# Toxicity Test for the Extract of Symbiont Bacteria Bacillus sp. as Anti-bacteria

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### **RESEARCH ARTICLE**

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## Toxicity Test for the Extract of Symbiont Bacteria Bacillus sp. as Anti-bacteria

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### Abstract

Increased antibiotic resistance spurs exploration of bioactive compounds as new antibiotic alternatives. *Bacillus sp.* is a symbiont bacterium which is a marine microorganism that has the potential to produce new bioactive compounds that can be developed as new antibiotics. This research is an experimental study aimed at identifying bioactive 33 pounds by thin layer chromatography methods and testing the activity of bioactive compounds by probit analysis EPA probit analysis program version 1.5. in *Artemia salina Leach*. Bioactive compounds identified were compounded from the alkaloid group with the category of highly toxic several-irritating base on 28 EPA toxicity category and the highly hazardous base on The WHO toxicity category based on the environmental protection agency probit analysis program used for calculating LC / EC value of LC 50 = 169,520 g / I. Symbiont *Bacillus sp.* produce secondary metabolites in the form of bioactive which have the ability as anti-bacteria. As a new antibiotic alternative to overcome resistance, especially in methicillin-resistant bacteria

**Keyword:** Bacillus sp., Artemia salina Leach, secondary metabolites, bioactive, highly toxic several – irritating, symbiont, highly hazardous, brine shrimp lethality toxicity.

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### INTRODUCTION

Indonesia as a maritime country has abundant biodiversity, especially the marine wealth that is both microorganisms and macroorganism. Macro-rich marine assets include various types of marine flora and fauna, while micro-species are various types of algae, yeast and marine bacteria, which have the potential to develop natural product materials<sup>1</sup>. As biotechnology develops, the utilization of marine riches and marine biodiversity as natural producting redients starts to be utilized in the field of pharmacology, especially in the field of finding new alternative antibiotics derived from the results of the isolation of secondary metabolites (bioactive compounds) produced by marine organism associations. Research on secondary metabolite compounds produced by symbiont bacteria associated with soft corals in the sea is the development of research in alternative new antibiotic discoveries. The isolated compounds were then tested for toxicity, to determine the effectiveness of secondary metabolites (bioactive) against pathogenic bacteria<sup>2</sup>. Toxicity tests for secondary metabolites can be tested by various 44 ethods, among others by the Brine Shirmps Lethality Test Method. Brine Shrimp Lethality Test Method is a method for determining the toxic nature of a secondary metabolite compound produced by certain organisms in the Arthemia salina test organism, if the secondary metabolite compound has biological activity, then the secondary metabolite is said to be a new alternative antibiotic bioactive compound3.

The symbiont bacteria are a community of bacteria that live in association with other biota, especially soft corals, hard corals and sponges in a variety of interaction patterns. In accordance with the characteristics of the two, the specific interactions between symbiont and host allow the potential for the same secondary metabolite product to occur, therefore drug development from bioactive compounds produced by soft corals, hard corals and sponges is more likely to be isolated from symbiont bacteria. Marine symbiont bacteria that are symbiotic with soft corals are alternatives that are more likely to be developed as sources of bioactive substances. Bacteriological research is easier and cheaper to rry out than research on high-level biota, this is because the breeding and isolation of marine bacteria are easier than isolation from high-level biota sources<sup>4</sup>.

Pressure conditions in the marine environment are greater than the terrestrial environment causing marine organisms to be more adaptive to extreme environments, so it is possible that the metabolite compounds produced will be better<sup>5</sup>. The various studies report on symbiont bacteria that live in a symbiotic interaction with soft corals and able to produce certain anti-bacterial compounds, namely the discovery of bacteria that live in symbiosis with hard corals, soft corals and sponges have the ability to inhibit *Escherichia coli*, *Streptococcus aureus*, *Streptococcus sp.* and *Aeromonas hidrophyla*<sup>6</sup>.

The soft coral Sarcophyton sp. is one type of soft coral that produces natural chemical compounds and is known as a natural product. These natural chemical compounds have the potential as a source of natural medicine. Active chemical compounds found in soft corals Sarcophyton sp. exhibits antibacterial, antifungal, antitumor, neurotoxic, and anti-inflammatory activities that are beneficial to the pharmaceutical industry<sup>7</sup>. This form of symbiotic interaction can stimulate the formation of bioactive compounds in symbiont organisms. Bioactive compounds from the marine environment have many unique chemical structures not found in terrestrial environments, in addition, bioactive compounds from the marine environment are potential agents as new antibiotic drug ingredients. Bacteria Bacillus sp. is one of the microorganisms that live in symbiosis with soft coral Sarcophyton sp. and several \$43 ies of Bacillus sp. known to be active against Methicillin-Resistant Staphylococcus aureus and resistant to Vancomycin-resistant Enterococcus and produce bacteriocin - type antibiotics8. In connection with this, it is necessary to conduct research on the symbiont bacteria Bacillus sp. from soft coral Sarcophyton sp., to determine the level of toxicity of secondary metabolites that it produces against the Artemia salina Leach test animals with the brine shrimp lethality test methods9.

### MATERIALS AND METHODS

Toxicity test for bioactive extract of Bacillus sp.

a) Experimental research with a post

37t-only control group design approach<sup>10</sup>, The aim of this study was to examine the potential toxicity of extracts of *Bacillus sp.* against *Artemia salina Leach* larvae. Potential toxicity of extracts of *Bacillus sp.* on *Artemia salina* Leach larvae was declared toxic if the *LC* value <50 1000µg/ml after an acute toxicity test was performed<sup>11</sup>.

b) Standard mortality indicator for Artemia salina Leach larvae if Artemia salina Leach larvae do not show movement for several seconds after observation.

c) Bacillus sp. bacteria used is a collection of the Faculty of Public Health, University of Diponegoro which is a symbiont soft coral bacterium

### Identification of bioactive compounds

Identification of Bioactive Compounds by separating the chemical content of the most active fractions from the partition results is done by thin layer chromatography<sup>12</sup>.

### Antimicrobial activity testing

Anti-microbial activity was measured in vitro to determine the potential of antibacterial substances and the sensitivity of a bacterium to the concentration of the test mat 5 al used, then analyzed by probit analysis of the EPA Probit Analysis Program version 1.5 (Used for calculating LC/EC values)<sup>13</sup>.

### RESULTS

Based on research conducted to obtain data as follows:

Brine ship lethality toxicity test14

The number of Artemia salina Leach

larvae mortality in each test tube at various concentrations of extracts of *Bacillus sp.* (Ta42: 1). Observation results show different effects on the death of 6 temia salina leach larvae, as follows:

The number of Artemia sali 27 Leach larvae in each test tube is 30 so that the total number of Artemia salina Leach larvae use 27 \$ 180 larvae, carried out with 3 replications. The total number of Artemia salina Leach larvae that died in each treatment tube was counted, volume the average death of Artemia salina Leach was obtained by dividing the total larvae mortality at each concentration by the number of replications performed. Then the percentage of larvae deaths were calculated from the average death at each concentration.

Toxicity Test Results of Bacillus sp. ex 30: against Artemia salina leach based on the EPA Probit analysis program use for caculating LC/EC Values Version 1.5, as follows (Table 2)

- 1. Extract of *Bacillus sp.* with a concentration of 0 ppm in 10 *Artemia salina Leach* test animals, there is 1 *Artemia salina Leach* test animal that responds (dies), with proportion responding = 0.0300 (3%).
- 2. Extract of *Bacillus sp.* with a concentration of 10 ppm in 10 *Artemia salina Leach* test animals, there are 2 *Artemia salina Leach* test animals that respond (die), with proportion responding = 0.2000 (20%)
- 3. Extract of *Bacillus sp.* with a concentration of 100 ppm in 10 *Artemia salina Leach* test animals, there were 4 *Arthemia salina Leach* test animals that responded (died), with proportion responding

**Table 1.** Test Results for the Brine Shrimp Lethality Test extract of *Bacillus sp.* in *Artemia salina leach* larvae at 48 hours

No.	Concen.	U	1	ι	J2	U3		Aver	rage	%
		+	-	+	-	+	-	+	-	
21										
1	1000 ppm		10		10		10	0	10	100%
2	500 ppm		10		10		10	0	10	100%
3	250 ppm	1	9	5	5	5	5	3,67	6,33	63%
4	100 ppm	7	3	6	4	4	6	5,56	4,33	43%
5	10 ppm	4	6	10		10		8	2	20%
6	Control	10		10		9	1	10	0,33	0%

Information:

+ : Life
- : Dead
\*\* : Percentage

Table 2. Probit Test Toxicity Test Extract of Bacillus sp. against Artemia salina Leach

			Propo	ortion		
Observed			Resp	onding	Predicted	
Number	Numbe	er	Proportion responding	Adjusted for control	Proportion responding	
Control	10	0	0.003	0	0.1241	
10,000	10	2	0.2	0.0866	0.0001	
100,000	10	4	0.533	0.3526	0.2365	
250,000	10	6	0.567	0.5056	0.7013	
500,000	10	8	1,000	1.00	0.9293	
1,000,000	10	10	1,000	1.00	0.9921	
8   - Square for Chi - Square for	-	neity (ta	ilculated) bular value at 0.0	= 3.246 5 levels) = 7.815 = 2.2292 = 0.3194		
Parameter	Es	timate	Std. Err.	95% Confidence	Limits	
Intercept	-1.	978474	2.543889	(-6.964497, 3.00	07549)	
Slope	3.3	130453	1.063219	(1.046543, 5.2	14363)	
Spontaneous Response Rate	0.1	124136	0.073508	(-0.019939, 0.20	68211)	

= 0.4330 (43.3%)

- 4. Extract of Bacillus sp. with a concentration of 250 ppm in 10 Artemia salina Leach test animals, there were 6 Artemia salina Leach test animals that responded (died), with proportion responding = 0.5670 (56.7%).
- 5. Extracts of *Bacillus sp.* with a concentration of 500 ppm in 10 *Artemia salina Leach* test animals, there are 10 *Artemia salina Leach* test animals that respond (die), with proportion responding = 1 (100%).
- 6. Extract of Bacillus sp. with a concentration of 1000 ppm in 10 Arthemia salina Leach test animals, there are 10 Artemia salina Leach test animals that respond (die), with proportion responding = 1 (100%).

Proportion responding is the proportion of test animals that respond to active compounds which are described in terms of a percentage. The higher the proportion of proportion responding, the greater the test animals that die due to the active compounds exposed. Chi-8 uare for Heterogeneity (calculated) = 3,246 and Chi-Square for Heterogeneity (tabular value at 0.05 levels) = 7,815 so the research is said to be homogeneous because Chi-Square for Heterogeneity (calculated)

= 4,847 < Chi-Square for heterogeneity (calculated) tabular value at 0.05 level = 7.815). Concentrations of potentially toxic substances in environmental media that cause death after a certain period of exposure are denoted by LC.  $LD_{50}$  is a statistic that is derived statistically, to express a single dose of a compound that is thought to be deadly or cause significant toxic effects in 50% of experimental animals after treatment.  $LD_{so}$  is a quantitative benchmark 17t is often used to express the lethal dose range. In general, the smaller the LD<sub>so</sub> value, the more toxic the compound is and the greater the LD value, the lower the 5 xicity. The results of probit analysis using the EPA Probit Analysis Program Version 1.5 (Used For Calculating LC/EC Values) show LC values of extracts of Bacillus sp. is

**Table 3.** Results of  $LC_{so}$  analysis of secondary metabolite compounds extracts of Bacillus sp. against Artemia salina Leach for 48 hours

Expo	sure	95 % Co Lim		
Point	Conc	Lower	Upper	
LC/EC <sub>50</sub>	169.52	58.337	269.048	

Table 4. EPA Toxicity Category<sup>17</sup>

Hazard		<b>Toxicity Category</b>		
Indicator	1	2	3	4
Oral <i>LD<sub>50</sub></i>	Up to end including	From 50 thru	From 500 thru	Greater than
	50 mg/kg	400 mg/kg	5000 mg/kg	500 mg/kg
Inhalation LC_	Up to end including	From 0,2 thru	From 2 thru	Greater than
30	0,2 mg/L	2 mg/L	20 mg/L	20 mg/L
Dermal <i>LD<sub>50</sub></i>	Up to end including	From 200 thru	From 2200 thru	Greater than
30	200 mg/kg	2000 mg/kg	20000 mg/kg	20000 mg/Kg
ye Effect	Corrosive corneal	Corneal opacity	No corneal opacity	No irritation
	opacity not reversible	severe irritation	irritation reversible	
	within 7 days	at 7 days	within 7 days	
Skin effect	Corrosive		Moderate irritation	Mild or slight
			at 72 hours	irritation
				at 72 hours

Source: 40 CFR 156,62

169,520 g/l. this refers to the bioactive compounds produced by *Bacillus sp.* categorized as follows:

- Highly toxic several dermal irritating according to EPA toxicity category standards<sup>15</sup>.
- 2. Highly hazardous based on WHO toxicity category<sup>16</sup>.

Output data from the results of probit analysis can be seen in appendix 1, while group toxicity based on EPA toxicity category, WHO toxicity category and Loomis toxicity category, as follows (Table 4)

Toxicity category 1: Highly toxic; severely irritating.

Toxicity category 2: Moderately toxic; moderately irritating.

Toxicity category 3: Slightly toxic; slightly irritating.

Toxicity category 4: Practically non-toxic; not an irritant.

To assign a signal word, use the highest

hazard shown by any of the indicators for the product

Danger – category 1. In addition, if the product is in Category 1 because of its oral  $LD_{50'}$  inhalation  $LC_{50'}$  or Dermal  $LD_{50'}$  the word "Poison" along with a skull and crossbones will be on the label

Warning – category 2 Caution – category 3 or 4

# Identification of dots using thin layer chromatography

Based on the identification results of the observation point extract of *Bacillus sp.* by Thin Layer Chromatography method, the following results are obtained:

Based on the results of qualitative analysis, secondary metabolites can be identified based on Rf (retardation factor) values follows:

The extraction toxicity test of Bacillus sp. was performed using the Artemia salina Leach

Table 5. WHO toxicity category<sup>16</sup>

	39					
Class	Class LD 50 for the rate (mg/kg body wight)					
		Oral				
	Solid	Liqiud	Solid	Liqiud		
-3						
1a. Extremely hazardous	5 or less	20 or less	10 or less	40 or less		
1b. Highy hazardous	May-50	20-200	10-100	40-400		
II. Moderateely hazardous	50-500	200-2000	100-1000	400-4000		
III. Slightly hazardous	over '500	over '2000	over '1000	over '4000		
2						

test using the Brine Shrimp Lethality method. Test of a bioactive compound is stated to have the acute toxic ability if it is able to kill 50% or more of the test animal population in a short interval of time. Based on research results that the bioactive compounds produced are alkaloids with  $LC_{c}$  169,520 (concentration = g / I) based on the EPA probit analysis program version 1.5 - used for calculating LC / EC values). The bioactive compounds are categorized as highly toxic several - irritating (EPA toxicity category) and highly hazardous (WHO toxicity category)17. The method of identification of bioactive compounds uses the ultra violet ray irradiation method with a wavelength of 254 nm and the thin layer chromatography method. The identification results show bioactive compounds with R, value = 0.8720.975, chromatogram dot brown color, bioactive compounds bound to polar compounds. Criteria for these compounds are included in the category of alkaloids<sup>12</sup>.

Toxicity test using the brine shrimp letality test with Artemia salina Leach test animals. This method was chosen because it is easy to implement, does not require a large cost, can be done in a short time and is easy to analyze. Artemia salina Leach is one of the widely used test animals because Artemia salina Leach has a short life cycle, has the ability to adap 260 high salinity and extreme temperatures, has a short life cycle, high adaptability to extreme environmental conditions, small body size and body organs which is simple and has a simple cell wall 18. Exposure to toxicity of bioactive compounds alkaloids to the

Table 6. Identification of nodes using Thin Layer Chromatography<sup>12</sup>

No	Chromatography Thin Layer Samples	Rf-value	Staining	UV 235 nm
1	Thin Layer Chromatography 1 The Bacillus sp. extract is bottled without concentrating with chloroform eluent	Negative	Negative	Negative
2	Thin Layer Chromatography 2 The Bacillus sp. extract is bottled by concentrating with chloroform eluent	Negative	Chocolate	Positive
3	Thin Layer Chromatography 3 The Bacillus sp. extract was concentrated dissolve in ether bottled without concentration with chloroform eluent without saturation	0.972	Chocolate	Positive
4	Thin Layer Chromatography 4 The Bacillus sp. extract concentrated dissolved in acetone bottled without concentrating with chloroform eluent without saturation	0.9	Chocolate	Positive
5	Thin Layer Chromatography 5 Extract of <i>Bacillus</i> sp. concentrated dissolved in hexan bottled without concentrating with chloroform eluent without saturation	0.875	Chocolate	Positive
6	Thin Layer Chromatography 6 The Extract of Bacillus sp. concentrated dissolved in ether bottled without concentrating with chloroform eluent without saturation	0.91	Chocolate	Positive
7	Thin Layer Chromatography 7 The extract of <i>Bacillus</i> sp. concentrated dissolved it in aceton is bottled without concentrated with chloroform eluents without saturation	0.9	Chocolate	Positive

Table 7. Secondary Metabolite Screening Results of Bacillus sp. with thin-layer chromatography<sup>12</sup>

No.	Туре	Value (Rf)		Chromato	gram Color	
	Identification	Retardation	Reactor	UV 30	65 nm	DPPH
		factor value		before	after	method
1	Alkaloids	0.8	Brown	-	-	Yellow
		0.87		Light blue	Light blue	
				(flouresens)	(flouresens)	
2	Flavonoids	0.13	Yellow Brown	-	-	Yellow
		0.72	Yellow Brown	-	-	Yellow
		0.78		Light blue	Blue	
				(flouresens)	(flouresens)	
3	Polyfenol	0.41	Black	-	-	-
		0.84		Light blue	-	-
				(flouresens)		
4	Terpenoid/ Steroid	0.06	Blackish purple	Light blue (flouresens)	-	-
		0.16	Reddish purple	-	-	-
		0.24	Dark purple	Light blue	-	-
				(flouresens)		
		0.37	Purple	Light blue	-	-
			•	(flouresens)		
		0.74	Purple	-	-	-

exoskeleton wall Artemia salina causes cell wall damage Artemia salina Leach19. Artemia Salina Leach is an osmoregulator type organism so that Artemia salina Leach will continue to ingest the surrounding media both toxic and non-toxic. with this osmoregulation system, alkaloids as secondary matabolites (bioactive) produced by Bacillus sp. in the Br 24 Shrimp Letality Test are toxic and can easily enter the body of Artemia salina Leach and cause death<sup>20</sup>. A 24 oids have the potential for acute toxicity and can cause larval death of Artemia salina Leach. The mechanism of larval death is related to the function of alkaloid compounds which can inhibit the eating power of larvae (antifeedants) which are stomach poisoning<sup>21</sup>. Bioactive components alkaloids cause also disruption of enzymatic function. Enzymes cannot work because of inhibition both competitive and non-competitive by alkaloids, this causes inhibition of metabolic processes and cellular respiration, causing death of Arthemia salina Leach<sup>17</sup>.

### CONCLUSION

Extract of the symbiont *Bacillus sp.* which is associated with rarely soft produce biocative secondary metabolites which have the ability as anti-bacteria. The resulting bioactives have the potential as an alternative alternative to new antibiotics in overcoming antibiotic resistance, so that further research is needed, especially in the development of research on the scale of the application of alkaloids.

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### AUTHORS' CON12 IBUTION 12

This research was conducted in collaboration between the two authors namely SI and S. The SI writer conducted the research design, analyzing the results of research, writing a draft of the initial script. Authors SI and S manage

the research analysis. Author S manages the search literature and makes final draft corrections. Both 19 hors have read and agreed to the final draft of this article.

### CONFLICT OF INTEREST

The authors declare that there is no conflict of interest.

### FUNDING

None.

### **AVAILABILITY OF DATA**

All datasets generated or analyzed during this study are included in the manuscript.

### **ETHICS STATEMENT**

Not applicable.

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