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Table of contents

Volume 217

2019

[◀ Previous issue](#) [Next issue ▶](#)

The 12th Congress of Indonesian Soc. for Biochemistry and Molecular Biology in Conjunction With The 2nd Int. Conf. "Collaboration Seminar of Chemistry and Industry (CoSCI)" and AnMicro Workshop 11-12 October 2018, Universitas Airlangga, Indonesia

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Preface

OPEN ACCESS 011001

The 12th Congress of Indonesian Society for Biochemistry and Molecular Biology in Conjunction With The 2nd International Conference "Collaboration Seminar of Chemistry and Industry (CoSCI)" and AnMicro Workshop

[+ Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS 011002

Committee

[+ Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS 011003

Conference Photographs

[+ Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS 011004

Peer review statement

[+ Open abstract](#) [View article](#) [PDF](#)

Papers

Chemistry

OPEN ACCESS

012001

Facile Sol-Gel Synthesis of Calcium Phosphates: Influence of Ca/P Ratio and Calcination Temperature

A J Permana, A T Utami, U S Handajani and H Setyawati

[+ Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012002

Determination of Brønsted Acid Sites In Porous Aluminosilicate Solid Catalysts Using Volumetric And Potentiometric Titration Method

A Purwaningsih, A N Kristanti, D Z Mardho, D W Saraswati, N M Putri, N H Saputri and Hartati

[+ Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012003

Carbon Paste Electrode Modified Imprinted Zeolite as a Selective Sensor for Creatine Analysis by Potentiometry

A. Athiroh, T Fadillah, D F Damayanti, A A Widati, A Abdulloh and M Khasanah

[+ Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012004

Voltammetric Study of Ascorbic Acid Using Polymelamine/Gold Nanoparticle Modified Carbon Paste Electrode

A N Farida, E Fitriany, A Baktir, F Kurniawan and M Harsini

[+ Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012005

Synthesis of Silver Nanoparticles and the Development in Analysis Method

H I Badi'ah, F Seede, G Supriyanto and A H Zaidan

[+ Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012006

Two Flavonoids From Stem Bark of *Casimiroa edulis* and Their Antidiabetic and Antioxidant Activities

K N W Tun, N S Aminah, A N Kristanti, R Ramadhan and Y Takaya

[+ Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012007

Graphene Oxide from Bagasse/Magnetite Composite: Preparation and Characterization

M Jannatin, G Supriyanto, Abdulloh, W A W Ibrahim and N K Rukman

[+ Open abstract](#) [View article](#) [PDF](#)

-
- OPEN ACCESS** 012008
GO-Fe₃O₄ Nanocomposite from coconut shell: Synthesis and characterization
N K Rukman, M Jannatin, G Supriyanto, M Z Fahmi and W A W Ibrahim
[+](#) Open abstract [View article](#) [PDF](#)
-
- OPEN ACCESS** 012009
First Order Kinetics of Salicylamide Release from κ -Carrageenan Hard Shell Capsules in Comparison with Gelatin
P Pudjiastuti, E Hendradi, S Wafiroh, H Darmokoesoemo, M A R D Fauzi, L Nahar and S D Sarker
[+](#) Open abstract [View article](#) [PDF](#)
-
- OPEN ACCESS** 012010
Chromanone Acid Derivatives from the Stem Bark of *Calophyllum incrassatum*
U Hasanah, T S Tjahjandarie and M Tanjung
[+](#) Open abstract [View article](#) [PDF](#)
-
- OPEN ACCESS** 012011
Preparation Hydrophobic Fabric Coated by TiO₂ and Hexadecyltrimethoxysilane
U S Handajani, A A Widati and I N Yusbainika
[+](#) Open abstract [View article](#) [PDF](#)
-
- OPEN ACCESS** 012012
Kecombrang (*Etligeria elatior*) Leaves Ethanol Extract Effect to Lens and Erythrocyte Aldose Reductase Activity in Wistar strain white rats (*Rattus norvegicus*) Streptozotocin induced
S Handayani, H Notopuro and G I Prabowo
[+](#) Open abstract [View article](#) [PDF](#)
-
- OPEN ACCESS** 012013
Adsorption of Isopropyl Alcohol (IPA) in Water Using Activated Bentonite
A Abdulloh, G Supriyanto and O W Ningsih
[+](#) Open abstract [View article](#) [PDF](#)
-
- OPEN ACCESS** 012014
Production of Nanopropolis Using High Pressure Ball Mill Homogenizer
D Hamdi, A Wijanarko, H Hermansyah, S C Asih and M Sahlan
[+](#) Open abstract [View article](#) [PDF](#)
-
- OPEN ACCESS** 012015
Synthesis of ZnO-TiO₂/Chitosan Nanorods By Using Precipitation Methods and Studying Their Structures and Optics Properties at Different Precursor Molar Compositions
Y Rilda, D Damara, Syukri, Y E Putri, Refinel and A Agustien

[+ Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012016

Phytochemical Screening and Antioxidant Activity of Ethanol Extract of Leilem (*Clerodendrum minahassae* Teijsm. & Binn) as an Antihyperlipidemic and Antiatherosclerotic Agent

C F Kairupan, F R Mantiri and R R H Rumende

[+ Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012017

Concentration of Some Metals in Water and Soil Samples at Some Locations near the Hotmud Flow at Porong Disaster Area, Sidoarjo, East Java, Indonesia.

A Wiryawan, R Suntari, Z Kusuma and Syekhfani

[+ Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012018

The Effect of Roselle (*Hibiscus sabdariffa* Linn) Flower Extract To The SGPT Activity In Male Wistar Rats (*Rattus Norvegicus*) Induced By High Dose Paracetamol

D Halim, E J Sihning and Tehupuring

[+ Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012019

Antioxidant Exploration in Cardamom Rhizome Potential as a Functional Food Ingredient

H Winarsi, A Yuniaty and Warsinah

[+ Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012020

Effect of Gambir Catechin Isolate (*Uncaria Gambir* Roxb.) Against Rat Triacylglycerol Level (*Rattus novergicus*)

Y Alioes, R R Sukma and S L Sekar

[+ Open abstract](#) [View article](#) [PDF](#)

Biochemistry and Molecular Biology

OPEN ACCESS

012021

Exploration of Cellulolytic Microorganism as A Biocatalyst Candidate for Liquid Fertilizer Production

N Halimah, A Baktir and P Purkan

[+ Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012022

Antibody Titers in The Sheep which were Immunated Antigen of *Whole* Protein from Third Instar Larvae *Musca domestica*

B Ariantini, H Ratnani, E M Luqman and P Hastutiek

[+](#) [Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012023

Lemon (*Citrus limon*) Juice Has Antibacterial Potential against Diarrhea-Causing Pathogen

ER Ekawati and W Darmanto

[+](#) [Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012024

Genetic Relationship of *Hibiscus* spp. Based on DNA bands Using RAPD Technique

Hamidah and A Z Muhtadi

[+](#) [Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012025

Effect of *Sticophus hermanii* extract on fasting blood glucose and skeletal muscle glut4 on type 2 diabetes mellitus rats model

I Safitri, B Purwanto, L Rochyani, G I Prabowo and D Sukmaya

[+](#) [Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012026

Callus Induction and Bioactive Compounds from *Piper betle* L. var nigra

Junairiah, A Mahmuda, Y S W Manuhara, Ni'matuzahroh and L Sulistyorini

[+](#) [Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012027

Antimicrobial Activity of Ethanol Extract of *Abrus precatorius* L. Roots against Planktonic Cells and Biofilm of Urine and Blood Methicillin Sensitive *Staphylococcus aureus* (MSSA) Isolate

B Mutmainnah, Ni'matuzahroh and A Baktir

[+](#) [Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012028

Utilization of Rice Straw Hydrolysis Product of *Penicillium* sp. H9 as A Substrate of Biosurfactant Production by LII61 Hydrocarbonoclastic Bacteria

Ni'matuzahroh, S K Sari, N Trikurniadewi, A D Pusfita, I P Ningrum, S N M M Ibrahim, T Nurhariyati, Fatimah and T Surtiningsih

[+](#) [Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012029

Carbon and Nitrogen Sources for Lipase Production of *Micrococcus* sp. Isolated from Palm Oil Mill Effluent-Contaminated Soil

S. Sumarsih, S. Hadi, D.G.T. Andini and F.K. Nafsihana

[+](#) Open abstract [View article](#) [PDF](#)

OPEN ACCESS

012030

Cytotoxicity of Combination Chitosan with Different Molecular Weight and Ethanol Extracted *Aloe vera* using MTT Assay

Sularsih, Soetjipto and Retno Pudji Rahayu

[+](#) Open abstract [View article](#) [PDF](#)

OPEN ACCESS

012031

Hepatoprotective Effect of Gamma-mangostin for Amelioration of Impaired Liver Structure and Function in Streptozotocin-induced Diabetic Mice

S A Husen, D Winarni, Salamun, A N M Ansori, R J K Susilo and S Hayaza

[+](#) Open abstract [View article](#) [PDF](#)

OPEN ACCESS

012032

Utility of *Saccharomyces cerevisiae* As Probiotics to Induce Protease Production For Worms Feed Improvement

R Arissirajudin, S Hadi, Abdillah Safa and P Purkan

[+](#) Open abstract [View article](#) [PDF](#)

OPEN ACCESS

012033

Induction of Angiogenesis Process in Mandible Using *Anadara granosa* Shell Graft (Experimental Laboratory Study on *Rattus norvegicus*)

Widyastuti, M Rubianto and Soetjipto

[+](#) Open abstract [View article](#) [PDF](#)

OPEN ACCESS

012034

Dehalogenase enzyme activity of *Bacillus* sp. D1 isolated from pharmaceutical waste

K Primasari, D W Sawitri, R Fikri, N Trikurniadewi, Ni'matuzahroh and G Supriyanto

[+](#) Open abstract [View article](#) [PDF](#)

OPEN ACCESS

012035

The impact of conditioned medium of umbilical cord-derived mesenchymal stem cells toward apoptosis and proliferation of glioblastoma multiforme cells

Novi Silvia Hardiany, Yohana and Septelia Inawati Wanandi

[+](#) Open abstract [View article](#) [PDF](#)

OPEN ACCESS

012036

Utilization of Bromelain Enzyme from Pineapple Peel Waste on Mouthwash Formula Against *Streptococcus mutans*

H Rahmi, A Widayanti and A Hanif

[+ Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012037

Michaelis-Menten Parameters Characterization of Commercial Papain Enzyme "Paya"

Mathias Elsson, Anondho Wijanarko, Heri Hermansyah and Muhamad Sahlan

[+ Open abstract](#) [View article](#) [PDF](#)

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012038

The effect of cytoglobin gene inhibition on fibroblast keloid cells proliferation

S W A Jusman, F M Siregar, M Sadikin and N S Hardiany

[+ Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012039

Effect of IPTG Concentration on Recombinant Human Prethrombin-2 Expression in *Escherichia coli* BL21(DE3) ArcticExpress

S Silaban, S Gaffar, M Simorangkir, I P Maksum and T Subroto

[+ Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012040

Exploration of *Chlorella sp.* as antibacterial to *Aggregatibacter actinomycetemcomitans* biofilm

P F Christabel, M V Hernando, C A Sutanto and K Parisihni

[+ Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012041

The Influence of Ethanolic Root Extracts of *Ruellia tuberosa L.* on Pancreatic Protease Activity and MDA Level of Rats (*Rattus norvegicus*) Induced by MLD-STZ

A Roosdiana, Sutrisno, C Mahdi and A Safitri

[+ Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012042

The Effect of spirulina on Apoptosis (Stored Biology Materials) To Pregnant Rat Wistar in the Second Trimester Wich is Induced By IL-6

Y Rani, H Gondo and N K Indahsari

[+ Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012043

Revealing the important role of allosteric property in sucrose phosphate synthase from sugarcane with N-terminal domain deletion

W D Sawitri and B Sugiharto

[+ Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS 012044
Potential of marine chitinolytic *Bacillus* isolates as biocontrol agents of phytopathogenic fungi
E Kurniawan, S Panphon and M Leelakriangsak
[+](#) Open abstract [View article](#) [PDF](#)

OPEN ACCESS 012045
Identification of α -amylase gene by PCR and activity of thermostable α -amylase from thermophilic *Anoxybacillus thermarum* isolated from Remboken hot spring in Minahasa, Indonesia
F R Mantiri, R R H Rumende and S Sudewi
[+](#) Open abstract [View article](#) [PDF](#)

OPEN ACCESS 012046
Broccoli Extract (*Brassica oleracea*) Decrease Periarticular Malondialdehyde Level and Disease Activity Score in Rats (*Rattus norvegicus*) with Adjuvant Arthritis
S Prabowo
[+](#) Open abstract [View article](#) [PDF](#)

OPEN ACCESS 012047
Synthesis of Aldehyde-Silica Nanoparticle for Matrix Immobilization of Endo- β -1,4-D-xylanase
A A I Ratnadewi, S Trissa, Suwardiyanto, W Handayani, A B Santoso and Sudarko
[+](#) Open abstract [View article](#) [PDF](#)

Medicine

OPEN ACCESS 012048
Counselling and Screening of Hepatitis B Virus Infection In Dukuh Kupang Community, Dukuh Pakis District, Surabaya
C D K Wungu, S Khaerunnisa, I Humairah, L Lukitasari, E Qurnianingsih, G I Prabowo, Sudarno, R Handajani and Suhartati
[+](#) Open abstract [View article](#) [PDF](#)

OPEN ACCESS 012049
Correlation Between Oxidative Stress With Clinical Symptoms In Chronic Schizophrenic Patients In Psychiatric Unit of Dr Soetomo General Hospital Surabaya
G I Prabowo, M M Maramis, E Yulianti, A Zulaikha, Z B Syulthoni, C D K Wungu, H M Margono and R Handajani
[+](#) Open abstract [View article](#) [PDF](#)

OPEN ACCESS 012050

Antigenic Protein Profile of *Streptococcus mutans* Biofilm For Developing of Dental Caries and Periodontal Disease Risk Biomarker

M Ni'mah, I L Kriswandini and A Baktir

[+](#) [Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012051

Detection Of Hepatitis C Virus (Hcv) Infection And Its Genotype In Patients At Hepatology Outpatient Clinic, Dr Soetomo General Hospital, Surabaya.

R Handajani, C D K Wungu, I Humairah, G I Prabowo, U Cholili, M Amin, P B Setiawan and Soetjipto

[+](#) [Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012052

Endothelial Dysfunction Improvement Mechanism By Hyperbaric Oxygen In Sprague Dawley By High-Cholesterol Diet

H Setianingsih, Soetjipto, I K Sudiana and G Suryokusumo

[+](#) [Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012053

Correlation of Homocysteine Levels With Folate Acid, Cyanocobalamine, and Pyridoxine Serum Levels In Acute Infark Miocard Patients

D Pertiwi and R Yaswir

[+](#) [Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012054

Taurine Intakes Increase Superoxide Dismutase Activity in Knee Osteoarthritis

A A E W Saraswati, D Sunardi, A M T Lubis, F Heru and N Mudjihartini

[+](#) [Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012055

Association Between the Ratio of Omega-6/Omega-3 Fatty Acids Intake to Plasma Malondialdehyde Level in Patients with Knee Osteoarthritis

S R Angelia, N R M Manikam, A M T Lubis, C Siagian and N Mudjihartini

[+](#) [Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012056

Enhance of IL-22 expression in Oral Candidiasis Immunosupressed Model with *Acanthus ilicifolius* Extract Therapy

D Andriani and A F Pargaputri

[+](#) [Open abstract](#) [View article](#) [PDF](#)

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012057

Expression Of Runx2 And Osteoblast Cell On The Periodontal Of Diabetes Mellitus Wistar Rat With Diet Extract Lemuru Fish Oils Treatment

[+ Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012058

Stichopus hermanii stimulation to Runx2 expression as Periodontal Remodeling Biomarkers to accelerate Orthodontic Tooth Movement

N Prameswari and B Handayani

[+ Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012059

The Differences of Effectivness HBO 2,4 ATA Between 7 and 10 Days In Bone Remodelling of Tension Area of Orthodontic Tooth Movement

A Brahmanta, D Mulawarmanti, F Z Ramadhani and W Widowati

[+ Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012060

The Effect of Sticopus Hermanii-Hyperbaric Oxygen Therapy to Inflammatory Response of Diabetic Periodontitis

D Mulawarmanti, K Parisihni and Widyastuti

[+ Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012061

Identification of *Mycobacterium tuberculosis* Bacteria with TB Antigen MPT64 Rapid Test Against Patients with Suspect Pulmonary Tuberculosis in Lubuk Alung Pulmonary Hospital, Padang Pariaman

E Bahar and A E Putra

[+ Open abstract](#) [View article](#) [PDF](#)

OPEN ACCESS

012062

Hypoxia increased malondialdehyde from membrane damages is highly correlated to HIF-1 α but not to renin expression in rat kidney

A R Prijanti, F C Iswanti, F Ferdinal, S W A Jusman, R R Soegianto, S I Wanandi and M Sadikin

[+ Open abstract](#) [View article](#) [PDF](#)

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18

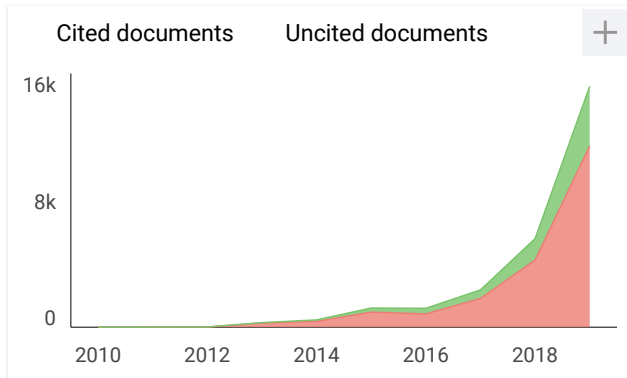
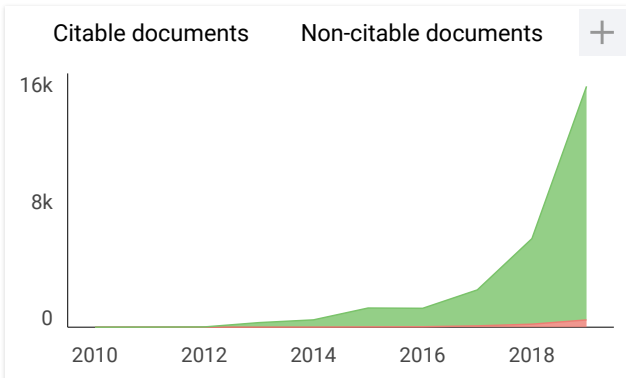
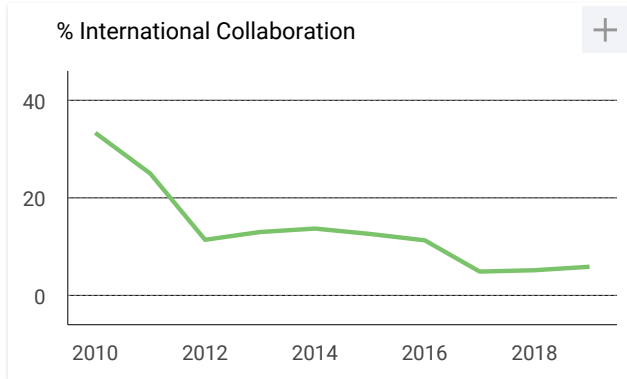
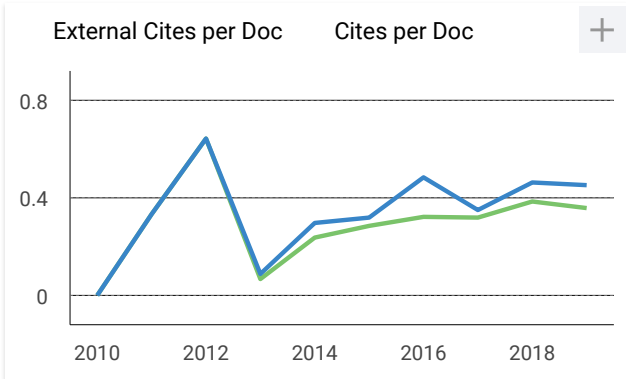
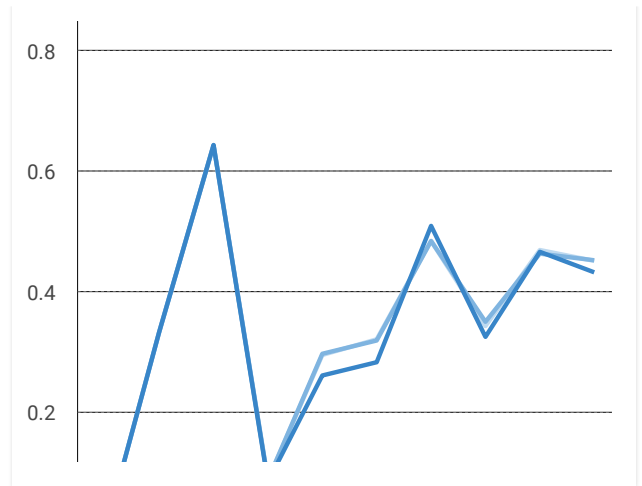
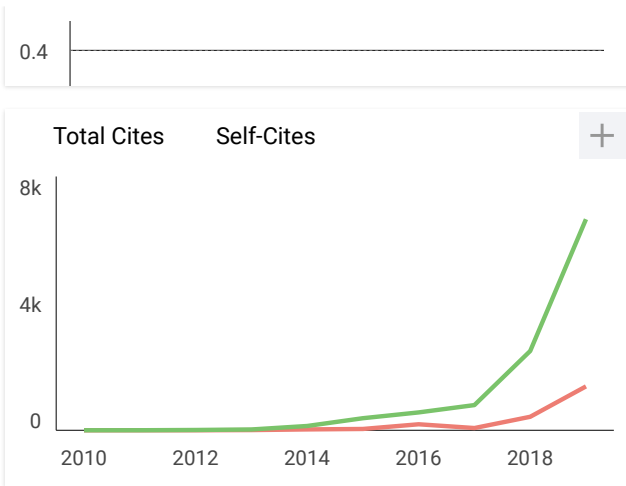
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Utilization of Rice Straw Hydrolysis Product of *Penicillium* sp. H9 as A Substrate of Biosurfactant Production by LII61 Hydrocarbonoclastic Bacteria

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Utilization of Rice Straw Hydrolysis Product of *Penicillium* sp. H9 as A Substrate of Biosurfactant Production by LII61 Hydrocarbonoclastic Bacteria

Ni'matuzahroh^{1*}, S K Sari¹, N Trikurniadewi¹, A D Pusfita¹, I P Ningrum¹, S N M M Ibrahim¹, T Nurhariyati¹, Fatimah¹, T Surtiningsih¹

¹ Department of Biology, Faculty of Science and Technology, Universitas Airlangga

*nimatuzahroh@fst.unair.ac.id

Abstract. This study aims to reveal the prospect of rice straw as a biosurfactant production substrate by hydrocarbonoclastic bacteria. Rice straw can be hydrolysed enzymatically into simple sugar by *Penicillium* sp. H9. Hydrolysis products are used as growth medium and biosurfactant production by LII61 bacteria. The concentration sugar from hydrolysis product was analyzed by Nelson method. In this study, molasses were used as a comparison substrate in biosurfactant production. The growth response of LII61 bacteria was observed by measuring the turbidity of the culture. Biosurfactant products were evaluated by measuring the emulsification activity (%) and surface tension (mN/m). Acquisition of sugar from rice straw hydrolysis product (RSHP) was 209.25 µg/mL. The optimum growth both of RSHP and molasses substrates were obtained on the 5th day of incubation with a culture turbidity value of OD_{λ650} 0.201 and OD_{λ650} 0.157 respectively. The lowest surface tension obtained in culture of RSHP was 48.85 mN/m. It was better than biosurfactant product on the molasses substrate on the 3rd day incubation. However, during the incubation time both substrates did not show emulsification activity. Biosurfactants produced have certain characteristics on variations in pH and temperature.

Keyword: Rice Straw, *Penicillium* sp. H9, Biosurfactan, LII61 Hydrocarbonoclastic Bacteria

1. Introduction

Indonesia is a tropical country with most of its terrain is the agricultural area. Agricultural yields in Indonesia always leave agricultural wastes that are accumulating in the environment. Rice straw is one of the agricultural wastes in Indonesia. According to the Central Statistics Agency of Indonesia [1], rice production in 2015 amounted to 75.40 million tons with rice straw production reaching 12-15 tons per hectare per harvest. Rice straw contains polysaccharides in the form of cellulose, hemicellulose, lignin, and pectin [2]. Lignocellulose content in rice straw consists of cellulose by 32%, hemicellulose 24%, and lignin 14% [3]. The high content of cellulose in rice straw has the potential to be transformed by microorganisms into materials that can be reused.

The use of residual agricultural products is a breakthrough in handling environmental organic waste. Organic material in the agricultural waste can be a source of nutrition for microorganisms [4]. The enzymatic specifications possessed by microbes will determine the conversion of ingredients into desired products [5]. Hydrolysis is the first step in utilizing agricultural waste. Hydrolysis product is obtained from enzymatic alteration of organic waste, in which enzymatic hydrolysis have the advantage such as no toxic products [6], no corrosion and higher sugar production than acid hydrolysis [7]. According to Pratama [8], *Penicillium* sp. H9 has a cellulolytic activity which is able to hydrolyze cellulose. *Penicillium* sp. H9 has the potential as an agent to hydrolyze rice straw into sugar as a result of hydrolysis (reducing sugars), these products can be used as carbon sources and substrates for biosurfactant production.

Biosurfactants are amphipathic compounds produced by microorganisms. These compounds can reduce surface tension and emulsify hydrocarbons. Biosurfactants have the potential to be applied in



the oil industry, such as oil tank washing and processing of oil sludge waste [9] and for the health aspect which can be used as an antimicrobial compound [10]. Previous research shows that LII61 bacteria are able to grow and produce biosurfactants in sugarcane (molasses) waste substrates [11]. Molasses contain simple sugars like glucose, sucrose, and fructose which support bacterial growth. However, the presence of molasses is increasingly scarce as demand in the food industry increases. Rice straw hydrolysis product (RSHP) is one of the alternative substrates for biosurfactant production by bacteria.

Ni'matuzahroh et al, [12] has succeeded in isolating potential bacteria which decompose hydrocarbons known as hydrocarbonoclastic bacteria. These bacteria also have the potential to produce biosurfactants [11]. One strain of hydrocarbonoclastic bacteria that has activities to produce biosurfactants is LII61. According to Pratiwi [13], LII61 produces lipase as well as biosurfactant enzymes so as to reduce the surface tension of the growth medium. This study aims to determine the use of RSHP of *Penicillium* sp. H9 as a substrate of biosurfactant production by hydrocarbonoclastic bacteria LII61. Optimization and characteristics of biosurfactants were also carried out in this study. LII61 biosurfactant products on RSHP compared with biosurfactant products on molasses substrate. Utilization of RSHP for biosurfactant substrate is an effort to reduce agricultural waste.

2. Experimental Method

2.1 Microorganism

The bacteria used in this study was *Micrococcus* sp. L II (61) obtained from Pegirian slaughterhouse (RPH) Surabaya East Java which is a collection from the Laboratory of Microbiology, Faculty Science and Technology, Airlangga University, Surabaya. Bacterial isolates were grown in Broth Nutrient media and incubated using a shaker incubator at room temperature for 24 hours. The bacteria used was 2% (v/v) of the total culture volume of 0.4 mL at OD_{650} 0.5.

2.2 Hydrolysis of rice straw by *Penicillium* sp. H9

Organic agricultural waste, rice straw, obtained from local farmers. The rice straw is mechanically delignified by the grinding process and chemically delignified using 1% NaOH for 1 hour at 100 °C. The pretreatment rice straw was enzymatically hydrolyzed by *Penicillium* sp. H9 for 6 days of incubation time. The results of hydrolysis was mechanically sterilized using a filter syringe and the sugar content was measured by the Somogy-Nelson method.

2.3 Screening of biosurfactant production by LII61 on RSHP substrate

Screening of biosurfactant production was carried out on Synthetic Mineral Water media (SMW) with the addition of 2% (v/v) RSHP. Media Synthetic mineral water used in this study is a modification from Pruthi and Cameotra (1997). The composition of synthetic mineral water are (g/L): $(NH_4)_2SO_4$ (3.0 g/L), NaCl (10 g/L), $MgSO_4 \cdot 7H_2O$ (0.2g/L), $CaCl_2$ (0.01g/L), $MnSO_4 \cdot H_2O$ (0.001 g/L), H_3BO_3 (0.001 g/L), $ZnSO_4 \cdot 7H_2O$ (0.001 g/L), $CuSO_4 \cdot 5H_2O$ (0.001 g/L), $CoCl_2 \cdot 6H_2O$ (0.005g/L) and $NaMoO_4 \cdot 2H_2O$ (0.001 g/L). SMWbuffer consists of (g/50mL): KH_2PO_4 (5 g), K_2HPO_4 (2.62047 g) and Fe (g / 50mL) Fe_3O_4 (0.0006 g). The total culture volume was 20 mL, microbial culture was incubated for 0, 1, 3, and 5 days in a rotary shaker 120 rpm at room temperature.

2.4 Optimization biosurfactant production by LII61

Optimization of biosurfactant production was executed with a variation of incubation time 0, 1, 3 and 5 days, incubation was carried out in a rotary shaker 120 rpm at room temperature. The media used are synthetic mineral water (SMW) with the addition of RSHP substrate as much as 1.5 mL of rice straw hydrolysis with the concentration 229 ppm. The total culture volume used was 20 mL.

2.5 Detection of biosurfactant product from LII61

Bacterial cultures that were incubated during the incubation time measured for the final pH value. Bacterial cultures were centrifuged at 9000 rpm for 15 minutes and the supernatant that contains biosurfactant measured surface tension and emulsification activity.

Surface tension was measured with tensiometer Du-Nuoy, Surface tension value was stated in mN/m or dyne/cm and distilled water was used for control in this measurement. Surface tension was measured with this formula.

$$r = r_o \frac{\theta}{\theta_o} \quad (1)$$

Which are r = surface tension of sample, r_o = surface tension of distilled water in t°C, θ = surface tension of surface sample that reads on the tool, and θ_o = surface tension of surface distilled water that reads on the tool

Emulsification activity was measured with 1 ml of supernatant from centrifugation. The centrifugation was carried out for 5 minutes with 3000 rpm. 1 ml of supernatant was added with 1 ml of kerosene. The mixture was compound with vortex for 2 minute, and the emulsification activity can be observed after 1 hour and 24 hours. Emulsification activity was observed by measuring the high of emulsy (cm) toward high liquid total (cm) multiplied 100% [14].

2.6 Extraction of biosurfactant from LII61

Crude biosurfactant was obtained with acid deposition. 1 Litre of supernatant from bacteria culture was incubated until 72 hours and centrifugation for 15 minutes with 6000 rpm. The supernatant from centrifugation was deposited with HCl 6N until the final pH was 2,0. Then, supernatant was incubated in the refrigerate for overnight and was centrifugated again to obtained the sediment of crude biosurfactant.

2.7 Characterization of crude biosurfactant extract from LII61 towards temperature and pH

Crude biosurfactant extract was characterized with CMC (Critical Micelle Concentration). CMC measurement was carried out by making various concentration from crude biosurfactant. The surface tension of various concentration of crude biosurfactant were measured with Tensiometer Du-Nuoy. The measurements were stopped if the critical point was reached, which is the value of surface tension was remained. The critical point was the CMC value of crude biosurfactant extract.

The measurement of stability from crude biosurfactant extract towards temperature change was obtained by making a various concentrations of crude biosurfactant extract at CMC value, and then the solutions were incubated to a various temperature (30 °C, 50 °C, and 70 °C) for 1 hour. The solutions were measurement for surface tension and emulsification activity with kerosene. The measurement of stability of crude biosurfactant extract towards pH change was obtained by making a various concentrations of crude biosurfactant extract at CMC value, and then the solutions were incubated to a various pH (4, 7, and 10) for 1 hour. The solutions were measurements for surface tension and emulsification activity with kerosene.

3. Results and Discussion

3.1 Hydrolysis of rice straw by *Penicillium sp. H9*

The lignocellulose content in organic waste has the potential to be hydrolyzed into simple sugar using *Penicillium sp. H9*. The cellulose, hemicellulose, and lignin contents of some organic wastes are shown in Table 1. The delignification process was needed to break the bonds between lignin and cellulose. The results showed the success of delignification was shown by an increase of cellulose, hemiselulose, and lignin levels in rice straw, 52.8%, 30.4% and 4.3% respectively. Rice straw hydrolized acquisition from the hydrolysis of rice straw by *Penicillium sp. H9* was 199 µg / mL for 6 days of incubation. The rice straw hydrolized obtained that used as a growth substrate and biosurfactant production by LII61.

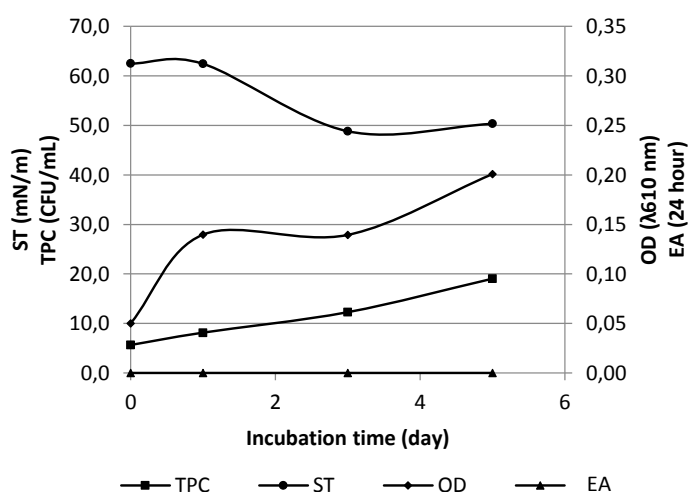
Table 1. Content of cellulose, hemiselulose and lignin in organic wastes

Organic wastes	Content (%)			References
	Cellulose	Hemicellulose	Lignin	
Palm oil	71.5	9.9	19.9	[15]
Wheat straw	38.2	21.2	23.4	[16], [17]
Bagasse	38.2	27.1	20.2	[16], [18]
Sorghum Biomass	22.2	19.4	21.4	[19]
Molasses	25.0	17.0	12.0	[16]
Rice straw	35.0	18.0	20.9	[20]
Rice straw	42.2	21.3	4.4	This work

Hydrolysis enzymatically will break the polymer chain specifically on certain branch while breaking the polymer chain with chemical and physical hydrolysis occur randomly [21]. *Penicillium* sp. H9 produces cellulase that play a role in the hydrolysis [22]. The mechanism of cellulose hydrolysis is through enzymatic (1) endo- β -1,4-glucanase breaks the β -1,4-glucoside bond randomly, especially amorf part of polysaccharide chains to produces cellobiose (2) ekso- β -1,4-cellobiohydrolase split of cellobiose unit from non-reduce polysaccharide chain terminal (3) β -glucosidase hydrolyze cellobiose and small chain of cello-oligosaccharide for producing glucose [23]. Concentration of rice straw, microbial enzyme, and environment condition influenced the hydrolysis product.

3.2 Screening of biosurfactant production by LII61 on RSHP substrate

Based on Fig. 1, enhancement of OD value and TPC were showed by LII61 during 5 days incubation (0.20 and 19.05 CFU/mL). The growth of LII61 suggested that RSHP can be a carbon source converted to energy for growth process. LII61 also showed existence of biosurfactant which was showed by lowering the surface tension value. The optimum lowering of surface tension was obtained in three days of incubation time amount 48,85 mN/m. Bioemulsifier from LII61 in the RSHP not detected because the concentration of biosurfactant is low.

**Figure 1.** Screening of biosurfactant product by LII61 in rice straw substrate

3.3 Optimization of biosurfactant production by LII61

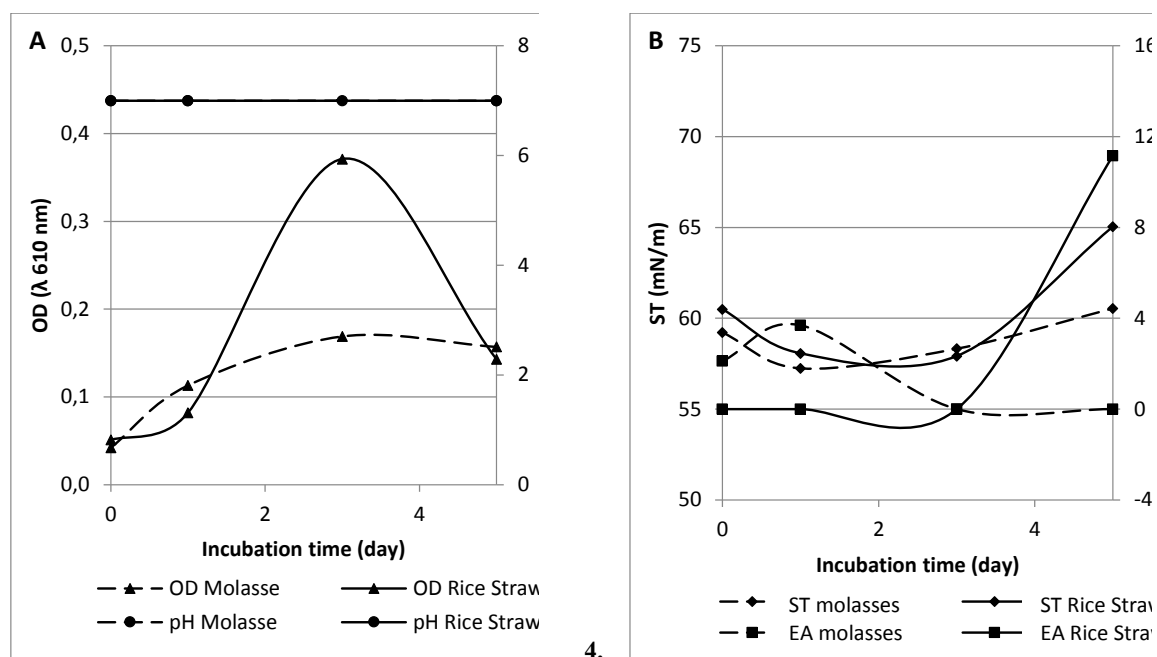


Figure 2. Optimalization of the biosurfactant production from LII61 in the RSHP substrate with molasses as control. A) growth and pH during incubation B) the activity of biosurfactant product

The results show that LII61 entered the log phase from the first day of incubation. In the molasses substrate and RSHP, LII61 showed a different response. The growth rate of LII61 on the RSHP was twice as fast as the molasses substrate at the same concentration (Fig. 2A). At the end of incubation, LII61 growth on RSHP substrates experienced a significant decrease. This shows the presence of cell death in culture. Whereas, in molasses culture, the decrease in cell counts did not differ significantly from the 3rd day of incubation. In this case pH changes were not found. Emulsification activity was detected in both substrates with the highest yield on RSHP substrate on the 5th day of incubation by 10.35% (Fig. 2B). Surface tension reduction was also detected on both substrates with significant differences. The optimum time for biosurfactant production of LII61 in the RSHP substrate is 3rd day incubation supported by the highest of number cell and the lowest of surface tension of culture supernatant. Similar with other bacteria, biosurfactant from *Corynebacterium lepus* are produced during exponential growth phase [24]. In this case, the increasing of emulsification activity in the 5th day incubation was affected by the lysis of cells. Utilization of RSHP as substrate have succesfull be a trigger of LII61 to produce biosurfactant. This showed that at the same concentration, RSHP can be an alternative substrate to substitute molasses.

3.4 Extraction of biosurfactant from LII61

The amount of biosurfactant produced by LII61 in RSHP was 80 mg/L during 3 days incubation. The crude biosurfactant can reduce the surface tension from 72 mN/m to 53.3 mN/m (Fig. 3). The CMC value of extracted biosurfactant was 1000 mg/L. Based on the previous study, the CMC value of crude biosurfactant produced by LII61 in the molasses substrate was 6000 mg/L with the value of surface tension was 51.5 mN/m [25]. This discovery is better than molasses as substrate on biosurfactant production. At CMC, surfactant molecules aggregate and form micelles in polar or aqueous environment [26]. The value of surface tension unable to significantly decrease at CMC.

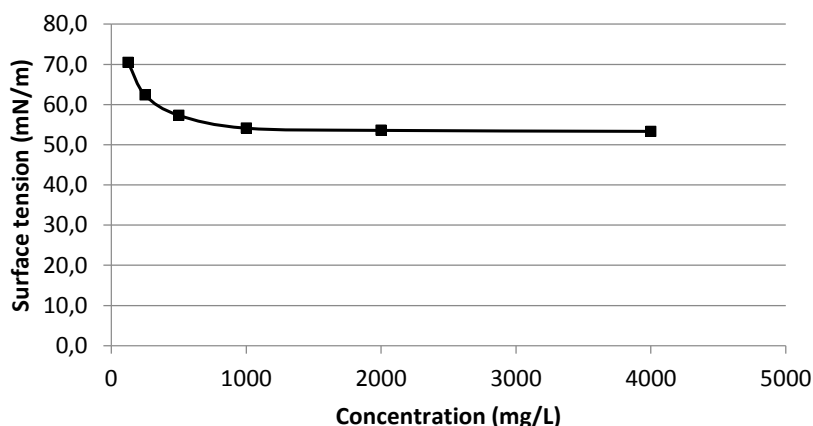


Figure 3. Critical Micelle Concentration (CMC) of crude biosurfactant in RSHP

Characterization of crude biosurfactant from LII61 towards temperature and pH

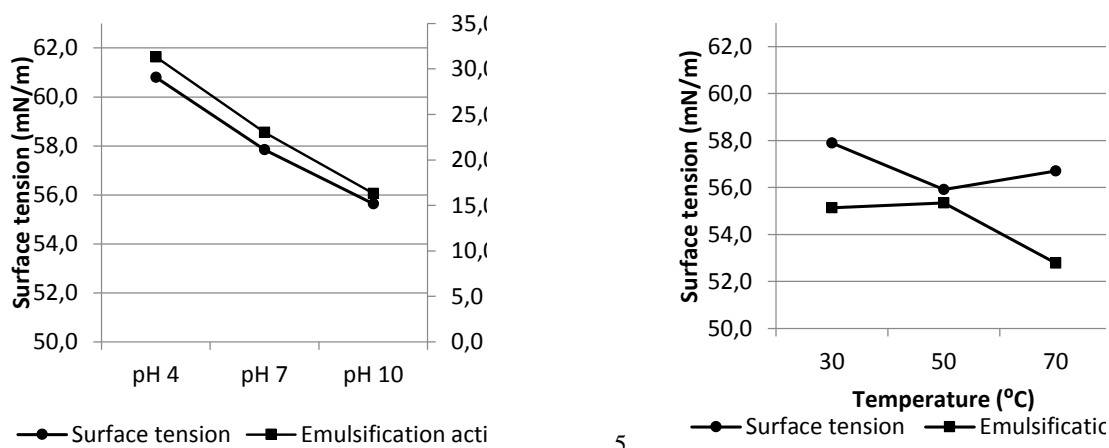


Figure 4. Characterization of crude biosurfactant from LII61 at various of condition A) pH B) temperature

Crude biosurfactant was tested at various pH and temperature. The surface tension value is showed at figure 4. The value of surface tension and emulsification activity at CMC of crude biosurfactant unstable in the various of pH. The surface tension decreased along with increased of pH (alkali) and the emulsification activity increased along with decreased of pH (acid). The optimum lowering of surface tension of LII 61 crude biosurfactant in the pH of 10 up to 55.6 mN/m approaching surface tension of CMC. This could be caused by a better stability of fatty acids surfactant micelles in the presence of sodium hydroxide and the precipitation of secondary metabolites at higher pH values [27]. The crude biosurfactant in the range of temperature 30°C - 70°C still show biosurfactant activities through emulsification activity and lowering surface tension. The crude biosurfactant capable to decrease the surface tension and show the emulsification activity in 30°C until 70 °C. The RSHP has the potential as substrate to produce biosurfactant from LII61.

4. Conclusions

The level of RSHP is 229 µg/mL in 6 days incubation time. RSHP can be used by LII61 to grow and produce biosurfactant as indicated by increasing of total plate count and optical density until 5 days. The best result of obtaining biosurfactant on RSHP are indicated by the surface tension 48.85 mN/m. Biosurfactants produced by LII61 have CMC value at 1000 mg/L and have the certain characteristic at

various pH and temperature. The use of rice straw hydrolysis product as a substitute for molasses in the manufacture of biosurfactant by LII61 shows good results and has favorable prospects.

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