Use of supercritical CO2-aided and conventional melt extrusion for enhancing the dissolution rate of an active pharmaceutical ingredient

Zsombor Kristof Nagy, Martial Sauceau, Katalin Nyul, Elisabeth Rodier, Balzs Vajna, György Marosi, Jacques Fages

To cite this version:


HAL Id: hal-01618290
https://hal.archives-ouvertes.fr/hal-01618290
Submitted on 24 Nov 2017

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.
Use of supercritical CO2-aided and conventional melt extrusion for enhancing the dissolution rate of an active pharmaceutical ingredient

Zsombor Kristóf Nagya*, Martial Sauceaub, Katalin Nyúla, Elisabeth Rodierb, Balázs Vajnaa, György Marosia and Jacques Fagesb

Dispersing at the molecular level a drug in a polymer matrix is a major challenge to be addressed by the pharmaceutical industry to enhance its bioavailability or to control its release. Melt extrusion and supercritical CO2-aided melt extrusion of solid pharmaceutical formulations were performed to enhance the dissolution rate of carvedilol, taken as a model of poorly soluble drug. The presence of the drug improved the processability of the polyacrylate matrix (Eudragit E) through its plasticizing effect. The supercritical method was found gentle compared with melt extrusion owing to the shorter residence time and lower processing temperature and melt viscosity. No traces of decomposition of the drug could be detected after the supercritical extrusion process based on capillary electrophoresis results. This extrusion process resulted in effective homogenization of the components and amorphization of the drug according to Raman mapping, Fourier transform infrared spectrometry, X-ray diffraction, and polarized light microscopy. The kinetics of dissolution can be dramatically improved.

Keywords: extrusion; supercritical CO2 extrusion; solid dispersion; solid solution; amorphization; enhanced drug dissolution rate

INTRODUCTION

The term “hot-melt extrusion” (HME) is used in pharmaceutical technology for describing the process that converts a raw material into a product of uniform shape and density by forcing it through a die under controlled temperature, pressure, and shear rate.[1] Pharmaceutical extrusion represents an efficient manufacturing technology to disperse drugs in a melt up to a true molecular solution of a pharmaceutical ingredient into a polymeric matrix.[2] Most of the applications describe the preparation of solid dispersions by HME either to increase the aqueous solubility and oral bioavailability of the active substance or to control its release. The mostly reported advantages of HME are as follows: (i) avoidance of organic solvent; (ii) reduction of processing steps (one-pot method); (iii) elimination of the good compressibility requirement for the active ingredients and the excipients; (iv) high level of drug content; (v) even dispersion of the drug throughout the matrix; and (vi) improved bioavailability through drug solubilization or molecular-level dispersion in water-soluble matrix.[3]

One of the major drawbacks of HME is that it can be applied only for producing thermally stable pharmaceuticals. For a number of drugs, proteins, and excipients, HME cannot be used unless a plasticizer is added in order to reduce the viscosity of the mixture and to lower the processing temperature applicable in the extruder. However, these additives remain in the product, altering its properties and performance.

In this context, supercritical carbon dioxide (scCO2) represents an interesting alternative. The injection of scCO2 in the extrusion process modifies the rheological properties of the polymer, and also, scCO2 acts as a blowing agent at the die exit.[4] In the barrel of the extruder, the reduction of viscosity decreases the mechanical constraints and the operating temperature. At the die exit, the pressure drop induces a thermodynamic instability in the polymer matrix, generating a large number of bubbles, which can grow until the foam is rigidified when temperature drops below the glass transition temperature, \( T_g \). Thus, combining extrusion with scCO2 allows using relatively fragile or thermally sensitive molecules, like pharmaceutical molecules, without any residue in the final material.

For the extrusion of pharmaceuticals, copolymers derived from acrylic acid and methacrylates, such as Eudragit types, are well-known matrices. Because of their multifunctional nature, unique properties, and good biocompatibility, these polymers are considered to be of common grades for the coating of pharmaceuticals.[5] In particular, Eudragit E, because of its tertiary...
amine groups, forms a coating soluble in gastric fluid in acidic pH.[6] Eudragit E has already been successfully implemented for HME. Six et al.[7] described the HME of the poorly water-soluble itraconazole with Eudragit E100 polymeric carrier. Melt extrusion of this drug with Eudragit E polymer resulted in significantly enhanced dissolution rate. More recently, Verreck et al. showed that pressurized carbon dioxide, acting as a removable plasticizer and foaming agent, reduces the melt extrusion temperature of several pharmaceutically acceptable polymers, including Eudragit E100.[8] Lately, Nikitine et al. studied the correlation between operating conditions and foam structures of Eudragit E100.[9] This work was also completed by the study of the influence of scCO₂ on the polymer flow of Eudragit E100 in the extruder.[10–12]

Lipid soluble chemicals, such as carvedilol (CAR), are almost insoluble in water. Therefore, they are poorly absorbed through the gastrointestinal tract. CAR is a nonselective β-adrenergic blocking agent with an α-blocking activity.[13] According to the Biopharmaceutics Classification System, it belongs to the class II drug having low solubility and good permeability. However, if CAR is well absorbed after oral administration, it is subject to first-pass metabolism in the liver resulting in only about 25% absolute bioavailability.[14] Eudragit E100 has already been used to improve the bioavailability of CAR. Recently, E100 nanoparticles containing CAR were prepared by nanoprecipitation.[15] Finally, high encapsulation efficiency was obtained, and the release of CAR from the nanoparticles was achieved within 5 min.

In this work, melt extrusion and scCO₂-coupled extrusion was used to generate solid dosage form based on Eudragit E100 with CAR drug according to the method elaborated for producing foams of this polymer.[16] The aim was to obtain similar drug release rate of CAR as it was achieved by nanoprecipitation and to investigate the potential advantages of applying scCO₂ during the extrusion process. For achieving rapid dissolution, the feasibility of an amorphous solid dispersion or solution of CAR and Eudragit E100 was investigated in order to avoid the necessity of overcoming the crystal lattice energy of CAR. In the case of solid solution, the rate-determining step of CAR release is the dissolution of Eudragit E, which is quite rapid in acidic media such as the gastric fluid. The properties of the extrudates were investigated by several methods to study the morphology, the crystallinity, the degradation, the mixing, and the dissolution profile.

**MATERIALS AND METHODS**

**Materials**

Carvedilol (Fig. 1) from Sigma-Aldrich (Budapest, Hungary) with purity ≥ 98% was used in this work. The melting point of CAR is 117°C.[16] CO₂ was purchased from Air Liquide (Paris, France) and Eudragit E (E100, Fig. 2) from IMCD (Saint-Denis La Plaine, France). Eudragit is an amorphous polymer with an average molecular weight of about 150 000 g mol⁻¹. Its glass transition temperature, measured by differential scanning calorimetry (ATG DSC 111, SETARAM, Caluire, France), is in the range 45–52°C. A rheological study at atmospheric pressure was performed,[17] and the polymer exhibited a pseudoplastic behavior with a viscosity similar to those of molten polymers (around 10⁵ Pa s). These previous characterizations helped in choosing the operating conditions to process Eudragit E100 by extrusion.

**Extrusion coupled with supercritical CO₂**

Figure 3 shows the experimental set up, which has previously been detailed elsewhere.[9,10] The single-screw extruder has a 30 mm screw diameter and a length to diameter (L/D) ratio of 35 (Rheoscam, SCAMEX, Crosne, France). A large L/D ratio generally allows increasing not only the capacity of mixing and melting but also energy consumption. The first conical part of the screw has an L/D ratio of 20 and allows the transport of solid polymers and then their melting and plasticizing. Then, the screw has a cylindrical geometry from the first gastight ring to the die with an L/D ratio of 15. Two restriction rings have been fitted out in order to prevent scCO₂ from backflowing. The diameter of the die is 1 mm and its length is 11.5 mm.

The temperature inside the barrel is measured at five locations: T_a and T_b before the CO₂ injection, T_e and T_o after the injection, and T_c in the die. In this study, the temperatures of the different zones were fixed as follows: T_a = 125°C, T_b = T_e = T_o = 135°C and T_c = 130°C and the measured temperatures were stabilized at the predetermined values (±2°C) during the experiments. They were kept in all cases higher than the melting point of CAR (117°C[16]), allowing it to be melted and thus increasing the possibility for obtaining solid solution.

The temperature and the pressure of the polymer inside the extruder were measured by means of three pressure gauges (P_1, P_2, and P_3) and two temperature sensors (T_1 and T_2). Errors associated to pressure and temperature sensors were about 0.2 MPa and 3.3°C, respectively.

CO₂ was pumped from a cylinder by a syringe pump (260D, ISCO, Lincoln, NE, USA) regulated at 7°C. It was then introduced at a constant volumetric flow rate and at a pressure slightly higher than P_1. The CO₂ injector was positioned at an L/D ratio of 20 from the feed hopper. The pressure and the volumetric CO₂ flow rate were measured within the syringe pump.

At the beginning of the extrusion process, pure E100 was fed in the extruder. Before the injection of scCO₂, the torque was controlled by the rotating speed of the screw (higher rotating
speed resulted in higher torque) and was kept at high values close to the established alarm limit (120 Nm) to ensure high production rate and short residence time. Once steady-state conditions had been reached with E100, drug polymer mixture was introduced through the hopper. supercritical CO2 was introduced into the barrel once the drug polymer mixture had reached the die. During the scCO2 injection, the torque decreased, and the rotation speed could therefore be increased to reach higher production rates and shorter residence time. When steady-state conditions had been reached again, extrudates were collected and air cooled at ambient temperature. The samples were obtained under the operating conditions given in Table 1. The sampling time and the mass of the collected samples were measured in order to determine the mass flow rate. Extrudates were then ground by an IKA MF10 microfine (MF) grinder equipped with a MF 10.2 impact grinding head (Staufen, Germany). The applied rotation speed was 3000 rpm and sieve was MF 2.0 mm, thus the obtained particle size was less than 2 mm. The grinding was carried out using the same method in all cases. Based on optical and scanning electron microscopy analyses, the sizes of majority of the particles were between 100 and 1500 µm.

Compositions and operating conditions for the different samples are presented in Table 1. The CO2 flow rate was fixed at 0.25 cm3 min⁻¹, and two drug mass contents were tested: 5% and 20%.

**Scanning electron microscopy**

Morphology of the samples was investigated by a JEOL 6380LV type scanning electron microscope (Tokyo, Japan). Each specimen was fixed by conductive double-sided carbon adhesive tape and gold-coated using a JEOL 1200 apparatus (Tokyo, Japan) in order to avoid electrostatic charging.

**Fourier transform infrared spectrometry**

KBr tablet sample preparation method was used for the Fourier transform infrared (FTIR) spectrometry measurement of untreated materials and ground extrudates. The obtained tablets were

<table>
<thead>
<tr>
<th>Exp</th>
<th>CAR (mass%)</th>
<th>CO2 (cm³ min⁻¹)</th>
<th>Pressure (P₁) in the barrel at CO2 injection (bar)</th>
<th>Screw rotation speed (rpm)</th>
<th>Flow rate (g min⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1CAR0</td>
<td>0</td>
<td>0</td>
<td>165</td>
<td>5</td>
<td>5.5</td>
</tr>
<tr>
<td>2CAR5</td>
<td>5</td>
<td>0</td>
<td>135</td>
<td>5</td>
<td>5.5</td>
</tr>
<tr>
<td>3CAR5SC</td>
<td>5</td>
<td>0.25</td>
<td>78</td>
<td>5</td>
<td>5.5</td>
</tr>
<tr>
<td>4CAR5SC</td>
<td>5</td>
<td>0.25</td>
<td>85</td>
<td>7</td>
<td>7.7</td>
</tr>
<tr>
<td>5CAR5SC</td>
<td>5</td>
<td>0.25</td>
<td>112</td>
<td>10</td>
<td>11</td>
</tr>
<tr>
<td>6CAR20</td>
<td>20</td>
<td>0</td>
<td>115</td>
<td>5</td>
<td>5.5</td>
</tr>
<tr>
<td>7CAR20SC</td>
<td>20</td>
<td>0.25</td>
<td>57</td>
<td>5</td>
<td>5.5</td>
</tr>
<tr>
<td>8CAR20SC</td>
<td>20</td>
<td>0.25</td>
<td>74</td>
<td>7</td>
<td>7.7</td>
</tr>
<tr>
<td>9CAR20SC</td>
<td>20</td>
<td>0.25</td>
<td>90</td>
<td>10</td>
<td>11</td>
</tr>
</tbody>
</table>

CAR, carvedilol.
analyzed by Bruker Tensor 37-type spectrometer equipped with deuterated triglycine sulfate detector (Ettingen, Germany) in the range of 4000 – 400 cm\(^{-1}\) with a resolution of 4 cm\(^{-1}\).

**Polarized light microscopy**

The crystalline and amorphous phases of the samples were distinguished by Amplital, Carl Zeiss Jena polarized light microscope (Jena, Germany).

**X-ray diffraction**

Powder X-ray diffraction patterns were recorded by a PANalytical X'pert Pro MDP X-ray diffractometer (Almelo, The Netherlands) using Cu Kα radiation (1.542 Å) and Ni filter. The applied voltage was 40 kV whereas the current was 30 mA. The untreated materials, physical mixture (mixed in a mortar by a pestle) and ground extrudates were analyzed for angles 2θ between 2° and 42°.

**Capillary electrophoresis**

The chemical stability of the extruded samples was characterized by capillary electrophoresis. The experiments were performed using a HP10CE fully automated instrument, Agilent Technologies (Santa Clara, CA, USA), equipped with diode-array detector and ChemStation software for data handling. The fused silica capillary (inner diameter 50 µm, effective length 30 cm) was thermostated at 25°C and was preconditioned prior all runs by flushing with Milli-Q water (0.5 min), 0.1 M NaOH (1 min) Milli-Q water (0.5 min) and the buffer (2.5 min, 60 mM NaH₂PO₄, pH = 2.5 adjusted with H₃PO₄). The samples were dissolved with 0.1 M HCl and were injected hydrodynamically (5 kPa, 4 s). The applied electrostatic potential was 20 kV during the measurement. Three parallel measurements of each sample were carried out.

**Raman microspectrometry**

Raman-mapping spectra were collected using a Horiba Jobin Yvon LabRAM system (Longjumeau, France) coupled with a frequency doubled Nd:YAG laser source (532 nm) and an Olympus BX-40 optical microscope. Objectives of ×10 and ×100 magnifications were used for optical imaging and spectrum acquisition.

The Raman spectra of the pure materials were collected with a ×100 objective without any sample preparation. The surface of the cut cross-section of each sample was mapped by Raman microscope in order to investigate the mixing efficiency. In the case of foamed samples, the tablet was compressed (Camilla OL95, Manfredi, Torino, Italy) to achieve a flat surface for Raman mapping. The reference sample was also a compressed tablet of the physical mixture (mixed in a mortar by a pestle) of CAR and E100 (mass ratio 1:4). The measured area was approximately 1 mm × 1 mm in all cases. The component concentrations were estimated with classic least squares method. Visualized score maps were created with LabSpec 5.41 (Horiba Jobin Yvon).

**Determination of drug content**

Samples equivalent to 10 mg of CAR were placed into a volumetric flask (1 dm\(^3\)) and were diluted with 0.1 M HCl. After complete dissolution and homogenization, the concentration of CAR was determined by UV spectrophotometry at 242 nm (Hewlett-Packard HP 8452A, Palo Alto, CA, USA). Ten parallel tests were analyzed statistically.

**In vitro dissolution measurement**

The dissolution studies were performed using an Erweka DT6 dissolution tester (USP II apparatus, Heusenstamm, Germany). Samples equivalent to 12.5 mg of CAR were added as a powder directly in the dissolution vessel containing 900 ml of 0.1 M HCl maintained at 37 ± 0.5°C and stirred at 50 rpm. Samples (5 ml) were collected periodically and the concentration of CAR was determined by UV spectrophotometry at 242 nm (Hewlett-Packard HP 8452A) using a diode-array detector at 242 nm. Six parallel tests were performed and analyzed statistically.

**RESULTS AND DISCUSSION**

**Comparison of the melt and supercritical CO₂-aided extrusion techniques**

To improve the dissolution rate of CAR by forming solid dispersion or solution of the drug in the E100 matrix, both melt extrusion and scCO₂-aided melt extrusion processes were performed (Table 1). The viscosity of the material in the barrel was correlated with the torque. Without the injection of scCO₂, the maximum applicable rotation speed was 5 rpm. In contrast, while injecting scCO₂, the torque significantly decreased, and the rotation speed could be increased stepwise up to 10 rpm. The torque at 10 rpm was lower than the value corresponding to 5 rpm without scCO₂ (Fig. 4). This is demonstrated by experiments 2 and 5 at 5% of CAR and 6 and 9 at 20% of CAR. A gain in the maximal throughput, from 5.5 to 11 g/min (cf. Table 1), was also observed as well as a shortened residence time. These effects are consistent with a lower viscosity, which can be attributed to the plasticization effect of carbon dioxide. The application of scCO₂ in the field of pharmaceutical extrusion has two noteworthy advantages as for the CAR stability point of
view. Because of the dissolution of CO\textsubscript{2} in the material processed, operating temperature can be decreased\textsuperscript{[8]}, and the mass flow rate can be increased because of the decrease of the viscosity. If higher mass flow is achieved, the residence time of the material can be significantly shortened; furthermore, the productivity is increased. Higher throughput and lower temperature result in a lower risk of decomposition and a lower amount of impurities (decomposition products), which can be kept under the permitted limit.

In addition to the effect of scCO\textsubscript{2}, the introduction of the active ingredient provided a plasticizing effect too, as evidenced by the decreasing torque, from 104 N m to 96 N m with 5% drug and 83 N m with 20% drug (cf. experiments without scCO\textsubscript{2} in Fig. 4). CAR, being a rather small molecule, proved to play the role of a plasticizer. E100 CAR mixtures exhibit lower torque than pure E100, which suggests a decrease of the viscosity.

The photographs of E100 CAR extrudates are shown in Fig. 5. The extrudates made by classic extrusion process (A) were glassy and transparent. In contrast, the extrudates made by scCO\textsubscript{2}-aided extrusion (B) were opaque because of the reflections caused by the internal surfaces of small bubbles formed during the extrusion.

**Scanning electron microscopy**

The scanning electron microscopy images of extrudates are shown in Fig. 6. The specific surface area is clearly larger in the case of the foamed samples (with CO\textsubscript{2}) providing extended

![Figure 5. E100 CAR extrudates; (A) 6CAR20 and (B) 9CAR20SC.](image1)

![Figure 6. Scanning electron microscopy images of the extrudates; (A) milled 1CAR0, (B) milled 5CAR5SC, and (C) 9CAR20SC.](image2)
availability to the dissolving medium, which may increase the dissolution rate. This effect may be particularly effective for the sample with 20% of CAR, for which an important opened porosity is present in coarser particles.

**Fourier transform infrared spectrometry**

The extruded products were analyzed by FTIR spectrometry in order to check their possible degradation under the processing conditions. The results are shown in Fig. 7. Compared with the untreated pristine components, their spectra are combined in the spectrum of the extrudate, in which no new peak could be detected. It suggests that no detectable decomposition occurred during the extrusion.

Carvedilol has two common polymorphic crystalline forms. These are CAR form I and form II.\(^{18,19}\) In the absorption region of O―H and N―H stretching vibration, CAR form II has one strong band at ~3340 cm\(^{-1}\), and CAR form I has two strong bands at ~3250 cm\(^{-1}\) and 3450 cm\(^{-1}\). E100 has only two small peaks in this region.

In the spectra of E100 CAR extrudates, beside the appearing peaks of CAR around 1600 cm\(^{-1}\), one broader peak was observable from ~3300 to 3450 cm\(^{-1}\), which demonstrates the intermolecular interaction between E100 and CAR. It suggests that CAR is not in crystalline form because the characteristic peaks of crystalline phases of the drug (form I and form II) are not present in the spectra. The broad peak indicates probably that CAR is present in amorphous form. This deduction corresponds to the results of Pokharkar et al.\(^{20}\) Further analyses were made by polarized light microscopy and X-ray diffraction in order to confirm that CAR is really in amorphous form in the extrudates.

**Polarized light microscopy**

The birefringent CAR crystals are well observable using polarized light microscopy because of the different color of the background (Fig. 8), which is the result of the double refraction of the polarized light. In contrast, the amorphous materials (E100 and amorphous CAR) exhibit the same color as the background.

The investigations of crystalline CAR and the extrudates made by polarized light microscope confirmed that there was no detectable crystalline phase in the extrudates. Moreover, the extrudates did not display any apparent inhomogeneities. It suggested again that the extrudates are amorphous.

![Figure 7. Fourier transform infrared spectra; (A) 9CAR20SC milled, (B) E100 powder, (C) CAR form II, and (D) CAR form I.](image1)

![Figure 8. Polarized light microscopic images; (A) CAR form II and (B) 6CAR20.](image2)
**X-ray diffraction**

Being a well-established method to characterize and distinguish crystalline and amorphous materials, X-ray diffraction was used to characterize the morphological changes of CAR induced by extrusion. After grinding, the plots of the extrudates prepared in experiments 6CAR20 and 9CAR20SC were compared with those of the individual components and their physical mixtures (Fig. 9). Although CAR preserved its form II polymorphic structure in the physical mixtures, it is transformed to fully amorphous form when extruded with amorphous E100 matrix. There was no sharp peak of crystalline CAR in the diffractograms of the extrudates in contrast to the physical mixture of the polymer matrix and the crystalline drug.

**Capillary electrophoresis**

The dissolved drug was investigated by capillary electrophoresis. Information is provided by number, integral, and appearance time of peaks of the UV spectra recorded by the diode-array detector.

The results showed that there is just one peak in the electropherogram of the extrudates, as well as in the reference (pure CAR), which confirms the chemical stability of CAR during the extrusion process (Fig. 10). The migration time changed, which is probably due to presence of E100. Indeed, E100 is a cationic polymer, which can modify the surface charges of the silica capillary and the electro-osmotic flow.

The UV spectra of each studied sample at the peak of the analyzed substance were the same, proving that there is no (detectable) degraded CAR in the sample (Fig. 11).

---

**Figure 9.** X-ray diffractograms; (A) CAR form II, (B) physical mixture of E100 CAR 5%, (C) 9CAR20SC milled, (D) 6CAR20 milled, and (E) E100 powder.

**Figure 10.** Electropherograms; (A) pure CAR, (B) 9CAR20SC, and (C) 6CAR20.
Raman microspectrometry

Raman mapping was performed to characterize the homogeneity of CAR in the E100 matrix. The reference sample was again the physical mixture of the E100 powder and CAR. The inhomogeneous distribution of CAR in the physical mixture is very well observable and is not surprising as each component preserved its size during the mixing process (Fig. 12B). The homogeneous distribution observable in the Raman maps of the extrudates seems to be, however, more uniform than expected based on other extrusion products.[21] Figure 12(A) illustrates that the distribution of CAR is almost perfectly homogeneous because of the extrusion process. Both components can be seen at the same time on the measured area. The figure on the left corresponds to the CAR, with spectral concentrations between 28% and 33% in all measured points (this is why it is green in every pixel, see color scaling). The figure on the right shows the distribution of the E100 on the same area, which is again almost perfectly homogeneous. The color scaling is the same in each figure so the pictures of the physical mixture and the extrudate can be directly compared with each other.

The relative standard deviation (RSD) of the calculated ratio of the substances was found to be less than 3% in cases of the conventional extrudate and scCO2‐aided extrudate, whereas the RSD for the physical mixture was more than 45% (Table 2). The slight difference between the RSDs of two types of extrudates is comparable to the error of the Raman‐mapping method. Compared with recent relevant published results, the RSD value of the samples prepared by melt extrusion is significantly lower than in the case of high‐shear granulation (6%) and fluid‐bed granulation (17%).[22] It means that extrusion technology is very effective in homogenization compared with the widely used traditional processes of granulation. Achieving appropriate homogeneity is a serious challenge especially in the case of the solid dosage forms with low dosage rate (<2%), such as hormone‐containing drugs. Application of extrusion technology for formulation of these types of drugs might be useful because of very effective homogenization.

Drug content

The drug content of the extrudates, determined with UV spectrophotometer, corresponded in all cases to the initially introduced CAR, and the RSD (1.31%) of the measured contents was under the established limit of U.S. Pharmacopeia (6%).[23]

In vitro dissolution and drug content determination

The in vitro dissolution of the different samples exhibited significant differences. The dissolution curves are shown in Fig. 13. Unprocessed crystalline CAR (curve C) shows moderate dissolution rate, which could be dramatically improved by HME with E100. 100% was dissolved within 5 min (curves A and B). It was found that the E100 matrix governs the dissolution rate of the extrudates. CAR molecules dissolved simultaneously with the E100 matrix because the drug, being in amorphous form, does not need any additional action to overcome the crystal lattice energy. It is consistent with the fact that CAR is molecularly dispersed or nearly molecularly dispersed in the E100 matrix.

In the very first minutes, the dissolution of the milled foamed sample is even slightly more rapid in comparison with the unfoamed samples. This is probably because there is an inner porosity, which results in an increased surface area accessible to the solvent: the dissolution rate depending on the surface area of the material exposed to the dissolution medium.[24]

It appears that the use of HME is a very efficient way to improve the dissolution rate of a drug dispersed in a polymer matrix. In addition, the use of scCO2 gives a tool to enhance this dissolution rate.

CONCLUSION

In this work, fast‐dissolving formulation of a poorly water‐soluble drug against hypertension, CAR, was achieved by conventional and scCO2‐assisted HME process using Eudragit E100 as a polymer matrix. The manufactured fine solid dispersions of CAR

![Figure 11. The UV spectra recorded at the peaks detected during capillary electrophoresis analyses; (A) CAR Form II, (B) 9CAR20SC, and (C) 6CAR20.](image-url)
in Eudragit E stabilize the drug in its amorphous form. This was demonstrated by means of several analytical methods, polarized light microscopy, Raman microspectrometry, FTIR spectrometry, and X-ray diffractometry. The dissolution-improving efficiency was also due to the matrix of Eudragit E: its dissolution rate governing the rate of the drug release. Moreover, mild operating conditions and high productivity of the extrusion technology could be achieved by decreasing the viscosity with the aid of scCO₂ as a temporary plasticizer. Upon processing, no degradation of the drug could be observed.

A potential future perspective of scCO₂-aided melt extrusion is the introduction of lipophilic drug into the extruder together with scCO₂ as a solution. This process can be promising for preparation of solid formulations with low dosage such as hormones where achieving appropriate homogeneity and dissolution is a serious challenge for pharmaceutical technologists.

<table>
<thead>
<tr>
<th>Statistical attribute</th>
<th>Extrudate (6CAR20)</th>
<th>Supercritical extrudate (9CAR20SC)</th>
<th>Physical mixture (CAR 20%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean</td>
<td>0.324</td>
<td>0.330</td>
<td>0.319</td>
</tr>
<tr>
<td>Standard deviation</td>
<td>0.00605</td>
<td>0.00979</td>
<td>0.14426</td>
</tr>
<tr>
<td>Relative standard deviation (%)</td>
<td>1.87</td>
<td>2.95</td>
<td>45.25</td>
</tr>
<tr>
<td>Minimum observed score</td>
<td>0.28410</td>
<td>0.29439</td>
<td>0.06996</td>
</tr>
<tr>
<td>Maximum observed score</td>
<td>0.33283</td>
<td>0.35228</td>
<td>0.86459</td>
</tr>
<tr>
<td>Range (maximum minimum)</td>
<td>0.04873</td>
<td>0.05790</td>
<td>0.79463</td>
</tr>
</tbody>
</table>

CAR, carvedilol.
Acknowledgements

This research was supported by OTKA Research Fund (code K76346) and ERA Chemistry (code NN 82426). The authors would like to express their special thanks to Zsuzsa Szabó for her generous support of Zsombor Kristóf Nagy’s scholarship. This work is connected to the scientific program of the “Development of quality-oriented and harmonized R + D + I strategy and functional model at BME” project. This project is supported by the New Hungary Development Plan (Project ID: TÁMOP-4.2.1/B-09/1/KMR-2010-0002).

REFERENCES