PROCEEDING

The International Conference on Pharmacy and Advanced Pharmaceutical Sciences

Faculty of Pharmacy UGM Yogyakarta Indonesia October 2009









PROCEEDING

The International Conference on Pharmacy and Advanced Pharmaceutical Sciences Yogyakarta, Indonesia, 2009

Editors:

Pudjono
Hilda Ismail
Ronny Martien
Triana Hertiani
Ritmaleni

Published by: Faculty of Pharmacy Universitas Gadjah Mada Sekip Utara, Yogyakarta, 5281, Indonesia

ISBN: 978-979-95107-7-8

First Edition, 2010

No part of this publication may be reproduced, stored in a retrieval system or transmitted in any form or by any means, electronic, mechanical, photocopying, recording or otherwise, wi thout the prior written permission of the publisher, Faculty of Pharmacy, Universitas Gadjah Mada, Yogyakarta, Indonesia.

No responsibility is assumed by the publisher for any injury and/or damage to persons or property as a matter of product l iability, negligence or otherwise, or from any use or operation of any methods, products, instructions or ideas contained in the material herein.

Printed in Yogyakarta, Indonesia

Preface from the Editor

The proceeding was produced based on papers and posters presented at the international Conference on Pharmacy and Advanced Pharmaceutical Sciences, held in Yogyakarta, Indonesia, 5 – 6 October 2009.

The proceeding clearly reflects broad interest; from there are participants coming from all around the world. Many contributions on Pharmaceutical Sciences there are quite a substantial number of papers on Pharmacist role in general. The papers presented file into a broad spectrum in Pharmaceutical sciences including Pharmacology, Toxicology, Analytical Chemistry and Drug Design, Drugs Synthesis, Formulation of Drugs, Pharmacy Social, Pharmacoepidemy, Traditional Medicine Natural Product Chemistry and Phytochemistry, etc.

In addition there are substantial numbers of paper deal with professional aspect of Pharmacist in general health care.

In this an opportunity, I would like to express my appreciation to the editorial team of the proceeding who have been working hard to review manuscripts, and making the first edition of this proceeding be possible.

I would like also to thanks to all invited speakers and presenters who participated in the International Conference on Pharmacy and Advanced Pharmaceutical Sciences and your contribution to this proceeding.

Finally, I hope this proceeding will give contribution to the advanced scientific research in the field of pharmaceutical sciences

Yogyakarta, July 2010

Dr. rer. nat. Pudjono, SU., Apt.

Organizing Committee

Steering committee

Prof. Dr. Marchaban, DESS, Apt (Pharmacy, UGM, INDONESIA)
Prof. Dr. Umar Anggara Jenie, M.Sc., Apt (LIPI, INDONESIA)
Prof. Dr. Retno S. Sudibyo, M.Sc., Apt. (Pharmacy, UGM, INDONESIA)
Prof. Dr. Syed Azhar S. Sulaiman (USM, MALAYSIA)
Prof. Masashi Kawaichi, Ph.D (NAIST JAPAN)
Dr. Edy Meiyanto, M.Si., Apt (CCRC, UGM INDONESIA)
Prof. Dr. Zullies Ikawati, Apt (Pharmacy, UGM, INDONESIA)

Chairman

Dr. Hilda Ismail, M.Si., Apt.

Secretary

Dr. rer. nat. Ronny Martien, M.Si

Treasurer

Dr. Ratna Asmah S., M.Si., Apt

Scientific Committee

Prof. Dr. Sismindari, SU., Apt.
Prof. Dr. Lukman Hakim, M.Sc., Apt
Prof. Dr. Suwidjijo Pramono, Apt.
Prof. Dr. Suwaldi, M.Sc., Apt

Welcome Message From the committee

Welcome to Yogyakarta

On behalf of the Scientific and Organizing Committees, it is a great pleasure for me to welcome all participants to Yogyakarta, to the International Conference on Pharmacy and Advanced Pharmaceutical Science 2009.

The international conference is organized by the faculty of Pharmacy UGM to celebrate its 63th anniversary and the Lustrum XII of Gadjah Mada University, as a collaboration work between the Faculty of Pharmacy UGM with the Nara Institute of Science and Technology (Japan) and the Universiti Sains Malaysia (Malaysia). In this conference 15 lectures within the field of Pharmaceutical Care and Advanced Pharmaceutical Science will be given by invited speakers. Besides, 55 posters and 75 paper will be presented in the parallels presentation sessions. Herewith, we express our gratitude to all speakers and presenter, who would like to share their advance knowledge in this scientific event.

The Organizing Committee gratefully acknowledges the Nara Institute of Science and Technology and the Universiti Sains Malyasia, for the nice collaboration in bringing forth this conference. A special acknowledgment is addressed to the Rector of Gadjah Mada University and the sponsors, for all supports that make this symposium possible. Furthermore, personally, I want to express my deep appreciation to the members of the Organizing Committee, for the good teamwork and their great effort given in the preparation for this symposium.

Finally, I wish all participants a scientifically rewarding and an enjoyable meeting in Yogyakarta.

Chairman

Dr. Hilda Ismail, M.Si., Apt.

Remark of the Dean Faculty

Assalamu'alaikum wr. wb.
Distinguished ladies & gentlemen.

First of all, on be half of the Faculty of Pharmacy Universitas Gadjah Mada, I would like come to all of you in Yogyakarta, thank you very much for your attention to come and to attend the international Symposium on Pharmacy and Advanced Pharmaceutical Sciences. I hope we are all in health condition.

Ladies and gentlemen.

The symposium is organized by the Faculty of Pharmacy UGM in collaboration with the Faculty of Pharmaceutical Sciences Universiti Sains Malaysia and the Nara Institute of Science and Technology Japan, and held as part to celebrate the 63th anniversary of the Faculty of Pharmacy UGM.

In the symposium, I hope we can communicate our recently information concerning social / clinical pharmacy and pharmaceutical sciences. I hope the symposium will be very fruitfull, very useful for all of us.

I addres special thanks to the plenary speakers both from domestic and aboard, the oral and poster presenters, as well as to those who come just to know the development of clinical or social pharmacy and pharmaceutical science. Your willingness to come , to communicate and to share your experiences is highly appreciated.

Special thanks also I address to my colleague the Dean of Faculty of Pharmacy USM who has been coordinating USM students to attend this symposium. The hope is not to set up networking between the pharmacy students of USM and UGM.

Therefore, during almost whole day discussing scientific matter related to human health and welfare, I hope we can make a wonderful opportunity to make a scientific closer relationship while we enjoy the cultural performances of Yogyakarta presented by our pharmacy student.

Finally, I hope that this meeting will give benefits to all of us, and we may see each other again in a similar event in the near future.

I look forward to thank you all for attending this event.

Wassalamu'alaikum wa rahmatullahi wa barakatuh, Dean of Faculty of Pharmacy UGM

Prof. Dr. Marchaban, DESS., Apt.

Speech of the Senior Vice Rector For Education, Research and Community Services, Gadjah Mada University

Assalamu'alaikum wa rahmatulLahi wa barakatuh,

On behalf of the Rector, I would like to welcome all of you to our campus Gadjah Mada University and to our home town Yogyakarta. It is a great honor for me and Gadjah Mada University to host the Two-day International Conference on Pharmacy and Pharmaceutical Sciences that is conducted by the Faculty of Pharmacy, Gadjah Mada University. The increasing problems and new cases of some diseases in the world, both the infectious and the degenerative diseases, have demanded the development of medical and pharmaceutical sciences and technologies for supporting the developments of early detection methods of the diseases, the accurate diagnoses, as well as the appropriate and effective medications or therapy. Pharmaceutical Science and Technology have been developing very fast within recent years. The development trend shows using much more biotechnological approach in both diagnose establishment and medication administrations. For examples the usage of some serums, enzymes, hormones, vaccines, etc., and their recombinant products. The science and technology for finding prevention method against infectious diseases or degenerative diseases now have been developing so amazing, for example the usage of growth hormones, vaccines, and stem cells for it.

Gadjah Mada University has been committed to become World Class University; therefore international networking in education, research and publication is much needed. I really support to this international conference on Pharmaceutical Science and Technology which can keep us in touch with the state of the art of pharmaceutical science. I do believe that by conducting this kind of international meeting, we can get and exchange new information and best practices on pharmaceutical science and technology, and it is very important to inspire our young researchers and enhance our research networking internationally. In this occasion, I would like to express my great gratitude to all the guest speakers and speakers, who have contributed their advanced presentations in this international conference. I also would like to extend my gratitude to the Organizing Committee from the Faculty of Pharmacy, Gadjah Mada University, who has already successfully arranged this international conference. I would also thank to all institutions or companies who have sponsored and supported this conference.

Finally, have a fruitful conference and enjoy Yogyakarta. Thank you Wassalamu'alaikum wa rahmatulLahi wa barakatuh,

Senior Vice Rector for Education, Research and Community Service Gadjah Mada University

Prof. Dr. Retno Sunarminingsih, M.Sc., Apt.

CONTENTS

Preface from the Editor	i
Organizing Committee	ii
Welcome Message Proceeding International Conference on Pharmacy and Advanced Pharmaceutical Sciences	
From the committee	iii
Remark of the Dean Facultyi	V
Senior Vice Rector For Education	v
CONTENT	vi
Pharmacogenetics: in case of cytochrome P450 oxidases (CYPS) related to adverse drug reactions Arum Pratiwi, Harianto Lim and Ronny Martien	1-4
Interaction of turmeric and garlic extract combination against free radical scavenging activity Patonah, Daryono H. Tjahjono, Elin Yulinah S. and I Ketut Adnyana	5 – 6
Influenced of Kojic Acid and B-Cyclodextrin on SPF Value Sunscreen Product Contained Oxybenzone and Octyl Dimetyl Paba (3:7) (In vanishing cream base formulation) Diana, Tristiana Erawati, Widji Soeratri and Noorma Rosita	7 – 14
Isolation and Antimicrobial activity of endophytic fungi <i>Kabatiella caulivora</i> var B isolated from <i>Alyxia reinwardtii</i> BL Noor Erma Sugijanto, Dian Anggraeny and Noor Cholies Zaini	15 – 17
Rapid and Simple Luciferase Reporter Gene Assays for the Discovery of Peroxisome Proliferator-Activated Receptor α and γ Agonists and Nuclear Factor- κ B Inhibitors from Medicinal Plants. N. Fakhrudin, S. Vogl, P. Picker, E. H. Heiss, J. Saukel, G. Reznicek, B. Kopp, A. G. Atanasov and V. M. Dirsch	18 – 24
Identification of components of essential oil from <i>Cananga odorata</i> which penetrated into the rat skin /(wistar strain) in the practice of <i>Timung</i> (development of <i>Timung</i> as alternative healing) Mangestuti Agil, Esti Hendradi and Budiastuti	25 – 29
In Vivo Antihyperglycemic Test of Albedo Durian (<i>Durio zibethinus</i> M) Extract on Aloxan-Induced Diabetic White Rat (<i>Rattus norvegicus</i>) F. M. Cahyani, I. Susanti, R. Ratna, Y. D. Panggi and Y. Pravitasari	30 – 33
Effect of Pasak Bumi's Root (<i>Eurycoma longifolia</i> , Jack) on Sperm Output in Rats Farida Hayati and Mustofa	34 – 37

The Influence of Arbutin 3% and Sesame Oil (3,5,7 % w/w) on SPF Values of Oxybenzon and Padimate O (3:7% w/w) in carbomer Gel Base Noorma Rosita, Tristiana Erawati and Rafi Jikrona	38 – 43
Sulochrin as α -glucosidase inhibitor <i>lead compound</i> Rizna Triana Dewi, Ahmad Darmawan, Sofna D.S Banjarnahor, Hani Mulyani, Marisa Angelina and Minarti	44 – 48
The Practice of Complementary Indigenous Malay Therapies In Rural Areas: Do Users' Attitudes, Beliefs And Perceptions Significantly Differ From Non-Users? Pei Lin Lua, Rohayu Izanwati Mohd Rawi, Suffian Mohamad Tajudin, Norlida Mamat and Ahmad Zubaidi Abdul Latif	49 – 54
An Interventional Pilot Study: Effect Of Dark Chocolate Consumption On Anxiety Level Among Female Nursing Students Sok Yee Wong, Pei Lin Lua, Rohayu Izanwati Mohd Rawi, Rokiah Awang, Ahmad Zubaidi and Abdul Latif	55 – 62
The Anti-proliferation Assay of Bioactive Fraction from <i>Curcuma zedoaria</i> Rhizome Ros Sumarny, Priyosoeryanto B. P., letje W., Latifah K. D. and Chairul	63 – 67
Studies of Sub-acuteToxicity Assay from <i>Acorus calamus</i> L. in Experimental Animal Models Banjarnahor S.D.S, Sri Hartati and Megawati	68 – 71
Antioxidant Properties and Phenolics Content of <i>Mikania scandens</i> L.(Wild) Sumi Wijaya, Ting Kang Nee, Khoo Teng Jin and Christophe Wiart	72 – 77
The Influence of Olive Oil Addition on Increasing of Arbutin Penetration in the Carbomer-940's Gel (Observation on Inhibition of Enzyme Tyrosinase Activity) Widji Soeratri, Tristiana Erawati, Noorma Rosita and Fahriyatul Wahyuni	78 – 81
The difference of antioxidant activity of various tea (<i>Camellia sinensis</i> L.) methanol extract Wahyu Widowati, Tati Herlina and Hana Ratnawati	82 – 88
Chemical Stability of Cisplatin and Ondansetron During Simulation of hemotherapy Administration Yahdiana Harahap, Rizka Andalusia and Armon Fernando	89 – 94
The Effects of Cassava Starch (<i>Manihot utilissima</i> , Pohl.) as a Binder on Physicochemical Characteristics of Acetaminophen Tablet Formulation Yandi Syukri, Tri Rahayu Ningsih and M. Hatta Prabawa	95 – 98
Drug Interaction Study in Hospitalized Hepatic Cirrhosis Patient in Dr. Ramelan Navy Hospital Amelia Lorensia, Aziz Hubeis, Widyati and Hary Bagijo	99 – 102
The Effect of Cold Storage in Krebs-Henseleit Buffer in the Viability and Metabolic Activities of Precision Cut Intestinal Slices Dewi Setyaningsih, AA Khan and GMM Groothuis	103 – 110

The Effect Of b-Cyclodextrin And Oxybenzone-Octyl Dimethyl Paba (3:7% W/W) Addition On The Penetration Of Kojic Acid In Vanishing Cream (Based on Activity Inhibition of Tyrosinase)	111 – 116
Diana Winarita, Tristiana Erawati, Noorma Rosita and Widji Soeratri	
Uniderstand Decree Leading 1	
The profile of knowledge and self-medication in handling cough symptoms by students of pharmacy at Airlangga university Elida Zairina, Liza Pristianty and Lestriana Kusumasari	117 – 120
Section 2015 to the following the contract of	
The Characteristics and Release of Diclofenac Sodium of Niosome System in Carbomer 940 Gel Base Preparation (Niosome System of Diclofenac Sodium-Span 60-Cholesterol with Molar Ratio 1:5:5)	121 – 128
Esti Hendradi, Tutiek Purwanti, Bety Nurfia Puspitarini and Bianda Ida Kurnia	
Red Betel Vine (Piper Crocatum) Essential Oil as Antituberculosis Farida Juliantina Rachmawaty	128 – 133
Effect of Pasak Bumi's Root (<i>Eurycoma longifolia</i> , Jack) on Sperm Output in Rats Farida Hayati and Mustofa	134 – 137
The Influence of Sesame Oil Addition on Arbutin Release from Carbomer-940 Gel Bases Hanifa Rahma, Tristiana Erawati and Noorma Rosita	138 – 141
Phytochemical Screening and Determination of Antioxidant Activity of Fractions from Ethyl Acetate Extract of Phyllanthus acidus (L.) Skeels Leaf Hindra Rahmawati, Hesty Utami and Moordiani	142 – 145
Study on Antihyperglicaemic Activitiy of Ethyl Acetate Extract of Sidaguri (<i>Sida rhombifolia</i> L.) Stem onAlloxan-Induced Diabetic Mice (<i>Mus musculus</i> L.) Irma Ratna K, Muktiningsih, Suhartono, Natalia Elisabeht and Muhammad Ali Zulfikar	146 – 152
The Influence of Arbutin and Olive Oil as an Enhancer in Characteristic and SPF Value of Sunscreen (Combination of Oxybenzone and Octyldimethyl Paba in <i>Carbomer</i> 940 Gel Base) Josephine Paramita Ayuningtyas, Tristiana Erawati, Noorma Rosita and Widji Soeratri	153 – 160
The Effect of Secondary Emollients Triethylhexanoate, Isopropyl myristate, and Propyleneglycol Isostearate on in-vitro skin penetration of tocopheryl acetate cream using Franz-diffusion cell Joshita Djajadisastra, Sutriyo and Fraida Aryani	161 – 165
Termina o Jeja di da di a di a di a di a di a di a d	
Immunomodulatory activity of Plantago major L. on IgM titer of mice Kartini, A. Kirtishanti, Dessy, Fauziah and Isnaini	166 – 169
Antibacterial activities of <i>Aleurites moluccana</i> (Euphorbiaceae) Othman Abd Samah and Rasyidah Mohamad Razar	170 – 178
Total synthesis and revised structure of benzophenone glucopyranosides from phaleria macrocarpa	179 – 185
Phebe Hendra, Yukiharu Fukushi and Yasuvuki Hashidoko	

Influence of Tween 80 Concentration in Carbomer/ Tween 80 Aggregate on Kojic Acid Penetration (Observed on Inhibiting Tyrosinase Activity in Vanishing Cream) Siti Evi Jayanti, Tristiana Erawati and Noorma Rosita	186 – 191
Validation for Result Degradation of Nifedipine Residue with Thin Layer Chromatography- Densitometry and Thin Layer Chromatography-Spectrophotometry Sitti faika and Sudibyo Martono	192 – 195
Synthesis and Biological Activity Test of Antibiotic UK-3 Analogues, 2-Hydroxynicotinyl-Butyl-Serine-Ester and Its Derivatives Ade Arsianti, Kiyomi Kakiuchi, Tsumoru Morimoto, M.Hanafi and Endang Saefudin	196 – 198
Vitamin e content in the dragon fruit Established by high performance thin layer chromatography—densitometry Any Guntarti and Warsi	199 – 204
Drug interaction study in hospitalized hepatic cirrhosis patient in Dr. Ramelan navy hospital Amelia Lorensia, Aziz Hubeis, Widyati and Hary Bagijo	205 – 208
PGV-1 inhibits G2M phase progression in WIDr colon cancer cell Endah Puji Septisetyani, Edy Meiyanto, Masashi Kawaichi and Muthi' Ikawati	209 – 212
Proceeding International Conference on Pharmacy and Advanced Pharmaceutical Sciences The influence of oleic acid pre-treatment on transport of epigallocathecin gallat in green tea (<i>Camellia sinensis</i> , L) extract Across mice skin in vitro Nining Sugihartini, Achmad Fudholi, Suwidjiyo Pramono and Sismindari	213 – 215
Development and Production of Anti Tuberculosis Fixed Dose Combinations (FDCs) Barokah Sri Utami, Syamsul Huda, Nurliya Irfiani and Badrus S.	216 – 218
The Characteristics and Release of Diclofenac Sodium of Niosome System in Carbomer 940 Gel Base Preparation (Niosome System of Diclofenac Sodium-Span 60-Cholesterol with Molar Ratio 1:5:5) Esti Hendradi, Tutiek Purwanti, Bety Nurfia Puspitarini and Bianda Ida Kurnia	219 – 224
The Characteristics and Release of Diclofenac Sodium of Niosome System in Carbomer 940 Gel Base Preparation (Niosome System of Diclofenac Sodium-Span 20-Cholesterol with Molar Ratio 1:5:5) Esti Hendradi, Tutiek Purwanti, Anditasari and Srimaryati	225 – 231
An Interventional Pilot Study: Effect Of Dark Chocolate Consumption On Anxiety Level Among Female Nursing Students Sok Yee Wong, Pei Lin Lua, Rohayu Izanwati Mohd Rawi, Rokiah Awang and Ahmad Zubaidi Abdul Latif	232 – 239
Antiemetics utilization in cancer patients with high emetogenic cytotoxic drugs in two governmental hospital in indonesia Dyah Aryani Perwitasari and Ana Hidayati	240 – 243
KEY WORDS INDEX	244
DISCUSSION	246

The Effect Of β -Cyclodextrin And Oxybenzone-Octyl Dimethyl Paba (3:7% W/W) Addition On The Penetration Of Kojic Acid In Vanishing Cream (Based on Activity Inhibition of Tyrosinase)

Diana Winarita*, Tristiana Erawati, Noorma Rosita, Widji Soeratri
Pharmaceutics Department, Faculty of Pharmacy Airlangga University, Surabaya-Indonesia
E-mail: deez_imoets@yahoo.com

Abstract

This research was aimed to investigat the effects of the addition of β -cyclodextrin (BCD) and a combined of oxybenzone and octyl dimethyl PABA (3:7% w/w) on the penetration of kojic acid based on inhibion activity of tyrosinase. The presence of BCD was expected to make the preparation effective, while the addition of oxybenzone and octyl dimethyl PABA was intended to overcome photosensitive properties of kojic acid. Effectiveness determination of vanishing cream preparation was conducted by testing activity inhibition of tyrosinase that was expressed as percentage inhibition. Tyrosinase was an enzyme that plays an important role in the process of melanin formation in human skin. Inhibition activity of tyrosinase by kojic acid was determined in vitro by observing absorption values of dopachrome (an intermediate product of melanin formation) using spectrophotometer. Based on results of effectiveness tests of kojicacid vanishing cream preparation by means of measurement of activity inhibition of tyrosinase, it was found that addition of sunscreen (oxybenzone and octyl dimethyl PABA without BCD) had no significant effect on the activity inhibition of tyrosinase. While, addition of BCD or combined of BCD-sunscreen significantly reduced inhibition of tyrosinase activity in vitro.

Keyword : kojic acid, β-cyclodextrin, oxybenzone, octyl dimethyl PABA, tyrosinase, enzym, inhibition

Introduction

Kojic acid is a skin lightening agent which has very small molecular size, so easily absorbed up through the basal membrane. Because of its small molecule, kojic acid hardly absorbed through the lipid membrane of its target sites, the melanocytes (Manosroi et al, 2005) and can penetrate into the systemic (Nakayama et al., 2005). It is likely absorbed through voids between cells on the skin. Losses due to kojic acid to the systemic is not able to work effectively inhibits melanin formation took place in the basal membrane of skin that results are not optimal lightening effect. Addition of β -cyclodextrin reported can decrease penetration of kojic acid (Nakayama et al., 2005). BCD can be used in a number of 0,5-10 parts per 1 weight part of kojic acid, which is the optimal composition of 4 parts cyclodextrin every a part of kojic acid (Hatae et al, 1988).

Generally, skin lightening maybe caused the skin more photosensitive, so in cosmetics usage needed the sun-rays filter (Zulkarnain, 2003). To obtain maximum protection, frequently sunscreen used in combination of anti UV-A with anti UV-B, ei: oxybenzone and octyl dimethyl PABA. In this study used the composition (3:7) % w/w which is the optimum composition in accordance with the results of previous studies (Wati, 2005).

This research investigated the effects of the addition of β -cyclodextrin and a combined of oxybenzone and octyl dimethyl PABA (3:7% b/b) on the penetration of kojic acid based on activity inhibition of tyrosinase. Effectivity determination of vanishing cream preparation was conducted by testing activity inhibition of tyrosinase that was expressed as percentage inhibition. Tyrosinase was an enzyme that played an important role in the process of melanin formation of human skin. Activity inhibition of tyrosinase by kojic acid was determined in vitro by observing absorption values of dopachrome (an intermediate product of melanin formation) using spectrophotometer.

Methodology

Materials

The materials used in this study if no others have mentioned pharma-ceutical grade standards. Materials for the manufacture of test preparations including Kojic acid, Oxybenzone, Octyl dimethyl PABA, BCD (p.a), stearic acid, Cetyl alcohol, Span 80, Tween 80, and 70% sobitol solution. While material for inhibition of tyrosinase activity test is the Mushroom tyrosinase, L-Tyrosine, $NaH_2PO_4.2H_2O$ (p.a) and Na_2HPO4 (p.a).

Instrument

Double beam UV-VIS Spectrophoto-meter, Fourier Transform Infrared Spectrophotometer, Micropipet various volumes, Pumpkin measure, Research Hanson dissolution testing, thermo-meters, pH meter CG SCHOOT Glass 842 Mainz, Milipore Membrane Filters, Tue diffusion, Analytical scales, Vortex, light microscopy, Electrothermal Melting Point Apparatus.

Preparations:

Vanishing cream base

14% stearic acid, 2% cetyl alcohol, Methyl paraben 0.1%, Propyl paraben 0.05%, 0.5% Span 80, 4.5% Tween 80, 3% sorbitol solution 70%, Aqua ad 100%.

Table 1 Formulation

	Percentage (% b/b)				
Component	F ₁ (control)	F ₂	F ₃	F4	
Kojic acid	1	1	1	1	
β-cyclodextrin	-	4	_	4	
Oxybenzone	_	_	3	3	
Octil dimethyl PARA	_	_	7	7	
Vanishing cream base(F ₀)	Ad 100	Ad 100	Ad 100	Ad 100	

Characteristics Determination of the cream preparation:

Determination of the physical quality of cream preparations include organoleptic (shape, colour, odour, texture), pH and spreading-ability.

Determination of the spreading-ability:

Determination of cream spreading-ability was performed using a pair of glass plate (20 X 20 cm). The cream preparation (1 gram) was put in the middle of the first glass plate that given the scale. Then put the second glass plate on the first glass plate and measured the diameter of cream spreading. After that put ballast on the second glass plate then measured the diameter spreading-ability of the cream. The weight of ballast that put on the second plate was increased until spreading-ability of the cream was constant.

Determination of tyrosinase activity inhibition Preparation of a solution of the reaction components.

The solution should be prepared for the implementation of this study was 0.1 M phosphate buffer pH 6.5, solution of tyrosinase (5370 units / mg solid in 0.1 M phosphate buffer pH 6.5 to 100.0 ml volume), a solution of 5.52 mM L-tyrosine, and 30% TCA.

Preparation of test sample solution.

The cream (around 3 grams) was put in the diffusion cell then covers with the Millipore membrane which was impregnated with isopropyl-myristate as modified lipid membrane. Then the preparation of cream in diffusion cell was put into the penetration chamber contain 500 ml of phosphate buffer pH 6.5 \pm 0.05 at 37 \pm 0.5°C as diffusion medium, and then the paddle was stirred 100 rpm. The sample solution around 3 ml was collected at 6 hours after it penetrated.

Determination of the dopakrom maximum wavelength

L-tyrosine solution of 5.52 mM has taken a number of prepared and then added 0.5 ml 3 ml of buffer solution pH 6.5 \pm 0.05. Then the mixture add with 1.0 ml of tyrosinase and oxygenated for 5 minutes. The mixture was incubated for 15 minutes at 26 \pm 0.5 °C. Then added 0.5 ml of 30% TCA. The solution is then inserted into cuvet and placed on the sample position in the spectro-photometer. Used as a blank solution 0.1 M phosphate buffer pH 6.5. Then do the reading of absorbance values from a wavelength of 400 nm to 500 nm, and the selected wavelength that gives the greatest absorption.

Inhibition Tyrosinase Activity Determination

L-tyrosine solution 5.52 mM added with 0.5 ml 3 ml of sample solution. Then the mixture added with 1.0 ml of tyrosinase and oxygenated for 5 minutes. The mixture was incubated for 15 minutes at 26 ± 0.5 °C. Then add 0.5 ml of TCA 30% and observed the absorbance at dopacrom maximum wavelength. Inhibition tyrosinase activity is the percent inhibition values obtained from the later absorbance value calculated by the equation:

- A = Absorbance at the maximum with the skin lightening
- B = Absorbance at maximum λ without the skin lightening

Results and Discussions

Characteristics Test Results

Based on the results of organoleptic test, all formulas have the same smell and texture. Meanwhile, in terms of different colours. Formula that contains sunscreen cream white yellowish colours, while others are white.

Table 2. Organoleptic test results

Formula	Replication	Organoleptic			
		Color	Odor	Texture	
	1		-	Semisolid,	
$\mathbf{F_1}$	2	White			
	3	white		smooth	
F ₂	1	XX71-04	White -	Semisolid, smooth	
	2				
	3	white			
\mathbf{F}_3	1				
	2	White yellowish		Semisolid,	
	3	yenowish		smooth	
\mathbf{F}_4	1	White yellowish	-	Semisolid, smooth	

Cream Type

The results of cream type test using water-soluble reagents (metylene blue) showed that the type of cream is O/W.

The pH Test Result

Based on the pH results in table 3, it was known that all formulas in the skin pH range (4 - 6.8).

Table 3. The pH test result

Formula	*pH	% CV
F_1	$4,46 \pm 0,01$	0,34
F_2	$4,24 \pm 0,03$	0,68
F_3	$4,29 \pm 0,03$	0,70
F_4	$4,04 \pm 0,02$	0,62

^{*)} The average obtained from 3 replicates it, each determined 3 times at different places.

Spreading-ability

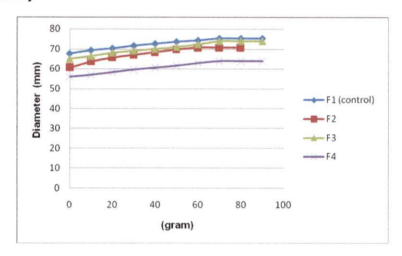


Figure 1. Profile of spreading-ability kojic acid cream on various formulas

Table 4. Sreading-ability result

	Replication	Dispersability (mm/g)	Average ± SD	
	1	0,1179		
F ₁ (Cantral)	2.	0,1083	0,1067±0,01	
	3	0,0940		
	1	0,1786		
F ₂	2	0,1393	0,1595 ± 0,02	
	3	0,1607		
	1	0,1083		
F ₃	2	0,1369	0,1226 ± 0,01	
	3	0,1226		
F4	1	0,1179		
	2	0,0964	0,1155 ± 0,02	
	3	0,1321		

Based on the results of one way ANOVA test, the addition of BCD (F_2) decreased its spreading-capacity, as well as addition of BCD on combination kojic-sunscreen (F_4)

Table 5. Sreading-capacity results

	Regli cation	Diameter at 60 g (mm)	Average ± SD
	1	75	
F ₁ (Control)	2	74	74,33 ± 0,58
3. 40.	3	74	
	1	70	
F ₂	2	71	70,67 ± 0,58
	3	71	1
	1	73	
F5	2	74	74,00 ± 1,00
	3	75	
F ₄	1	64	
	2	62	63,00 ± 1,00
	3	63	

Test Rresults Activity Inhibition of Tyrosinase

Table 6. Inhibition percent tyrosinase activity

Formula	% Inhibition Replikation		Average (%)±SD	CV (%)	
	I	II	III		
\mathbf{F}_1	61,73	62,10	63,07	62,3 ± 0,69	1,11
\mathbf{F}_2	28,39	28,40	29,28	$28,69 \pm 0,51$	1,78
F_3	60,50	60,65	62,26	$61,14 \pm 0,98$	1,60
F ₄	23,69	23,33	24,52	$23,63 \pm 0,61$	2,56

Based on one way ANOVA test, known that the addition of BCD decreased inhibition percent compared to the control constraints. Nevertheless the addition of sunscreen alone does not affect the inhibition percent of tyrosinase activity.

Addition of BCD on combination kojic-sunscreen preparations showed the same phenomena such as decreased its inhibition percent of tyrosinase activity.

Decreasing of inhibition percent of tyrosinase activity after addition of BCD alone or BCD-sunscreen was might be caused by decreasing of kojic acid penetration. It was probably caused by some of the things that the release process of kojic acid from base and in the process of penetration through the membrane. Before reach basal membrane where the kojic acid site of action, kojic molecule should released from base further more penetrate through the skin.

From spreading-ability test was known that the addition of BCD alone and also BCD on combination with sunscreen preparations reduced spreading-capacity, it means increasing viscosity. Increasing viscosity will decrease the mobility of active ingredient molecules that caused barriers against the release of kojic acid.

After the release process, kojic acid has to penetrate through trans-epidermal route through diffusion mechanism and trans-appendageal. One of the factors affecting the diffusion of the active ingredient is a molecule characteristic, which is the size of molecules. In the research on the use of drug fendeline with BCD, showed that the diffusion through a semi-permeable membrane decreased which was caused by increasing its molecule size

(Duchene et al, 1986). In this case decreasing kojic acid penetration by BCD addition might be caused by increasing of molecules the size of kojic acid. There for it expected would inhibit its penetration to the systemic, and make it survive longer in basal membrane, which is synthesis of melanin site. While the addition of BCD on combination kojic-sunscreen (F_4) reducing inhibition tyrosinase enzyme activity significantly compared with the control, as well as formula with the addition of a single BCD (F_2). This is probably due to the interaction between these materials so that the amount of kojic acid penetrated decreased and tyrosinase activity in vitro also declined.

Conclusion

Based on research results can be concluded that:

- 1. Combination of sunscreen oxybenzone and octyl dimethyl PABA (3:7) % w/w in the preparation of the skin lightening 1% kojic acid in a vanishing cream base does not affect the penetration kojic acid based on inhibition of tyrosinase enzyme activity in invitro.
- 2. β-cyclodextrin decrease penetration kojic acid based on inhibition of tyrosinase enzyme activity in *invitro*.
- 3. The existence of β -cyclodextrin and combination sunscreen oxybenzone and octyl dimethyl PABA (3:7)% w / w decrease penetration kojic acid based on inhibition of tyrosinase enzyme activity in *invitro*.

Acknowledgement

This study was supported financially by Project Grant of Faculty of Pharmacy Airlangga University.

References

- Duchene ,D., Vaution., & Glomot., 1986. Cyclodextrins, Their Value in Pharmaceutical Technology, France: Chatenay Malabry.
- Hatae, S & Kazuo, N., 1988. Whitening Cosmetic, United States patent.
- Manosroi., Wongtrakul ., Midorikawa., Hanyu., Yuasa., Sugawara., Sakai., Abe, 2005. The Entrapment of kojic oleate in bilayer vesicles, International Journal of Pharmaceutics.
- Nakayama, H., Ebihara, T., Satoh, N., dan Jinnai, T., 2000. Depigmentation Agents. In:Elsner, P., dan Maibach, H.I. (eds.). Cosmeceuticals and Active Cosmetics: Drug vs Cosmetics, Boca Raton: Taylor & Francis, p. 185-204.
- Wati, E.K., 2005. Penentuan Komposisi Optimum Bahan Tabir Surya Oksibenson-Oktildimetil PABA dalam Basis Vanishing Cream secara In vitro. Skripsi. Surabaya: Fakultas Farmasi Universitas Airlangga.
- Zulkarnain I, 2003. Kosmetika Pemutih Kulit dan Permasalahannya. Berkala Ilmu Penyakit Kulit dan Kelamin, Vol. 15, No 1, April 2003, p 47-53.