

Volume 2055

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# Proceedings of the Europe/Africa Conference Dresden 2017 – Polymer Processing Society PPS



**Dresden, Germany**  
27-29 June 2017

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## • Table of Contents

### PROCEEDING OF INTERNATIONAL BIOLOGY CONFERENCE 2016: Biodiversity and Biotechnology for Human Welfare

< PREV NEXT >



Conference date: 15 October 2016

Location: Surabaya, Indonesia

ISBN: 978-0-7354-1528-7


Editors: Michael Murkovic, Gianfranco Risuleo, Endry Nugroho Prasetyo, Maya Shovitri and Gibson S. Nyanhongo

Volume number: 1854

Published: Jun 26, 2017

DISPLAY : 20 50 100 all

## PRELIMINARY


 Free . June 2017

### **Preface: Proceeding of International Biology Conference 2016 Biodiversity and Biotechnology for Human Welfare**

AIP Conference Proceedings **1854**, 010001 (2017); <https://doi.org/10.1063/1.4985390>

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
### **The Committee of 3rd International Biology Conference (IBOC) 2016 & 10th Korea-ASEAN Biomass Symposium**

AIP Conference Proceedings **1854**, 010002 (2017); <https://doi.org/10.1063/1.4985391>

---



## ARTICLES

 Free . June 2017

### **Lipase production in lipolytic yeast from Wonorejo mangrove area**

Nur Hidayatul Alami, Liziyatin Nasihah, Rurin Luswidya Artaty Umar, Nengah Dwianita  
Kuswytasari, Enny Zulaika and Maya Shovitri

AIP Conference Proceedings **1854**, 020001 (2017); <https://doi.org/10.1063/1.4985392>

---

SHOW ABSTRACT



## The production and activity test of cellulases using bagasse substrate on *Aspergillus niger* isolated from Clove field, Kare, Madiun


Muh. Waskito Ardhi, Ani Sulistyarsi and Pujiati

AIP Conference Proceedings **1854**, 020002 (2017); <https://doi.org/10.1063/1.4985393>

---

SHOW ABSTRACT



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## Lead (Pb) bioaccumulation; genera *Bacillus* isolate S1 and SS19 as a case study


Achmad Arifiyanto, Fitria Dwi Apriyanti, Puput Purwaningsih, Septian Hary Kalqutny, Dyah Agustina, Tini Surtiningsih, Maya Shovitri and Enny Zulaika

AIP Conference Proceedings **1854**, 020003 (2017); <https://doi.org/10.1063/1.4985394>

---

SHOW ABSTRACT



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## Local community knowledge and participation for animal diversity conservation in SSWP IV Sidoarjo, East Java, Indonesia

Nova Maulidina Ashuri, Dwi Oktafitria, Indra Wirawan, Zainul Muttaqin, M. Ulya Alfarisy, Abdul Azis, Sherly Eka Argiyanti and Via Nur Fadilah

AIP Conference Proceedings **1854**, 020004 (2017); <https://doi.org/10.1063/1.4985395>

---

SHOW ABSTRACT



## Potency of *Nicotiana tabacum* as anti - microfouling


Aunurohim, Dian Ahmada Nurilma and Nengah Dwianita Kuswytasari

AIP Conference Proceedings **1854**, 020005 (2017); <https://doi.org/10.1063/1.4985396>

---

SHOW ABSTRACT



 Free . June 2017

## The effect of pomelo citrus (*Citrus maxima* var. Nambangan), vitamin C and lycopene towards the number reduction of mice (*Mus musculus*) apoptotic hepatocyte caused of ochratoxin A


Badriyah and Utami Sri Hastuti

AIP Conference Proceedings **1854**, 020006 (2017); <https://doi.org/10.1063/1.4985397>

---

SHOW ABSTRACT



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## Potency of bio-charcoal briquette from leather cassava tubers and industrial sludge


Nita Citrasari, Tety A. Pinatih, Eko P. Kuncoro, Agoes Soegianto, Salamun and Bambang Irawan

AIP Conference Proceedings **1854**, 020007 (2017); <https://doi.org/10.1063/1.4985398>

---

SHOW ABSTRACT



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## Distribution study on migratory bird (*Scolopacidae: Numenius*) in Surabaya, Indonesia: Estimating the effect of habitat and

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## climate change


Iska Desmawati, Indah Trisnawati D. T., Ory Kurnia, Albi Hamdani, Mahsun Fahmi and Mirza Fahmi

AIP Conference Proceedings **1854**, 020008 (2017); <https://doi.org/10.1063/1.4985399>

---

SHOW ABSTRACT



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### **Effect of gamma $^{60}\text{Co}$ irradiation on the lipid content and fatty acid composition of *Nannochloropsis* sp. microalgae**


Dini Ermavitalini, Ika Puspita Sari, Endry Nugroho Prasetyo, Nurlita Abdulgani and Triono Bagus Saputro

AIP Conference Proceedings **1854**, 020009 (2017); <https://doi.org/10.1063/1.4985400>

---

SHOW ABSTRACT



 Free . June 2017

### **Effects of gamma irradiation dose-rate on sterile male *Aedesaegypti***


Beni Ernawan, Usman Sumo Friend Tambunan, Irawan Sugoro and Hadian Iman Sasmita

AIP Conference Proceedings **1854**, 020010 (2017); <https://doi.org/10.1063/1.4985401>

---

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
### **Bioremediation of oil sludge using a type of nitrogen source and the consortium of bacteria with composting method**

AIP Conference Proceedings **1854**, 020011 (2017); <https://doi.org/10.1063/1.4985402>

---

SHOW ABSTRACT



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## **Analysis of *Hylocereus* spp. diversity based on phenetic method**


Hamidah, Husnus Tsawab and Rosmanida

AIP Conference Proceedings **1854**, 020012 (2017); <https://doi.org/10.1063/1.4985403>

---

SHOW ABSTRACT



 Free . June 2017

## **Briquettes of rice husk, polyethylene terephthalate (PET), and dried leaves as implementation of wastes recycling**


Sucipto Hariyanto, Mohammad Nurdianfajar Usman and Nita Citrasari

AIP Conference Proceedings **1854**, 020013 (2017); <https://doi.org/10.1063/1.4985404>

---

SHOW ABSTRACT



 Free . June 2017

## **Histopathological assessment of cadmium effect on testicles and kidney of *Oreochromis niloticus* in different salinity**


Alfiah Hayati, Hanna Pratiwi, Inayatul Khoiriyah, Dwi Winarni and Sugiharto

AIP Conference Proceedings **1854**, 020014 (2017); <https://doi.org/10.1063/1.4985405>

---

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 Free . June 2017

## **The influence of snakehead (*Channa striata*) fish extract to increase hyperglycemic mice fertility based on spermatogenic cell composition**


Dewi Hidayati, Nurlita Abdulgani, Nova Maulidina Ashuri, Noor Nailis Sa'adah and Maharani Lukitasari

AIP Conference Proceedings **1854**, 020015 (2017); <https://doi.org/10.1063/1.4985406>

---

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## **Analysis of protein profiles in diabetic rat blood plasma that induced by alloxan**


Dewi Hidayati, Nurlita Abdulgani, Hengki Setiyawan, Indah Trisnawati, Nova Maulidina Ashuri and Noor Nailis Sa'adah

AIP Conference Proceedings **1854**, 020016 (2017); <https://doi.org/10.1063/1.4985407>

---

SHOW ABSTRACT



 Free . June 2017

## **Effect of methionine and lactic acid bacteria as aflatoxin binder on broiler performance**

Lusty Istiqomah, Ema Damayanti, Hardi Julendra, Ade Erma Suryani, Awistaros Angger Sakti and Ayu Septi Anggraeni


AIP Conference Proceedings **1854**, 020017 (2017); <https://doi.org/10.1063/1.4985408>

---



SHOW ABSTRACT



 Free . June 2017

## **Growth and physiological responses of some *Capsicum frutescens* varieties to copper stress**


Nurul Jadid, Rizka Maziyah, Desy Dwi Nurcahyani and Nilna Rizqiyah Mubarokah

AIP Conference Proceedings **1854**, 020018 (2017); <https://doi.org/10.1063/1.4985409>

---

SHOW ABSTRACT



 Free . June 2017

## **Antioxidant activities of different solvent extracts of *Piper retrofractum* Vahl. using DPPH assay**


Nurul Jadid, Dewi Hidayati, Sylviana Rosyda Hartanti, Byan Arasyi Arraniry, Rizka Yuanita Rachman and Wiwi Wikanta

AIP Conference Proceedings **1854**, 020019 (2017); <https://doi.org/10.1063/1.4985410>

---

SHOW ABSTRACT



 Free . June 2017

## **Adsorption of cadmium from aqueous solution using algae waste based adsorbent**


Eko Prasetyo Kuncoro, Thin Soedarti, Trisnadi Widyalaksono Catur Putrato and Nurul Alvia Istiqomah

AIP Conference Proceedings **1854**, 020020 (2017); <https://doi.org/10.1063/1.4985411>

---

SHOW ABSTRACT



 Free . June 2017

## **Mixing of acacia bark and palm shells to increase caloric value of palm shells white charcoal briquette**


Edy Wibowo Kurniawan, Rudianto Amirta, Edy Budiarmo and Enos Tangke Arung

AIP Conference Proceedings **1854**, 020021 (2017); <https://doi.org/10.1063/1.4985412>

---

SHOW ABSTRACT



 Free . June 2017

## **Keratinase from newly isolated strain of thermophilic Bacillus for chicken feed modification**


Ditya Larasati, Nur Tsurayya, Maharani Pertiwi Koentjoro and Endry Nugroho Prasetyo

AIP Conference Proceedings **1854**, 020022 (2017); <https://doi.org/10.1063/1.4985413>

---

SHOW ABSTRACT



 Free . June 2017

## **Revegetation increase bird diversity in coastal area of Socorejo, Tuban, East Java – Indonesia**

Yeni Indah Lestari, Wasito Edi, Alkautsar Alivvy, Acib Setia Ibadah, Fadina Yuliana Sari, Finda Nuraini, Ahmad Yanuar, Agus Satriyono, Citra Fitrie Riany, Dian Saptarini and Farid Kamal Muzaki

AIP Conference Proceedings **1854**, 020023 (2017); <https://doi.org/10.1063/1.4985414>

---

SHOW ABSTRACT



## Growth of vegetative explant *Moringa oleifera* on different composition of auxin and cytokinin and its synthetic seed germination


Wirdhatul Muslihatin, Nurul Jadid, Ika D. Puspitasari and Chusnul E. Safitri

AIP Conference Proceedings **1854**, 020024 (2017); <https://doi.org/10.1063/1.4985415>

---

SHOW ABSTRACT



 Free . June 2017

## Community structure of fish larvae in mangroves with different root types in Labuhan coastal area, Sepulu – Madura


Farid Kamal Muzaki, Aninditha Giffari and Dian Saptarini

AIP Conference Proceedings **1854**, 020025 (2017); <https://doi.org/10.1063/1.4985416>

---

SHOW ABSTRACT



 Free . June 2017

## Biodegradation of naphthalene and phenanthren by *Bacillus subtilis* 3KP


Ni'matuzahroh, N. Trikurniadewi, A. R. A. Pramadita, I. A. Pratiwi, Salamun, Fatimah and Sri Sumarsih

AIP Conference Proceedings **1854**, 020026 (2017); <https://doi.org/10.1063/1.4985417>

---

SHOW ABSTRACT



 Free . June 2017

## **Manganese (Mn) stress toward hyperaccumulators plants combination (HPC) using *Jatropha curcas* and lamtoro gung (*L. leucocephala*) in mychorrizal addition on soybean (*Glycine max*) seedling stage**


Tania Sylviana Darmawan, Tata Taqiyyatuz Zahroh, Mirza Merindasya, Ririn Masfaridah, Dyah Ayu Sri Hartanti, Sekar Arum, Sri Nurhatika, Anton Muhibuddin, Tini Surtiningsih and Achmad Arifiyanto

AIP Conference Proceedings **1854**, 020027 (2017); <https://doi.org/10.1063/1.4985418>

---

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 Free . June 2017

## **The potency of curing fish waste pellet for growth and protein level of African sharptooth catfish (*Clarias gariepinus*)**


Awik Puji Dyah Nurhayati and Asti R. Febiyani

AIP Conference Proceedings **1854**, 020028 (2017); <https://doi.org/10.1063/1.4985419>

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
 Free . June 2017

## **Reducing the level of leaves damage of (*Brassica rapa*) caused by armyworm (*Spodoptera litura* F.) through liquid bioinsecticide formulation of bintaro (*Cerbera odollam*) leaves extract**

Kristanti Indah Purwani, Sri Nurhatika, Dini Ermavitalini, Triono Bagus Saputro and Dwi Setia Budiarti

AIP Conference Proceedings **1854**, 020029 (2017); <https://doi.org/10.1063/1.4985420>

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 Free . June 2017

## **Molecular characters of melon (*Cucumis melo* L. “Tacapa”) in response to karst critical land**


Yuanita Rachmawati, Budi Setiadi Daryono and Ganies Riza Aristya

AIP Conference Proceedings **1854**, 020030 (2017); <https://doi.org/10.1063/1.4985421>

---

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## **Analysis of lipid profile and atherogenic index in hyperlipidemic rat (*Rattus norvegicus* Berkenhout, 1769) that given the methanolic extract of Parijoto (*Medinilla speciosa*)**


Noor Nailis Sa’adah, Kristanti Indah Purwani, Awik Puji Dyah Nurhayati and Nova Maulidina Ashuri

AIP Conference Proceedings **1854**, 020031 (2017); <https://doi.org/10.1063/1.4985422>

---

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## **White syndrome on massive corals: A case study in Paiton power plant, East Java**


Farid Kamal Muzaki, Dian Saptarini and Aida Efrini Riznawati

AIP Conference Proceedings **1854**, 020032 (2017); <https://doi.org/10.1063/1.4985423>

---

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## **Genetic diversity of improved salt tolerant calli of maize (*Zea mays* L.) using RAPD**


Triono Bagus Saputro, Siti Dianawati, Nur Fadlillatus Sholihah and Dini Ermavitalini

AIP Conference Proceedings **1854**, 020033 (2017); <https://doi.org/10.1063/1.4985424>

---

SHOW ABSTRACT



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## **Spicule size variation in *Xestospongia testudinaria* Lamarck, 1815 at Probolinggo-Situbondo coastal**


Iwenda Bella Subagio, Edwin Setiawan, Sucipto Hariyanto and Bambang Irawan

AIP Conference Proceedings **1854**, 020034 (2017); <https://doi.org/10.1063/1.4985425>

---

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## **Soil burial method for plastic degradation performed by *Pseudomonas* PL-01, *Bacillus* PL-01, and indigenous bacteria**


Maya Shovitri, Risyatun Nafi'ah, Titi Rindi Antika, Nur Hidayatul Alami, N. D. Kuswytasari and Enny Zulaikha

AIP Conference Proceedings **1854**, 020035 (2017); <https://doi.org/10.1063/1.4985426>

---

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## **Production of secondary metabolites *trimethyl xanthina* by *Camellia sinensis* L suspension culture**


Sutini, Mochamad Sodik, Wirdhatul Muslihatin and Mochamad Rasjad Indra

AIP Conference Proceedings **1854**, 020036 (2017); <https://doi.org/10.1063/1.4985427>

---

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## **Effects of the combination between bio-surfactant product types and washing times on the removal of crude oil in nonwoven fabric**


Agus Triawan, Ni'matuzahroh and Agus Supriyanto

AIP Conference Proceedings **1854**, 020037 (2017); <https://doi.org/10.1063/1.4985428>

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 Free . June 2017

## **The effectiveness of habitat modification schemes for enhancing beneficial insects: Assessing the importance of trap cropping management approach**


Indah Trisnawati and Abdul Azis

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
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
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
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**Publication type** Conferences and Proceedings

**ISSN** 0094243X, 15517616

**Coverage** 1974-1978, 1983-1984, 1993, 2000-2001, 2003-2020

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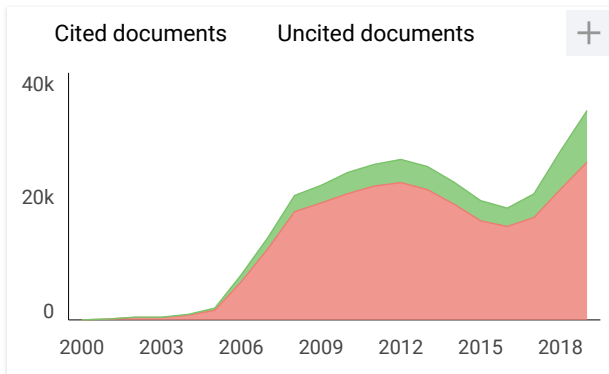
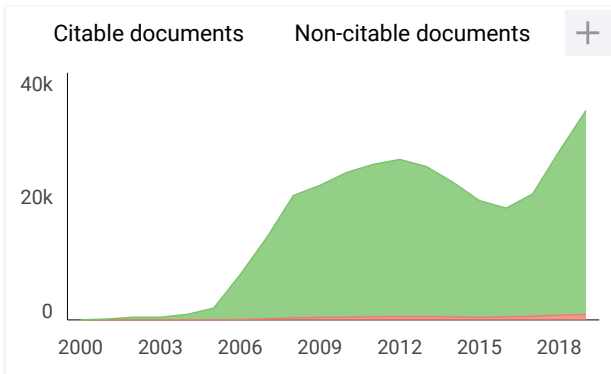
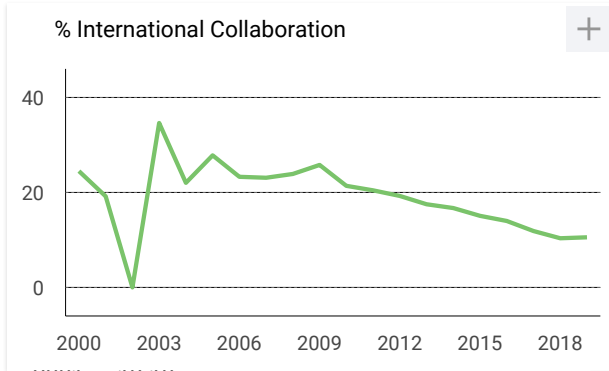
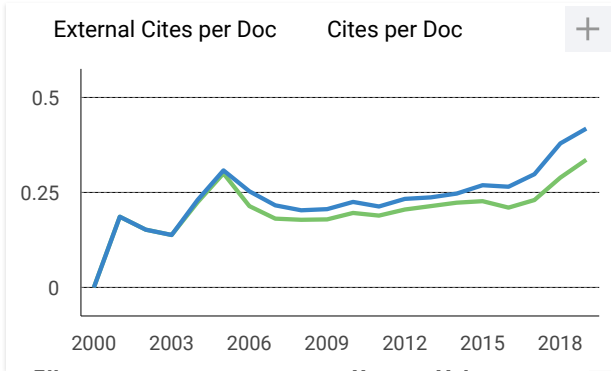
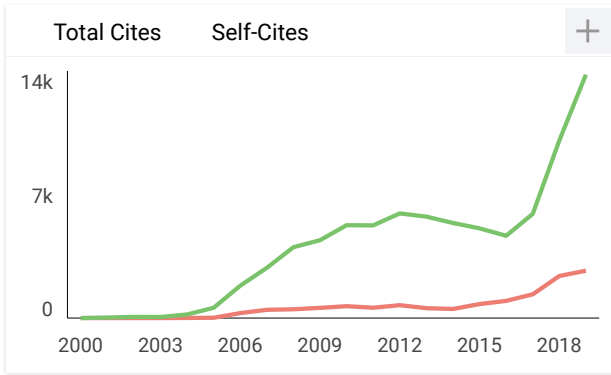
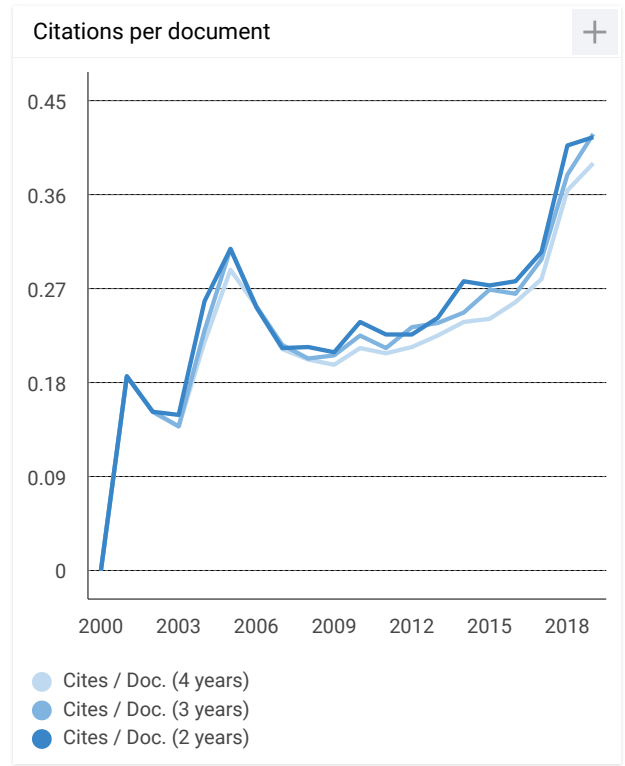
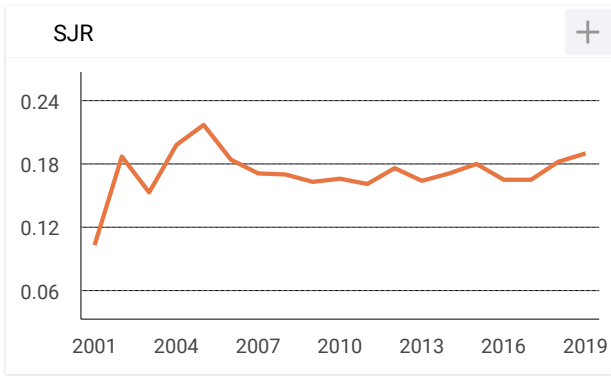
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Cite as: AIP Conference Proceedings 1854, 020026 (2017); <https://doi.org/10.1063/1.4985417>  
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# Biodegradation of Naphthalene and Phenanthrene by *Bacillus subtilis* 3KP

Ni'matuzahroh<sup>1,a)</sup>, Trikurniadewi N.<sup>1,b)</sup>, Pramadita A. R. A.<sup>1</sup>, Pratiwi I. A.<sup>1</sup>,  
Salamun<sup>1</sup>, Fatimah<sup>1</sup>, Sumarsih Sri<sup>2</sup>

<sup>1</sup> Department of Biology, Faculty of Science and Technology, Universitas Airlangga, 60115 Surabaya, Indonesia.

<sup>2</sup> Department of Chemistry, Faculty of Science and Technology, Universitas Airlangga, 60115 Surabaya, Indonesia.

a) Corresponding author: [nimatuzahroh@fst.unair.ac.id](mailto:nimatuzahroh@fst.unair.ac.id)

b) [nastiti.trikurniadewi-2015@fst.unair.ac.id](mailto:nastiti.trikurniadewi-2015@fst.unair.ac.id)

**Abstract.** The purposes of this research were to know growth response, degradation ability, and uptake mechanism of naphthalene and phenanthrene by *Bacillus subtilis* 3KP. *Bacillus subtilis* 3KP was grown on Mineral Synthetic (MS) medium with addition of 1% yeast extract and naphthalene and phenanthrene respectively 200 ppm in different cultures. *Bacillus subtilis* 3KP growth response was monitored by Total Plate Count (TPC) method, the degradation ability was monitored by UV-Vis spectrophotometer, and the uptake mechanism of hydrocarbon was monitored by emulsification activity, decrease of surface tension, and activity of Bacterial Adherence to Hydrocarbon (BATH). *Bacillus subtilis* 3KP was able to grow and show biphasic growth pattern on both of substrates. Naphthalene and phenanthrene were used as a carbon source for *Bacillus subtilis* 3KP growth that indicated by the reduction of substrate concomitant with the growth. At room temperature conditions ( $\pm 30^{\circ}\text{C}$ ) and 90 rpm of agitation for 7 days, *Bacillus subtilis* 3KP could degrade naphthalene in the amount of 70.5% and phenanthrene in the amount of 24.8%. Based on the analysis of UV-Vis spectrophotometer, three metabolites, 1-hydroxy-2-naphthoic acid, salicylic acid, and pyrocatechol were found in both cultures. The metabolite identification became basis of propose degradation pathway of naphthalene and phenanthrene by *Bacillus subtilis* 3KP. The results of hydrocarbon uptake mechanism test show that *Bacillus subtilis* 3KP used all of the mechanism to degrade naphthalene and phenanthrene.

Keyword: *Bacillus subtilis* 3KP, Biodegradation, Biosurfactant, Phenanthrene, Naphthalene

## INTRODUCTION

Naphthalene and phenanthrene are types of hydrocarbon that classified as polycyclic aromatic hydrocarbon (PAH). This kind of compound can cause damage on mammal tissue when it is exposure in high dose (Patnaik, 1994). The existence of PAH, as pollutant, is often yield from exploration and production in petroleum industry. Petroleum waste sludge (oil sludge) consists of hydrocarbon compounds such as aliphatic hydrocarbons and polycyclic aromatic hydrocarbons (PAH), water, metals and non-hydrocarbon compounds such as nitrogen, sulfur, oxygen, and asphalt (Connell and Miller, 1995). PAH almost reaches amount of 13.24% in petroleum (Yuliani, 2014).

A method called bioremediation has become an important method to solve the petroleum waste by using indigenous or exogenous microbes (Helmy et al., 2010). The progress on degradation study showed how difficult to degrade PAH because of its stability and recalcitrantly in soil. Less of solubility also limits bioavailability and efficiency of bioremediation process.

In biodegradation process of PAH, it is often mentioned about the role of biosurfactant on biodegradation. Biosurfactant could increase solubility and availability of substrate for bacteria (Li, 2009). Besides

biosurfactant, some enzymes also have been linked with biodegradation process of hydrocarbon, especially PAH. The existence of enzyme enable bacterial cell degrade hydrocarbon polymer into simpler compound. So, it can be used as carbon source. Various types of enzyme also related with hydrocarbon uptake mechanism of bacteria.

*Bacillus subtilis* 3KP was bacteria which known as hydrocarbon degrading bacteria. These bacteria are also able to produce biosurfactant. According to Ni'matuzahroh et al. (2013), *Bacillus subtilis* 3KP was able to produce biosurfactant in the amount of 12.3 g/L using molasses substrate. *Bacillus subtilis* 3KP was potential to degrade PAH compound, but the research has not ever done yet. This research aims to observe the bacterial growth, biodegradation percentage of PAH, and mechanism of biodegradation that was probably developed by *Bacillus subtilis* 3KP.

## MATERIALS AND METHODS

### Chemicals

All chemicals were analytical grade or better. Naphthalene (99%), phenanthrene (99%), 1-hydroxy-naphtoic acid (99%), salicylic acid (99%), and pyrocatechoid acid (99%) were purchased from Sigma-Aldrich. Stock solution of 1-hydroxy-naphtoic acid, salicylic acid, and pyrocatechoid acid were prepared in ethyl acetate (MERCK). Naphthalene and phenanthrene were prepared in mineral salts medium (MS, see below) and sterilized by autoclaving before the bacteria were inoculated.

### Microorganism and Growth Medium

The following bacteria used are *Bacillus subtilis* 3KP which isolated from petroleum-contaminated environment in Donan River, Cilacap, Indonesia. *Bacillus subtilis* 3KP was grown in Nutrient Broth and Nutrient Agar before inoculation.

All media used in these studies were prepared in distilled water. The MS medium from the method by Pruthi and Cameotra (1997) contained ( $L^{-1}$ ) 3 g  $(NH_4)_2SO_4$ , 0.2  $MgSO_4 \cdot 7H_2O$ , 10 g NaCl, 0.01  $CaCl_2$ , 0.001 g  $MnSO_4 \cdot H_2O$ , 0.001 g  $H_3BO_3$ , 0.001 g  $ZnSO_4 \cdot 7H_2O$ , 0.001 g  $CuSO_4 \cdot 5H_2O$ , 0.005 g  $CoCl_2 \cdot 6H_2O$ , 0.001 g  $Na_2MoO_4 \cdot 2H_2O$ , and 1% yeast extract. The buffer composition contained ( $L^{-1}$ )  $K_2HPO_4$  and  $KH_2PO_4$  for respectively 2.6207 g and 5 g in 50 ml. It was also added 0.0006 g  $FeSO_4 \cdot 7H_2O$  which sterilized separately in 50 ml distilled water. The pH of the final medium was adjusted to 7.0. For preparing inoculation, bacterial cultures were suspended in 0.85% salt solution.

### Determination of Residual Substrat

Biodegradation tests were performed in 250 ml flask and run in duplicate. *Bacillus subtilis* 3KP (5%) (v/v) was grown in 30 ml MS medium. Naphthalene and phenanthrene were added to the medium 200 ppm respectively. The flasks were shaken on a rotary shaker with 90 rpm at 30°C for 7 days. All cultures were started with cell density  $OD_{610\text{ nm}}$  of 0.5. Growth profile was observed by common serial dilution. The residual hydrocarbons and the bacterial growth were investigated at days 0, 1, 3, 5, and 7.

At the same time of interval, analysis of hydrocarbon residue in culture was observed by UV-vis spectrophotometer (Kumar et al., 2010). Before analysis, the culture was extracted by adding ethyl acetate to the separating funnel in ratio 1:1. Then it was shaken twice for 15 minute in each shaking. Separating of liquid phase, emulsion, and solvent phase was done by adding absolute ethanol. The data was shown as absorbance value at optimal wavelength 276 nm for naphthalene, 293 nm for phenanthrene, 339 nm for 1-hydroxy-2-naphtoic acid, 304 nm for salicylic acid, and 278 nm for pyrocatechol. Standard curve was used to determine concentration of naphthalene or phenanthrene in the culture. Hydrocarbons biodegradation percentage was calculated with the formula:

$$\text{Biodegradation (\%)} = \left( \frac{\text{initial concentration of hydrocarbon} - \text{final concentration of hydrocarbon}}{\text{initial concentration of hydrocarbon}} \right) \times 100\%$$

### Cell surface Hydrophobicity

According to Rosenberg et al. (1980), the percentage of the cells adhering to an oily phase was investigated by using a modification method. The bacterial cells were harvested each 48 hours by centrifugation (3000 rpm, 15 minute). The cell pellets were washed twice and suspended in phosphate buffer ( $g\ L^{-1}$ ; 7.3  $KH_2PO_4$  and 16.9  $K_2HPO_4$ ), then diluted to an optical density (OD) of 0.5 at 610 nm. Into the cell suspension (4 mL) in test tube,

100  $\mu\text{L}$  of hexadecane was added and vortex-shaken for 3 minute. After forming the suspension, it was allowed to separate for 60 minute, the absorbance  $\text{OD}_{610}$  in the aqueous phase was measured directly. The clearing measurement of the aqueous phase indicated the capacity of adherence cell to the oily phase. The result is presented as percentage with formula:

$$H (\%) = \left(1 - \frac{A}{A0}\right) \times 100\%$$

H was hydrophobicity; A was cell suspension absorbance after added by hexadecane; A0 was initial cell suspension absorbance.

### Surface Tension Reduction

All experiments were performed using two separately grown bacterial cultures. The cultures were centrifuged at 3000 rpm for 15 minute. Then, the surface tension of supernatant was measured by Du-Nouy tensiometer. The result is presented as  $\text{mN m}^{-1}$  and reported as an average of triplicate each sample.

### Emulsification Activity Assay

Emulsification activity was measured using the method by Cooper and Goldenberg (1987). The supernatant (1 mL) as yield of centrifugation was added to a test tube containing 1 ml kerosene. The mixture was homogenized by vortexing for 2 minute and the emulsion was allowed to stand for 60 minute to be measured. Emulsification index was determined as ratio of the emulsion height to total height of the mixture, then it was calculated their percentage.

## RESULTS AND DISCUSSIONS

Biodegradation process of PAH is affected by several factors, those are microbial character, environmental factor that support the growth, and characteristic of substrate. Microbial ability in degradation of hydrocarbons depends on its adaptation, toxic resistance, and gene expression of the microbe (Abbasnezhad et al., 2011). Microbial ability in degrade hydrocarbon could be known through the growth, the use of hydrocarbon as sole carbon, and the secondary metabolites that produced by microbe.

*Bacillus subtilis* 3KP grew up and showed positive responses to naphthalene and phenanthrene, 200 ppm respectively. *Bacillus subtilis* 3KP was able to adapt and use PAH as a sole carbon (Fig.1). This is proved by decreasing of concentration of substrate accompanied by increasing of the cell amount. The growth pattern of *Bacillus subtilis* 3KP on naphthalene and phenanthrene showed a biphasic pattern. In compare to control study that showed a normal phase, this kind of pattern indicated that *Bacillus subtilis* 3KP has adapted to the content of PAH in the medium.

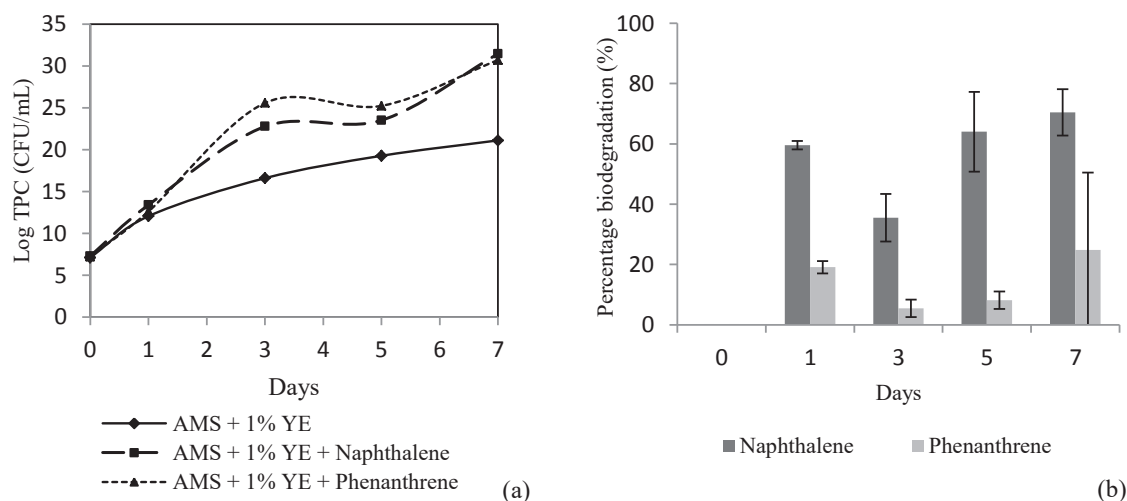


FIGURE 1. (a) *Bacillus subtilis* 3KP growth response in the naphthalene and phenanthrene substrates, (b) Percentage of naphthalene and phenanthrene biodegradation at several days

Stationary phase occurs when cell growth was inhibited by cell density, accumulation of secondary metabolites that could be toxic and limited oxygen (Pelczar, 1986). In this study, accumulation of intermediate compound was found in the end of exponential phase, those are 1-hydroxy-2-naphthoic acid and salicylic acid. According to metabolic pathway of phenanthrene degradation by bacteria, 1-hydroxy-2-naphthoic acid would be transformed to 1-2-dihydroxynaphthalene before it became salicylic acid. Pumphrey and Madsen (2007) have explained that 1-2-dihydroxynaphthalene could be oxidized to a toxic 1-2-naphthaquinon which inhibited the growth of *Polaromonas naphthalenivorans*. The intermediate compounds of naphthalene and phenanthrene metabolism can be accumulated in culture because of their easy-soluble character and more toxic while they have increased continuously.

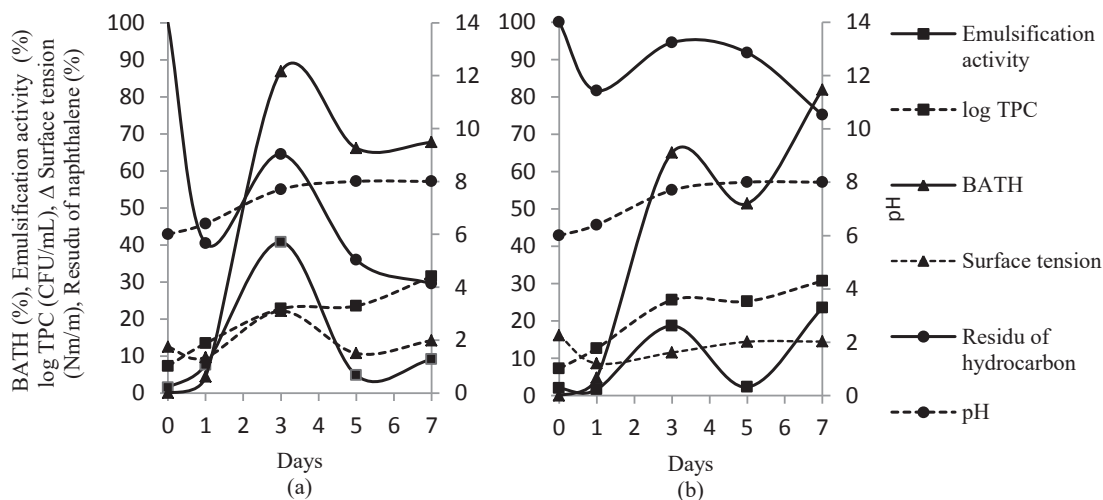
Some microorganisms are able to hold out from naphthalene and phenanthrene and to transform them to enter the TCA (*tricarboxylic acid*) cycle, *Bacillus subtilis* 3KP did it. At day 7, *Bacillus subtilis* 3KP has degraded 70.5% naphthalene and 24.8% phenanthrene. Intermediate compounds have been observed in the study, those are 1-hydroxy-2-naphthoic acid and salicylic acid. *Pyrocatechol* was used along with the use of main substrate. It showed that *Bacillus subtilis* 3KP has degraded naphthalene and phenanthrene in a same metabolic pathway. Data existence of intermediate was shown in table 1.

**TABLE 1.** The existence of intermediate compounds during the process of degradation of naphthalene and phenanthrene by *Bacillus subtilis* 3KP

Days	Existence of intermediate compounds					
	1-hydroxy-2-naphthoicacid		Salicylic acid		Pyrocatechol	
	Naphthalene	Phenanthrene	Naphthalene	Phenanthrene	Naphthalene	Phenanthrene
0	+	-	+	-	+	+
1	+	-	+	-	+	+
3	+	-	+	-	+	+
5	+	-	+	-	+	+
7	+	-	+	+	+	+

Biodegradation process of naphthalene and phenanthrene depends on the ability of microorganism to produce enzymes. It was assumed that there was an important role of *dioxygenase*, *hydrogenase*, and *monoxygenase* in biodegradation process of naphthalene and phenanthrene by *Bacillus subtilis* 3KP. Enzyme regulation in naphthalene and phenanthrene is affected by existence of gen. They were nah and phn which take the role in biodegradation of naphthalene and phenanthrene (Laurie and Jones, 1999). For this research, the existence of catabolic gen in *Bacillus subtilis* 3KP should be investigated as advanced research to affirm its ability of PAH degradation.

Moreover, hydrocarbon properties also affect biodegradation process. Complexity of structure, molecular weight, and solubility of the hydrocarbon become a limiting factor microbe's access to hydrocarbons. *Bacillus subtilis* 3KP was released biosurfactant to contact with the hydrocarbons. There was a similar growth pattern and uptake mechanism of substrate by *Bacillus subtilis* 3KP that indicated in Figure 2.



**FIGURE 2.** Mechanism of naphthalene (a) and phenanthrene (b) biodegradation by *Bacillus subtilis* 3KP



At the early stationary phase, emulsification activity and adherence activity increased, while surface tension reduced. Extracellular biosurfactant of *Bacillus subtilis* 3KP is able to reduce the surface tension of culture supernatant at the early stationary phase. It was in accordance to Cooper and Goldenberg (1987), the lysis of *Bacillus cereus* in sucrose was increasing the biosurfactant activity as a *surface active agent* because of its polysaccharide (emulsifier) and lipid reduced the surface tension. So that, at the early of stationary phase, emulsification activity increased together with the reducing of surface tension. It was assumed that *Bacillus subtilis* 3KP was secreted polypeptide-type biosurfactant.

Biosurfactant consists of surfactant monomer that would form stable micelle from about 10 – 200 molecules (Makkar and Rockne, 2003). Scippers et al. (2000) explained about the role of biosurfactant in biodegradation of PAH. The first role, micelle was used as a PAH porter to microbial cell wall surface, then PAH was entered diffusely the cell. The second role, micelle could increase PAH solubility in the medium. The last role, without micelle, surfactant still could facilitate the use of PAH by a cell-surfactant-PAH contact directly (bacterial adherence to hydrocarbons).

In pH 6 – 8, *Bacillus subtilis* 3KP are able to grow. Emulsification activity and surface tension reduction were not affected by increasing in pH. Therefore, at day 7 when the value of pH showed 8.0, emulsification activity was still rising. While in surface tension, there was not a significant difference.

Biosurfactant takes role in biodegradation process in accessing not soluble substrate. It was showed from the result that emulsification activity and surface tension reduction in culture with naphthalene was higher than with phenanthrene. However, the adherence activity was decreased after the end of exponential for culture with naphthalene. While in culture with phenanthrene, adherence activity has increased much higher at day 7. It was assumed that in culture with more complex structure, cells were adapted to transform its cell wall to be more hydrophobic. Prabhu and Phale (2003) have learned about uptake mechanism of aliphatic and aromatic hydrocarbons by *Pseudomonas* sp. strain PP2. Cell hydrophobicity raised after the decreased of emulsification activity in stationary phase. Level of cell hydrophobicity was higher in aromatic which is a lower soluble compound than aliphatic one.

At the same relatively environmental conditions and with the same type of microbe, phenanthrene degradation was longer than the naphthalene degradation. It proves that the type of substrate affects the biodegradation process. Although naphthalene and phenanthrene are classified as a some group of polycyclic aromatic hydrocarbons, naphthalene has much higher solubility than phenanthrene. Low substrate solubility in the medium can inhibit the bacterial access to the substrate. Limited of available substrate is the most important factor that involved in the slow degradation of PAHs (Makkar and Rockne, 2003).

## ACKNOWLEDGMENTS

The authors thank to the Direktorat Riset dan Pengabdian Masyarakat (DRPM) KemenristekDikti, Indonesia for funding and to Universitas Airlangga for the support of the facility during the research.

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