





6^{TH} ASIA PACIFIC INTERNATIONAL CONGRESS OF ANATOMY (6^{TH} APICA)

&

 13^{TH} NATIONAL CONGRESS OF INDONESIAN ANATOMIST ASSOCIATION (13^{TH} PIN-PAAI)

Proceeding Book

THE FUTURE OF ANATOMY

Clinical Anatomy Biomolecular and Cellular Anatomy Anatomy in Radiology and Imaging



GrahaBIK-IPTEKDOK Faculty of Medicine of Airlangga University Surabaya, 22nd-23rd July 2011 Indonesia

PROGRAM DAY-2

July 23rd, 2011

07.00-08.00 am	VENUE: GRAHABIK-IPTEKDOK (GRABIK) Re-registration				
07.00-08.00 am	Re-registration				
08 00-08 20 am	Re-registration				
08 00-08 20 am	Venue: Grabik 2 ^{au} floor				
UA UU-UA /U am	Moderator: Prof. H. Bambang Rahino S, dr.				
00.00 00.20	Keynote speaker I: Prof. H. Ari Gunawan, dr., MS., PhD (Indonesia). (KS II.1)				
08 20.08 40 am	"The Role of Anatomy in Supporting Medical Sciences" Koungte speaker II: Prof Madya, Dr Srijit Das (Malaysia) (KS II II)				
08.20-00.40 am	"Active Research in Anatomy : Do We Really Care for the Needy"				
	Venue: GRABIK 2nd floor				
	Moderator: Prof. Jeanne A Pawitan, dr., PhD				
08.40-09.00 am	Keynote Speaker III: Dr. Parker B. Antin (AAA,USA)(KS II.III)				
09.00-09.20 am	Keynote Speaker IV: Assc/ Prof. Christopher Bri	ggs, PhD (ANZACA, Australia)(KS II.IV)			
09.20-09.40 am	Keynote Speaker V: Djoko Santoso, dr., PhD., Sp	b.PD, K-GH (Indonesia) (KS II.V)			
09.40-10.00 am	Keynote Speaker VI : Prof. Dr. Teddy Untoseno,	dr., Sp.A (K)., Sp.JP (Indonesia) (KS II.VI			
10.00-10.30 am	Discussion				
10.30-10.50 am	Coffee break				
and the second second	V	Wanney BK Anotomi			
And And	Plenary session IV: Cellular & Biomolecular	Plenary session V: Clinical anatomy			
A Constant	Anatomy (Moderator: Prof. Dr. Yanwirasti, dr.)	(Moderator: Prof. Dr. Nancy M. Rehatta			
A. L.	Thatomy (Housing) This Die Tallinash, aly	SpAn K.IC)			
10.50-11.10 am	Speaker XIII: Prof. In-Sun Park (Chairman of	Speaker XVII: Prof. Dr. Doddy M. Soebad			
	KAS) (PSIV.I)	dr, Sp.U (Indonesia) (PSV.III)			
11.10-11.30 am	Speaker XIV: M.H. Nasr-Esfahani, B.Sc, PhD	SpeakerXVIII:Prof.PasukMahakkanukrauh			
and the second second	(Iran) (PSIV.II)	(Thailand) (PSV.I)			
11.30-11.50 am	Speaker XV: Assc/ Prof. Heidari M. Hassan,	Speaker XIX: Prof. Chang-Seok Oh, MD, I			
	PhD (Iran) (PSIV.III)	(Korea) (PSV.IV)			
11.50-12.10 am	Speaker XVI: Prof. Dr. Gayatri Rath (India)	Discussion			
12 10 12 10	(PSIV.IV)				
12.10-12.40 am	Discussion				
	Venue: Grabik 2 nd floor				
Sec. Sec.	Lunch & poster presentation (even numbers)				
12.40-02.00 pm	Jury : Teddy H. Wardhana, dr., Sp.OT, dr. Ni W	ajan T, dr., MS, PA, Prof. Dr. Nasronuddin.			
	Sp.PD., K-PTI., FINASIM, Prof. Pasuk Mahakka	nukrauh, Assc/ Prof. Heidari M. Hassan, P			
	Visiting Prof. Yoshiyuki Tohno. Prof. Purnomo S,	dr, MS, Dr., Dr. Kumkum Rana			
	Note: Presenter for poster competition must stand beside his/her poster from 01.00-01.45 pm				
	jury to be able to mark (Q&A)				
13 12-	Cmall warm masting				
- Martin mine in	Moderator : Prof. Dr. Nancy M. Rehatta, dr., SpAn K.IC				
2.13	Agenda : Discussion of potential networking in Reasearch and Educational Training				
1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1	Participant : University representative. Dean and	Vice Dean III of Medical Faculty Airlan			
Star and	University, Head of Department of Anatomy and Histology, Senior Lecturers of Department				
and the second	Anatomy and Histology FMAU, Head of UPPM	FMAU, Invited guests, APICA, and ANZA			
	Representative members				
	Venue : VIP Room GRABIK Lt 2.				

	Parallel session II Oral presentation for competition (@ 10 mins presentation, 5 mins Q&A)				IN	Venue: Ruang Praktikum Anatomi		Venue: Ruang Sidang Anatomi	
IV) II.VI)	Venue: GRABIK 2nd floor Moderator: dr. Ni Wajan T, dr., MS, PA Jury : Prof.H. Bambang Rahino S, dr., Prof. Jong Eun Lee	Venue: RK Anatomi Moderator: Teddy H. Wardhana, dr., Sp.OT Jury : Prof. Dr. Teddy Ontoseno, dr., Sp.A (K)., Sp.JP, Prof. Changman Zhou	Venue: RK Histologi Moderator: Prof. Dr. Bambang Sektiari L, drh., DEA Jury:Dr., Dra. Toetiek Koesbardiati, Prof. Fedik A. Rantam, drh., M.Kes, PhD	Venue: R Khusus Moderato Ferdiansy dr., SP.OT Jury : Pro Sudjono Aswin, dr PhD, Prof Dr. Abdul Hafid Bajamal, dr., Sp.BS	r: ah, Γ f. 	Moderator: S drg., MS, I Anita Tuli Forensic Anthropolog Maciej Hennet MSc (summa laude), PhD, I FAIBiol (Austra Facial Reconnstruct Demonstration Myrtati DA Dra.,MA., Ph (Indonesia)	Susy, Prof. Sy Derg, cum DSc., alia) ion on , D	PIC: Prof. Rio Sofwanhadi, dr., PA(K) APICA organization meeting Moderator: Prof. Kyung Ah Park Prof Joghataei, Prof Yun-Qing Li, Prof. Park (representative member from KAA), Dr Abdurrachman, APICA members	
Soebach.	Participant OB17- OB20	Participant OB21,OC1- OC3	Participant OC4-OC7	Participa OC8-OC	int 11				
NO. P	Parallel session III							& A)	
	Venue: GRABIK 2 nd floor Moderator : Yan Efrata S, dr.,Sp.BTKV Jury: Prof P. Gopalakhrisnakone MBBS, PhD., FAMS., DSc, Prof. Dr. Yanwirasti, dr		Venue: RK Anatomi Moderator : N Tomy Lesmana, dr., M SpB. KBD E Jury : Paulus Rahardjo, J dr., Sp.Rad (K), Myrtati DA,Dra.,MA.,Ph.D L d k		M	Venue: RK Histologi Moderator : Dr. Mustofa Helmi Pur Effendi, drh., MS Jur; Jury: Prof.Dr. Dar Gayatri Rath, Dr. Djoko Santoso, dr., PhD., Sp.PD, K-GH		enue: RK Khusus loderator : r. H. Bambang	
nuddin, c ssan, Ph .45 pm					Mu Effi Jur Ga Dja dr. K-			nomo, drh., MS y : Prof. Eryati rwin, dr., Parker B. Antin	
Airlans	Participant C	0C12-0C17	Particpant OC	8-OC23	Par OC	rticipant OC24- C26, OB22	Part OC2	icipant OC27- 29, OB23	
ANZAC	Coffee Break & Social time		2.13	Ve Jur wir	nue: RK Khusu y meeting for ora nning	s/ RS al & p	Anatomi oster award-		
and shares pm	Venue: Grabik 2 nd floor Award winning ceremony, PA & PA(K) Brevet award Closure						a na kuna an tan t		

ix

Biomolecular and Cellular Anatomy (PB23)

VIRGIN COCONUT OIL (VCO) AND POVIDONE IODINE ON THE DENSITY OF NAND WOUND CONTRACTION IN WOUND HEALING PROCESS IN RATS

Darwin, E¹, Jamil M²

scology, ²School of Nursing, Faculty of Medicine, University of Andalas, Padang, Indonesia Email : eryatidarwin@fk.unand.ac.id

ABSTRACT

Excition: Indonesia is an archipelago of biodiversity with around 30,000 plant species, and more have been used for treatment. Virgin Coconut Oil (VCO) derived from palm that grows along indonesian contains lauric, that induce wound healing process. Objective: To know the effect Oil (VCO) on wound healing on male and healthy Wistar rast. Methods and material: ded into three groups, there are group I (control), group II smeared with a VCO, and in group in povidone iodine for 10 days. After day 10, wound contraction was measured, and examination of wound tissue to determine the density of collagen. Results: The result shows grificant differences in wound contraction between group II and II compared with control group histopstologic appearance, dense collagen density seen in the group who get the VCO, medium agen density in the group of rats who received povidone iodine, and the density of collagen was cess in the control group. Conclusion: The results of this study show that the VCO play a role in process such as povidone iodine.

CO, wound, collagen

NTION

kin is a greater total mass than any other organ in the body. Its made up of outer layer, the s composed of stratified squamous keratinizing epithelium. The deeper layer, the dermis is vascularized and irregularly arranged fibroelastic connective tissue. The epidermis contains 5 bottom to top the layers are named: stratum basale, stratum spinosum, stratum granulosum, stratum stratum corneum. The bottom layer, the stratum basale, has cells that are shaped like columns. In cells divide and push already formed cells into higher layers. As the cells move into the higher flatten and eventually die. The top layer of the epidermis, the stratum corneum, is made of dead, flat that shed about every 2 weeks. There are three types of specialized cells in the epidermis, the produces pigment (melanin), the Langerhans' cell is the frontline defense of the immune system in the Merkel's cell's function is not clearly known

the back. The dermis is composed of three types of tissue that are present throughout - not in layers. If the back is composed of three types of tissue that are present throughout - not in layers. If tissue are: collagen, elastic tissue, and reticular fibers. The two layers of the dermis are the and reticular layers: the upper, papillary layer, contains a thin arrangement of collagen fibers, and the collar layer, is thicker and made of thick collagen fibers that are arranged parallel to the surface of the

The skin has many important functions, such as aesthetics and communication, sensation, regulation, storage and synthesis, contlor of evaporation and primarily for protection: an anatomical barrier from and damage between the internal and external environment in bodily defense. Once the protective broken, the normal (physiologic) process of wound healing is immediately set in motion and repair in injury. Wound healing, which requires the concerted effort of numerous cell types, involves cell on proliferation, differentiation, and apoptosis³.

chanisms of Wound Healing

The goal of wound-healing process is to repair the damaged skin. The classic model of wound healing is and into three or four sequential, yet overlapping phases: considered a (1) inflammatory, (2) proliferative maturation and remodeling. When tissue is first wounded, blood comes in contact with collagen, and blood platelets to begin secreting inflammatory factors. Platelets also express glycoproteins on their membranes that allow them to stick to one another and to aggregate, forming a mass. Fibrin and fibronectin link together and form a plug that traps proteins and particles and prevents further blood loss. This fibrinmectin plug is also the main structural support for the wound until collagen is deposited. Migratory cells use and as a matrix to crawl across, and platelets adhere to it and secrete factors. The clot is eventually lysed eplaced with granulation tissue and then later with collagen. Platelets, the cells present in the highest

301

numbers shortly after a wound occurs, release a number of things into the blood, including ECM p cytokines, including growth factors. Growth factors stimulate cells to speed their rate of division release other proinflammatory factors like serotonin, bradykinin, prostaglandins, prostacyclins, and histamine, which serve a number of purposes, including to increase cell proliferation and mena area and to cause blood vessels to become dilated and porous⁴.

In the inflammatory phase, bacteria and debris are phagocytosed and removed, and factors at that cause the migration and division of cells involved in the proliferative phase. The proliferative characterized by angiogenesis, collagen deposition, granulation tissue formation, epithelialization, a contraction. In angiogenesis, new blood vessels are formed by vascular endothelial cells. In fibre granulation tissue formation, fibroblasts grow and form a new, provisional extracellular matrix excreting collagen and fibronectin. Concurrently, re-epithelialization of the epidermis occurs, in what cells proliferate and 'crawl' atop the wound bed, providing cover for the new tissue⁵.

In contraction, the wound is made smaller by the action of myofibroblasts, which establish a prowound edges and contract themselves using a mechanism similar to that in smooth muscle cells maturation and remodeling phase, collagen is remodeled and realigned along tension lines and cells maturation longer needed are removed by apoptosis.

The final stage of wound healing is maturation and remodeling, in which the granulation is fibroplasia recede. In this phase, collagen is remodeled and realigned along tension lines and cells a longer needed are removed by apoptosis. During this stage, the epidermis regenerates by undergoing a of transient hypertrophy, while the provisional matrix is replaced by a dermal matrix of collagen and a low cellularity scar. Degradation of the collagen matrix is mediated by matrix metalloproteinases, secreted by the epidermal cells, fibroblasts, endothelial cells, and macrophages. Eventually, the replaced by a new functional tissue. However, this process is not only complex but fragile, and sace interruption or failure leading to the formation of chronic non-healing wounds. Factors which may complet this include diabetes, venous or arterial disease, old age, and infection^{6,7}.

The process of wound healing consists of integrated cellular and biochemical events least reestablishment of structural and functional integrity with regain of strength of injured tissue. Clinical often encounters non-healing, under-healing or over healing. Therefore the aim of treating a wound at the shorten the time required for healing or to minimize the undesired consequences. Plants and their extract immense potential for the management and treatment of wounds. Various herbal products have been us many countries in management and treatment of wounds such as Aloe vera, Azardica indica, Lantane en Linn, Tridax procumbens, Hydnocarpus wightiana, Chromolaena odorata, Helianthus annus Linn, Jan auriculatum, Ginkgo biloba, Cedrus deodara, Centella asiatica, VCO etc.^{8,9}.

Virgin Coconut Oil (VCO)

As a tropical country Indonesia is rich in various species of flora, from about forty thousand species flora that grows in the world, there are thirty thousand species have grown in Indonesia and 26% have cultivated and the rest still grows wild. About 940 species used as traditional medicine. One of that for coconut tree, it has the world's most extensive coconut tree field (3712 million ha), and there are is scattered the island of Java, Sumatra and Sulawesi¹⁰.

OBJECTIVES

To determine the effect of VCO on the wound healing process, we conducted the research on more were wounded. The wound were given with VCO and Povidone iodine. The wound contraction and the de of collagen in the wound were obvserved, and were compared with self healing wound.

METHODS AND MATERIAL

Three sets of experiments with three groups of male Wistar rats each consisting of 7 animals were for studying wound healing. This research was recommended by Research Ethic Comitee of Facar Medicine, Andalas University. Health Wistar rats, aged 8-12 weeks and 150-250 gram weight, were wou and evaluated for 10 days. Group 1 treated with NaCl 0.9% was the control, group 2 was treated with VCC group 3 was treated with Povidone iodine.

After the experimental period, the healing property of VCO was evaluated by monitoring the time a for contraction of the wound, collagen solubility pattern and histopathology of the tissue were also analyzed.

Data were taken based on the comparison between two experimental groups. Data analysis for parameters was calculated by ANOVA test. As for non-parametric data using Kruskal-Wallis test. The approach is based experimental research designs.



realuation of wound contraction, it was showed there were statistically significant difference real group (group I) and treated group (group II and III) with p value <0.05. While the comparison intraction between group II and group III showed no statistically significant difference (p>0.05)

resentation of wound contraction on control group rats which reciefed NaCl (group I) and group II vCO and group III treated with povidone iodine in day 10 after wounded

Group I (NaCl)		Group II	Group III
	92,85	97.07	98,76
	91.76	99,33	99,84
	97,50	100,00	97,66
	98,82	99,73	98,30
	96,47	100,00	98,30
	92,50	99,69	99,00
	92,50	100,00	98,42
1	94,62	99,40	98,67
	2,87	1,05	0,68
= 0,02		p(I):p(III)=0,011.	p(II): p(III) = 0.073

E I: Comparison of wound healing in control group rats given (a), group II which treated with VCO (b) roup III which treated with povidone iodine (c) at day-10.

Histolpathological appearance of wound healing process on control group 10 days after wounded, ed incomplete epithelization, abcess with many capillaries. The density of collagen were moderate to table 2: figure 2).

On the group I which treated with VCO, the histopathological treatment showed complete elization, there are few capillaries and few inflammatory cells. The density of collagen in all of animal d preparat are dens. While in group III which treated with povidone iodine the density of collagen are dens most animal wound preparat (figure 3).

e 2: The density of colagen of wound on control group rats which reciefed NaCl (group I) and group II and group II and group II and group II treated with povidine iodine in day 10 after wounded

No	Group I	Group II	Group III
I	I second a second	2	2
2	and the set of the second second	2	2
3	2	2	2
4	2	$\overline{2}$	$\overline{2}$
5	2	2	$\overline{2}$
6	2	2	See al the bar and a second
-	-	. .	,

sity of collagen on wound

: few collagen fiber predominantly with loss conective tissue

: moderat collagen fiber with moderate loss conective tissue

: dens collagen fiber with a few connective tissue

303



Figure 2: Histopathological appearance of wound healing process on control group a (40x) dat Collagen fiber and inflammatory cell (arrow)





Figure 3 : Histopathological appearance of wound healing process in treated group. Group II treated (a) and (b) and group III treated with povidone iodine (c)

DISCUSSION

Wound healing is a complex process that involves the organization of cells, chemical extracellular matrix to repair the tissue. In turn, the treatment of wounds tries to quickly close the obtain a functionally and esthetically satisfactory scar. To that end, it is indispensable to he understanding of the biological process involved in the healing of wounds and tissue regeneration¹¹.

One of fibroblasts' most important duties is the production of collagen. Collagen deposition is because it increases the strength of the wound; before it is laid down, the only thing holding the wound the fibrin-fibronectin clot, which does not provide much resistance to traumatic injury. Also, cells is inflammation, angiogenesis, and connective tissue construction attach to, grow and differentiate on the matrix laid down by fibroblasts. Type III collagen and fibronectin are generally beginning to be preasured amounts at somewhere between approximately 10 hours and 3 days, depending maining size. Their deposition peaks at one to three weeks. They are the predominating tensile substances unphase of maturation, in which they are replaced by the stronger type I collagen¹².

This study shows that day 10 after wounded is the final stage of wound healing, there a remodeling phase. During this phase, collagen is deposited by fibroblasts and formed into an organized Initially, the collagen strands laid down in the wound are thin and run parallel to the wound surface remodeling, however, collagen production increases. At the same time, some destruction of the original making room for the formation of new collagen, which is thicker and tends to be oriented along the lines within the wound.



• 4: Histopathologis appearant of collagen fiber on wound process healing in rat, which treated with VCO,

The anti-inflammatory property and the presence of *Lauric acid* of VCO is in the early synthesis of in fibers by mimicking. Farmacological activity of *Lauric acid* in VCO increase in blood pantioxidant, membrane stabilizing, improvement in cognition and pro-healing. Its can also promote relation without altering wound contraction. The other activity of *lauric acid* is to lysis of microbial rane that inactivate various microba

The use of VCO as phyto-medicines for wound healing are not only cheap and affordable but are also redly safe as hyper sensitive reactions are rarely encountered with the use of these agents. These natural s induce healing and regeneration of the lost tissue by multiple mechanisms. However, there is a need for fic validation, standardization and safety evaluation of plants of the traditional medicine before these the recommended for healing of the wounds

CLUSION

Wound is defined simply as the disruption of the cellular and anatomic continuity of a tissue. Wound reproduced by physical, chemical, thermal, microbial or immunological insult to the tissue. VCO is cheap, and natural active agent that can induce wound healing by stimulate cellular and colagen fiber. There is advance research to use VCO for treatment of wound healing.

RENCES

Cormack DH, Essential Histology. 2nd ed. LW &W, 2001

Inqueira LC. Basic Histology.9th ed. Appleton and Lange, 2007

Kana R, Prawez S, Verma PK Pankaj NK. Medicinal Plants and their Role in Wound Healing. VetScan, 2008.3(1): 1-7.

Filer BJ et al. Transforming Growth factor-B and wound healing. Focus on Basic Science. Perspectives execular surgery and endovascular therapy :2006 : 55-56.

Gelitzer R and Goebeler M.. Chemokines in cuteneus wound healing. Journal of Leukocyte Biology. 203; 513-519.

ello YM and Phillips TJ. Recent Advances in Wound Healing. JAMA. 2000; 283:716-718.

Seanes SR, Dang C, Soo C and Ting K. The phases of cutaneous wound healing. Expert Reviews in Indecular Medicine: 2003.

Reguna L. Efficacy of Butea monosperma on dermal wound healing in rats. Int J Biochem Cell Biol. 1998; 566–573

Scalan P.C. The role of honey in the management of wounds. Journal of Wound Care. 1999: 8 (8):415-419 Scalar B dan Prayogo S Membuat VCO Berkualitas Tinggi. Penebar Swadaya :2006.

Stegelmenn RF and Evans MC. Wound Healing : An Overview of Acute, Fibrotic and Delayed Healing. Fontiers in Bioscience 9 : 2004, 283-289.

Broughton G 2nd, Janis JE, Attinger CE. The basic science of wound healing: retraction. Surg Clin North an 2006; 77:509-528.

305



THE EFFECT OF VIRGIN COCONUT OIL (VCO) AND POVIDONE IODINE ON THE DENSITY OF COLLAGEN AND WOUND CONTRACTION IN WOUND HEALING PROCESS IN RATS

<u>Darwin Eryati</u>* and Mohammad Jamil **. * Departement Histology and **School of Nursing, Faculty of Medicine, University of Andalas. Padang, Indonesia

Background

The skin has many important functions, such as aesthetics and communication, sensation, regulation, excretion, storage and synthesis, control of evaporation and primarily for protection. Once the protective barrier is broken, the physiologic process of wound healing is immediately set in motion and repair itself after injury, which requires the concerted effort of numerous cell types, involves cell migration, proliferation, differentiation, and apoptosis. Clinically, one often encounters non-healing, underhealing or over healing. Therefore the aim of treating a wound is to either shorten the time required for healing or to minimize the undesired consequences.Plants and their extracts have immense potential for the management and treatment of wounds. Various herbal products have been used by many countries in management and treatment of wounds such as *Aloe vera, Azardica indica, Lantana camara* Linn, *Tridax procumbens, Jasminum auriculatum, Ginkgo biloba, VCO* etc. To determine the effect of VCO on the wound healing process, the wound contraction and the density of collagen in the experimental animal wound were obvserved.

Methods

Three groups of male Wistar rats each consisting of 7 health Wistar rats, aged 8-12 weeks and 150-250 gram weight, were wounded and evaluated for 10 days. Group 1 treated with NaCl 0.9% was the control, group 2 was treated with VCO, and group 3 was treated with Povidone iodine. After the experimental period, the healing property of VCO was evaluated by monitoring the time taken for contraction of the wound, collagen solubility pattern and histopathology of the tissue were also analyzed.

Results:

In evaluation of wound contraction, it was showed there were statistically significant difference between control group (group I) and treated group (group II and III) with p value <0.05. While the comparison of wound contraction between group II and group III showed no statistically significant difference (p>0.05) (table 1).On the group I which

treated with VCO, the histopathological treatment showed complete epithelization, there are few capillaries and few inflammatory cells. The density of collagen in all of animal wound preparat are dens. While in group III which treated with povidone iodine the density of collagen are dens in almost animal wound preparat (Figure:1)

This study shows that during the skin-remodeling phase collagen is deposited by fibroblasts and formed into an organized network. Initially, the collagen strands laid down in the wound are thin and run parallel to the wound surface. At the same time, some destruction of the original collagen occurs, making room for the formation of new collagen, which is thicker and tends to be oriented along the lines of stress within the wound.

Table 1: Percentage of wound cintraction on control group(Group I), treated group (group II) and treated with povidine iodine (group III) 10 days after wounded

No.	Group I (NaCl)	Group II	Group III		
1	92,85	97,07	98,76		
2	2 91,76 99,33		99,84		
3	97,50	100,00	97,66 98,30		
4	98,82	99,73			
5 96,47		6,47 100,00			
6	6 92,50 99,6		9 99,00		
7	92,50	92,50 100,00			
Mean	94,62±2,87	99,40±1,05	98,67±0,68		



Figure 1: Histopathological appearance of wound healing process in control group (a) and treated with VCO (a) and (b) treated with povidone iodine (c). Arrow: collagen fiber

Conclusion

Wound is defined simply as the disruption of the cellular and anatomic continuity of a tissue. Wound may be produced by physical, chemical, thermal, microbial or immunological insult to the tissue. VCO is cheap, safe, and natural active agent that can induce wound healing by stimulate cellular and colagen fiber. There is need advance research to use VCO for treatment of wound healing





ASIA-PACIFIC INTERNATIONAL CONGRESS OF ANATOMY (APICA) INDONESIAN ANATOMIST ASSOCIATION (PERHIMPUNAN AHLI ANATOMI INDONESIA-PAAI)

This certificate is presented to:

Prop. Ervalt Darwin, dr

As Speaker/Participant/Moderator/Jury/Editor boards/Committee in 6th Asia-Pacific International Congress of Anatomy 13th National Congress of Indonesian Anatomist Association

(6th APICA & 13th PIN-PAAI)

22nd-23rd of July, 2011 Faculty of Medicine Airlangga University Surabaya, Indonesia

Dr. Abdurachman, dr., M.Kes, PA(K)