

## FORMULATION AND EVALUATION OF *IN VITRO* TRANSDERMAL PATCH DICLOFENAC SODIUM USING CHITOSAN POLYMER AND POLYVINYL ALCOHOL CROSS-LINKED TRIPOLYPHOSPHATE SODIUM

ERNAWATY GINTING\*, JULIA REVENY, SUMAIYAH

Department of Pharmacy, Universitas Sumatera Utara, Indonesia. Email: Ernawatyg@yahoo.com

Received: 06 February 2018, Revised and Accepted: 24 April 2018

### ABSTRACT

**Objective:** The aim of this study was to investigate diclofenac sodium patches using chitosan (Ch) and polyvinyl (PVA) alcohol cross-linked tripolyphosphate sodium (TPP) to increased transdermal permeation of the drug from the matrix system across rabbit skin.

**Materials and Methods:** The chemical characterization of diclofenac sodium was done by ultraviolet-visible spectrophotometry. Formulation of diclofenac sodium patches using solvent evaporation method with cross-link technique. Evaluation of physical character of the film includes organoleptic observation, weight test, thickness, % moisture absorption, fold resistance, interaction between materials used Fourier transform infrared (FTIR), and active substance levels. The drug release was determined using Franz diffusion cells in phosphate buffer (pH 7.4).

**Results:** The result of physiochemical parameters of the transdermal patch were found satisfactory. Formula F1, F2, F3, and F4 produce patches with fine texture, F5 and F6 formulas produce patches with coarse texture. Formula F1, F2, F5, and F6 are flexible and fulfill the multiplier test requirements, while the F3 and F4 formulas are not flexible and do not meet the multiplier test requirements. The patch weight, thickness, and drug content were uniform. The release of patches with following the zero-order release. The optimal formula with the total sodium diclofenac released as much as 17.89  $\mu\text{g}$ , the release flux of 89.42  $\mu\text{g}/\text{cm}^2/\text{h}$  of permeation time of 10 h and the moisture absorption rate of  $1.07 \pm 0.193$ . The FTIR data of rabbit skin indicated Ch and PVA alcohol cross-linked TPP increase transdermal permeation of diclofenac sodium in the stratum corneum.

**Conclusion:** The diclofenac sodium can be prepared by cross-linked method, resulted in a better discharge profile.

**Key words:** Diclofenac sodium, Chitosan, Polyvinyl alcohol, Tripolyphosphate sodium, Glycerin, Release kinetics.

© 2018 The Authors. Published by Innovare Academic Sciences Pvt Ltd. This is an open access article under the CC BY license (<http://creativecommons.org/licenses/by/4.0/>) DOI: <http://dx.doi.org/10.22159/ajpcr.2018.v11i8.25145>

### INTRODUCTION

The transdermal patch is designed to be able to deliver the drug in a controlled manner directly to the blood vessels [1]. The transdermal advantages such as to avoid absorption problem in the gastrointestinal tract, prevent the first cross effect, may be the choice for drugs not to be orally administered, may control the drug [29]. Transdermal can be formulated in ointment, cream, gel emulsion, and patch form [5]. The controlled delivery of drugs in the blood vessels can be achieved through the matrix system [36]. The basic components of the matrix system consist of polymers [7], drugs, enhancers, and other excipients [34]. Variations of polymer compositions, enhancers, and plasticizers will result in different release and permeation profiles [37]. The polymer in the matrix system will control the release rate of the drug from the patch matrix [3].

This study used diclofenac sodium as a test model of *in vitro* release and permeation using rabbit skin [33]. Diclofenac sodium includes non-steroidal anti-inflammatory drugs that act by inhibiting cyclooxygenase in a non-selective manner, resulting in inhibition of prostaglandin synthesis, which is a pain mediator [2]. The diclofenac sodium has a half-life of 1–2 h so that it is orally administered in a few times, leading to adverse side effects on the stomach and reducing patient compliance [4]. Stratum corneum is a limiting factor in the speed of percutaneous drug taking [23]. The diclofenac sodium has a fairly high lipophilicity with a low partition coefficient value, octanol/water log P at 25°C that is 4.17 causing low diclofenac sodium permeability [46].

This study used chitosan (Ch) polymer excipients, tripolyphosphate sodium (TPP), alcohol polyvinyl (PVA), and glycerol plasticizer [35]. Ch is widely used because it has the characteristics of biocompatibility, non-

toxic, and antimicrobial [13]. Ch and alcohol PVA that are hydrophilic will increase patch permeability and drug diffusion [12,32]. Ch with alcohol PVA produces cross-linking hydrogels with good swelling activity. The combination of Ch and alcohol PVA anhydrous maleic cross-links by alprazolam model resulted in patches with physical characteristics, permeability, and release profiles [29]. Penetration test of Ch cross-linked tripolyphosphate sodium (TPP) with diclofenac diethylamine model showed 30-min cross-link time yielding the best permeation through mouse skin membrane [39]. The combination of Ch and alcohol PVA cross-linked tripolyphosphate sodium using bovine serum albumin produced a controlled release film that could be characterized by Fourier transform infrared spectroscopy (FTIR), scanning electron microscope (SEM), and release test. Studies on the use of glycerin plasticizer using Ch - alcohol PVA polymer resulted in a controlled patch [52].

### MATERIALS AND METHODS

#### Materials

##### Chemicals

The materials used in this study were diclofenac sodium (Kimia Farma, Indonesia), PVA alcohol, TPP, Ch, glycerin, acetic acid, hydrogen phosphate sodium, bromide potassium, and chloride potassium (Merck).

##### Animals

Adult male New Zealand albino rabbits (1.5–2 kg) were supplied by the Animal House of Pharmacology, Laboratory of Pharmacy Department at Universitas Sumatera Utara was inhabited under standard laboratory condition with proper diet. The animal were accepted after the proposed



AQ7

**Table 1:** The composition of the diclofenac sodium patch

| Formula | Cross-linked time | Glycerin (%) | TPP (%) | Ch-PVA (2 and 8%) | Diclofenac sodium (mg) |
|---------|-------------------|--------------|---------|-------------------|------------------------|
| F1      | 10                | 5            | 1       | 30:70             | 20                     |
| F2      | 20                | 5            | 1       | 30:70             | 20                     |
| F3      | 10                | 5            | 1       | 50:50             | 20                     |
| F4      | 20                | 5            | 1       | 50:50             | 20                     |
| F5      | 10                | 5            | 1       | 70:30             | 20                     |
| F6      | 20                | 5            | 1       | 70:30             | 20                     |

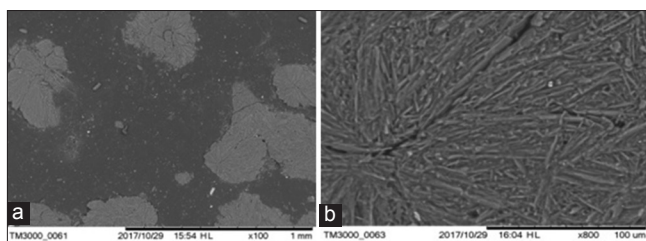
TPP: Tripolyphosphate sodium, Ch: Chitosan, PVA: Polyvinyl

**Table 2:** Drug content and physical parameters of different formulation of patch

| Formulation code | Uniformity of weight (mg) | Film thickness (mm) | Folding endurance | % Drug content |
|------------------|---------------------------|---------------------|-------------------|----------------|
| F1               | 172±0.574                 | 0.253±0.0010        | 631               | 95.08±0.626    |
| F2               | 177±0.812                 | 0.262±0.0066        | 631               | 96.37±0.102    |
| F3               | 184±0.812                 | 0.274±0.0014        | 8                 | 97.54±0.376    |
| F4               | 181±1.000                 | 0.266±0.0016        | 8                 | 95.74±0.359    |
| F5               | 184±0.812                 | 0.266±0.0016        | 724               | 97.54±0.376    |
| F6               | 185±1.153                 | 0.267±0.0010        | 722               | 97.46±0.343    |

**Table 3:** *In vitro* cumulative percentage of diclofenac sodium release

| No | Time | F1    | F2    | F3    | F4    | F5    | F6    |
|----|------|-------|-------|-------|-------|-------|-------|
| 1  | 10   | 4.99  | 3.53  | 4.25  | 1.17  | 5.42  | 1.48  |
| 2  | 20   | 10.11 | 7.58  | 10.49 | 2.4   | 11.43 | 3.16  |
| 3  | 30   | 15.52 | 12.21 | 15.63 | 5.31  | 17.7  | 5.96  |
| 4  | 60   | 21.83 | 17.22 | 21.9  | 8.2   | 24.57 | 7.51  |
| 5  | 120  | 27.72 | 23.46 | 28.73 | 11.31 | 31.81 | 10.09 |
| 6  | 180  | 34.6  | 30.37 | 35.97 | 14.78 | 39.3  | 12.9  |
| 7  | 240  | 42.27 | 37.82 | 43.78 | 18.2  | 47.07 | 16.34 |
| 8  | 300  | 50.35 | 45.62 | 52.23 | 21.21 | 55.47 | 19.11 |
| 9  | 360  | 58.81 | 54.06 | 62.82 | 23.1  | 64.01 | 22.89 |
| 10 | 420  | 68.12 | 63.97 | 70.22 | 25.68 | 73.24 | 26.93 |
| 11 | 480  | 77.62 | 74.88 | 77.99 | 28.69 | 82.64 | 31.13 |
| 12 | 540  | 85.94 | 84.24 | 87.91 | 30.15 | 91.8  | 34.36 |
| 13 | 600  | 94.07 | 92.82 | 95.76 | 32.69 | 99.81 | 37.24 |

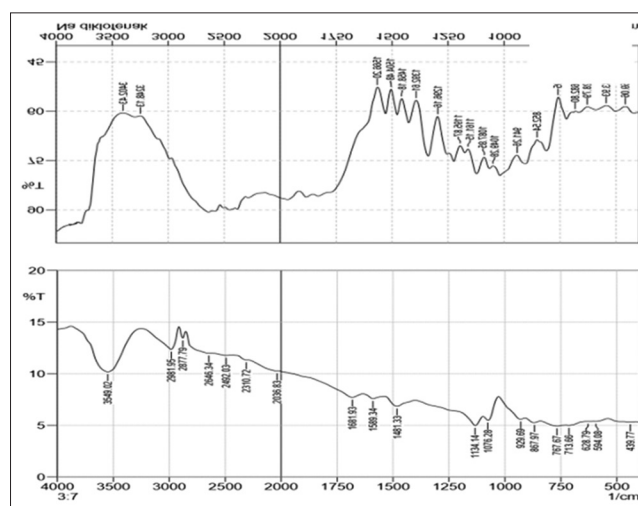
**Fig. 1:** The morphology of patch sodium natrium diclofenac at ×100 (a) and ×800 (b)

3437.15/cm in the presence of the  $\text{NH}_2$  and OH groups in the 1600.92/cm, 1666.50/cm region in the presence of C=O.

Observation of FTIR PVA shows peak spectrum at 3375.43/cm region with the presence of OH group. FTIR observations of TPP showed peak spectrum at 3430.03/cm, 3383.14/cm, 3217.27/cm, and 1157.29/cm. Ch interaction TPP resulted in a shift of absorption peak at 1549.02/cm, 1481.33/cm, and 1134/cm. Peak at 3549.02/cm indicated no interaction between sodium diclofenac and the excipient polymer.

#### Permeation study

All the six formulations were subjected to *in vitro* permeation studies across rabbit skin for establishing the permeation parameters [15,19]. The results of permeation study were shown on Table 3 and Fig. 3. According to the penetration test, cumulative value of penetrated sodium diclofenac was found highest in F5 (cumulative % 99.81 time

**Fig. 2:** Infrared spectra of pure of sodium diclofenac and patch sodium diclofenac

10 h), followed by F3, F1, and F2. Cumulative value of penetrated sodium diclofenac was found lowest in F6 and F4 with cross-linked time 20 min. **The release of patches with formulas was following zero-order release** (Table 4). Permeation study was showed F1, F3, and F5 with cross-linked time 10 min has bigger % cumulative and better release than F2, F4, and F6 with cross-linked time 20 min.

AQ2

#### CONCLUSION

In the present study, various formulation of transdermal sodium diclofenac patches were prepared using cross-linked method of Ch polymer, polyvinyl alcohol, and tripolyphosphate sodium. This research showed that the formulation of transdermal diclofenac sodium patches using cross-linked method with cross-linked time of 10 min has better profile release than cross-linked time of 20 min. Formulation of transdermal sodium diclofenac patches using cross-linked method needed optimal drying to make good characteristic and profile release.

#### AUTHOR'S CONTRIBUTION

The first author has carried out the research and provided study conception. The second and third authors have provided study conception, the design of work, drafting of the manuscript, and critical revision.

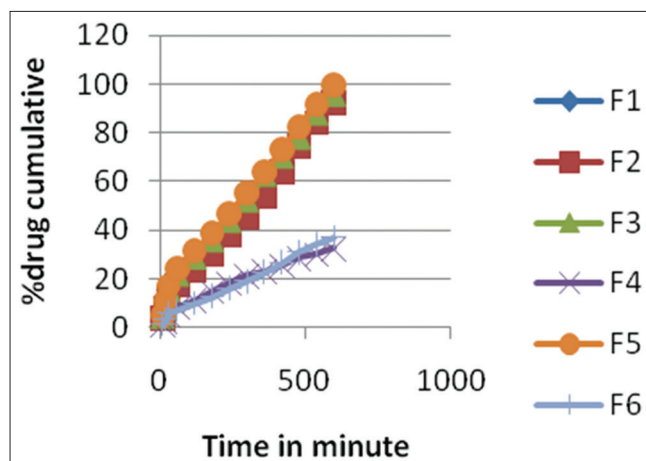


Fig. 3: In vitro profile permeation of various patches

Table 4: R<sup>2</sup> value from of zero order, Higuchi and Korsmeyer plot

|                  | F1    | F2    | F3    | F4    | F5    | F6    |
|------------------|-------|-------|-------|-------|-------|-------|
| Orde nol         | 0.994 | 0.994 | 0.993 | 0.973 | 0.991 | 0.994 |
| Korsmeyer-Peppas | 0.982 | 0.982 | 0.977 | 0.973 | 0.979 | 0.974 |
| Higuchi          | 0.965 | 0.95  | 0.971 | 0.996 | 0.973 | 0.959 |

#### CONFLICT OF INTEREST

The authors declare that there is no conflict of interest regarding the publication of this article.

AQ5

#### REFERENCES

- Agoes G. Sistem Penghantaran Obat Pelepasan Terkendali. 2<sup>nd</sup>. Bandung: Penerbit ITB; 2008. p. 373.
- Agustin R, Ratih H. Dissolution Profile of Sustained Release Tablet of Diclofenac Sodium Using Metolose 90 Sh 4000. J Sains Farm Klin 2015;1:176-83.
- Alexander A, Dwivedi S, Ajazuddin, Giri TK, Saraf S, Saraf S, et al. Approaches for breaking the barriers of drug permeation through transdermal drug delivery. J Control Release 2012;164:26-40.
- Allen L, Ansel HC. Ansel's Pharmaceutical Dosage Forms and Drug Delivery Systems. Philadelphia: Lippincott Williams & Wilkins; 2013. p. 295.
- Ansel HC. Pengantar Bentuk Sediaan Farmasi. Edisi Keempat. Jakarta: UI-Press; 2005. p. 176-81.
- Banakar UV, Pharmaceutical Dissolution Testing. New York: Marcel Dekker, Inc. 1991. J Pharm 1992;151:26-32.
- Barhate SD, Potdar MB. Formulation of transdermal patch of Carvedilol by using novel polymers. Pharmacia 2012;2:185-9.
- Bartosova L, Bajgar J. Transdermal drug delivery *in vitro* using diffusion cells. Curr Med Chem 2012;19:4671-7.
- Bazigha K, Rasool A, Eman F, Fahmy A, Sadan S. Development and evaluation of ibuprofen transdermal gel formulation. Trop J Pharm Res 2010;9:355-63.
- Bhattarai N, Gunn J, Zhang, M. Chitosan hydrogels for controlled, localized drug delivery. Adv Drug Deliv Syst 2010;62:83-99.
- Bhavani PP, Kumar PR, Shankar RK, Santosh T. Formulation and evaluation studies on transdermal dosage forms of diclofenac sodium. World J Pharm Pharm Sci 2015;4:1043-63.
- CanAS, Erdal MS, Gungor S, Ozsoy Y. Optimization and characterization of chitosan film for transdermal delivery of ondansetron. Molecules 2013;18:5455-71.
- Cheung RC, Ng TB, Wong JH, Chan WY. An update potential biomedical and pharmaceutical application. Mar Drug 2013;13:5156.
- Das A, Ahmed AB. Formulation and evaluation of transdermal patch of indomethacin containing of pathcouli oil as natural penetration enhancer. Asian J Pharm Clin Res 2017;10:320-5.

- Dash S, Murthy PN, Nath L, Chowdhury P. Kinetic modeling on drug release from controlled drug delivery system. Acta Pol Pharm 2010;67:217-23.
- Devissaguet J, Aiache JM. Farmasetika 2 Biofarmasi. 2<sup>nd</sup> ed. Surabaya: Widji Soerarti. Airlangga University Press; 1993. p. 443-58, 172.
- Flyn GL, Stewart B. Percutaneous drug penetration choose candidates for transdermal drug development. Drug Dev Res 1988;13:169-85.
- Fox SC. Remington Education Pharmaceutics. London: Pharmaceutical Press; 2014. p. 432-4.
- Gadhekar R, Saurabh MK, Thakur GS, Saurabh A. study of formulation, characterisation and wound healing potential of transdermal patches curcumin. Asian J Pharm Clin Res 2012;5:225-30.
- Giri TK, Thakur A, Alexander A, Badwaik H, Tripathi DK. Modified chitosan hydrogels as drug delivery and tissue engineering system, present status and application. Acta Pharm Silaica B 2012;2:439-49.
- Goci EN, Haloci EN, Xhulaj SK, Malaj LE. Formulation and *in vitro* evaluation of diclofenac sodium gel. Int J Pharm Pharm Sci 2014;6:259-61.
- Goswami DS, Uppad N, Gloyal S, Mehta N, Gupta AK. Permeation enhancer for transdermal drug delivery from natural and synthetic sources. J Biomed Pharm Res 2013;2:19-29.
- Gupta JR, Irchiya R, Garud N, Priyanka T. Formulation evaluation of matrix type transdermal patches of glibenclamide. Int J Pharm Sci Drug Res 2011;1:46-50.
- Jadhav RT, Kasture PV, Gattani SG, Surana SJ. Formulation and evaluation of transdermal films of diclofenac sodium. Int J Pharm Tech Res 2009;1:1507-11.
- Jahan L, Ferdaus R, Shabeen SM, Sultan MZ, Mazid MA. *In vitro* transdermal delivery of metformin from HPMC/PVA based transdermal drug delivery patch et different pH. J Sci Res 2011;3:651-7.
- Jatav VS, Jitendra S, Rakesh KJ, Sharma K, Ravindra PS. Recent advances in development of transdermal patches. Pharmacopore 2011;2:287-97.
- Liang S, Liu L, Huang Q, Yam KL. Preparation of single or double network chitosan/poly(Vinyl Alcohol) gel film through selectively cross-linking method. Carbohydrate Polym 2009;77:718-24.
- Kouchak M, Ameri A, Naseri B, Boldag SK. Chitosan and polyvinyl alcohol composite film containing nitrofurazone, preparation and evaluation. J Basic Med Sci 2014;17:14-20.
- Malji P, Gandhi A, Jena S, Maji N. Preparation and characterization of maleic anhydride cross linked chitosan polyvinyl alcohol hydrogel matrix transdermal patch. J Pharm Sci Tech 2013;2:62-7.
- Patel H, Bhimani B, Patel G. Transdermal drug delivery system. as prominent dosage form for the highly lipophilic drugs. Int J Pharm Biosci 2012;1:42-65.
- Patil SS, Gupta RM, Gupta KS, Dodayya H. Formulation and characterization of tpp cross-linked chitosan microspheres loaded with lornoxicam. J Biomed Pharm Res 2014;3:51-8.
- Pouranvari S, Ebrahimi F, Javadi G, Maddah B. Chemical cross-linking of chitosan/polyvinyl alcohol electrospun nanofibers. Original Sci Article 2016;50:663.
- Raj AR. Formulation evaluation and *in vitro* permeation studies of transdermal nifedipine from matrix type patch. Int J Pharm Pharm Sci 2013;6:185-8.
- Raza R, Mittal A, Kumar P, Alam S, Prakash S, Chauhan N. Approach and Evaluation Of Transdermal Drug Delivery System. Int J Dev Res 2015;7:222-33.
- Rowe RC, Sheskey PJ, Owen SC. Handbook of Pharmaceutical Excipients. 5<sup>th</sup> ed. London: The Pharmaceutical Press; 2006. p. 217-611.
- Sachan R, Bajpai M. Transdermal drug delivery system a review. Int J Dev Pharm Life Sci 2013;3:748-65.
- Saroha K, Yadav B, Sharma, B. Transdermal patc, a discrete dosage form. Int J Curr Pharm Res 2011;3:98-108.
- Setyawan EI. Pengaruh Kombinasi HPMC dan MC Terhadap Karakter Fisik dan Pelepasan Ketoprofen dari Matrik patch Transdermal. Yogyakarta: Tesis Universitas Gajah Mada; 2013. p. 73.
- Sharma K, Singh V, Arora A. Natural biodegradable polymer as matrices in transdermal drug delivery. Int J Drug Dev Res 2011;3:85-103.
- Sherwood L. Human Physiology, From Cells to Systems. 6<sup>th</sup> ed. Belmont, CA: Cengage Learning; 2012. p. 485-6.
- Ng SF, Rouse JJ, Sanderson FD, Meidan V, Eccleston GM. Validation of static franz diffusion cell system for *in vitro* permeation studies. AAPS Pharm Sci Tech 2010;2:1432-41.
- Singh I, Sri P. Percutaneous penetration enhancement in transdermal drug delivery. Asian J Pharm 2010;4:92-101.
- Sinko PJ. Martin's Physical Pharmacy and Sciences. 6<sup>th</sup> ed. Philadelphia: Lippincott William, Philadelphia Pharmaceutical; 2016. p. 223.

44. Soujanya C, Lakshmi Satya B, Lokesh Reddy M, Manogna K, Ravi Prakash P, Ramesh A. Formulation and *in vitro* and *in vivo* evaluation of transdermal patches of lornoxicam using natural permeation enhancers. *Int J Pharm Pharm Sci* 2014;6:282-6.
45. Subrata M, Kumar PD, Mukherjee B, Sengupta S, Pattnaik S, Chakraborty S. Optimization of *in-vitro* permeation pattern of ketorolac tromethamine transdermal patches. *Iran J Pharm Res* 2011;10:193-201.
46. The Board of Trustees. United States Pharmacopeia. 27<sup>th</sup> ed. National Formulary Rockville: The Board of Trustees; 2007.
47. Varshosaz J, Karimzadeh S. Development of cross-linked chitosan for oral mucosal delivery of lidocaine. *Res Int Pharm Sci* 2007;2:43-52.
48. Wasitaatmaja SM, Penuntun Ilmu Kosmetik dan Medik. Indonesia: Penerbit Universitas Jakarta; 1997. p. 3-5.
49. Watkinson AC. Transdermal and topical drug delivery today, Transdermal and Topical Drug Delivery. Principles and Practice. Hoboken, NJ, USA: John Wiley Sons Inc.; 2012. p. 357-66.
50. Wester RC, Maibach HI, Regional Variation in Percutaneous Absorption Drugs, Cosmetic, Mechanism, Methodology. New York: Marcel Dekker; 1997. p. 107.
51. Wise DL. Handbook of Pharmaceutical Controlled Release Technology. New York: Marcel Dekker; 2000. p. 321.
52. Wang Q, Du Y, Fan L. Properties of chitosan/poly(vinyl alcohol) film for drug controlled release. *J Appl Polim Sci* 2004;96:808-13.
53. Xiong Y, Liu QL, Zhang QG, Zhu AM. Synthesis and characterization of cross-linked quaternized poly(vinyl alcohol)/chitosan composite anion exchange membranes for fuel cells. *J Power Sour* 2008;183:447-53.

#### Author Queries???

AQ1:Please check the edit made throughout the file.

AQ2:Kindly review the sentence.

AQ3:Please review the sentence for more clarity.

AQ4:Please check the term.

AQ5:Kindly cite references 9,10, 17,18,20,22,24,25,28,30,38,40,41,42,47,48,49,50,51 in the text part and also chronological order