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3rd International Conference on Tropical and Coastal Region Eco Development 2017

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FOREWORD FROM THE CHAIR OF THE 3RD ICTCRED 2017



On behalf of the Organizing Committee, I would like to extend our warmest welcome to you at the International Conference on Tropical and Coastal Region Eco Development (ICTCRED) 2017. This annual conference is the third event after the second has been successfully conducted in 2016 at Bali. This conference is organized by Research and Community Services Institute (LPPM), Diponegoro University. The conference aims to provide a forum for researchers, academicians, professionals, and industries to expose and exchange innovative ideas, methods, and experience in the areas related to tropical life sciences and coastal development. This conference also provides forum for researchers

and scientists to exchange ideas and their current achievements.

In this year, 215 abstracts from various universities and research centers from many countries have been received. However, after in-depth review, only 147 high quality papers are accepted for oral and poster presentation in this conference. In addition, we cordially invite seven highly respected researchers in various fields as keynote speakers in this conference, to share their knowledge and expertise. I am grateful of each one of them for setting aside their valuable time to participate in this conference.

Moreover, I would like to announce that the ICTCRED 2017 Committee has signed an agreement with the Institute of Physics (IOP) to publish the conference proceeding in their Scopus-indexed *IOP Conference Series: Earth and Environmental Sciences (EES)* after a series of review. We do hope that the collaboration with IOP will increase the visibility of this conference papers to international levels which also give benefits to authors and also their institutions.

Finally the success of this conference lies not only in the quality of papers but also on the dedicated team efforts of the organizing committee. We thanks to the keynote speakers for the participation in this conference. I would like to acknowledge Institute of Physics (IOP) for the collaboration in publishing the conference proceedings. Indeed, I would like to thank the Scientific Committee members for their effort in reviewing and evaluating the papers for maintaining the quality of the conference. Last but not least, all staffs of The Research and Community Services Institute, Diponegoro University, deserves our great appreciation for their unlimited supports.

To all delegates, I hope that the 3rd ICTCRED 2017 event will be memorable not only from the scientific perspective but in the joy of meeting with other scientists for mutual collaboration. I wish you enjoy the conference as well as the beautiful nature and great traditions of Yogyakarta.

Chair, Organizing Committee of ICTCRED 2017

Dr. Agus Trianto



FOREWORD OF THE DIRECTOR

Dear distinguished speakers, delegates, ladies, and gentlemen

I am very pleased to welcome you all to this international conference, Tropical and Coastal Region Eco-development, which acts as a forum for those interested in tropical and coastal development issues. Diponegoro University commits to provide an opportunity for scientific society to always play important role in disseminating ideas and research results especially in the area of coastal and tropical development, which is the main research field of our university. Hence, this conference offers a platform for extensive sharing and exchange of knowledge for the development of coastal and tropical areas.

The topics presented in this conference cover marine biodiversity, sustainable marine utilization and development, climate change on coastal and marine ecosystems and coral reef ecosystems and coastal management. In the tropical field, this conference deals with relevant ideas and knowledge addressing vital life sciences, tropical health and nutrition, tropical diseases and tropical food and energy. In addition, this conference also covers socio-economic aspects such exploration of tropical rainforests, deforestation, rising immigration, etc. Thus, it is clear that the International Conference on Tropical and Coastal region eco-development is a unique blend of coastal and tropical that nicely fits the current interest among the community concerned with sustainable coastal and tropical ecosystems.

Finally, we would like to express our gratitude to our distinguished keynote speakers, Prof. Susilo Wibowo, Dr. Hadiyanto, Prof. Gerard Pals, Prof. Junichi Tanaka, Dr. Roel H. Bosma, Prof. Tao Liu, and Prof. Soottawat Benjakul, who had been traveling all the way to Yogyakarta. Certainly we will have an important benefit of preparing the next generation of Indonesian scientists with international exposure. We thank our participants to present their research papers, to share extensively and exchange of ideas thoughts and discussions so that this conference facilitates the formation of networks among participants. We thank all invited guests who have shown their interests in coastal and tropical region development field. Many thanks to the organizing and scientific committee of ICTCRED 2017 who have work very hard to run the conference.

I wish you all a productive and successful conference.

Yours sincerely Director of Research and Community Services Institute Diponegoro University

Prof. Heru Susanto

WELCOME ADDRESS OF THE RECTOR OF DIPONEGORO UNIVERSITY



It's a great pleasure and honour for our University to be the host of the 3rd International Conference on Tropical and Coastal Region Eco Development 2017 organized by Research and Community Service Institute, Diponegoro University. The special acknowledgement, I address to the distinguished speakers Prof. Susilo Wibowo and Dr. Hadiyanto from Diponegoro University-Indonesia, Prof. Gerard Pals from VU University Medical Center – Netherlands, Prof. Junichi Tanaka from University of the Ryukyus – Japan, Dr. Roel H. Bosma from Wageningen University – Netherlands, Prof. Tao Liu from Ocean University of China, and Prof. Soottawat Benjakul from Prince of Songkla University. Thank you for the valuable time to deliver knowledge and share scientific information at this

conference. I believe that this opportunity will provide the valuable information for us and deliberate some new research ideas for participants of this conference. For all the participants, I would also like to welcome you at this conference.

The origin of the conference theme is reflected from the idea of our Center of Excellence (CoE) which was established in 2012 representing our priority as a research university. Since the declaration of Diponegoro University as a research university, the main theme of every research result will be enhanced to the level of international benchmarking.

Diponegoro University, consists of 13 faculties, has strong human resources and research background related to the coastal development and tropical life sciences. It is also supported by integrated laboratory of marine and fisheries, which are located at Teluk Awur Jepara.

Coastal development and tropical life sciences are two important issues in Indonesia and need to be actualized by the government. The enormous potencies of Indonesia, large resources of marine area and their potential value of natural marine products extract, provide an opportunity to contribute for health energy and food. Recent issue has been arisen that the food and energy also can be exploited from the sea. Indeed, the exploration and exploitation of marine products must be considered on the impact of the environmental devastation. These issues are interesting topics which are reflected by large number of abstracts submitted to this conference. These interesting issues need to be discussed in this conference by sharing research finding and ideas. I am gratefull to see that this conference has enormous responds from the participants either from domestic or from other countries such Japan, China, US, and France, as reported by organizing committee.

Number of publication indexed by reputable database has been set as an indicator for world university rank including Indonesia. Therefore, Diponegoro University also encourages all scientists and academic staffs to increase their publication records in these international reputation journals. Currently, Diponegoro University is in the 6th position among universities in Indonesia for the number of

publications in reputable International journals. I sincerely express appreciation to the organizing committee for their effort to realize this conference.

By the end of my short welcome address, I hope our foreign guests take advantage of their stay here to enjoy Yogyakarta and its wonderful places. It is a beautiful and historical city to visit with a wonderful and unique traditional art dance, stunning sunset, great sceneries and interesting shopping.

Once again, it is my great pleasure to welcome you all to the 3rd International Conference on Tropical and Coastal Region Eco Development 2017. I wish you a pleasant two fully scientific days of conferences and I hope you can get a fruitfull share with other scientists on current developed knowledge and perhaps seeking for potential collaboration of your interested field.

Thank you for your kind attention.

Prof. Yos Johan Utama Rector

KEYNOTE SPEAKERS



Prof. Susilo Wibowo

Universitas Diponegoro – Indonesia "Indonesia got obese; do we care? Genetics, epigenetic, and environmental point of view"



Dr. Hadiyanto Universitas Diponegoro – Indonesia *"Effects of sugar addition on the thermal degradation of phycocyanin from Spirulina sp."*



Prof. Gerard Pals VU University Medical Center – Netherlands *"Cancer and the environment"*



Prof. Junichi Tanaka University of the Ryukyus – Japan *"Exploration of coral reef organisms for bioactive molecules and related issues"*



Dr. Roel H. Bosma Wageningen University& Research – Netherlands "Investing in climate change mitigation and adaptation on mangrove and aquaculture doubles benefits."



Prof. Tao Liu Ocean University of China - China *"Research on complete mitochondrial genome of marine algae"*



Prof. Soottawat Benjakul Prince of Songkla University *"Valorization of fish processing by product"*

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Table of contents

Volume 116

2018

◆ Previous issue Next issue ▶

3rd International Conference on Tropical and Coastal Region Eco Development 2017 2–4 October 2017, Yogyakarta, Indonesia

Accepted papers received: 31 January 2018 Published online: 08 March 2018

Open all abstracts

Preface

OPEN ACCESS			011001
3rd International	Conference on Trop	pical and Coastal Region Eco Development 2017	
+ Open abstract	View article	🔁 PDF	
OPEN ACCESS			011002
Note from Editor	S		
+ Open abstract	View article	🔁 PDF	
OPEN ACCESS			011003
Peer review state	ment		
+ Open abstract	Tiew article	🔁 PDF	
Papers			
OPEN ACCESS			012001
Social Cognitive	Predictors of Intere	st in Research Among Life Sciences Academics	
Dian R. Sawitri, Ha	arlina Nurtjahjanti and	Anggun R. Prasetyo	
+ Open abstract	View article	PDF	
OPEN ACCESS			012002
Library Develop	ment Strategy for T	he Community at Coastal Areas	
Putut Suharso, Ban	i Sudardi, Sahid Teguh	a Widodo and Sri Kusumo Habsari	

 \odot

	Tiew article	PDF	
OPEN ACCESS The Effects of Tr on The Resistant Banana (<i>Musa ba</i>	eatments on Batu B Starch, In Vitro Sta <i>Ilbisiana</i> Colla) Flo	anana Flour and Percentage of Wheat Substitution arch Digestibility Content and Palatability of Cookies M ur	012003 ade with
D Ratnasari, N Rus	tanti, F Arifan and DN	l Afifah	
+ Open abstract	View article	PDF	
OPEN ACCESS Stock Analysis o Central Java, Ind	f <i>Metapenaeus affir</i> onesia	uis (H.Milne Edwards, 1837) on the North Coast of	012004
Suradi Wijaya Sapu	tra, Anhar Solichin an	d Wiwiet Teguh Taufani	
+ Open abstract	Tiew article	🔁 PDF	
OPEN ACCESS Potential of L-fue compare to Carbo	cose isolated from I oxymethyl Cellulos	Brown Seaweeds as Promising Natural Emulsifier e (CMC)	012005
A N Al-Baarri, A M	I Legowo, Widayat, S	B M Abduh, F P Lestari, D Desnasari and I P M Santoso	
	View article	PDF	
OPEN ACCESS Copigmentation (Using Ferulic Ac	Of Anthocyanin Ex id And Tannic Acic	tract of Purple Sweet Potatoes (Ipomea Batatas L.) 1	012006
▲ Open abstract	View article		
+ Open abstract	[≡] view article		
OPEN ACCESS Growth Performa Density on the Bi	nce of Catfish (Cla iofloc System	rias Gariepinus Burchell, 1822) Cultured in High	012007
Fajar Basuki, Tristia	ana Yuniarti, Dicky Ha	arwanto and Titik Susilowati	
	Tiew article	🔁 PDF	
OPEN ACCESS Sex Diversity Ap Southern Waters	proach of Spiny Lo of Java	obster (Panulirus spp) to Marine Oil Spill Pollution in	012008
F E D Haryono, An	bariyanto and I Sulist	уо	
+ Open abstract	Tiew article	PDF	
OPEN ACCESS Effect of The Phy	ztase Enzyme Addit	tion in The Artificial Feed on Digestibility of Feed	012009
Feed Conversion Stadia I	Ratio and Growth	of Gift Tilapia Saline Fish (<i>Oreochromis niloticus</i>) Nurs	sery

Diana Rachmawati,	Istiyanto Samidjan and	d Tita Elfitasari	
+ Open abstract	Tiew article	🔁 PDF	
OPEN ACCESS Engineering Tech (Cromileptes Altiv	nology Of Fish Farı velis) Used Artificia	ming Floating Nets Cages On Polka Dot Grouper al Feed Enriched Phytase Enzyme	012010
Istiyanto Samidjan a	nd Diana Rachmawati	i	
+ Open abstract	View article	🔁 PDF	
OPEN ACCESS The Diversity of V <i>vannamei</i>) from E	Vibrios Associated v extensive Shrimp Po	with Vibriosis in Pacific White Shrimp (<i>Litopenaeus</i> ond in Kendal District, Indonesia	012011
Sarjito, Alfabetian H	larjuno Condro Hadito	omo, Desrina, Ali Djunaedi and Slamet Budi Prayitno	
	Tiew article	🔁 PDF	
OPEN ACCESS Factors Affecting	Husband Participat	ion in Antenatal Care Attendance and Delivery	012012
R Rumaseuw, S M E	Berliana, N Nursalam,	F Efendi, R Pradanie, P D Rachmawati and G E Aurizki	
	View article	🔁 PDF	
OPEN ACCESS Nitrate and Phosp Mangrove <i>Rhizop</i> E Supriyantini, A Sa	hate Contents on Se <i>hora</i> Sp. in Mangro ntoso and N Soenardje	ediments Related to The Density Levels of ove Park Waters of Pekalongan, Central Java o	012013
+ Open abstract	View article	▶ PDF	
OPEN ACCESS Analysis of the Su Annisa Nur Islami W Eka Misbahatul M H	urvival of Children Varrohmah, Sarni Man Ias, Elida Ulfiana and	Under Five in Indonesia and Associated Factors iar Berliana, Nursalam Nursalam, Ferry Efendi, Joni Haryanto, Sylvia Dwi Wahyuni	012014
+ Open abstract	View article	PDF	
OPEN ACCESS Formulation of Er (<i>Panicum miliace</i>) R B K Anandito, S F + Open abstract	nergency Food in B um) and Snakehead Kurniawan, E Nurha I View article	Biscuit-Form Made From Proso Millet Flour Fish (<i>Channa striata</i>) –Tempeh Flour Koya rtadi and Siswanti PDF	012015
OPEN ACCESS The Relation of E	nvironmental Quali	ty and Fishery Sector in Indonesia	012016
Shanty Oktavilia, Re	eikha Habibah Yusfi, F	irmansyah and FX Sugiyanto	
+ Open abstract	View article	PDF	

OPEN ACCESS	012017
Oceanographic Factors in Fishing Ground Location of Anchovy at Teluk Cenderawasih National Park, West Papua : Are These Factors Have an Effect of Whale Sharks Appearance Frequencies?	
Evi Nurul Ihsan, Siti Yasmina Enita, Kunarso and Anindya Wirasatriya	
+ Open abstract 🔄 View article 🏝 PDF	
OPEN ACCESS Probiotic Candidates from Fish Pond Water in Central Java Indonesia	012018
Alfabetian Harjuno Condro Haditomo, Desrina, Sarjito and S. Budi Prayitno	
← Open abstract	
OPEN ACCESS Flavor Enhancer From Catfish (<i>Clarias batrachus</i>) Bekasam Powder and Angiotensin-I- Converting Enzyme (ACE) Inhibitory Activity in Various Dishes	012019
+ Open abstract 🔄 View article 🎦 PDF	
OPEN ACCESS The Use of Different Diets for Feeding Rate and Growth of Shortfin Eel (<i>Anguilla bicolor bicolor</i>)	012020
N Taufiq-Spj, S Sunaryo, A Wirasatria, I Pratikto, D H Ismunarti and M I Syaputra	
← Open abstract	
OPEN ACCESS Impact of Monsoon to Aquatic Productivity and Fish Landing at Pesawaran Regency Waters	012021
Kunarso, Muhammad Zainuri, Raden Ario, Bayu Munandar and Harmon Prayogi	
+ Open abstract 🔄 View article 🏷 PDF	
OPEN ACCESS Business Profile of Boat Lift Net and Stationary Lift Net Fishing Gear in Morodemak Waters Central Java	012022
Trisnani D Hapsari, Bogi B Jayanto, Aristi D P Fitri and I Triarso	
+ Open abstract 🔄 View article 🏷 PDF	
OPEN ACCESS Impact of Fishery Policy on Fishery Manufacture Output, Economy and Welfare in Indonesia	012023
Firmansyah, Shanty Oktavilia, F.X. Sugiyanto and Ibnu N Hamzah	
+ Open abstract 🔄 View article 🄁 PDF	

OPEN ACCESS			012024
Cytotoxicity and F Bacteria	Phytochemical Prop	filing of Sargassum Sp. Extract As Anti-Mdr	012024
Wilis A. Setyati, Rin	i Pramesti, Muhamac	l Zainuddin, Maya Puspita and Person P Renta	
	Tiew article	PDF	
OPEN ACCESS			012025
Mapping of HABs	s Contaminated In	Green Shells (Perna viridis) in Semarang Bay	
Churun A'in, Suryan	ti Suryanti and Haeru	ıddin Haeruddin	
	Tiew article	🔁 PDF	
OPEN ACCESS		· T I C I D ·	012026
Nutritional Compo	Disition Changes Di	uring Tempeh Gembus Processing	
Ruth Nazaretha Sanc	lessy Damanik, Dwi	Yanti Winda Pratiwi, Nurmasari Widyastuti, Ninik Rustanti,	
Gemala Anjani and I	Diana Nur Afifah		
+ Open abstract	View article	🔁 PDF	
OPEN ACCESS			012027
Characterizations liquid smoke addition	of milkfish (<i>Chan</i> d tion	os chanos) meatballs as effect of nanoencapsulation	
Fronthea Swastawati	, Ambaryanto, Bamb	ang Cahyono, Ima Wijayanti and Diana Chilmawati	
	Tiew article	🔁 PDF	
OPEN ACCESS Clinical Outcome by Snakehead Fish	And Arginine Seru	um of Acute Ischemic Stroke Patients Supplemented	012028
Dwi Pudjonarko, Ret	tnaningsih and Zainal	l Abidin	
+ Open abstract	View article	🔁 PDF	
OPEN ACCESS			012029
Implementation of Semarang City, In	f Water Safety Plar donesia	ns (WSPs): A Case Study in the Coastal Area in	
Budiyono, P Ginand	jar, L D Saraswati, D	R Pangestuti, Martini and S P Jati	
	Tiew article	🔁 PDF	
OPEN ACCESS Effect of Different Microcapsules fro	t Coating Materials m <i>Caulerpa racen</i>	s on The Characteristics Of Chlorophyll	012030
R A Kurniasih, E N	Dewi and L Purnama	yati	
+ Open abstract	Tiew article	🔁 PDF	
OPEN ACCESS			012031

Effect of Melanin Free Ink Extracted From Squid (*Loligo* sp.) on Proximate and Sensory Characteristics of Soft-Bone Milkfish (*Chanos chanos*) During Storage

Tri Winarni Agustini, Hadiyanto, Ima Wijayanti, Ulfah Amalia and Soottawat Benjakul

OPEN ACCESS			012032
Nutrition Quality Vitamin D ₃ durin	v and Microbiology ng Storage	of Goat Milk Kefir Fortified with Vitamin B_{12} and	012002
EP Dianti, G Anjan	ii, DN Afifah, N Rusta	nti and B Panunggal	
+ Open abstract	View article	🔁 PDF	
OPEN ACCESS			012033
Total Lactic Acid Yoghurt with Bir	l Bacteria (LAB), A nahong Leaf Extract	ntioxidant Activity, and Acceptance of Synbiotic (Anredera cordifolia (Ten.) Steenis)	
R P Lestari, C Niss	a, D N Afifah, G Anja	ni and N Rustanti	
+ Open abstract	View article	PDF	
OPEN ACCESS The Effectivenes Used as Antibact	s of Heterotrophic l erial against Pathog	Bacteria Isolated from Dumai Marine Waters of Riau, gens in Fish Culture	012034
F Feliatra, Nursyirv	vani, A Tanjung, DS A	Adithiya, M Susanna and I Lukystyowati	
+ Open abstract	View article	🔁 PDF	
OPEN ACCESS Preliminary Stud <i>australis</i> from Ko Widianingsih Widia	y On Gonad Maturi enjeran Water, Sura aningsih, Muhammad	ity Stages of the Sea Cucumber <i>Paracaudina</i> baya, Indonesia Zaenuri, Sutrisno Anggoro, Hermin Pancasakti Kusumaningrur	012035 n and
Retno Hartati			
+ Open abstract	View article	🔁 PDF	
OPEN ACCESS EXPLORING TI City Pattern	HE POSITION OF	OLD SEMARANG SEA PORT: Based on Javanese	012036
R. Siti Rukayah, Er	ndang Sri Susilo, Muh	ammad Abdullah and Siddhi Saputro	
+ Open abstract	View article	🔁 PDF	
OPEN ACCESS Total lactic acid l red ginger extrac	bacteria, antioxidan t (<i>Zingiberofficinal</i> a	t activity, and acceptance of synbiotic yoghurt with <i>e var. rubrum</i>)	012037
B A Larasati, B Par	nunggal, D N Afifah, (G Anjani and N Rustanti	
+ Open abstract	View article	🔁 PDF	

OPEN ACCESS	1 2 4		012038
Edco-tourism; A Coasta	l Managemei	nt Program to Improve Social Economics	
Arsi Rakhmanissazly, Angg	gun Intan Perm	atasari and Ely Chandra Peranginangin	
+ Open abstract	iew article	PDF	
OPEN ACCESS			012039
Exploration of Sea Cucu Production of Marine Bi	mbers <i>Stiche</i> ological Res	opus hermanii from Karimunjawa Islands as ources	
Delianis Pringgenies, Siti R	udiyanti and E	rvia Yudiati	
+ Open abstract	iew article	🔁 PDF	
OPEN ACCESS			012040
Microbiological Charact Vitamin D ₃ Fortification	eristic and N Time	lutrition Quality of Goat Milk Kefir Based on	
F Fauziyyah, B Panunggal,	D N Afifah, N	Rustanti and G Anjani	
+ Open abstract	view article	🔁 PDF	
OPEN ACCESS Effect of Protein-Based Cooked Shrimp	Edible Coati	ng from Red Snapper (Lutjanus sp.) Surimi on	012041
I Rostini, B Ibrahim and W	Trilaksani		
+ Open abstract	iew article	🔁 PDF	
OPEN ACCESS The Environmentally So Metal of Lead (Pb) on So Tegal City	ound Aquacul eaweed of G	lture Strategies Based on Bioaccumulation of Heavy <i>racilaria verrucosa</i> on Aquaculture Areas of Muararej	012042 aVillage,
Nurjanah, Ambariyanto, Su	priharyono and	d Bambang Yulianto	
+ Open abstract	view article	PDF	
OPEN ACCESS Determination Hypoiodo	ous Acid (HI	O) By Peroxidase System Using Peroxidase Enzyme	012043
A N Al-Baarri, A M Legow	o, Widayat, S	B M Abduh, M Hadipernata, Wisnubroto, D K Ardianti, M N S	Susanto,
M Yusuf and E K Demasta			
+ Open abstract	view article	🔁 PDF	
OPEN ACCESS			012044
Antimicrobial activity of	t tempeh gem	<i>ibus</i> hydrolyzate	
A Noviana, F F Dieny, N R	ustanti, G Anja	ani and D N Afifah	
+ Open abstract	iew article	🔁 PDF	
OPEN ACCESS			012045

IOP Conference Series: Earth and Environmental Science, Volume 116, 2018 - IOPscience

The Abudance O Area	of Makrozoobenthos	s On Different Break Water In Semarang And Demak C	oastal
A Kristiningsih, D	N Sugianto, Munasik,	R Pribadi and J Suprijanto	
+ Open abstract	View article	🔁 PDF	
OPEN ACCESS			012046
Wastewater Trea	tment from Batik Ir	ndustries Using TiO2 Nanoparticles	
Fahmi Arifan, FS N	Nugraheni, Hafiz Rama	a Devara and Niken Elsa Lianandya	
	Uiew article	🔁 PDF	
OPEN ACCESS The Application Quality of Jelly (of Microencapsulat Candy	ed Phycocyanin as a Blue Natural Colorant to the	012047
E N Dewi, R A Kur	rniasih and L Purnama	ayati	
	Tiew article	🔁 PDF	
OPEN ACCESS Mapping of tropl in Jatibarang Res	hic states based on i servoir	nutrients concentration and phytoplankton abundance	012048
Siti Rudiyanti, Sutr	isno Anggoro and Ari	f Rahman	
	Tiew article	🔁 PDF	
OPEN ACCESS Characterization (terasi)	of Lactic Acid Bac	teria (LAB) isolated from Indonesian shrimp paste	012049
U Amalia, Sumardi	ianto and T W Agustir	ni	
	View article	🔁 PDF	
OPEN ACCESS Effect of Time La Based	enght Fermentation	to Katsuobushi Oxidation Rate As Fish Flavor	012050
U Amalia, L Rianir	ngsih and I Wijayanti		
	View article	PDF	
OPEN ACCESS Community Parti Karimunjawa Isla	icipation Of Coasta and	l Area On Management Of National Park,	01205
Bambang A Wibow	vo, Aryo B Aditomo a	nd Kukuh E Prihantoko	
	View article	🄁 PDF	
OPEN ACCESS Study of AUTO- in Semarang, Cer	LION (Automatic I ntral Java	Lighting <i>Rumpon</i>) on Fisheries of Stationary Lift Net	012052

S	Chairunnisa.	Ν	Setiawan.	Irkham	Κ	Ekawati	A	Anwar	and	А	DP	Fitri
0	Chan unniba	. T.A	Setta wan.	insnam	17	LINGWALL	, <i>i</i> i	1 MI W CI	anu	17	\mathbf{D}	TIUI

+ Open abstract	View article	🔁 PDF	
OPEN ACCESS			012053
Biodiversity of C Pocillopora sp. V	ryptofauna (Decapo /olume at Bunaken	ods) and Their Correlation with Dead Coral Island, North Sulawesi	
Muhammad Danie	Al Malik, Nenik Khol	ilah, Eka Maya Kurniasih, Andrianus Sembiring, Ni Putu Dian Pe	ertiwi,
Ambariyanto Amba	riyanto, Munasik Mu	nasik and Christopher Meyer	
+ Open abstract	Tiew article	PDF	
OPEN ACCESS			012054
Identification of <i>White Spots</i> Dise	Antipathogenic Bac	cterial Coral Symbionts Against <i>Porites Ulcerative</i>	
Nor Sa'adah, Agus	Sabdono and dan Diał	n Permata Wijayanti	
+ Open abstract	View article	PDF	
OPEN ACCESS			012055
Community Strue	cture of Decapod In	habit Dead Coral Pocillopora sp. in Pemuteran, Bali	
N P D Pertiwi, M D	A Malik, N Kholilah	, E M Kurniasih, A Sembiring, A W Anggoro, Ambariyanto and	
C Meyer			
+ Open abstract	Tiew article	🔁 PDF	
OPEN ACCESS			012056
Amino Acid Prof (Pangasius hypop	file and Volatile Fla <i>phthalmus</i>) and Nar	vour Compounds of Raw and Steamed Patin Catfish row-barred Spanish Mackerel (<i>Scomberomorus commers</i>	son)
Rusky I Pratama, I	Rostini and E Rochim	a	
	Tiew article	PDF	
OPEN ACCESS			012057
Anti-Pathogenic Dipsastraea from	Activity of Coral B Tengah Island, Kar	acteria Againts White Plaque Disease of Coral rimunjawa	
Sakti Imam Muchli	ssin, Agus Sabdono ar	nd Diah Permata W	
+ Open abstract	Tiew article	🔁 PDF	
OPEN ACCESS			012058
Seagrass Paramet	ter Affect the Fish A	Assemblages in Karimunjawa Archipelago	
Endang Sri Susilo, I	Denny Nugroho Sugia	into, Munasik, Nirwani and Chrisna Adhi Suryono	
+ Open abstract	Uiew article	🔁 PDF	
OPEN ACCESS	od for infants (0 (months old) in moduress results have done	012059
transcultural nurs	sing theory	months old) in madurese people based on	

Eka Mishbahatul M. Has, M. Syaltut, Tiyas Kusumaningrum and Ferry Efer	ndi
--	-----

	Tiew article	🔁 PDF	
OPEN ACCESS Genetic Diversity Lobophyllia corys	and Geographical <i>mbosa</i> in The Sulay	Gene Flow Patterns of Spawning Broadcast Coral wesi Waters as A Coral Triangle Area	012060
Widyastuti Umar, Ja	maluddin Jompa and	Asmi Citra Malina A.R. Tassakka	
+ Open abstract	View article	PDF	
OPEN ACCESS Vertical Distribut Western Pacific	ion of Temperature	in Transitional Season II and West Monsoon in	012061
Hikari A H Pranoto,	Kunarso and Endro S	Soeyanto	
	View article	🔁 PDF	
OPEN ACCESS The Quality of Ec with Addition of	lible Film Made fro Different Type Seav	om Nile Tilapia (<i>Oreochromis niloticus</i>) Skin Gelatin weed Hydrocolloid	012062
H Deanti, J M Hulu	, A. Setyaji, R N Eliya	anti, K Aliya and E N Dewi	
	View article	PDF	
OPEN ACCESS Effect of ENSO of	on the variability of	SST and Chlorophyll-a in Java Sea	012063
Anindya wirasatriyaOpen abstract	View article	PDF	
OPEN ACCESS Molecular Identif Deer Island, Raja	ication and Genetic Ampat, West Papu	e Diversity of <i>Acropora hyacinthus</i> from Boo and a	012065
 Open abstract 	View article	PDF	
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2020	IOP Conference	Series: Earth and Environmental Science, Volume 116, 2018 - IOPscience	
+ Open abstract	Tiew article	PDF	
OPEN ACCESS Inclusive blue sw Indonesia	imming crab fisher	y management initiative in Betahwalang Demak,	012068
A Ghofar, S Redjek	i, H Madduppa, M At	bbey and N Tasunar	
	Tiew article	PDF	
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L K Pinandita, I Rin	niatsih and I Irwani		
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R W Ariyati, S Reje	ki and R H Bosma		
	View article	PDF	
OPEN ACCESS			012072
Modelling Appro	ach	lying Nutrients (N and P) in Jepara Using Numerical	
Lilik Maslukah, Sri	Yulina Wulandari and	I Indra Budi Prasetyawan	
	Tiew article	PDF	
OPEN ACCESS Seasonal Variatio <i>affinis</i>) Catches in	ns of Oceanograph n the North Indram	ic Variables and Eastern Little Tuna (<i>Euthynnus</i> ayu Waters Java Sea	012073
Mega Syamsuddin,	Sunarto and Lintang	Yuliadi	
+ Open abstract	View article	PDF	
OPEN ACCESS			012074
The Influenced of Fermentation Pro Yogyakarta	f <i>Lactobacillus plan</i> cess on The Activit	<i>ntarum</i> Starter Addition and The Length Time of ty of Seaweed Antioxidant <i>Ulva lactuca</i> from Krakal Be	each,

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+ Open abstract	View article	🔁 PDF	
OPEN ACCESS Contribution of C Acids Contents in	Golden Apple Snail n Weaning Food	Flour to Enhance Omega- 3 and Omega-6 Fatty	012075
D D Marsyha, H S	Wijayanti, Nuryanto a	nd G Anjani	
	View article	🔁 PDF	
OPEN ACCESS Phycocyanin stab inlet temperature	oility in microcapsu	les processed by spray drying method using different	012076
L Purnamayati, EN	Dewi and R A Kurnia	asih	
	View article	PDF	
OPEN ACCESS Freshwater Clam Weaning Food to	s (<i>Pilsbryoconcha</i>) Overcome Stunting	<i>Exilis</i>) as an Potential Local Mineral Sources in g in Grobogan, Central Java, Indonesia	012077
S R Putri, Gemala	Anjani, Hartanti Sandi	Wijayanti and Nuryanto	
+ Open abstract	View article	PDF	
OPEN ACCESS Content Heavy M D Suprapto S Surv	Aetal Pb, Cd In <i>Peri</i> anti and N Latifah	na viridis And Sediments In Semarang Bay	012078
Open abstract	View article	PDF	
OPEN ACCESS Safely Intake Nu Semarang, Centra	mber of <i>Macridisci</i> al Java, Indonesia	us sp. (Kerang Ceplos) from Tambak Lorok Waters,	012079
Eduard Meirenno T	ielman, Jusup Suprija	nto and Ita Widowati	
	View article	🔁 PDF	
OPEN ACCESS Water Quality Im Cleaner Producti	nprovement of Med	ia Culture for Tilapia (Oreochromis niloticus) with	012080
Haeruddin, Supriha	ryono and S Febrianto		
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OPEN ACCESS Epidemiology of Center Semarang	Child Tuberculosis City, Indonesia)	s (A Cross-Sectional Study at Pulmonary Health	012081
L D Saraswati, P G	inandjar, B Widjanark	o and R A Puspitasari	

IOP Conference Series: Earth and Environmental Science, Volume 116, 2018 - IOPscience

+ Open abstract	View article	🔁 PDF	
ODEN ACCESS			012002
Vaccines Cold Ch	ain Monitoring: A	Cross Sectional Study at Three District In Indonesia	012082
L D Saraswati. P Gi	nandiar. Budivono. N	lartini. A Udivono and Kairul	
+ Open abstract	View article		
• Open abstract			
OPEN ACCESS			012083
The Evidence of	mposex in Turbo s	p. from Ujungpiring Waters of Jepara	
RAT Nuraini, R Har	tati, H Endrawati, Wi	dianingsih, MJ Rachma and RT Mahendrajaya	
+ Open abstract	View article	🔁 PDF	
OPEN ACCESS Antibacterial Act <i>Sydowii</i> on Four (ivity Symbiotic Fu Growth Medium	ngi of Marine Sponge Axinella sp., Aspergillus	012084
S Widyaningsih, A	Γrianto, OK Radjasa a	and K Wittriansyah	
	View article	🔁 PDF	
 OPEN ACCESS The Biophysical (olivacea) Eggs In Suryono, R Ario, E + Open abstract 	Characteristics Of I Turtle Conservatio Wibowo and G Hando	Hatching Habitat Of Lekang Turtle (<i>Lepidhochelys</i> on And Education Center, Bali oyo PDF	012085
Mapping of Nitra Massive Coral Re	te, Phospat And Zo ef in Karimunjawa	ooxanthelae With Abundance Of Sea Urchins on Island	012086
S Suryanti, C Ain ar	nd N Latifah		
	View article	PDF	
OPEN ACCESS The Assessment of Emas Semarang	of Biological and P	ollution Index of Estuaries Around Port of Tanjung	012087
A Tjahjono, O Wah	yuni and S Purwantini	i	
+ Open abstract	View article	PDF	
OPEN ACCESS			012088
Prospective Source associated with B	ce of Antimicrobial rown Alga (Phaeo	Compounds From Pigment Produced by Bacteria phyceae) Isolated from Karimunjawa island, Indonesia	
A T Lunggani, Y S	Darmanto, O K Radja	sa and A Sabdono	
	Tiew article	🔁 PDF	

OPEN ACCESS			012089
The Effect of Ferm	entation Time wit	h Probiotic Bacteria on Organic Fertilizer as	.1
Daphnia magna C Performance Enha	ultured Medium to ncement	wards Nutrient Quality, Biomass Production and Grow	'th
Vivi Endar Herawati,	Ristiawan Agung Nu	ugroho, Pinandoyo, YS Darmanto and Johannes Hutabarat	
+ Open abstract	View article		
+ Open abstract			
OPEN ACCESS			012090
Behind the Slow R Relatively High M	oad to Progress: A aternal Mortality i	ddressing Myriad Causes of the Persistence of n Brebes Regency after the Post EMAS Program	
Sri Kusumo Habsari,	Sofiah Sofiah and Su	imardiyono Sumardiyono	
+ Open abstract	Tiew article	🔁 PDF	
OPEN ACCESS			012091
Proximate content	of wild and culture	ed eel (Anguilla bicolor) in different part of body	
I Wijayanti and E S S	usilo		
+ Open abstract	View article	🔁 PDF	
OPEN ACCESS			012092
Determination of S by Resistivity Met	Soft Lithology Cau hods	ses The Land Subsidence in Coastal Semarang City	
Sugeng Widada, Sidh	i Saputra and Hariad	i	
+ Open abstract	View article	🔁 PDF	
OPEN ACCESS			012093
The Preliminary St Fishing Ground at	tudy of Organochlo Eastern Part of Co	orine Pesticide Residues on Sediments of Bivalvia pastal Semarang	
Chrisna Adhi Suryon	o, Subagyo, Wilis Ar	i Setyati, Endang Sri Susilo, Baskoro Rochaddi and	
Robertus Triaji Mahe	ndrajaya		
+ Open abstract	View article	🔁 PDF	
OPEN ACCESS Hierarchical Synth Marine Park	esis of Coastal Eco	osystem Health Indicators at Karimunjawa National	012094
Johan Danu Prasetya,	Ambariyanto, Supri	haryono and Frida Purwanti	
+ Open abstract	View article	🔁 PDF	
OPEN ACCESS Reef Development Java	on Artificial Patcl	h Reefs in Shallow Water of Panjang Island, Central	012095
Munasik, Sugiyanto,	Denny N Sugianto ar	nd Agus Sabdono	
+ Open abstract	Tiew article	🔁 PDF	

2020	IOP Conference	Series: Earth and Environmental Science, volume 116, 2018 - IOPScience	
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+ Open abstract	Uiew article	🎦 PDF	
OPEN ACCESS Effect of Tourist (Value Perceptions	Characteristic, Mari	ine Tourism Demand, and Number of Visits to the o Pay to Environmental Marine Tourism in Ambon City	012097
Renoldy L Papilaya			
	View article	PDF	
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Agus Trianto, Agus	Sabdono, Baskoro Ro	ochaddi, Desy Wulan Triningsih and Dewi Seswita Zilda	
	Tiew article	🔁 PDF	
OPEN ACCESS Preliminary Study Metal (Arsenic) in	y Contamination of n Shallow Groundw	Organochlorine Pesticide (Heptachlor) and Heavy vater Aquifer of Semarang Coastal Areas	012099
Baskoro Rochaddi,	Chrisna Adhi Suryonc	, Warsito Atmodjo and Alfi Satriadi	
	View article	PDF	
OPEN ACCESS Proliferative Acti Receiving Ethano	vity of Mammary C l Extract of Sponge	Carcinoma Cells by AgNOR Count in C3H mice e Haliclona sp	012100
Lanceria Sijabat, Ne	eni Susilaningsih, Agu	is Trianto and Retno Murwani	
	View article	PDF	
OPEN ACCESS Ethanol Extract o Adenocarcinoma	f <i>Haliclona</i> sp. Imp in C3H Mice	proved Histological Grade of Mammary	012101
R Murwani, A Triar	nto, E Wijayanti, A Rie	dlo and N Susilaningsih	
	Tiew article	PDF	
OPEN ACCESS Reduction of feca Steenis leaves pow	Il parasites by <i>Arecl</i> wder in laying hens	ha catechu L. seed and Anredera cordifolia (Ten)	012102
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OPEN ACCESS			012103

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OPEN ACCESS The Ethanolic Ex Carcinoma Grow	tracts The Gorgonia th In C3H Mice	n Isis hippuris Inhibited the Induced Mammary	012104
Agus Trianto, Yogi	Andriyas, Ali Ridlo, Si	ri Sedjati, Neni Susilaningsih and Retno Murwani	
	View article	PDF	
OPEN ACCESS Effectivity Test C Sub Unit Vaccine	of Crude Protein Spo To Prevent the Gol	ore of <i>Myxobolus koi</i> as Materials Development For d Fish (<i>Cyprinus carpio</i> , Linn) Dead by Myxobolusis	012105
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Cytotoxicity and Phytochemical Profiling of *Sargassum Sp.* Extract As Anti-Mdr Bacteria

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Cytotoxicity and Phytochemical Profiling of Sargassum Sp. **Extract As Anti-Mdr Bacteria**

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Abstract. Sargassum sp. contains bioactive compounds having the potential as an antibacterial agent. Sargassum sp. was collected from five different locations, i.e., Teluk Awur, Panjang Island, Bandengan, Ujung Piring, and Bondo. There were several different species of Sargassum sp. identified from each sampling locations. The collected seaweeds were washed, naturally dried, and ground to the powder-sized dry material. Dry seaweed was extracted gradually using n-hexane, ethyl acetate, and methanol (1:3 w/v). The antibacterial analysis was conducted based on Agar Diffusion method using Zobell as media. Resistance analysis was performed to evaluate the resistance of pathogen against commercial antibiotics, namely, chloramphenicol, ampicillin, erythromycin, amoxycillin, and tetracycline. Each Sargassum sp. extract was tested against three candidates of MDR bacteria, i.e., Staphylococcus aureus, Escherichia coli, and S. epidermidis. Results showed that S. aureus was resistant towards four out of five commercial antibiotics. E. coli and S. epidermidis were not susceptible to two and three out of five commercial antibiotics, respectively. N-hexane, ethyl acetate and methanol yielded 0.1-0.3%, 0.3-0.7 % and 0.8-4.7% of dry extract. Ethyl acetate extract of Sargassum from Teluk Awur performed the best antibacterial activity and contained an alkaloid, flavonoid, and phenolic compounds. Toxicity analysis showed that this ethyl acetate extract had LC₅₀ at 463 ppm and categorized as chronic toxicity.

1. Introduction

Applications of drugs particularly synthetic antibiotics have been commonly used to control the disease. The excessive use of antibiotics becomes the single most important factor leading to a resistance increase of bacteria towards antibiotics. This is due to the adaptability of disease to the toxicity of antibiotic and bacterial resistance [1]. Bioactive compounds from marine resources offer a promising solution to overcome the resistance problem. Also, it also becomes an alternative replacement of synthetic drugs to treat infection disease. World Health Organization (WHO) has recommended the use of natural products in maintaining the well being of the community, in preventing and resolving chronic and degenerative disease [2]. Sargassum sp. is one of the marine resources that can be proposed as alternative natural products for disease control. Sargassum sp. has been reported to show interesting and promising pharmacological properties serving as anticancer, anti-inflammatory, antibacterial, and antivirus [3], antioxidant [4], antifouling [5]. And antifungal [6]. S. fulvellum and S. thunbergii showed antipyretic, analgesic, and anti-inflammatory activities in mice [7]. S. echinocarpum, S. duplicatUM and S. polycystum performed potential antioxidant [8]. S. aquifolium, S. ilicifolium and S. polycystum [9]. showed antibacterial activity against Bacillus subtilis, Pseudomonas aeruginosa and Escherichia coli [9]. S. pallidum can serve as antioxidant and antihemolysis agents exhibiting potential for further exploration as functional food or complementary

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medicine [10]. S. tenerrimum from India is reported to have antibacterial activity against Klebsiella pneumonia, Escherichia coli, Staphylococcus aureus, Proteus sp., Streptococcus sp., Pseudomonas aeruginosa, Vibrio parahaemolyticus, Salmonella sp., Shewanella sp., Vibrio fluvialis, Vibrio splendidus, Vibrio cholera, Shigella flaxneri, Staphylococcus epidermidis, Aeromonas liquefaciens and Bacillus subtilus. Furthermore, Sargassum sp. also have antifungal activity against Aspergillus niger, A. flavus, A stetreus, Candida albicans and Penicillum sp. [11]. This study aims to evaluate the inhibition ability of Sargassum sp. extracts against pathogenic bacteria categorized as multidrug resistant bacteria. Using conservation and biotechnological approach, the study is expected to resolve the resistance of pathogenic bacteria towards synthetic drugs and to prevent more infections.

2. Materials And Methods

The samples *Sargassum sp*.were collected from five different locations, i.e. Teluk Awur (6° 7' 36.48" S, 110° 24' 0" E), Panjang Island (06° 34" 30' LS, 110° 37" 44' BT), Bandengan (°07' 36" LS 110° 24' 00" BT / 6,1268° LS 110,4° BT), Ujung Piring (6° 30' 40.91" S, 110° 40' 06.42" E) and Bondo (6° 28' 26.79" S, 110° 42' 33.13" E) at Jepara, Central Java.

The solvents used : n-hexane, ethyl acetate, and methanol as solvents to extract the bioactive compounds in *Sargassum sp.* Antibacterial activity against *S. aureus, E. coli*, and *S. epidermis* was evaluated through the Agar Diffusion method by using Zobell as media. The media used consisted of peptone, yeast, and agar [12].

2.1 Samples preparation

The collected seaweeds were washed with tap water to remove the epiphytes and the remaining debris. A whole thallus of seaweed was taken for taxonomic identification based on morphology study. Before natural drying under the shade, seaweeds were cut to pieces (\pm 5 cm). Further, the dried seaweeds were ground with a multi use blender to obtain powder material.

2.2 Extraction of Sargassum sp.

Dry seaweed of 25 gr was extracted gradually by using 100 mL n-hexane, ethyl acetate, and methanol, respectively. The extraction was performed for 24 h at room temperature in obscurity. The extracted seaweed was filtered by Whatman Filter Paper to separate the solute from the residues. The remaining residues of dry seaweed material were re-extracted using the same process and solvents. Filtrates obtained from the extraction process were pooled. N-hexane, ethyl acetate, and methanol filtrates were evaporated to dryness with vacuum rotary evaporator at 40 °C in 500 mmHg. n-hexane, ethyl acetate and methanol extracts of *Sargassum sp.* were dried using N₂ before storage in -20 °C [13]. Extraction yield was determined with following formula :

$$Ce = \left(\frac{W_2}{W_1}\right) \times 100\%$$

Where :

Ce = Extraction yield (%)

 W_2 = Weight of extract (gr)

 W_1 = Weight of initial sample (gr)

2.3 Resistance analysis of Multidrug-Resistant (MDR) bacteria

Staphylococcus aureus, Escherichia coli, and *S. epidermidis* were analyzed against five commercial antibiotics. The antibiotics were chloramphenicol, ampicillin, erythromycin, amoxicillin, and tetracycline.

2.4 Antibacterial activity analysis

Bacterial culture in liquid media was centrifuged at 5000 rpm for 10 min and was washed with PBS.

0.1 mL of bacterial solution was read with a spectrophotometer at 600 nm to obtain optical density between 0.6-0.8 [14].

MDR bacterial solution of 0.1 mL previously cultured in liquid media was spread evenly on to solid media in the petri dish using spread technique [14]. The inoculated bacterial solution was left for 30 min to diffuse on the media. Extract and chloramphenicol served as a positive control was prepared in distilled water at 2500 ppm. The sterile paper disc was impregnated with 20 μ L of antibiotics solution at 50 μ g/disc [13]. And 20 μ L of solvents without extracts as a negative control. Paper discs were placed on the solid media in Petri dish previously inoculated with MDR bacteria. All Petri dishes were incubated at 37 °C for 24-48 h.

2.5 Artemia salina Brine Shrimp Lethality Test (BSLT)

BSLT was performed to evaluate the toxic effect and cytotoxicity of chemical compounds against the larvae of shrimp *Artemia salina*. The test was based on [15], [16], [17]. *A. salina* was immerged in fresh water for 15-30 min to clean the attached feces. 0.25 gr of eggs were put in a box filled with 500 mL seawater at 25-30 °C with pH at 7-8. Irradiation from TL 40 watt light was placed above the box to light it during the hatching for 48 h. After 48 h, the eggs hatched to nauplii instar III/IV and were ready to use for BSLT test [18].

Ten larvae of *A. salina* were introduced into a test tube filled with extract showing the best antibacterial activity. Concentrations of extracts applied for the BLST were: 1000, 500, 100, 50 and 10 ppm with ethanol 2% [15]. Negative control used was methanol concentrated at 6, 4, 2 and 1%. The observation was made at 1, 3, 6, 12, 18, 24 and 36 h after the introduction of larvae into extract by using a loop. Determination of time interval was according to the lethal concentration of a compound whether it was acute or chronic lethal. LC_{50} was calculated from the linear regression equation based on the observation made. All tests were performed in triplicate.

2.6 Phytochemical profiling of Sargassum sp. extracts

This analysis was meant to examine the presence of certain chemical compounds in the tested samples, in this case, *Sargassum sp.* By determining a presence of certain compounds in plant matrix, a correlation can be made between the compounds and the biological activities performed. Thus it facilitates the phytopharmacological procedures [12]. Phytochemical profiling of *Sargassum sp.* extracts was carried out to evaluate the presence of alkaloid, flavonoid, saponin, steroid, and triterpenoid. *Sargassum sp.* dry material of 0.05 gr was dissolved in 10 drops of sulfuric acid 2 N, and three drops of this solution was introduced to Marquis, Dragendorff, Meyer and Wagner reagent. These reagents, previously prepared in separated test tubes, were designated to test the alkaloid content in a sample. For flavonoid, 0.05 g of dry seaweed material was added with 0.1 mg magnesium and 0.4 mL amyl alcohol, a mixture of HCl 37% and ethanol 95% in the same volume. Further, four mL of alcohol was added to the solution. To detect the presence of saponin, 0.05 g of dry seaweed material was dissolved in hot water. This test was also called as foam test due to the foam established on the surface of the solution. For the steroid/triterpenoid test, 0.05 g of dry seaweed material was dissolved in 2 mL of chloroform in test tube added with Lieberman Burchard reagent.

2.7 Statistical analysis

Data from resistance and antibacterial analysis were statistically analyzed using SPSS 16. Test of normality by Shapiro-Walk and homogeneity by Levene test with the significant level at 0.05 were performed. Multivariable analysis of variance (MANOVA) was used to analyze the interaction between variables, *i.e.*, sampling locations, *Sargassum sp.*, type of solvents and bacteria. Post-hoc test of Tuckey was applied to acknowledge the significant difference between tested variables (p<0.05).

3. Results And Discussion

Use of different solvents in extracting the bioactive compound from Sargassum sp. yielded dried extract differently. N-hexane resulted the lowest yield of extraction with $0.23 \pm 0.05\%$ of dry extract compared to ethyl acetate and methanol with $0.56 \pm 0.12\%$ and $2.18 \pm 1.23\%$ of dry extract, respectively. It indicates that the bioactive contained in Sargassum sp. tend to dissolve in polar solvents than in semi or no-polar one, implying that the extracted compounds are polar. Basic principle of extraction is the appropriate selection of solvents used to extract the solute. Non-polar solvents dissolve non-polar solutes while polar solvents dissolve polar solutes [19]. Bactericidal activity of Sargassum sp. extracts showed a diverse activity based on different locations. It might be suggested that there is a geographical variation in biological activity of Sargassum sp. Inhibition zone of extracts from Sargassum sp. collected at Bondo, Ujung Piring, and Bandengan were significantly different one to each other (p < 0.05). Also, these same samples were also significantly different to those from Panjang Island and Teluk Awur (p < 0.05). It is possible that these two stations, Panjang Island and Teluk Awur, have similarity in their water qualities. Furthermore, extracts of Sargassum sp. from Panjang Island and Teluk Awur showed the most active against the MDR bacteria than the other extracts. It was based on the biggest inhibition zone shown by the extracts from these two locations. It leads to an assumption that the environmental conditions in Panjang Island and Teluk Awur are favorable for the production of bioactive compounds.

Table 1 shows the statistical analysis of the antibacterial activity of *Sargassum sp.* based on different stations, species, and solvents. As can be seen from Table 1, antibacterial activity was best performed by *S. crassifolium* extracted with ethyl acetate. Moreover, samples from Panjang Island showed the best activity.

Variance	df	F	р	<i>Tukey p</i> <0.05
Station	4	103 1	Δ	Bondo ^a < Ujung piring ^b < Bandengan ^c < Teluk
Station 4 493.4	475.4	0	awur ^d < Pulau panjang ^d	
Species	6	99.65	0	Bin ^a < Echi ^b < Plag ^b < Cine ^b < Poly ^c < Dupli ^{cd} < Cras ^d
Solvent	2	4602	0	n-hexana ^a < Metanol ^b < Etil asetat ^c
Df. Januar	of fund	E = E	1-	and an analysister Dias C his day Cines C since and

Table 1. Analysis of variance from each variable analyzed

Df: degree of freedom; *F*: F value; *p*: probability; Bin: *S. binder*; Cine: *S. cinereum*; Cras: *S crassifolium*; Dupli: *S. duplicatum*; Echi: *S. echinocarpum*;Plag: *S .plagyophyllum*; Poly: *S. polycystum*; Sa: *S. aureus*. The order of variables was based on the smallest of the biggest average value (<); different letters indicate significant difference (p > 0.05).

Factors influencing the difference in inhibition zone from the antibacterial assay are the sensitivity of organisms, the media of culture, the condition of incubation, and the agar diffusion rate. Meanwhile, the agar diffusion rate depends on the concentration of tested samples, the composition of media, temperature and period of incubation [20]. Aside from these factors, it can also be suggested that different type of bioactive compounds extracted from *Sargassum sp.* play an important role in their bactericidal effects towards the MDR bacteria.

Type of solvents also contributes to the antibacterial effect of *Sargassum sp.* extract. Ethyl acetate and methanol extracts exhibited the highest inhibition zone; meanwhile, the n-hexane extracts had the smallest inhibition zone. Such difference might be due to different polarity of bioactive compounds from *Sargassum sp.* As non-polar solvents, n-hexane attracts non-polar compounds. As for semi-polar to polar compounds, they prefer to dissolve in ethyl acetate and methanol. Ethyl acetate is an aromatic compound characterized by its semipolar with $CH_3CH_2OC(O)CH_3$ as its chemical structure. This

compound can dissolve analytes tending to be semipolar and polar [21]. It indicates that the bioactive compounds extracted from *Sargassum sp.* are semipolar.

From 42 extracts tested, one best extract with a wide spectrum of antibacterial effect and the biggest inhibition zone was chosen. It was *S. duplicatum* from Teluk Awur extracted with ethyl acetate. This extract was further tested for its toxic effect using BSLT and results are presented in Table 2.

ta; S. duplicatum; ea							
Time				Correlation		LC ₅₀	
(hours)	$\mathbf{y} = \mathbf{a} + \mathbf{b}^* \mathbf{x}$	R ²	R	(%)	Concentration	Crit	orio
1					(ppiii)	CII	lella
1	-	-	-	-	-	-	-
	y = 0.017 x -					Not	
3	1.051	0.989	0.9945	99.45	3003	Toxic	-
	y = 0.025 x +					Not	
6	2.130	0.904	0.9508	95.08	1915	Toxic	-
	y = 0.034 x +					Not	
12	5.311	0.891	0.9439	94.39	1314	Toxic	-
	y = 0.036 x +					Not	
18	13.27	0.906	0.9518	95.18	1020	Toxic	-
	y = 0.082 x +						
24	12.04	0.946	0.9726	97.26	463	Toxic	Chronic
	y = 0.076 x +						
30	28.47	0.930	0.9644	96.44	283	Toxic	Chronic
	y = 0.143 x +						
36	30.42	0.915	0.9566	95.66	137	Toxic	Chronic

 Table 2. BSLT represented by the value of LC₅₀ of S. duplicatum in different time interval

Toxicity of *S. duplicatum* extracted in ethyl acetate was evaluated towards the larvae of *A. salina* through the BSLT method. It seems that the toxicity of this extract started 24 h after introducing the larvae of *A. salina* to ethyl acetate extract of *S. duplicatum*. Value of LC₅₀ after 24 h was 463 ppm implying that this concentration could cause lethal effect towards 50% of the larvae population.

Table 2 displays that ethyl acetate extract *S. duplicatum* is considered as active or toxic because of its ability to cause 50% of mortality to the tested organisms with less than 1000 ppm [23]. Based on the category of toxicity from a compound (15). It generates toxic effect during the time interval 18 h. BSLT method is widely used as a method to evaluate a toxicity of natural product as mentioned by several studies [22]; [23]. The advantages of this method are the simplicity. The high reproductivity (24 h). cheap and can be used to determine the intensity of cytotoxicity.

Table 3. Phytochemical profiling of S. duplicatum extracted with ethyl acetate

Phytochemical profile	S. duplicatum	Standard (color)
Alkaloid		
Dragendorf	+	Red to orange
Meyer	+	Murky white
Flavonoid	+	Layer of amyl alcohol is red/yellow/green

Steroid	-	Green/blue
Phenolic	+	Green or greenish blue
Triterpenoid	-	Violet or purple
Saponin	-	Foaming on top of the layer
Tanin	-	Greenish brown or dark blue (bluish black)

Table 3 shows that the ethyl acetate extract of *S. duplicatum* contained alkaloid. Flavonoid and phenolic compounds. These compounds are assumed to be responsible for the antibacterial effect of *S. duplicatum* against *S.aureus*. *E. coli* and *S. epidermidis*. Inhibition of bacterial growth in the presence of antibacterial agents is provoked by the combination of these agents with bioactive compounds. These bioactive compounds might be from the secondary metabolites such as alkaloids. Peptides. Terpenes. Pigments and sterols [24]. Mechanisms of antibacterial from a compound can take place in several ways, which are : 1) the destruction of bacteria cell wall; 2) the change of permeability in a membrane cell and 3) prevention of protein and nucleic acid synthesis. Furthermore, concentration of extracts and bacteria species of bacteria the availability of organic material, temperature, and pH have been suggested as factors affecting the antibacterial effect of a compound [25].

Alkaloid serves as antibacterial agents and provokes the death of bacteria cell due to the presence of alkali groups containing nitrogen in its structure [26]. It leads to a change in amino acid structure and the genetic balance experiences a lysis. Moreover, flavonoid could obstruct the growth of bacteria by destroying the permeability of bacteria cell wall. Flavonoid can release transduction energy to the cytoplasmic membrane of bacteria and to inhibit the motility of bacteria. Hydroxyl group in flavonoid contributes to the alteration of the organic component and transport of nutrition generating a toxic effect towards the bacteria. Phenolic compounds also have a hydroxyl group in their chemical structure. The mechanism to eliminate the bacteria contain more lipids, then higher concentration is needed to rupture the bacteria cell. The mechanism of phenolic activity as an antimicrobial is the ability of phenolic to cause shrinkage. also known as astringent [27]. Phenols can form a complex with a microbial enzyme to reach the membrane cell of bacteria through the cell wall. Bacteria cell wall consists of different polysaccharides and proteins; thus. it allows part of phenols to penetrate the cell wall.

4. Conclusion

The present study reveals that *Sargassum* sp. extract can inhibit the growth MDR bacteria. It implies that *Sargassum* sp. contains bioactive compounds having the property as an antibacterial agent. of this extract. Results obtained unlocks the potential of *Sargassum* to replace commercial antibiotic against the MDR bacteria. Nevertheless. it is important to further consider the type of solvent used to extract the target compounds that are active against the MDR bacteria. Apparently. certain solvents might have a negative impact towards the living cell. as presented in this study. Eventhough ethyl acetate extract of *Sargassum* sp. showed to be the most effective against MDR bacteria, this extract was also toxic. As a consequence, it is crucial to perform further research related to the application of a more eco-friendly method to extract the target compound from *Sargassum* sp.

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M513