


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



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


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


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SCREENING AND ENZYMATIC–BIOCHEMICAL CHARACTERIZATION OF PROBIOTIC CANDIDATE BACTERIA ISOLATED FROM THE DIGESTIVE TRACT OF LAKE TOBA NILE TILAPIA (*Oreochromis niloticus*)

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ABSTRACT

Intensive Nile tilapia (*Oreochromis niloticus*) culture in floating net cages in Lake Toba has increased organic waste accumulation and reduced feed digestibility, contributing to water quality degradation and mass fish mortality. Probiotics containing bacteria that produce digestive enzymes are considered a potential solution to improve feed digestibility. However, information on indigenous probiotic bacteria from the digestive tract of tilapia cultured in Lake Toba is limited, particularly regarding their enzymatic and biochemical characteristics. This study aimed to isolate, screen, and characterize enzyme-producing bacteria from the digestive tract of Lake Toba Nile tilapia as early probiotic candidates. Bacterial isolation was performed using serial dilution and culture on tryptic soy agar, followed by purification and morphological characterization. Enzymatic screening for amylolytic, proteolytic, and lipolytic activities was conducted using substrate-specific agar media, and selected isolates were identified using biochemical tests. Seven bacterial isolates were obtained, of which three exhibited extracellular enzymatic activity. Two isolates showed proteolytic activity with clear-zone diameters ranging from 17.4 ± 0.7 to 22.6 ± 0.7 mm, while one isolate demonstrated amylolytic activity with a clear-zone diameter of 1.57 ± 0.5 mm. No lipolytic activity was detected. Based on morphological and biochemical characteristics, the enzyme-producing isolates were identified as belonging to the genera *Micrococcus*, *Staphylococcus*, and *Streptococcus*. These results indicate that the digestive tract of Lake Toba Nile tilapia harbors indigenous enzyme-producing bacteria with potential as probiotic candidates. Further molecular identification, safety evaluation, and *in vivo* assessment are required prior to their application in aquaculture.

KEYWORDS: Lake Toba, mass mortality; organic waste; probiotic; tilapia

ABSTRAK: *Skrining dan Karakterisasi secara Enzimatis–Biokimia Bakteri Kandidat Probiotik yang Diisolasi dari Saluran Pencernaan Ikan Nila (*Oreochromis niloticus*) Danau Toba*

Budidaya ikan nila (*Oreochromis niloticus*) secara intensif dalam sistem keramba jaring apung di Danau Toba telah meningkatkan akumulasi limbah organik dan menurunkan daya cerna pakan, sehingga berkontribusi terhadap penurunan kualitas perairan dan terjadinya kematian ikan secara massal. Probiotik yang mengandung bakteri penghasil enzim pencernaan dipandang sebagai salah satu solusi potensial. Namun, informasi mengenai bakteri probiotik indigenous yang berasal dari saluran pencernaan ikan nila yang dibudidayakan di Danau Toba masih terbatas, khususnya terkait karakteristik enzimatis dan biokimianya. Penelitian ini bertujuan untuk mengisolasi, menyeleksi, dan mengkarakterisasi bakteri penghasil enzim dari saluran pencernaan ikan nila Danau Toba sebagai kandidat probiotik tahap awal. Isolasi

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52 bakteri dilakukan menggunakan metode pengenceran bertingkat dan penanaman pada media
53 tryptic soy agar, diikuti dengan pemurnian dan karakterisasi morfologi. Skrining aktivitas
54 enzimatik meliputi aktivitas amilolitik, proteolitik, dan lipolitik menggunakan media agar
55 dengan substrat spesifik, sedangkan isolat terpilih diidentifikasi melalui uji biokimia. Sebanyak
56 tujuh isolat bakteri berhasil diperoleh, dengan tiga isolat menunjukkan aktivitas enzim
57 ekstraseluler. Dua isolat menunjukkan aktivitas proteolitik dengan diameter zona bening
58 berkisar antara $17,4 \pm 0,7$ hingga $22,6 \pm 0,7$ mm, sedangkan satu isolat menunjukkan aktivitas
59 amilolitik dengan diameter zona bening sebesar $1,57 \pm 0,5$ mm. Aktivitas lipolitik tidak
60 terdeteksi. Berdasarkan karakteristik morfologi dan biokimia, isolat penghasil enzim tersebut
61 diidentifikasi berasal dari genus *Micrococcus*, *Staphylococcus*, dan *Streptococcus*. Hasil ini
62 menunjukkan bahwa saluran pencernaan ikan nila Danau Toba mengandung bakteri
63 indigenous penghasil enzim yang berpotensi sebagai kandidat probiotik. Identifikasi molekuler
64 lebih lanjut, evaluasi keamanan, dan pengujian *in vivo* diperlukan sebelum aplikasinya dalam
65 sistem akuakultur.

66

67 **KATA KUNCI:** Danau Toba; ikan nila; kematian masal; limbah organik; probiotik

68

69 INTRODUCTION

26 Freshwater fish farming using floating net cages, or *keramba jaring apung* (KJA), has

71 been well developed in Lake Toba and contributes to the livelihood of the local community
72 around the lake (Garno *et al.*, 2020). However, the rapid expansion of floating net cages in
73 Lake Toba, combined with its limited carrying capacity, has adversely affected the aquatic
74 environment. High levels of waste from uneaten feed and fecal matter have been consistently
75 observed, resulting in poorer water quality (Panjaitan & Manullang, 2022; Prasetya *et al.*,
76 2023).

77 It is well established that high concentrations of dissolved organic matter in fish-rearing
78 media increase oxygen consumption during the aerobic biodegradation of these compounds.
79 This results in a decline in dissolved oxygen concentration, after which anaerobic
80 decomposition occurs, producing substances such as H₂S and ammonia and increasing toxic
81 nitrogen and phosphorus (Ashari *et al.*, 2023). The accumulation of these toxic materials will
82 cause mass mortality of fish and other aquatic organisms cultured in KJA and ponds (Chong,
83 2022). In Lake Toba, Garno *et al.* (2020) reported that BOD and COD values are regularly
84 classified as high, exceeding Class I and Class II water quality thresholds. Their study

85 suggested that the elevated BOD values in Lake Toba's waters likely originate from KJA's
86 operations within the lake.

87 One of the primary causes of high concentrations of dissolved matter in water bodies is
88 uneaten fish feed (Nathanailides *et al.*, 2023; Wisnu *et al.*, 2021). Preventing the accumulation
89 of dissolved materials is challenging, given prevailing water quality conditions and fish-
90 feeding behaviors. Moreover, it is concerning that, even after feeding, nutritional value may be
91 lost due to the fish's low digestive capacity and limited digestive enzyme production
92 (Nathanailides *et al.*, 2023). To mitigate eutrophication, improving feed digestibility and the
93 production of digestive enzymes in fish by supplementing feed with bacteria that exhibit
94 amylolytic, proteolytic, and lipolytic activities is recommended (Munguti *et al.*, 2020;
95 Nathanailides *et al.*, 2023).

96 Tilapia is an omnivorous fish that produces protease, lipase, and carbohydrase enzymes
97 (Mawardi *et al.*, 2023), and probiotic bacteria could improve these enzymatic processes.
98 However, not all microbes living in the digestive tract of tilapia are capable of being probiotics;
99 Therefore, it is necessary to precisely identify the characteristics of probiotic bacteria found in
100 the fish digestive tract (Thepnarong *et al.*, 2024). This approach ensures that native probiotics
6 collected from the tilapia digestive tract function optimally, adapt to, and survive in the
102 digestive tract of the same species (Kuebutornye *et al.*, 2020).

22 Research on the isolation of probiotics from the digestive tract of fish, including gourami
104 (Dalahi *et al.*, 2014), catfish (Kurniasih *et al.*, 2014), and tilapia (Efendi & Yusra, 2014), has
105 been widely published. Their application in feed has demonstrated significant effects on fish
39 growth and feed efficiency. The findings of those studies underscore the importance of
19 establishing a framework for the isolation, selection, and identification of probiotic bacteria
108 from the digestive tracts of tilapia cultured in Lake Toba. Such a framework is needed due to
12 limited specific information on the morphological and biochemical characteristics of probiotic

110 bacteria in the digestive tract of tilapia cultured in floating net cages in the lake, particularly
111 their amylolytic, proteolytic, and lipolytic activities. Therefore, this study aimed to determine
112 the morphological and biochemical characteristics of bacteria from the digestive tract of tilapia
113 cultured in Lake Toba, including the identification of isolates exhibiting amylolytic,
114 proteolytic, and lipolytic activities.

115

116 MATERIALS AND METHODS

117 Sampling Location

118 Sampling was conducted in the waters of Lake Toba, North Sumatra, Indonesia, where
119 Nile tilapia (*Oreochromis niloticus*) are intensively cultured in floating net-cage systems.
120 Laboratory analyses, including bacterial isolation, enzymatic screening, and biochemical
121 identification, were performed at the Fisheries Laboratory, Faculty of Fisheries,
122 Dharmawangsa University, Medan, North Sumatra, Indonesia.

123

124 Sample Collection

125 Healthy Nile tilapia aged 2–3 months, weighing approximately 100–150 g, were
126 randomly collected from floating net cages. Fish showing physical deformities (e.g., spinal or
127 jaw deformities and fin erosion) or clinical signs of disease in accordance with standard fish
128 health assessment criteria (e.g., skin lesions, hemorrhages, fin rot, abnormal swimming
129 behavior, lethargy, or visible ectoparasites) were excluded. All samples were transported to the
130 laboratory in sterile containers and placed in an insulated cooler with ice packs to maintain a
131 stable temperature of approximately $4 \pm 1^\circ\text{C}$ until immediate processing in the laboratory
132 (Linscott & Wang, 2023).

133

134 Experimental Design

11 This study used an experimental–descriptive exploratory design to isolate, screen, and
136 identify probiotic candidate bacteria from the digestive tract of tilapia. The research was carried
137 out in three consecutive stages: (1) field survey and sampling, (2) bacterial isolation and
138 morphological characterization, and (3) screening of enzymatic activity followed by
139 biochemical identification of selected isolates (Makuachukwu & George-Okafor, 2025).

140

141 **Bacterial Isolation**

142 A total of ten healthy Nile tilapia were used in this study. The digestive tract (stomach
143 and intestine) of each fish was aseptically excised and homogenized in sterile physiological
10 saline (0.85% NaCl) for 1 minute. A serial dilution was prepared up to 10^{-6} , and 0.1 mL of each
145 dilution was spread onto tryptic soy agar (TSA; Merck, Germany) plates. The inoculated plates
146 were incubated at 36°C for 48 hours. Bacterial colonies exhibiting distinct morphological
147 characteristics were selected for further analysis (Firdus *et al.*, 2022).

148

149 **Purification and Morphological Characterization**

150 Selected colonies were repeatedly subcultured on TSA plates until pure isolates were
151 obtained. Morphological characterization was conducted by observing colony shape, color,
152 margin, and elevation. Cell morphology was examined by Gram staining under a binocular
153 microscope (Olympus, Japan) to determine the Gram reaction and cellular morphology
154 (Agustina *et al.*, 2022; Yulikasari *et al.*, 2024).

155

156 **Screening of Enzymatic Activity**

157 Pure bacterial isolates were screened for extracellular enzymatic activity, including
158 amylolytic, proteolytic, and lipolytic activities. Amylolytic activity was tested using TSA
159 (Merck, Germany) supplemented with 2% soluble starch and visualized by iodine staining

160 (Merck, Germany); proteolytic activity was assessed on TSA supplemented with 4% skim milk
161 powder (Oxoid, United Kingdom); and lipolytic activity was evaluated using TSA
162 supplemented with 2% olive oil (Sigma-Aldrich, United States).

15 The formation of clear zones around bacterial colonies indicated positive enzymatic
24 activity. The diameter of the hydrolytic zones was measured using a caliper and expressed in
165 millimeters (mm).

166

167 Biochemical Identification

168 Bacterial isolates exhibiting positive enzymatic activity were subjected to biochemical
169 characterization, including the catalase and oxidase tests, glucose and lactose fermentation
170 tests, motility tests, spore-formation tests, and Ziehl–Neelsen staining. The biochemical
14 profiles were compared with standard identification keys in *Bergey's Manual of Determinative
172 Bacteriology* and *Cowan and Steel's Manual for the Identification of Medical Bacteria to
173 determine bacterial genus* (Lee, 2023).

174

29 Data Analysis

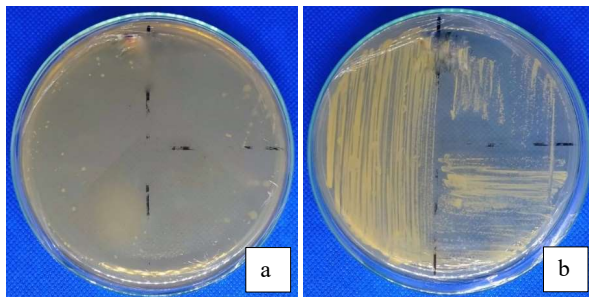
176 Morphological and biochemical characteristics of bacterial isolates were analyzed
177 descriptively. Enzymatic activity was quantified from the mean diameter of the hydrolytic clear
178 zones and expressed as mean \pm standard deviation. The identified probiotic candidate bacteria
179 were classified according to their morphological, biochemical, and enzymatic profiles
180 (Kumiasih *et al.*, 2014).

181

182 RESULTS AND DISCUSSION

183 Colony and Cell Morphology of Potential Probiotic Bacteria

184 Based on research conducted on bacteria found in the digestive tract (intestine) of tilapia,
 185 ten intestinal samples collected from tilapia yielded seven bacterial isolates that could grow
 186 and develop on TSA. Colony morphology was observed, including shape, edge, elevation, and
 187 color, which is illustrated in Table 1. Overall, the morphology of these colonies is nearly
 188 identical, differing only in shape, color, edge, and convex elevation. Each colony that showed
 189 predominant growth from the tilapia's digestive tract samples was transferred and re-grown on
 6 TSA media for 24 hours. The outcomes of the dilution and purification of bacteria isolated from
 191 the digestive tract of tilapia are displayed in Figure 1.



192
 5 Figure 1. Bacterial colonies from the digestive tract of Nile tilapia grown on tryptic soy agar:
 193 initial isolation (a) and purified isolates (b)
 196

197 Conventional methods identify bacteria by enriching microbes to obtain a pure culture.
 198 From this pure culture, microbial identification is achieved by comparing the characteristics of
 199 the bacterial isolate with those of previously identified bacteria. If no bacterium shows 100%
 200 similarity, the focus shifts to the bacteria that exhibit the highest similarity. As a result,
 201 conventional identification techniques will always identify a previously known bacterium and
 202 struggle to recognize a new species.

203
 2 Table 1. Morphological characteristics of bacterial colonies isolated from the digestive tract of
 204 Lake Toba Nile tilapia during probiotic screening

Isolate	Colony shape	Colony edge	Elevation	Color
US1.1	Small circular	Entire	Convex	Yellowish white
US1.2	Small circular	Entire	Convex	Milky white

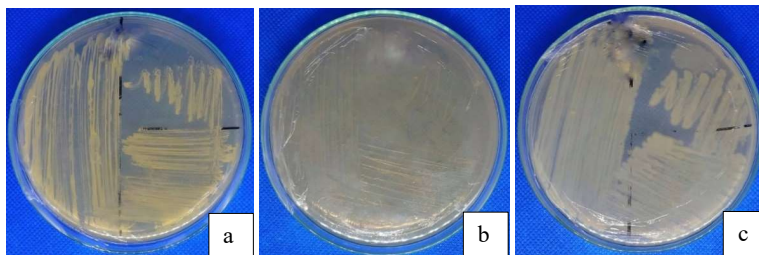
US2.1	Small	circular	Entire	Convex	Milky white
US2.2	Small	circular	Entire	Convex	Translucent white
US2.3	Moderate	circular	Entire	Convex	Milky white
US3.1	Moderate	circular	Entire	Convex	Milky white
US3.2	Small	circular	Entire	Convex	Milky white

206

207 **Table 1** shows that the seven probiotic bacterial isolates exhibit similarities, characterized
 208 by entire edges and convex elevations. Isolates US2.3 and US3.1 exhibit a moderate-circular
 209 colony shape, while the other five isolates display a small-circular colony shape. Similarly,
 210 isolates US1.1 and US2.2 have yellowish-white and translucent-white colonies, respectively,
 211 whereas the other isolates have milky-white colonies (Figure 2).

212 Microbes exist in numerous forms. To study their growth, morphology, and physiological
 213 properties, it is necessary to isolate each microbe from others to obtain a pure culture. A pure
 214 culture consists of cells from a single species or strain of microbes. Depending on the type of
 215 microorganism involved, bacterial isolation can be performed using serial dilution followed by
 216 the spread plate technique to obtain discrete colonies from mixed microbial samples (Gupta,
 217 2023).

218



219

220 Figure 2. Colony color variation of bacterial isolates from the digestive tract of Lake Toba Nile
 221 tilapia: yellowish white (a), translucent white (b), and milky white (c)
 222

223 **Hydrolysis of Starch (Amylum), Fat (Olive Oil), and Casein (Protein) in Bacterial Isolates**
 224 **and the Resulting Clear Zone**

225 The hydrolysis test of starch (amylum), fat (olive oil), and casein (protein) was conducted
 226 to assess the ability of bacteria to produce amylase, lipase, and protease enzymes, marking the
 227 initial stage in identifying bacterial isolates with probiotic potential for tilapia. This stage
 228 involved inoculating each bacterial colony onto media containing starch (amylum), fat (olive
 229 oil), and casein (protein). Seven isolates were selected as probiotic candidates based on their
 230 capacity to hydrolyze starch, fat, and casein. The results were indicated by the presence of clear
 231 zones around the isolates grown on enriched TSA (Table 2 and Figure 3).

232
 233 Table 2. Enzymatic hydrolysis activity of bacterial isolates from the digestive tract of Lake
 234 Toba Nile tilapia assessed using starch, casein, and lipid substrates during probiotic
 235 screening

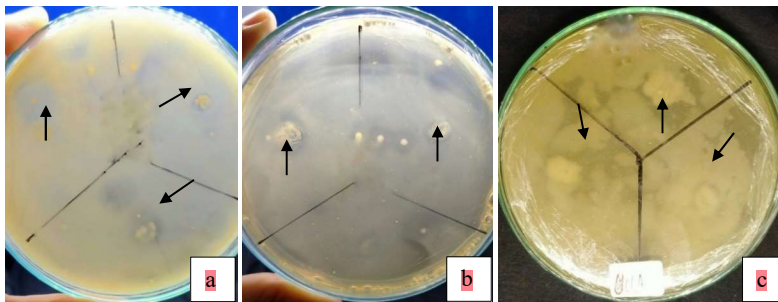
Isolate	Hydrolysis			Enzyme activity (mm; diameter of hydrolytic zone)
	Starch	Fat	Casein	
US1.1	-	-	+	22.6 ± 0.7
US1.2	-	-	+	17.4 ± 0.7
US2.1	-	-	-	-
US2.2	-	-	-	-
US2.3	-	-	-	-
US3.1	+	-	-	1.57 ± 0.5
US3.2	-	-	-	-

236 Note: no detectable hydrolytic activity (-); detectable hydrolytic activity (+).

237
 238 Based on the results of proteolytic, amylolytic, and lipolytic activity tests conducted on
 239 the seven samples, three isolates were identified as potential probiotic bacteria. Isolates US1.1
 240 and US1.2 exhibited hydrolytic activity on casein, resulting in clear zones around the isolates,
 241 and are therefore classified as proteolytic bacteria. Isolate US3.1 demonstrated activity in
 242 hydrolyzing starch, indicating that it is an amylolytic bacterium. However, no lipolytic bacteria
 243 were detected at this stage, as no clear zones formed around isolates inoculated onto lipid-
 244 substrate media. Typically, lipolytic bacterial isolates produce clear zones around their colonies
 245 (Ado *et al.*, 2025).

15 No lipolytic activity was detected in this study, as indicated by the absence of clear zones
247 around the bacterial colonies on olive oil-supplemented agar. This negative result may be
38 attributed to several factors. First, lipase-producing bacteria are often less prevalent in the
249 digestive tract of Nile tilapia, where carbohydrate- and protein-degrading enzymes are more
250 dominant due to the omnivorous feeding habit of the species (Bairagi *et al.*, 2002). Second, the
251 qualitative plate assay using olive oil as a substrate has limited sensitivity, as lipase diffusion
252 in solid media is relatively poor and clear zone formation may not be evident despite low-level
253 enzyme production (Sirisha *et al.*, 2010). Furthermore, lipase expression is highly dependent
254 on environmental conditions, including substrate concentration, incubation time, and pH,
255 which may not have been optimal in the present screening assay (Al-Haidari *et al.*, 2021).
256 Therefore, the absence of detectable lipolytic activity should be interpreted as a limitation of
257 the preliminary screening method rather than definitive evidence of the absence of lipase-
258 producing bacteria.

259



36
261
262 **Figure 3.** Clear-zone formation indicating proteolytic and amylolytic activities of selected
35 bacterial isolates from the digestive tract of Lake Toba Nile tilapia during probiotic
264 screening. Proteolytic activity of isolates US1.1 (a) and US1.2 (b), and amylolytic
265 activity of isolate US3.1 (c)

7
266 Measurements of the diameter of the hydrolytic clear zone showed that the proteolytic
268 bacterial isolate US1.1 exhibited the highest extracellular protease activity, with an average

269 clear zone diameter of 22.6 ± 0.7 mm. A clear zone of this size indicates significant proteolytic
270 potential, as many proteolytic probiotic candidates in recent studies have demonstrated protein
271 clear-zone diameters exceeding approximately 15–20 mm during qualitative testing for enzyme
272 secretion. Proteolytic LAB strains showing clear zones > 20 mm are indicative of substantial
273 protease production (Elkased, 2024). Protease production by probiotic bacteria is important
274 because these enzymes degrade proteins in the host gastrointestinal tract, thereby improving
275 nutrient digestion and feed utilization, which are desirable traits for effective probiotic strains
276 in aquaculture systems. In comparison, isolate US1.2 had an average diameter of 17.4 ± 0.7
277 mm, slightly smaller than that of isolate US1.1. On the other hand, the isolated US3.1 showed
278 amylolytic activity, with a clear zone. For isolates with amylolytic activity, the average
279 hydrolytic zone diameter is 1.57 ± 0.5 mm.

280 The amylolytic activity observed in isolate US3.1 produced a relatively small clear zone
281 diameter (1.57 ± 0.5 mm), indicating low extracellular amylase activity under the conditions
282 of the qualitative plate assay. This limited zone formation may be influenced by several factors,
283 including low enzyme secretion, diffusion constraints in starch-agar media, or suboptimal
284 incubation conditions for amylase expression. Similar observations have been reported in
285 preliminary screening studies, in which amylase-producing bacteria exhibited small hydrolytic
286 zones despite measurable enzymatic activity (Kusmiyati *et al.*, 2025). Therefore, the amylolytic
287 activity observed in this study should be interpreted as an initial indication rather than a
288 definitive measure of enzymatic efficiency.

21 Tests for starch, lipid, and casein hydrolysis were conducted to identify the presence of
290 amylase, lipase, and protease activities in the bacteria. Starch and casein hydrolysis tests are
37 crucial for selecting probiotic candidates. The activities of amylase, lipase, and protease
27 enzymes can enhance the function of endogenous enzymes in the digestive tract of fish,
2 suggesting that the presence of these bacteria may indirectly benefit their host. This aligns with

294 the findings of Kusmiyati *et al.* (2025), who indicate that the action of protease, lipase, and
17 amylose enzymes stimulates the production of endogenous enzymes by bacteria in the digestive
296 tract. Therefore, the presence of these bacteria may indirectly benefit their host.

3 The starch hydrolysis test results are characterized by the formation of a clear zone after
3 iodine application to bacterial isolates. This occurs because starch molecules are water-soluble
298 and produce a blue color when mixed with iodine solution, but they create a clear zone during
299 starch hydrolysis. This complex forms when amylose coils around the iodine molecule in a
300 helical conformation. When the amylose polymer is cut into shorter lengths, the complex
301 formed with iodine changes, causing the color to lighten and turn red or brown (Pesek &
302 Silaghi-Dumitrescu, 2024). Based on this finding, only isolate US3.1 exhibited the ability to
303 hydrolyse starch, indicating the presence of amylase activity, which is capable of degrading
304 starch into glucose, maltose, and dextrin.
305

23 A positive casein hydrolysis test is indicated by the formation of a clear zone around
307 bacteria grown on skim milk agar. Based on this study's findings, isolates US1.1 and US1.2
308 could hydrolyze casein, indicating that these isolates possess protease activity that breaks down
309 proteins into amino acids. According to Gurina & Mohiuddin (2020), protein hydrolysis yields
310 individual amino acids, which can be used for protein synthesis, other cellular processes, or
311 energy production.
312

313 Characterization and Identification of Potential Probiotic Bacteria

314 After testing the isolates for probiotic activity, morphological and biochemical tests were
315 conducted. Based on the results of previous tests on seven isolates obtained from the digestive
316 tracts of tilapia collected from Lake Toba, two potential isolates as proteolytic bacteria and one
317 as amylolytic bacteria were identified. Furthermore, the three bacterial types exhibiting
318 hydrolytic activity were examined at the genus level based on similarities to previously

319 identified characteristics. Bacterial identification is performed using conventional techniques
 320 by comparing the bacteria to previously identified species. In the absence of an exact
 321 phenotypic match, bacterial isolates are identified based on the most closely related known
 322 species. As a result, conventional identification approaches inherently favor the recognition of
 323 previously described bacteria and have limited capacity to reveal novel species (Selim *et al.*,
 2024). The results of the biochemical tests for the three bacterial isolates are presented in Table
 3.

Table 3. Morphological and biochemical characteristics of probiotic candidate bacteria isolated from the digestive tract of Lake Toba Nile tilapia

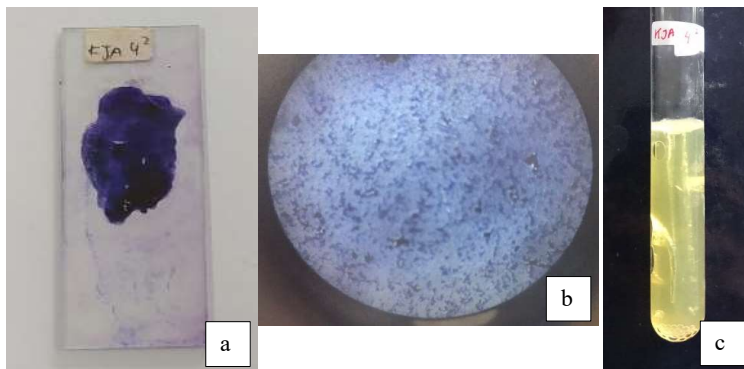
Morphology & biochemical test	Bacterial isolates		
	US1.1	US1.2	US3.1
Cell morphology			
Gram	+	+	+
Shape	coccus	coccus	coccus
Biochemistry			
Ziel Nielsen	-	-	-
Catalase	+	+	+
Glucose fermentative	+	+	-
Lactose fermentative	+	+	+
Motility	+	+	+
Oxidation	-	+	+
Bacteria genus	<i>Micrococcus</i>	<i>Staphylococcus</i>	<i>Streptococcus</i>

Note: (+) positive, (-) negative.

The characterization and identification of the three bacterial isolates revealed that isolate US1.1 exhibited Gram-positive cell morphology with a spherical (coccus) shape. Biochemical analysis showed that this isolate was motile, catalase-positive, oxidase-negative, and capable of fermenting glucose and lactose. Based on these characteristics, isolate US1.1 was identified as belonging to the genus *Micrococcus*. Isolate US1.2 exhibited characteristics largely similar to those of isolate US1.1, differing primarily in its oxidase reaction. Based on its biochemical profile, isolate US1.2 was classified as a member of the genus *Staphylococcus*. The

1 characteristics of isolates US1.1 and US1.2 are consistent with the findings of Kurniasih *et al.*
339 (2014), who reported that *Micrococcus* and *Staphylococcus* belong to the family
340 Staphylococcaceae. The third isolate, US3.1, was identified as a member of the genus
341 *Streptococcus* based on its biochemical characteristics. The results of Gram staining, cell
342 morphology, and motility tests, which formed part of the bacterial identification process, are
343 presented in Figure 4.

344



345
346 Figure 4. Gram-positive staining (a), coccoid cell morphology (b), and motility test results (c)
348 of selected probiotic candidate bacterial isolates from the digestive tract of Lake Toba
349 Nile tilapia

350 The results of morphological and biochemical analyses indicated that the bacterial
352 isolates considered as potential probiotic candidates from the digestive tract of Nile tilapia
354 belonged to the genera *Staphylococcus*, *Micrococcus*, and *Streptococcus*. Similar genera have
356 previously been reported from the digestive tract of tilapia and other cultured fish species
(Afrianto & Liviawaty, 2019; Kurniasih *et al.*, 2014), suggesting that these bacteria are
commonly associated with the fish gut microbiome. Comparable findings were also reported
by Ginting *et al.* (2018), who identified *Staphylococcus*, *Micrococcus*, and *Lactobacillus*
species as potential probiotic candidates in the digestive tract of milkfish (*Chanos chanos*).

358 Despite their frequent occurrence in the fish gut, it is important to note that certain species
359 within the genera *Staphylococcus* and *Streptococcus*, such as *S. aureus*, *S. iniae*, and *S.*
360 *agalactiae*, are well-documented fish pathogens with established virulence factors and disease
13 associations in aquaculture systems (Maulu *et al.*, 2021; Wang *et al.*, 2022). Therefore,
362 although the isolates identified in this study exhibited extracellular enzymatic activities
363 associated with probiotic potential, they cannot yet be classified as probiotics. Additional safety
364 evaluations, including hemolytic activity, virulence-associated traits, and antibiotic resistance
365 profiling, are required to ensure biosafety prior to any *in vivo* application.

366 Isolate US1.1, identified as belonging to the genus *Micrococcus*, exhibited Gram-positive
367 coccoid morphology with yellow-pigmented colonies and convex, circular edges. The cells
368 appeared singly, in pairs, or in tetrads and demonstrated catalase-positive, oxidase-negative
369 reactions, consistent with the characteristics commonly reported for *Micrococcus* species
370 (Selim *et al.*, 2024; Tizabi & Hill, 2023).

371 Isolate US1.2 was classified within the genus *Staphylococcus* based on its Gram-positive
372 coccoid morphology, creamy-white colonies, and biochemical properties, including catalase
373 and oxidase positivity. These traits are consistent with previously described *Staphylococcus*
374 species commonly isolated from aquatic environments and fish-associated microbiota (Selim
375 *et al.*, 2024).

376 The third isolate, US3.1, was identified as a member of the genus *Streptococcus*. This
377 isolate exhibited Gram-positive coccoid cells arranged in pairs or short chains, a morphology
378 typical of *Streptococcus* species. The observed physiological characteristics are consistent with
379 those reported for mesophilic streptococci commonly associated with aquatic organisms.

380

381 **CONCLUSIONS**

382 The probiotic bacterial candidates identified in this study consist of indigenous, enzyme-
383 producing gut bacteria from the genera *Micrococcus*, *Staphylococcus*, and *Streptococcus*,
5 isolated from the digestive tract of Lake Toba Nile tilapia. These bacteria were selected as
32 early-stage probiotic candidates based on their ability to produce extracellular digestive
386 enzymes, particularly proteases and amylases, which are associated with improved feed
387 digestibility and nutrient utilization in aquaculture systems. However, this study is limited by
388 the use of conventional morphological, biochemical, and qualitative enzymatic assays, which
389 do not allow strain-level identification or precise evaluation of functional efficiency. In
390 addition, genera such as *Staphylococcus* and *Streptococcus* include species known to be
391 opportunistic or pathogenic in fish, highlighting the need for careful biosafety evaluation.
392 Future studies should therefore apply molecular identification methods (e.g., 16S rRNA gene
393 sequencing), quantitative enzymatic analyses, and comprehensive safety assessments, followed
394 by *in vivo* feeding trials, to confirm the probiotic suitability and practical applicability of these
395 bacterial candidates.

396

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408

409 AUTHOR CONTRIBUTION

ES: investigation, data curation, formal analysis, validation, writing – original draft, and
writing – review and editing. DTA: investigation and writing – review and editing.

412

413 DECLARATION OF COMPETING INTEREST AND USE GENERATIVE AI

414 The authors declare no competing interests.

415

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