Relationships between pond water and tilapia skin microbiomes in 1 2 aquaculture ponds in Malawi

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23 Highlights

- Fish skin and pond water communities differ structurally, but share common taxa
- Pond locations have a stronger influence on water versus fish skin microbiome community structure
 - Selected skin-associated taxa could be used to monitor dysbiotic events in aquaculture •
- Taxa with opportunistic pathogen potential were identified at low abundance •

29 Abstract

30 Intensification of fish farming practices is being driven by the demand for increased food 31 production to support a rapidly growing global human population, particularly in lower-middle 32 income countries. Intensification of production, however, increases the risk of disease 33 outbreaks and thus the likelihood for crop losses. The microbial communities that colonise the 34 skin mucosal surface of fish are poorly understood, but are important in maintaining fish health 35 and resistance against disease. This skin microbial community is susceptible to disruption 36 through stressors associated with transport, handling and the environment of intensive 37 practices, and this risks the propagation of disease-causing pathogens. In this study, we 38 characterised the microbial assemblages found on tilapia skin — the most widely farmed finfish 39 globally — and in the surrounding water of seven earthen aquaculture ponds from two pond 40 systems in distinct geographic regions in Malawi. Metabarcoding approaches were used to 41 sequence the prokaryotic and microeukaryotic communities. We found 92% of prokaryotic 42 amplicon sequence variants were common to both skin and water samples. Differentially 43 enriched and core taxa, however, differed between the skin and water samples. In tilapia skin, 44 Cetobacterium, Paucibacter, Pseudomonas and Comamonadaceae were enriched, whereas, the 45 cyanobacteria Cyanobium, Microcystis and/or Synechocystis, and the diatom Cyclotella, were most prevalent in pond water. Ponds that clustered together according to their water prokaryotic 46 47 communities also had similar microeukaryotic communities indicating strong environmental 48 influences on prokaryotic and microeukaryotic community structures. While strong site-49 specific clustering was observed in pond water, the grouping of tilapia skin prokaryotes by

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50 pond site was less distinct, suggesting fish microbiota have a greater buffering capacity against

51 environmental influences. The characterised diversity, structure and variance of microbial 52 communities associated with tilapia culture in Malawi provide the baseline for studies on how

52 communities associated with thapia culture in Malawi provide the baseline for studies on now 53 future intensification practices may lead to microbial dysbiosis and disease onset.

- 54
- 55 Keywords

56 Aquaculture, skin microbiome, tilapia, pond, bacterial community, eukaryotic community

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58 **1. Introduction**

59 Capture fisheries will not be able to satisfy the demand for seafood products from an ever-60 increasing human population with rising living standards (Henchion et al., 2017) combined 61 with plateauing, and in some cases declining, wild fish stocks due to overfishing and ecosystem 62 degradation (Link and Watson, 2019). Seeking to meet this demand for aquatic products, many 63 aquaculture farming practices are undergoing intensification. Shifting from extensive to 64 intensive and semi-intensive practices in aquaculture, however, is often associated with 65 increased incidence of infectious disease (Hinchliffe et al., 2020; Pulkkinen et al., 2010). Intensification can cause chronic stress that adversely impacts fish physiology resulting in 66 67 reduced growth and impaired disease resilience. Increasing pond stocking rates and levels often 68 occurs with insufficient amounts of clean water, leading to the deterioration of water quality, 69 including dissolved oxygen, pH and ammonia (Abdel-Tawwab et al., 2014; Sundh et al., 2019), 70 which in turn impacts negatively on fish growth and health, and renders the fish more 71 susceptible to diseases. Regular restocking of ponds with fish of uncertain health status to 72 compensate for mortalities, in turn, increases the likelihood of repeated introductions of sub-73 clinical infections (Bondad-Reantaso et al., 2005; Murray and Peeler, 2005).

74 Disease remains a huge challenge for aquaculture, particularly in Asia where 89% of 75 global aquaculture production occurs (FAO, 2020c). Successful management of disease risk 76 and intensification of aquatic species production requires a better understanding of the 77 relationships between the microbial systems (microbiomes) of both the cultured aquaculture 78 species and of the environments in which they are grown (Bass et al., 2019). The study of 79 microbiomes in aquaculture is gaining momentum and recent studies have investigated how 80 pond and fish treatments (e.g. antibiotics, dietary supplements, probiotic treatments and pond 81 fertilisers) affect fish microbiomes (Limbu et al., 2018; Minich et al., 2018; Suphoronski et al., 82 2019; Tan et al., 2019). Much of this research has focused on the gut microbiome due to its 83 intricate role in gut health, which when optimised can maximise feed conversion, growth, and 84 overall aquaculture productivity (Perry et al., 2020). When considering disease resistance 85 and/or susceptibility in fish aquaculture, however, arguably the microbial communities harboured on/in the skin and gills are likely to be equally if not more important. 86

87 These outer facing mucosal surfaces are in continuous contact with the aquatic 88 environment and provide a primary barrier against invading pathogens (Legrand et al., 2018; 89 Rosado et al., 2019b). The microbes colonising this skin niche include those specifically 90 adapted to the host mucosal surface, as evidenced by host-species specificity of microbiome 91 composition (Doane et al., 2020), but also microbes derived from the surrounding water 92 community (Krotman et al., 2020). Relatively little is known about the environmental and host 93 contributions to these microbial assemblages, particularly in aquaculture ponds. It is known, 94 however, that skin colonisers have a direct connection with the host immune system helping to 95 shape its function and responses (Kanther et al., 2014). Equally, the immune system provides 96 feedback in sculpting the microbial community structure (Kelly and Salinas, 2017; Tarnecki et 97 al., 2019). If these finely balanced communities are disrupted, to a state known as dysbiosis, 98 resulting health complications and disease may occur. The fish skin microbiome has been 99 reported to change following stressful events, such as high stocking densities and hypoxia

100 (Boutin et al., 2013), in fish showing clinical signs of gastrointestinal enteritis (Legrand et al., 2018) and also following viral infection (by salmonid alphavirus; see Reid et al., 2017), 101 102 bacterial infection (by Photobacterium damselae; see Rosado, Xavier, et al., 2019b) and 103 macroparasitism (by the sea lice *Lepeophtheirus salmonis*; see Llewellyn et al., 2017). In all 104 of these cases, there was a decrease in abundance of reputedly beneficial taxa, concurrent with 105 an increase in opportunistic pathogens. The resulting theory is that dysbiosis within the skin 106 microbiome causes fish to become more susceptible to secondary bacterial infections. This has 107 been shown for exposure to the antimicrobials rifampicin in Gambusia affinis Baird & Girard 108 (Carlson et al., 2015) and potassium permanganate (Mohammed and Arias, 2015) in *Ictalurus* 109 *punctatus* Rafinesque, where increased mortality occurred for dysbiotic fish compared with 110 controls when challenged with the disease-causing Edwardsiella ictaluri and Flavobacterium 111 columnare, respectively.

112 A limitation in the majority of microbiome studies, regardless of host species, is a focus 113 on the bacterial community only with little or no attention given to the remaining microbial 114 community members. This includes microeukaryotes, a taxonomic group that encompasses 115 protists, microfungi, microalgae, and microbial metazoans (Bass and del Campo, 2020; del 116 Campo et al., 2019), as well as viruses that infect an expansive host range including 117 microeukaryotes, bacteria and the animal host (Gadoin et al., 2021). Microeukaryotic 118 communities are well described in some settings, such as the contribution of microalgae to 119 primary production in the ocean (Benoiston et al., 2017). The relationships between 120 microeukaryotes and animal hosts have predominantly focussed on parasitism and 121 pathogenesis, yet microeukaryotes play an intricate role in the broader microbial community 122 of host-associated niches. One of the best described examples is *Blastocystis*, a protist 123 commonly found to colonise the gut of humans and other animal hosts. Its presence is thought 124 to correlate with protection against several gastrointestinal inflammatory diseases by 125 interacting with the bacterial community to promote a healthy microbiome (Laforest-Lapointe 126 and Arrieta, 2018), specifically via an associated increase in bacterial diversity and strong co-127 occurrence patterns with reputed beneficial bacteria (Audebert et al., 2016; Beghini et al., 128 2017). The full role *Blastocystis* plays in human health remains unresolved and controversial. 129 The extent of interactions occurring between bacteria and microeukaryotes and/or viruses in 130 the fish skin microbiome is largely unknown and unreported.

131 Tilapia are the most widely farmed finfish in global aquaculture, produced in over 170 132 countries. Numerous species of tilapia are farmed, dominated by Nile tilapia (Oreochromis 133 niloticus L.), and predominantly in lower-middle income countries (LMICs) across the 134 Southeast Asian, African and South American continents (FAO, 2020b). Given their fast 135 growth, adaptability to a variety of environmental culture conditions, and resilience against 136 both disease and poor water quality, tilapia are now a production staple for many LMICs, and 137 colloquially is often referred to as the aquatic chicken (FAO, 2020a). While some aquaculture 138 species grown in LMICs, such as shrimp, are high-value products for export, the bulk of tilapia 139 production is for domestic markets. As a consequence, fewer regulations exist for tilapia 140 production (El-Sayed, 2019) and there has been far less scientific research for optimising 141 sustainable production compared to some other high-value teleost species, such as Atlantic 142 salmon.

Aquaculture in Malawi is in its relative infancy compared with other countries in Africa and Asia. Nevertheless, production has seen on average a 24% yearly growth between 2006 and 2016 (CASA, 2020). The levels of intensification or disease incidence seen in Malawi are low compared with Asia, but as demand increases, disease levels will inevitably increase also. Tilapia species cultured in Malawi include *Coptodon rendalli* Boulenger and *Oreochromis shiranus* Boulenger, with the notable absence of Nile tilapia, which is considered an invasive species. To fully elucidate the influence of microbiomes on fish health during disease 150 processes, we need to better understand the relationships between the microbial diversity, 151 community variance and structure in the mucosal surfaces of fish and those in the aquatic 152 environment, including microeukaryotes (often excluded from microbiome studies), for 153 disease-free populations. In this study therefore, we applied high throughput DNA sequencing 154 for metabarcoding of the 16S and 18S ribosomal RNA (rRNA) small subunit (SSU) marker 155 genes (which are conserved within prokaryotes and eukaryotes, respectively), to characterise 156 the microbial communities of pond water and tilapia skin (C. rendalli and O. shiranus) from 157 earthen aquaculture ponds in Malawi. With these data, we investigated the relationships 158 between the pond water and skin microbiome. We identified differentially enriched and core 159 taxa within the tilapia skin microbiome that are likely to play an important biological role for 160 the host and may provide notable taxa for future studies to interpret disease events.

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162 **2. Materials and methods**

163 **2.1. Sample collection**

164 Seven tilapia aquaculture earthen ponds were sampled in October 2017 from two pond systems 165 in Malawi. Two ponds from a commercial farm were located in Maldeco, and a five ponds 166 from a community pond syndicate were located 200 km further south in Blantyre 167 (Supplementary Fig. 1). Two sample types were collected: pond water and tilapia skin swabs 168 (Table 1). Pond surface water was collected from five locations within each pond by passing 169 200 mL of water through a polycarbonate filter (0.4 µm pore, 47 mm diameter, Whatman). The 170 volumes of water filtered were affected by the amount of organic/particulate matter in the 171 samples such that volumes were sampled until filters became saturated and prevented further 172 filtration. Mucosal skin samples of tilapia flanks (C. rendalli and O. shiranus) were collected 173 by swabbing three times along the entire length of the lateral line (Delamare-Deboutteville et 174 al., 2021) with sterile polyester swabs (Texwipe). Filters were preserved in 1.8 ml of 100% 175 molecular grade ethanol (FisherScientific), while swabs were preserved in 1.8 ml of RNAlater 176 (Qiagen), and stored at ambient temperature until transferred to the UK for prolonged storage 177 at -20 °C, until used for DNA extraction and sequencing.

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179 Table 1. Details of pond sites and samples for the pond water and fish skin swabs obtained180 from Malawian tilapia aquaculture ponds.

Pond system location	Pond site	No. pond water samples	No. fish skin swabs *	Cultured species	Mean fish length (mm) ± SD	No. fish measured for length
Blantyre, Malawi	1	5	2	Coptodon rendalli	115 ± 11.8	5
Blantyre, Malawi	2	5	1	Coptodon rendalli	125 ± 19.1	5
Blantyre, Malawi	3	5	7 (5)	Coptodon rendalli, Clarias gariepinus	149 ± 9.3 N/A	8
Blantyre, Malawi	4	5	8	Coptodon rendalli, Oreochromis shiranus	137 ± 4.2 122 ± 6.6	5 3
Blantyre, Malawi	5	5	5	Coptodon rendalli	162 ± 18	5

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Total	7	35	33 (28)				
Maldeco, Malawi	7	5	4 (3)	Oreochromis shiranus	212 ± 12.7	5	
Maldeco, Malawi	6	5	6 (4)	Oreochromis shiranus	222 ± 30.5	5	

*Numbers in parentheses refer to the number of 18S rRNA samples successfully sequenced,
where this differs from 16S rRNA samples.

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184 **2.2. DNA extraction**

Ethanol was removed from pond water filters by freeze-drying (ScanVac CoolSafe Pro, 4 L 185 186 condenser at -110 °C) and filters were then stored at -80 °C. RNA*later* was removed from fish 187 swab samples by vortexing the swabs for 30 seconds in 23 mL of 1x sterile phosphate buffered 188 saline (Sigma) to allow detachment of microbes. The swab and solution were transferred to a 189 syringe for filtration with a 0.22 µm Sterivex (Millipore) filter unit. Following ethanol and 190 filters DNA RNAlater removal from and swabs, was extracted with а 191 CTAB/EDTA/chloroform method adapted from Bramwell et al. (1995) and Lever et al. (2015), 192 and is available in full at (https://dx.doi.org/10.17504/protocols.io.bw8gphtw).

193 Briefly, for DNA extraction, filters were first suspended in 570 µl lysis buffer (30 mM 194 Tris, 30 mM EDTA, pH 8, FisherScientific), freeze-thaw lysed in liquid nitrogen and 195 homogenised by bead-beating with Lysing Matrix A Bulk Beads (Garnet) on the Qiagen 196 TissueLyser II for 40 seconds at 30 Hz. The sample suspension was digested with 1 µL Ready-197 Lyse lysozyme (1000 U/ µL, Epicentre), and 3 µL proteinase K (20 mg/mL, Sigma) in 30 µL 198 SDS (10% w/v, FisherScientific) for 1 hour at 55 °C. Samples were then incubated for 10 199 minutes at 65 °C in 120 µL NaCl (5 mM, Sigma) and CTAB solution 200 (hexadecyltrimethylammonium bromide, 96 μL, 10% w/v, Sigma). An equal ratio of sample 201 and 24:1 chloroform: isoamyl alcohol (Acros Organics) were used for extractions, with 202 centrifugation at 14,000 x g, 4 °C for 5 minutes. The aqueous layer was retained for a second 203 extraction, after which 1 µL of linear polyacrylamide solution (GenElute LPA, Sigma) was 204 added to aid precipitation with 0.7 volumes isopropanol (Acros Organics). Following overnight 205 incubation at 4 °C, samples were centrifuged at 21,000 x g, 4 °C for 30 minutes and the 206 resulting pellet was washed with 70% ethanol (FisherScientific). After 10 minutes of 207 centrifugation at 21,000 x g and pipetting off the ethanol, DNA pellets were resuspended in TE 208 buffer (10 mM Tris-HCl, 1 mM EDTA, pH 8, Sigma) and stored at -20 °C until used for 209 sequencing.

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211 2.3. Metabarcoding

212 Metabarcoding of prokaryotic and microeukaryotic SSU rRNA marker genes was performed 213 by PCR amplification with the Earth Microbiome Project recommended primers. The 16S hypervariable 214 V4region was targeted by 515F rRNA (Parada) 5'-215 GTGYCAGCMGCCGCGGTAA-3' (Parada et al., 2016); 806R (Apprill) GGACTACNVGGGTWTCTAAT (Apprill et al., 2015), and the 18S rRNA V9 hypervariable 216 217 region was targeted by 1391f 5'-GTACACCGCCGTC-3' (Lane, 1991) and EukBr 5'-218 TGATCCTTCTGCAGGTTCACCTAC-3' (Medlin et al., 1988). Amplification conditions for 219 16S V4 were 98 °C for 30 seconds; 30 cycles of 98 °C for 10 seconds, 55 °C for 30 seconds, 220 72 °C for 30 seconds; and a final extension of 72 °C for 2 minutes. 18S V9 conditions were the 221 same, with the exception of an annealing temperature of 60 °C. Samples were amplified and 222 multiplexed in a 1-step PCR with a dual-indexing scheme (Kozich et al., 2013). Individual 223 samples were run as 50 µL reactions with 2 ng starting DNA, 25 µL NEBNext High-Fidelity 224 PCR Master Mix (New England Biolabs), 0.5 µM forward and reverse primers, prior to pooling

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and sequencing by the University of Exeter Sequencing Service on the Illumina MiSeq, using
 v2 chemistry (250 bp paired-end for 16S and 150 bp paired-end for 18S). The sequencing runs
 included four positive controls (ZymoBIOMICS® Microbial Community DNA standard, lot
 number ZRC190811) and six negative controls comprising nuclease free water carried through
 the entire DNA extraction and PCR amplification.

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231 2.4. Bioinformatics processing

232 All bioinformatics and statistical analyses were performed in R v3.6.3. Following sample 233 demultiplexing, reads were quality controlled and processed by the DADA2 pipeline v1.14 234 (Callahan et al., 2016). Briefly, quality profiles of paired reads were inspected and forward and 235 reverse reads were truncated at 200 bp and 160 bp, respectively, for prokaryotes, and 100 bp 236 for both reads of microeukaryotes. Amplicon sequence variants (ASVs) were then inferred with 237 DADA2's pooling method to enhance the detection of rare ASVs. Paired reads were merged if 238 they achieved a minimum overlap of 100 bp for prokaryotes and 25 bp for microeukaryotes. 239 To remove off-target sequencing artefacts, final ASVs were only retained for the lengths 250 240 -256 bp for prokaryotes and 90 - 150 bp for microeukaryotes. Chimeras were removed and 241 taxonomy assigned to each ASV against the SILVA SSU v138 taxonomic database (Quast et 242 al., 2012). For the microeukaryotic dataset, only ASVs classified by SILVA as eukaryotic were 243 retained and final taxonomic classifications of these ASVs were made by the PR2 v4.12 244 taxonomic database (Guillou et al., 2012). Accuracy of the taxonomic assignment was assessed 245 in positive controls, with all members of the ZYMO mock community present, as expected 246 (Supplementary Fig. 2).

A phylogenetic tree of ASVs was constructed with Phangorn v2.5.5 (Schliep, 2010) by first generating a neighbour-joining tree, followed by fitting a generalised time reversible substitution model to generate a maximum likelihood tree. The statistical tool Decontam v1.6 (Davis et al., 2018) was used to identify contaminating ASVs by looking at the prevalence in negative controls, with standard parameters baring a 0.5 prevalence threshold. Thus, all sequences found at greater prevalence in negative controls than positive samples were classed as contaminants and were removed from the ASV table.

254 ASVs and sample data were parsed to Phyloseq v1.30 (McMurdie and Holmes, 2013) 255 for all subsequent quality control and data analyses. To remove sequencing noise, only ASVs 256 that reached a 2% prevalence threshold across samples were retained. Furthermore, any ASVs 257 taxonomically assigned as chloroplasts, mitochondria, eukaryotic or unclassified at kingdom 258 level were removed from the prokaryotic dataset. Additionally, for the microeukaryotic dataset, 259 14 sequences classified as Craniata were removed, as these most likely represented fish 260 sequences. As a result, the characterisation of fish skin microeukaryotes was limited due to the 261 high levels of contaminating host 18S rRNA sequences (98.6%) in swab samples.

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263 **2.5. Statistical and data analysis**

Alpha diversity metrics were calculated with Phyloseq on counts rarefied to the minimum sequencing depth. The difference between pond sites was statistically tested by Welch's ANOVA and *post-hoc* pair-wise Games-Howell test, following confirmation of normality. Further testing between sample types utilised ImerTest v3.1-3 (Kuznetsova et al., 2017) to perform a linear mixed-effects model that accounted for pond site as a random effect. A Pearson's correlation coefficient was used to test for correlation of Chao1 richness and Shannon diversity between sample types.

Beta diversity analysis was performed with compositional data analysis principles.
These comprise log-based transformations, which cannot be performed on zero values.
Therefore, ASV counts were subjected to a count zero multiplicative replacement method in
zCompositions v1.3.4 (Palarea-Albaladejo and Martín-Fernández, 2015). A centred log-ratio

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(CLR) transformation was then applied to ASV counts with the CoDaSeq package v0.99.6
(https://github.com/ggloor/CoDaSeq). Euclidean distance was calculated on log-ratios and
ordinated by PCoA biplot with FactoExtra v1.0.7 (https://github.com/kassambara/factoextra).
Statistical differences between pond site and sample type groups were conducted on the
Euclidean distance matrix by permutational multivariate analysis of variance (PERMANOVA)
and permutation tests for homogeneity of multivariate dispersions, implemented in Vegan
v2.5-6 (Dixon, 2003).

282 Community composition was presented as heat trees of taxon relative abundance with 283 Metacoder v0.3.4 (Foster et al., 2017), utilising a Davidson-Harel layout algorithm. Differential 284 abundance between sample types was assessed by CornCob v0.1 (Martin et al., 2020), utilising 285 the Wald Chi-Squared test and accounting for pond site as a random effect. Core microbiome 286 analysis was performed on ASVs amalgamated to genus level and rarefied to the minimum 287 sequencing depth. Classification of the fish skin core genera was performed with the 288 Microbiome package v2.1 (Lahti and Shetty, 2017) based on a prevalence threshold of 80% 289 and a detection threshold of 0.01% in all swab samples. Heatmaps of core genera and 290 discriminant taxa were depicted as heatmaps of CLR abundance of non-rarefied counts by 291 pheatmap v1.0.12 (https://github.com/raivokolde/pheatmap).

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The significance level and false discovery rate of 0.05 was set for all statistical analyses.

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294 **2.6. Data availability**

Raw sequencing reads were deposited in the European Nucleotide Archive under the accession
 PRJEB46984. Data processing, analysis scripts and final ASV tables are accessible at
 <u>https://github.com/jamiemcm/Malawi_Tilapia_Microbiomes.</u>

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3. Results

Following quality control and filtering, the final prokaryotic dataset contained 969,562 reads and 5782 ASVs from all skin swab and pond water filter samples, respectively, collected in this study (67 samples). The eukaryotic dataset comprised 94,611 reads, 1659 ASVs from the 62 samples collected. Full read counts per library, including break down between skin swabs and pond water filters, are available in Supplementary Table 1.

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306 **3.1. Phytoplankton communities**

307 Compositional approaches (CLR) to beta diversity were applied to explore variation in 308 microbial community composition and abundance of pond water between sites are shown in 309 Figures 1A and 1B. Clear clustering of water samples by pond site was evident, with the 310 position of mean group centroids corresponding to site shown to be significantly different from 311 each other according to PERMANOVA for both prokaryotic (F(6,28) = 34.29, $R^2 = 0.88$, p < 0.88312 0.001) and microeukaryotic (F(6,28) = 15.12, $R^2 = 0.76 p < 0.001$) communities. Dispersion of 313 the pond water samples collected within each site was relatively small, particularly with respect 314 to prokaryotes (Fig. 1B). However, largely due to pond site 2, dispersion in prokaryotes 315 differed significantly according to permutation tests for homogeneity of multivariate 316 dispersions (Prokaryotes F(6,28) = 3.95, p = 0.003; Eukaryotes F(6,28) = 0.87, p = 0.53).

While pond location had a strong influence on the separation of pond water samples, the clustering observed in prokaryotes of the tilapia skin was less distinctive (Fig. 1C). There was a significant difference between the mean centroid position of each pond site by PERMANOVA F(6,25) = 4.19, $R^2 = 0.50$, p < 0.001) and significant dispersion between fish within the same pond (F(6,25) = 5.32, p = 0.009).

Specific taxa were associated with driving community separation between pond sites.
 Figure 1 shows the top 15 contributing taxa plotted as arrows on each biplot, with their CLR

324 abundance depicted in the accompanying heatmaps. In pond water, microeukaryotes included 325 several diatoms (ASV6: Cyclotella and ASV16: Aulacoseira), the presence of which separated 326 pond site clusters 2,3 and 6,7 from 1,4,5. ASV47: Eukaryota was the major taxon - found at 327 high abundance - discriminating pond site cluster 1,4,5 from the remaining ponds, and a 328 BLASTn search of this ASV revealed 90% similarity to the microalgae Cryptomonas. In the 329 pond water prokaryotic community, photosynthetic Cyanobacteria were particularly prevalent 330 in pond site clusters 2,3 and 6,7, with apparently differing *Cyanobium* ASVs (ASV3, ASV4) 331 in each cluster, and a shared Synechocystis (ASV1). Pond site cluster 1,4,5 was distinguished 332 by typical freshwater planktonic Proteobacteria (ASV9: Polynucleobacter, ASV111: 333 Limnohabitans and ASV22: Comamonadaceae), among others. For fish skin prokaryotes, three 334 out of the top 15 discriminant taxa (ASV43: Aquabacterium, and ASV16, ASV74: 335 Comamonadaceae) explained the separation of pond cluster 1,4,5 only. Many of these 336 identified taxa shared taxonomic affiliation to the aforementioned prokaryotes of pond water, 337 but were represented by separate ASVs than those previously identified, such as ASV17, 338 ASV24: Cyanobium, ASV16, ASV74: Comamonadaceae and ASV6, ASV73, ASV135: 339 Actinobacteria hgcl clade (Warnecke et al., 2004).

340 Alpha diversity metrics gave an insight into species diversity of the pond water samples 341 from different pond sites as determined through assessing community richness (Chao1) and 342 evenness (Shannon diversity and Inverse Simpson diversity) (Fig. S3A,B). Applying Welch's 343 ANOVA showed a significant difference between both prokaryotic and microeukaryotic 344 communities of each pond site for all diversity metrics (Tab. S2). No correlation was found 345 between prokaryotic and microeukaryotic communities for the mean richness/diversity metric 346 of each pond site (Fig. S3D) (Pearson's correlation: Chao1 richness R = 0.56, p = 0.19; Shannon 347 diversity R = 0.35, p = 0.44; InvSimpson R = -0.024, p = 0.96).

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349 **3.2. Microbial niche separation**

350 We used measures of alpha and beta diversity to explore the influence of the environment (pond 351 water) in shaping tilapia skin prokaryotic microbiota. When controlling for pond site as a random effect in linear mixed-effects modelling, ASV richness of the fish skin was found to 352 353 be significantly lower than pond water (by 503 ± 59.49 ASVs, $R^2 c = 0.61$, p < 0.001) (Fig. 2A). 354 Shannon diversity of fish skin and pond water varied according to pond site, however, there 355 was no overall clear separation between the sample types when the aforementioned statistical 356 model was applied (pond water 4.96 ± 0.12 , fish skin 4.72 ± 0.15 , $R^2 c = 0.11$, p = 0.115) (Fig. 357 2B). Additionally, neither richness nor diversity were correlated between the fish skin and pond 358 water when comparing between pond sites, according to Pearson correlation tests (Chao1 359 richness R = 0.18, p = 0.71; Shannon diversity R = 0.11, p = 0.81) (Fig. 2C,D). Pair-wise 360 comparisons were made of the beta diversity (Aitchison distance) between samples within each 361 pond site (Fig. 2E) and this showed pond water samples clustered closely together, but greater 362 dispersion was apparent between fish skin samples. The largest Aitchison distance values were 363 seen in the comparisons between pond water and fish skin samples, indicating different 364 prokaryotic community structures between these niches. Although these structures were made 365 up of shared taxa, albeit at different abundances, with 4020 of a total 5782 ASVs detected in 366 both pond water and fish skin (Fig. 2F).

Depicting taxonomic composition of prokaryotic and microeukaryotic communities from skin swab and pond water samples as phylogenetic heat trees (see Fig. S4) illustrates much of the diversity for individual samples is accounted for by rare taxa found at low abundance. The prokaryotic community composition at coarse taxonomic levels was overall very similar between the pond water and skin environments, although divergence emerges at finer taxonomic resolution. For the microeukaryotic community, a far greater overall taxonomic diversity was observed in pond water than on skin, with numerous rare taxa.

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However, skin diversity was artificially under-sampled due to the over-amplification of tilapia
 host 18S RNA gene copies.

376 Taxonomic relative abundance (depicted as dot plots of prokaryotes at class level and 377 microeukaryotes at division level) highlights the differences between pond water and fish skin 378 niches (Fig. 3). According to differential abundance statistical testing, the bacterial classes 379 Gammaproteobacteria and Clostridia were enriched (FDR <0.05) in the fish skin. Pond water 380 by contrast had enriched abundances of Cyanobacteria, Actinobacteria, Bacteroidia, 381 Verrucomicrobiae, Planctomycetes, Kapabacteria and Chloroflexia. Differential abundance 382 testing controlled for pond site as a random effect, however, the degree and consistency of 383 enrichment did vary between pond sites.

384 Within these high level taxa, individual prokaryotic taxa were (FDR <0.05) 385 differentially abundant between pond water and the skin (Fig. 4). In general, these taxa 386 followed phylogenetic trends of enrichment, whereby if a taxon was found to be differentially 387 abundant, all other identified taxa within the same phylum were enriched in the same sample 388 type. The pond water was differentially enriched with several taxa associated with key nutrient 389 cycling processes in the aquatic environment, such as the photoautotrophs Cyanobium, 390 Synechocystis and Microcystis, and the methanotroph Methylocystis. Meanwhile, selected 391 ASVs found to be differentially enriched at the skin surface included taxa previously reported 392 as fish microbiome commensals, such as Cetobacterium, as well as additional fish related taxa, 393 which in some cases can be associated with diseases, such as Aeromonas, Pseudomonas, 394 Staphylococcus, and Streptococcus.

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396 3.3. Tilapia species differences

397 This study featured two tilapia species commonly cultured in Malawi (Coptodon rendalli and 398 Oreochromis shiranus). No significant difference of prokaryotic community alpha diversity 399 were observed between species (Fig. S5) and while beta diversity did showed potentially 400 unique community structures between species, this only explained 11% of variance. Pond site 401 in contrast explained 50% of variance in beta diversity. Additionally, intra-species dispersion 402 of C. rendalli prokaryotic communities (Average distance to median 54.49) was similar to any 403 inter-species dispersion observed between C. rendalli and O. shiranus at Blantyre (Average 404 distance to median 53.79).

405

406 **3.4. Tilapia skin core microbiome**

To further explore specific taxa prevalent within the skin microbial communities we identified 14 prokaryotic core genera of tilapia skin. Abundances of these core genera are depicted for both fish skin and pond water samples in Figure 5. Two of the prokaryotic core genera had a clear enrichment of abundance in the fish skin versus to pond water, namely ASV47: *Pseudomonas* and ASV8731: *Sphingomonas*. The remaining prokaryotic genera were found at high abundance in both pond water and skin samples, despite being classified as part of the tilapia skin core microbiome.

414

415 **4. Discussion**

Previous work has highlighted the collective contributions of microbial symbionts, the host and the environment to fish health and disease susceptibility under the pathobiome concept (Bass et al., 2019). Applying this framework to aquaculture production of finfish, the skin mucosal surface microbiome and its direct interface with the environment is likely to play a role in the maintenance of fish health and disease resilience. However, relationships between the microbial assemblages on the skin of fish in culture and their aquatic environment remain poorly established. Here, we characterised the prokaryotic and microeukaryotic communities

423 of the tilapia skin mucosal surface and accompanying water in aquaculture ponds of southern 424 Malawi in the absence of detectable disease to develop a holistic understanding of the 425 relationships between these microbial communities and niches in healthy animals and 426 environments, and against which future studies may assess how microbial dysbiosis contributes 427 to disease onset.

428 In this study, biogeographic factors played a key role in determining the diversity and 429 structure of pond water microbial communities. Pond location explained 88% of prokaryotic 430 and 76% of microeukaryotic beta diversity separation in microbial abundance profiles. 431 Significant differences in richness and alpha diversity were observed between the seven pond 432 sites. In freshwater ecosystems, both neutral and deterministic processes contribute to the 433 separation of microbial assemblages (Lear et al., 2014; Lee et al., 2013). Interestingly, just over 434 1% of prokaryotic pond water ASVs were detected in all seven pond sites, suggesting limited 435 species dispersal and/or distinct micro-ecologies between ponds. Numerous environmental 436 selective pressures could play a role in the divergence in ASVs between ponds, such as 437 alternative feeding regimes (Deng et al., 2019), differences in water physiochemistry (Qin et 438 al., 2016) and differences in pond treatments, that can include the use of probiotics (Wu et al., 439 2016) and manure fertilisers (Minich et al., 2018). Within a pond complex, some of these 440 factors will be conserved, such as weather and water source. Yet microbial community 441 divergence was still observed between ponds in the Blantyre pond complex, with two notable 442 clusters of pond sites (1,4,5 and 2,3).

443 The tight clustering of pond water samples was concurrent between both prokaryotic 444 and microeukaryotic communities suggesting cross-domain relationships shaped by ecological 445 or environmental processes. This connection has recently been observed in shrimp culture 446 ecosystems, with the deterministic process of homogenous selection largely responsible (Zhou 447 et al., 2021). In this theory, each pond site cluster represents a comparable set of environmental 448 conditions (be it nitrogen, phosphorous or oxygen availability) that exerts strong selective 449 pressures on both prokaryotic and microeukaryotic communities (Zhou and Ning, 2017). 450 Additionally, direct cross-domain ecological interactions may contribute to the observed 451 trends. For instance, phagotrophic protists and their prokaryotic prey have negative interactions 452 (Sherr and Sherr, 2002), while microalgae and bacteria can show all manner of symbiotic 453 relationships, including extensive cross-feeding (Fuentes et al., 2016; Ramanan et al., 2016).

454 The close proximities of microbial communities of pond water with those in the fish 455 outer mucosal surfaces mean they are physically closely interconnected, yet pond and skin 456 microbiomes clearly differ. Our results demonstrate these differences in prokaryotic 457 community structure, with ASV richness differing significantly, and separation by beta 458 diversity. However, there was no significant difference in alpha diversity, a finding previously 459 reported in freshwater and marine environments (Chiarello et al., 2015; Reinhart et al., 2019; 460 Webster et al., 2018). At finer taxonomic scales further separation between the skin and pond 461 water profiles was seen, and conserved across all ponds sites, with 25 ASVs differentially 462 enriched at the fish skin mucosal surface. The abundances assessed at coarse taxonomic 463 classifications reflected previous reports, namely that Proteobacteia (and in particular 464 Gammaproteobacteria) dominated the fish skin mucosal surface, as seen in a variety of 465 freshwater cichlids (Krotman et al., 2020); reviewed in depth by Gomez and Primm (2021). 466 The next most abundant bacterial classes in the fish skin were Verrucomicrobiae, Bacteroidia 467 and Clostridia. The pond water was similarly dominated by Proteobacteria, followed by 468 Cyanobacteria and Planctomycetes, which is in accordance with a previous report of the 469 bacterioplankton community in Nile tilapia aquaculture ponds in China (Fan et al., 2016).

470 Despite divergent abundance profiles, there was a high number of taxa shared between
471 pond water and skin mucosa. Only 8% of the total fish skin ASVs were unique to the skin,
472 which contrasts with that for reports on some other fish species. For example, in freshwater

473 river-dwelling mature northern pike, Esox lucius L., 36% of skin taxa were not detected in 474 samples of the surrounding water (Reinhart et al., 2019). In a study on freshwater Atlantic 475 salmon, Salmo salar L., this figure was 73% (Webster et al., 2018), where fry (8-9 months 476 post-hatch) were sampled from both wild rivers and hatcheries. Both of these studies were in 477 natural aquatic environments and flow-through systems with high water exchange rates which 478 is very different from typical carp and tilapia earthen aquaculture ponds, where daily water 479 exchange rates tend to be very limited, typically a maximum of 20% total pond volume (Nhan 480 et al., 2008). In fact, often in Africa and Asia during dry seasons, due to the lack of water 481 availability, there is no daily water exchange at all in tilapia earthen aquaculture ponds. Such 482 static conditions and high stocking densities may be reflected in a greater microbial crossover 483 between fish skin and pond water. Given the common taxa seen between the tilapia skin and 484 pond water environments, it is noteworthy that we found no correlation in ASV richness or 485 Shannon diversity between the pond and skin niches within each pond site. This finding 486 supports the hypothesis that the skin and pond water niches support uniquely structured 487 microbial communities.

488 The core microbiome refers to taxa found in the majority of samples which, by 489 inference, may therefore play an important functional role in the microbiome. Fourteen 490 prokaryotic core genera (from a total 770 genera) were identified in tilapia skin, consistent with 491 previously published findings from other studies of fewer than 20 core OTUs on fish skin 492 (reviewed by Gomex and Primm, 2021; Rosado, Pérez-Losada, et al., 2019a). Among the core 493 genera found in the tilapia skin, *Cetobacterium* has been widely reported as a core genus in the 494 gut of freshwater fish (Liu et al., 2016; Sharpton et al., 2021), including tilapia (Bereded et al., 495 2020; Elsaied et al., 2019). This genus may represent an important functional symbiont, and is 496 reputed to synthesise vitamin B12 and antimicrobial metabolites (Tsuchiya et al., 2007). Other 497 core genera and differentially enriched taxa of the skin are listed in Supplementary Table 3. 498 Ten of the fish skin core genera were also detected at relatively high abundance in pond water. 499 These include Cvanobium and Methylocystis (two of the most abundant and differentially 500 enriched phytoplankton in pond water), which may have resulted from swab sampling 501 incorporating some residual pond water. Some studies propose only retaining ASVs unique to 502 swab samples and those statistically enriched from water samples to avoid this possible 503 complication (Krotman et al., 2020). This approach, however, risks underestimating diversity 504 and missing key taxa of the fish skin that through mucosal sloughing may still be detected at 505 high abundances in water. The majority of studies make no corrections; instead, acknowledging 506 crossover is inevitable and representative of these niches.

507 While the current study included two different species of tilapia, geographic location 508 (and the associated environmental factors) of each pond site appeared to be a stronger influence 509 of prokaryotic fish skin communities than any species differences observed between *C. rendalli* 510 and *O. shiranus*. This suggests species is a complicating factor in our study but is of lesser 511 importance when considering the broader trends of microbial community separation between 512 pond water and fish skin. The importance of habitat over host taxonomy has previously been 513 demonstrated for a large-scale study of marine fish gut microbiomes (Kim et al., 2021).

514 Several of the bacterial genera we found to be differentially abundant in fish skin 515 contained species pathogenic to tilapia, including Aeromonas (hydrophila) (Dong et al., 2017), 516 Streptococcus (agalactiae) (Zhang, 2021) and Pseudomonas (fluorescens) (Hal and El-517 Barbary, 2020). In addition to these fish skin enriched genera, further potentially pathogenic 518 taxa were detected in pond water and fish skin. Namely, Plesiomonas (shigelloides) (Liu et al., 519 2015), Flavobacterium (columnare) (Dong et al., 2015) and Acinetobacter spp. an emerging 520 group of freshwater fish pathogens (Malick et al., 2020). Likewise, among the detected 521 microeukaryotic genera, there were species pathogenic to tilapia, including the parasitic ciliate 522 Ichthyophthirius multifiliis (El-Dien and Abdel-Gaber, 2009) and two skin-targeting

523 pathogenic oomycetes, Aphanomyces invadans (OIE, 2013) and Saprolegnia parasitica 524 (Ellison et al., 2018) but the metabarcoding of short hypervariable regions of marker genes 525 does not allow us to accurately assign species or strain level classifications to determine their 526 pathogenicity. The detected genera also contain numerous non-pathogenic species. The above 527 'pathogens' were all at very low (typically less than 1%) relative abundances in fish skin, and 528 indeed none of these ponds had any reported incidence of disease. This work does not preclude 529 the fact that other pathogens may be present below the limits of detection thresholds or 530 taxonomic resolution. Presence may raise the risk of opportunistic disease as primary or 531 secondary pathogens if environmental stressors create a state of dysbiosis in the fish skin to 532 favour pathobiont propagation, leading to disease onset (Bass et al., 2019).

533 Contrary to pathogenesis, many of the detected fish skin microbes will exhibit 534 commensal or mutualist relationships with their fish host. For instance, symbiotic bacteria can 535 provide colonisation resistance against pathogens through competition for nutrients and 536 adhesion sites (Legrand et al., 2018). In the microeukaryotic kingdom, ciliates were among the 537 most widely detected taxa of tilapia skin and may offer beneficial roles to the fish by predating 538 upon other microorganisms (Pinheiro and Bols, 2013). Although, the precise functional roles 539 played by symbiotic bacteria and protists of the fish skin remain almost entirely unresolved.

540 Fish skin microbiomes are inherently variable between populations (Webster et al., 541 2018), species (Chiarello et al., 2018), individuals in the same environment, and even across 542 different areas of skin (anal, caudal, dorsal and pectoral fins) of the same individual (Chiarello 543 et al., 2015). We observed separation of fish skin communities according to environment (pond 544 site), however inter-individual dispersion within pond sites was considerable, and the degree 545 of dispersion between pond sites was significantly different. This compares to pond water 546 microbiomes showing strong similarities within sites, suggesting the fish skin microbiome is 547 less subjected to environmental influences. This may be due to host factors enabling greater 548 buffering tolerance against environmental directed microbial community assembly. Host 549 genetics is further known to contribute to the inter-individual variation of fish skin communities 550 (Boutin et al., 2014). Additionally, fish age has been seen to influence individual taxa 551 abundances but offers a limited explanation of inter-individual variation at the microbial 552 community level (Rosado et al., 2021). To account for the observed inter-individual variation 553 of fish skin microbiomes we recommend increased fish numbers (6 or more) per 554 treatment/location during sampling campaigns.

555 In contrast to the variability of fish skin, pond water communities were more consistent 556 across sample sites. At most pond sites, one photoautotroph (Cyanobium or Synechocystis) was 557 dominant, at up to 20% relative abundance. While Synechocystis is well studied as a model 558 organism, little is known of *Cyanobium* and its large contribution to primary production despite 559 being among the most abundant taxa in carp aquaculture ponds (Marmen et al., 2021) and 560 freshwater lakes (Rogers et al., 2021). Additionally, the harmful algal bloom agent Microcystis 561 was detected at very high abundance in pond sites 5, 6 and 7, which is concurrent with 562 observations of rich blue-green algae during sampling. *Microcystis* (see Marmen et al., 2021, 563 2016; Zimba and Grimm, 2003), and its toxin microcystin, are frequently detected in 564 aquaculture ponds and can have toxic effects in tilapia (Abdel-Latif and Khashaba, 2017). 565 Conversely, eukaryotic microalgae, in particular diatoms, contribute positively to the 566 freshwater ecosystem as key primary producers and stabilisers of water quality (Guedes and 567 Malcata, 2012; Li et al., 2017). The barbed spines of some diatoms (*Chaetoceros* spp.), 568 however, can cause gill haemorrhage in saltwater aquaculture (Yang and Albright, 1992). Pond 569 sites 2, 3 and 6 were dominated by several diatoms, including Cyclotella, Nitzschia and 570 Aulacoseira. In other pond sites, many high abundance ASVs remained unclassified beyond 571 kingdom level, however, BLASTn searches suggested several of these were photosynthetic 572 microalgae and likely contribute to oxygen cycling.

13

573

574 **5. Conclusions**

575 This study highlights the diversity, structure and variance of the microbial communities found 576 in tilapia skin and pond water, and characterises the microbiomes for 'healthy' earthen 577 aquaculture ponds in Malawi. Future studies seeking to establish relationships between 578 dysbiosis and disease states need to take into account the inter-individual variation between 579 fish, and community variance across pond sites that also occurs within the same pond complex. 580 We found a large degree of taxa crossover between fish skin and pond water, some of which 581 may be reflective of swab sampling bias, but also unique microbial communities supported by 582 each niche. The identified core genera and differentially enriched taxa may represent conserved 583 markers of tilapia skin, whose presence and abundance should be considered in future dysbiosis 584 events, albeit in most cases the functional host relation of these taxa at the level of fish skin 585 remains to be determined. Developing a deeper understanding on the microbial communities, 586 particularly those that interface between the aquatic environment and culture species from 587 different geographies, is critical for understanding health risks in aquaculture species as 588 production expands and intensifies, bringing with it an increased risk of dysbiosis and 589 incidence of disease.

590

591 6. Author contribution statement

592 Conceptualisation CRT, DB, BT; Fieldwork CRT, DB, CG, JN; Formal analysis JM, DLC;
593 Funding acquisition CRT and DB; Methodology SA, DLC; Supervision CRT, DLC, JDD, BT,
594 JC; Visualisation JM, DLC; Writing JM & CRT; Editing JM, SA, DLC, DB, CG, JDD, CVM,
595 JC, BT and CRT.

596

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606

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611

612 9. References

- Abdel-Latif, H.M.R., Khashaba, A.M.A., 2017. Subchronic toxicity of Nile tilapia with
 different exposure routes to *Microcystis aeruginosa*: Histopathology, liver functions,
 and oxidative stress biomarkers. Vet. World 10, 955–963.
- 616 https://doi.org/10.14202/vetworld.2017.955-963
- Abdel-Tawwab, M., Hagras, A.E., Elbaghdady, H.A.M., Monier, M.N., 2014. Dissolved
 Oxygen Level and Stocking Density Effects on Growth. Feed Utilization, Physiology.
- 618 Oxygen Level and Stocking Density Effects on Growth, Feed Utilization, Physiology, 619 and Innate Immunity of Nile Tilapia, *Oreochromis niloticus*. J. Appl. Aquac. 26, 340–
- 620 355. https://doi.org/10.1080/10454438.2014.959830
- 621 Apprill, A., McNally, S., Parsons, R., Weber, L., 2015. Minor revision to V4 region SSU

622	rRNA 806R gene primer greatly increases detection of SAR11 bacterioplankton. Aquat.
623	Microb. Ecol. 75, 129–137. https://doi.org/10.3354/ame01753
624	Audebert, C., Even, G., Cian, A., Loywick, A., Merlin, S., Viscogliosi, E., Chabé, M., 2016.
625	Colonization with the enteric protozoa <i>Blastocystis</i> is associated with increased diversity
626	of human gut bacterial microbiota. Sci. Rep. 6, 25255. https://doi.org/10.1038/srep25255
627	Bass, D., del Campo, J., 2020. Microeukaryotes in animal and plant microbiomes: Ecologies
628	of disease? Eur. J. Protistol. 76, 125719. https://doi.org/10.1016/j.ejop.2020.125719
629	Bass, D., Stentiford, G.D., Wang, HC., Koskella, B., Tyler, C.R., 2019. The Pathobiome in
630	Animal and Plant Diseases. Trends Ecol. Evol. 34, 996–1008.
631	https://doi.org/10.1016/j.tree.2019.07.012
632	Beghini, F., Pasolli, E., Truong, T.D., Putignani, L., Cacciò, S.M., Segata, N., 2017. Large-
633	scale comparative metagenomics of <i>Blastocystis</i> , a common member of the human gut
634	microbiome. ISME J. 11, 2848–2863. https://doi.org/10.1038/ismej.2017.139
635	Benoiston, AS., Ibarbalz, F.M., Bittner, L., Guidi, L., Jahn, O., Dutkiewicz, S., Bowler, C.,
636	2017. The evolution of diatoms and their biogeochemical functions. Philos. Trans. R.
637	Soc. B Biol. Sci. 372, 20160397. https://doi.org/10.1098/rstb.2016.0397
638	Bereded, N.K., Curto, M., Domig, K.J., Abebe, G.B., Fanta, S.W., Waidbacher, H.,
639	Meimberg, H., 2020. Metabarcoding Analyses of Gut Microbiota of Nile Tilapia
640	(<i>Oreochromis niloticus</i>) from Lake Awassa and Lake Chamo, Ethiopia. Microorganisms
641	8, 1040. https://doi.org/10.3390/microorganisms8071040
642	Bondad-Reantaso, M.G., Subasinghe, R.P., Arthur, J.R., Ogawa, K., Chinabut, S., Adlard, R.,
643	Tan, Z., Shariff, M., 2005. Disease and health management in Asian aquaculture. Vet.
644	Parasitol. 132, 249–272. https://doi.org/10.1016/j.vetpar.2005.07.005
645	Boutin, S., Bernatchez, L., Audet, C., Derôme, N., 2013. Network analysis highlights
646	complex interactions between pathogen, host and commensal microbiota. PLoS One 8,
647	1–16. https://doi.org/10.1371/journal.pone.0084772
648	Boutin, S., Sauvage, C., Bernatchez, L., Audet, C., Derome, N., 2014. Inter Individual
649	Variations of the Fish Skin Microbiota: Host Genetics Basis of Mutualism? PLoS One 9,
650	e102649. https://doi.org/10.1371/journal.pone.0102649
651	Bramwell, P.A., Barallon, R. V., Rogers, H.J., Bailey, M.J., 1995. Extraction and PCR
652	amplification of DNA from the rhizoplane, in: Molecular Microbial Ecology Manual.
653	Springer Netherlands, Dordrecht, pp. 89–108. https://doi.org/10.1007/978-94-011-0351-
654	0 9
655	Callahan, B.J., McMurdie, P.J., Rosen, M.J., Han, A.W., Johnson, A.J.A., Holmes, S.P.,
656	2016. DADA2: High-resolution sample inference from Illumina amplicon data. Nat.
657	Methods 13, 581.
658	Carlson, J.M., Hyde, E.R., Petrosino, J.F., Manage, A.B.W., Primm, T.P., 2015. The host
659	effects of <i>Gambusia affinis</i> with an antibiotic-disrupted microbiome. Comp. Biochem.
660	Physiol. Part C Toxicol. Pharmacol. 178, 163–168.
661	https://doi.org/10.1016/j.cbpc.2015.10.004
662	CASA, 2020. Aquaculture Sector Strategy – Malawi.
663	Chiarello, M., Auguet, JC., Bettarel, Y., Bouvier, C., Claverie, T., Graham, N.A.J.,
664	Rieuvilleneuve, F., Sucré, E., Bouvier, T., Villéger, S., 2018. Skin microbiome of coral
665	reef fish is highly variable and driven by host phylogeny and diet. Microbiome 6, 147.
666	https://doi.org/10.1186/s40168-018-0530-4
667	Chiarello, M., Villéger, S., Bouvier, C., Bettarel, Y., Bouvier, T., 2015. High diversity of
668	skin-associated bacterial communities of marine fishes is promoted by their high
669	variability among body parts, individuals and species. FEMS Microbiol. Ecol. 91, 1–12.
670	https://doi.org/10.1093/femsec/fiv061
671	Davis, N.M., Proctor, D.M., Holmes, S.P., Relman, D.A., Callahan, B.J., 2018. Simple

672	statistical identification and removal of contaminant sequences in marker-gene and
673	metagenomics data. Microbiome 6, 226. https://doi.org/10.1186/s40168-018-0605-2
674	del Campo, J., Bass, D., Keeling, P.J., 2019. The eukaryome: Diversity and role of
675	microeukaryotic organisms associated with animal hosts. Funct. Ecol. 1–10.
676	https://doi.org/10.1111/1365-2435.13490
677	Delamare-Deboutteville, J., Khor, L., Chadag, V., 2021. Quick fish sampling guide for
678	disease diagnostics - Microbiome sampling guide. WorldFish.
679	Deng, Y., Zhou, F., Ruan, Y., Ma, B., Ding, X., Yue, X., Ma, W., Yin, X., 2019. Feed Types
680	Driven Differentiation of Microbial Community and Functionality in Marine Integrated
681	Multitrophic Aquaculture System. Water 12, 95. https://doi.org/10.3390/w12010095
682	Dixon, P., 2003. VEGAN, a package of R functions for community ecology. J. Veg. Sci. 14,
683	927–930. https://doi.org/10.1111/j.1654-1103.2003.tb02228.x
684	Doane, M.P., Morris, M.M., Papudeshi, B., Allen, L., Pande, D., Haggerty, J.M., Johri, S.,
685	Turnlund, A.C., Peterson, M., Kacev, D., Nosal, A., Ramirez, D., Hovel, K., Ledbetter,
686	J., Alker, A., Avalos, J., Baker, K., Bhide, S., Billings, E., Byrum, S., Clemens, M.,
687	Demery, A.J., Lima, L.F.O., Gomez, O., Gutierrez, O., Hinton, S., Kieu, D., Kim, A.,
688	Loaiza, R., Martinez, A., McGhee, J., Nguyen, K., Parlan, S., Pham, A., Price-Waldman,
689	R., Edwards, R.A., Dinsdale, E.A., 2020. The skin microbiome of elasmobranchs
690	follows phylosymbiosis, but in teleost fishes, the microbiomes converge. Microbiome 8,
691	93. https://doi.org/10.1186/s40168-020-00840-x
692	Dong, H.T., LaFrentz, B., Pirarat, N., Rodkhum, C., 2015. Phenotypic characterization and
693	genetic diversity of Flavobacterium columnare isolated from red tilapia, Oreochromis
694	sp., in Thailand. J. Fish Dis. 38, 901–913. https://doi.org/10.1111/jfd.12304
695	Dong, H.T., Techatanakitarnan, C., Jindakittikul, P., Thaiprayoon, A., Taengphu, S.,
696	Charoensapsri, W., Khunrae, P., Rattanarojpong, T., Senapin, S., 2017. Aeromonas
697	jandaei and Aeromonas veronii caused disease and mortality in Nile tilapia,
698	Oreochromis niloticus (L.). J. Fish Dis. 40, 1395–1403.
699	https://doi.org/10.1111/jfd.12617
700	El-Dien, A.H.S., Abdel-Gaber, R.A., 2009. Ichthyophthirius multifiliis infection induces
701	apoptosis in different species of Tilapia. J. Egypt. Soc. Parasitol. 39, 665–678.
702	El-Sayed, A.F.M., 2019. Tilapia Culture, 2nd ed. Academic Press.
703	Ellison, A.R., Uren Webster, T.M., Rey, O., Garcia de Leaniz, C., Consuegra, S., Orozco-
704	terWengel, P., Cable, J., 2018. Transcriptomic response to parasite infection in Nile
705	tilapia (Oreochromis niloticus) depends on rearing density. BMC Genomics 19, 723.
706	https://doi.org/10.1186/s12864-018-5098-7
707	Ellison, A.R., Wilcockson, D., Cable, J., 2021. (Accepted/In press). Circadian dynamics of
708	the teleost skin immune-microbiome interface. Microbiome.
709	https://doi.org/10.1101/2021.01.29.428758
710	Elsaied, H.E., Soliman, T., Abu-Taleb, H.T., Goto, H., Jenke-Kodam, H., 2019. Phylogenetic
711	characterization of eukaryotic and prokaryotic gut flora of Nile tilapia, Oreochromis
712	niloticus, along niches of Lake Nasser, Egypt, based on rRNA gene high-throughput
713	sequences. Ecol. Genet. Genomics 11, 100037.
714	https://doi.org/10.1016/j.egg.2019.100037
715	Fan, L.M., Barry, K., Hu, G.D., Meng, S. long, Song, C., Wu, W., Chen, J.Z., Xu, P., 2016.
716	Bacterioplankton community analysis in tilapia ponds by Illumina high-throughput
717	sequencing. World J. Microbiol. Biotechnol. 32, 10. https://doi.org/10.1007/s11274-015-
718	1962-7
719	Food and Agriculture Organization of the United Nations (FAO), 2020a. The State of World
720	Fisheries and Aquaculture 2020. Sustainability in Action. Rome.
721	https://doi.org/10.4060/ca9229en

722	Food and Agriculture Organization of the United Nations (FAO), 2020b. Fishery and
723	Aquaculture Statistics. Global aquaculture production 1950-2018 (FishStatJ) [WWW
724	Document]. FAO Fish. Aquac. Dep. URL
725	www.fao.org/fishery/statistics/software/fishstatj/en
726	Food and Agriculture Organization of the United Nations (FAO), 2020c. Cultured Aquatic
727	Species Information Programme Oreochromis niloticus (Linnaeus, 1758) [WWW
728	Document]. URL http://www.fao.org/fishery/cultured species/Oreochromis_niloticus/en
729	(accessed 7.16.20).
730	Foster, Z.S.L., Sharpton, T.J., Grünwald, N.J., 2017. Metacoder: An R package for
731	visualization and manipulation of community taxonomic diversity data. PLOS Comput.
732	Biol. 13, e1005404. https://doi.org/10.1371/journal.pcbi.1005404
733	Fuentes, J., Garbayo, I., Cuaresma, M., Montero, Z., González-del-Valle, M., Vílchez, C.,
734	2016. Impact of Microalgae-Bacteria Interactions on the Production of Algal Biomass
735	and Associated Compounds. Mar. Drugs 14, 100. https://doi.org/10.3390/md14050100
736	Gadoin, E., Desnues, C., Monteil-Bouchard, S., Bouvier, T., Auguet, JC., Roque
737	d'Orbcastel, E., Bettarel, Y., 2021. Fishing for the Virome of Tropical Tuna. Viruses 13,
738	1291. https://doi.org/10.3390/v13071291
739	Gomez, J.A., Primm, T.P., 2021. A Slimy Business: the Future of Fish Skin Microbiome
740	Studies. Microb. Ecol. https://doi.org/10.1007/s00248-020-01648-w
741	Guedes, A.C., Malcata, F.X., 2012. Nutritional Value and Uses of Microalgae in
742	Aquaculture, in: Muchlisin, Z. (Ed.), Aquaculture. IntechOpen.
742	https://doi.org/10.5772/30576
743 744	· ·
744	Guillou, L., Bachar, D., Audic, S., Bass, D., Berney, C., Bittner, L., Boutte, C., Burgaud, G.,
	de Vargas, C., Decelle, J., del Campo, J., Dolan, J.R., Dunthorn, M., Edvardsen, B.,
746	Holzmann, M., Kooistra, W.H.C.F., Lara, E., Le Bescot, N., Logares, R., Mahé, F.,
747	Massana, R., Montresor, M., Morard, R., Not, F., Pawlowski, J., Probert, I., Sauvadet,
748	AL., Siano, R., Stoeck, T., Vaulot, D., Zimmermann, P., Christen, R., 2012. The
749 750	Protist Ribosomal Reference database (PR2): a catalog of unicellular eukaryote Small
750	Sub-Unit rRNA sequences with curated taxonomy. Nucleic Acids Res. 41, D597–D604.
751	https://doi.org/10.1093/nar/gks1160
752	Hal, A.M., El-Barbary, M.I., 2020. Gene expression and histopathological changes of Nile
753	tilapia (Oreochromis niloticus) infected with Aeromonas hydrophila and Pseudomonas
754	<i>fluorescens</i> . Aquaculture 526, 735392.
755	https://doi.org/10.1016/j.aquaculture.2020.735392
756	Henchion, M., Hayes, M., Mullen, A., Fenelon, M., Tiwari, B., 2017. Future Protein Supply
757	and Demand: Strategies and Factors Influencing a Sustainable Equilibrium. Foods 6, 53.
758	https://doi.org/10.3390/foods6070053
759	Hinchliffe, S., Butcher, A., Rahman, M.M., Guilder, J., Tyler, C., Verner-Jeffreys, D., 2020.
760	Production without medicalisation – risk practices and disease in Bangladesh
761	aquaculture. Geogr. J. geoj.12371. https://doi.org/10.1111/geoj.12371
762	Kanther, M., Tomkovich, S., Xiaolun, S., Grosser, M.R., Koo, J., Flynn, E.J., Jobin, C.,
763	Rawls, J.F., 2014. Commensal microbiota stimulate systemic neutrophil migration
764	through induction of Serum amyloid A. Cell. Microbiol. 16, 1053–1067.
765	https://doi.org/10.1111/cmi.12257
766	Kelly, C., Salinas, I., 2017. Under Pressure: Interactions between Commensal Microbiota and
767	the Teleost Immune System. Front. Immunol. 8.
768	https://doi.org/10.3389/fimmu.2017.00559
769	Kim, P.S., Shin, NR., Lee, JB., Kim, MS., Whon, T.W., Hyun, DW., Yun, JH., Jung,
770	MJ., Kim, J.Y., Bae, JW., 2021. Host habitat is the major determinant of the gut
771	microbiome of fish. Microbiome 9, 166. https://doi.org/10.1186/s40168-021-01113-x

772	Krotman, Y., Yergaliyev, T.M., Alexander Shani, R., Avrahami, Y., Szitenberg, A., 2020.
773	Dissecting the factors shaping fish skin microbiomes in a heterogeneous inland water
774	system. Microbiome 8, 1–15. https://doi.org/10.1186/s40168-020-0784-5
775	Kuznetsova, A., Brockhoff, P.B., Christensen, R.H.B., 2017. ImerTest Package: Tests in
776	Linear Mixed Effects Models. J. Stat. Softw. 82. https://doi.org/10.18637/jss.v082.i13
777	Laforest-Lapointe, I., Arrieta, MC., 2018. Microbial Eukaryotes: a Missing Link in Gut
778	Microbiome Studies. mSystems 3. https://doi.org/10.1128/mSystems.00201-17
779	Lahti, L., Shetty, S., 2017. Tools for microbiome analysis in R.
780	Lane, D.J., 1991. 16S/23S sequencing, in: Stackerbrant, E., Goodfellow, M. (Eds.), Nucleic
781	Acid Techniques. Wiley, New York, pp. 115–175.
782	
	Lear, G., Bellamy, J., Case, B.S., Lee, J.E., Buckley, H.L., 2014. Fine-scale spatial patterns in
783	bacterial community composition and function within freshwater ponds. ISME J. 8,
784	1715–1726. https://doi.org/10.1038/ismej.2014.21
785	Lee, J.E., Buckley, H.L., Etienne, R.S., Lear, G., 2013. Both species sorting and neutral
786	processes drive assembly of bacterial communities in aquatic microcosms. FEMS
787	Microbiol. Ecol. 86, 288-302. https://doi.org/10.1111/1574-6941.12161
788	Legrand, T.P.R.A., Catalano, S.R., Wos-Oxley, M.L., Stephens, F., Landos, M., Bansemer,
789	M.S., Stone, D.A.J., Qin, J.G., Oxley, A.P.A., 2018. The inner workings of the outer
790	surface: Skin and gill microbiota as indicators of changing gut health in Yellowtail
791	Kingfish. Front. Microbiol. 8, 1–17. https://doi.org/10.3389/fmicb.2017.02664
792	Lever, M.A., Torti, A., Eickenbusch, P., Michaud, A.B., Šantl-Temkiv, T., Jørgensen, B.B.,
793	2015. A modular method for the extraction of DNA and RNA, and the separation of
794	DNA pools from diverse environmental sample types. Front. Microbiol. 6.
795	https://doi.org/10.3389/fmicb.2015.00476
796	Li, X., Marella, T.K., Tao, L., Peng, L., Song, C., Dai, L., Tiwari, A., Li, G., 2017. A novel
797	growth method for diatom algae in aquaculture waste water for natural food
798	development and nutrient removal. Water Sci. Technol. 75, 2777–2783.
799	https://doi.org/10.2166/wst.2017.156
800	Limbu, S.M., Zhou, L., Sun, S.X., Zhang, M.L., Du, Z.Y., 2018. Chronic exposure to low
801	environmental concentrations and legal aquaculture doses of antibiotics cause systemic
802	adverse effects in Nile tilapia and provoke differential human health risk. Environ. Int.
802	115, 205–219. https://doi.org/10.1016/j.envint.2018.03.034
	Link, J.S., Watson, R.A., 2019. Global ecosystem overfishing: Clear delineation within real
804	
805	limits to production. Sci. Adv. 5, eaav0474. https://doi.org/10.1126/sciadv.aav0474
806	Liu, H., Guo, X., Gooneratne, R., Lai, R., Zeng, C., Zhan, F., Wang, W., 2016. The gut
807	microbiome and degradation enzyme activity of wild freshwater fishes influenced by
808	their trophic levels. Sci. Rep. 6, 1–12. https://doi.org/10.1038/srep24340
809	Liu, Z., Ke, X., Lu, M., Gao, F., Cao, J., Zhu, H., Wang, M., 2015. Identification and
810	pathological observation of a pathogenic Plesiomonas shigelloides strain isolated from
811	cultured tilapia (Oreochromis niloticus). Wei Sheng Wu Xue Bao 55, 96-106.
812	Llewellyn, M.S., Leadbeater, S., Garcia, C., Sylvain, F.E., Custodio, M., Ang, K.P., Powell,
813	F., Carvalho, G.R., Creer, S., Elliot, J., Derome, N., 2017. Parasitism perturbs the
814	mucosal microbiome of Atlantic Salmon. Sci. Rep. 7, 1–10.
815	https://doi.org/10.1038/srep43465
816	Malick, R.C., Bera, A.K., Chowdhury, H., Bhattacharya, M., Abdulla, T., Swain, H.S.,
817	Baitha, R., Kumar, V., Das, B.K., 2020. Identification and pathogenicity study of
818	emerging fish pathogens Acinetobacter junii and Acinetobacter pittii recovered from a
819	disease outbreak in <i>Labeo catla</i> (Hamilton, 1822) and <i>Hypophthalmichthys molitrix</i>
820	(Valenciennes, 1844) of freshwater wetland in. Aquac. Res. 51, 2410–2420.
020	https://doi.org/10.1111/org.14594

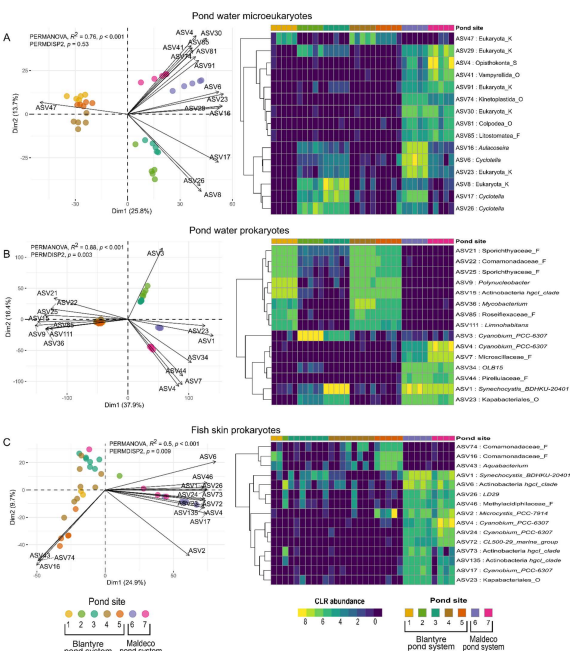
821 https://doi.org/10.1111/are.14584

822	Marmen, S., Aharonovich, D., Grossowicz, M., Blank, L., Yacobi, Y.Z., Sher, D.J., 2016.
822	Distribution and Habitat Specificity of Potentially-Toxic <i>Microcystis</i> across Climate,
823	Land, and Water Use Gradients. Front. Microbiol. 7.
825	https://doi.org/10.3389/fmicb.2016.00271
825	Marmen, S., Fadeev, E., Al Ashhab, A., Benet-Perelberg, A., Naor, A., Patil, H.J., Cytryn, E.,
820	Viner-Mozzini, Y., Sukenik, A., Lalzar, M., Sher, D., 2021. Seasonal Dynamics Are the
827	Major Driver of Microbial Diversity and Composition in Intensive Freshwater
828	Aquaculture. Front. Microbiol. 12. https://doi.org/10.3389/fmicb.2021.679743
830	Martin, B.D., Witten, D., Willis, A.D., 2020. Modeling microbial abundances and dysbiosis
831	with beta-binomial regression. Ann. Appl. Stat. 14, 94–115. https://doi.org/10.1214/19-
832	AOAS1283
833	McMurdie, P.J., Holmes, S., 2013. phyloseq: An R Package for Reproducible Interactive
834	Analysis and Graphics of Microbiome Census Data. PLoS One 8, e61217.
835	Medlin, L., Elwood, H.J., Stickel, S., Sogin, M.L., 1988. The characterization of
836	enzymatically amplified eukaryotic 16S-like rRNA-coding regions. Gene 71, 491–499.
837	https://doi.org/10.1016/0378-1119(88)90066-2
838	Minich, J.J., Zhu, Q., Xu, Z.Z., Amir, A., Ngochera, M., Simwaka, M., Allen, E.E., Zidana,
839	H., Knight, R., 2018. Microbial effects of livestock manure fertilization on freshwater
840	aquaculture ponds rearing tilapia (<i>Oreochromis shiranus</i>) and North African catfish
841	(<i>Clarias gariepinus</i>). Microbiologyopen 7, 1–15. https://doi.org/10.1002/mbo3.716
842	Minniti, G., Hagen, L.H., Porcellato, D., Jørgensen, S.M., Pope, P.B., Vaaje-Kolstad, G.,
843	2017. The skin-mucus microbial community of farmed Atlantic salmon (<i>Salmo salar</i>).
844	Front. Microbiol. 8, 1–11. https://doi.org/10.3389/fmicb.2017.02043
845	Mohammed, H.H., Arias, C.R., 2015. Potassium permanganate elicits a shift of the external
846	fish microbiome and increases host susceptibility to columnaris disease. Vet. Res. 46, 1–
847	13. https://doi.org/10.1186/s13567-015-0215-y
848	Murray, A.G., Peeler, E.J., 2005. A framework for understanding the potential for emerging
849	diseases in aquaculture. Prev. Vet. Med. 67, 223–235.
850	https://doi.org/10.1016/j.prevetmed.2004.10.012
851	Nhan, D.K., Verdegem, M.C.J., Milstein, A., Verreth, J.A. V, 2008. Water and nutrient
852	budgets of ponds in integrated agriculture-aquaculture systems in the Mekong Delta,
853	Vietnam. Aquac. Res. 39, 1216–1228. https://doi.org/10.1111/j.1365-
854	2109.2008.01986.x
855	OIE, 2013. Infection with Aphanomyces invadans (Epizootic Ulcerative Syndrome), in:
856	Manual of Diagnostic Tests for Aquatic Animals. World Organisation for Animal
857	Health.
858	Palarea-Albaladejo, J., Martín-Fernández, J.A., 2015. zCompositions — R package for
859	multivariate imputation of left-censored data under a compositional approach. Chemom.
860	Intell. Lab. Syst. 143, 85–96. https://doi.org/10.1016/j.chemolab.2015.02.019
861	Parada, A.E., Needham, D.M., Fuhrman, J.A., 2016. Every base matters: assessing small
862	subunit rRNA primers for marine microbiomes with mock communities, time series and
863	global field samples. Environ. Microbiol. 18, 1403–1414. https://doi.org/10.1111/1462-
864	2920.13023
865	Perry, W.B., Lindsay, E., Payne, C.J., Brodie, C., Kazlauskaite, R., 2020. The role of the gut
866	microbiome in sustainable teleost aquaculture. Proc. R. Soc. B Biol. Sci. 287.
867	https://doi.org/10.1098/rspb.2020.0184
868	Pinheiro, M.D.O., Bols, N.C., 2013. Use of Cell Cultures to Study the Interactions of Ciliates
869	with Fish. Springer Sci. Rev. 1, 95–113. https://doi.org/10.1007/s40362-013-0008-5
870	Pulkkinen, K., Suomalainen, LR., Read, A.F., Ebert, D., Rintamäki, P., Valtonen, E.T.,
871	2010. Intensive fish farming and the evolution of pathogen virulence: the case of

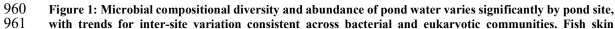
872	columnaris disease in Finland. Proc. R. Soc. B Biol. Sci. 277, 593–600.
872	https://doi.org/10.1098/rspb.2009.1659
873	Qin, Y., Hou, J., Deng, M., Liu, Q., Wu, C., Ji, Y., He, X., 2016. Bacterial abundance and
875	diversity in pond water supplied with different feeds. Sci. Rep. 6, 35232.
875	https://doi.org/10.1038/srep35232
870	Quast, C., Pruesse, E., Yilmaz, P., Gerken, J., Schweer, T., Yarza, P., Peplies, J., Glöckner,
878	F.O., 2012. The SILVA ribosomal RNA gene database project: improved data
878 879	processing and web-based tools. Nucleic Acids Res. 41, D590–D596.
879	https://doi.org/10.1093/nar/gks1219
881	Ramanan, R., Kim, BH., Cho, DH., Oh, HM., Kim, HS., 2016. Algae–bacteria
882 883	interactions: Evolution, ecology and emerging applications. Biotechnol. Adv. 34, 14–29. https://doi.org/10.1016/j.biotechadv.2015.12.003
884	Reid, K.M., Patel, S., Robinson, A.J., Bu, L., Jarungsriapisit, J., Moore, L.J., Salinas, I.,
885	2017. Salmonid alphavirus infection causes skin dysbiosis in Atlantic salmon (Salmo
886	salar L.) post-smolts. PLoS One 12, 1–17. https://doi.org/10.1371/journal.pone.0172856
887	Reinhart, E.M., Korry, B.J., Rowan-Nash, A.D., Belenky, P., 2019. Defining the Distinct
888	Skin and Gut Microbiomes of the Northern Pike (Esox lucius). Front. Microbiol. 10, 1–
889	12. https://doi.org/10.3389/fmicb.2019.02118
890	Rogers, J.E., Devereux, R., James, J.B., George, S.E., Forshay, K.J., 2021. Seasonal
891	Distribution of Cyanobacteria in Three Urban Eutrophic Lakes Results from an
892	Epidemic-like Response to Environmental Conditions. Curr. Microbiol. 78, 2298–2316.
893	https://doi.org/10.1007/s00284-021-02498-6
894	Rosado, D., Pérez-Losada, M., Pereira, A., Severino, R., Xavier, R., 2021. Effects of aging on
895	the skin and gill microbiota of farmed seabass and seabream. Anim. Microbiome 3, 10.
896	https://doi.org/10.1186/s42523-020-00072-2
897	Rosado, D., Pérez-Losada, M., Severino, R., Cable, J., Xavier, R., 2019a. Characterization of
898	the skin and gill microbiomes of the farmed seabass (Dicentrarchus labrax) and
899	seabream (Sparus aurata). Aquaculture 500, 57–64.
900	https://doi.org/10.1016/j.aquaculture.2018.09.063
901	Rosado, D., Xavier, R., Severino, R., Tavares, F., Cable, J., Pérez-Losada, M., 2019b. Effects
902	of disease, antibiotic treatment and recovery trajectory on the microbiome of farmed
903	seabass (Dicentrarchus labrax). Sci. Rep. 9, 1-11. https://doi.org/10.1038/s41598-019-
904	55314-4
905	Schliep, K.P., 2010. phangorn: phylogenetic analysis in R. Bioinformatics 27, 592–593.
906	https://doi.org/10.1093/bioinformatics/btq706
907	Sharpton, T.J., Stagaman, K., Sieler, M.J., Arnold, H.K., Davis, E.W., 2021. Phylogenetic
908	Integration Reveals the Zebrafish Core Microbiome and Its Sensitivity to Environmental
909	Exposures. Toxics 9, 10. https://doi.org/10.3390/toxics9010010
910	Sherr, E.B., Sherr, B.F., 2002. Significance of predation by protists in aquatic microbial food
911	webs. Antonie Van Leeuwenhoek 81, 293–308.
912	https://doi.org/10.1023/A:1020591307260
913	Sundh, H., Finne-Fridell, F., Ellis, T., Taranger, G.L., Niklasson, L., Pettersen, E.F.,
914	Wergeland, H.I., Sundell, K., 2019. Reduced water quality associated with higher
915	stocking density disturbs the intestinal barrier functions of Atlantic salmon (Salmo salar
916	L.). Aquaculture 512, 734356. https://doi.org/10.1016/j.aquaculture.2019.734356
917	Suphoronski, S.A., Chideroli, R.T., Facimoto, C.T., Mainardi, R.M., Souza, F.P., Lopera-
918	Barrero, N.M., Jesus, G.F.A., Martins, M.L., Di Santis, G.W., de Oliveira, A.,
919	Gonçalves, G.S., Dari, R., Frouel, S., Pereira, U.P., 2019. Effects of a phytogenic, alone
920	and associated with potassium diformate, on tilapia growth, immunity, gut microbiome
921	and resistance against francisellosis. Sci. Rep. 9, 6045. https://doi.org/10.1038/s41598-

922	019-42480-8
923	Tan, H.Y., Chen, S.W., Hu, S.Y., 2019. Improvements in the growth performance, immunity,
924	disease resistance, and gut microbiota by the probiotic Rummeliibacillus stabekisii in
925	Nile tilapia (Oreochromis niloticus). Fish Shellfish Immunol. 92, 265–275.
926	https://doi.org/10.1016/j.fsi.2019.06.027
927	Tarnecki, A.M., Brennan, N.P., Schloesser, R.W., Rhody, N.R., 2019. Shifts in the Skin-
928	Associated Microbiota of Hatchery-Reared Common Snook Centropomus undecimalis
929	During Acclimation to the Wild. Microb. Ecol. 77, 770–781.
930	https://doi.org/10.1007/s00248-018-1252-7
931	Tsuchiya, C., Sakata, T., Sugita, H., 2007. Novel ecological niche of Cetobacterium somerae,
932	an anaerobic bacterium in the intestinal tracts of freshwater fish. Lett. Appl. Microbiol.
933	071018031740002-??? https://doi.org/10.1111/j.1472-765X.2007.02258.x
934	Warnecke, F., Amann, R., Pernthaler, J., 2004. Actinobacterial 16S rRNA genes from
935	freshwater habitats cluster in four distinct lineages. Environ. Microbiol. 6, 242–253.
936	https://doi.org/10.1111/j.1462-2920.2004.00561.x
937	Webster, T.M.U., Consuegra, S., Hitchings, M., de Leaniz, C.G., 2018. Interpopulation
938	variation in the Atlantic salmon microbiome reflects environmental and genetic
939	diversity. Appl. Environ. Microbiol. 84, 1–14. https://doi.org/10.1128/AEM.000691-18
940	Wu, D.X., Zhao, S.M., Peng, N., Xu, C.P., Wang, J., Liang, Y.X., 2016. Effects of a probiotic
941	(Bacillus subtilis FY99-01) on the bacterial community structure and composition of
942	shrimp (Litopenaeus vannamei, Boone) culture water assessed by denaturing gradient
943	gel electrophoresis and high-throughput sequencing. Aquac. Res. 47, 857-869.
944	https://doi.org/10.1111/are.12545
945	Yang, C., Albright, L., 1992. Effects of the harmful diatom Chaetoceros concavicornis on
946	respiration of rainbow trout Oncorhynchus mykiss. Dis. Aquat. Organ. 14, 105–114.
947	https://doi.org/10.3354/dao014105
948	Zhang, Z., 2021. Research Advances on Tilapia Streptococcosis. Pathogens 10, 558.
949	https://doi.org/10.3390/pathogens10050558
950	Zhou, J., Ning, D., 2017. Stochastic Community Assembly: Does It Matter in Microbial
951	Ecology? Microbiol. Mol. Biol. Rev. 81. https://doi.org/10.1128/MMBR.00002-17
952	Zhou, R., Wang, H., Wei, D., Zeng, S., Hou, D., Weng, S., He, J., Huang, Z., 2021. Bacterial
953	and eukaryotic community interactions might contribute to shrimp culture pond soil
954	ecosystem at different culture stages. Soil Ecol. Lett. https://doi.org/10.1007/s42832-
955	021-0082-6
956	Zimba, P. V, Grimm, C.C., 2003. A synoptic survey of musty/muddy odor metabolites and
957	microcystin toxin occurrence and concentration in southeastern USA channel catfish
958	(Ictalurus punctatus Ralfinesque) production ponds. Aquaculture 218, 81-87.
959	https://doi.org/10.1016/S0044-8486(02)00519-7

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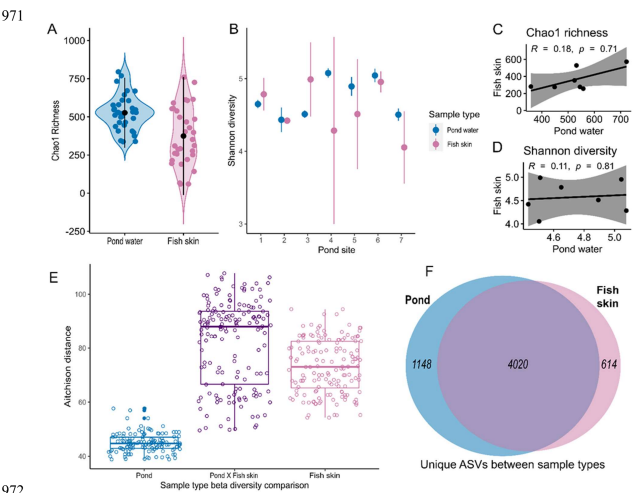


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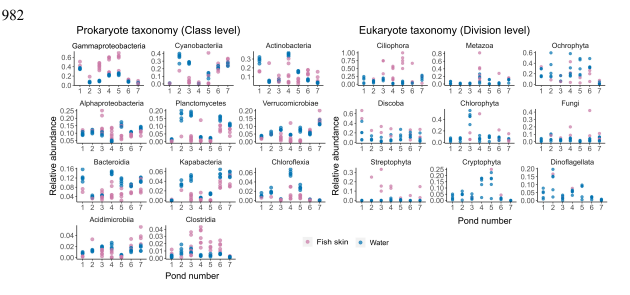


- 962 samples show significant dispersion within pond sites. Left panel: Ordination by PCoA biplots on Euclidean 963 distance of log-ratios (Aitchison distance). Points represent samples, coloured by pond site, with arrows for top
- 964 15 Amplicon Sequence Variants (ASV) explaining variation between samples. ASV abundances increase in the
- 965 direction of arrows and arrow length represents magnitude of change. Angles between arrows denote correlation
- 966 between ASVs (approximately 0° = correlated, < 90° positive correlation, > 90° negative correlation, 90° no 967 correlation).
- 968 Right panel: The centred log-ratio (CLR) abundances of top 15 discriminant ASVs are plotted as accompanying 969 heatmaps, with ASVs ordered according to a hierarchical clustering dendrogram. Labels include ASV number,
- 970 lowest available taxonomic classification and rank of this classification e.g. "_F" = Family.

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972 973 Figure 2: Comparisons of prokaryotic richness and diversity between pond water and fish skin. 974 A. Chaol richness estimates of ASVs per sample, with sample type group means and standard 975 deviations. B. Shannon diversity was calculated for each sample and plotted for each pond site as group 976 means and standard deviations. C, D. Chaol richness and Shannon diversity showed no correlation 977 between fish skin and pond water, with points plotted for the mean of each pond site. A regression line 978 of Pearson's correlation coefficient is plotted, with 95% confidence intervals. E. Beta diversity pairwise 979 comparisons of Aitchison distance between samples of pond water, pond water vs fish skin and fish 980 skin, within each pond site. F. Number of ASVs unique or shared between pond water and fish skin. 981



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Figure 3: Relative abundance of bacterial (16S) and eukaryotic (18S) taxonomic communities.
Each plot reveals the relative abundance for every sample collected at pond sites 1-7, with faceting at the taxonomic levels class (16S) and division (18S). Fish skin and pond water samples are represented in pink and blue respectively. Each facet is scaled independently.

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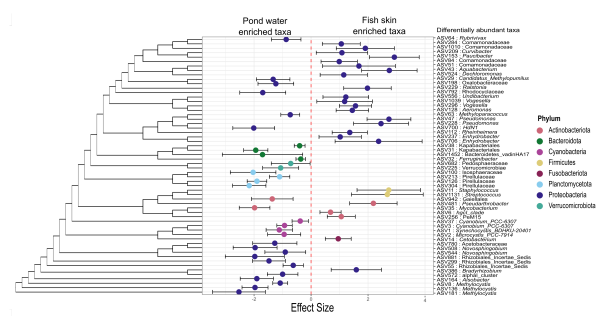
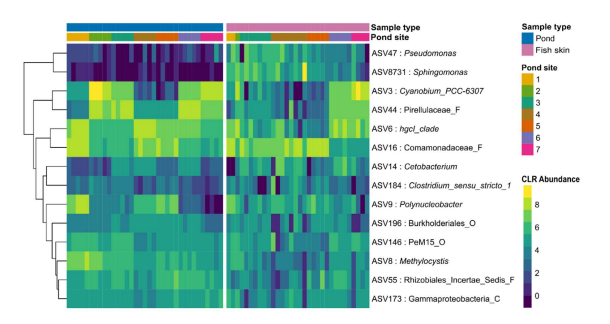




Figure 4: Differentially abundant prokaryotes of pond water and fish skin show phylogenetically conserved trends. The effect sizes and 95% prediction intervals of significant differential abundant taxa (FDR < 0.05) are plotted. Taxa to the left with a negative effect size are enriched in pond water, while taxa with a positive effect size are enriched in the fish skin. Taxa are ordered according to the phylogenetic tree, with labels included for the highest available taxonomic classification of each ASV.

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999 Figure 5: Core bacterial genera of tilapia skin communities. ASVs were amalgamated at genus level,

rarefied and core genera were classified across fish samples at an 80% prevalence threshold and 0.01%
 detection threshold. Abundance counts of core genera were transformed and are presented as non-

rarefied compositional log ratios for both pond water and fish skin samples. Abundances were utilisedfor ordering by a hierarchical clustering dendrogram.