

# Efficacy of Biopesticide Formula containing *Streptomyces* sp. and *Trichoderma* sp. Against Southern Green Stink Bug (*Nezara viridula*) on Soybean (*Glycine max* L.)

## ABSTRACT

*Streptomyces* sp. and *Trichoderma* sp. are soil microorganisms isolated from shallot fields that can act as biological agents and increase crop production. *Nezara viridula*, the southern green stink bug, is the leading pest of soybean during the generative period, which can cause damage up to 80%. This study aimed to determine the efficacy of a liquid biopesticide formula using a mixture of coconut water and potato extract containing *Streptomyces* sp. and *Trichoderma* sp. This study used a randomized block design. The first factor was the time of application and the second factor was the concentration level. There were 8 treatment combinations and 2 controls. Each treatment combination was repeated three times. Probit  $LC_{50}$  and  $LT_{50}$  were performed to determine the effectiveness of biopesticides. The calculation of probit analysis obtained results of 84,443 ppm or about 84% for  $LC_{50}$ , while the  $LT_{50}$  analysis obtained results of 4.7 days.

Keyword : Lethal concentration, Lethal time, time of application, *Nezara viridula*

## 1. INTRODUCTION

Soybean is a ingredient that has a high carbohydrate content of 14 grams and is used by Asian people as an essential ingredient for daily meal. The high population level accompanied by the availability of soybeans must be sufficient so that there is no gap, based on data from BPS (2017) [1] showing that there was a decline in production in 2015, reaching 344.998 tons, but in 2016 (274.317) and 2017 (200.916) there was a decrease of about 70.000 tons, one of the factors causing a decrease in soybean production is the attack of the Southern green stink bug (*Nezara viridula*).

*Nezara viridula* is a pest that attacks soybeans during the generative period when pods begin to form. *N. viridula* can cause 80% damage. This is due to the high level of mobility and the ability to produce many offspring, Radiyanto (2010) [2] reported that females are able to lay eggs up to 104 – 470, which are placed in groups during their lifetime, *N. viridula* is also able to act as a vector of plant diseases.

*Trichoderma* sp. and *Streptomyces* sp. is a soil microbe that is often found because the presence of this microbe they can provide a good impact on plants because *Trichoderma* sp. able to protect plants from pests and diseases while encouraging soybean production based on Hasibuan's research (2022) [3] *T. harzianum* was able to increase the number of pods, dry stover, and dry weight per plant, Saputri's research (2015) [4] showed that the administration of *T. harzianum*, *T. honingii*, and *T. viridae* were able to inhibit the growth of *S. rolfsii* disease, Ritongga's study (2022) [5] showed that administration of *T. harzianum* could give mortality as much as 83% within 24-26 hours after application, while Hidayah's study (2019) [6] where *Streptomyces* sp. can control larvae of *Lepidiotia stigma* and *Streptomyces* sp. able to play a role as PGPR (Plant Growth Promoting Rhizobium) Vurukonda's research (2018) [7] reported that *Streptomyces* sp. capable of producing antibiotics and volatile organic compounds in soil and in planta. The presence of diverse microorganisms can encourage the resistance of these plants.

Microorganism such as *Streptomyces* sp. and *trichoderma* sp. can work as BCA (Biological Control Agents) because it can produce enzyme or compounds that can damage the cells such as chitinase enzyme, this enzyme can degrade chitin on the shell or cuticle of an insect. Sidabutar's research (2022) [8] showed that by applying *Trichoderma viridae* 60 g/ 10 L was able to control the population of *Oryctes rhinoceros* larvae as much as 91.67 % with an  $LT_{50}$  15 days, Safri *et al* (2017) [9] showed that *Streptomyces* sp. able to hinder the fruit fly (*Bactrocera* sp.) pupation process with a spore density of  $10^{-2}$ . Several research used a combination of *Streptomyces* sp. dan *Trichoderma* sp. as an entomopathogens such as Fitriana *et al* (2019) [10] that can cause the reduce feeding activity of *Spodoptera litura* also Avrianto *et al* (2022) [11] research showed that using a single microorganism *Streptomyces* sp. can give better result than using a combination of *Streptomyces* sp. and *Trichoderma* sp. to repel the soybeans pest such as *Aphid* spp, *Bemisia tabaci*, and *Nezara viridula*.

*Streptomyces* sp. and *Trichoderma* sp. both can be grown on potato extract liquid media but by adding another supplement such as coconut water it can boost its nutritional value Mayaserli *et al* (2015) [12] stated that coconut water have 4 % carbohydrate, 0.1 % fat, 0.02 calcium, 0.01 % phosphor, 0.5 % iron, and 9 g/l of protein that can support the growth of *Pseudomonas fluorescens* colony, both of the BCA have an antagonistic characteristic that can rival each other but according to Breza's research (2016) [13] *Streptomyces* sp. and *Trichoderma* sp. can be compatible this was done by inoculate them in a PDA

46 media pH 6 and 7. Combining 2 BCA types can give an astonishing result because by using two it can  
 47 cover each other flaw, *Trichoderma* sp. have a rapid growth rate but have a low production of chitinase  
 48 enzym while *Streptomyces* sp. have a slow growth rate but have a high production of chitinase enzym, by  
 49 combining these two BCA it can be achieve a high level of chitinase enzym.

50 Calculation of probit analysis  $LC_{50}$  and  $LT_{50}$  was performed to determine how effective these biopesticides  
 51 were in controlling pests, the main objective of this efficacy test was to determine the effectiveness of  
 52 biopesticides with active ingredients *Streptomyces* sp. and *Trichoderma* sp. in a liquid formula  
 53 against *Nezara viridula* attack on soybean plants.

## 54 55 2.2 MATERIAL AND METHOD

### 56 2. EXPERIMENTAL DETAILS

#### 57 2.1 Research Methode

58 The study were conducted at January 2022 to February 2022 in Plant protection labororium at  
 59 agriculture faculty at UPN "Veteran" east java, the study used a factorial randomized block design with 2  
 60 factors. The first factor was the application time before *N. viridula* infestation (S0) and after *N. viridula*  
 61 (S1) infestation. The second factor is the level of concentration consist of 25% (K1), 50% (K2), 75% (K3),  
 62 and 100% (K4), 0 % (K0) as a control, 200 ml of biopesticide will be administered on each replication.  
 63 Mortality rate of *nezara viridula* will the main focus for this research

64

S <sub>0</sub> K <sub>1</sub> (1)	S <sub>1</sub> K <sub>2</sub> (1)	S <sub>0</sub> K <sub>2</sub> (1)	S <sub>1</sub> K <sub>3</sub> (1)	S <sub>0</sub> K <sub>3</sub> (1)	S <sub>1</sub> K <sub>4</sub> (1)	S <sub>0</sub> K <sub>4</sub> (1)	S <sub>1</sub> K <sub>0</sub> (1)	S <sub>0</sub> K <sub>0</sub> (1)	S <sub>1</sub> K <sub>1</sub> (3)
S <sub>1</sub> K <sub>1</sub> (1)	S <sub>0</sub> K <sub>1</sub> (2)	S <sub>1</sub> K <sub>2</sub> (2)	S <sub>0</sub> K <sub>2</sub> (2)	S <sub>1</sub> K <sub>3</sub> (2)	S <sub>0</sub> K <sub>3</sub> (2)	S <sub>1</sub> K <sub>4</sub> (2)	S <sub>0</sub> K <sub>4</sub> (2)	S <sub>1</sub> K <sub>0</sub> (2)	S <sub>0</sub> K <sub>0</sub> (2)
S <sub>0</sub> K <sub>0</sub> (3)	S <sub>1</sub> K <sub>2</sub> (2)	S <sub>0</sub> K <sub>1</sub> (3)	S <sub>1</sub> K <sub>2</sub> (3)	S <sub>0</sub> K <sub>2</sub> (3)	S <sub>1</sub> K <sub>3</sub> (3)	S <sub>0</sub> K <sub>3</sub> (3)	S <sub>1</sub> K <sub>4</sub> (3)	S <sub>0</sub> K <sub>4</sub> (3)	S <sub>1</sub> K <sub>0</sub> (3)

65 Figure 1. Layout of eperiment design

66

67 Table 1. Treatments used in the experiments

Codes	Treatments	Concentration level
S <sub>0</sub> K <sub>1</sub>	Before infestation application	25 %
S <sub>0</sub> K <sub>2</sub>	Before infestation application	50 %
S <sub>0</sub> K <sub>3</sub>	Before infestation application	75 %
S <sub>0</sub> K <sub>4</sub>	Before infestation application	100 %
S <sub>0</sub> K <sub>0</sub>	control	0 %
S <sub>1</sub> K <sub>1</sub>	After infestation application	25 %
S <sub>1</sub> K <sub>2</sub>	After infestation application	50 %
S <sub>1</sub> K <sub>3</sub>	After infestation application	75 %
S <sub>1</sub> K <sub>4</sub>	After infestation application	100 %
S <sub>1</sub> K <sub>0</sub>	control	0 %

68 Note : 200 ml of biopesticide containing *Streptomyces* sp. and *Trichoderma* sp. will be givin once each replicant.

#### 69 2.2 *Streptomyces* sp. and *Trichoderma* sp. isolation

70 Exploration was carried out in Pare, Kediri on healthy shallot farming land, 500 grams of soil samples  
 71 were taken at random and then 1 gram was taken as an isolating material. Isolation of *Streptomyces* sp.  
 72 and *Trichoderma* sp. used soil platting method by Dhingra and Sinclair (2017) [12]. It was carried using  
 73 10<sup>-5</sup> and 10<sup>-6</sup> dilutions for *Streptomyces* sp. then inoculated on GNA (*Glucose Nutrient Agar*) media and  
 74 incubated for 2 weeks. *Trichoderma* sp. diluted in 10<sup>-4</sup> and 10<sup>-5</sup> dilutions then inoculated on PDA media  
 75 (*Potato Dextrose Agar*) and incubated for 3 days.

#### 76 2.3 Provision of Biopesticide Concentration

77 The process of dissolving the concentration of biopesticides using sterile distilled water as a solvent.  
 78 Concentration 25% consist of 125 ml biopesticide with 375 ml aquadest, concentration 50% consist of  
 79 250 ml biopesticide with 250 ml aquadest, concentration 75% consist of 375 ml biopesticide with 125 ml  
 80 aquadest, concentration 100% consist of 500 ml biopesticide, while 0% concentration was control for this  
 81 treatment.

82 **2.4 *Nezara viridula* Rearing**

83 *Nezara viridula* was placed in a rearing box measuring 50 cm x 40 cm X 60 CM, which was covered by a  
 84 net. Long beans were used as feed for the test insects and replaced every 2 days. *N. viridula* Imago will  
 85 be used as a test insect because it has the highest Feeding Activity (Bowling, 1980) [14]

86 **2.5 Application and Infestation**

87 The study used soybean that are in generative stage where its already produced pods, each polybag  
 88 containing 3 soybean plants and there are 30 polibag. The infestation process used 300 imagos of *N.*  
 89 *viridula* total. every treatment need 10 imagos *N.viridula*, biopesticide which will be applied for 200 ml  
 90 once and observed for around 10 days.

91 **2.6 Statistical Analysis**

92 Data analysis will be using ANOVA (Analysis of Variance) and DMRT (Duncan Multiple Range Test) with  
 93 a probability level of 5 % while probit analysis of  $LC_{50}$  and  $LT_{50}$  will be conducted by using regression  
 94 linear, data were submitted to Microsoft excel 2019.

96 **3. RESULT AND DISCUSSION**

97 **3.1 Symptom and Mortality Percentage of *Nezara viridula***

98 The symptom cause by *streptomyces* sp. is shown by the mophological changes in its cuticle thus  
 99 resulting colour changing in the abdomen, thorax, and head (figure. 1) this is aligned with [trisanawati's](#)  
 100 [research \(2018\) \[15\]](#) that the colour change in the abdominal area is the result of chitin degradation thus  
 101 changing the colour to dark black compared to the control.



114 Figure 1. Morphological deformation of *Nezara*  
 115 *viridula*

116 *Nezara viridula* has a thick cuticle layer based on research by [Rawda \(2015\) \[16\]](#) showed that *N.*  
 117 *viridula* has a chitin content of 2% in its cuticle so that it can affect the infection process of a  
 118 microorganism. *Streptomyces* sp. is able to produce chitinase enzyme, but the amount of chitinase  
 119 enzyme is influenced by the type of species, the age of the isolate, and the environment [Sowmya \(2012\)](#)  
 120 [\[17\]](#) reported that *Streptomyces* sp. is able to produce as much as 0.4% (w/v) at the optimum state of pH  
 121 7 with a temperature of 30°C

122  
 123 Table 1. The mortality percentage of *Nezara viridula* on various concentration and application time for 10  
 124 days

Days	Before infestation (S <sub>0</sub> )					After infestation (S <sub>1</sub> )				
	25% (K <sub>1</sub> )	50% (K <sub>2</sub> )	75% (K <sub>3</sub> )	100% (K <sub>4</sub> )	0% (K <sub>0</sub> )	25% (K <sub>1</sub> )	50% (K <sub>2</sub> )	75% (K <sub>3</sub> )	100% (K <sub>4</sub> )	0% (K <sub>0</sub> )
1	0,00%	0,00%	0,00%	0,00%	0,00%	13,33%	0,00%	0,00%	23,33%	0,00%
2	0,00%	0,00%	0,00%	0,00%	0,00%	23,33%	23,33%	23,33%	43,33%	0,00%
3	0,00%	0,00%	10,00%	0,00%	10,00%	26,67%	26,67%	30,00%	50,00%	0,00%
4	0,00%	13,33%	10,00%	0,00%	10,00%	26,67%	26,67%	30,00%	50,00%	0,00%
5	0,00%	13,33%	26,67%	20,00%	16,67%	26,67%	26,67%	30,00%	50,00%	0,00%
6	16,67%	23,33%	30,00%	30,00%	16,67%	26,67%	26,67%	30,00%	50,00%	0,00%
7	26,67%	56,67%	30,00%	33,33%	16,67%	26,67%	26,67%	30,00%	50,00%	0,00%

128  
 129  
 130  
 131  
 132  
 133  
 134  
 135  
 136  
 137

Table 2. time of infestation and concentration level effect on *Nezara viridula* mortality

No	treatment (time of infestation, concentration)	Mortality rate	Notasi
1	Before infestation. concentration 25 %	0,36	bcd
2	Before infestation. concentration 50 %	0,6	e
3	Before infestation. concentration 75 %	0,3	abcd
4	Before infestation. concentration 100 %	0,43	cd
5	Before infestation. concentration 0 % (control)	0,16	ab
6	After infestation. concentration 25 %	0,26	abc
7	After infestation. concentration 50 %	0,26	abc
8	After infestation. concentration 75 %	0,3	abcd
9	After infestation. concentration 100 %	0,5	d
10	After infestation. concentration 0 % (control)	0,13	a

Means within column for each treatment followed by the same letter are not significantly different at  $P \leq 0.05$

138  
 139  
 140  
 141  
 142  
 143  
 144  
 145  
 146  
 147  
 148  
 149  
 150  
 151  
 152  
 153  
 154  
 155

Table 1. showed that the time of death occurred at different times this is because of the before and after infestation treatment, the treatment makes the biopesticide act as stomach poison and contact poison this is aligned with [Djojsumarto \(2008\) \[18\]](#) statement that contact poison works through the insect cuticle while the stomach poison works if the part affected by the biopesticide is being eaten, the research by [Fitriana et al \(2019\) \[19\]](#) showed that by giving a combination of *streptomyces* sp. and *trichoderma* sp. will act as stomach poison for *spodoptera litura* that resulted in increasing feeding activity according to [Lu et al \(1994\) \[20\]](#) insect require a lot of energy to neutralize the poison insect their abdomen.

The highest mortality results were obtained by the 50% concentration treatment, which obtained total mortality of 60% from 30 test insects, while the lowest result was in the 75% treatment, on average the treatment before infestation (S0) obtained a mortality result of 37.33% while after infestation (S1) obtained 29.33%. The factor that affects the effectiveness of contact poison is the level of chitin in the cuticle because *N. viridula* has a chitin level of 5 percent, while *N. viridula* has a piercing-sucking mouth type, so the amount of fluid taken is not maximal.

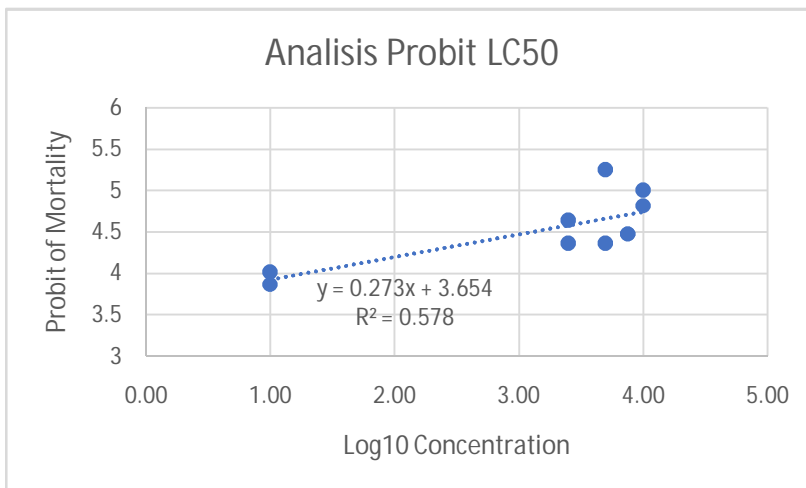
Before infestation (S0) mortality occurred in 4 DAA in 50% and 75% of treatment, respectively. It was recorded that 13% and 10% of mortality, the highest mortality rate occurred in 7 DAA in 75% treatment, which reached 50% normality. After infestation (S1) mortality occurred in 1 DAA in 25% and 100% treatment, respectively recorded at 13% and 23%. The highest mortality rate occurred in 3 DAA, reaching

156 50%. Mortality rate shown that the S<sub>0</sub>K<sub>2</sub> have the highest result of 60 % rather than the S<sub>0</sub>K<sub>4</sub> (43,33 %)   
 157 this was cause due the enviroment temperature (33°C) that can cause the evaporation of the biopesticide,   
 158 some will evaporize while some of it absorbed by the seed pods this can be act as a stomach poison   
 159 because the liquid that containg the BCA's have an enzyme that can degrade the chitin on insect, seed   
 160 pods tend to hold the moisture around it to keep the seed inside moist and cool. S<sub>1</sub>K<sub>4</sub> treatment that have   
 161 a concentration level of 100 % gives an instant result, in under 12 hours it already killed nearly 23,33 %   
 162 after 48 hours it was already at 50 % then it became stagnant, the reason it became stagnant varies from   
 163 evaporation, *N. viridula* might have a thicker cuticle, or it needs another dose of biopesticed in order to   
 164 keep the mortality rate rising. Based on the mortality results, it was concluded that the biopesticide with   
 165 the active ingredient *Streptomyces* sp. and *Trichoderma* sp. It can be used as a stomach poison and a   
 166 contact poison.

### 167 3.2 LC<sub>50</sub> and LT<sub>50</sub> Analysis

#### 168 3.2.1 Probit Analysis of LC<sub>50</sub>

169 Probit analysis on the mortality of *N. viridula* after 10 days of application of the biopesticide formulation   
 170 containing *Streptomyces* sp. and *Trichoderma* sp. is presented in figure 2. probit analysis is carried out to   
 171 determine the lc<sub>50</sub> of the biopesticide the result showed a linier equation which is the correlation between   
 172 the probit of mortality (Y) and the logarithm of concentration (X) as follows :  $y = 0.2731x + 3.6547$  with a   
 173 correlation value (R) 0.7605 while the regression (R<sup>2</sup>) 0.5785, so it can be concluded that there is a   
 174 correlation between the administration of biopesticide and the mortality of *N. viridula*. correlation value (R)   
 175 is close to 1 therefore there's a strong connection that resulted in the percentage mortality of 52.35 % the   
 176 results of probit analysis showed that the LC<sub>50</sub> value with a time period of 10 days was 84,453 ppm or 84   
 177 % is the concentration required to control the population of *N. viridula*.

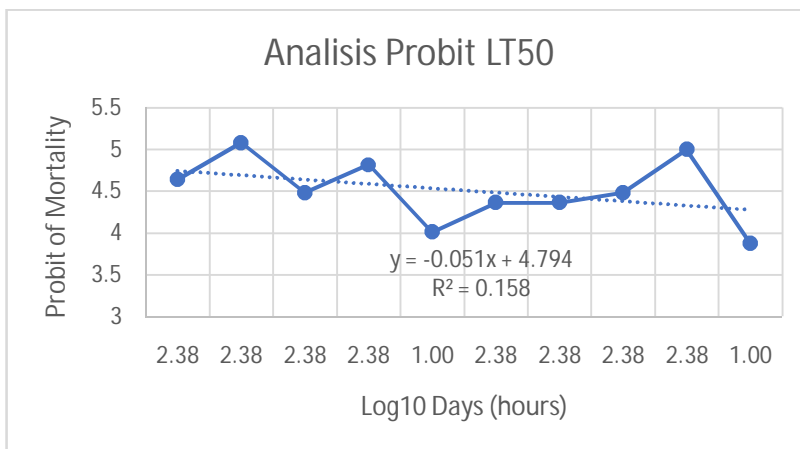


180  
181  
182  
183  
184  
185  
186  
187  
188  
189  
190  
191  
192 Figure 2 shows an increase of correlation between the concentration and probit of mortality, so it can be   
 193 concluded that the higher the biopesticide concentration, the higher the mortality rate. This is in   
 194 accordance with the results of [Handayani's research \(2019\) \[21\]](#) giving a concentration of 16 ppm of   
 195 ethanol extract of *Tabernaemontana macrocarpa jack* leaves can control the population of *Artemia*   
 196 *salina*, LC<sub>50</sub> results showed 0.7440 g/ml, [Hasyim's research \(2019\) \[22\]](#) showed that the administration of   
 197 botanical insecticides from *Cerbera manghas* leaf extract was able to control *S. exigua* as much as 85%   
 198 with an LC<sub>50</sub> of 1.002.67 Ppm and an LT<sub>50</sub> of 46.98 hours.

#### 199 200 3.2.2 Probit Analysis of LT<sub>50</sub>

201 Probit LT<sub>50</sub> analysis was used to determine on what days this biopesticide was able to control the pest   
 202 population as much as 50%, Figure 3. shows a linear equation which is the correlation between probit   
 203 mortality percentage (y) and the logarithm of days (hours) (x) as follows:  $y = -0.0516 x + 4.794$  with a   
 204 correlation value of (R) is 0.398. The calculation of probit analysis shows that the regression (R<sup>2</sup>) obtained   
 205 is 0.1589, and the correlation value (R) is 0.3986 with a value (R) close to 0 so it can be concluded that   
 206 there is no a strong relationship this is due to the presence of unknown factors. The trendline shows that   
 207 the decline is due to the weakening influence between variables, causing a decrease. The analysis of

208 probit LT50 with a period of 10 days was obtained in the form of 113 hours. If converted in days, there are  
209 4.7 days needed to control the insect pest population *N. viridula*.



221  
222  
223  
224  
225  
226  
227  
228  
229  
230  
231  
232  
233  
234  
235  
236  
237  
238  
239  
240  
241  
242  
243  
244  
245  
246  
247  
248  
249  
250  
251  
252  
253  
254  
255  
256  
257  
258  
259

Figure 3. Probit Analysis of LT<sub>50</sub> againts *Nezara viridula* population

Probit analysis LC<sub>50</sub> and LT<sub>50</sub> can be affected by the type and age of micro organisme, and the target insect, [Nia's reasearch \(2016\) \[23\]](#) showed that by administrated *Metharizium* on *M. Anisopliae* on *Lepidiota stigma* 3<sup>rd</sup> instar it can give result of LT<sub>50</sub> 7.7 days while the LC<sub>50</sub> is 8.2 x 10<sup>8</sup> conidia/ml this was due to shedding or molting which could inhibit the infection process, [Cahyani's research \(2022\) \[24\]](#) showed that by administrated *Aspergillus niger* to *Aedes aegypti* larvae it give a result of LC<sub>50</sub> 6.1 x 10<sup>-7</sup> and LT<sub>50</sub> 1.919 hours.

#### 4. CONCLUSSION

Efficacy test of biopesticide containing *Streptomyces* sp. and *Trichoderma* sp. in controlling the insect pest *N. viridula* showed symptoms of death caused by the degradation of chitin in the abdomen and thorax, mortality rate showed that S0K2 (60.00 %) treatment have the highest mortality rate while the lowest is from S1K1 and S1K2 (26.67 %). LC<sub>50</sub> analysis showed a concentration of 84% while the LT<sub>50</sub> analysis showed of 4.7 days, this biopesticide can also be used as a stomach poison and contact poison. Therefore this biopesticide is effective but not efficient.

#### ACKNOWLEDGEMENT

Thank you to the Universitas Pembangunan Nasional "Veteran" Jawa Timur who have provided research grants to Dr. Ir. Penta Suryaminarsih and PKKMM Fund (Program Kompetisi Kampus Merdeka) Ministry of Education and culture, research and technology, Noni Ramadhini. SP. M.Sc. and Ramadhani Mahendra Kusuma, S.P., M.P., M.Sc.for providing support and guidance on writing and grammar revision, my friend Amri, Sofi, Mandayu, and Firda for creating this biopesticide, Adis, Indah, Teha, Ana, Axcel, Adi, Sukma, Rey, and Retno for giving me support and love.

#### REFERENCES

1. Central Bureau of Statistics. (2017). Harvest Area, Productivity, and Production of Soybeans in East Java 2002 – 2017 at <https://jatim.bps.go.id/statictable/2018/10/31/1342/wide-panen-productivity-dan-hasil-kedelai-di-jawa-timur-2002-2017.html>. Accessed on July 3, 2021.

- 260  
261  
262  
263  
264  
265  
266  
267  
268  
269  
270  
271  
272  
273  
274  
275  
276  
277  
278  
279  
280  
281  
282  
283  
284  
285  
286  
287  
288  
289  
290  
291  
292  
293  
294  
295  
296  
297  
298  
299  
300  
301  
302  
303  
304  
305  
306  
307  
308  
309  
310
2. Radiyanto, B., Sodiq, M., & Nurcahyani, N. M. (2010). Diversity of insect pests and natural enemies on soybean plantations in Balong-Ponorogo District. *Indonesian Journal of Entomology*, 7(2), 116-116.
  3. Hasibuan, S. A., Zurni, T. C., & Syamsuddin, S. (2022). The Effectiveness of BioPriming *Trichoderma harzianum* and Compost Application of Oil Palm Empty Fruit Bunches on Growth and Production of Soybean (*Glycine max* L. Merr.) Plants. *Journal of Agrista*, 26(1), 9-16.
  4. Saputri, E., Lisnawita, L., & Pinem, M. I. (2015). Encapsulation of Several Types of *Trichoderma* sp. on Soybean Seeds to Control Disease *Sclerotium rolfsii* Sacc. *Journal of Agroecotechnology*, University of North Sumatra, 3(3), 105478.
  5. Ritonga, N. F., Nuraida, N., & Sari, A. (2022). Pathogenicity of *Trichoderma harzianum* to Larval Pests of Horn Beetle (*Oryctes rhinoceros*) in Oil Palm Plants (*Elaeis guineensis* Jacq.) in the Laboratory. *Journal of Agrofolium*, 2(2), 98-107.
  6. Hidayah, A. R., Harijani, W. S., Widajati, W., & Ernawati, D. (2019). Potential entomo pathogenic fungi *Metarhizium anisopliae*, *Beauveria bassiana* and *Streptomyces* sp. on mortality of *Lepidiotia stigma* in sugarcane. *Plumula: Agrotechnology Scientific Periodic*, 7(2), 64-72.
  7. Vurukonda, S. S. K. P., Giovanardi, D., & Stefani, E. (2018). Plant growth promoting and biocontrol activity of *Streptomyces* spp. as endophytes. *International journal of molecular sciences*, 19(4), 952.
  8. Sidabutar, M., Nuraida, N., & Sofian, A. (2022). Pathogenicity of *Trichoderma viride* Fungus against Horn Beetle Larvae Pests on Oil Palm Plants. *Journal of Agrofolium*, 2(2), 135-141.
  9. Safri, M., Harijani, W. S., & Suryaminarsih, P. (2017). Pupa Viability Test of Fruit Flies (*Bactrocera* sp.) Become Imago by Giving *Streptomyces* sp Biological Agent. *Agrotechnology-PLUMULA Scientific Periodic*, 5(1).
  10. Fitriana, I. N., Suryaminarsih, P., & Mujoko, T. (2018). Potential of Multientomopa *Streptomyces* sp. and *Trichoderma* sp. in Potato Extract Broth and Glucose Nitrate Broth Media on Pests (*Spodoptera litura*) Eating Behavior by in Vitro Test. *Nusantara Science and Technology Proceedings*, 270-276.
  11. AVRIANTO, N. I. (2021). The Effect of Application of the Biological Agent Formula of *Streptomyces* sp. and *Trichoderma* sp. On the Presence of Insect Pests on Soybean Plants (*Glycines Max* L. Merril) Vegetative Phase (Doctoral dissertation, "Veteran" National Development University, East Java).
  12. Mayaserli, D.P. and Renowati, R., 2015. Utilization of Coconut Water as a Growth Media for *Pseudomonas fluorescens* and its Application as a Liquid Plant Fertilizer. *Pioneer Health Journal*, Vol. 2 (2): 19-22.
  13. Breza Boruta, B., &Paluszak, Z. (2016). The antagonistic activity of actinomycetes of *Streptomyces* genus in relation to *Trichoderma koningii*. *Journal of Ecological Engineering*, 17(1), 106-113.
  14. Sinclair, J. B., & Dhingra, O. D. (2017). *Basic plant pathology methods*. CRC press.
  15. Bowling, C. C. (1980). The stylet sheath as an indicator of feeding activity by the southern green stink bug on soybeans. *Journal of Economic Entomology*, 73(1), 1-3
  16. Trisnawati, Didin Julia, Wiwik Sri Harijani, and Penta Suryaminarsih. 2018. "Concentration Test of *Streptomyces* sp. Biological Agent. against the Pupae of the *Bactrocera* sp Fruitfly." 6(1):41-48.
  17. Rawda M. Badawy and Hadeer I. Mohamed. Chitin extration, Composition of Different Six Insect Species and Their Comparable Characteristics with That of the Shrimp. *J Am Sci* 2015;11(6):127-134].
  18. Sowmya, B., Gomathi, D., Kalaiselvi, M., Ravikumar, G., Arulraj, C., & Uma, C. (2012). Production and Purification of Chitinase by *Streptomyces* sp. from Soil. *Journal of Advanced Scientific Research*, 3(3).
  19. Djojsumarto, P. (2008). *Complete guide to pesticides & their application*. agromedia.
  20. Fitriana, I. N., Suryaminarsih, P., & Mujoko, T. (2018). Potential of Multientomopa *Streptomyces* sp. and *Tripchoderma* sp. in Potato Extract Broth and Glucose Nitrate Broth Media on Pests

- 311 (*Spodoptera litura*) Eating Behavior by in Vitro Test. Nusantara Science and Technology  
312 Proceedings, 270-276.
- 313 21. Lu, H., Rajamohan, F., & Dean, D. H. (1994). Identification of amino acid residues of *Bacillus*  
314 *thuringiensis* delta-endotoxin CryIAa associated with membrane binding and toxicity to *Bombyx*  
315 *mori*. *Journal of Bacteriology*, 176(17), 5554-5559.
- 316 22. Handayani, F. F., Sentat, T., & Rahim, A. (2019). Acute Toxicity Test of Ethanol Extract of Selutui  
317 Puka Leaves (*Tabernaemontana macrocarpa* Jack.) on (*Artemia salina* Leach) larvae. *Journal of*  
318 *the World of Pharmacy*, 4(1), 1-7.
- 319 23. Hasyim, A., Lukman, L., & Marhaeni, L. S. (2019). Evaluation of lethal concentration and lethal  
320 time of botanical insecticides on onion caterpillar (*Spodoptera exigua*) in the laboratory. *Journal of*  
321 *Horticulture*, 29(1), 69-80.
- 322 24. Niâ, L., Himawan, T., & Mudjiono, G. (2016). Pathogenicity test of entomopathogenic fungus  
323 *Metarhizium anisopliae* (Moniliales: Moniliaceae) against *Lepidiota stigma* F. (Coleoptera:  
324 Scarabaeidae). *Journal of Plant Pests and Diseases*, 4(1), 24-31.
- 325 25. Cahyani, N. D. K. S., Wiadnya, I. B. R., Khusuma, A., & Getas, I. W. (2022). Analysis of Lethal  
326 Concentration and Lethal Time of *Aspergillus Niger* Fungus Isolate Against *Aedes Aegypti*.  
327 *Journal of Bioscience Medical Analysts (JAMBS)*, 9(2), 78-86.

328