1 First detection of Infectious Spleen and kidney Necrosis Virus (ISKNV)

2 associated with massive mortalities in farmed tilapia in Africa

3	Short running tittle: First detection of ISKNV in farmed tilapia in Africa
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18 Summary

19 In late 2018, unusual patterns of very high mortality (>50% production) were reported in intensive 20 tilapia cage culture systems across Lake Volta in Ghana. Samples of fish and fry were collected 21 and analysed from two affected farms between October 2018 and February 2019. Affected fish 22 showed darkening, erratic swimming and abdominal distension with associated ascites. 23 Histopathological observations of tissues taken from moribund fish at different farms revealed 24 lesions indicative of viral infection. These included haematopoietic cell nuclear and cytoplasmic 25 pleomorphism with marginalisation of chromatin and fine granulation. Transmission electron 26 microscopy showed cells contained conspicuous virions with typical Iridovirus morphology i.e. 27 enveloped, with icosahedral and or polyhedral geometries and with a diameter c.160 nm. PCR 28 confirmation and DNA sequencing identified the virions as Infectious Spleen and Kidney 29 Necrosis Virus (ISKNV). Samples of fry and older animals were all strongly positive for the 30 presence of the virus by qPCR. All samples tested negative for TiLV and Nodavirus by qPCR. All 31 samples collected from farms prior to the mortality event were negative for ISKNV. Follow up 32 testing of fish and fry sampled from 5 additional sites in July 2019 showed all farms had fish that 33 were PCR positive for ISKNV, whether there was active disease on the farm or not, demonstrating 34 the disease was endemic to farms all over Lake Volta by that point. The results suggest that 35 ISKNV was the cause of disease on the investigated farms and likely had a primary role in the 36 mortality events. A common observation of coinfections with *Streptococcus agalactiae* and other 37 tilapia bacterial pathogens further suggests that these may interact to cause severe pathology. 38 particularly in larger fish. Results demonstrate that there are a range of potential threats to the 39 sustainability of tilapia aquaculture that need to be guarded against.

40 Keywords: Tilapia; Africa; Tilapia Viral Diseases; fish virus; Infectious Spleen and Kidney
41 Necrosis Virus; ISKNV;

42 Introduction

The farming of tilapia species (*Oreochromis* spp.) is one of the most important sectors in aquaculture worldwide with total global production estimated at more than 6 686 000 tonnes in 2016 (FAO, 2018). In Africa, production is still dominated by Egypt with more than 1 000 000 tonnes produced in 2017 (1). However, tilapia culture has become increasingly important in several other African countries, where it boosts the local economy and constitutes an affordable source of animal protein for human consumption.

In Ghana, Nile tilapia production in 2016 had reached more than 50 000 tonnes, from only 2 000 tonnes per year in 2006 (FAO, 2016) with more than 90% of the production derived from high stocking density floating cage systems in Lake Volta. However, as production systems have intensified in the area, the industry has been increasingly affected by a range of disease issues (Jansen, Cudjoe, & Brun, 2018; Verner-Jeffreys et al., 2018).

In 2017 (Verner-Jeffreys et al., 2018) conducted the first comprehensive disease investigation in tilapia farmed in Lake Volta Ghana. *Streptococcus agalactiae* multilocus sequence type 261 was shown to be a major cause of mortality for farmed Nile tilapia and a range of other bacterial and parasitic pathogens including *Aeromonas* spp., *Streptococcus iniae*, *Flavobacterium columnare* and a *Myxobolus* sp. were also detected.

Jansen et al. (Jansen et al., 2018) conducted follow up studies, including a broad-ranging epidemiological investigation and suggested that mortalities caused by bacteria in Lake Volta were not a major concern for the local economy as farmers had managed to sustain losses by increasing production of fingerlings, treatment with antibiotics and use of autogenous vaccines.

Historically, bacterial infections were the major threat for the health of farmed tilapia (Plumb &
Hanson, 2011). However, in recent years a number of viral diseases have emerged worldwide with

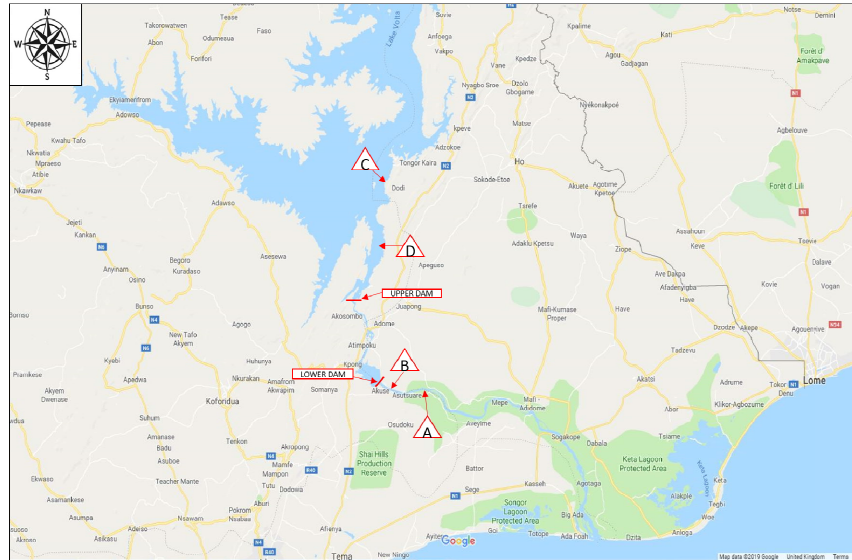
65 devastating effects for the industry(Machimbirike et al., 2019). Tilapia Lake Virus (TiLV) is 66 considered the main viral challenge to tackle, as it has spread to many producing countries, causing high mortalities in all production stages. Although TiLV has been detected elsewhere in 67 68 Africa (Hounmanou et al., 2018), the initial study by (Verner-Jeffreys et al., 2018), diagnostic 69 investigations undertaken by Ridgeway Biologicals Ltd. (Ramirez et al. unpublished data), or the 70 more recent survey undertaken by (Jansen et al., 2018), all failed to provide any evidence of TiLV 71 in diseased tilapia reared in Lake Volta. Up until September 2018, with the exception of the 72 detection of a nodavirus sequence by (Verner-Jeffreys et al., 2018), no significant association 73 between viral agents and tilapia mortality events have been demonstrated in Ghana.

74 From September 2018 to March 2019 outbreaks of disease with very high levels of morbidity and 75 mortality (60-90%), were experienced in both vaccinated and unvaccinated tilapia by farmers in 76 Lake Volta. In late-September 2018, a farm located below the lower dam in the region of 77 Asutsuare, was the first to suffer episodes of massive acute mortalities (https://goo.gl/LmqbG2 78 and <u>https://bit.ly/2NwDEbD</u>). Approximately a week after the first report, a second farm located 79 in the Akuse region (~5 km upstream of Asutsuare, but still below the lower dam) also 80 experienced acute mortalities. By mid-October, multiple floating cage-based farms in the Dodi 81 region (above the upper dam) reported losses of more than 10 tonnes per day 82 https://goo.gl/yj4oT4. In late November, farmers that had been unaffected in the Asikuma region 83 (downstream of Dodi but still above the upper dam) also started to suffer episodes of massive 84 acute mortality. (Figure 1).

By the end of 2018, most tilapia farmers in Lake Volta had reported mortalities that they were not able to contain by increased production of fingerlings or treating with antibiotics (**Supplementary File 1**, <u>https://bit.ly/2KZFFuw</u> and <u>https://goo.gl/cPmpSE</u>). Mortality events continued into and throughout 2019. We report the results of a comprehensive disease investigation, conducted at

- 89 seven affected farms from two different regions of Lake Volta, to gain insights into the causes of
- 90 these

mortalities.



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Figure 1 Map of lower region of Lake Volta in Ghana, West Africa. Red triangles indicate the regions in chronological order (A to D) where the outbreaks of mortality occurred.

94 2 Materials and Methods

95 2.1 Sampling

96 Farm 1, a medium size (approximately 600-800 tonnes per annum production) cage culture
97 operation on upper Lake Volta, was visited on the 18/10/18. Samples from 10 fish (average
98 weight 200g) were taken for bacteriology. In addition five sets of samples containing liver
99 and brain from fish from different cages were collected for molecular diagnostics as detailed
100 in Table 1.

101 Farm 2, a large cage culture operation (>2000 tonnes per annum), was visited for sampling on 102 28/11/18, 17/12/18 and 20/02/19. For the first two visits, moribund fish between 40-646g 103 collected from cages on the main lake were sampled. In the first visit for bacteriology and 104 virology on the second visit for virology only. For the third visit, responding to reports that 105 they were now experiencing very heavy mortalities in their fry production units (>70%), 106 samples of moribund fry and juveniles from both nursery cages on the main lake and from 107 their onshore hatchery supplied with water pumped from the main lake, were analysed for 108 virology. From this farm material was also taken for histological and molecular diagnostic 109 investigations during the visits as detailed in **Table 1 and Supplementary File 2**. During the 110 visits, semi-structured interviews were carried out with farm managers and/or workers. 111 Interview questions were primarily constructed to ascertain trends in mortality levels since 112 September 2019, any observed clinical signs in this time and whether there were any 113 differences in impact associated with fish life stage and/or system. Additionally, relevant 114 information on potential risk factors, mitigation measures and biosecurity practices, as well as 115 any further farmer concerns, were discussed.

A further 5 farms in the Akosombo, Atimpoku and Dasasi regions were visited and sampledfor virology, bacteriology, histology and molecular diagnostic investigation from 9-

10/07/2019. These included fry, on-growing and broodstock fish from farms of varying
capacity and with either no reported disease, fish that had survived recent mortality events on
farms or fish with ongoing clinical signs or mortalities.

121

All moribund fish from the visited farms were humanely euthanised with a lethal overdose of
tricaine methanesulfonate 1,000 mg/g (Pharmaq, Hampshire, UK) followed by brain
destruction prior to the necropsy.

125 2.2 Bacteriology

Samples for bacteriology were collected from the brain, liver, kidney and spleen with sterile cotton swabs and inoculated onto tryptone soya agar (TSA), Columbia blood agar (CBA), Tryptone yeast extract salts agar (TYES) (Southern Group Laboratory, Corby, UK) and cystine heart agar with 2% bovine haemoglobin (CHAH) (Becton Dickinson, Oxford, UK).

130 All inoculated agar plates were incubated at 28 °C for 24-72 hours. Colonies assessed as 131 significant based on occurrence and dominance were subcultured to purity on similar media. 132 Pure relevant isolates were initially identified by morphology and Gram staining. The partial 133 16S rRNA genes of the Gram negative isolates identified were PCR amplified and sequenced 134 using the method described by (Klindworth et al., 2013). Gram positive cocci forming chains 135 were screened using a Streptococcus agalactiae specific capsular typing multiplex PCR 136 developed by (Shoemaker, Xu, Garcia, & Lafrentz, 2017). The Gram negative isolates strains 137 confirmed as Aeromonas spp. based on partial 16S rRNA gene sequence analysis were 138 further characterised based on partial gyrB sequencing analysis for identification at the 139 species level as described by (Yáñez, Catalán, Apráiz, Figueras, & Martínez-Murcia, 2013).

140 **2.3 Histopathology and Electron Microscopy**

141 Tissues were fixed in neutral buffered formalin (NBF) for a minimum of 24 hr before being 142 placed in glycerol diluted 50:50 with phosphate buffered saline (PBS) for transportation to 143 Cefas. On receipt, tissues were rinsed in 70% alcohol and placed again in NBF for a final 144 period of fixation prior to processing using standard protocols in a vacuum infiltration 145 processor. Tissue sections were cut at a thickness of 3-4 µm on a Finnese® microtome, 146 mounted on glass slides and stained with haematoxylin and eosin using an automated staining 147 protocol. Stained sections were examined for general histopathology by light microscopy (Nikon Eclipse E800). Digital images and measurements were obtained using the LuciaTM 148 149 Screen Measurement software system (Nikon, UK).

150 For electron microscopy, small samples of tissues fixed in NBF as above were rinsed three 151 changes of 0.1 M sodium cacodylate buffer, followed by post fixation in 2.5% glutaraldehyde 152 in the same buffer for 1 hour prior to a second post fixation for 1 hour in 1 % osmium 153 tetroxide in 0.1 M sodium cacodylate buffer. Subsequently, fixed tissues were dehydrated in 154 an ascending acetone series acetone series and embedded in epoxy resin 812 (Agar Scientific 155 pre-Mix Kit 812, Agar scientific, UK) and polymerised at 60 °C overnight. Semi-thin $(1 \ \mu m)$ 156 survey sections were stained with 1 % Toluidine Blue and examined by light microscope to 157 identify areas of interest. Ultrathin sections (70-90 nm) of the targeted areas were placed on 158 uncoated copper grids and stained with uranyl acetate and Reynold's lead citrate (Reynolds 159 1963). Grids were examined using a JEOL JEM 1400 transmission electron microscope and 160 digital images captured using a GATAN Erlangshen ES500W camera and Gatan Digital MicrographTM software. 161

162 **2.4 Molecular diagnosis (viral and bacterial detection)**

163 The samples collected for molecular diagnosis were washed twice in 750µl of sterile 1x PBS
164 to remove the RNA-*later*® and homogenised. Total nucleic acids were extracted using

165 nanomagnetic beads i.e. Genesig Easy DNA/RNA Extraction Kit (Primerdesign,

- 166 Southampton, UK) and stored until further use.
- 167 *Multiplex PCR for detection of Streptococcus* spp.
- 168 Nucleic acids extracted were used as a template on a multiplex PCR (unpublished data) to
- 169 confirm the presence of Streptococcus spp. that had been previously reported as fish
- 170 pathogens including Streptococcus agalactiae (Delannoy et al., 2013), Streptococcus uberis
- 171 (Luo et al., 2017) and *Streptococcus dysgalactiae* (Abdelsalam, Asheg, & Eissa, 2013).
- 172 *qPCR for detection of TiLV and Nodavirus*
- 173 Nucleic acids were used for the detection of tilapia lake virus and nodavirus by quantitative
- 174 PCR using the commercial kits: Path-TiLV-EASY and Path-Betanodavirus-EASY
- 175 (Primerdesign, Southampton, UK) in the platform Genesig q16® (Primerdesign,
- 176 Southampton, UK) as per the protocols suggested by the manufacturer.
- 177 Conventional PCR for detection of megalocytiviruses

178 The generic PCR protocol for notifiable aquatic megalocytiviruses (Red Seabream Iridoviral 179 disease / Infection spleen and kidney necrosis virus (J Kurita, Nakajima, Hirono, & Aoki, 180 1998; World Organisation for Animal Health OIE, 2018)) was initially used, to screen the 181 samples of fish collected from visit 2 at Farm 2. For this, genomic DNA was extracted as 182 follows: the RNA-later® was removed and the tissue samples weighed. Depending on the 183 weight of the tissue available the samples were diluted in RTL buffer (Qiagen) to provide 184 either a 1:10 w/v or a 1:5 w/v and homogenised per fish i.e. all the organs of each fish into an 185 individual pool using Matrix A and the FastPrep apparatus (MP Biomedicals). Following 186 homogenisation, the samples were diluted further with RTL buffer to give a 1:60 w/v homogenate and total nucleic acid was extracted from $300 \,\mu$ l of the clarified sample using the

188 RNA tissue mini kit without DNase (Qiagen) and eluted in a 60 µl volume.

RT was performed at 37°C for 1 h in a 20 µl volume consisting of 1× M-MLV RT reaction
buffer (50 mM) Tris pH 8.3, 75 mM KCl, 10 mM DTT, 3 mM MgCl2) containing 1 mM
dNTP, 100 pmol random primers, 20 U M-MLV reverse transcriptase (Promega,
Southampton UK) and 4µl of the nucleic acid extracted above.

193 PCR was performed in duplicate in a 50 µl reaction volume with 2.5 µl of cDNA of total 194 nucleic acid, 25 mM dNTPs, 1 x GoTaq® buffer (2.5 mM MgCl2 solution), 5 pmol of each 195 (C1105 5'-GGTTCATCGACATCTCCGCG-3' C1106 5'primer and 196 AGGTCGCTGCGCATGCCAATC-3') and 1.25 units of GoTag® DNA polymerase 197 (Promega). The cycling conditions were as follows: 40 temperature cycles (1 min at 95°C, 1 198 min at 55°C and 1 min at 72°C) after an initial denaturing step (5 min at 95°C) followed by a 199 final extension step of 10 min at 72°C.

200 Amplified products were electrophoresed in a 2% (w/v) agarose/TAE (40 mM Tris-acetate, 201 pH 7.2, 1 mM EDTA) gel containing 1.0 μ g ml-1 ethidium bromide at 120 v for 30 mins and 202 viewed under UV light.

203 PCR products were excised from the gel and the DNA was extracted and purified by ethanol 204 precipitation. Both strands of the PCR product were sequenced using the ABI PRISM Big 205 Dye Terminator v3.1 cycle sequencing kit and the same primers used for the amplification. 206 The forward and reverse sequences were aligned and a consensus sequence generated using 207 the CLC software (Qiagen). Generated consensus sequences were compared with sequences 208 from GenBank using BLASTN (Altschul et al., 1997) and aligned using the MUSCLE 209 application of the MEGA software version 6 (Tamura, Stecher, Peterson, Filipski, & Kumar, 210 2013).

211	In addition, the OIE recommended PCR protocol for notifiable aquatic megalocytiviruses
212	(World Organisation for Animal Health OIE, 2018) developed by (J Kurita et al., 1998) was
213	used was used to screen total nucleic acids extracted from the rest of the samples fixed for
214	viral molecular analyses.

215 In analyses from farms where clinical disease was not observed, a second round PCR using

216 the nested primers C1073 5'-AATGCCGTGACCTACTTTGC-3' and C1074 5'-

- 217 GATCTTAACACGCAGCCACA -3' (15) was employed.
- 218 *qPCR for detection and quantification of megalocytiviruses*

219 The amount of virus present in the samples was also investigated by qPCR. For this the 220 homogenised tissues were subjected to total nucleic acids extraction (~20mg of each organ) 221 using the Genesig Easy DNA/RNA Extraction Kit (Primerdesign) as described earlier. The 222 extracted nucleic acids were tested using the commercial kit Path-ISKNV-EASY 223 (Primerdesign, Southampton, UK) in the platform Genesig q16[®] (Primerdesign) as per the 224 manufacturer instructions. This detects both red sea bream iridovirus and ISKNV variants. In 225 all cases fish were individually analysed either by pooling liver, spleen and brain or screening 226 individual tissues.

227 *Retrospective analyses of archived samples by qPCR*

A total of 16 samples of archived tissue homogenates from 5 different farms (that included Farms 1 and 2) were retrospectively screened for ISKNV by qPCR with the commercial kit Path-ISKNV-EASY as described before. From these 7 had been collected during 2017 and the rest in March 2018 (**Supplementary File 5**). All the samples had been previously confirmed as negative for TiLV and Nodavirus using the commercial kits Path-TiLV-EASY and Path-Betanodavirus-EASY.

234 Virus isolation

235	Frozen spleen and kidney tissue or whole fry of fish showing clinical signs taken from 2 farm
236	sites on Lake Volta on 10 July 2019 were homogenised with sand and pestle and mortar in
237	1:10 w/v cell culture transport medium (L-15 plus 1% antibiotic antimycotic solution, Gibco).
238	Homogenate was clarified by centrifugation for 10 min at 3000g, inoculated at 1:100 and
239	1:1000 final dilutions onto GF, BF-2 and E-11 cells in 24 well cell culture plates (Gibco) and
240	incubated at 25°C. After 7 days cells were blind passaged and incubated for a further 7 days.
241	Cells were observed for cytopathic effect (CPE) by light microscopy with phase contrast
242	(IX83 inverted microscope, Olympus, UK).

243

3 Results

245 3.1 Farm visits

On the first visit to Farm 2 on the 27/11/18, farm staff reported very high and ongoing mortalities (Figure 2) in fish bigger than 20g, including broodstock, but no significant losses in fingerlings. Losses reportedly peaked at about 670 crates (equating to approximately 40 tonnes per day) shortly after this visit on the 2/12/2018 (**Figure 2**).

Losses were so severe that accurate estimation was not possible, with more than 50 additional labourers recruited locally just to remove dead and moribund fish during the peak period. By the second visit to Farm 2, losses of ongrowing fish had reportedly declined back to the background 10-20 crates per day more typically observed i.e. less than 1-2 tonnes per day (**Figure 2**).

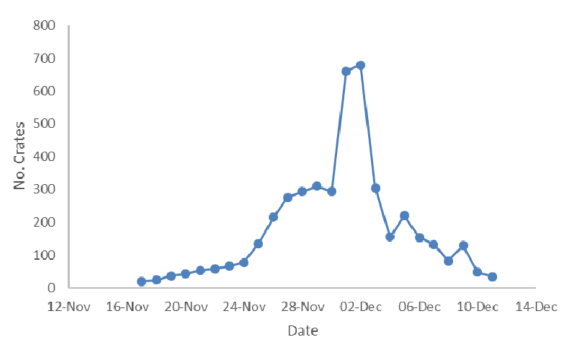
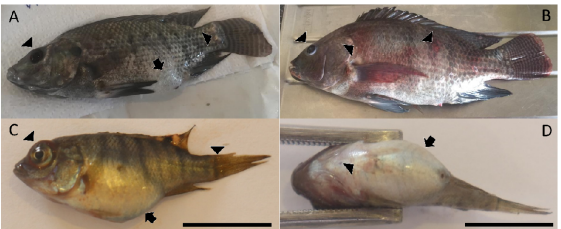


Figure 2. Number of crates of dead/ moribund fish removed from Farm 2 during period of maximum losses. Daily mortalities were estimated based on numbers of crates of rejected ongrowing tilapia collected each day by farm staff. Each crate typically contained approximately 60kg tilapia collected from the cages and rejected because they were either dead, moribund or displayed other adverse signs that prevented their sale.

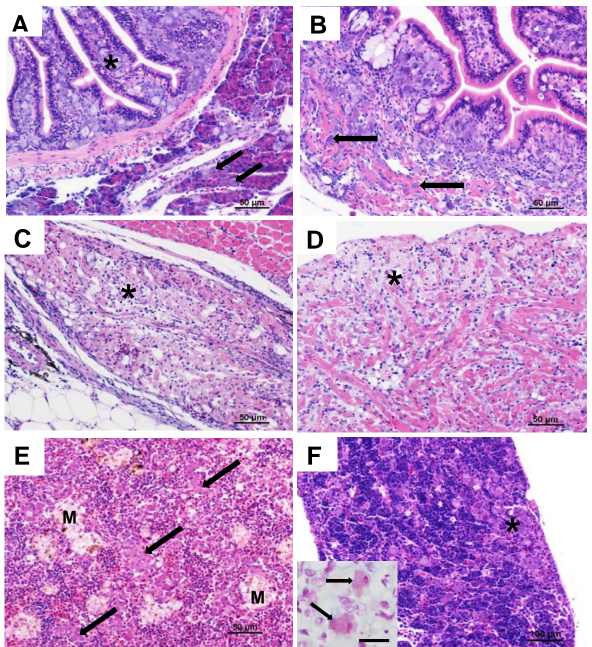
- During the first two visits to the farms, but particularly to Farm 2, diseased fish were observed swimming away from the school with erratic swimming i.e. on one side, in circles, lethargic, with no equilibrium, upside down etc. (**Supplementary File 3**).
- Externally, the fish displayed a range of clinical signs, including skin nodules, frayed fins, loss of eyes, opaque eyes, loss of scales, exophthalmia, anorexia, decolouration or darkened skin, excess of mucous, skin haemorrhages and distended abdomen (**Figure 3**). At necropsy, fish from both the first visits to Farm 1 and 2 presented with marked ascites, enlarged and haemorrhagic organs including the spleen, heart, brain, gills, but most notably liver and kidney. The gastrointestinal track was empty of solids but contained transparent fluid similar to that also seen in the peritoneal cavity.

In contrast to the earlier visits, when Farm 2 was visited on 20/02/19, there were very high and ongoing mortalities in the fry production systems (>70%). This was both in their onshore hatchery (supplied with water pumped from the main lake) where eggs were hatched and held until the fry were approximately 20g, and in the nursery cages on the main lake to which fry had been transferred. As with the larger fish sampled, affected fry showed erratic swimming behaviour, skin haemorrhages and severe ascites as the main clinical signs (**Figure 3**).



279 280 Figure 3 External lesions and clinical signs in diseased Nile tilapia in Lake Volta. A 281 Ongrowing fish with emaciation slight ascites (arrow), endophthalmia (left arrow head) and 282 skin purulent abscess (right arrow head). **B** Broodstock with microphthalmia left arrow head), 283 skin haemorrhages (middle arrow head) and skin ulcers (right arrow head). C juvenile with 284 exophthalmia (left arrow head), ascites (arrow) and loss of scales, excess of mucus and 285 haemorrhages (right arrow head). Bar = 1 cm. **D** Ventral view of juvenile fish presenting 286 severe ascites black arrow and skin haemorrhages (arrow head). Bar= 1cm. 287

- 288
- 289 3.2 Megalocytivirus and bacterial infections in Farm 2 during period of high mortalities
- 290 Four out of the ten fish examined histologically showed mild tissue necrosis in the spleen and
- 291 renal haematopoietic tissue with the presence of large numbers of cells showing relative
- 292 eosinophilia cytoplasmic and nuclear pleomorphism with margination of chromatin in some
- affected nuclei suggestive of a viral infection diffused throughout the tissue (Figure 4).



294 Figure 4 Histopathological cross sections of tissues from diseased Nile tilapia in Lake Volta. All 295 sections stained with H&E unless otherwise stated. A. Section through the intestine of an infected fry 296 showing the presence of large numbers of megalocytes in the lamina propria (*) but not affecting the 297 mucosa or underlying muscularis. Note the presence of isolated megalocytes in the pancreatic acinar 298 tissue (arrows). Bar = 50 μ m. **B.** Same fish as in A, showing infiltration and disruption of the 299 muscularis associated with the presence of megalocytes (arrows). Bar=50 µm. C. Pseudobranch 300 showing extensive necrosis (*). Bar= 50μ m. **D.** Heart showing necrosis of the ventricular muscle (*), 301 particularly in the peripheral regions associated with inflammation. Bar=50µm. E. Spleen from a fish 302 from Farm 2 during a mortality episode. Affected cells show pronounced eosinophilia and are 303 distributed throughout the section (*). Numerous vacuolated macrophage aggregates are also present 304 (M). Bar = $50\mu m$. F. Pronephros showing extensive distribution of megalocytes without associated 305 tissue necrosis (*). Bar=100µm. Inset shows pale staining of DNA in the cytoplasm of affected cells 306 (arrows). Feulgen stain. Bar=25µm.

307 The ultrastructure of affected cells revealed the presence of conspicuous viral particles. Some 308 cells showed the presence of numerous diffusely spread virions within the hypertrophied 309 nucleus of affected cells, which also showed degradation of the nuclear membrane (Figure 310 5). Virions were approximately 160 nm in diameter (Figure 5D). Virion morphology 311 showing icosahedral symmetry with an external double membrane and internal core was 312 consistent with that of viruses from the genus Megalocytivirus. In other cases, the nucleus of 313 affected cells appeared condensed and densely stained in histological and resin sections. 314 TEM showed that in these cases the nucleus was tightly packed with virions in various stages 315 of maturation and with some evidence of formation of 'arrays' (Figure 5). In two of these 316 cases, a concomitant Gram +ve bacterial infection was also present in the gill and liver. 317 Incidental findings of gill parasites, myxozoan cysts and monogeneans, both present in low 318 numbers as well as low grade epitheliocystis were observed. The brain of a single fish 319 harboured small cysts containing necrotic debris. Gram and Ziehl-Neelsen staining did not 320 demonstrate the presence of bacteria. Other tissues appeared normal. For the set of samples 321 collected at the height of the mortalities from Farm 2 on 28/11/18, Aeromonas jandaei, 322 Aeromonas veronii (from skin), Streptococcus agalactiae capsular type Ib biotype 2 (non 323 haemolytic) and *Edwardsiella tarda* (from liver and kidney) were recovered.

324 3.3 PCR and sequence confirmation of Megalocytivirus infection

Within the 7 individuals collected from Farm 2 on the second visit that were analysed with the protocol proposed by (Rimmer, A. E., Becker, Tweedie, & Whittington, 2012), a single fish (fish 1) was clearly positive by PCR for RSIV/ISKN and a second very weak product of the correct size was also seen in tissues from Fish 6 (**Supplementary File 4**) The consensus sequence generated from the PCR product from Fish 1 was confirmed as ISKNV sharing 100% nucleotide identity with ISKNV accession no AF371960.1. In the phylogenetic

- analysis the sequence was assigned to the same lineage as the bulk of the ISKNV sequences
- **332** (**Figure 6**).
- 333 The samples collected from Farm 1 and Farm 2 at the height of the mortalities were found to
- be strongly positive when they were retrospectively tested using the (J Kurita et al., 1998;
- 335 World Organisation for Animal Health OIE, 2018) current OIE recommended PCR method.

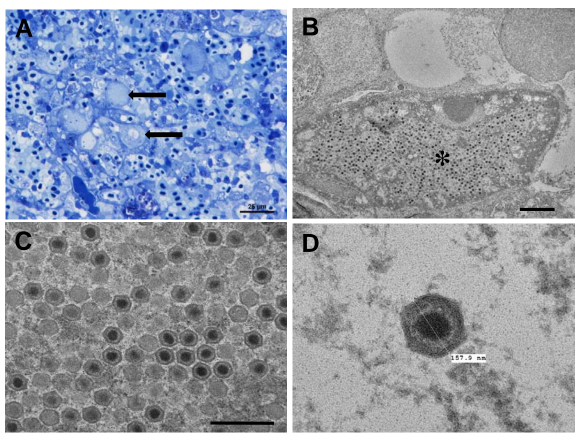


Figure 5 Micrographs of the kidney of diseased Nile tilapia in Lake Volta. A Semithin 336 337 section of affected renal tissue showing characteristic cellular hypertrophy (black arrows). 338 Toluidine Blue stain. Scale bar = $25 \mu m$. B. Electron micrograph of an individual infected cell 339 with numerous viral particles (*). Adjacent cells appear uninfected. Scale bar = $2\mu m$. C. 340 Numerous lightly stained developing virions with mature virions. The outer membranes and 341 central electron lucent core are clearly visible. Scale bar = 500nm. **D.** Mature icosahedral 342 virion showing detail of the outer caspid and inner membrane with central electron lucent 343 core.

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349 350 3.4 qPCR results for megalocytivirus from ongrowing tilapia samples

351	The qPCR results confirmed that all the fish sampled from Farm 1 in the Akuse region were
352	positive for megalocytivirus. Also, as expected, the samples collected from the first Farm 2
353	visit during the peak of mortality (28/11/18), were also positive and presented the highest
354	viral titres in grow out fish with some containing over 3 x 10^6 copies per sample reaction. In
355	contrast, the samples collected from grow out fish during the second visit to Farm 2 had
356	much lower viral copy numbers.

357 All the archived samples collected in 2017 and March 2018 were negative for ISKNV, when 358 tested by qPCR (Supplementary File 5).

359 3.5 Fry samples had characteristic megalocytivirus pathology and high copy numbers of 360 virus

361 All the fry samples collected from Farm 2 on 20/02/19, when there were reportedly very high 362 (>90%) losses in that part of their system, were positive for ISKNV by qPCR and these presented the highest titres in the study with some containing up to 1.5×10^7 copies per 363 364 sample reaction (Table 1). All fry showed moderate to marked histological and pathological 365 features of infection with megalocytivirus. Splenic tissues were necrotic and associated with 366 the presence of megalocytes characterised by light sometimes granular cytoplasmic 367 basophilia and hypertrophied nuclei. Kidney also showed the presence of megalocytes but 368 usually with only mild cellular necrosis. The lamina propria in the intestine of a single fish 369 was packed with megalocytes (Figure 4), although necrosis appeared to be absent and the 370 epithelial layer remained intact. Gills showed only minimal focal necrosis, usually affecting 371 the underlying connective tissues. In some fish the choroidal rete was affected with mild 372 necrosis and variable numbers of megalocytes and most fish showed mild myositis with few

373	megalocytes in the skeletal muscle. However, a single fish showed extensive inflammation
374	and myofibrillar necrosis (Figure 4). Connective tissues of the head and in particular around
375	the pharyngeal teeth were often infiltrated with megalocytes. Brain and spinal cord appeared
376	normal. Liver samples were not examined as they were used for virus quantification.

377

378 3.6 Other pathological observations

Megalocytivirus -like pathology was not observed in any of the samples taken from the second visit to Farm 2, two weeks after the peak of mortalities had passed. As with the samples taken at the first visit, there was evidence of bacterial infection in some individuals, particularly fish 5, had marked bacterial infection of the spleen, liver and brain (meningitis). All the samples were positive for the presence of *Streptococcus agalactiae* and negative for *Streptococcus uberis* and *Streptococcus dysgalactiae* by PCR (**Supplementary File 6**).

A range of different potential bacterial species, including *Aeromonas jandaei* and *Plesiomonas shigelloides* were recovered from fish from Farm 1 (**Table 1**), but not as pure growths or high quantities, suggesting they had a limited role in observed disease in these animals.

All the samples from Farm 1 and Farm 2 tested for TiLV and from Farm 2 for nodavirus by
qPCR were all negative (Table 1 and Supplementary File 4).

391

392 The results for all the individual fish tested are shown in Supplementary File 2.

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394 3.7 Follow up testing in July 2019

395 A further 5 farms sampled in July 2019 were tested using nested conventional PCR (15). 396 Fish on farms Farms 6 (fry) and 7 (lake ongrowers) were experiencing ongoing mortality and 397 showing typical clinical signs described above at time of sampling. All samples were 398 strongly positive in a single round assay and virus was isolated in cell culture from both 399 farms. For samples from farms 4 (fry) and 5 (lake ongrowers) which both were 400 recovering from recent mortality events, but had no remaining observable clinical disease, 401 between 40-100% of samples tested positive but these were in second round only of nested 402 assay and in some cases only in one of duplicate reactions indicating low levels of virus. 403 From farm 3 (all age groups) for which no mortality events had been reported between 30% 404 to 100% of samples were positive but all in the second round only. On sequencing 405 representatives from all positive farms showed identical sequence (data not shown).

Sampling date & Farm	No. of Fish and size	Farm observations and ‡clinical signs	Bacteriology	Histopathology	Virology Results [¥] [] – qPCR _{RSIV/ISKNV} copy number/sample reaction range
18/10/2018 †Farm 1	n=5; 100-300g	Wide scale ongoing mortalities in on growing fish in cages.	Aeromonas veronii; Aeromonas jandaei; Plesiomonas shigelloides; Chryseobacterium sp; Acinetobacter johnsonii;	Not done	*qPCR _{TILV} 5/5 -ve *cPCRa _{ISKNV} 4/5 +ve *qPCR _{RSIV/ISKNV} 5/5 +ve [7.5 x 10 ¹ – 4.60 x 10 ⁵]
28/11/2018 †Farm 2	n=14; 74-401g	Wide scale ongoing mortalities in on growing fish in cages (Figure 2). On shore fry production unit: no unusual mortalities.	Aeromonas veronii; Aeromonas jandaei; Streptococcus agalactiae capsular type Ib biotype 2 (non-haemolytic); Edwardsiella tarda	Evidence of Gram negative and Gram positive bacterial and viral# infection (including in same fish)	$\begin{array}{c} qPCR _{TiLV} 14/14 -ve \\ qPCR _{NODA} 14/14 -ve \\ *cPCRa _{ISKNV} 12/14 +ve \\ *qPCR _{RSIV/ISKNV} 14/14 +ve \\ [2.48 x 10^1 - 3.3 x 10^6] \end{array}$
17/12/2018 †Farm 2	n=7; 40-646g	Much lower mortalities in on growing fish in cages than previous sampling visit. On shore fry production unit: no unusual mortalities.	Not done	Evidence of bacterial infection but not of viral infection	$\begin{array}{c} qPCR_{TiLV} 7/7 \text{ -ve} \\ qPCR_{NODA} 7/7 \text{ -ve} \\ \textbf{cPCRb}_{RSIV/ISKNV} 1/7 \text{ +ve} \\ \textbf{megalocytivirus, sequence} = \textbf{ISKNV} \\ ^{*}qPCR_{RSIV/ISKNV} 6/7 \text{ +ve} \\ [1 \ x \ 10^{0} \ - 24 \ x 10^{0}] \end{array}$
20/02/2019 †Farm 2	n=14; 6-9 cms	Limited mortalities in on growing fish in cages. Very severe mortalities in on shore fry production unit.	Not done	Very severe virus- associated pathology (in tissue samples from all fish).	$\begin{array}{c} qPCR \\ TiLV not done \\ qPCR \\ NODA not done \\ cPCRa \\ ISKNV \\ 7/7 + ve \\ qPCR \\ RSIV/ISKNV \\ 13/13 + ve \\ [5.1 x 10^5 - 1.51 x 10^7] \end{array}$
09/07/2019 Farm 3	Grp 1, n=3; 60-80g Grp 2, n=3; 20-40g Grp 3, n=5; 1-2g fry Grp 4, n=5 pools of \leq 0.2g fry	No mortality events reported at site. Grp 1 on-growing in lake, Grps 2-4 in ponds.	Grps 1 & 2 - No significant colonies Grps 3-5 Not done	Evidence of metacercaria, bacterial gill epitheliocystis and myxospordiosis in Grps 1 & 2 No evidence of viral infection.	cPCRb _{RSIV/ISKNV} Grp 1 - $2/3$ +ve (nested only) Grp 2 - $3/3$ +ve (nested only) Grp 3 - $5/10$ +ve (nested only) Grp 4 - $3/10$ +ve (nested only)

09/07/2019 Farm 4	n=5 pools of ≤0.2g fry	Reported recent mortality event now recovering. No clinical signs	Not done	Myxosporidian cysts in cranial sub- epithelial and connective tissue	cPCRb _{RSIV/ISKNV} 2/5 +ve (nested only and in one of duplicates only)
10/07/2019 Farm 5	n=5 x3 fish organ pools; ~40g	Ongrowing fish on lake, recent mortality event in adjacent cage cleared previous month, no clinical signs in sampled fish	No significant colonies	not done	cPCRb _{RSIV/ISKNV} 5/5 +ve (nested only, one of which weak in 1st round)
10/07/2019 Farm 6	n=10 x3 fish pools; 0.2-0.75g	Ongoing losses of approx. 30% in fry pools. Fry observed with distended abdomen and disturbed swimming.	Not done	Evidence of viral infection, occasionally severe	cPCRb _{RSIV/ISKNV} 10/10 +ve Virus isolation – BF cells
10/07/2019 Farm 7	n=5 x2 fish organ pools; 20-40g	Internal haemorrhaging, ascites, pale kidney, anaemia, reduced spleen and enlarged liver	No significant colonies	Evidence of viral infection	cPCRb _{RSIV/ISKNV} 5/5 +ve Virus isolation – BF cells

406 **Table 1 Summary of sampling and results**

407 † See Materials and Methods for farm descriptions

408 ‡ All moribund fish, including fry, presented similar signs: erratic swimming, lethargy, ascites, swollen, dark spleen, haemorrhagic livers and other organs.

409 # Haematopoietic cell nuclear and cytoplasmic pleomorphism with marginalisation of chromatin and fine granulation.

410 ¥ qPCR _{TiLV}, qPCR _{NODA}, qPCR _{RSIV/ISKNV}: results of qPCR testing for tilapia lake virus, nodavirus and ISKNV respectively. cPCRa & cPCRb : conventional PCR using

411 OIE recommended protocols by Kurita et al., 1998 (a) (J Kurita et al., 1998) and Rimmer et al., 2012 (b) (Rimmer, A. E. et al., 2012) protocols, results in bold represent

412	index	finding,	*represents	retrospective	testing	post	index	finding.
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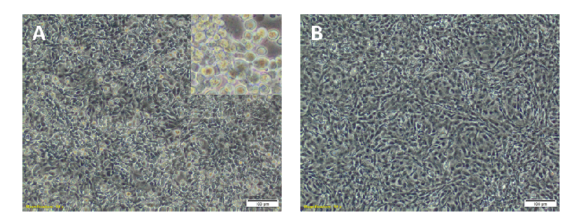


462 numbers and the host species are included where available.

463 3.8 Virus isolation

Fish material from Farms 6 and 7 was inoculated onto BF-2, E-11 and GF cell lines. Cytopathic effects (CPE) were observed in BF-2 and E-11 cells but not GFs on first inoculation (Figure 7). On passage (P1) in the same cell types CPE was only observed for BF-2 cells and intensity of CPE was diminished. Isolated virus from clarified harvested cell culture supernatant from the P1 BF-2 cells was confirmed positive for ISKNV by PCR with sequence identical to that obtained by PCR direct from tissue homogenate (data not shown).

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- 471



473 Figure 7 Virus isolation and cytopathic effects in BF-2 cells. (A) BF-2 cells at 5 days post
474 inoculation (-3 dilution) showing enlarged, rounded, phase bright cells. Inset, advanced
475 extensive CPE from -2 dilution of BF-2 at same time point. (B) Negative control BF-2 cells
476 at 5 days.

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479 4 Discussion

480 The results suggest ISKNV had a significant role in the high mortalities experienced by the 481 two farms that were investigated. Fish sampled from the second farm at the height of the 482 disease outbreak showed severe clinical and pathological signs typically associated with

483 infection by the virus (including visualisation of distinctive megalocytes with characteristic 484 virions). Both, these fish and those sampled earlier from Farm 2, had extremely high copy 485 numbers of RSIV/ISKNV Megalocytivirus-like virions and ISKNV was confirmed by gene 486 sequencing. Further investigations later in 2019 of five further farms from different areas of 487 lake Volta showed by then the disease was widely established (endemic) in farms across the 488 lake, with both symptomatic and asymptomatic fish positive by PCR for the virus. Farmers 489 emphasised how the disease had a devastating effect on the industry during discussions on 490 this latter visit. They reported how the disease continued to have an impact on the broodstock 491 and growout fish at a lower rate, however the mass mortalities were now predominantly 492 observed in juvenile fish (1-5g in weight). Survival rates to the growout stage, at that time, 493 were estimated to be as low as 5-20%. Juvenile mortalities were reported to be episodic in 494 nature, occurring a few days after the sex reversal process or translocation to lake cages, and 495 lasting up to 3 to 4 weeks. These stress triggered mortality outbreaks may be indicative of 496 either widespread persistence of virus in the environment or a latent ISKNV stage.

497 To improve survival rates, some farmers had trialled reductions in juvenile stocking densities. 498 Large reductions in stocking density were only associated with small increases in survival, 499 therefore this practice was not considered a viable solution. Instead, farmers resorted to a 500 substantial increase in the level of juvenile production and stocking rates, trading off higher 501 overall mortality with some guarantee of a small but not inconsequential harvest. The 502 economic impact of ISKNV has been significant. The higher production costs and reduced 503 harvests resulted in most farms having to either temporarily or permanently halt production. 504 As larger farms can be the primary employers of some villages, the disease also had a direct 505 impact on the livelihoods of local community members. It was also reported that tilapia 506 market price had more than doubled due to the production shortages and that the feed sales of 507 Raanan Fish Feeds had reduced by 70% (https://thefishsite.com/articles/ghanaian-fish-

508 <u>farmers-seek-help-from-big-business</u>), both potential indicators of the virus having an impact
 509 on a much larger scale.

510 The results would be consistent with a recent introduction of the virus onto one or more farms 511 prior to October 2018, that then extended upstream of the dam to other farms resulting in the 512 unusual, widespread and significant mortalities observed. Firstly, there were no observations 513 of typical ISKNV-associated pathology in any previous disease investigations, on the affected 514 or other farms. Secondly, the limited PCR screening in this study of archived samples of 515 diseased tilapia recovered from before the major mortality event including both farms, were 516 all negative. Moreover, (Verner-Jeffreys et al., 2018) also screened for the presence of 517 RSIV/ISKNV by PCR in 2016, including samples from the affected farms, without detecting 518 the virus, or associated pathology.

519 Iridioviruses are large icosahedral cytoplasmic double stranded DNA viruses, which can 520 infect a wide range of hosts, including invertebrates and poikilothermic vertebrates. The 521 family Iridoviridae includes five genera: Iridovirus, Chloriridovirus, Ranavirus, 522 Lymphocystivirus and Megalocytivirus (Jancovich et al., 2012). Fish pathogenic iridoviruses 523 Lymphocystivirus are representatives of *Ranavirus*, and *Megalocytivirus* genera 524 (Subramaniam, Shariff, Omar, & Hair-Bejo, 2012; Whittington, Becker, & Dennis, 2010). 525 Infectious spleen and kidney necrosis virus (ISKNV), is a member of the genus 526 Megalocytivirus (Jun Kurita & Nakajima, 2012), and causes disease in a range of freshwater 527 and marine fish species (Subramaniam, Shariff, Omar, & Hair-Bejo, 2012; Whittington, 528 Becker, & Dennis, 2010). ISKNV is closely related to red sea bream iridovirus and both 529 viruses are listed by the OIE as notifiable pathogens (World Organisation for Animal Health 530 OIE, 2018). Although tilapia is not listed as a susceptible species by the OIE at present 531 (World Organisation for Animal Health OIE, 2018), recent reports from the USA

(Subramaniam et al., 2016) and Thailand (Suebsing et al., 2016) suggest that it is a susceptible species likely to suffer significant mortality. The change in the known host range of virus needs to be communicated to the international community to prevent future transboundary spread through movement of infected tilapia.

It was interesting to note that many of the ISKNV positive fish were actively co-infected with *Streptoccocus agalactiae* and other bacterial pathogens, presenting severe bacteraemia / meningitis, as well as ISKNV-associated pathology. The high mortalities on the farms in the larger on-growing fish may well have been exacerbated by these coinfections.

540 Although the mortalities were initially confined to on-growing fish in cage culture systems, 541 the later observations of very high ISKNV associated mortalities in fry, associated clinical 542 signs and high viral copy numbers shows the virus likely affects all life stages. As fry are 543 often reared in onshore facilities below the dam and then translocated to on-growing cages on 544 the main lake, this may have been one of the routes that disease was rapidly spread after it 545 first emerged. Anecdotally, at the time of writing, farmers on the sites visited report that 546 mortalities in on-growing facilities have declined, while fry production continues to be badly 547 affected. It is possible that surviving fry have been exposed to the virus and then protected 548 against subsequent exposure. This suggests that immunisation of fry, or use of previously 549 exposed individuals, could represent a practical disease management strategy. Vaccination as 550 a control strategy may be used to control red sea bream iridiodovirus (Nakajima et al., 1999; 551 Shimmoto, Kawai, Ikawa, & Oshima, 2010) and there are also encouraging reports of its 552 potential effectiveness for protection against ISKNV in mandarin fish (Dong, Weng, He, & 553 Dong, 2013). As some of these reports showed efficacy using formalin killed virus infected 554 cells (Dong et al., 2013), rapid development and testing of vaccines based on the direct use

of the strain(s) of ISKNV circulating in Lake Volta farms (e.g. autogenous vaccines) should
be possible.

557 Outbreaks of disease that cause significant morbidity and/or mortalities in an aquaculture 558 operation is always a major concern. This is exacerbated when this appears to represent the 559 incursion of a new agent into a system, or region (country or zone in a country) which has not 560 previously been affected. A stark example of this is the epidemic of infectious salmon 561 anaemia virus (ISAV) which reduced production by three quarters and resulted in severe 562 economic and social crisis in the developing Chilean Atlantic salmon industry between 2007 563 and 2010 (Godoy et al., 2008; Mardones, Martinez-Lopez, Valdes-Donoso, Carpenter, & 564 Perez, 2014; Vike, Stian Nylund, & Nylund, 2009).

The Ghanaian authorities have for some time been concerned that the, to date, successful, expansion of its industry on Lake Volta may be affected by such disease incursions. Partly for this reason, and to safeguard the genetic integrity of Lake Volta Nile tilapia strains, they have tried to limit the culture to locally reared Nile tilapia stocks. However genetic testing by the Ghanaian Fisheries Commission (Ziddah *et al.*, Unpublished Observations) showed that fish on some of the farms on Lake Volta were likely of imported GIFT strain origin (Ponzoni et al., 2011) or hybrids of GIFT and indigenous strains.

572 If farmers have been illegally sourcing broodstock from Asia and other areas, that would be 573 an ideal method of translocating pathogens from one region to another. It should be noted 574 though that ISKNV has also been detected in internationally traded freshwater ornamental 575 species, theoretically posing another possible introduction route (Jung-Schroers et al., 2016)

576 It is very possible that this is not the first time disease introduction has taken place in Lake 577 Volta Ghana. The study by (Verner-Jeffreys et al., 2018) showed that outbreaks of S. 578 *agalactiae* investigated in 2016 were all caused by genetically indistinguishable isolates of

579 ST 261, with closest genetic identity to Asian isolates. Discussions with affected farmers at 580 the time suggested, that S. agalactiae associated mortalities were a relatively recent 581 phenomenon, although the disease is now clearly endemic to all the areas in the Volta area. 582 Other studies have shown that S. agalactiae ST261 has likely been translocated around the 583 world in association with farmed tilapia (Kawasaki et al., 2018). As a single large 584 epidemiological unit, it will be difficult to control transmission of virus between farms on 585 lake Volta. It is important to try to prevent spread from Lake Volta to surrounding 586 watersheds and the wider African continent by control over movement of live fish and 587 equipment. The development of biosecure offline hatcheries with borehole water or UV 588 treatment of water, to facilitate production of juveniles which survive to a size they can be 589 vaccinated will likely be key to vaccine control. Additionally, recent technological advances 590 in rapid selective breeding (Houston et al., 2020; Robledo, Palaiokostas, Bargelloni, 591 Martínez, & Houston, 2018) should be employed to develop ISKNV disease resistant 592 populations, or strains of tilapia to enable the industry to recover.

Although most attention to date has focussed on the emergence and spread of TiLV within the tilapia industry world-wide, these results also demonstrate that there is a range of other potential threats to the sustainability of tilapia aquaculture.

596 Conclusion

597 This is the first report of Infectious Spleen and Kidney Necrosis Virus (ISKNV) in farmed 598 tilapia in Africa. ISKNV was found in co-infection with *Streptococcus agalactiae* and other 599 bacterial pathogens in Lake Volta, Ghana. The correlations seen between the mortality 600 events, histopathology and viral loads in the tissues suggest that ISKNV was a major cause of 601 mortalities during the outbreaks. In general, the results support continued efforts to improve 602 the biosecurity of the industry in Ghana. There is a clear need to strengthen domestic

603 capability to rapidly diagnose and control emerging disease threats caused by ISKNV and 604 other pathogens. Further work is also needed to map the distribution of the virus and its 605 impact, including potential effects on wild fish species, and to implement practical control 606 strategies.

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