Preparation of Solid Dispersion for Ethanazamide–Carbopol and Theophylline–Carbopol Systems Using a Twin Screw Extruder

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Received January 21, 2002; accepted March 8, 2002

In the present study, we prepared solid dispersions of water-insoluble and soluble drugs (ethanazamide (ETZ) and theophylline (THEO)) by the twin screw extruder method, which made it possible to control both kneading and heating at the same time under the fusion point of each drug, using three types of the controlled-release high-molecular-weight substance Carbopol (CAR) as the carrier. The solid dispersions obtained were evaluated and compared with those prepared by the organic solvent method. These products showed significantly increased solubility of ETZ, but the solubility of THEO was reduced indicating that CAR slows the release of THEO. It is important not only to simply knead under high pressure but to select the optimal operation temperature to bring these drugs into a semi-fusion state. Solid dispersions obtained by this method showed X-ray diffraction and differential scanning calorimetry (DSC) patterns similar to those obtained by the organic solvent method indicating that the former can be used as a simple and effective method for preparation of solid dispersions.

Key words solid dispersion; Carbopol; twin screw extruder; ethanazamide; theophylline

The use of poorly soluble drugs has a number of drawbacks such as increasing the dosage, administration frequency and the resultant occurrence of side effects. Furthermore, as solubility is a rate-limiting factor for absorption in the human body, it is important to increase the solubility of a drug to improve its bioavailability. Various pharmaceutical approaches have been adopted to improve solubility using micelles and synthesis of soluble derivatives and salts.

The solid dispersion method, by which a drug is dispersed in a carrier to make it amorphous, is one of the pharmaceutical approaches most commonly employed to increase bioavailability of poorly soluble drugs. Various methods for preparation of solid dispersions including coprecipitation, lyophilization, spray drying, solvent evaporation, fusion and powder mixing methods have been reported. The solvent evaporation method and fusion method have a number of drawbacks, including residual organic solvent and decomposition of the drug. A pressurization/kneading/extruding method using a twin screw extruder is one of the methods proposed for this purpose. This extruder, originally designed as an extraction/casting device for polymer alloys in the plastic industry, is now used to process cereals and “functionalize” food materials such as tissue products from the plastic industry. It is now used to process cereals and “functionalize” food materials such as tissue products from the plastic industry. This device has already been used successfully to prepare solid dispersions of nifedipine and hydroxypropylmethylcellulosephthalate, and floating dosage forms of nicardipine hydrochloride and hydroxypropylmethylcellulose acetate succinate.

In the present study, the properties of a solid dispersion of ethanazamide (ETZ), which is poorly soluble in water, and that of theophylline (THEO), which is readily soluble in water, were prepared using a twin screw extruder and compared with their physical mixtures and a solid dispersion obtained using organic solvents. The solid dispersions were prepared using 3 grades of Carbopol® (CAR), a cross-linked polymer of acrylic acid used as a drug release controlling carrier, biological mucous membrane adherent, viscosity donor and emulsifier with the fishnet gel structure (CAR971P) and fuzzball gel structure (CAR974P and 934P) shown in Table 1. The effects of temperature on the degree of crystallinity of drugs in solid dispersions were also evaluated.

Experimental

Materials The physical properties of CAR® (CBC Co., Ltd., BF Goodrich), ETZ (Yoshitomi Fine Chemicals Co., Ltd.) and THEO (Wako Pure Chemical Industries, Ltd.) are listed in Tables 1 and 2. All other reagents used were analytical grade and were obtained.

Twin Screw Extruder The twin screw extruder (KEX-25, Kurimoto, Ltd.) used in the present study consisted of a hopper, barrels, a die, a kneading screw, and heaters. The thread interval of the feed screw decreases from the hopper side to the die side. A physical mixture introduced into the hopper is carried forward by the feed screw, kneaded under high pressure by the kneading screw, and extruded from the die. Temperature inside the barrels can be accurately controlled from 30 to 300 °C with 4 independent heaters, and water can be introduced from the liquid injection port if necessary.

Preparation of Materials Twin Screw Extruder Method ETZ or THEO was mixed with CAR at the ratios shown in Table 3 at 42 rpm for 20 min using a V-type mixer (VM-5, Fuji Powdal Co., Ltd.). Mixtures were then treated with the twin screw extruder at a screw rotation rate of 100 rpm, powder supply rate of 10 g/min, water supply rate of 1 ml/min, and a barrel temperature of 85 °C. Treated mixtures were dried for 10 h using a dryer (inside temperature: 50 °C), pulverized using a QUADRO COMIL (QC-197S, Powlex Co., Ltd.) equipped with a grater-type screen (2.39 mm), dried overnight in a dryer, pulverized again using an automatic mortar (Model ANM200 E, Nitto Kagaku Co., Ltd.), dried overnight again in a dryer to remove water completely, and sieved through a 150–250 μm sieve to obtain a sample for granule dissolution. Samples 55 μm or finer were used for powder X-ray diffraction and differential scanning calorimetry (DSC) analysis.

Mixtures of designated ratios were put into 50 ml sample bottles and mixed for 5 min using a test tube shaker (Vortex-Genie-2, SM Kiki Co., Ltd.) to obtain physical mixtures.

Organic Solvent Method ETZ (50 g) was dissolved in 200–500 ml of a 1:1 (w/w) mixture of ethanol and dichloromethane, and CAR (50, 150, 250 g) of 3 different grades was suspended in the resultant solutions. For establishment of equilibrium, the suspensions thus obtained were allowed to remain at room temperature and pressure, and the solvent was spontaneously evaporated. The solvent was then evaporated under reduced pressure for 24 h at 60 °C using a thermostatic reduced pressure dryer (Isihai Vacuum Sample Oven, Isihai Labo Works Co., Ltd.). The dry materials obtained after evaporation were pulverized, dried and sieved as described above and their physical properties were evaluated.

Evaluation of Solid Dispersions Powder X-Ray Diffraction Powder X-ray diffraction was conducted using a linear X-ray diffraction system (RAD-IIVC, Rigaku Denki Co., Ltd.) in which CuKα-rays (40 kV, 20 mA) were used as X-rays. The degree of diffraction was measured at 4°/min every 0.02° between 4° and 40° (2θ).
DSC  Fusion temperature of each sample (10 mg) was measured at a
temperature gradient of 10 °C/min in atmospheric air. A differential scan-
ning calorimeter (DSC-3100, Mac Science Co., Ltd.) was employed.

Fourier Transformation IR Spectroscopy (FT-IR)  Drug–carrier inter-
actions in samples were determined based on IR spectra using a FT -IR spec-
troscope (FT -200, Horiba Ltd.) by the KBr method.

Dissolution Test  The amount of drug released from samples was deter-
mined using a dissolution tester (NTR-3000, Toyama Sangyo Co., Ltd.) ac-
cording to dissolution test method 2 (paddle method) described in the Japan-
ese Pharmacopoeia (JP XIII). The paddle was rotated at 100 rpm using
50 mg of treated powder and, as test fluid, 900 ml of purified water adjusted
at 37°±0.2 °C and the 1st and 2nd fluids used in the disintegration test ac-
cording to JP XIII. Test fluids (about 1 ml) were sampled at regular time in-
tervals through a glass filter and immediately filtered through a 0.45
mm membrane filter (DISMIC-25CP , Toyo Filter Paper Co., Ltd.). The amount of
drug in the filtrate was assayed by HPLC (LC-9A, Shimadzu Corp.) under
the following conditions: column, Shim-pack VP-ODS (4.6×150 mm,
5 µm); flow rate, 1 ml/min; wavelength, 293 nm for ETZ and 340 nm for THEO. A
Chromatopack C-R4A (Shimadzu Corp.) was employed for data processing.

The Relationship between Operation Temperature and the Degree of
Crystallinity  A 1:1 mixture of ETZ and CAR-934P was treated by the
twin screw extruder method at barrel temperatures ranging from 40 to
115 °C to evaluate the effects of barrel temperature on the properties of ex-
truder-treated samples. Integrated intensity of ETZ (I_{ETZ}) and solid disper-
sions of each temperature (I_{SD}) were determined from the
diffraction intensity. The integrated intensity of CAR-934P treated with the
twin screw extruder (I_{CAR}) was subtracted from (I_{SD}), and the ratio of the
value obtained to the integrated intensity of ETZ was calculated as the de-
gree of crystallinity at each temperature using Eq. 1.

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\text{degree of crystallinity} = \frac{I_{SD} - I_{CAR}}{I_{ETZ}} \times 100
\]  

Results and Discussion  Evaluation of Solid Dispersion. Changes in Crystal-
linity  Powder X-ray diffraction patterns obtained with ETZ alone, CAR alone, their 1:1 physical mixture, and ETZ-
CAR-P-NF treated by the twin screw extruder method and the organic solvent method are shown in Fig. 1, and that of
THEO alone, a 1:1 physical mixture of THEO and CAR, and mixture of THEO and CAR treated by the twin screw ex-
truder method are shown in Fig. 2.

ETZ alone showed marked crystallinity with many diffraction
peaks. The high-molecular weight carrier CAR showed a
broad diffraction pattern. The physical mixture showed ETZ-de-
rived diffraction peaks. Solid dispersions prepared by the
twin screw extruder method showed small peak intensity at a
ratio of 1:1 (THEO:CAR). Refraction peaks disappearing completely, showing a
completely hollow pattern that of the solid dispersion.

In evaluation of solid dispersions prepared by the twin screw extruder method using CAR of different grades, CAR-971P was found to give less diffraction peaks than CAR-974P and CAR-934P at a mixing ratio of 1 : 1 (ETZ : CAR) indicating that CAR-971P have more ability to form the solid dispersion than others.

**DSC Curves** DSC curves of ETZ, CAR, their physical mixture, and solid dispersions obtained by the twin screw extruder method and the organic solvent method are shown in Fig. 3 and those of THEO, physical mixture of THEO and CAR, and solid dispersions obtained by the twin screw extruder method are shown in Fig. 4.

ETZ showed a sharp endothermic peak at 139 °C due to fusion, while CAR showed a broad endothermic peak. The physical mixture showed an ETZ-derived endothermic peak.

Solid dispersions prepared by both the twin screw extruder method and the organic solvent method showed broad curves in the ETZ-derived endothermic point. No differences between CAR of different grades were seen.

The observation that similar patterns were obtained with samples obtained by the twin screw extruder method and the organic solvent evaporation method confirmed that ETZ changed to an amorphous state in mixtures of ETZ and CAR and solid dispersions were formed when treated by the twin screw extruder method.

When the twin screw extruder method was employed, the solid dispersion prepared using CAR-971P showed least crystallinity, probably because CAR of this grade is less cross-linked when swollen, and for this reason the drug is more readily dispersed into macromolecules and undergoes reactions such as hydrogen bonding when treated by the twin screw extruder method. In contrast, CAR-934P and CAR-974P show a dense cross-linkage pattern when they come into contact with water, preventing the drug from penetrating deeply into the CAR structure and inhibiting reactions such as hydrogen bonding.

THEO showed a sharp endothermic peak at 275 °C due to fusion. The physical mixture also showed a THEO-derived endothermic peak. The 1 : 1 (THEO : CAR) solid dispersion prepared by the twin screw extruder method showed a slight endothermic peak shifted to a lower temperature, but the 1 : 5 (THEO : CAR) solid dispersion showed a broad curve with no endothermic peak.

These findings indicated that the twin screw extruder method made THEO amorphous in THEO-CAR samples, as well as ETZ in ETZ-CAR samples, and formed solid dispersions.

**Confirmation of Interactions by IR Spectrum** FT-IR spectra obtained with samples prepared by the twin screw extruder method and the organic solvent method using CAR-971P are shown in Fig. 5 and those obtained with THEO alone and samples prepared by the twin screw extruder method are shown in Fig. 6.

As ETZ has a primary alkylamide group, binding of its crystals is thought to be fortified by dimer chain association. ETZ and the physical mixture of ETZ and CAR-971P showed N–H stretching vibrations at 3371 and 3178 cm$^{-1}$ due to the primary amide of ETZ, while samples prepared by the twin screw extruder method and the organic solvent method showed a new peak at 3455 cm$^{-1}$, but no more peaks
due to primary amide at 3369 or 3176 cm$^{-1}$. This was though to be because of interactions between the primary amide of ETZ and CAR during formation of the solid dispersion, which resulted in various associations such as hydrogen bonding between the –NH$_2$-group of ETZ and –COOH group of CAR.

THEO alone and the physical mixture showed an N–H deformation vibration at 1664 cm$^{-1}$ due to the secondary amine of THEO, while samples prepared by the twin screw extruder method showed a shift of the N–H deformation vibration to 1625 cm$^{-1}$.

This finding suggested that the formation of solid dispersions of THEO and CAR by the twin screw extruder method was due to interactions of the two components and that hydrogen bonding between the N–H of THEO and COOH of CAR is involved in these interactions.

**Dissolution Test** Dissolution profiles of ETZ from solid dispersions in purified water, the 1st fluid and the 2nd fluid are shown in Fig. 7 and those of THEO are shown in Fig. 8.

As a solid dispersion was fully formed when 1 : 3 (ETZ : CAR) and 1 : 5 (ETZ : CAR) ratios were used for the mixture, samples prepared by the twin screw extruder
method showed more rapid dissolution than ETZ alone or the 1:1 (ETZ:CAR) mixture regardless of the CAR grade used. Slow dissolution, especially that observed in the 1:1 (ETZ:CAR) mixture in purified water and in the 2nd fluid compared to that seen in the 1st fluid, was thought to be because granules are more likely to adhere to each other immediately after the addition of samples to purified water and the 2nd fluid. Rapid dissolution as observed in the 1:3 and 1:5 (ETZ:CAR) mixture was thought to be because ETZ was dispersed completely and solid dispersions were formed. In the 1st fluid, granules were less likely to adhere to each other because CAR shows less swelling and the solubility of ETZ alone was higher at a low pH, and this is thought to result in rapid dissolution.

To evaluate differences in the dissolution rate according to the grade of CAR, the times needed for 70% drug release were defined as the T70, and the relationship between T70 of ETZ and THEO, and CAR contents was evaluated. In neutral and alkaline solutions such as purified water and the 2nd fluid, only samples with a 50% CAR concentration showed marked adhesion of granules during dissolution. In purified water and the 2nd fluid, however, marked differences were observed when the CAR content was 50%. The total volume of expanded channel in the purified water, 1st and 2nd fluid differed in the structure differences of CAR grades which affected dissolution rate.

Relationship between Operation Temperature and the Degree of Crystallinity

The degree of crystallinity at each temperature is shown in Fig. 11. The degree of crystallinity of ETZ decreased as barrel temperature increased, suggesting that when preparing solid dispersions of ETZ and CAR using the twin screw extruder method (ETZ:CAR=1:1) the release of drug was delayed due to the interactions between the N–H of THEO and COOH of CAR released the drug more slowly and in some cases decreased because re-entrapped by CAR.

Fig. 7. Dissolution Profiles of ETZ from Solid Dispersions in 900 ml at 37 °C (CAR-971P)

A) water, B) 1st fluid, C) 2nd fluid. ○, ETZ; ●, twin screw extruder method (ETZ:CAR=1:1); ×, twin screw extruder method (ETZ:CAR=1:3); □, twin screw extruder method (ETZ:CAR=1:5). Each point represents the mean and S.D. (n=3).

Fig. 8. Dissolution Profiles of THEO from Solid Dispersions in 900 ml at 37 °C (CAR-971P)

A) water, B) 1st fluid, C) 2nd fluid. ○, THEO; ×, twin screw extruder method (THEO:CAR=1:1); ●, twin screw extruder method (THEO:CAR=1:5). Each point represents the mean and S.D. (n=3).
Conclusions

(1) Powder X-ray diffraction and DSC evaluation showed that when ETZ-CAR and THEO-CAR mixtures are treated with a twin screw extruder, ETZ and THEO are made amorphous and solid dispersions can be formed as by the organic solvent method.

(2) When preparing ETZ-CAR solid dispersions by the twin screw extruder method, it is important not only to simply knead under high pressure but to select optimal operation temperature to bring the drug into a semi-fusion state.

(3) Solid dispersions obtained using the twin screw extruder showed significantly increased ETZ release, but the release rate of THEO decreased when the same treatment was conducted indicating that CAR slows the release rate of THEO.

(4) Solid dispersions obtained by the twin screw extruder method and those obtained by the organic solvent method showed similar X-ray diffraction and DSC patterns, indicating that the former can be used as a simple and effective method for preparation of solid dispersions.

Acknowledgements The author is grateful to Kenichi Harada, Professor, and Kiyonaga Fujii, Research Associate, of the Department of Instrumental Analysis, for their help in the determination of FT-IR spectra.

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