Original research paper

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Phytochemical components and antibacterial activity of *Tamarindus indica* Linn. extracts against some Pathogens

5 ABSTRACT

Aim: to determine the phytochemical composition and antimicrobial properties of tamarind extracts on
 some aquatic pathogenic bacteria.

8 Study design: Completely Randomized Design (CRD)

9 Place and duration of the study: Department of Animal Production, Fisheries and Aquaculture,
 10 Kwara State University, Malete, Nigeria, between August 2014 and April, 2015.

11 **Methodology**: The phytochemical constituents in ordinary, warm and hot water as well as ethanol 12 extracts of tamarind seed coat, pulp and leaves were screened. The Zone of Inhibition (ZOI) diameter 13 (mm) and Minimum Inhibitory Concentration (MIC) and Minimum Bactericidal Concentration (MBC) 14 against some aquatic pathogenic bacteria were determined. Data were analyzed using ANOVA at P =15 .05.

16 Results: The result revealed presence of reducing sugar, flavonoid, saponin and terpenoids in all 17 tamarind extracts. The synthetic antibiotics used had significantly higher ZOI than the tamarind 18 extracts for all the test organisms. Tamarind pulp hot water extract significantly inhibited Aeromonas hydrophila and Hafnia alvei than other extracts while leaf warm and hot water extracts had 19 significantly higher zone of inhibition against Pseudomonas putida. The best MIC was obtained for 20 21 oxytetracycline and erythromycin against Enterobacter gergovia and Escherichia coli respectively. Pulp extracts and erythromycin exhibited the same MIC, 2.56mg/ml, for Bacillus subtilis and H. alvei 22 23 while the former had lower MIC (2.56mg/ml) against Salmonella typhi than the MIC (5.12mg/ml) of the later. Oxytetracycline and tamarind extracts also demonstrated the same MIC (2.56mg/ml) against S. 24 typhi. Pulp extracts exhibited MBC for most of the test organisms. 25

26 **Conclusion**: warm and hot aqueous tamarind pulp and leaf extracts demonstrated better 27 antimicrobial activities against some bacteria used in this study and hence the extracts could be used 28 to control such microbes associated with the aquatic environment and fish products.

29 Key words: Tamarind, antibacterial activity, phytochemical, minimum inhibitory

30 concentration, synthetic antibiotic

31 **1. INTRODUCTION**

32 Some of the major challenges facing fish culturists are adequate sources of low cost quality feed,

33 availability of quality fish feed and promotion of fish health. As aquaculture becomes more and more

34 intensive, feeds and disease prevention are significant factors in increasing the productivity and

35 profitability of aquaculture. Hence, investing in disease prevention and treatment is crucial in aqua

36 ventures to stay profitable (1). Intensive aquaculture has led to growing problems with bacterial

37 diseases and so intensive treatment with antimicrobials is required to reduce the economic losses.

38 There are several opportunistic and pathogenic microbes that infect fish, resulting in great morbidity

- 39 and mortality. Amongst such microbes are bacteria such as Aeromonas hydrophila, Edwardsiella
- 40 tarda, Flavobacterium columnare, Francisella spp., Pseudomonas spp., Mycobacterium marinum,
- 41 Mycobacterium fortuinum, Streptococcus iniae, Staphylococcus aureus (2)

42 Antibiotics at therapeutic or growth promoting levels are usually administered for short periods of 43 time orally to sets of fish that share the same culture facility. Oxytetracycline, florfenicol, and 44 ulfadimethoxine/ormetoprim are the antimicrobials authorized in United States of America for use in 45 aquaculture (3). Emerging antimicrobial resistance, due to use of antimicrobials, is a public health 46 concern in human and animal medicine worldwide. In fish farming industry, the widespread use of 47 the limited synthetic antibiotics for treating bacterial diseases has been associated with development 48 of antibiotic resistance in Aeromonas hydrophila, A. salmonicida, Edwardsiella tarda, E.icttaluri, 49 Pseudomonas spp., Vibrio anguillarum, V. salmonicida, Pasteurella piscida and Yersinia ruckeri (4, 50 5, 6). The European Union banned their use because of the risk of chemical residues in food, the 51 development of resistant pathogen strains which can be transferred from animals to humans, 52 immune suppression, destabilization of helpful bacterial populations as well as the environmental 53 pollution because up to 70-80 percent of the drug ends up in the environment (7, 8, 9, 10, 11, 12, 13, 54 14).

55 Scientists have been searching for efficacious natural alternatives to antibiotics aimed at promoting 56 animal health. Such alternatives include phytobiotics, probiotics, synbiotics and organic acid. The 57 antimicrobial activities of phyobiotics (phytogenics) such as tamarind (15, 16, 17), black pepper, 58 curry leaf, coriander (18) turmeric and ginger (19), onion and walnut leaf (20), essential oil from 59 Pakistani spices (21) and leaves, bark and root of guava among others (22) have been investigated 60 as possible alternatives to the synthetic antibiotics. The antibacterial activities from plant origin have 61 been linked to the presence of bioactive phytochemicals in such plants. Phytochemicals contain 62 secondary metabolites such as alkaloid, saponin, tannin, terpenoids and phenolic compounds which 63 have been associated with antimicrobial, antioxidants and antiinflammatory properties (23, 24).

64 Tamarindus indica Linn (tamarind), a multipurpose tree widely available in the tropics, is of great 65 importance in traditional medicine. The leaves and bark of the plants have been utilized for the 66 treatment of body pain, yellow fever and stomach disorders traditionally (15). Compounds such as 67 carvacrol, cinnamaldehyde, epicathechin, lupeol, tartaric acid are components of tamarind (25, 26, 68 and 27). (28 and 29) also reported antibacterial, antifungal, antiviral, antioxidant, carminative, 69 digestive and laxatives activities of tamarind. Most of the earlier researchers on the use of natural 70 alternatives to antibiotics had focused mainly on pathogens relating to human and terrestrial live 71 stocks. Therefore, the aim of this study was to determine the phytochemical composition and 72 antimicrobial properties of tamarind extracts against some aguatic pathogenic organisms.

73 2. MATERIAL AND METHODS

74 **2.1 Source of plant materials and preparation**

Tamarind leaf and fruit were obtained from the environment of Teaching and Research Farm
College of Agriculture, Kwara State, University, Malete. The plant parts were taken to the herbarium
of the Department of tany, University of Ibadan and the plant was identified as *Tamarindus indica*

Linn. The fruit husk was carefully removed, the pulp was scrapped from the seeds, remnant of pulp
was washed and the seed coat removed. The leaves, pulp and seed coats were air-dried under
shade.

81 2.2 Plant Extraction

82 Both the leaves and seed coats of tamarind were ground with blender, while the pulp was blend with 83 small volume of the solvent for extraction and later top up to the required volume. The extraction of 84 tamarind leaf, seed coat and pulp was carried out using mamtion method with distilled water and ethanol. Each sample was mixed with ordinary distilled water, warm distilled water at 50°C, Hot 85 distilled water at 80°C (30) and 96% ethanol at a ratio 1:10 (w/v) (31). The mixtures of plant parts 86 87 were homogenized and the kept on rotary shaker (32) for 2 days. The homogenized mixtures were 88 centrifuged (SE-CF-TDZ-WS, Labkits, U-Therm International (Hong Kong) Limited) at 4000 rpm for 30 minutes at room temperature and the supernatant collected, sieved with double layer of muslin 89 90 cloth after which it was filtered through Whatman No.4 filter paper. The solvents were removed under vacuum using a rotary evaporator (IKA[®] RV10, Artisan Technology Group, Champaign, US) at 91 92 60°C for ethanol and 90°C for water. The concentrated extracts were further dried in freeze-drier 93 (LYOTRAP, LTE Scientific Ltd., Great Britain) and kept in freezer before use.

94 2.3 Qualitative Phytochemical screening of tamarind extracts

The extract of the seed coat, the pulp and the leaves of tamarind were evaluated for qualitative
determination of major phytoconstituents which include reducing sugar, terpenoids, alkaloids,
cardiac glycosides, flavonoids, saponins and tannins as described by (33 and 34).

98 2.4 *In vitro* screening of antimicrobial activity of tamarind extracts.

99 2.4.1 Source of microorganisms

100 Pure isolates of Escherichia coli, Staphylococcus aureus, Bacillus substilis, Salmonella typhi,

101 Pseudomonas putida, Enterobacter gergovia, Hafniaalvei and Aeromonas hydrophila were obtained 102 from the laboratory stock of the Departments of Microbiology and Veterinary Medicine, University of 103 Ibadan, Nigeria. The organisms were sub- cultured on nutrient agar in plates within 24hrs at 37^oC and 104 thereafter the isolates were grown on nutrient agar slants and preserved in refrigerator at 4^oC during 105 the study.

106 2.4.2 Agar well diffusion assay

- The antimicrobial activity of aqueous and ethanolic extracts of tamarind leaf, seed coat and pulp against the aforemer ad microbes was determined as described by 35 and 36 standards. The bacteria were sub-cultured from the preserved slants for 24 hour before use. Mueller-Hinton Agar was prepared, sterilized, allowed to cool to room temperature and then poured into plates to about 4mm depth under an aseptic condition. 24-hour old culture of each test organisms was standardized to 0.5 McFarland standards (10⁶ CFU/ml). About 100µl of the standardized cell suspensions was spread on
- 113 Mueller-Hinton agar plates in triplicates. Four wells were bored on each plate with a sterile 6mm

- 114 diameter cork borer; 100 µl of the crude extracts at 10mg/ml were introduced into the wells, allowed to
- stand at room temperature for about 30 minutes. The plates were then incubated at 37^oC for 24h.
- 116 Controls were set up in parallel using the solvent used for extraction as well as two synthetic
- 117 antibiotics, Oxytetracycline and Erymycin commonly used in aquaculture and livestock imptry.
- 118 The plates were observed for inhibition zone diameter (mm).

119 2.4.3 Minimum Inhibitory Concentration of tamarind Extracts

- 120 Estimation of Minimum Inhibitory Concentration (MIC) of the tamarind extracts was carried out using
- agar dilution method. Two-fold dilutions of antimicrobial agents were prepared as described by (36)
- 122 from 10.24mg/ml of each using distilled water as diluents. Briefly, 18mls Mueller Hinton Agar (MHA)
- 123 was prepared in McCartney bottles & sterilized. The sterilize MHA was allowed to cool to 50°C in
- 124 water bath after which 2mls of each diluted antimicrobial agent was gently mixed with MHA and
- 125 poured into sterilized petri- dishes under aseptic condition. This was allowed to gel and cooled for 1
- hour. A 24-h old culture of each of the test organisms was serially diluted in 0.85% sterilized saline
- 127 water to standardize the organisms to 0.5 McFarland standards (10⁶ CFU/ml). 1ml syringe was used
- 128 to deliver 2 drops of the standardized inoculums to 100mm diameter plate equivalent to approximately
- 129 40µl per plate. The inoculum was spread on the agar surface and the plates were allowed to stand at
- 130 room temperature for about 30 minutes to ensure the moisture in the inoculum is absorbed into the
- agar. The plates were then inverted and incubated at 37°C. The plates were thereafter observed after
- 132 20 to 24-hour incubation period for growth of organism. The lowest concentration of tamarind extracts
- 133 and the synthetic antibiotics that completely inhibits growth of the inoculum was recorded as MIC.

134 2.4.4 Minimum Bactericidal Concentration (MBC) of tamarind extract

- 135 Sterile inoculating loop was used to pick from the MIC plates and streak on a sterilized MHA plate
- 136 surfaces. The inoculated plates were incubated at 37^oC for 24hour. The lowest concentration in which
- tamarind extracts and the synthetic antibiotics did not allow growth of organisms on the MHA plates
- 138 was recorded as MBC.

139 2.5 Statistical analysis

- 140 One-way Analysis of Variance (ANOVA) was used to analyze the data on zones of inhibition. Duncan
- 141 multiple range tests was used to compare differences among means at 5% probability level using
- 142 statistical software SAS (Statistical Analysis System, 2010).

143 3. RESULTS AND DISCUSSION

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145 3.1 Phytochemical constituents in tamarind extracts

- The result on phytochemical screening of tamarind extracts (Table 1) revealed presence of reducingsugar, flavonoid, saponin, terpenoids while tannin and cardiac glycosides were absent.
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Samples			Phytochemicals						
	Alkaloid	Cardiac glycosides	Flavonoids	Reducing sugar	Saponin	Tannin	Terpenoid		
LOW	+	-	+	+	+	-	+		
LWW	+	-	+	+	+	-	+		
LHW	+	-	+	+	+	-	+		
LET	+	-	+	+	+	-	+		
POW	+	-	+	+	+	-	+		
PWW	+	-	+	+	+	-	+		
PHW	+	-	+	+	+	-	+		
PET	+	-	+	+	+	-	+		
SOW	+	-	+	+	+	-	+		
SWW	+	-	+	+	+	-	+		
SHW	+	-	+	+	+	-	+		
SET	+	-	+	+	+	-	+		

153 Table 1: Results of qualitative phytochemical screening of tamarind extract

 154
 LOW = Leat Ordinary Water
 LWW = Leat Warm Water LHW = Leat Hot Water
 LE = Leat Ethanol
 POW = Pulp Ordinary Water
 PWW = Pulp Warm

 155
 Water
 PHW = Pulp Hot Water
 PET = Pulp Ethanol
 SOW = Seed
 Coat Ordinary Water
 SWW = Seed
 Coat Hot Water
 SCET =

 156
 Seed
 Coat
 Ethanol
 SCET =
 Seed
 Coat
 Ethanol

157 3.2 Antimicrobial activities of tamarind extracts

158 3.2.1 Zones of inhibition of tamarind extracts

159 Table 2 shows the results of zone of inhibition of the tamarind extracts compared to the synthetic

160 antibiotics. The synthetic antibiotics used had significantly higher (P = .05) zones of inhibition than the

161 tamarind extracts for all the test organisms. Tamarind Pulp Hot Water (PHW) extract significantly

162 inhibited Aeromonas hydrophila better than other extracts while Leaf Ethanolic (LET) extract had the

163 lowest zone of inhibition against A. hydrophila. Leaf Warm Water (LWW) extracts had significantly

164 higher (*P* = .05) zone of inhibition against *Pseudomonas putida*. Higher significant zone of inhibition

165 was also exhibited by PHW extract against *Hafnia alvei*. The zones of inhibition of LWW, LHW, PHW

166 extracts against *Escherichia coli* were significantly higher (*P* = .05) than other extracts while the seed

167 coat showed no antimicrobial activities against *E. coli* and *Bacillus subtilis*. Pulp Ethanol Extract (PET)

had significantly higher (*P* = .05) inhibition (12.00mm) against *Salmonella typhi*.

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TE/A			Path	logens				
	A. hydrophila	P. putida	E. gergovia	H. alvei	E. coli	S. aureus	B. subtilis	S. typhi
Solvents	0.00 [†]	0.00 ^h	0.00'	0.00 ^ĸ	0.00 [†]	0.00 ^c	0.00 ^g	0.00 ^h
ERY	21.33 ^ª	22.00 ^a	26.00 ^a	15.00 ^b	32.67 ^a	28.33 ^a	26.67 ^a	25.67 ^b
OTC	22.33 ^ª	13.33 [⊳]	15.00 ^b	21.00 ^a	25.67 ^b	27.67 ^a	22.00 ^b	26.33 ^ª
LOW	9.67 ^{de}	9.33 ^{etg}	9.67 ^{de}	11.00 ^g	9.33 ^e	9.67 ^b	10.00 ^c	10.00 ^e
LWW	11.67 ^c	12.00 ^c	11.00 ^c	11.67 ^e	10.67 ^{cd}	11.00 ^b	11.00 ^c	10.00 ^e
LHW	10.67 ^{cd}	10.00 ^{de}	11.00 ^c	11.00 ^g	11.00 ^{c.}	11.00 ^b	11.67 ^c	11.00 ^d
LET	8.67 ^e	8.67 ^{tg}	9.33 ⁹	10.00 ^j	9.00 ^e	9.00 ^b	9.00 ^t	9.00 ^g
POW	10.33 ^{cde}	9.67 ^{def}	10.00 ^e	11.33 ^t	9.00 ^e	9.00 ^b	9.67 ^{ef}	10.00 ^e
PWW	10.67 ^{cd}	10.00 ^{de}	10.00 ^d	11.67 ^e	9.33 ^e	9.00 ^b	9.67 ^c	11.00 ^d
PHW	13.33 ^b	10.67 ^d	11.00 ^c	13.33°	11.33 [°]	9.00 ^b	9.33°	11.00 ^d
PET	9.67 ^{de}	9.00 ^{efg}	10.00 ^d	12.00 ^d	9.67 ^e	9.00 ^b	9.00 ^c	12.00 ^c
SOW	9.33 ^{de}	8.33 ^g	9.00 ^t	10.00 ^j	0.00 [†]	9.00 ^b	0.00 ^e	9.67 [†]
SWW	10.33 ^{cde}	9.00 ^{efg}	9.00 ^t	10.67 ^h	0.00 ^t	9.00 ^b	0.00 ^e	9.67 [†]
SHW	11.00 ^{cd}	9.67 ^{det}	10.00 ^d	10.33'	0.00 ^t	9.00 ^b	6.00 ^d	10.00 ^e
SET	11.00 ^{cd}	9.00 ^{etg}	11.33 [°]	10.00 ^J	0.00 [†]	9.00 ^b	0.00 ^e	9.00 ^g
SEM	0.578	0.423	0.158	0.274	0.428	0.765	0.942	0.318

175 **Table 2: Antagonistic activity (mm) of synthetic antibiotics and tamarind extracts at 10mg/ml** 176 **against some pathogens**

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Means with the same letter on the same row are not significantly different at ${\cal P}$ =

 TE/A = Tamarind extracts/Antibiotics ERY= Erythromycin
 OTC = Oxytetracycline
 LOW = Leaf Ordinary Water
 LWW = Leaf Warm Water
 LHW = Leaf Hot Water
 LET

 179
 = Leaf Ethanol
 POW =
 Pulp Ordinary Water
 PWW = Pulp Warm Water
 PHW =
 Pulp Hot Water
 PET =
 Pulp Ethanol
 SOW =
 Seed
 Coat Ordinary Water
 SWW =

 180
 Seed
 Coat
 Hot Water
 SET =
 Seed
 Coat
 Enterobacter
 H =
 Hafnia
 E =

 180
 Seed
 Coat
 Hot Water
 SET =
 Seed
 Coat
 Enterobacter
 H =
 H =
 Hafnia
 E =

 181
 Escherichia
 S =
 Salmonella

182 3.2.2 Minimum Inhibitory Concentration (MIC) and Minimum Bactericidal 183 Concentration (MBC) of tamarind extract

184 MIC and MBC of aqueous and ethanolic extracts of tamarind against the test organisms in

185 comparison to erythromycin and oxytetracycline are shown in Table 3. The lowest MIC, 0.64mg/ml,

186 was obtained for oxytetracycline and erythromycin against *E. gergovia* and *E. coli* respectively.

187 Amongst the leaf extracts, LWW extract exhibited lower MIC against all the test organisms except for

188 *P. putida* and *S. typhi* against which higher values were obtained. Similar value of MIC, 2.56mg/ml,

189 was also exhibited by all tamarind pulp extracts against the test organisms except for *P. putida* while

190 higher values of MIC were obtained from seed coat extracts. MBC value of 1.28mg/ml was obtained

191 for oxytetracycline against A. hydrophila and MIC of 2.56 against S. typhi. MBC value of 2.56mg/ml

- 192 was similarly exhibited by POW and PWW extracts against *S. typhi* and PHW and PET extracts
- against *E. coli* and *S. aureus* but MBC was not exhibited by the synthetic antibiotics against these
- 194 three pathogens.
- 195 Table 3: Minimum Inhibitory Concentration and Minimum Bactericidal Concentration (mg/ml)
- 196 of two synthetic antibiotics and tamarind extracts against some pathogens

TE/A			Patho	ogens				
	A. hydrophila	P. putida	E. gergovia	H. alvei	E. coli	S. aureus	B. subtilis	S. typhi
ERY	1.28	1.28	1.28	2.56	0.64	1.28	2.56	5.12
OTC	1.28*	2.56	0.64	1.28	1.28	1.28	1.28	2.56
LOW	10.24	10.24	5.12	5.12	5.12	5.12	5.12	5.12
LWW	2.56	10.24	2.56	2.56	2.56	2.56	2.56	5.12*
LHW	5.12	5.12	2.56	2.56	2.56	5.12	5.12	5.12*
LET	2.56	5.12	2.56	2.56	5.12	10.24	5.12	5.12*
POW	2.56	5.12	2.56	2.56	2.56	2.56	2.56	2.56*
PWW	2.56*	5.12	2.56	2.56	2.56	2.56	2.56*	2.56*
PHW	2.56	5.12	2.56*	2.56	2.56*	2.56*	2.56	2.56
PET	2.56	5.12	2.56	2.56	2.56*	2.56*	2.56	2.56
SOW	10.24	10.24	5.12	10.24	10.24	NA	10.24	NA
SWW	10.24	10.24	5.12	5.12	5.12	5.12	5.12	5.12
SHW	5.12	5.12	5.12	5.12	5.12	5.12	5.12	5.12
SET	10.24	10.24	10.24	10.24	10.24	NA	10.24	10.24*

 197
 "Minimum Bactericidal Concentration NA- Not Active at the highest concentration used TE/A = Tamarind extracts/Antibiotics EHY= Erythromycin OTC = Oxytetracycline

 198
 LOW = Leaf Ordinary Water LWW = Leaf Warm Water LHW = Leaf Hot Water LET = Leaf Ethanol
 POW = Pulp Ordinary Water PWW = Pulp Warm Water PHW

 199
 = Pulp Hot Water PET = Pulp Ethanol
 SOW = Seed Coat Ordinary Water SWW = Seed Coat Warm Water SHW = Seed Coat Hot Water SET = Seed Coat

 200
 Ethanol A = Aeromonas P = Pseudomonas E = Enterobacter H = Hafnia E = Escherichia S = Staphylococcus B = Bacillus S = Salmonella

202 **3.2.3 Discussion**

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203 This result of the phytoconstituents of tamarind in this study is similar to what has been reported by 204 other researchers (15, 16, and 37) on the phytoconstituents of tamarind. However, the absent of 205 tannins from this study is contrary to other reports. Antibacterial activity obtained from tamarind 206 extracts in this study coincided with the reports of other researchers who studied antibacterial 207 activities of phytogenics. (15) similarly reported higher zones of inhibition from synthetic antibiotic 208 compared with tamarind extracts; however higher zone of inhibition and lower MIC values were 209 obtained from leaf extract in this study. Also in contrast to (15) not all extracts in this study exhibit 210 bactericidal (MBC) activity. (16) also reported better antibacterial activity of aqueous tamarind pulp

211 extract compared to the ethanolic extract against *P. aeroginosa*. Furthermore, lack of clear or lower

- 212 zone of inhibition discovered in this study against *S.typhi* and *S.aureus* coincided with the observation
- 213 of (10)

Higher zones of inhibition obtained from oxytetracycline compared to tamarind extracts against the 214 215 test organisms is in agreement with the report of (17) but lower MIC values were obtained from the 216 tamarind aqueous extracts against A. hydrophila and S. aureus than what the researchers reported 217 from clove aqueous extract. Contrary to the absence of antimicrobial activity reported (19) on turmeric 218 and ginger root aqueous extract, tamarind pulp and leaf aqueous extract exhibit antibacterial activities 219 against Pseudomonas and E. coli. The absence of antibacterial activities of the seed coat extracts 220 against E. coli and B. subtilis is similar to the observation of (20) on lack of antibacterial activities of 221 onion bulb and walnut leaf extracts against B. subtilis and E. coli respectively. 222 Lower MIC values were discovered in this study from all tamarind pulp extracts as well as warm and

- hot aqueous leaf extracts than what was reported (21) on Pakistani spices against *E. coli* and *S*
- aureus. The MIC value of the plant extracts examined (22) against *E. coli* is similar to 2.5mg/ml
- obtained in this study while lower value was obtained in this study against *A. hydrophila* than what the
- authors reported. Generally, the isolates investigated in this study were more sensitive to warm and hot water extracts than to ordinary water extracts and ethanolic extracts. The higher antibacterial
- 228 activity of the warm and hot aqueous extracts in this study might be an indication of higher solubility of
- 229 phytoconstituents in water at higher temperature than lower temperature. The demonstration of better
- antibacterial activity from warm and hot aqueous extract provides the scientific basis for the boiling of
- 231 herbs by the traditional folks in disease treatments. The use of the aqueous extracts is of better
- economic advantage for fish farmer because ethanol is more costly than distilled water.

233 4. Conclusion

The antibacterial activity demonstrated by tamarind extracts in this study shows that the extracts could be used to control bacterial associated with the aquatic environment and fish products. The discovery from this study is an indication that tamarind pulp and leaf warm/hot extract could be used as possible phytogenic to control *Aeromonas* and *Pseudomonas* infection in fish as well as protect fish products from poisoning organisms. Further study is however needed on the concentration of tamarind extracts that would be as effective as synthetic antibiotics and the *in vivo* investig

- warm and hot aqueous extract of tamarind leaf and pulp.

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