>60</td>
<td>6.0±2.8c,b,e</td>
<td>26.0±8.7e</td>
<td>10.3±1.3b,c,e</td>
<td>302.6±139.8b,c,e</td>
<td>111.1±51.4b,c,g</td>
</tr>
<tr>
<td>with 17% HB750</td>
<td>60</td>
<td>12.0±8.5c,b,c,e</td>
<td>22.4±10.0e</td>
<td>10.7±2.8b,e</td>
<td>303.9±116.1b,c,e</td>
<td>111.8±42.7b,c,e</td>
</tr>
<tr>
<td>with 20% HB750</td>
<td>60</td>
<td>7.2±3.6b,c,e</td>
<td>12.0±5.0b,c,d</td>
<td>12.2±1.2b,c,d</td>
<td>165.6±55.8b,c,d</td>
<td>60.9±20.5b,c,d</td>
</tr>
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</table>

Results are expressed as the mean±S.D. of 4—5 experiments. Relative bioavailability ($BA$) was calculated by the following equation: relative $BA=\frac{AUC}{DOE}$ of Witepsol H15 suppository $\times$ dose of morphine for Witepsol H15 suppository (30 mg)/Dose of morphine for each mixed base suppository (60 mg). $a,b,c,d,e,p<0.05$, compared with the Witepsol H15 alone, 10% HB70, 15% HB70, 17% HB70, 20% HB70, respectively.
could not be experimentally observed. In this case, \( C_{\text{max}} \) would be between 2 and 4 h after dosing. On the other hand, a mixed base suppository containing 15 or 17% HB750 did not decrease the relative bioavailability of morphine with prolonging MRT. However, the rectal administration of 20% HB750-containing suppository would lead to the decrease in bioavailability of morphine, although MRT was significantly prolonged. The decrease in the relative BA of morphine might be caused by the release rate that was too slow because of the high apparent viscosity of the mixed base. Furthermore, the effective surface area might be smaller for the mixed base suppository containing 20% HB750 than other suppositories, because the base with higher viscosity and higher melting point could maintain the shape of suppository longer and retain in the original position of administration as reported by Yahagi et al.\(^{15}\)

These results clearly indicate that Witepsol H15-HB750 mixed base suppositories containing 15—17% HB750 could be the best preparation to maintain the plasma concentration of morphine by controlling the release of morphine from suppositories. Moreover, the in vivo absorption behavior of morphine could be reflected by the in vitro release characteristics of suppositories (Fig. 5) to some extent.

CONCLUSIONS

In the present study, controlled-release morphine suppositories were prepared by utilizing mixed bases of Witepsol H-15 with PGEF such as PS500, DAS750 and HB750. HB750 showed the slowest release rate of morphine in the in vitro release study, and the release rate decreased with the increase in the apparent viscosity of the mixed bases, which could be ascribed to the decrease in the diffusion of morphine through the mixed bases. The in vivo absorption of morphine from suppositories was successfully prolonged with HB750, but Witepsol H-15-mixed base suppositories containing 15—17% HB750 could be promising ones because they sustained the plasma concentration of morphine without decreasing BA of morphine.

REFERENCES