Original Research Article

2	BIOCONTROL POTENTIAL OF BACILLUS THURINGIENSIS ISOLATED FROM
3	SOIL
4	AGAINST MOSQUITO LARVA

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ABSTRACT

A major challenge for achieving successful mosquito control is overcoming insecticide 7 resistance. This study was carried out to assess the larvicidal activity of *Bacillus thuringiensis* 8 isolated from different soil samples within Sokoto metropolis using standard methods. 9 Confirmatory identification of the organism was made based on biochemical characterization 10 and microscopic observation. The larvicidal activity of *Bacillus thuringiensis* isolated was tested 11 against the larva of mosquito using three dilutions of the Bacillus culture in a bioassay. The 12 isolated organisms were confirmed as Bacillus thuringiensis. The result of the bioassay showed 13 variation in the level of efficacy of the bacteria and depended on time of the exposure. Mortality 14 rate greater than 20% was observed after 60 minutes and increased to 100% after time of 15 exposure was increased for all dilutions of B. thuringiensis used. The results showed that 16 Bacillus thuringiensis toxins can be bacteriocidal to mosquito larvae in a matter of minutes 17 depending on the concentration ingested by the larvae. This in essence proved that Bacillus 18 thuringiensis is an effective bio-larvicide that can be used to reduce and possibly eradicate the 19 nuisance of disease-causing mosquitoes and aid in the rollback of malaria. 20

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Keywords: Mosquito control, *Bacillus thuringiensis*, bioassay, larvicidal activity, add mosquito
 scientific name

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27 INTRODUCTION

28 Mosquitoes are associated with the transmission of pathogens to humans and other vertebrates. 29 Some of these include the causative agents of malaria, filariasis and dengue, as well as other 30 mosquitoe-borne zoonotic aborvirus, like Saint Louis Encephalitis Virus (SLEV), West Nile Virus (WNV) and Eastern Equine Encephalitis Virus (EEEV) [1]. Significant morbidity and 31 32 mortality are recorded as results of these diseases are as a result of the inherent difficulty of 33 controlling mosquitoes [2, 3]. Increase in the distribution of mosquitoes was associated with the 34 emergence of viruses and diseases in new areas [4]. WHO [3] reported malaria to be the world most important vector-borne disease. Cases of these diseases have been reported in more than 35 100 countries, with approximately more than 3 billion people living in endemic areas. More than 36 37 200 million cases of malaria and eight hundred thousand (800,000) malaria-related deaths are recorded every year [5]. The increase in number of malaria cases is as a result of deteriorating 38 health systems, increase resistance of anopheline to insecticides, time taken to develop an 39 40 effective vaccine and as well resistance of plasmodium to antimalarial drugs [1].

Dengue, including dengue hemorrhagic fever and Dengue Shock Syndrome (DSS) transmitted by *Aedes* mosquitoes is rapidly becoming a worldwide disease, threatening a third of the world population, with an estimate of 50-100 million cases every year [6, 7]. So also, *Lymphatic filariasis* caused by *Wuchereria bancropti*, which is transmitted by mosquitoes, affects more than 120 million people around the world [8]. Lack of effective vaccine against these diseases has left the control of mosquito population as the most effective way to prevent vector-borne diseases [9, 10].

Chemical insecticides have been used in the last century to successfully control mosquitoes of
the genus *Aedes* and *Anophelles*. Current ecological and environmental protection standards halt

the use of these chemicals, because of their adverse effects on non-target species, including 50 humans, environmental impact, contamination soil and water and development of mosquito 51 resistance to insecticides [11]. New strategies were created to replace the use of chemical 52 insecticides. They include Integrated Pest management (IPM) that has guidelines. Guidelines of 53 which are based on environmental planning, public awareness and biological control that control 54 the mosquitoes more efficiently while preserving the environment from contamination [1]. 55 Commercial preparations of Bacillus thuringiensis (Bt) as a biocontrol agent has been the 56 greatest success in microbial pesticides, with more than 95% of the microbial pesticides sold 57 being of this bacterial agent [12]. Bacillus thuringiensis (Bt) is a gram-positive, rod-shaped and 58 spore forming bacteria that is mostly found in the soil and produces polymorphic crystal proteins 59 [13]. The insecticidal activity of Bt is due to the proteic parasporal inclusions that are produced 60 during sporulation [14]. The insecticidal proteinaceous crystals (ICPs) comprised one or more 61 crystal (Cry) and Cytolytic (Cyt) proteins recognized as δ-endotoxin. When ingested by the 62 target insect, the ICPs dissolve in the midgut of the larva releasing proto-toxins that eventually 63 lead to the formation of pores that causes cell-cytolysis [15, 16]. 64

Despite the use of *Bacillus thuringiensis* as a biocontrol agent for over 30 years, no significant resistance was recorded. However, the search for natural *Bt* isolates with increase activity against mosquito and other insect is still encouraged. Recently, *Bt* with increased activity against *Aedes caspius* and *Culex pipiens* were isolated [17]. In this study, the larvicidal activity of *Bt* isolated from soil samples will be evaluated on mosquitoes.

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73 MATERIALS AND METHODS

74 Sample Collection

75 A total of 4 soil samples were collected from Tamaje, Mabera, Arkilla and Kantin Daji areas of

76 Sokoto state, where there is no previous record of application of *Bacillus thuringiensis* based

⁷⁷ insecticides. The soil samples were collected aseptically from top of 5cm depth. The samples

78 were placed immediately in plastic bags and labeled appropriately [18]. The soil samples were

79 transported to the laboratory and stored at room temperature. In the same vein, mosquito larva

80 was collected from stagnant waters around Sokoto metropolis.

81 Media Preparation

82 The media to be used to culture the bacteria such as Nutrient agar and Luria-Bertani will be

83 prepared according to the manufacturer's instruction.

84 Nutrient Agar

Twenty-eight grams (28g) of nutrient agar was weighed and dissolved in 1000ml of distilled water in a sterile conical flask. The mixture was heated using a hot plate to dissolve the medium. The conical flask was plugged with cotton wool stoppers and wrapped with aluminum foil. It was then sterilized using an autoclave at 121°C for 15 minutes. The medium was cooled at about 45 - 50°C after sterilization and then poured into sterile petri dishes (about 20ml per plate) under aseptic conditions. The plates were allowed to solidify and incubated at 37°C for 24 hours and the sterility of the medium was checked [19].

92 3.2.2 Luria-Bertani Agar

93 This media is made up of Tryptone, Yeast extract, NaCl, NaOH and Bacto agar for jelling. Ten 94 grams (10g) of Tryptone, ten grams (10g) of NaCl, five grams (5g) of NaOH and ten grams 95 (10g) of yeast extract were weighed and dissolved in 950ml of distilled water in a sterile conical **Comment [P4]:** Add latitude and longitude of soil sample collection areas

Comment [P5]: Please mention the room temperature and humidity of the sample.

Comment [P6]: Add latitude and longitude of mosquito larvae collection areas.

Comment [P7]: Please add ingredients of Bacterial culture medias

flask. The mixture was heated using a hot plate to dissolve. The final volume was added up to 1000ml and fifteen grams (15g) of Bacto agar was added. The conical flask was plugged with cotton wool stoppers and wrapped with aluminum foil. It was then sterilized using an autoclave at $\frac{121^{9}\text{C}-121^{\circ}\text{C}}{15}$ for 15 minutes. The medium was cooled at about $45 - \frac{50^{9}\text{C}-50^{\circ}\text{C}}{2}$ after sterilization and then poured into sterile petri dishes (about 20ml per plate) under aseptic conditions. The plates were allowed to solidify and incubated at $\frac{37^{9}\text{C}-37^{\circ}\text{C}}{2}$ for 24 hours and the sterilizy of the medium was checked [20].

103 Isolation of *Bacillus thuringiensis* from Soil

Five grams (5g) of each soil sample was weighed and added to 100ml of distilled water. The samples were heated on a hot plate for 10 minutes to eliminate all bacteria incapable of producing endospores. Since it is known that *Bacillus thuringiensis* produces spores, it will be safe to assume that if it was present in the soil, it would be in our heated sample. The samples were then diluted 5 fold to eliminate all humic materials within the samples and to reduce the overall colony forming units within each sample [18].

110 Culturing of Bacillus thuringiensis

The diluted samples were cultured on nutrient agar plates for 24 hours at $37^{\circ}C_{37^{\circ}C}$ in to order to give the spores chance to germinate on media with adequate nutrients and optimal temperature [18]. The media, however, offers favorable growth for a wide range of bacteria as well as *Bacillus thuringiensis*. The colonies were sub-cultured onto Luria-Bertani plates and incubated at $37^{\circ}C_{37^{\circ}C}$ for 24 hours, so as to obtain pure cultures of *B. thuringiensis*. Series of tests which include gram staining and biochemical tests were further employed to identify *Bacillus thuringiensis* after formation of colonies with smooth round shape and earthy odor.

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119 Gram Staining Techniques

A smear of colonies isolated after the identification were made on a clean glass slides using a 120 121 sterile wire loop. They were air dried and fixed. The smears were flooded with crystal violet for about 60 seconds and were washed with tap water. They were then tipped off with lugol's iodine 122 for 30 seconds and then washed with tap water. They were decolorized with acetone and washed 123 off with tap water. The fixed smears were counterstained with safranin and allowed for 60 124 seconds and then washed off with tap water and allow to air dry. Oil immersion was added to the 125 stained slides and viewed under a microscope using x100 objective for the morphological 126 characteristics of the isolates [21]. 127

127 characteristics of the isolates [21].

128 Characterization of the Isolated Bacteria

129 The colonies that form on the T3 agar will again be confirmed by biochemical tests based on:

Indole test, Catalase test, Triple Sugar Iron test (T.S.I), Methyl Red test (M.R.), VoguesProskauer (V.P.).

132 Indole Test

A test tube of sterile peptone water, enriched with 1% tryptophan will be inoculated with young culture of isolates and incubated at 37° C for 48hrs. About 4 drops of kovac's reagent will be added and shaken gently. A red color will occur immediately at upper part of the test tube indicating a positive test. A yellow color at the surface will denote a negative result [21].

137 Catalase Test

The container containing 3% hydrogen peroxide solution will be shaken to expel the dissolved oxygen. One drop of the solution will be placed on a clean glass slide. Presence of gas bubbles indicates a positive test while absence of gas bubbles indicates negative reaction [19].

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142 Triple Sugar Iron Agar (T.S.I test)

A speck of the isolate will be inoculated by streaking and stabbing into the medium and will be incubated at 37^oC for 24 hours. Fermentation of any of the sugar will be indicated by a change in color, from red to yellow and crack or raised in the medium indicates gas production [21].

146 Methyl Red Test (M.R.)

147 A speck of the isolate will be inoculated into the medium, which would be incubated at $37^{\circ}C$ for

148 48 hours. Few drops of methyl red would be added to the culture. M.R positive test will indicate

red color while no changes denotes negative [21].

150 Vogues-Proskauer Test (V.P)

A speck of the isolate will be inoculated into glucose phosphate water medium and incubated at 37^oC for 2 days. Ethanoic solution of 5% α -naphthol (1.2 ml) and 0.4 ml potassium hydroxide solution will be added to 2ml of culture and will be shaken vigorously. It will be placed in a sloping position (for maximum exposure of the culture to air) and will be examined after 30 to 60 minutes. The evolution of red color indicates a positive test for Voges-Proskauer [21].

156 Coagulase Test

About 2 or 3 colonies was emulsified in 0.05ml of saline contained in a serological tube. 1ml of plasma was added and incubated at 35^{0} - 37^{0} C and it was checked after 1hour, 2hours, 3hours and 4hours of incubation for signs of clotting of the plasma. Increase in viscosity or complete clotting indicates a positive coagulase test, while absence of viscosity or clotting indicates a negative coagulase test [21].

162 Motility Test

A small quantity of each isolate was stabbed into triple sugar iron agar and incubated at 37° C for 4 A small quantity was observed by spread of the organism outwards from the stabbed area.

165 Urease Test

A speck of each isolate was inoculated into Christensen's urea agar and incubated at 37°c for 24 166 hours. Liberation of red color indicates urease positive test while initial yellow color indicates 167 168 negative test. **Citrate Utilization Test** 169 A speck of each isolate was inoculated into Koser's citrate medium and was incubated at 37°C 170 for 72 hours. A positive citrate is confirmed by the promotion of blue color while the initial 171 green color denotes a negative result [21]. 172 **Bioassay** 173 The Bacillus thuringiensis isolates selected were tested against larva of mosquito. The stock 174 cultures of Bacillus thuringiensis from slant bottles were picked using a sterile wire loop and 175 diluted five-fold 10^{-1} - 10^{-5} in sterile distilled water in five test tubes. Five (5) ml each of the 176 cultures in the first, third and fifth test tubes was added to three (3) disposable cups containing 177 45ml of sterile distilled water, providing each cup with different dilution factors. Twenty-five 178 (25) larvae were transferred into each of the disposable cups. The cups were kept at $\frac{25^{\circ} - 25^{\circ} - 25^{$ 179 30°C for 6 hours. At intervals of 30 minutes, each cup was observed for larval presence and 180 larval mortality rate was calculated. 181 RESULTS 182 The microscopic and biochemical characteristics of the isolated organisms are shown in table 183

4.1. The characteristics of which confirmed the isolated organism to be *Bacillus thuringiensis*.
Table 4.2 shows the result for the bioassay of 10⁻¹ diluents of *Bacillus thuringiensis* against
mosquito larvae. The mortality rate was found to increase as the incubation time increases. A
mortality rate of 52% was recorded after 150 minutes and a 100% mortality rate was recorded

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Comment [P12]: Which mosquito larvae??? Comment [P13]: What is size of the bioassay cups??

Comment [P14]: How to calculated larval mortality???

188	after 330 minutes. The result illustrated in table 4.3 shows the bioassay for the 10^{-3} diluents of <i>B</i> .
189	thuringiensis on mosquito larvae. Mortality rate of 52% was recorded after 180 minutes, after
190	which a 100% mortality rate recorded after 360 minutes. Illustrated in table 4.4 is the bioassay of
191	the 10 ⁻⁵ diluents of <i>B. thuringiensis</i> against mosquito larvae. Mortality rate of 60% was recorded
192	after 210 minutes, after which a mortality rate of 100% was recorded after 360 minutes.
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V Isolates	Gram Reaction	Catalase	Coagulase	Glucose	Sucrose	Lactose	Gas	Motility	Citrate	MR	VP	Urease	+ Indole
Α	+ve Rod	+	-	-	-	-	-	+	+	+	-	+	+
В	+ve Rod	+	+	-	-	-	-	+	+	+	-	+	+
С	+ve Rod	+	-	-	-	-	$\overline{\mathbf{a}}$	+	+	+	-	+	+
D	+ve Rod	+	+	-	-	Ń		+	+	+	-	+	+
Key:	+ = Positive				Ŕ								
	- = Negative				F								
			4		<i>.</i>								
Time	(min)	0	30	60	90	120	150	180 21	10 2	40 27	70 300	330	36

196Table 4.1: Biochemical and morphological characteristics of Bacillus thuringiensis

	No. of larva	25	24	2	0	17	14	12	10	7	4	2	1	0	0	_
	Control	25	25	2	5	25	25	25	25	25	25	25	25	25	25	
	Mortality	0	1	5		8	11	13	15	18	21	23	24	25	25	
	Mortality rate (%)	0	4	2	0	32	44	52	60	72	84	92	96	100	100	
200																_
201								1								
202	Table 4.2: Bioassay of 10 ⁻¹ Diluents of Bacillus thuringiensis Culture Against Mosquito Larvae.												Comment [P16]: This concentrations contain how much bacterias???			
203														Comment [P17]: Which mosquitoes you have used		
204	Table 4.3: Bioassay of 10 ⁻³ Diluents of <i>Bacillus thuringiensis</i> Culture Against Mosquito Larvae.												Comment [P18]: This concentrations contain how much bacterias???			
	Time (min)	0	30	60	90	120		150	180	210	240	270	300	330	360	Comment [P19]: Which mosquitoes you have used
	No. of larva	25	24	23	21	19		16	12	9	7	6	3	1	0	
	Control	25	25	25	25	25		25	25	25	25	25	25	25	25	
	Mortality	0	1	2	4	6		9	13	16	18	19	22	24	25	
	Mortality rate (%)	0	4	8	14	24		36	52	64	72	76	88	96	100	

Table 4.4: Bioassay	of <mark>10⁻⁵</mark>	Diluen	ts of <i>Ba</i>	cillus th	uringien	esis Cultu	ire Again	st <mark>Mosq</mark> ı	uito Lar	vae.				Comment [P20]: This concentrations how much bacterias???
Time (min)	0	30	60	90	120	150	180	210	240	270	300	330	360	Comment [P21]: Which mosquito lau you has taken the how you identify that mosquito or not???
No. of larva	25	25	24	23	23	21	18	10	6	3	1	0	0	-
Control	25	25	25	25	25	25	25	25	25	25	25	25	25	
Mortality	0	0	1	2	2	4	7	15	19	22	24	25	25	
Mortality rate (%)	0	0	4	8	8	16	28	60	76	88	96	100	100	
			0											-

WORLD REFERENCE

216

217 **DISCUSSION**

Mosquitoes are a great nuisance and they pose a serious threat to human health in the society. 218 219 Many chemical insecticides have been produced for the control of mosquitoes in the past years, some of which have been very effective while others have done little or no good at all. Most of 220 the insecticides used are made of synthetic chemicals and were found to have negative effect on 221 the diversity of many insects, well being of humans and as well the environment. These 222 therefore, call the need for the search of biological control methods that cause less harm to 223 human health, diversity of the insects and the environment. The efficacy of Bacillus 224 thuringiensis as a larvicide for controlling mosquito larva yielded great results. The microscopic 225 and the biochemical characteristics of the organisms as shown in Table 4.1 confirmed the basic 226 227 characteristics of Bacillus thuringiensis, being Gram positive and having a rod shape. The biochemical characteristics, showed the motile nature of the organism among others. These 228 characteristics are similar to what was reported by Ahmed et al. [22]. 229 In the bioassay for the control of mosquito larvae, different diluents used showed varying degree 230 of effectiveness with 100% mortality rate recorded after 360 minutes. This might be attributed to 231 the ability of the organism to produce binary toxin (Bin) which is a primary insecticidal 232 component produced during sporulation and vegetative stage of B. thuringiensis in controlling 233 the growth of the mosquito larvae. This is in agreement with what was reported by Oei et al. 234 [23]. In all the diluents, very low mortality was recorded after 30 minutes of incubation, which 235 could be attributed to the time of exposure of the larvae as well as the number of the organisms 236 present in the container. But more than half of the larvae were death after 240 minutes, with a 237 high mortality rate of over 70% recorded in all the diluents of the B. thuringiensis. This could 238

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239	also be attributed to the time of exposure of the larvae and as well the increase in number of the
240	cells in the medium that could be attributed to the increase in the number of organisms ingested
241	by the mosquitoe larvae, which causes damage in the midgut of the larvae [24]. Thereby
242	releasing the crystallized binary toxins, that in turns are solubilized in the midgut, releasing two
243	proteins [25], that are cleaved by endogenous proteins to form active toxins [26]. This is in
244	agreement with what was reported by Aissaoui and Boudjelida [27].
245 246	Conclusion
247	Bacillus thuringiensis naturally found in the soil has proved to be a good larvicidal agent against
248	mosquito larvae in the laboratory. The organism and it product can be further studied to search
249	for novel compounds that can be use in control of mosquito borne diseases such as malaria. Comment [P24]: The conclusion part not written properly so please read some other
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