THE RELEASE OF SODIUM DICLOFENAC
FROM MATRIX TYPE OF TRANSDERMAL PATCH

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ABSTRACT

The sodium diclofenac released from transdermal patch of combination of ethyl cellulose (EC) N-20 and polyvinyl pyrrolidone (PVP) K-30 was investigated. In this study, matrix-type of transdermal patch containing diclofenac sodium were prepared using polymeric combination of EC N-20 and PVP K-30 in various ratios (9:1 (Formula I); 8:2 (Formula II); and 7:3 (Formula III)). The percentage of moisture content was calculated as a difference between initial and final weight with respect to final weight. The homogeneity of patch surface was determined using fluorescence microscope and scanning electron microscope (SEM). Released test using dissolution apparatus were carried out in 500 mL of phosphate buffer saline pH 7.4 ± 0.05 at temperature 37 ± 0.5°C with speed of swirl 50 rpm. Results were analyzed by statistic programmed of SPSS using one way analysis of variance with degree of believed 95% (α = 0.05). It can be concluded that the combination of EC N-20 and

Kata Kunci: Pelepasan Sodium diclofenak, plasticizer, kombinasi matrix dari EC N-20 dan PVP K-30
PVP K-30 at ratio 7:3 was the best choice for manufacturing transdermal patch based on physicochemical and the release profile.

**Keywords:** Diclofenac sodium released, plasticizer, combination matrix of EC N-20 and PVP K-30

**INTRODUCTION**

These last few years, the transdermal route has become one of the most successful and innovative focus for research in drug delivery. One dosage form is provided by transdermal patch dosage forms. Patch is a dosage form that aims to transport drugs through the skin into the blood circle [1]. Patch will be effective when the drug must obviously be able to penetrate into the skin barrier and reach the target [2]. Criteria these are typically used for selection include parenteral dose less than 20 mg/day, short half-life (requiring multiple daily doses in current dosage form), acceptable skin toxicity (little or no skin irritation or sensitization), molecular weight less than 500 Daltons, log octanol-water partition coefficient of approximately 0–3 and solubility in mineral oil and water greater than 1 mg/mL³.

Some of the drug when given in oral form, drugs undergoes substansial hepatic first-pass metabolism and give side effects such as irritation of the stomach, so better if given in the patch dosage form [3]. Patch preferred over intravenous administration for not causing pain, tissue damage and removes the fear of patients [3].

Sodium diclofenac is reported used for topical applications. The drug undergoes substansial hepatic first-pass metabolism and only 50% of administered dose reaches systemic circulation [4].

The aim of the present study is to know how the influence of a combination of ethyl cellulose polymer and polyvinylpyrrolidone in the ratio 9:1, 8:2 and 7:3 in increasing the flux of diclofenac sodium release from matrix 100 mg/50 cm², added menthol as an enhancer with levels of 1% and polyethylene glycol 400 as plasticizer with levels of 20%, with a method of making a matrix controlled, so as to obtain a combination of polymers with a precise comparison to diclofenac sodium dosage patch matrix type.

**METHOD**

**Material**

Sodium diclofenac (Aarti Drugs Limited), PVP K-30 (SP (Singapore) Pte Ltd), EC N-22 (Dow Chemical Company), and menthol were purchased from PT Bratachem. All ingredients of these were pharmaceutical grade, alcohol 96% was analytical grade.

**Preparation**

Dosage sodium diclofenac patch matrix type controlled by the ratio of different polymers made planned to produce an average weight of 0.500 grams with a diameter of 3 cm, surface area 7.065 cm² (table 1).PVP K-30 and EC N-20, respectively - were dissolved in ethanol 96%. Solution of PVP K-30 included in the solution of N-20 EC, stirring until evenly mixed. Previously menthol solution was made by dissolving 0.050 grams of menthol in ethanol 96% to 100 mL.
The Release of Sodium Diclofenac from Matrix Type of Transdermal Patch

Table 1. Formula of the sodium diclofenac patch base

<table>
<thead>
<tr>
<th>Material</th>
<th>Concentration (mg/7.065 cm²)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Formula 1</td>
</tr>
<tr>
<td>Sodium diclofenac</td>
<td>14.13 mg/7.065 cm²</td>
</tr>
<tr>
<td>EC N-20</td>
<td>342.78</td>
</tr>
<tr>
<td>PVP K-30</td>
<td>38.09</td>
</tr>
<tr>
<td>Polyethylenglikol 400</td>
<td>100</td>
</tr>
<tr>
<td>Menthol</td>
<td>5</td>
</tr>
</tbody>
</table>

Figure 1. The results of organoleptics evaluation of the sodium diclofenac patch, A=formula 9:1; B= formula 8:2 and C= formula 7:3

For each - each formula, 1.0 mL of menthol solution was poured into the sodium diclofenac, stirred until dissolved. Mixture of diclofenac sodium and menthol added to the mixture of PVP K-30 and EC N-20. Stirred carefully until all the ingredients mixed. Polyethylene glycol 400 and the remaining 96% ethanol solutions were added gradually while continue to be stirring until homogeneous. After a homogeneous preparation was poured into a mold and dried at room temperature for 24 hours.

Homogeneity of the patch surface

Homogeneity of patch surface was analyzed using a fluorescence microscope and a scanning electron microscope (SEM).

In Vitro Release Studies

The in vitro release of the patches was performed using USP dissolution apparatus 5-paddle over disk with completely cell diffusion.

The patches (in cell diffusion) were placed respective in the dissolution chamber. All in vitro released study were performed at 50 rpm, with each medium of dissolution (phosphate buffer saline pH 7.4 ± 0.05 at temperature 37 ± 0.5°C) was 500 mL. The samples withdrawn at different time intervals were analyzed for drug content using spectrophotometer Double Beam UV-VIS Recording UV 160 A (Shimadzu). All of the results were analyzed by statistic programmed of SPSS using one way analysis of variance with degree of believed 95% (α = 0.05).
The Release of Sodium Diclofenac from Matrix Type of Transdermal Patch

![Figure 2](image1)

**Figure 2.** Histogram of the moisture content patch sodium diclofenac. Each column represents the mean ± SD (n=3).

![Figure 3](image2)

**Figure 3.** The results of Fluorescence microscope images of diclofenac sodium patch using electron microscope Olympus DX 41 Nokuler Three types of TF with 100x magnification. A: patch type polymer matrix with a ratio of ethyl cellulose (EC) N-20-polyvinylpyrrolidone (PVP) K-30 in a row is: A1 = 9:1; A2 = 8:2; A3 = 7:3; control patch type polymer matrix with a ratio of ethyl cellulose (EC) N-20-polyvinylpyrrolidone (PVP) K-30 in a row are: B1 = 9:1; B2 = 8:2; B3 = 7:3
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RESULT AND DISCUSSION

The results of organoleptics evaluation showed that the composition of patch formulation using combination polymer EC N-20 and PVP K-30 at ratio 9:1 and 8:2 more rigid and not easy to breaking compared with that of ratio 7:3 (figure 1). Physicochemical parameter of the prepared formulation were evaluated in this study. The percentage moisture content of the formulation was determined (figure 2). The result of moisture content studies showed that the %value ± SD of moisture content from 3 replication of formula 1 was 8.67 ± 1.33; formula 2 12.30 ± 3.42 and formula 3 was 13.30 ± 1.65. The results showed with an increasing percentage of hydrophilic polymers, PVP K-30, in the formulation, moisture content increased.

Fluorescence microscope and a Scanning electron microscope (SEM) study showed the surface of the sodium diclofenac matrix (figure 3 dan 4). Observations using electron microscopy at 100x magnification showing the addition of polymers PVP-K30 diclofenac sodium patch causes the surface to be uneven, and this is because the formation of pores on the surface of the matrix that will assist the release of diclofenac sodium. To clarify the results observed with electron microscope photographs, images can be compared with surface patches using a scanning Electron Microscope (SEM) with 500x magnification. Results from SEM images show the formation of pores on the surface of diclofenac sodium patch. The greater the levels of polyvinylpyrrolidone (PVP) K-30, the pores formed are larger and spread out evenly across the surface of the patch.

The results of in vitro drug release are presented in figure 5, showed the released process of sodium diclofenac from patch base. The cumulative amount of the drug released from patch base was plotted as a function root of time and a linear regression analysis was used to determine the flux of sodium diclofenac.

Figure 6 shows the flux of sodium diclofenac from the patch base. Based on the in vitro released study, the flux of diclofenac from patch base were 37.482 ± 15.529 µg/cm²/minute½; 34.587 ± 4.027 µg/cm²/minute½; and 43.540 ± 2.555 µg/cm²/minute½ respectively for Formula I, Formula II and Formula III. The result of statistic using ANOVA one way showed that Fcalculated (0.712) > Ftable (5.14). It means that there was no significant difference in increased concentration of PVP K-30 in formulation of sodium diclofenac patch. But the highest flux of piroxicam was 43.540±2.555 µg/cm²/minute½ with the highest concentration of PVP K-30 and the smallest concentration of the EC N-22 between all of the formula. PVP K-30 in matrix would form pores for released sodium diclofenac. This phenomena was similar with result of the preview research.
that said with increasing the hydrophilic polymer the formed of pores in the matrix increased, so that the released of the drug increased [5].

Based on this study, the using of polymer combination between PVP K-30 and EC N-22 could increase flux released of sodium diclofenac from patch base. The ratio of 7:3 for PVP K-30 and EC N-30 at Formula III was the optimal composition for base of patch sodium diclofenac type matrix with menthol as enhancer and PEG 400 as plasticizer.

CONCLUSION

The polymeric combination PVP K-30 and EC N-22 at ratio 7:3 (Formula III) was the best choice for manufacturing transdermal patch base of sodium diclofenac among the formulations studies with the highest value of flux (43.540±2.555 μg/cm²/minute$.^{1/2}$).

Figure 5. Profil of Sodium diclofenac released from patch bases to phosphate buffer saline solution pH 7.4±0.05 at 37±0.5 °C, with speed of swirl 50 rpm. Each column represents the mean ± SD (n=4).

Figure 6. Histogram of the flux of Sodium diclofenac released from patch bases to phosphate buffer saline solution pH 7.4±0.05 at 37±0.5 °C, with speed of swirl 50 rpm. Each column represents the mean ± SD (n=4).

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