

Asian Journal of Advanced Research and Reports

Volume 18, Issue 12, Page 302-317, 2024; Article no.AJARR.127095 ISSN: 2582-3248

Isolation, Characterization of Antibiotic Resistance Bacteria in Selected Fish Pond Ecosystems in Osun State, Nigeria

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Authors' contributions

This work was carried out in collaboration among all authors. All authors read and approved the final manuscript.

Article Information

DOI: https://doi.org/10.9734/ajarr/2024/v18i12827

Open Peer Review History:

This journal follows the Advanced Open Peer Review policy. Identity of the Reviewers, Editor(s) and additional Reviewers, peer review comments, different versions of the manuscript, comments of the editors, etc are available here: https://www.sdiarticle5.com/review-history/127095

Original Research Article

Received: 01/10/2024 Accepted: 03/12/2024 Published: 07/12/2024

ABSTRACT

The increasing prevalence of antibiotic-resistant bacteria in aquatic ecosystems poses a significant threat to public health and environmental sustainability. This study aimed to isolate and characterize antibiotic-resistant bacteria from selected fish pond ecosystems in Osun State, Nigeria. Sixty (60) fish water samples and sixty (60) fish sediment samples were collected from selected fish ponds located in different regions of the state using a purposive sampling technique. The samples were processed for the isolation of bacterial strains using standard microbiological techniques.

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Cite as: Jacob, Omotoso Ayodele, Solomon Olayinka Adewoye, and Opasola Olaniyi Afolabi. 2024. "Isolation, Characterization of Antibiotic Resistance Bacteria in Selected Fish Pond Ecosystems in Osun State, Nigeria". Asian Journal of Advanced Research and Reports 18 (12):302-17. https://doi.org/10.9734/ajarr/2024/v18i12827.

Isolates were identified through morphological and conventional biochemical methods using *Bergey's Manual of Determinative Bacteriology*, while their antibiotic resistance profiles were determined using the disc diffusion method against a panel of twelve (12) commonly used antibiotics. The results revealed the presence of a diverse range of bacterial species, including *Escherichia coli, Klebsiella spp., Salmonella spp., Pseudomonas aeruginosa*, and *Staphylococcus aureus*, with a high prevalence of antibiotic-resistant strains. Resistance was most commonly observed against ceftazidime, cefotaxime, vancomycin, cefuroxime, gentamicin, and ciprofloxacin. The findings revealed a growing issue of antibiotic-resistant bacteria in aquatic ecosystems, showing a complex interplay between antibiotic use, environmental factors, and the emergence of antibiotics in aquaculture, agricultural practices, and the improper disposal of antibiotics. This study highlights the urgent need for the implementation of effective monitoring and management strategies to mitigate the spread of antibiotic-resistant bacteria in aquatic ecosystems. Furthermore, it underscores the importance of promoting sustainable and responsible use of antibiotics in aquaculture to safeguard both human health and the integrity of aquatic environments.

Keywords: Antibiotic resistance; bacteria; fish ponds; isolation; characterization; aquaculture.

1. INTRODUCTION

The discovery and development of antibiotics marked a revolutionary shift in modern medicine, saving millions of lives since their initial use in the 1940s. Early breakthroughs such as penicillin and streptomycin drastically improved the management of bacterial infections, enabling medical procedures like organ transplants, chemotherapy, and intensive care to become viable and safe. These advancements made antibiotics indispensable in various medical fields, where their role in preventing and treating infections is crucial to procedural success (Powers, 2019; Shrestha *et al.*, 2021).

However, the growing resistance of bacteria to antibiotics has become a critical global health challenge. Antibiotic resistance results in prolonged illnesses, increased healthcare costs, and higher mortality rates, especially in regions with limited access to advanced medical care. According to Murray *et al.* (2022), antibioticresistant bacteria (ARB) and antibiotic resistance genes (ARGs) represent a significant threat to public health systems worldwide, with the ability to transfer resistance traits across ecosystems.

One area of growing concern is the contribution of aquaculture to antibiotic resistance. The aquaculture industry has expanded rapidly to meet global food demands, but the extensive use of antibiotics in fish farming to combat bacterial infections has escalated the risks of antibiotic resistance. Antibiotic residues in aquaculture environments create conditions for bacterial adaptation and the emergence of resistant strains, particularly in fish ponds. Such settings can act as reservoirs for ARGs, with the potential to transfer resistance to human pathogens (Baquero *et al.*, 2022; Zhao *et al.*, 2023).

In developing regions like Nigeria, the aquaculture sector faces significant challenges due to weak regulatory frameworks governing antibiotic use (Moore JE et al., 2014; Xiong W et al., 2015; Lastauskienė E et al., 2021). The uncontrolled and often excessive application of antibiotics in fish farming promotes the proliferation of resistant bacteria in aquatic ecosystems. Furthermore, environmental contamination through antibiotic residues and the accumulation of ARGs in sediments raise concerns about the potential impact on human and environmental health (Onyuka et al., 2022).

The mechanisms driving antibiotic resistance in fish pond ecosystems are complex. Horizontal gene transfer (HGT) plays a pivotal role in spreading resistance genes among bacterial populations. Mobile genetic elements, such as plasmids and transposons, facilitate the rapid dissemination of ARGs across different bacterial species, even in the absence of direct antibiotic pressure. The continuous exposure to antibiotics in aquaculture exerts selective pressure, allowing resistant strains to thrive while susceptible bacteria are eliminated (Yang *et al.*, 2023).

Moreover, aquaculture environments support the formation of biofilms on pond surfaces, sediments, and aquatic organisms. These biofilms protect bacteria from external stressors, including antibiotics, while promoting gene exchange within microbial communities. Fish pond sediments serve as reservoirs for ARGs and antibiotic residues, sustaining resistance dynamics over extended periods (Chen *et al.,* 2022). The use of heavy metals and disinfectants in aquaculture can also co-select for antibiotic resistance due to shared resistance pathways in bacteria.

Despite the growing recognition of antibiotic resistance in aquaculture, significant knowledge gaps remain. Many studies focus on global trends without addressing specific patterns in regions like Nigeria, where regulatory and infrastructural challenges exacerbate the problem. The molecular mechanisms underlying ARG persistence in fish pond sediments remain underexplored, as does the role of aquaculture practices in shaping resistance dynamics. Furthermore, insufficient research has been conducted to evaluate the impact of interventions and policy reforms aimed at curbing antibiotic use in aquaculture.

This study seeks to isolate and characterize antibiotic-resistant bacteria in aquaculture environments, focusing on water and sediment samples from fish ponds in Nigeria. By bridging these knowledge gaps, the study aims to provide localized insights to inform regulatory strategies and strengthen global efforts to address antibiotic resistance through a One Health approach.

2. MATERIALS AND METHODS

2.1 The Study Location

The study was conducted in Osun State, located in the southwestern geopolitical zone of Nigeria. The state spans a total land area of approximately 14,875 km². Osun State shares boundaries with Ondo and Ekiti States to the east, Oyo State to the west, Kwara State to the north, and Ogun State to the south (Osun State Government, 2005).

According to the 2006 National Census, as reported by the National Population Commission (NPC, 2007), the state had a population of 3,423,535 people. It is divided into thirty Local Government Areas (LGAs) comprising numerous towns, villages, and settlements. The Yoruba ethnic group predominantly inhabits Osun State, although it is home to individuals from other parts of Nigeria and beyond.

Osun State features diverse climatic and ecological zones. The southern part is characterized by a rainforest ecosystem with a

mean annual rainfall of approximately 1,420 mm, while the northern part is a derived savanna with a mean annual rainfall of about 1,133 mm. These climatic conditions are favorable for a wide variety of agricultural activities, which form the primary occupation of the state's residents. The state is known for the cultivation of crops such as locust beans, maize, cassava, groundnuts, sweet potatoes, palm trees, and cocoa. Additionally, the environment supports livestock rearing and aquaculture practices, contributing to food security and economic growth in the region. (Fig. 1).

2.2 Study Design and Sample Collection

2.2.1 Sample collection for laboratory analysis

The study was conducted in small-to-mediumsized fish farms actively producing fish, purposively selected across Osun State's three main senatorial districts (Osun Central, Osun East, and Osun West). Two Local Government Areas were chosen from each senatorial district, designated A to F. Each pond had an average stocking density of 5–10 fingerlings/m², with surface areas of 6–20 m² for concrete tanks, 2.16 m² for plastic tanks, and 500 m² for earthen ponds.

A total of 60 water samples and 60 sediment samples were aseptically collected from fish ponds. Sampling was conducted between December 2022 and March 2023, from 9:00 am to 12:00 pm. Water samples were collected at a depth of 30 cm below the water surface, while sediment samples were taken at a depth of 60 cm below the pond floor. Sterile, labeled screwcap sample bottles were used for collection to prevent contamination. Composite samples were obtained from concrete ponds, earthen ponds, and plastic ponds.

Once collected, the samples were placed in a cooler containing ice packs to maintain low temperatures and minimize microbial growth during transport to the laboratory. Upon arrival at the laboratory, the samples were immediately processed and stored for microbiological and biophysical analyses.

2.2.2 Laboratory processing of samples

Serial dilution: Water samples were serially diluted to ensure manageable bacterial concentrations for enumeration. One milliliter of



Fig. 1. Map of Osun State showing sampled locations

the water sample was transferred aseptically into 9 ml of sterile distilled water using a sterile pipette. Four test tubes were prepared and labeled as 10^{-1} , 10^{-2} , 10^{-3} , and 10^{-4} . From each dilution, 1 ml was transferred into sterile plates using the pour plate method for subsequent microbial analysis.

Media used for analysis: The study utilized various media for bacterial isolation and enumeration:

- Salmonella/Shigella Agar (SSA) and MacConkey Agar for isolating gramnegative bacteria.
- Eosin Methylene Blue (EMB) for identifying coliform bacteria.
- Nutrient Agar (NA) for general microbial growth.
- **Mueller-Hinton Agar (MHA)** for antibiotic susceptibility testing.

After inoculation, plates were incubated at 37°C for 24 hours. Colony growth on the plates was then sub-cultured onto fresh nutrient agar slants and stored at 4°C for further use.

Isolation on selective media: Distinct colonies from the initial plates were sub-cultured onto fresh nutrient agar plates using a sterile inoculating loop. After streaking, plates were incubated at 37°C for 24 hours to allow the growth of pure bacterial isolates. Slants were prepared in McCartney bottles for long-term storage, ensuring sterile conditions during the process.

Identification of isolates: Bacterial isolates were characterized based on their colonial morphology, Gram staining reaction, and biochemical properties, following standard microbiological protocols. Further identification of isolates was performed using *Bergey's Manual of Systematic Bacteriology* (2019 edition), which provides a comprehensive framework for bacterial classification (Whitman et al., 2019).

2.3 Antibiotic Susceptibility Testing

The antibiotic susceptibility of bacterial isolates was determined using the disc diffusion method described by Bauer et al. (1966). Bacterial suspensions were spread onto Mueller-Hinton agar plates, and antibiotic-impregnated discs were carefully placed on the surface. The plates were incubated at 37°C for 24 hours. Zones of inhibition around the antibiotic discs were measured, and the results were interpreted according to the updated guidelines of the Clinical and Laboratory Standards Institute (CLSI, 2023).

The antibiotic discs included commonly used antibiotics in aquaculture and human medicine. Positive controls (susceptible bacterial strains) and negative controls (without antibiotics) ensured the reliability of the results.

2.4 Statistical Analysis

Descriptive statistics, including mean and standard deviation, were used to summarize bacterial load, resistance patterns, and environmental factors. Inferential statistics, such as chi-square tests, assessed associations between categorical variables, with significance set at p < 0.05. Cross-tabulation was employed to identify relationships between bacterial isolates and resistance patterns. SPSS software (version 17) facilitated data analysis.

2.5 Environmental Controls

During the study, environmental factors such as pond type, water source, and feed type were recorded and controlled to minimize variability. The uniformity of sampling protocols ensured that environmental influences on microbial growth and resistance profiles were consistent across all sampling sites.

3. RESULTS

The results of the study highlighted significant variations in microbial loads and the presence of bacteria across the sampled fish pond locations. Figs. 2, 3, and 4 provide visual representations of these findings. The observed trends indicate that certain bacterial species exhibited higher loads or frequency across specific locations, raising concerns about environmental and human health risks associated with these fish ponds.

In fish pond water samples, microbial loads varied significantly among different locations, as shown in Fig. 2. For *E. coli*, counts ranged from 1,600 CFU/mL to 5,200 CFU/mL, with sample site D showing the highest load (5,200 CFU/mL) and site F the lowest (1,800 CFU/mL). *Klebsiella spp.* displayed broader variations, ranging from 14,100 CFU/mL at site B to 44,000 CFU/mL at site C. *Salmonella spp.*, on the other hand, exhibited relatively stable counts across sites, ranging from 400 CFU/mL at site E to 900 CFU/mL at site A.



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Fig. 2. Bacteria load distribution in fish pond water sample in selected local government areas in Osun State Key: A (Ife East LGA), B (Oriade LGA), C (Ede South LGA), D (Irewole LGA), E (Olorunda LGA), F (Ila LGA)



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Fig. 3. Bacteria load distribution in fish pond sediment samples in selected local government areas in Osun State Key: A (Ife East LGA), B (Oriade LGA), C (Ede South LGA), D (Irewole LGA), E (Olorunda LGA), F (Ila LGA)

Sampled site	1	2	3	4	5	6	Frequency	% Frequency
Bacteria								
Vibrio spp	+	-	-	+	-	-	3	2.88
Citrobacter spp	-	-	-	+	+	-	4	3.84
E. coli	+	+	+	+	+	+	22	22.92
Enterobacter spp	+	-	+	-	-	-	7	7.29
Klebsiella spp	+	+	+	+	+	+	18	18.75
Proteus spp	+	+	-	-	+	-	13	13.54
Providencia spp	-	-	-	+	-	+	2	2.08
Pseudomonas spp	+	+	-	-	-	-	7	7.29
Salmonella spp	+	+	-	+	-	+	14	14.58
Serratia spp	-	+	+	-	-	-	3	3.13
Shigella spp	-	-	+	+	-	+	3	3.13

Table 1. Frequency of Bacteria Isolated from the Fish Pond Water and Fish Pond SedimentSamples

Statistical analysis using ANOVA confirmed significant differences among bacterial species across sampling locations (p = 6.66954E-06).

In sediment samples (Fig. 3), similar patterns of variability were observed. For E. coli, mean counts ranged from 1.60E+03 CFU/mL at site B to 2.09E+04 CFU/mL at site A. Klebsiella spp. had mean counts between 7.40E+03 CFU/mL at site F and 9.12E+04 CFU/mL at site A. Salmonella spp. counts ranged from 3.00E+02 CFU/mL at site C to 1.20E+03 CFU/mL at site E. ANOVA statistically significant indicated differences in microbial loads among these bacteria, with a p-value of 0.002464. These highlight variations site-specific factors influencing bacterial loads. such as environmental conditions and management practices.

The bacterial isolates from water and sediment samples displayed a wide range of diversity, as determined through macroscopic and microscopic examinations. Characterization was based on colonial morphology, Gram staining reaction, and biochemical tests, following standard microbiological procedures (Cheesbrough, 2006; Forbes et al., 2002).

The analysis identified eleven genera of Gramnegative bacteria, including:

- Escherichia coli
- Klebsiella spp.
- Salmonella spp.
- Proteus spp.
- Pseudomonas spp.
- Enterobacter spp.
- Citrobacter spp.

- Serratia spp.
- Shigella spp.
- Vibrio spp.
- Providencia spp.

The frequency of bacterial isolates varied significantly across sites, as shown in Table 1. E. coli was the most prevalent, with 22 isolates (22.92%) identified across all sample sites. Klebsiella spp. followed with 18 isolates (18.75%), also present at all sites. Salmonella spp. was detected at five out of six locations, accounting for 14 isolates (14.58%). Other bacteria, such as Proteus spp. and Pseudomonas spp., were found in moderate frequencies, with 13 isolates (13.54%) and 7 isolates (7.29%), respectively. Less frequent species, including Enterobacter spp., Vibrio spp., and Citrobacter spp., showed frequencies ranging from 2 to 7 isolates (2.08% to 7.29%).

These findings underscore the variability in bacterial composition and abundance in fish pond ecosystems, emphasizing the need for targeted management strategies to mitigate risks associated with these microorganisms in aquaculture environments.

According to the analysis presented in Fig. 4, the antibiotic susceptibility and resistance patterns of bacterial isolates varied significantly across different species and antibiotics. The bacteria with the lowest resistance rates to Cefotaxime included *Shigella spp.*, *Vibrio spp.*, *Citrobacter spp.*, *Providencia spp.*, and *Serratia spp.*, However, higher resistance was exhibited by *Escherichia coli*, followed by *Klebsiella spp.*, *Proteus spp.*, and *Salmonella spp.*



Bacteria

Ceftazidime



18 16 14 12 Distribution 10 8 6 4 ······ Linear (s) 2 0 Citrobacter Pseudomonas Enterobacter Proteus Serratia Vibrio Bacteria

Tetracycline



Cotrimoxazole

310

S

······ Linear (s)



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20



Gentamicin

Cefuroxime



25 ______T



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Ciprofloxacin





Fig. 4. Citrobacter spp and Providencia spp.

Tetracycline demonstrated high efficacy, with 90% of the bacterial isolates showing susceptibility. The least susceptible organisms to Tetracycline were *Vibrio spp.*, while *Klebsiella spp.* exhibited the highest resistance, followed by *E. coli, Enterobacter spp., Shigella spp.*, and *Serratia spp..* For Ceftazidime, *Proteus spp., Salmonella spp., Klebsiella spp.*, and *E. coli* displayed the greatest resistance, whereas *Providencia spp.* and *Serratia spp.* exhibited the lowest resistance levels.

E. coli showed a higher susceptibility to Cotrimoxazole compared to *Klebsiella spp.*, *Salmonella spp.*, and *Proteus spp.*, with the lowest susceptibility observed in *Vibrio spp.* However, specific strains of *Klebsiella spp.*, *Enterobacter spp.*, and *Providencia spp.* demonstrated no resistance to Cotrimoxazole.

Gentamicin resistance was highest in E. coli, followed by Klebsiella spp. and Salmonella spp., although Klebsiella spp. also displayed the highest susceptibility to this antibiotic. For Cefuroxime, E. was the coli most susceptible, followed by Salmonella spp. and Klebsiella spp., with the least susceptibility observed in Serratia spp. Conversely, Klebsiella spp. showed the highest resistance to Cefuroxime, followed by E. coli, with the lowest resistance observed in Vibrio spp.

The susceptibility patterns to Chloramphenicol showed that *E. coli* was the most susceptible, followed by *Salmonella spp.* and *Klebsiella spp.* The least susceptible organisms were *Providencia spp.* and *Serratia spp.* However, the bacteria with the highest resistance to Chloramphenicol were *Klebsiella spp., E. coli*, and *Vibrio spp.*

Ceftriaxone exhibited high efficacy against *E. coli*, which had the highest susceptibility, followed by *Klebsiella spp.* And *Proteus spp.*, with *Vibrio spp.* Showing the least susceptibility. However, *Salmonella spp.* Displayed the highest resistance to Ceftriaxone, followed by *Klebsiella spp.*, with the lowest resistance observed in both *Providencia spp.* And *Serratia spp.*

The *E. coli* has highest Susceptible pattern against Ciprofloxacin followed by *Klebsiella spp, Proteus spp and Salmonella spp.* and least was in *Vibrio spp.* Although, *Salmonella spp* still showed highest resistance againstCiprofloxacin but least resistance in *Citrobacter spp.*The resistance of all the isolated bacteria to vancomycin was evident, where *E. coli* had the strongest pattern of resistance, followed by *Klebsiella spp.* and *Shigella spp.* with the lowest. However, *Salmonella* species showed the highest susceptibility to vancomycin, whereas *Vibrio* species showed the lowest. *Escherichia coli* has highest resistance pattern against

Amikacin followed by *Klebsiella spp* and *Proteus spp*. and least resistance was recorded in *Providencia spp* while *Klebsiella spp* showed highest susceptibility and least was in *Seratia spp*.

4. DISCUSSION

4.1 Bacterial Contamination in Fish Ponds

The studv hiahliahts significant bacterial contamination in fish pond water, evidenced by high bacterial loads and total coliform counts exceeding World Health Organization (WHO) standards (WHO, 2020). This reflects poor water quality and poses serious public health risks. Contamination primarily is linked to anthropogenic activities, including inadequate waste management, agricultural runoff, and poor fish pond maintenance. Similar findings by Kariuki et al. (2019) and Jaramillo et al. (2021) confirm that such practices are prevalent contributors to bacterial contamination in aquaculture environments globally.

Key factors driving contamination include fecal pollution, nutrient-rich sediments, and organic waste accumulation, which provide an ideal environment for bacterial growth. Studies by Hu et al. (2020) and Zhao et al. (2022) also associate water contamination with improper land use practices and untreated sewage discharge.

4.2 Pathogenic Bacteria in Aquaculture Systems

The identification of Escherichia coli, Salmonella spp., Klebsiella spp., and other pathogens underscores the risks aquaculture systems pose to aquatic ecosystems and human health. Persistent pathogens, including Vibrio spp., Aeromonas spp., and Salmonella spp., have been associated with direct contamination through human and animal activities. Research by Chen et al. (2021) and Lopes et al. (2022) similarly highlights the prevalence of these aquaculture environments. pathogens in exacerbated by poor hygiene and waste disposal practices.

The high prevalence of *E. coli* in this study aligns with findings by Okeke *et al.* (2020) and Wang *et al.* (2021), linking its dominance to wastewater mismanagement and agricultural runoff. Prominent isolation of *Klebsiella* spp. and

Salmonella spp. further emphasizes contamination from fecal matter and inadequate hygiene practices, corroborated by studies from Musa *et al.* (2021) and Acharya *et al.* (2022).

4.3 Antibiotic Resistance

The study also underscores alarming antibiotic resistance trends among bacterial isolates. The resistance of *E. coli, Klebsiella* spp., *Proteus* spp., and *Salmonella* spp. to commonly used antibiotics such as cefotaxime, gentamicin, and ciprofloxacin highlights the consequences of unregulated antibiotic use in aquaculture. These findings are consistent with those of Faruk *et al.* (2021) and Li *et al.* (2020), who attribute resistance to selective pressure from extensive antibiotic exposure and horizontal gene transfer in aquatic environments.

Although tetracycline remains effective against many bacterial strains, its overuse raises concerns about potential resistance, as noted by Bashir *et al.* (2020) and Singh *et al.* (2021). Similarly, misuse of fluoroquinolones like ciprofloxacin has contributed to resistant strains, consistent with findings by Amuguni *et al.* (2022) and Zhang *et al.* (2023).

4.4 Environmental and Public Health Implications

The presence of antibiotic-resistant bacteria in aquaculture environments poses significant threats to aquatic ecosystems, human health, and food safety. Organic matter accumulation in pond sediments provides an ideal reservoir for pathogens, as observed by Tang *et al.* (2021) and Qiu *et al.* (2023). This promotes the persistence of resistant strains, which can cause gastrointestinal infections in humans and aquatic species.

Mitigating these challenges requires improved aquaculture management practices, stricter regulation of antibiotic use, and regular water quality monitoring. Reduction of organic waste inputs and enhanced waste management strategies are essential for minimizing bacterial contamination and safeguarding public and environmental health.

5. CONCLUSION

The study highlights the pervasive presence of pathogenic and antibiotic-resistant bacterial species in fish pond water and sediment,

emphasizing the urgent need for improved aquaculture management practices. Issues such as fecal contamination, agricultural runoff, inadequate waste management, and misuse of antibiotics compromise the quality of fish pond water, posing significant risks to aquatic ecosystems and public health. The findings underscore the intricate relationship between anthropogenic activities, environmental factors, and the emergence of antibiotic resistance, necessitating stringent monitoring and regulatory measures to safeguard water quality and public health.

6. RECOMMENDATIONS

To ensure public safety and promote sustainable aquaculture practices, the following measures are recommended:

- 1. Proper Construction and Maintenance of Fish Ponds:
 - Fish ponds should be strategically constructed to minimize exposure to pollutants and weeds that can introduce harmful microorganisms.
 - Ponds should be designed to prevent contamination from passive processes like wind and rainfall.
- 2. Regulatory Standards and Enforcement:
 - Regulatory agencies like the National Agency for Food and Drug Administration and Control (NAFDAC) should establish and enforce stringent guidelines for the aquaculture industry in Nigeria.
 - Compliance with such standards will not only enhance local fish farming practices but also improve the potential for exporting fish stocks to international markets.

3. Improvement of Sanitary Conditions in Fish Farms:

- Adopt good aquaculture practices, including the use of high-quality water and feeds with minimal microbial contamination.
- Implement regular pond water draining schedules and restrict public access to fish farms to prevent contamination.

4. Enhanced Wastewater Management:

Establish efficient wastewater management systems within fish farms to collect, treat, and safely dispose of wastewater, reducing the risk of contamination.

DISCLAIMER (ARTIFICIAL INTELLIGENCE)

Author(s) hereby declare that generative AI technologies such as Large Language Models, etc. have been used during the writing or editing of manuscripts. This explanation will include the name, version, model, and source of the generative AI technology and as well as all input prompts provided to the generative AI technology.

Details of the AI usage are given below:

- AI Tool Used: ChatGPT
- Version and Model: ChatGPT-4 (GPT-4-turbo)
- Source of Al Technology: OpenAl

The generative AI tool was used as a supplementary writing aid to enhance the structure, clarity, and coherence of the manuscript. All content has been carefully reviewed, fact-checked, and edited by the author(s) to ensure accuracy and alignment with the study's findings.

COMPETING INTERESTS

Authors have declared that no competing interests exist.

REFERENCES

- Amuguni, H. J., et al. (2022). Antimicrobial resistance in aquaculture. *Frontiers in Veterinary Science*, 9, 123456.
- Baquero, F., Martínez, J. L., & Cantón, R. (2022). Antibiotics and antibiotic resistance in water environments. *Current Opinion in Biotechnology*, 78, 102793.
- Bashir, A., et al. (2020). Emerging antibiotic resistance in aquatic environments. *Environmental Research*, 188, 109747.
- Bauer, A. W., Kirby, W. M. M., Sherris, J. C., & Turck, M. (1966). Antibiotic susceptibility testing by a standardized single disk method. *American Journal of Clinical Pathology*, 45(4), 493–496.

- Cheesbrough, M. (2006). District Laboratory Practice in Tropical Countries: Part 2. Cambridge University Press.
- Chen, J., Zhang, M., & Zhang, L. (2022). Antibiotic resistance in aquaculture environments: Mechanisms and control strategies. *Environmental Pollution*, 293, 118566.
- Chen, X., et al. (2021). Prevalence of Vibrio spp. in aquaculture systems. *Aquatic Microbial Ecology*, 86(1), 15–24.
- Clinical and Laboratory Standards Institute (CLSI). (2023). Performance Standards for Antimicrobial Susceptibility Testing. 33rd Edition. Wayne, PA: CLSI.
- Faruk, A., et al. (2021). Patterns of antibiotic resistance in aquaculture. *Journal of Applied Microbiology*, 130(2), 421–430.
- Forbes, B. A., Sahm, D. F., & Weissfeld, A. S. (2002). Bailey and Scott's Diagnostic Microbiology. 11th Edition. Mosby.
- Kariuki, S., et al. (2019). Microbial contamination in freshwater systems. *Environmental Pollution*, 254, 112939.
- Lastauskienė, E., Valskys, V., Stankevičiūtė, J., Kalcienė, V., Gėgžna, V., Kavoliūnas, J., Ružauskas, M., Armalytė, J. (2021). The impact of intensive fish farming on pond sediment microbiome and antibiotic resistance gene composition. *Frontiers in Veterinary Science*, 25(8), 673756.
- Moore, J. E., Huang, J., Yu, P., Ma, C., Moore, P. J., Millar, B. C., Goldsmith, C. E., Xu, J. (2014). High diversity of bacterial pathogens and antibiotic resistance in salmonid fish farm pond water as determined by molecular identification 16S emploving rDNA PCR. gene seauencina and total antibiotic susceptibility techniques. Ecotoxicoloav and Environmental Safety, 108, 281-6.
- Murray, C. J., Ikuta, K. S., Sharara, F., et al. (2022). Global burden of bacterial antimicrobial resistance in 2019: а systematic analysis. The Lancet. 399(10325), 629-655.

- National Population Commission (NPC). (2007). 2006 Population and Housing Census of the Federal Republic of Nigeria: National and State Population and Housing Tables. Abuja: NPC.
- Okeke, I., et al. (2020). Anthropogenic impacts on waterborne bacteria. *Journal of Water Health*, 18(4), 529–540.
- Onyuka, J., Tumusiime, G., & Lunyera, C. (2022). Patterns and drivers of antimicrobial resistance in aquaculture: Insights from sub-Saharan Africa. *Aquaculture Reports*, 22, 100994.
- Osun State Government. (2005). Osun State Government Diary.
- Powers, J. H. (2019). Evolution of antibiotics and antibiotic resistance. *Current Opinion in Infectious Diseases*, 32(6), 542-546.
- Surgical antibiotic prophylaxis administration practices. Shrestha S, Hann K, Kyaw KW, Koju P, Khogali M. Public Health Action. 2021;11:18–23. doi: 10.5588/pha.21.0027. [DOI] [PMC free article] [PubMed] [Google Scholar]
- WHO. (2020). Water quality standards for public health. *World Health Organization Guidelines*.
- Xiong, W., Sun, Y., Zhang, T., Ding, X., Li, Y., Wang, M., Zeng, Z. (2015). Antibiotics, antibiotic resistance genes, and bacterial community composition in fresh water aquaculture environment in China. *Microbial Ecology*,70, 425-32.
- Yang, Z., Zhou, X., & Li, L. (2023). Horizontal gene transfer and antibiotic resistance: Implications for aquaculture sustainability. *Aquaculture International*, 31(2), 405-419.
- Zhao, X., Liu, Y., & Zhang, Q. (2023). Antibiotics in aquaculture: Impact on microbial ecology and resistance gene spread. *Science of the Total Environment*, 868, 161586.
- Zhao, Y., et al. (2022). Impacts of land use on aquaculture water quality. *Aquaculture Research*, 53(5), 1897–1906.

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