



Journal of the Hellenic Veterinary Medical Society

Vol 75, No 2 (2024)



To cite this article:

Kumru, S., Dengiz Balta, Z., Aliu, H., & Balta, F. (2024). Identification of Enterococcus Species Isolated from Commercial Fish Feeds and Infected Fish Specimens. *Journal of the Hellenic Veterinary Medical Society*, *75*(2), 7523–7530. https://doi.org/10.12681/jhvms.35199

Research article Ερευνητικό άρθρο

Identification of Enterococcus Species Isolated from Commercial Fish Feeds and Infected Fish Specimens

S. Kumru^{1*}^(b), Z. Dengiz Balta²^(b), H. Aliu³^(b), F. Balta⁴^(b)

¹Recep Tayyip Erdogan University, Faculty of Fisheries, Rize 53020, Turkey

²Recep Tayyip Erdogan University, Faculty of Fisheries, Rize 53020, Turkey

³University of Pristina "Hasan Prishtina", Faculty of Agriculture and Veterinary, Pristina, Kosovo

⁴Recep Tayyip Erdogan University, Faculty of Fisheries, Rize 53020, Turkey

ABSTRACT: The genus of lactic acid bacteria Enterococci are Gram-positive cocci bacterium that can survive in different environmental conditions such as water, plants, and soil. They are also bacteriological signs of fecal contamination. In aquaculture facilities, *Enterococcus* species have appeared as one of the crucial opportunistic fish pathogens. Enterococcus-caused fish diseases have been reported in different fish species like yellow tail, turbot, and tilapia. Even though Enterococcus species are used as probiotics and are members of the gastrointestinal flora, they also have pathogenic potential to produce septicemia, wound infections, urinary tract infections, and others. In this study, we isolated bacterial strains from affected rainbow trout and trout feed specimens. Based on the API 20 strep test kit, they were determined as Enterococcus faecium. While fish-isolated samples had 74.4%-99.9% similarity to E. faecium, trout feed isolated samples had 98.4%-99.9% similarity to E. faecium. In order to identify the isolates of the trout feed, PCR was performed using universal 16S rRNA primers. Sequence results indicate that the samples were E. faecium and E. faecalis. The phylogenetic tree was constructed with other Enterococcus species of 16S rRNA, and our samples were located in the E. faecium and E. faecalis species. In conclusion, there may be contamination of Enterococcus with food or other factors. Enterococcus sp. strains are opportunistic microorganisms and cause pathogenicity when the host immunity weakens. Even though all samples with API 20 strep test kit were identified as E. faecium, they had the lower percentage similarity, so they may be E. faecalis and other Enterococcus species. Thus, further studies are needed to understand their probiotic and pathogenicity functions in aquaculture production.

Keywords: Enterococcus; Enterococcus faecalis; Enterococcus faecium; trout; fish feeds

Corresponding Author: Salih Kumru, Recep Tayyip Erdogan University, Faculty of Fisheries, Rize 53020, Turkey E-mail address: salih.kumru@erdogan.edu.tr

Date of initial submission: 09-08-2023 Date of acceptance: 15-03-2024

INTRODUCTION

The Enterococcus genus members are Gram-pos-L itive cocci bacteria that can survive anywhere in nature and be found in a variety of environments including soil, sediments, marine water, freshwater, and different plants. They are generally isolated from contaminated water via fecal wastes or sewage so they are commonly used as bacteriological indicators for fecal contamination. Enterococcus species are typically known members of the regular gastrointestinal flora of both humans and livestock as well. There is a new insight that free-living birds may be a vector and reservoir of Enterococcus species, which impact animal and human health (Kwit et al. 2023). Currently the genus comprises of 66 species and many undefined groups. Enterococcus faecalis is one of the most common species that is isolated primarily from clinical cases (Jha et al., 2005; Lebreton et al., 2014; Zaheer et al., 2020). E. faecalis causes disease in humans and animals like Streptococcus (Akter et al., 2023).

In aquaculture facilities, opportunistic fish pathogens have been recently recognized as the causative mediators for several epidemics. Enterococcus species have appeared as essential fish pathogens (Martins et al., 2008). Enterococcus species have been reported as pathogens from important fish species such as yellow tail (Seriola quinqueradiata) and turbot (Scophthalmus maximus) (Nieto et al., 1995). Moreover, E. faecalis were isolated from streptococcosis like infection in tilapia (Oreochromis niloticus) in Egypt, India, and Bangladesh (Akter et al., 2023). Enterococcus spp. were isolated from infected and healthy fish in Bangladesh (Rahman et al., 2017). On the other hand, it was reported that E. faecium was used as a possible probiotic for ornamental cichlid fish (Pterophyllum scalare) and it facilitates nutrient uptakes from the feed (Dias et al., 2019).

Antimicrobial-resistant bacterial species have been identified in fish farming systems and water (Novais et al., 2018). *Enterococci* are members of the highlighted water-isolated bacteria among resistant microorganisms (Lebreton et al., 2014). Even though *Enterococcus* species are used as probiotic and are members of the gastrointestinal flora, they can cause the pathogenicity like septicemia, wound infections, urinary tract infections, and others as opportunistic microorganisms when the host immunity weakens (Kwit et al., 2023). Currently, bacterial species have been identified with sequencing approach. In this study, bacterial species isolated from infected rainbow trout and commercial fish feed were identified as *E. faecalis* and *E. faecium* with sequencing and API test.

MATERIAL AND METHODS

Bacterial culture

From trout food, bacteria were isolated from five trout food samples including different sizes and brands. Three replicates were used from each sample and nearly 1-gram (gr) food was transferred to 10 milliliter (ml) distilled water into 15 ml centrifuge tubes. The centrifuge tubes were vortexed and then samples were streaked on tryptic soy agar (TSA) via a loop. Similarly, bacterial samples were isolated from 20 kidneys of affected rainbow trouts, exhibiting signs described in detail in section "Clinical Symptoms and Gross findings of infected fish". The fish weights were between 100 and 150 grams and the fish age was 6-7 months. The bacterial samples were streaked on the surface of TSA with the help of a loop. They were incubated for 1-3 days at 20 °C. Bacteria showing colony characteristics of Enterococcus members, which occur singly or settled in pairs, in short chains, or as small irregular clusters (de la Maza et al., 2020; Qamer et al., 2003), were transferred to tryptic soy broth (TSB), and were incubated for 1 day at 20 °C. Then, enrichment culture was streaked on TSA agar again to see pure colonies.

Bacterial identification tests

To identify the bacterial species, conventional biochemical tests were performed on all purified colonies motility test, Gram staining, catalase, and oxidase activity (Balta, 2016). Also, rapid API 20 strep (bioMérieux, France) kit systems were used for Gram-positive bacteria. For API test kits and bacterial hemolysis activity, bacteria were inoculated on blood agar supplemented with 5% sheep blood. They were incubated on blood agar for 24-48 hours at 25 °C. They were checked whether the isolates formed hemolysis on blood agar. Isolates grown on blood agar were inoculated conferring to API 20 strep test kit instructions and incubated for 24 and 48 hours in an incubator at 25 °C. API 20 strep test kits were evaluated by adding reagents after 24 hours. The results of the biochemical tests for isolates were evaluated by making the final reading at the end of 48 hours for sugar tests (Balta and Karatay, 2021).

DNA extraction and PCR

From determined colonies, enrichment overnight

culture was prepared, and then DNA extraction was performed using the previously described protocol (Dashti et al., 2009). Briefly, bacterial cells were gathered from 1 ml of TSB culture, and then the cells were washed with distilled water three times. The cells were suspended in 1 ml distilled water, and then the suspensions were heated at 99 °C for 10 minutes. Finally, the suspension was centrifuged at 10.000 rpm for 5 minutes, and the supernatants were used as DNA templates for PCR. The PCR was carried out using Tag PCR Master Mix kit (Qiagen, Germany) with a pair of universal 16S rRNA bacterial primers 27F (5'-AGAGTTTGATCCTGGCTCAG-3') and 1392R (5'-GGTTACCTTGTTACGACTT-3') (Srinivasan et al., 2015). The thermal cycling conditions were as follows: Initial denaturation at 95 °C for 5 minutes followed by 30 cycles of 95 °C for 1 minute, 60 °C for 1 minute, 72 °C for 1 minute, then final extension at 72 °C for 10 minutes. PCR products then underwent an electrophoresis step using 1% agarose gel to identify specific bands of nearly 1450 bp for 16S RNA.

DNA sequencing and Phylogenetic tree

PCR products were sequenced by a sequencing company using Sanger sequencing (Macrogen-Europe, Netherlands). The sequence results were trimmed using the Sequencher 5.4.6 (Gene Code Corporation). Then, the sequences were analyzed via BLAST searches against *Enterococcus* genomes on the National Center for Biotechnology Information (NCBI). A phylogenetic tree was made using BLAST pairwise alignments via the Neighbor-Joining method (Zhang et al., 2000).

RESULTS

Clinical Symptoms and Gross findings of infected fish

The fish were collected at the water outlet and pool sides, and they were swimming lethargically on the water surface. Darkening of the skin and exophthalmia were determined. In the autopsy of the fish, the liver was pale and the peritoneum and air sacs had petechial hemorrhages. The stomach was generally empty and filled with liquid, and the intestines were hemorrhagic and filled with yellow exudate. The symptoms of the external and internal organs of the diseased fish were similar to previous studies (Athanassopoulou and Roberts, 2004) and are shown in Figure 1.

Bacterial isolates

Based on the API test results, 11 E. faecium isolates were identified. 5 of them (1BYB, 1KYB, 9BYGD, 11YGD, and 12YGD) were isolated from trout food. Additionally, 6 of them (B368, B369, B370, B384, B387, and B388) were isolated from diseased fish. Bacterial isolates' positive and negative reactions on the API 20 strep kit are shown in Table 1. Also, based on the API test kit company, bacterial isolates similarity is given in Table 2. From trout food isolates, 11YGD has the highest similarity (99.9%) to E. faecium while 1KYB has the lowest similarity (96.4%) to E. faecium. On the other hand, from diseased fish isolates, B387 and B388 have the highest similarity (99.9%), and B368 has the lowest similarity (74.4%) to E. faecium. From samples, an assessment of negative and positive results of the API 20 strep test kit for B384 identification is given in Figure 2.



Figure 1. Darkening of the skin and exophthalmia (A), Liver pale, petechial hemorrhage on peritoneal membrane and air sac (B).

API 20	Bacteria isolates										
Strep	1BYB	1KYB	9BYGD	11YGD	12YGD	B368	B369	B370	B384	B387	B388
VP	-	+	+	+	+	+	+	+	+	+	+
HIP	+	+	+	+	+	+	-	+	+	-	-
ESC	+	+	+	+	+	+	+	+	+	+	+
PYRA	+	+	+	+	+	+	+	+	+	+	+
αGAL	-	+	+	+	+	+	+	+	+	+	+
βGUR	-	-	-	-	-	-	-	-	-	-	-
βGAL	+	+	+	+	+	+	+	+	+	+	+
PAL	-	-	-	-	+	+	+	+	+	-	-
LAP	+	-	+	+	+	+	+	+	+	+	+
ADH	+	+	+	+	+	+	+	+	+	+	+
RIB	+	+	+	+	+	+	+	-	+	-	-
ARA	+	+	+	+	+	-	-	+	+	+	+
MAN	+	+	+	+	+	+	+	-	+	+	+
SOR	-	+	+	-	+	+	+	+	+	-	-
LAC	+	+	+	+	+	+	-	+	+	+	+
TRE	+	+	+	+	+	+	+	-	+	+	+
INU	-	-	+	-	-	-	-	-	-	+	+
RAF	-	+	+	+	+	+	+	+	+	+	+
AMD	-	+	+	-	+	+	+	+	+	+	+
GLYG	-	-	+	-	-	-	-	-	-	-	-
βΗΕΜ	-	-	-	-	-	-	-	-	-	-	-

Table 1. Assessment of positive and negative results of API 20 strep to identify bacterial samples from food and diseased fish isolates.

VP: Pyruvate, HIP: Hippurate hydrolysis, ESC: Esculin, PYRA: Pyrrolidonyl arylamidase, α GAL: α -galactosidase, β GUR: β -glucuronidase, β GAL: Galactosidase, PAL: Alkaline phosphatase, LAP: Leucine arylamidase, DH: Arginine dihydrolase, RIB: Ribose, ARA: L-Arabinose MAN: Mannitol, SOR: Sorbito, LAC: Lactose, TRE: Trehalose, IN U: Inulin, RAF: Raffinose, AMD: Starch, GLYG: Glycogen, β HEM: Beta hemolysis.

Table 2. Bacterial samples and their percentage similarity against the company database.

The second sumption and then percentage community against the company and active							
Bacterial Samples	Sum of Positive	Matching Bacterial Species	Similarity (%)				
1BYB	6157510	Enterococcus faecium	98.4				
1KYB	7317751	Enterococcus faecium	96.4				
9BYGD	7357773	Enterococcus faecium	98.4				
11YGD	7357550	Enterococcus faecium	99.9				
12YGD	7377751	Enterococcus faecium	99.8				
B368	7373751	Enterococcus faecium	74.4				
B369	5373551	Enterococcus faecium	98.9				
B370	5373551	Enterococcus faecium	94.9				
B384	7377751	Enterococcus faecium	99.8				
B387	5355571	Enterococcus faecium	99.9				
B388	5355571	Enterococcus faecium	99.9				



Figure 2. Assessment of positive and negative results API 20 strep for sample B384.

Phylogenetic tree

Bacterial samples, isolated from trout food, were identified as *E. faecium* and *E. faecalis* according to 16S rRNA sequence similarity. Their percentage identity similarities against the NCBI database are listed in table 3. All samples were submitted to NCBI. From samples, 1BYB (accession#: OQ13113.1) and

1KYB (accession#: OQ131137.1) were submitted as *Enterococcus* sp.. Moreover, 12YGD (accession#: OQ132535.1), 11YGD (accession#: OQ132524.1), and 9BYGD (accession#: OQ131217.1) were submitted to NCBI as *E. faecalis*. To show the relationship between samples and other *Enterococcus* species, the phylogenetic tree is shown in Figure 3.

Table 3. Bacterial samples and their percentage similarity against the NCBI database.								
		E. faecium		E. faecalis				
	DSM 20477	ATCC 19434	LMG 11423	ATCC 19433	JCM 5803	LMG 7937	EGM 183	
1BYB	98.5	98.4	97.9					
1KYB	95.7	95.6	95.1					
9BYGD				99.9	99.6	99.7	99.9	
11YGD				98.9	98.5	98.7	99.0	
12YGD				99.0	98.7	98.8	99.1	



Figure 3. The phylogenetic tree was constructed with a 16S rRNA sequence of bacterial samples from feed and other *Enterococcus* species (* indicates bacterial samples used in this study).

DISCUSSION

Aquaculture is influenced by bacterial agents that cause economic losses and reduce the efficiency of production worldwide (Kotob et al., 2016; Zorrilla et al., 2003). Recently, some bacterial fish pathogens in aquaculture facilities have been determined as contributory agents for severe outbreaks. Enterococci are important opportunistic fish pathogens that affect the aquaculture industry (Martins et al., 2008). Enterococcus sp. was previously described in yellow tail (Seriola quinqueradiata) as a fish pathogen in Japan and was determined as enterococcal septicemia in turbot (Scophthalmus maximus) in Spain (Nieto et al., 1995). Then, E. faecalis was stated as a tilapia pathogen causing streptococcal infection in Thailand and Egypt (Petersen and Dalsgaard, 2003). It was reported that Enterococcus sp. was often isolated from infected and healthy fish in Bangladesh (Rahman et al., 2017). Moreover, it was reported that the pathogenic E. faecalis was isolated from a tilapia suffering from streptococcosis in Bangladesh (Akter et al., 2020). On the other hand, in red tilapia (Oreochromis hybrid), experimental infection of E. faecalis showed low pathogenicity in producing streptococcosis (Rizkiantino et al., 2021). E. faecalis and E. faecium species, isolated from trout food and diseased fish, might be pathogenic to trout, and this pathogenicity may have been transferred from food contamination.

In Brazil, multi-resistant and virulent Enterococcus spp. were isolated from fish farming environments, and it was suggested that multi-resistant may be related to environmental pollution and aquaculture may be a reservoir for virulent and resistant Enterococci (Araujo et al., 2021). E. faecium was also isolated from fish mucus (El Jeni et al., 2020), so it may be related to environmental contamination as well. On the other hand, in vitro and in vivo, studies on trout showed that E. faecalis and its enterocin had a protective effect against the fish pathogen Lactococcus garvieae. In aquaculture, the enterocin may have a potential for alternatives to antibiotics to control diseases (Banos et al., 2019). Similarly, it was reported that heat-killed E. faecalis had stimulatory effects on cell-mediated immunity in crucian carp (Carassius auratus) (Matsuura et al., 2017). Even though, it was stated E. faecalis had protective effects against the fish pathogens in previous studies, further studies are needed to explore these bacterial species' protective effects against other pathogens.

It was reported that E. faecium, which was isolated

from the gastrointestinal tract of tilapia, had quorum sensing potential so it had a protective effect in controlling Aeromonas hydrophila infection in goldfish (Carassius auratus) when it was used as a probiotic (Vadassery and Pillai, 2020). E. faecium was used as a probiotic for angelfish (Pterophyllum scalare). The result was an improvement in juvenile angelfish growth when supplemented with feed (Dias et al., 2019). Additionally, E. faecium supplemented diet improved the immunological response, growth performance, and disease resistance to A. hydrophila in Cirrhinus mrigala production (Tilwani et al., 2022). The microencapsulated and herbal hydrogel-based encapsulated feed with E. faecium increases the resistance of tilapia against Streptococcus iniae and Streptococcus agalactiae infection (Kahieshesfandiari et al., 2021; Nami et al., 2022; Suphoronski et al., 2021). It was determined that E. hirae was isolated from the intestine of juvenile seabass, and it had new potential as a probiotic against pathogenic vibriosis (Masduki et al., 2020). It is not clear that isolated bacterial species, E. faecalis and E. faecium, have probiotic features against other fish pathogens. They may have new potential as protective effects and probiotics for other fish pathogens.

The pathogenicity of enterococcal infection is not clearly understood, and the infection is horizontally transferred via direct contact with infected fish or contaminated fish food (Athanassopoulou and Roberts, 2004). Multidrug-resistant E. faecium was isolated from feed, trout tanks, and upstream samples. It was reported that feed was an additional contamination source in aquaculture production (Novais et al., 2018). Similarly, rainbow trout feed and background environments was suitable sources to isolate enterococci as probable probiotics for aquaculture (Araujo et al., 2015). Even though the isolation of enterococci from trout feed, tanks, and rearing environments was reported as contamination and a possible probiotic for sustainable aquaculture, we isolated enterococci from trout food and infected fish. Our results indicate that the identified enterococci species may originate from contaminated feed.

CONCLUSION

In conclusion, *Enterococcus* spp., *E. faecalis*, and *E. faecium* were isolated from infected trout and commercial fish food and were identified via 16S rRNA sequence and API strep kit test results. Based on the previous studies, *E. faecium* can be used as a beneficial probiotic supplement while *E. faecalis* may be

determined as a fish pathogen. In addition, *Enterococcus* species are mostly resistant to antimicrobial agents and they may cause pathogenicity when the host immunity weakens because of stress and other infections. There may be a contamination of farm water and food via free-living birds or others. All samples were determined as *E. faecium* via API 20 strep test kit but some of them, isolated from fish and food, had lower coverage so they may be fish pathogen *E*. *faecalis* or other *Enterococcus* species as well. Thus, further studies are needed to identify the *enterococci* species isolated from fish food and infected fish to improve our understanding of their pathogenicity or probiotic functions in aquaculture production.

CONFLICT OF INTEREST

The author state there are no conflicts of interest.

REFERENCES

- Akter T, Haque MN, Ehsan R, Paul SI, Foysal MJ, Tay ACY, Islam MT, Rahman MM (2023) Virulence and antibiotic-resistance genes in *Enterococcus faecalis* associated with streptococcosis disease in fish. Sci Rep-Uk 13.
- Akter T, Rahman MM, Tay ACY, Ehsan R, Islam MT (2020) Whole-Genome Sequence of Fish-Pathogenic *Enterococcus faecalis* Strain BFFF11. Microbiol Resour Announc 9.
- Araujo AJG, Grassotti TT, Frazzon APG (2021) Characterization of *Enterococcus spp.* isolated from a fish farming environment in southern Brazil. Braz J Biol 81, 954-961.
- Araujo C, Munoz-Atienza E, Hernandez PE, Herranz C, Cintas LM, Igrejas G, Poeta P (2015) Evaluation of *Enterococcus spp*. from rainbow trout (*Oncorhynchus mykiss*, Walbaum), feed, and rearing environment against fish pathogens. Foodborne Pathog Dis 12, 311-322.
- Athanassopoulou F, Roberts RJ (2004) Streptococcal infections of farmed fish. Journal of The Hellenic Veterinary Medical Society 55, 136-144.
- Balta F (2016) Phenotypic, Serotypic and Genetic Characterization and Antimicrobial Susceptibility Determination of *Vibrio anguillarum*, Isolated from Cultured Sea Bass (*Dicentrarchus labrax L.*, 1758) in the Southeast Black Sea, Turkey. Fresenius Environmental Bulletin 25, 4393-4400.
- Balta F, Karatay I (2021) Lactococcus garvieae Infection and Treatment Observed in Rainbow Trout (Oncorhynchus mykiss). Journal of Anatolian Environmental and Animal Sciences 6, 651-661.
- Banos A, Ariza JJ, Nunez C, Gil-Martinez L, Garcia-Lopez JD, Martinez-Bueno M, Valdivia E (2019) Effects of *Enterococcus faecalis* UGRA10 and the enterocin AS-48 against the fish pathogen *Lactococcus garvieae*. Studies in vitro and in vivo. Food Microbiol 77, 69-77.
- Dashti AA, Jadaon MM, Abdulsamad AM, Dashti HM (2009) Heat Treatment of Bacteria: A Simple Method of DNA Extraction for Molecular Techniques. Kuwait Med J 41, 117-122.
- de la Maza L, Pezzlo M, Bittencourt C, Peterson E (2020) *Enterococcus*, In: Color Atlas of Medical Bacteriology, 3 ed. American Society for Microbiology, USA, pp. 24-29.
- Dias JAR, Abe HA, Sousa NC, Silva RDF, Cordeiro CAM, Gomes GFE, Ready JS, Mourino JLP, Martins ML, Carneiro PCF, Maria AN, Fujimoto RY (2019) *Enterococcus faecium* as potential probiotic for ornamental neotropical cichlid fish, Pterophyllum scalare (Schultze, 1823). Aquacult Int 27, 463-474.
- El Jeni R, Ghedira K, El Bour M, Abdelhak S, Benkahla A, Bouhaouala-Zahar B (2020) High-quality genome sequence assembly of R.A73 *Enterococcus faecium* isolated from freshwater fish mucus. BMC Microbiol 20, 322.
- Jha AK, Bais HP, Vivanco JM (2005) *Enterococcus faecalis* mammalian virulence-related factors exhibit potent pathogenicity in the Arabidopsis thaliana plant model. Infect Immun 73, 464-475.
- Kahieshesfandiari M, Nami Y, Lornezhad G, Kiani A, Javanmard A, Jaymand M, Haghshenas B (2021) Herbal hydrogel-based encapsulated *Enterococcus faecium* ABRIINW.N7 improves the resistance of red hybrid tilapia against *Streptococcus iniae*. J Appl Microbiol 131,

2516-2527.

- Kotob MH, Menanteau-Ledouble S, Kumar G, Abdelzaher M, El-Matbouli M (2016) The impact of co-infections on fish: a review. Vet Res 47, 98.
- Kwit R, Zajac M, Smialowska-Weglinska A, Skarzynska M, Bomba A, Lalak A, Skrzypiec E, Wojdat D, Koza W, Mikos-Wojewoda E, Pasim P, Skora M, Polak M, Wiacek, J, Wasyl D (2023) Prevalence of *Enterococcus spp.* and the Whole-Genome Characteristics of *Enterococcus faecium* and *Enterococcus faecalis* Strains Isolated from Free-Living Birds in Poland. Pathogens 12.
- Lebreton F, Willems RJL, Gilmore MS (2014) *Enterococcus* Diversity, Origins in Nature, and Gut Colonization, In: Enterococci: From Commensals to Leading Causes of Drug Resistant Infection, Boston.
- Martins ML, Mourino JLP, Amaral GV, Vieira FN, Dotta G, Jatoba AMB, Pedrotti FS, Jeronimo GT, Buglione-Neto CC, Pereira G (2008) Haematological changes in Nile tilapia experimentally infected with *Enterococcus sp.* Braz J Biol 68, 657-661.
- Masduki F, Y JM, Min CC, Karim M (2020) Characterization of *Entero*coccus hirae Isolated from the Intestine of Seabass (*Lates Calcarifer*) as a New Potential Probiotic against Pathogenic Vibrios. Curr Microbiol 77, 3962-3968.
- Matsuura Y, Takasaki M, Miyazawa R, Nakanishi T (2017) Stimulatory effects of heat-killed *Enterococcus faecalis* on cell-mediated immunity in fish. Dev Comp Immunol 74, 1-9.
- Nami Y, Kahieshesfandiari M, Lornezhad G, Kiani A, Elieh-Ali-Komi D, Jafari M, Jaymand M, Haghshenas B (2022) Administration of microencapsulated *Enterococcus faecium* ABRIINW.N7 with fructo-oligosaccharides and fenugreek on the mortality of tilapia challenged with *Streptococcus agalactiae*. Front Vet Sci 9, 938380.
- Nieto JM, Devesa S, Quiroga I, Toranzo AE (1995) Pathology of *Enterococcus sp.* Infection in Farmed Turbot, *Scophthalmus-Maximus* L. Journal of Fish Diseases 18, 21-30.
- Novais C, Campos J, Freitas AR, Barros M, Silveira E, Coque TM, Antunes P, Peixe L (2018) Water supply and feed as sources of antimicrobial-resistant *Enterococcus spp.* in aquacultures of rainbow trout (*Oncorhyncus mykiss*), Portugal. Sci Total Environ 625, 1102-1112.
- Petersen A, Dalsgaard A (2003) Antimicrobial resistance of intestinal *Aeromonas spp.* and *Enterococcus spp.* in fish cultured in integrated broiler-fish farms in Thailand. Aquaculture 219, 71-82.
- Qamer S, Sandoe JA, Kerr KG (2003) Use of colony morphology to distinguish different enterococcal strains and species in mixed culture from clinical specimens. J Clin Microbiol 41, 2644-2646.
- Rahman M, Rahman MM, Deb SC, Alam MS, Alam MJ, Islam MT (2017) Molecular Identification of Multiple Antibiotic Resistant Fish Pathogenic *Enterococcus faecalis* and their Control by Medicinal Herbs. Sci Rep-Uk 7.
- Rizkiantino R, Pasaribu FH, Soejoedono RD, Purnama S, Wibowo DB, Wibawan IWT (2021) Experimental infection of Enterococcus faecalis in red tilapia (Oreochromis hybrid) revealed low pathogenicity to cause streptococcosis. Open Vet J 11, 309-318.

Srinivasan R, Karaoz U, Volegova M, MacKichan J, Kato-Maeda M,

Miller S, Nadarajan R, Brodie EL, Lynch SV (2015) Use of 16S rRNA gene for identification of a broad range of clinically relevant bacterial pathogens. PLoS One 10, e0117617.

- Suphoronski SA, de Souza FP, Chideroli RT, Mantovani Favero L, Ferrari NA, Ziemniczak HM, Goncalves DD, Lopera Barrero NM, Pereira UP (2021) Effect of *Enterococcus faecium* as a Water and/or Feed Additive on the Gut Microbiota, Hematologic and Immunological Parameters, and Resistance Against Francisellosis and Streptococcosis in Nile Tilapia (*Oreochromis niloticus*). Front Microbiol 12, 743957.
- Tilwani YM, Sivagnanavelmurugan M, Lakra AK, Jha N, Arul V (2022) Enhancement of growth, innate immunity, and disease resistance by probiotic *Enterococcus faecium* MC-5 against *Aeromonas hydrophila* in Indian major carp, *Cirrhinus mrigala*. Vet Immunol Immunopathol 253, 110503.
- Vadassery DH, Pillai D (2020) Quorum quenching potential of *Enterococ*cus faecium QQ12 isolated from gastrointestinal tract of *Oreochromis*

niloticus and its application as a probiotic for the control of *Aero-monas hydrophila* infection in goldfish, *Carassius auratus* (Linnaeus 1758). Braz J Microbiol 51, 1333-1343.

- Zaheer R, Cook SR, Barbieri R, Goji N, Cameron A, Petkau A, Polo RO, Tymensen L, Stamm C, Song J, Hannon S, Jones T, Church D, Booker CW, Amoako K, Van Domselaar G, Read RR, McAllister TA (2020) Surveillance of *Enterococcus spp.* reveals distinct species and antimicrobial resistance diversity across a One-Health continuum. Sci Rep 10, 3937.
- Zhang Z, Schwartz S, Wagner L, Miller W (2000) A greedy algorithm for aligning DNA sequences. J Comput Biol 7, 203-214.
- Zorrilla I, Chabrillon M, Arijo S, Diaz-Rosales P, Martinez-Manzanares E, Balebona MC, Morinigo MA (2003) Bacteria recovered from diseased cultured gilthead sea bream (*Sparus aurata L.*) in southwestern Spain. Aquaculture 218, 11-20.