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2 **ANTIMICROBIAL POTENTIAL OF THE CRUDE**
3 **EXTRACTS AND PEPTIDE FRACTIONS OF TWO**
4 **MARINE MOLLUSCS: *Tympanotonus fuscatus***
5 **VAR RADULA (LINNEAUS) AND *Pachymelania***
6 ***aurita* (MULLER)**
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12 **ABSTRACT**
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The abstract should be concise and informative. It should not exceed 300 words in length. It should briefly describe the purpose of the work, techniques and methods used, major findings with important data and conclusions. Different sub-sections, as given below, should be used. No references should be cited in this part. Generally non-standard abbreviations should not be used, if necessary they should be clearly defined in the abstract, at first use.

Aims:

Study Design:

Place and Duration of Study: Department of Microbiology, Obafemi Awolowo University, Ile-Ife, Nigeria

Methodology: The antimicrobial activity of the whole body aqueous and acetone-methanol extracts of *T.fuscatus* Var Radula and *P.aurita*, collected from the Niger-Delta region of Nigeria, were evaluated based on inhibition zone diameter using the agar well diffusion method against ten bacterial isolates and *C.albicans*. These organisms were further used in the TLC bioautography experiment. The peptide fraction from the organic extracts of both organisms was obtained by Molecular sieve chromatography on Sephadex LH20. Peaks obtained were pooled and further analysed on TLC. A simple contact TLC bioautographic procedure was used to detect the number of antibacterial and antifungal peptides present in the extracts of both *T.fuscatus* and *P.aurita*.

Results: The aqueous extract of both *T.fuscatus* and *P.aurita* had no antimicrobial effect against the test microorganisms whereas the acetone-methanol extract showed broad-spectrum antibacterial activity against five bacterial isolates at the highest concentration (100 mg/ml). It also showed inhibition against *C. albicans* at this concentration (100mg/ml).

All the peptides exhibited bactericidal activity against the five test bacterial isolates and bacteriostatic activity against *C.albicans*. This activity was denoted by inhibition of growth in the region in which the peptides on the TLC plate made contact with the agar containing the isolates

Conclusion:

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17 **1. INTRODUCTION (ARIAL, BOLD, 11 FONT, LEFT ALIGNED, CAPS)**
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19 In general, marine molluscs are soft bodied and sessile and often live in microbe-
20 rich habitats. Hence, molluscs are quite exposed to pathogens and do not often possess a
21 proper physical barrier against microbial infection. This suggests that molluscs must have
22 evolved alternative biological defence strategies and systems, including the secretion of
23 mucus containing a range of different antibacterial, antifungal, antiparasitic and antiviral
24 secondary metabolites to protect themselves against an onslaught of microbial invasion from
25 their environment (Benkendorff, 2010 and Datta *et al.*, 2015). One of the defence strategies
26 used by mollusc, and indeed, a host of other organisms is the production of antimicrobial
27 peptides (AMPs) also known as host defence peptides (HDPs). AMPs play key roles in
28 innate immunity and they had been observed from a wide variety of organisms in the last few
29 years. Research has been focused on the discovery and exploitation for health benefits of
30 AMPs and other peptides with antimicrobial activity. This is partly driven by the need for new
31 antibiotics, which is in turn due to the emerging threat of antibiotic resistance.

32 AMPs are ribosomally synthesized from proteinogenic amino acids. They are short,
33 generally positively charged, potent, broad spectrum antibiotics and have been
34 demonstrated to kill both gram positive and gram negative bacteria (Mahlapuu *et al.*, 2016).
35 Unlike antibiotics, which target specific cellular activities, AMPs target the
36 lipopolysaccharide layer of cell membrane, which is unique to microorganisms. The high
37 cholesterol level and negative charge of the bacterial membrane ensure that eukaryotic cells
38 are usually not targets of many AMPs (Bahar and Ren, 2013; Bechinger and Gorr, 2016;
39 Mahlapuu *et al.*, 2016). Another important feature of AMPs is their fast killing ability. Some
40 AMPs can kill in seconds after the initial contact with cell membrane (Bahar and Ren, 2013).
41 In addition to their role as endogenous antibiotics, some AMPs contribute to inflammation as
42 well as exhibit immunomodulatory activities (Mahlapuu *et al.*, 2016). Hence, these AMPs act
43 indirectly to kill microorganisms by modulating the host defense systems. Some other AMPs
44 kill bacteria by inhibiting some important pathways inside the cell such as DNA replication
45 and protein synthesis (Bahar and Ren, 2013).

46 Their preferential attack on the cell membrane or cell wall of bacterial and fungi ensure that
47 AMPs should not cause widespread resistance. In cases where specific protein targets are
48 involved, the possibility exists for genetic mutations and bacterial resistance. However,
49 evidence suggests that this is a rare event which can be overcome by subtle structural
50 modifications made to the AMP (Bechinger and Gorr, 2016).

51 Other activities that AMPs have been shown to possess include: anti-biofilm, wound repair
52 and anticancer activities. They are also known to play a role in regulation of the adaptive
53 immune system (Datta *et al.*, 2015; Haney *et al.*, 2017). Hence, several AMPs are currently
54 being evaluated in clinical trials, not only as novel antibiotics (Bechinger and Gorr, 2016), but
55 also as new pharmacological agents to modulate the immune response, promote wound
56 healing, and prevent post-surgical adhesions.

57 This study was undertaken to evaluate the antimicrobial potential of the crude extracts as
58 well as the peptide fraction of *Tympanotonus fuscatus* Var Radula and *Pachymelania aurita*.
59 *Tympanotonus fuscatus* and *Pachymelania aurita* are two of the most common molluscs
60 species which inhabit the mangroves of the Niger Delta region of Nigeria although they are
61 also found along the coast of West Africa as well as in Angola and Gabon. Their flesh is
62 used in the preparation of delicacies and although related species feature in a range of
63 traditional natural remedies, including wound healing and stomach upsets, and have been
64 evaluated for their antimicrobial properties, these two species have not been assessed to
65 determine if they possess antimicrobial activity. This is the first known study to investigate
66 the antimicrobial potential of whole body extracts of *T.fuscatus* Var Radula and *P.aurita*,
67 obtained from the Niger Delta region of Nigeria.
68

69 2. MATERIAL AND METHODS

70 2.1. Microorganisms

71 Microbial isolates were obtained from the National Collection of Industrial Food and Marine
72 Bacteria (NCIB), UK and the American Type Culture Collection, ATCC, Rockville, MD, USA.
73 The bacteria strains used for this study were: *Proteus Vulgaris* (NCIB 67); *Pseudomonas*
74 *aeruginosa* (NCIB 950); *Bacillus Subtilis* (NCIB 3610); *Staphylococcus aureus* (ATCC
75 43300); *Escherichia coli* (NCIB 86); *Micrococcus luteus* (NCIB 196); *Klebsiella Pneumoniae*
76 (NCIB 418); *Clostridium sporogenes* (NCIB 532); *Bacillus Stearothermophilus* (NCIB 8222);
77 *Serratia marcescens* (NCIB 1377) while the yeast strain used was *Candida albicans*.

78 2.2. Sample Collection

79 Live *Tympanotonus fuscatus var radula* and *Pachymelania aurita* were purchased from the
80 Oron Beach Market, Oron, Akwa Ibom State, Nigeria (GPS coordinates: 4°49'37.6''N
81 8°14'04.4''E). The molluscs were washed thoroughly to remove mud and then deshelled to
82 collect both their flesh and hemolymph.

83 2.3. Preparation of acetone-methanol extracts

84 The alcohol extracts of *T.fuscatus* and *P.aurita* were prepared using the method
85 described by Eghianruwa et al., 2019. 200 g of mollusc flesh in its hemolymph was
86 macerated using a blender and extracted twice with 1 L acetone for both cycles. Each cycle
87 of extraction with acetone was carried out at room temperature for 12hrs with constant
88 stirring using a magnetic stirrer and the homogenate was filtered using a muslin cloth. After
89 acetone extraction, the biomass residue of the sample was subjected to two cycles of
90 extraction using a total of 1500 ml of methanol. The Acetone and methanol fractions were
91 combined and concentrated by evaporation using a rotary evaporator at 40 °C then stored at
92 4°C.

93 2.4 Preparation of the Aqueous Extracts

94 200 g of mollusc flesh in its hemolymph was homogenized with 2000 ml of Phosphate
95 buffered saline; PBS, pH 7.2 (0.1M Sodium chloride in 0.025M Sodium dihydrogen
96 orthophosphate with 0.1 M PMSF) using a blender. The homogenate was left to extract for
97 48 hours at 4 °C after which it was centrifuged at 10,000 g using a cold centrifuge, freeze
98 dried and stored at 4°C.

99 2.5 Sensitivity Test

100 The antimicrobial activity of the extracts was carried out using the agar-well diffusion
101 method as described by Adegoke *et al.*, 2010 with some modifications. The bacterial strains
102 used were first grown on nutrient agar for 18hrs before use. The turbidity of the 18 hr old
103 culture was adjusted to 0.5 McFarland Standards (106 cfu/mL) in sterile normal saline. The
104 inoculum was then seeded onto sterilized Mueller-Hinton agar using a sterile swab stick.
105 Wells were made in the seeded plates using a sterile 6mm cork borer. The wells were filled
106 up with known concentrations of the extracts (25, 50 and 100mg/ml) using a micropipette.
107 Care was taken to avoid spillage of the extract onto the surface of the medium. The plates
108 were allowed to stand on the work bench for 1hr to allow proper inflow of the extract
109 solutions into the medium before incubating in an incubator at 37°C for 24 hr following which
110 the plates were observed for zones of inhibition. The effects of the extracts of *T.fuscatus* and
111 *P.aurita* on the bacterial strains were compared to a standard antibiotic (Streptomycin).
112 Water and a mixture of acetone-methanol (1:1) were also used as control in the experiment
113 to confirm that any activity observed is as a result of the extract and not the solvents used in
114 extraction.

115 2.6. Molecular Sieve chromatography on Sephadex LH20

116 The peptide fractions from the alcohol extracts of *P.aurita* and *T.fuscatus* were obtained via
117 molecular sieve chromatography on a Sephadex LH20 column. Sephadex LH20 resin (40g)
118 was swollen at room temperature in 200 ml of absolute methanol for 5hrs. The slurry was
119 stirred every hour during this time and fine particles were removed by decantation. The
120 slurry was packed into a column (10 × 1.5 cm) according to instructions contained in the

121 Pharmacia laboratory techniques manual. The packed column was equilibrated with 300 ml
122 PBS, pH 7.2. Crude aqueous extract (2.5 ml) of either *P.aurita* and *T.fuscatus* were applied
123 on the column and eluted with 1 column volume of methanol and fractions (1 ml) were
124 collected at a flow rate of 10 ml/hr. Peptide-containing fractions were detected by means of
125 TLC

126 2.7 TLC Bioautography

127 Analytical thin layer chromatography was used to detect the peptide containing
128 fractions from the Molecular sieve experiment using the method as described by (Osoniyi
129 and Onajobi, 1998). The TLC was carried out on aluminium-backed silica 60 F254 gel plates
130 (10cm X 8cm), using a solvent system of butanol: acetic acid: water (3: 1: 1) as the mobile
131 phase while the detection stain was 0.2 % Ninhydrin in ethanol. Plates were activated in an
132 oven at 100-120 °C for 1hr and allowed to cool before use. The solvent system was
133 prepared fresh 15mins before each run, placed in the tank, swirled and allowed to saturate
134 the tank before the run. 5 µl of each fraction obtained from the molecular sieve procedure
135 was spotted on the plate and allowed to dry before placing in the tank for the run. After the
136 run, the plate was air dried, sprayed with the ninhydrin stain and then dried in the oven at
137 100 °C for 15mins to allow for colour development. Plates used in bioautography were not
138 sprayed with the stain.

139 The bioautography assay was carried out in order to determine which peptide band
140 exhibits antimicrobial activity. Agar plates inoculated with the microbial strains were prepared
141 as described above for the sensitivity testing with the exception of the holes. The alcohol
142 extracts of either *P.aurita* or *T.fuscatus* were run on a TLC as described above. The
143 unstained TLC plates were placed on the Agar plates and the plates were allowed to stand
144 on the work bench for 1hr to allow proper diffusion of the peptides on the TLC plates into the
145 medium before incubating in an incubator at 37°C for 24 hr following which the agar plates
146 were observed for zones of inhibition. The plates were incubated for a further 24 hr after
147 which the TLC plates were removed with a pair of forceps. After the removal of the TLC
148 plates, the agar plates were incubated for another 48 hrs to check for microbial growth.

151 3. RESULTS AND DISCUSSION

152 The results showing the length of the zones of inhibition of the various bacterial
153 species by the extracts are shown in Table 1. The aqueous extracts of *P.aurita* and
154 *T.fuscatus* do not exhibit any antimicrobial activity against the organisms tested, even at the
155 highest concentration tested (100 mg/ml). Alcohol extracts of both *P.aurita* and *T.fuscatus*,
156 on the other hand demonstrated antimicrobial activity. A zone of inhibition measuring
157 approximately 8 mm was observed in the culture of *S.aureus* and *B.stereothermophilus*,
158 when treated with 25 mg/ml of the alcohol extract of *P.aurita* (PAAC). At 100 mg/ml, longer
159 zones of inhibition were observed in the cultures of *S.aureus* and *B.stereothermophilus*,
160 *M.luteus*, *C.sporogenes* and *K.pneumoniae*. Zones of Inhibitions were also observed in the
161 cultures of *M.luteus*, *C.sporogenes*, *K.pneumoniae*, *B.stereothermophilus* and *S.aureus*,
162 treated with the alcohol extract of *T.fuscatus* (TFAC) but only at a concentration of 100
163 mg/ml of extract. Zones of inhibition (12mm) were also observed in the culture media of
164 *C.albicans* treated with the crude alcohol extracts of *P.aurita* and *T.fuscatus* at a
165 concentration of 100mg/ml of extract. Hence, the bioautography experiments were carried
166 out using cultures of *M.luteus*, *C.sporogenes*, *K.pneumoniae*, *B.stereothermophilus*,
167 *S.aureus* and *C.albicans*.

173 Table 1. Zones of inhibition values (mm) from the sensitivity testing of the crude aqueous
 174 and alcohol extracts of *P.aurita* and *T.fuscatus*. Values are expressed as mean \pm SEM, n=2.
 175 Streptomycin (1mg/ml) was used as the standard control.

Isolates	PAAC			TFAC			Control Streptomycin (1mg/ml)
	25 mg/m l	50 mg/ml	100 mg/ml	25 mg/m l	50 mg/ml	100 mg/ml	
<i>P.vulgaris</i> (NCIB 67)	0 \pm 00	0 \pm 00	0 \pm 00	0 \pm 00	0 \pm 00	0 \pm 00	25 \pm 00
<i>Ps. Aeruginosa</i> (NCIB 950)	0 \pm 00	0 \pm 00	0 \pm 00	0 \pm 00	0 \pm 00	0 \pm 00	25 \pm 0.6
<i>B.subtilis</i> (NCIB 3610)	0 \pm 00	0 \pm 00	0 \pm 00	0 \pm 00	0 \pm 00	0 \pm 00	27 \pm 0.4
<i>S.aureus</i> (ATCC 43300)	8 \pm 1.3	12 \pm 0.7	12 \pm 1.4	0 \pm 00	0 \pm 00	11.5 \pm 0. 7	27 \pm 1.7
<i>E.coli</i> (NCIB 86)	0 \pm 00	0 \pm 00	0 \pm 00	0 \pm 00	0 \pm 00	0 \pm 00	30 \pm 00
<i>M.luteus</i> (NCIB 196)	0 \pm 00	0 \pm 00	12 \pm 1.4	0 \pm 00	0 \pm 00	12.0 \pm 1. 4	30 \pm 00
<i>K.pneumoniae</i> (NCIB 418)	0 \pm 00	0 \pm 00	11.5 \pm 0. 7	0 \pm 00	0 \pm 00	11.5 \pm 0. 7	25 \pm 1.3
<i>C.sporogenes</i> (NCIB 532)	0 \pm 00	0 \pm 00	11.5 \pm 0. 7	0 \pm 00	0 \pm 00	11.0 \pm 00	25 \pm 0.7
<i>B.stereothermophilus</i> (NCIB 8222)	8 \pm 0.6	12 \pm 00	10.5 \pm 0. 7	0 \pm 00	0 \pm 00	10.5 \pm 0. 7	30 \pm 00
<i>S.marcescens</i> (NCIB 1377)	0 \pm 00	0 \pm 00	0 \pm 00	0 \pm 00	0 \pm 00	0 \pm 00	25 \pm 00
<i>c.albicans</i>	0 \pm 00	0 \pm 00	12 \pm 00	0 \pm 00	0 \pm 00	12 \pm 00	25 \pm 00

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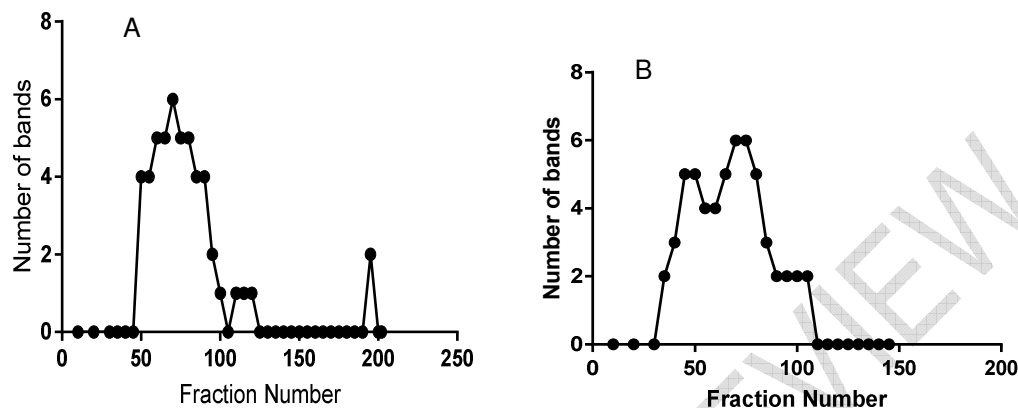
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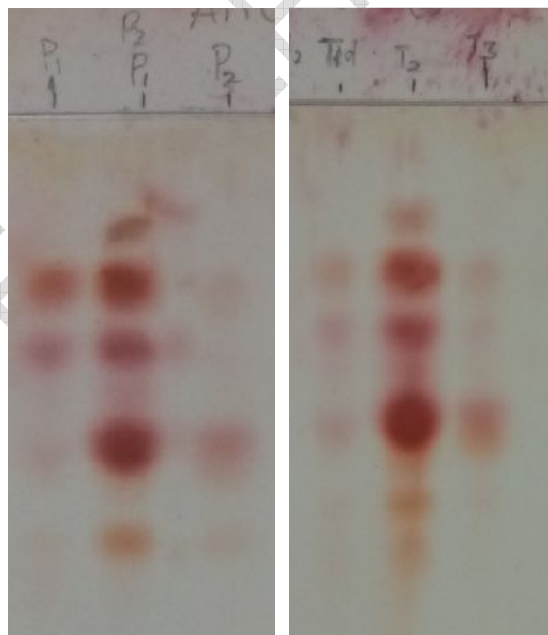
181 The results of the molecular sieve chromatography of the crude alcohol extract of
 182 *P.aurita* and *T.fuscatus* is illustrated in Figure 1. Three peptide peaks were detected by
 183 spotting each fraction in TLC. Each peak was observed to contain several peptide molecules
 184 (Figure 2). However, peak 2 obtained from both extracts (PAAC and TFAC) appear to
 185 possess similar peptides.



186

187 **Figure 1:** Elution profile from thin layer chromatography of fractions obtained from molecular
 188 sieve chromatography of the alcohol extract of *P.aurita* (A) and *T.fuscatus* (B) on Sephadex
 189 LH20. Eluant for the molecular sieve chromatography was methanol. The major peaks were
 190 pooled separately and used for the bioautography experiment

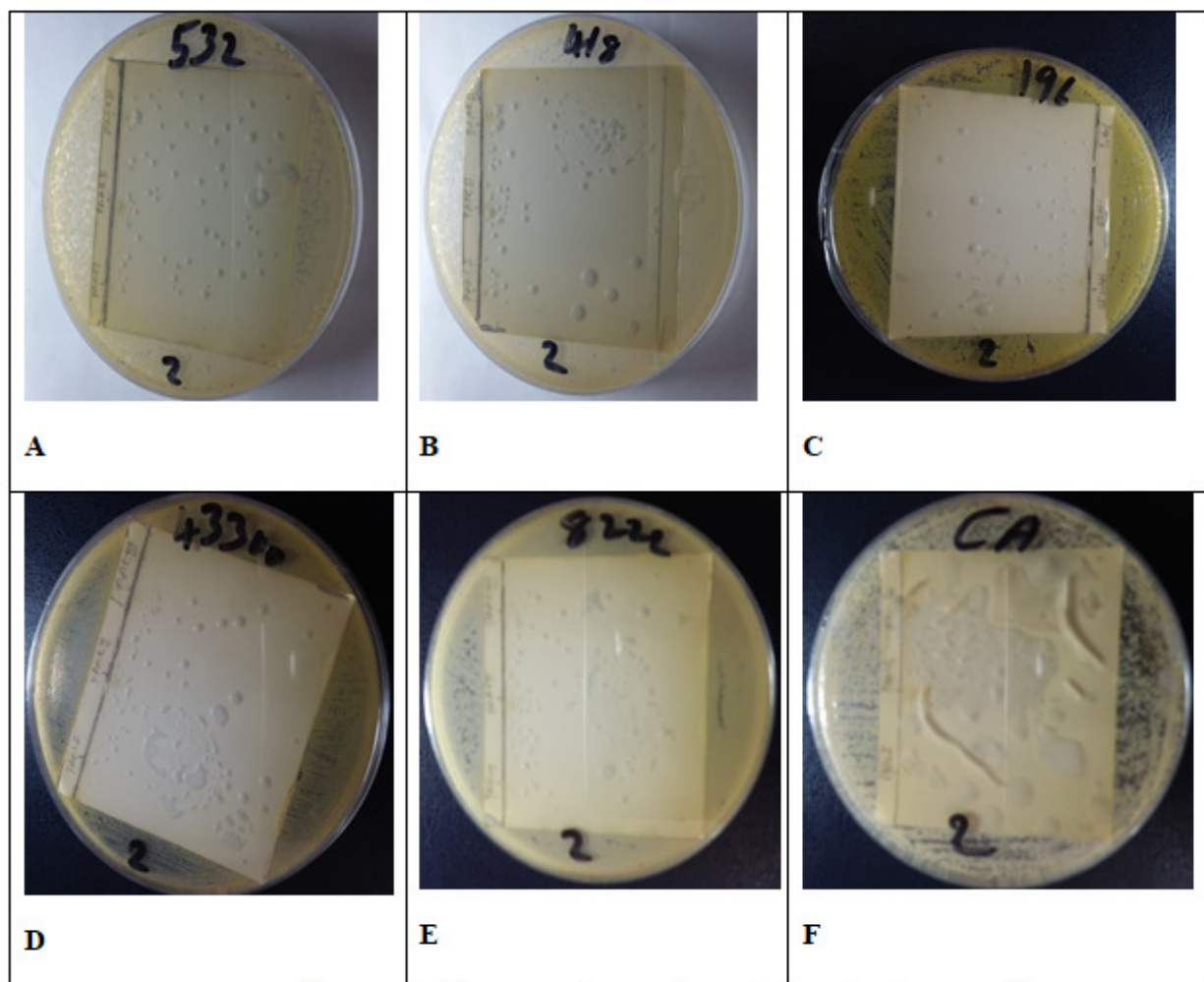
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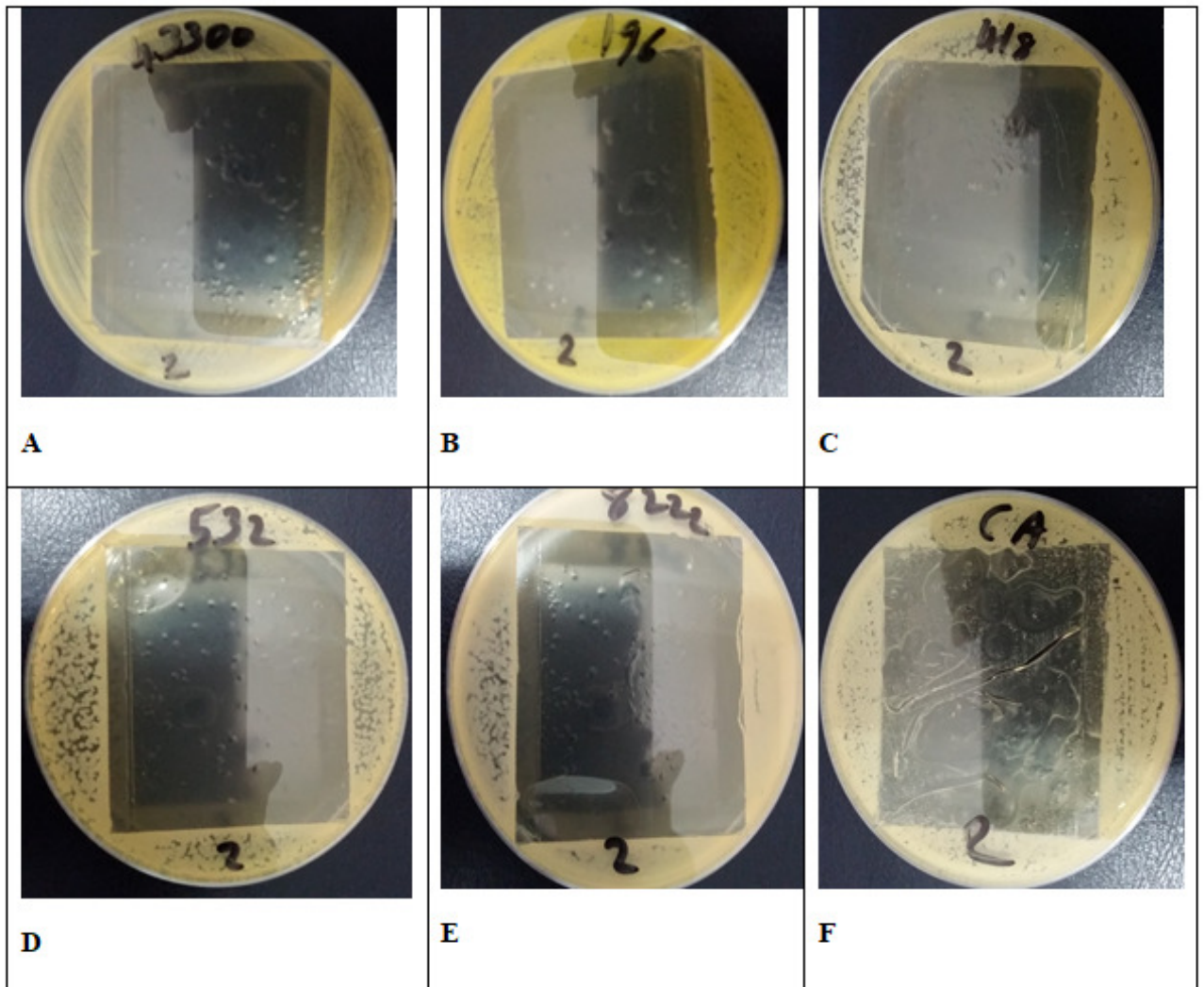
196 **Figure 2:** TLC chromatogram of the three peaks pooled from the molecular sieve
 197 chromatography separation of the alcohol extract of *P.aurita* (P1-P3) and *T.fuscatus* (T1-T3).
 198 Replicate plates, which was not stained after the TLC run, was used for the bioautography
 199 experiments

200 Resolved but unstained TLC plates of the analysed peptide peaks were placed in
201 cultures of *M.luteus*, *C.sporogenes*, *K.pneumoniae*, *B.stereothermophilus*, *S.aureus* and
202 *C.albicans*. After 24hrs of the start of the bioautography experiment, no microbial growth was
203 observed under the plates (Figure 3). Slight zones of inhibition were observed in the cultures
204 of *M.luteus* and *Bacillus Stearothermophilus* treated with TFAC. After 48 hrs, the TLC plates
205 were removed and observed for microbial growth (Figure 4). Microbial growth was observed
206 in the area previously covered by the TLC plate in the culture for *C.albicans*, although the
207 growth was slight and not as profuse as in the rest of the culture. The culture dishes were
208 incubated further for 48 hrs after the removal of the TLC plates (Figure 5). Slight microbial
209 growth was observed in most of the cultures albeit less than in the rest of the culture plate.
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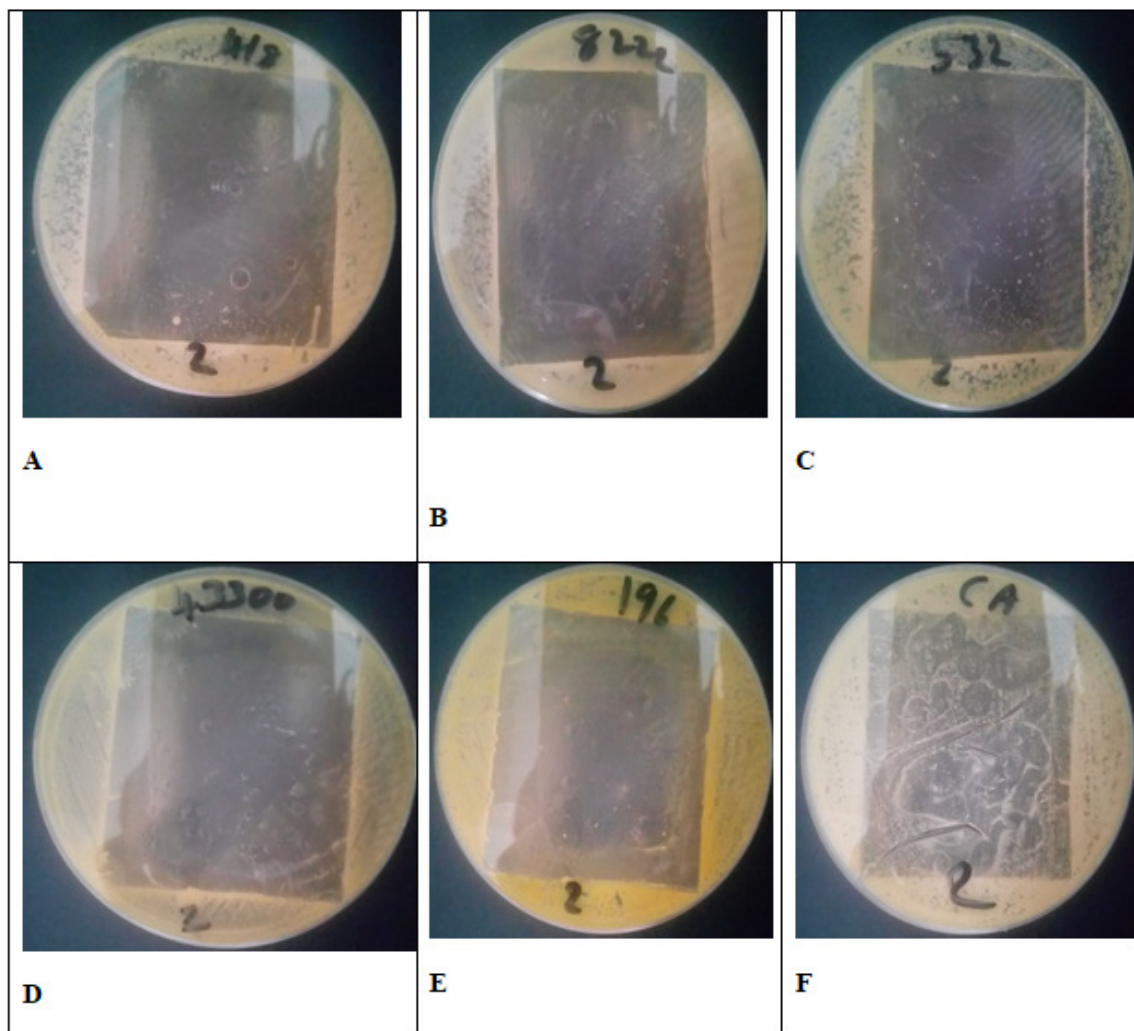
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212 **Figure 3:** Stage 1 of the contact bioautography experiment (24hrs after the start of the
213 experiment) of the peptides obtained from molecular sieve of the crude alcohol extract of
214 *P.aurita* on sephadex LH20. The Agar plates were inoculated with the microbial strains;
215 *S.aureus* (43300), *B.stereothermophilus* (8222), *M.luteus* (196), *C.sporogenes* (532),
216 *K.pneumoniae* (418) and *C.albicans* (CA). Similar results were observed for *T.fuscatus*



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Figure 4: Stage 2 of the contact bioautography experiment (48 hrs after the start of the experiment) involving the peptides obtained from molecular sieve of the crude alcohol extract of *P.aurita* on sephadex LH20 and peaks resolved on TLC. The Agar plates were inoculated with the microbial strains; *S.aureus* (43300), *B.stereothermophilus* (8222), *M.luteus* (196), *C.sporogenes* (532), *K.pneumoniae* (418) and *C.albicans* (CA). Similar results were observed for *T.fuscatus*



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Figure 5: Stage 3 of the bioautography experiment (96 hrs after the start of the experiment) involving the peptides obtained from molecular sieve of the crude alcohol extract of *P.aurita* on sephadex LH20 and peaks resolved on TLC. The Agar plates were inoculated with the microbial strains; *S.aureus* (43300), *B.stereothermophilus* (8222), *M.luteus* (196), *C.sporogenes* (532), *K.pneumoniae* (418) and *C.albicans* (CA). Similar results were observed for *T.fuscatus*

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239 To survive their microbe-rich environment, molluscs must have evolved alternative
240 biological defense strategies and systems, including the secretion of mucus containing a
241 range of different antibacterial, antifungal, antiparasitic and antiviral secondary metabolites
242 to protect themselves against an onslaught of microbial invasion from their environment
243 (Benkendorff, 2010 and Datta *et al.*, 2015).

244 The observation, from this study, that only the alcohol extracts exhibited
245 antimicrobial activity against any of the isolates tested, are in line with evidence from
246 previous studies which have reported that the compounds responsible for antimicrobial
247 activity are mainly non-polar in nature (Masoko and Eloff, 2006; Suleiman *et al.*, 2010),
248 hence alcohols (especially methanol) are a better solvent system for more reliable extraction
249 of antimicrobial molecules from natural sources compared to other solvents (Karaman *et al.*,
250 2003; Santhiya and Sanjeevi, 2014; Seleman and Amri, 2015). Indeed, in the case of marine
251 organisms, non-polar antimicrobial molecules would be most desirable as these molecules
252 would not easily interact with the surrounding water environment and be washed easily off
253 the organism.

254 In isolating molecules with antimicrobial activity, bioautography is a very useful
255 technique as the Rf of the active compounds can be used in bioassay guided fractionation
256 instead of requiring labour intensive determination of activity of several fractions (Suleiman
257 *et al.*, 2010; Dewanje *et al.*, 2014). This also ensures that the molecule isolated at the end is
258 the same one that was present in the extract and not an artefact of the isolation procedure
259 (Suleiman *et al.*, 2010). In contact bioautography, antimicrobial agents diffuse from a
260 developed TLC plate to an inoculated agar plate (Dewanje *et al.*, 2014). The complete lack
261 of bacterial growth in the area in contact with the peptides from the TLC plates indicates that
262 all the peptides present on the chromatogram from the extracts exhibit antibacterial activity
263 which can be said to be bactericidal in nature. The peptides also exhibited antimicrobial
264 potential against *C.albicans* but the slight but uniform growth observed suggests that the
265 peptides also exhibited a fungistatic, rather than fungicidal activity against *C.albicans*. After
266 removal of the TLC plates, the isolates were still incubated for a further 48hrs. At this time,
267 slight microbial growth was observed in the area of the of the Agar plate that had been in
268 contact with the peptides. As the growth of the microbes in this area was not as profuse as
269 other areas of the plate, this implies a sustained antimicrobial activity of the peptides.

270 One of the defence strategies used by molluscs, and indeed, a host of other
271 organisms is the production of antimicrobial peptides (AMPs) also known as host defence
272 peptides (HDPs). AMPs play key roles in innate immunity. Unlike antibiotics, which target
273 specific cellular activities, AMPs target the lipopolysaccharide layer of cell membrane, which
274 is unique to microorganisms. The high cholesterol level and negative charge of the bacterial
275 membrane ensure that eukaryotic cells are usually not targets of many AMPs (Bahar and
276 Ren, 2013; Bechinger and Gorr, 2016; Mahlapuu *et al.*, 2016). Hence, Selectivity is a very
277 important feature of the antimicrobial peptides and it can guarantee their function as
278 antibiotics in host defense systems. Another important feature of AMPs is their fast killing
279 ability. Some AMPs can kill in seconds after the initial contact with cell membrane (Bahar
280 and Ren, 2013). In addition to their role as endogenous antibiotics, some AMPs contribute to
281 inflammation as well as exhibit immunomodulatory activities (Mahlapuu *et al.*, 2016). Hence,
282 these AMPs act indirectly to kill microorganisms by modulating the host defense systems.
283 Some other AMPs kill bacteria by inhibiting some important pathways inside the cell such as
284 DNA replication and protein synthesis (Bahar and Ren, 2013). Their preferential attack on
285 the cell membrane or cell wall of bacterial and fungi ensure that AMPs should not cause
286 widespread resistance. In cases where specific protein targets are involved, the possibility
287 exists for genetic mutations and bacterial resistance. However, evidence suggests that this
288 is a rare event which can be overcome by subtle structural modifications made to the AMP
289 (Bechinger and Gorr, 2016).

290 Generally, AMPs are only effective against one class of microorganisms (e.g.,
291 bacteria or fungi). However, there are some notable exceptions like indolicidin, which can kill

292 bacteria, fungi, and viruses (HIV) (Bahar and Ren, 2013). The AMPs from the alcohol extract
293 of *P.aurita* and *T.fuscatus* have been shown in this study to be effective against both
294 bacteria and fungi species albeit by different mechanisms.
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298 **4. CONCLUSION**

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300 The results demonstrate that the acetone- methanol extract of *T.fuscatus* and *P.aurita*
301 possess multiple peptides with antibacterial and fungistatic properties which may be useful
302 as antimicrobial agents in new drugs for therapy of infectious diseases
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Authors declare that no competing interests exist.

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