Enhancement of solubility of dexibuprofen applying mixed hydrotropic solubilization technique

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Summary

Dexibuprofen, is a practically water-insoluble nonsteroidal anti-inflammatory drug which has a better anti-inflammatory effect than ibuprofen. A mixed hydrotropic solubilization technique was applied in order to improve the aqueous solubility and dissolution rate of dexibuprofen. Nine formulae were prepared using different concentrations of hydrotropic agents (sodium citrate dihydrate and urea). The prepared formulae were inspected visually for color and odor. Hygroscopicity, micromeric properties, solubility, and pH for 1% aqueous solutions were determined. In-vitro dissolution studies of the different prepared formulae were performed adopting the USP XXII dissolution method type I basket apparatus method. The prepared formulae were characterized by infrared (IR) spectroscopy and differential scanning calorimetry (DSC). The prepared formulae were a white color, odorless, slightly hygroscopic and exhibited good flow properties. Formulae containing higher amounts of hydrotropic agents exhibited an increase in the pH, solubility, rate and amount of dexibuprofen released from the dissolution medium. The highest dissolution rate was achieved from the F9 formula at drug:sodium citrate dihydrate:urea ratio (1:3:7.5). IR and DSC thermograph of dexibuprofen, hydrotropic agents and prepared formulae indicated the presence of intermolecular interaction between drug and hydrotropic agents which increased solubility and dissolution rate of drug, also, there is no chemical interaction confirming the stability of the drug with hydrotropic agents.

Keywords: Solubility, dexibuprofen, hydrotropic agents, formulae

1. Introduction

Dexibuprofen ((S)-(+) ibuprofen), which has better anti-inflammatory effects than ibuprofen and less gastric damage belongs to class II of the Biopharmaceutical Classification System (BCS) having low water solubility which is the rate limiting step in absorption of drug (1,2).

Poor solubility manifests many in vivo limitations like incomplete release, poor bioavailability, food effects, and higher inter-subject variability. However, different efforts have been demonstrated to improve bioavailability by increasing dissolution rate, for example: formulation of solid dispersions, solid solutions, micronization, nanosuspension, co-crystal molecular encapsulation with cyclodextrin, co-solvency, hydrotropy, spray drying, solubilization with surfactant, microemulsion, salt formation, polymorphism and combinations of effects (3).

Hydrotropy is the term originally put forward by Neuberg (4), to describe the increase in the solubility of a solute by the addition of fairly high concentrations of alkali metal salts of various organic acids. However, the term has been used in the literature to designate non-micelle-forming substances, either liquids or solids, organic or inorganic, capable of solubilizing insoluble compounds. The hydrotropic solubilization process involves cooperative intermolecular interaction with several balancing molecular forces, rather than either a specific complexion event or a process dominated by a medium effect, such as co-solvency or salting-in. Hydrotropic agents have been observed to enhance the aqueous solubility of poorly water-soluble drugs.

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The aim of our work is to prepare a soluble form of dexibuprofen which can be used in various dosage forms.

2. Materials and Methods

2.1. Materials

Dexibuprofen ((S)-(+)·ibuprofen) standard was purchased from Sigma-Aldrich (St. Louis, MO, USA). Dexibuprofen raw material was a gift from Future Pharmaceutical Industries (FPI) (Badr city, Cairo, Egypt). Distilled water was prepared in Central Nervous System (CNS) lab, National Organization of Drug Control and Research (NODCAR) (Dokki, Giza, Egypt). Methanol was purchased from Sigma-Aldrich (Taufkirchen, Germany). Sodium citrate dihydrate and urea were purchased from Medex (Medical Export Co Ltd, Near Naseby, Northamptonshire, UK).

2.2. Equipment


2.3. Preparation of formulae

Nine formulae were prepared using different concentrations of hydrotropic agents (sodium citrate dihydrate and urea) (Table 1). Urea and sodium citrate were accurately weighed. A minimum (possible) quantity of distilled water at 80-85°C contained in a 250 mL beaker was used to dissolve the urea and sodium citrate for quick dissolution. Then, dexibuprofen was added at 30-40°C and a teflon coated magnetic bead was dropped in it. Stirring of magnetic bead in beaker was started using a magnetic stirrer, maintaining the temperature at 30-40°C. Dexibuprofen was completely solubilized. Stirring was continued until a semisolid mass was obtained in the beaker (after evaporation of a large quantity of water). The semisolid mass obtained was spread on several watch glasses in thin layers for quick drying. The watch glasses were kept in an oven, maintained at 40°C for drying. When the mass became pulverizable, it was triturated with the help of mortar and pestle and again kept in oven for drying. After almost complete drying, the powder of the solid dispersion was passed through sieve #100 and was kept for 6 days in a desiccator containing silica gel and stored in air-tight containers.

2.4. Evaluation of the prepared formulae

2.4.1. Visual inspection

The prepared formulae were inspected visually for color and odor.

2.4.2. Determination of loading efficacy

A weighed amount of each of the prepared formulae equivalent to a theoretical content of 100 mg drug was accurately weighed and allowed to disintegrate completely in 100 mL distilled water, by sonicing for 20 min so as to dissolve drug and hydrotropic agents. One mL of each solution was diluted to 100 mL with distilled water, the absorbance of the solution was measured spectrophotometrically at λmax 222 nm using distilled water as blank. The concentration of the drug was calculated. Each sample was assayed in triplicate and an average of three determinations was calculated.

2.4.3. Solubility and pH

Solubility was determined by taking an excess quantity of each of the prepared formulae in fixed volumes of solvent (distilled water). The resulting solution was placed on a water bath sonicator for 24 h. After equilibrium the samples were centrifuged and then supernatant was filtered through a 0.45 μm filter membrane and the concentration of drug in the saturated solution in formulae under investigation was determined spectrophotometrically at λmax 222 nm after appropriate dilution of the filtrate using distilled water as a blank. Each experiment was done in triplicate and an average of three determinations was calculated. The pH for 1% aqueous solution was determined.

2.3.4. Hygroscopicity

Hygroscopicity was measured for 10 g powder (m1)
stored on a watch glass at 25°C, 80% R.H. for 24 h and the mass gain (m₂) was recorded (7). The percent mass gain (% m/m) was calculated from the following equation: 
\[
\text{% m/m} = \left( \frac{m_2 - m_1}{m_1} \right) \times 100.
\]

2.3.5. **In vitro dissolution studies**

*In vitro* dissolution studies of the different prepared formulae were performed adopting the USP XXII dissolution method type I basket apparatus method. The samples were separately encapsulated in transparent hard gelatin capsules. Five hundred mL of distilled water was used as dissolution media at 37 ± 0.5°C maintaining stirring speed at 50 rpm. The samples were withdrawn at specified time intervals and replaced with an equivalent volume of fresh dissolution medium at different time intervals to keep the volume constant and estimated spectrophotometrically at \(\lambda_{\text{max}}\) 222 nm for dissolving drug using distilled water as a blank (8). All dissolution tests were performed in triplicate and an average of three determinations was calculated.

2.3.6. **Micromeritic properties**

The prepared formulae were characterized by their micromeretic properties such as bulk and tapped densities, Hausner ratio, % compressability and angle of repose.

Determination of bulk and tapped densities: Five grams powder (m) in 250 mL graduated cylinder and the volume occupied (bulk volume \(v_b\)) recorded. The graduated cylinder was tapped till constant volume was obtained and the volume of powder (true or tapped volume \(v_t\)) was then recorded. The bulk density \(P_b\) and tapped density \(P_t\) were calculated in g/mL by dividing the weight over the corresponding volume. Hausner ratio and % compressibility (Carr’s index) were calculated using measured values for bulk density \(P_b\) and tapped density \(P_t\) according to the following equations (9): 
\[
\text{Hausner ratio} = \frac{P_t}{P_b}, \quad \text{Ci}\% = \left( \frac{P_t - P_b}{P_t} \right) \times 100.
\]
Ci% is the % compressability of the powder. Each value reported is an average of three determinations. Determination of powder flowability: The fixed highest cone method was adopted. A cut-stem glass funnel having an internal diameter of 0.6 cm was used. The funnel was held at 2.5 cm height (h) over a flat surface. The powder sample (prepared formulae) was allowed to flow gently through the funnel until the cone was formed and reached the funnel surface orifice, powder flow was stopped and the average diameter of the formed cone (D) was determined. The area of base of cone was taken as a measure of internal friction between particles. The angle of repose was calculated by the equation (10): 
\[
\tan (\alpha) = \frac{2h}{D}.
\]

The micromeritic properties were done in triplicate and an average of three determinations was calculated.

2.3.7. **Fourier transform infrared spectroscopy (FTIR)**

An amount of 2-3 mg of dexibuprofen, sodium citrate dihydrate, urea, and prepared formulae were mixed separately with 400 mg of dry potassium bromide powder, compressed into transparent discs and their IR spectra was recorded in the wave length region of 4,000-400 cm\(^{-1}\).

2.3.8. **DSC**

DSC was used to evaluate changes of dexibuprofen characteristic peak and heat enthalpy that might occur after mixing with hydrotropic agents. Thermograms of dexibuprofen, sodium citrate dihydrate, urea, and best formula that showed highest dissolution were recorded using Shimadzu DSC-50 calorimeter. The instrument was calibrated using purified indium (99.99%). Samples of (2-4 mg) of each substances as well as the best formula were heated under a nitrogen atmosphere as a carrier gas on an aluminum pan at a flow rate 25 mL/min and a heating rate of 10°C/min over a temperature range of 20 to 300°C.

3. Results and Discussion

3.1. **Visual inspection**

The prepared formulae were white in color and odorless.

3.2. **Determination of loading efficacy**

The entrapment efficacy (EE) of drug loading in the prepared formulae are presented in Table 2. It ranged from 96.67 to 99.79% indicating that there is no effect of hydrotropic agents on the drug.

3.3. **Solubility and pH**

Table 3 shows the phase solubility study and pH for 1% aqueous solution of the prepared formulae. It was noticed that increasing the amount of sodium citrate dihydrate and urea lead to increasing the solubility.

<table>
<thead>
<tr>
<th>Formula</th>
<th>Theoretical percent of dexibuprofen (%)</th>
<th>Entrapment efficiency (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>F1</td>
<td>22.22</td>
<td>96.67</td>
</tr>
<tr>
<td>F2</td>
<td>18.18</td>
<td>94.61</td>
</tr>
<tr>
<td>F3</td>
<td>15.38</td>
<td>98.31</td>
</tr>
<tr>
<td>F4</td>
<td>14.28</td>
<td>98.67</td>
</tr>
<tr>
<td>F5</td>
<td>12.5</td>
<td>97.76</td>
</tr>
<tr>
<td>F6</td>
<td>11.11</td>
<td>96.49</td>
</tr>
<tr>
<td>F7</td>
<td>10.53</td>
<td>99.62</td>
</tr>
<tr>
<td>F8</td>
<td>9.52</td>
<td>99.79</td>
</tr>
<tr>
<td>F9</td>
<td>8.69</td>
<td>99.42</td>
</tr>
</tbody>
</table>

Table 2. **Entrapment efficiency of the prepared formulae**
This is due to increasing the concentrations of alkali metal salts of various acids (4), and it is concluded that the solubility of dexibuprofen increases synergistically by mixed hydrotropy. The improvement of solubility may be due to changing the crystal forms, different habit, structure and surface modification (11). The pH for 1% aqueous solution ranges from 6.4 to 7.8 and that increasing the amount of sodium citrate and urea lead to increasing the pH. The solubility of dexibuprofen increases slightly with an increase in pH, which may be due to the acidic nature of dexibuprofen. Thus, it can be said that, the solubility enhancement of drug by hydrotropes is not entirely due to a pH effect, but it is largely due to hydrotropy (12).

### 3.4. Hygroscopicity

The % mass gain of the prepared formulae is presented in Table 3. It ranges from 0.161 to 0.568 so the prepared formulae are described as slightly hygroscopic (7).

### 3.5. In vitro dissolution studies

The dissolution profiles of dexibuprofen from the prepared formulae were done and graphically represented in Figures 1, 2, and 3. Table 4 shows the percent dexibuprofen dissolved from different formulae. Dissolution of drug depends on physicochemical and physicotechnical properties of drug particle. These forms directly affect the absorption kinetics of drug and bioavailability of formulae. This assumes greater importance for a drug exhibiting low solubility that makes absorption to be dissolution rate limited. It is established that the modification of the polymorphic state of a compound can bring an increase in solubility (13). It is obvious that the percent drug dissolved from the prepared formulae containing higher amounts of sodium citrate dihydrate and urea (F7, F8, and F9) (92.46, 92.67, and 97.57% after 10 min) which is higher than the percent drug dissolved from the prepared formulae containing lower amounts of sodium citrate dihydrate and urea (F1, F2, and F3) (53.47, 54.83, and 65.97% after 20 min). The improvement of dissolution

<table>
<thead>
<tr>
<th>Formula</th>
<th>Solubility (g/100mL) for dexibuprofen in the prepared formulae</th>
<th>pH (1% solution)</th>
<th>Percent mass gain (%m/m)</th>
</tr>
</thead>
<tbody>
<tr>
<td>F1</td>
<td>0.287</td>
<td>6.4</td>
<td>0.328%</td>
</tr>
<tr>
<td>F2</td>
<td>0.598</td>
<td>6.8</td>
<td>0.568%</td>
</tr>
<tr>
<td>F3</td>
<td>0.619</td>
<td>7.0</td>
<td>0.357%</td>
</tr>
<tr>
<td>F4</td>
<td>0.753</td>
<td>6.5</td>
<td>0.161%</td>
</tr>
<tr>
<td>F5</td>
<td>1.885</td>
<td>6.9</td>
<td>0.346%</td>
</tr>
<tr>
<td>F6</td>
<td>2.247</td>
<td>7.1</td>
<td>0.421%</td>
</tr>
<tr>
<td>F7</td>
<td>2.354</td>
<td>6.5</td>
<td>0.344%</td>
</tr>
<tr>
<td>F8</td>
<td>3.091</td>
<td>6.8</td>
<td>0.293%</td>
</tr>
<tr>
<td>F9</td>
<td>4.182</td>
<td>7.8</td>
<td>0.431%</td>
</tr>
</tbody>
</table>

Figure 1. In vitro release profile of dexibuprofen from F1, F2, and F3 formulae. 500 mL distilled water was used as dissolution media at 37 ± 0.5°C while maintaining stirring speed at 50 rpm. Samples were estimated spectrophotometrically at λ <sub>max</sub> 222 nm for dissolving drug in triplicate and an average of three determinations was calculated.

Figure 2. In vitro release profile of dexibuprofen from F4, F5, and F6 formulae. 500 mL distilled water was used as dissolution media at 37 ± 0.5°C while maintaining stirring speed at 50 rpm. Samples were estimated spectrophotometrically at λ <sub>max</sub> 222 nm for dissolving drug in triplicate and an average of three determinations was calculated.

Figure 3. In vitro release profile of dexibuprofen from F7, F8, and F9 formulae. 500 mL distilled water was used as dissolution media at 37 ± 0.5°C while maintaining stirring speed at 50 rpm. Samples were estimated spectrophotometrically at λ <sub>max</sub> 222 nm for dissolving drug in triplicate and an average of three determinations was calculated.
behavior could be attributed to solubilization of the drug using hydrotropic agents (8). The faster dissolution could be due to better solubility of the prepared formulae that contains higher amounts of hydrotropic agents because increasing the amount of hydrotropic agents in the formulae leads to increased solubility and dissolution rate.

3.6. Micromeritic properties

The results of micromeretic properties of the prepared formulae are shown in Table 5. Bulk densities ranged from 0.43 to 0.45 g/mL and tapped densities ranged from 0.48 to 0.51 g/mL. Hausner ratio values ranged from 1.13 to 1.15 indicating low to moderate inter-particle friction and thus good flow properties (14). They showed good flow with respect to average % compressibility which ranged from (11.09 to 11.71%). Also, the angle of repose ranged from 31.30° to 34.42° indicating good flow properties (15).

3.7. Fourier transform infrared spectroscopy (FTIR)

The collected FTIR of plain dexibuprofen drug, sodium citrate dihydrate, urea and the prepared formulae are shown in (Figure 4). The main peaks characteristic of the drug are identical and well identified in the prepared formulae. This can be summarized: prominent peaks at 3,087, 1,707, 1,406, 1,050 cm⁻¹ corresponding to O-H stretching, C=O stretching, C-C stretching, O-H bending respectively. The FTIR showed no changes occurred in the chemical nature of drug indicating absence of chemical interaction between drug and hydrotropic agents and so, confirming drug stability. From these results, it can be speculated that a drug-hydrotropic agent hydrogen bond existed in this formula and caused reduced drug recrystallization (16), also intermolecular interaction between drug and hydrotropic agents occurred (5,6).

3.8. DSC

DSC pattern of plain dexibuprofen, sodium citrate dihydrate, urea and the F9 formula are recorded in Figure 5. It is clear that, the DSC of plain dexibuprofen drug showed an endothermic peak at onset of 47.25°C reaching a peak at 56.51°C, an endset of 66.49°C and the enthalpy was 210.40 J/g while hydrotropic agents; sodium citrate and urea showed an endothermic peak at 169.8°C and 138.53°C, respectively. The endothermic peak of drug is related to its melting point. There was a slight difference in melting endotherms of selected formulae as well as decreasing intensity and loss of sharpness compared to that of pure drug (drug = 62.52°C, sodium citrate dihydrate = 143.65°C, urea = 103.05°C). These results might be explained in terms of a presence of intermolecular interaction between drug and hydrotropic agents. Basically the thermal process of any mixture is the sum of individual components. However, there was invariably very little
change in transition temperature when mixing two or more components. When two substances are mixed, the purity of each may be reduced and generally very slight lower melting endotherms may result (17). This explains the very slight shift drug hydrotropic agent peak. Previous work (18), accepts small events and considers them to not represent any determinable interaction. This could be due to differences in moisture content of samples (19). Also, some changes in the peak shape and height to width ratio can be seen because of possible differences in the mixture sample geometry (20). The observed change in enthalpy indicates a change in crystallinity of the drug (21), resulting in an increase in dissolution rate.

In conclusion, the mixed hydrotropic solubilization technique significantly influenced solubilization of dexibuprofen which is practically insoluble thus contributing to dissolution improvement. Increasing the amount of hydrotropic agents leads to an increase in pH, solubility, rate and amount of dexibuprofen release from the dissolution medium. The highest dissolution
rate was achieved from F9 formula at drug: hydrotropic agent ratio 1:3:7.5. The prepared formulae are slightly hygroscopic and show good flow properties. Drug stability with hydrotropic agents and intermolecular interaction between them has been confirmed by FTIR and DSC results.

Acknowledgements

This work was supported by a grant from National Organization of Drug Control and Research and Department of Pharmaceutical Technology, Faculty of Pharmacy, Tanta University.

References


(Received April 7, 2014; Revised August 13, 2014; Accepted August 22, 2014)