

Research Article**Seasonal variation of bacteriological parameters of rui (*Labeo rohita*) and kalibaus (*Labeo calbasu*) collected from the Hakaluki Haor, Bangladesh**Hasan M¹, Hossain MM¹, Hasan MN¹, Rahman ASMS¹, Bormon AKA³, Hasan MA², Kashem MA^{1*}¹Department of Fisheries Technology and Quality Control, Sylhet Agricultural University, Sylhet, Bangladesh.²Faculty of Animal and Biomedical Sciences, Sylhet Agricultural University, Sylhet, Bangladesh.³Department of Aquatic Resource Management, Sylhet Agricultural University, Sylhet, Bangladesh**ABSTRACT****Article History**

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The fish *in situ* bacterial flora correlated with the aquatic environment. The higher initial bacterial flora in fish ultimately affects the storage life and quality of the products. The present study aimed to collect 36 samples of Rui (*Labeo rohita*) and Kalibaus (*Labeo calbasu*) from three locations in four seasons (pre-monsoon, monsoon, post-monsoon and winter) of Hakaluki haor to investigate their seasonal bacteriological changes. The samples were investigated for total bacterial load in term of Total Viable Count (TVC) and Total Coliform Count (TCC) as sanitary index. The prevalence of *E. coli*, *Salmonella* spp. and *Vibrio* spp were also investigated. The higher viable bacterial loads (logarithmic scale) were observed in Kalibaus than Rui in all study seasons. In Rui, the bacterial load was 5.56 ± 0.078 , 5.51 ± 0.043 , 5.36 ± 0.065 , and 5.48 ± 0.091 (CFU/g \pm SD) whereas in Kalibaus, those were 5.68 ± 0.050 , 5.58 ± 0.057 , 5.52 ± 0.040 , and 5.63 ± 0.067 in monsoon, post-monsoon, winter, and pre-monsoon, respectively. For total coliform (TCC), Rui were 73, 72, 28, and 48 MPN/g, while in Kalibaus, the values were 78, 96, 65, and 86 MPN/g in monsoon, post-monsoon, winter, and pre-monsoon, respectively. *E. coli*, *Salmonella* spp., and *Vibrio* spp. were also present in both fish at each season. The results indicate that Rui had a lower prevalence of pathogenic bacteria than Kalibaus. The occurrence of *E. coli* and *Salmonella* spp. in both fish samples indicates fecal contamination, which may be linked to unhygienic environmental conditions.

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INTRODUCTION

Bangladesh is a riverine country. Fish and fisheries are closely related to the Bangladeshi people for their contribution to providing animal proteins at low prices as well as income. It is also well-praised for providing sources of foreign currency (Hossain, 2015; Azim *et al.*, 2003; Alam and Thomson, 2001). "Haor" fishery contributes a considerable portion of the total fish production of inland captured fishery. Haors are bowl-shaped shallow depressions covering about 25% of the northeastern part of Bangladesh (Hossain and Nishat, 1989). In the case of fishery products, biological hazards like pathogenic bacteria and spoilage associated with pathogenic bacteria are important for food safety and human health (Herrera *et al.*, 2006). Due to their

higher moisture and protein content, fish are very susceptible to spoilage. The initial freshness of fish is primarily hampered by enzymatic and chemical reactions, whereas microbial activity is accelerated the spoilage after death of fish (Pal and Mahendra, 2015; Rhea, 2009; Huss, 1998). Basically, live fish is sterile. The spoilage from enzymes and bacteria starts when it commences to death. Bacteria grow rapidly after the death of fish and accelerate the spoilage of fish (Gram and Huss, 1996; McMeekin *et al.*, 1992). Contamination in the fishery product mainly occurs due to the bursting and splitting of fish intestines during handling, abject processing, or unhealthy washing. The quality of harvested fish is greatly influenced by the degree of adulteration by pathogenic and spoilage bacteria during handling or processing (Adebayo-Tayo *et al.*, 2012). It is

reported that fish contaminated with pathogenic bacteria from different sources, such as intestinal microflora of enteric bacteria of human or animal origin, are responsible for spoilages ([Geldreich and Clarke 1966](#)). Both fresh and spoiled fish and seafood have been reported with many spoilage bacteria and fungi. Among spoilage bacteria, *Aeromonas*, *Alcaligenes*, *Bacillus*, *Enterobacter*, *Enterococcus*, *Escherichia coli*, *Listeria*, *Pseudomonas*, and *Shewanella* are well documented, while fungi *Aspergillus*, *Candida*, *Cryptococcus*, and *Rhodotorula* are also isolated from both fresh and spoiled fish and other sea foods ([Church, 1998](#)). Other spoilage organisms are *Pseudomonas putrifaciens*, *Pseudomonas fluorescens*, *Bacillus*, *Moraxella* etc; and pathogenic bacteria are *Escherichia coli*, *Salmonella* spp., *Staphylococcus* spp., *Vibrio* spp., *Aeromonas* spp., *Clostridium botulinum*, *Listeria monocytogenes* ([Abraham *et al.*, 2010](#); [David *et al.*, 2009](#); [Ayulo *et al.*, 1994](#); [Hood *et al.*, 1983](#)). Food-borne infection and intoxication are caused by *Salmonella* spp., *Staphylococcus* spp., *Vibrio* spp., and *Aeromonas* spp, which may also transmitted by fish if contaminated with those pathogens ([Emikpe *et al.* 2011](#); [Gold and Salit, 1993](#)). Last few decades, Bangladesh has been exporting fish and fishery products and has earned foreign currencies. Hence, the fisheries and aquaculture sectors have been reported as the third most important contributors to export earnings ([Ghose, 2014](#)). In a few cases, exported products also were returned back due to the low microbiological quality (e.g., coliform, fecal coliform, *Pseudomonas*, *Staphylococcus*, etc.), which have caused some controversy about Bangladeshi fishery products ([Noor *et al.*, 2013](#)). As a consequence, importers are much more concerned about food safety in terms of the microbial quality of the products ([Pal and Mahendra, 2015](#)). To ensure the best fishery products and consumer acceptance, maintaining the quality of fish and fish products is essential ([Okoro *et al.*, 2010](#); [Begum *et al.*, 2010](#); [Prabakaran *et al.*, 2011](#); [Anbudhasan *et al.*, 2012](#)). Fish a highly perishable foodstuff. It requires care and attention in every case of handling until it reaches the consumption table ([Czerwińska *et al.*, 2008](#); [Poli *et al.*, 2005](#); [Clucas and Ward, 1996](#)). During handling, bacteria present on the body surface or in the intestine can enter the fish flesh and cause contamination ([Nerín *et al.*, 2016](#); [Chowdhury *et al.*, 2010](#); [Ivanek *et al.*, 2004](#)). Nowadays, the pollution of waterbodies and the possible occurrence of different pollutants (physical, chemical, and biological) is a matter of concern for public health ([Adedeji *et al.*, 2012](#); [Ababouch 2006](#)). The Hakaluki *haor* is situated in the Sylhet district of Bangladesh and is one of the prominent sources of fish. Though a huge amount of fish is collected from the *haor* area, there is no fish quality (physical or microbiological) monitoring in the *haor* area. In Bangladesh, studies about microbial contamination of fish and information on the bacteriological population are available, but information about the distribution of bacteria in fish from the open environment is patchy. Among the captured fishes from the Hakaluki *haor*, Rui and Kalibaus are two popular and available fish species among fish lover because of their taste and reasonable price. In this study, Rui and Kalibaus were collected from Hakaluki *haor* for comparative bacteriological assessment not only for their local demand and export market. It can give a clear idea about the microbial status of these fishes related to the *haor* environment in terms of food safety and health concerns.

MATERIALS AND METHODS

The research investigated the bacterial load and isolation of bacteria from Rui and Kalibaus of Hakaluki *haor* in the Sylhet region of Bangladesh.

Study area, sample, and planning

Pre-selected three fishing points of the Hakaluki *haor*, namely Ghilachora of Fenchuganj Upazila, Islamganj area of Kulaura Upazila, and Kanungo area of Barlekha Upazila were used to collect samples. For the bacteriological status assessment of Rui and Kalibaus, a total of 72 individual fish (36 for each) were collected from this area. These fish species of Hakaluki Haor were selected because these species are the most valuable and highly demandable for the domestic. Four seasons (winter, pre-monsoon, monsoon, and post-monsoon) were also considered for the comparative analysis of the bacterial status of the selected fish samples. During this study period, temperature was recorded on all dates of sample collection.

Preparation of sample

A muscle sample of 20g, including skin, was collected for both fish. The samples were blended and homogenized in sterile alkaline peptone water aseptically. The samples were dilution for plate counts using serial dilutions. Briefly, 1mL of the samples were diluted into 9ml sterile distilled water. Hence, it is a ten-fold of dilution. Subsequently, dilution was prepared serially until 10^{-6} according to protocol ([ISO, 1995](#)). Then, 100µl of samples were plated into an agar plate, incubated in an incubator for 24h, and counted the colonies for total viable counts.

Calculation of Total Viable Count (TVC)

A sample of 100µl was taken from each serial dilution and inoculated into a plate using spread plate techniques. The plates were incubated in an inverted position in an incubator at 37°C for 24h. After 24h, the plates having 30 to 300 colonies were considered for counting. No. of bacteria per gram of the sample (CFU/g) was calculated by using the following formula:

$$\text{CFU/g} = \frac{\text{No. of colonies on petridish} \times 10 \times \text{dilution factor} \times \text{Volume of total sample solution}}{\text{Wt. of fish sample (g)}}$$

Total Coliform Count (TCC)

The total coliform of a sample is measured by the MPN (Most Probable Number) method, which estimates the viable population density of microorganisms in the sample. It was calculated based on the method described by [Benson, 2002](#). Gas production is the fermentation tubes indicated its positive growth response. Interestingly, fewer positive cultures should be in such a manner that results from higher dilutions of the sample.

Isolation and identification of bacteria

Morphological characteristics such as size, shape, arrangement, and motility were investigated by Gram's

staining reaction. The colony characteristics, biochemical reaction, catalase test, and motility test were also done for further isolation and identification of the bacteria. Briefly, the pure colonies from the streak agar plate were used as inoculum and subcultured in nutrient agar, VRB, TCBS, EMB, MacConkey, SS, and BGA to promote the growth of a particular type of bacterium. In all cases, the aseptic condition was maintained via inoculating the samples under laminar airflow. Afterward, the test results were analyzed, and the isolated bacteria present in the samples were identified. For Sugar fermentation, five basic sugars are lactose, dextrose, maltose, sucrose, and mannitol used. Change of color from red to yellow and the presence of gas bubbles were used for the test Acid and gas production, respectively. No color and gas indicated a negative reaction.

RESULTS

Bacterial load of Rui and Kalibaush in different seasons

Table 1: Bacterial load of Rui and Kalibaush in different seasons. Values showing different superscripts are significantly different ($P < 0.05$). Values are mean \pm standard deviation.

Season	Species	
	Rui (<i>Labeo rohita</i>) TVC (Log CFU/g \pm SD)	Kalibaush (<i>Labeo calbasu</i>) TVC (Log CFU/g \pm SD)
Monsoon	5.56 \pm 0.078 ^a	5.68 \pm 0.050 ^b
Post-monsoon	5.51 \pm 0.043 ^a	5.58 \pm 0.057 ^a
Winter	5.36 \pm 0.065 ^a	5.52 \pm 0.040 ^b
Pre-monsoon	5.48 \pm 0.091 ^a	5.63 \pm 0.067 ^b

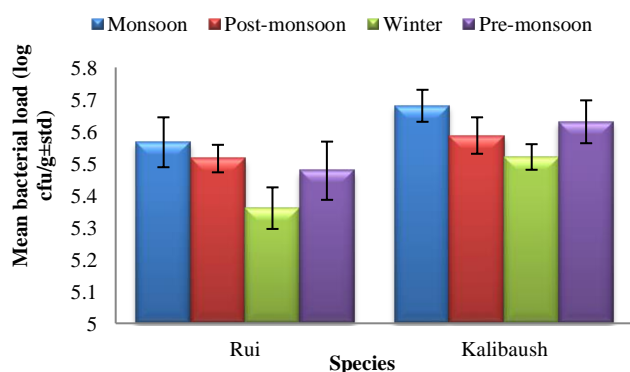


Figure 1: Bacterial load of Rui and Kalibaush in different seasons.

The mean of bacterial load was calculated on a logarithmic scale as CFU/g \pm SD. Table 3.1 shows the overall seasonal result of the bacterial load of both fish. Table 1 and Figure 1 described that the significantly highest bacterial load was found in Rui 5.56 \pm 0.078 CFU/gm in the monsoon season, and the lowest was 5.36 \pm 0.065 in the winter season. Likewise, in Kalibaush, the highest (mean Log CFU/gm \pm SD) bacterial load was 5.68 \pm 0.050 in monsoon, and the lowest bacterial load was 5.52 \pm 0.040 in winter. In post-monsoon, no significant differences were observed between Rui and Kalibaush.

3.2 Effect of temperature on bacterial load of Rui and Kalibaush

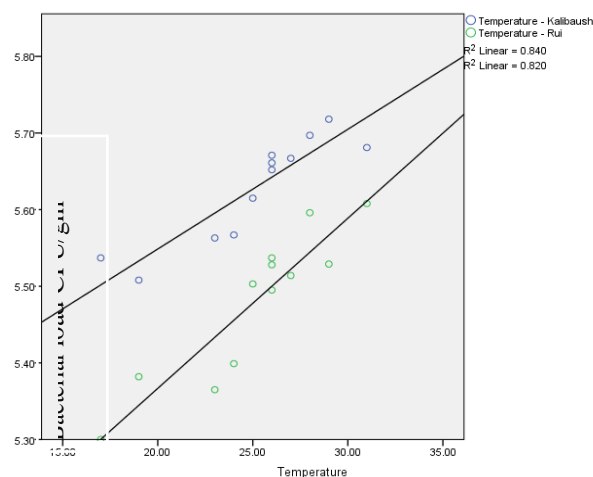


Figure 2: Temperature effect on bacterial load of Rui and Kalibaush

During this experiment period, the temperature remained (17°C-31°C) from May to April. Temperature showed an impact on bacterial load that occurred in both Rui (*Labeo rohita*) and Kalibaush (*Labeo calbasu*). Food, temperature, and other environmental conditions have an influence on the growth and proliferation of bacteria. In this study period, bacterial load was highest at the temperature of 31°C, and the lowest was at 17°C for both types of species. The temperature was correlated positively with the bacterial load of Rui (correlation, $r = 0.904$) and Kalibaush (correlation, $r = 0.916$). Temperature was highly correlated with microbial load in both fishes of Rui and Kalibaush (Fig. 2).

Seasonal Variation of Total Coliform Count (TCC) in Rui and Kalibaush

MPN index was used for calculating total coliform. In post-monsoon, the total coliform was higher in both species. However, the mean Total Coliform of Rui was higher (73 MPN/g) in the monsoon season, indicating contamination of hoar from the surface runoff and/or flood during the monsoon season (Fig. 3). On the other hand, in Kalibaush (*Labeo calbasu*), it was higher (96 MPN/g) in the post-monsoon season (Fig. 3). Interestingly, the Total Coliform of all seasons in both fish complied with the ICMSF standard.

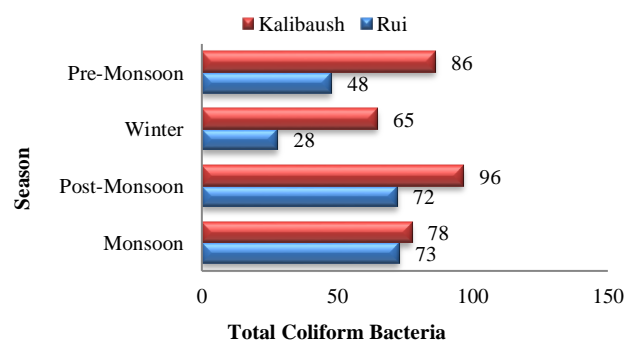


Figure 3: Seasonal variation of Total Coliform Count in Rui and Kalibaush

Pathogenic bacteria isolation and identification based on cultural characteristics, Gram staining, and biochemical test

Morphological characteristics

Three (3) types of bacteria were isolated from Rui and Kalibaus (Table 2). The isolated bacteria are *Escherichia coli*, *Salmonella* spp. and *Vibrio* spp. Gram staining was also done to identify *Escherichia coli*, *Salmonella* spp., and *Vibrio* spp. according to their staining characteristics (Table 3).

Table 2: Cultural characteristics (morphological) of *Salmonella* spp., *Escherichia coli*, and *Vibrio* spp.

Sl. no.	Name of bacteria	Selective media	Colony characteristics
1	<i>E. coli</i>	EMB agar	Metallic sheen (greenish black) colony.
2	<i>Salmonella</i> spp.	SS agar	Opaque, smooth, round with black centered.
3	<i>Vibrio</i> spp.	TCBSS agar	Yellow and green colony.

Table 3: Gram's staining properties of *Salmonella* spp., *Escherichia coli* and *Vibrio* spp.

Characteristics			Identified bacteria
Shape	Arrangements	Gram's staining reaction	
Short plump rods	Single, paired or in short chain	Negative	<i>E. coli</i> (d)
Very short plump rods	Single	Negative	<i>Salmonella</i> spp. (e)
Curved rod shaped	Single	Negative	<i>Vibrio</i> spp. (f)

3.4.2 Biochemical characteristics of *Escherichia coli*, *Salmonella* spp. and *Vibrio* spp.

Five basic sugars such as dextrose, maltose, lactose, sucrose, and mannitol were used for sugar fermentative test. Change of color from red to yellow and the presence of gas bubbles were used for the test Acid and gas production, respectively. No color and gas indicated a negative reaction. The results of the biochemical tests are presented in Table 4.

Table 4: Biochemical properties of *Salmonella* spp., *Escherichia coli* and *Vibrio* spp.

Carbohydrate fermentation test					TSI test	MR test	VP test	Indole test	Interpretation of results
DX	ML	L	S	MN					
AG	AG	AG	AG	AG	+	+	-	+	<i>E. coli</i>
A	A	-	-	A	+	+	-	-	<i>Salmonella</i> spp.
A	A	-	A	A	+	+	-	+	<i>Vibrio</i> spp.

Prevalence of pathogenic bacteria of Rui at different seasons

A spider web analysis was performed for the prevalence of pathogenic bacteria of Rui in different seasons. The availability of pathogenic bacteria in Rui was estimated by staining, motility test, catalase test, and different biochemical tests (sugar fermentation test, indole test, TSI test, Cimon Citrate test, MR-VP test). *E. coli* bacteria was found to be higher in monsoon and pre-monsoon seasons at 55.55% and lowest in winter at 33.33%. Whereas, *Salmonella* spp. steady in (the monsoon, post-monsoon and winter) seasons at 22.22% then marginally increase in pre-monsoon at 33.33%. On the other hand, *Vibrio* spp. was not found in the Winter season, but 22.22% in pre-monsoon. The percentage of *E. coli* was higher than *Salmonella* spp. and *Vibrio* spp. in Rui in all cases.

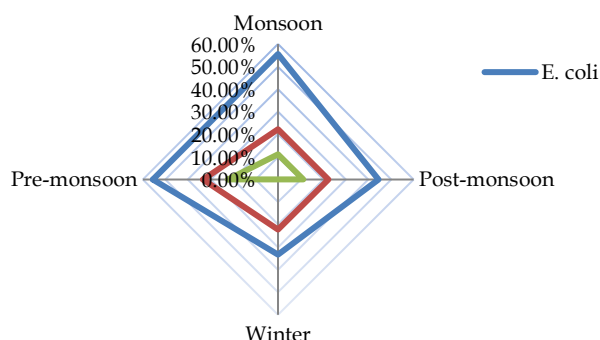


Figure 4: Prevalence of pathogenic bacteria in Rui (*Labeo rohita*)

Prevalence of pathogenic bacteria in Kalibaus at different seasons

A spider web analysis was performed for the prevalence of pathogenic bacteria of Rui in different seasons. The availability of pathogenic bacteria in Kalibaus was estimated by staining, motility test, catalase test, and different biochemical tests (sugar fermentation test, indole test, TSI test, Cimon Citrate test, MR-VP test).

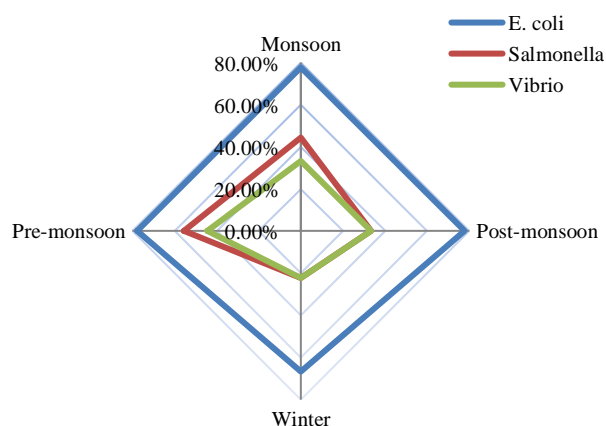


Figure 5: Prevalence of pathogenic bacteria in Kalibaus (*Labeo calbasu*)

E. coli bacteria was found static and higher in (monsoon, pre-monsoon and post-monsoon) season as 77.77%, similarly lower in winter as 66.66%. Whereas, *Salmonella* spp. was



found higher in pre-monsoon at 55.55% and lower in winter at 22.22%. On the other hand, *Vibrio* spp. was found in lower amounts than the other two bacteria. It was the lowest in winter at 11.11%. In winter, these three bacterial species' percentages were found to be lower (Fig. 5). Overall, *E. coli* is found with a high prevalence during the monsoon season.

DISCUSSION

The results indicate that there was a seasonal variation in bacterial load. It was higher in Kalibaus than in Rui during the study period (Table 1). Statistical analysis between these two fish of TVC shows that results were highly significant ($p < 0.05$). Though viable bacterial counts in Kalibaus were higher than the Rui, viable bacterial counts from both fishes comply with ICMSF standards (ICMSF 1986). For white fish, the standard plate count (SPC) accepted by ICMSF is 5×10^5 cfu/g, while it is 100 MPN/g and < 3 MPN/g for total coliform and fecal coliform, respectively. However, the samples must be absent with *Salmonella* spp. and *Vibrio cholera* (including O group-01/non O group-139) (ICMSF 1986). It was reported that the total viable count of bacteria ranged from 9.9×10^6 to 1.4×10^7 CFU/g of intestine in different age groups of Rui fish (Hossain, 1993). Alternatively, it showed that the month of July had the highest bacterial load while it was lowest in the month of January.

Bandekar *et al.* (2004) studied the bacteriological screening of processed Rui (24 samples) and whole Rui (6 samples) obtained from fish processing factories, where TVC in 3 samples of whole Rui out of 6 samples exceeded 5×10^5 cfu/g and 3 samples of processed Rui out of 24 samples exceeding 5×10^5 cfu/g. Seasonal variations of bacterial load between two selected fish species were observed. In the monsoon season, both Rui and Kalibaus showed higher bacterial load (log cfu/g \pm sd) 5.56 ± 0.078 and 5.68 ± 0.050 , respectively. The lowest bacterial load (Log CFU/g \pm SD) in both fishes was found in the winter season, 5.36 ± 0.065 in Rui and 5.52 ± 0.040 in Kalibaus, which are significantly ($P < 0.05$) different. In post-monsoon and pre-monsoon, bacterial load (Log CFU/g \pm SD) was observed as 5.51 ± 0.043 and 5.48 ± 0.091 in Rui, whereas 5.58 ± 0.057 and 5.63 ± 0.067 in Kalibaus. In post-monsoon, in the case of bacterial load between Rui and Kalibaush, non-significant differences were observed. Ahmed H. Al-Harbi (2003) mentioned in tilapia intestine that the total viable count based on seasons ranged from $1.6 \pm 2.0 \times 10^6$ to $5.1 \pm 2.5 \times 10^7$ cfu/g in summer, $6.8 \pm 1.9 \times 10^6$ to $7.5 \pm 1.4 \times 10^7$ cfu/g in early summer, and $8.9 \pm 1.8 \times 10^5$ to $1.3 \pm 0.9 \times 10^7$ cfu/g in winter. Bisth *et al.* (2014) found that sediment of pond (6.40×10^4 cfu) had higher (10 times) bacterial load in comparison to water of the pond (6.93×10^3 cfu/ml), while microbial load in the intestinal of winter season was higher around 100 times (6.67×10^5 cfu/g) while it was higher in 100 times (2.33×10^6 cfu/g) during summer season in comparison to the surficial skin of fish during winter and summer (3.39 and 8.87×10^3 cfu/cm²) respectively. Total coliform of Rui and Kalibaus in all seasons complied with the ICMSF standard. The highest TCC was found (73 MPN/g and 96 MPN/g) in Rui and Kalibaus in the Post-monsoon season. Whereas, the lowest (28 MPN/g and 65 MPN/g) was in Rui and Kalibaus in the winter season. The result of the present study is more or less similar to Begum *et al.* (2010), where she found the highest total coliform (> 240 MPN/g) in a number of samples from

local markets, whereas the lowest count (0.9 MPN/g) was in Rui fish of super shop. The load of TCC in this fish sample indicates a range of contamination. These results are also more or less similar to the other work of Das *et al.* (2007) and Fatema (2005), where they found a comparable range of counts. Rui and Kalibaus were identified with three different types of pathogenic bacteria and it was *E. coli*, *Salmonella* spp., and *Vibrio* spp. Our finding aligned with the findings of Adebayo-Tayo *et al.* (2012), Truong *et al.* (2008), and Zaman (2013). Eosin Methylene Blue (EMB) agar plate showed characteristics of colony color with a metallic green sheen, indicating the *E. coli* (Sharada *et al.* 1999; Shorif, 2013). Morphologically, *E. coli* is a single or paired gram-negative and motile bacteria (Thomas, 1998). *Salmonella* was examined using selective media and found in almost all sample fishes. In SS agar plate colonies were observed with opaque and translucent having black in the centre (Hossain 2002), indicating the colonies of *Salmonella* spp. In gram staining, bacteria were observed with short rods, gram-negative, motile, single, or paired arrangement under the microscope (Samad, 2005). present investigation, *Vibrio* spp. was found in three out of four seasons in Rui samples that were collected from the haor. The morphology of *Vibrio* spp. on the TCBS agar plate showed yellow and green color colonies in this study (Khan *et al.*, 2007). Begum *et al.*, (2010) also found the same bacteria in examined fish but not in Rui fish. The recent study highlighted the seasonal difference in standard bacterial count and occurrence of *E. coli*, *Salmonella* spp., and *Vibrio* spp. collected from open water bodies of Hakaluki haor. To tackle the situation, hygienic conditions should be maintained at every step of fish handling, such as catching, landing, transporting, processing, and marketing. This information will help in controlling bacterial contamination, reducing foodborne disease, and increasing the fish and fishery products' self-life.

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REFERENCES

- Ababouch L 2006: Assuring fish safety and quality in international fish trade. *Marine pollution bulletin* **53** 561-568.
- Abraham A, Sergelidis D, Kirkoudis I, Anagnostou V, Kaitsa-Tsiopoulou E, Kazila P, Papa A 2010: Isolation and antimicrobial resistance of *Staphylococcus* spp. in freshwater fish and Greek marketplaces. *Journal of Aquatic Food Product Technology* **19** 93-102.
- Adebayo-Tayo AC, Odu NN, Michael MU, Okonko IO. 2012: Multi-Drug resistant (MDR) organisms isolated from Sea-foods in Uyo, South-Southern Nigeria. *Nature and Science* **10** 61-70.
- Adediji OB, Okerentugba PO, Innocent-Adiele HC, Okonko IO, Ojeniyi SO, Adejoro SA, Mohamed SA 2012: Benefits, public health hazards and risks associated with fish consumption. *Benefits* 26-32.
- Alam MF, Thomson KJ 2001: Current constraints and future possibilities for Bangladesh fisheries. *Food policy* **26** 297-313.

- Al-Harbi AH, Uddin MN 2003: Quantitative and qualitative studies on bacterial flora of hybrid tilapia (*Oreochromis niloticus* × *O. aureus*) cultured in earthen ponds in Saudi Arabia. *Aquaculture Research* **34** 43-48.
- Anbudhasan P, Uma A, Rebecca G 2012: Assessment of bacteriological quality in selected commercially Important Processed Sea Foods by Polymerase Chain Reaction (PCR). *International Journal of Food, Agriculture and Veterinary Sciences* **2** 20-25.
- Ayulo AMR, Machado RA, Scussel VM 1994: Enterotoxigenic *Escherichia coli* and *Staphylococcus aureus* in fish and seafood from the southern region of Brazil. *International Journal of Food Microbiology* **24** 171-178.
- Azim ME, Wahab MA, Verdegem MCJ 2003: Status of aquaculture and fisheries in Bangladesh. *World Aquaculture* **67** 37-40.
- Bandekar JR, Kamat AS, Karani M, Dhokane V, Shashidhar R, Kakatkar A, Ghadge N, Bhat A, Venugopal V, Warner SB 2004: Bacteriological quality of farmed freshwater fish and shellfish meant for export. *Fishery Technology* **41** 57-62.
- Begum M, Ahmed ATA, Das M, Parveen S 2010: A Comparative Microbiological Assessment of Five Types of Selected Fishes Collected from Two Different Markets. *Advances in Biological Research* **4** 259-265.
- Benson HJ 2002: *Microbiological applications: a laboratory manual in general microbiology*. [McGraw-Hill].
- Bisth A, Singh UP, Pandey NN 2014: Comparative study of seasonal variation in bacterial flora concomitant with farm raised fingerlings of *Cyprinus carpio* at tarai region of Uttarkhand. *Journal of Environmental Biology* **35** 363-367.
- Chowdhury P, Viraraghavan T, Srinivasan A 2010: Biological treatment processes for fish processing wastewater—A review. *Bioresource Technology* **101** 439-449.
- Church N 1998: Feature MAP fish and crustaceans-sensory enhancement. *Food Science and Technology Today* **12** 73-82.
- Clucas IJ, Ward AR 1996: Post-harvest fisheries development: A guide to handling, preservation, processing and quality. Charthan Maritime, Kent ME4 4TB, United Kingdom. pp. 113-116.
- Czerwińska E, Piotrowski W, Sowa E 2008: Technological aspects of fish processing with determination of critical points and identification of microbial contamination (of fish material, devices and product). *Rocznik Ochrona Środowiska* **10** 57-71.
- Das M, Hafiz F, Ahmed MK, Parveen S 2007: Microbiological analysis of some raw fish samples. *Bangladesh Journal of Microbiology* **24** 67-69.
- David OM, Wandili S, Kakai R, Waindi EN 2009: Isolation of *Salmonella* and *Shigella* from fish harvested from the Winam Gulf of Lake Victoria, Kenya. *The Journal of Infection in Developing Countries* **3** 099-104.
- Emikpe BO, Adebisi T, Adedeji OB 2011: Bacteria load on the skin and stomach of *Clarias Gariepinus* and *Oreochromis niloticus* from Ibadan, South West Nigeria: Public health implications. *J. Microbiol. Biotech. Res.* **1** 52-59.
- Fatema K 2005: Bacteriological and heavy metal analysis of Kachki (*Corica soborna*) and Mola (*Amblypharyngodon mola*) fishes. Dept. of Zoology, University of Dhaka.
- Geldreich EE, Clarke NA 1966: Bacterial pollution indicators in the intestinal tract of freshwater fish. *Applied microbiology* **14** 429-437.
- Ghose B 2014: Fisheries and aquaculture in Bangladesh: Challenges and opportunities. *Annals of Aquaculture and Research* **1** 1-5.
- Gold WL, Salit IE 1993: *Aeromonas hydrophila* infections of the skin and soft-tissue: Report of 11 cases and review. *Clinical Infectious Diseases* **1**: 69-74.
- Gram L and Huss HH 1996: Microbiological spoilage of fish and fish products. *International journal of food microbiology* **33** 121-137.
- Herrera FC, Santos JA, Otero A, GarcíaLópez ML 2006: Occurrence of foodborne pathogenic bacteria in retail prepackaged portions of marine fish in Spain. *Journal of Applied Microbiology* **100** 527-536.
- Hood MA, Ness GE, Blake NJ 1983: Relationship among fecal coliforms, *Escherichia coli*, and *Salmonella* spp. in shellfish. *Applied and Environmental Microbiology* **45** 122-126.
- Hossain GM, Nishat A 1989: Planning considerations for water resources development in the haor areas. In AIT-BUET Workshops on Development and Technology, Bangladesh University of Engineering and Technology, Dhaka (Vol. 12).
- Hossain KM 2002: Characterization of bacteria isolated from diarrhoeic calves. MS Thesis, Department of Microbiology and Hygiene, Faculty of Veterinary Science. BAU, Mymensingh.
- Hossain MAR 2015. An overview of fisheries sector of Bangladesh. *Research in Agriculture Livestock and Fisheries* **1** 109-126.
- Hossain MM 1993: Studies on the intestinal microflora of Rohu fish (*Labeo rohita*). MS Thesis, Department of Fisheries Technology, Bangladesh Agricultural University, Bangladesh.
- Huss HH 1998: Control of indigenous pathogenic bacteria in seafood. Fish Inspection, Quality Control, and HACCP: A Global Focus. pp.163.
- ICMSF (International Commission of Microbiological Specification for Food) 1982. Microorganisms in food (Vol. 2). Sampling for microbiological analysis: principles and specific applications. International Commission on the Microbiological Specification of Foods. Univ. Toronto Press, Toronto, Canada.
- ICMSF (International Commission of Microbiological Specification for Food) 1986: Microorganisms in food (Vol. 2). Sampling for microbiological analysis: principles and specific applications. International Commission on the Microbiological Specification of Foods. Univ. Toronto Press, Toronto, Canada.
- ISO (International Standard Organization), 1995. Recommendation of the meeting of the subcommittee, International Organization for Standardization, on meat and meat products. ISO/TC-36/Sc-6. The Netherlands 10-18.
- Ivanek R, Groehn YT, Wiedmann M, Wells MT 2004: Mathematical model of *Listeria monocytogenes* cross-contamination in a fish processing plant. *Journal of food protection* **67** 2688-2697.
- Khan AW, Hossain SJ, Uddin SN 2007: Isolation, identification and determination of antibiotic susceptibility of *Vibrio parahaemolyticus* from shrimp at Khulna region of Bangladesh. *Res J Microbiol* **2** 216-227.

- McMeekin TA, Ross T, Olley J 1992: Application of predictive microbiology to assure the quality and safety of fish and fish products. *International journal of food microbiology* **15** 13-32.
- Nerín C, Aznar M, Carrizo D 2016: Food contamination during food process. *Trends in Food Science & Technology* **48** 63-68.
- Noor R, Acharjee M, Ahmed T, Das KK, Paul L, Munshi SK, Urmi NJ, Rahman F, Alam MZ 2013: Microbiological study of major sea fish available in local markets of dhaka city, Bangladesh. *Journal of Microbiology, Biotechnology and Food Science* **2** 2420-2430.
- Okoro CC, Aboaba OO, Babajide OJ 2010: Quality Assessment of a Nigerian Marine Fish, Mullet (*Liza falcipinnis*) under different Storage Conditions. *New York Science Journal* **3** 21-28.
- Pal M, Mahendra R 2015: Sanitation in Food Establishments. LAP Lambert Academic Publishing, Saarbruchen, Germany.
- Poli BM, Parisi G, Scappini F, Zampacavallo G 2005: Fish welfare and quality as affected by pre-slaughter and slaughter management. *Aquaculture International* **13** 29-49.
- Prabakaran P, Kannan SK, Anand M, Pradeepa A 2011: Microbiological quality assessment in a fish processing plant at Mandapam, Ramanathapuram District. *Archives of Applied Science Research* **3** 135-138.
- Rhea F 2009: Microbiology handbook: Fish and seafood. Leatherhead Food International Ltd. Surrey, UK.
- Samad MA 2005: *Avian salmonellosis*. In: Poultry science and medicine. 1st Ed. The Lyric- Epic Prokasone, BAU campus, Mymensingh, Bangladesh. pp. 504-514.
- Sharada R, Krishnappa G, Raghavan R, Gowda RS, Upendra HA 1999: Isolation and serotyping of *Escherichia coli* from different pathological conditions in poultry. *Indian Journal of Poultry Science* **34** 366-369.
- Shorif M. 2013: Identification of different Bacterial diseases in Jalali Pigeon in Sylhet district. MS Thesis, Department of Microbiology and Immunology, Faculty of Veterinary and Animal Science. Sylhet Agricultural University (SAU), Sylhet.
- Thomas CGA 1998: Gram-negative Bacilli. In: *Medical microbiology*. 6th Ed. Bailliere Tindall, Oxford, UK. pp: 273-274.
- Truong TH, Areechon NS, Wasde MS 2008: Identification and antibiotic sensitivity test of the bacteria isolated from Tra Catfish (*Pangasianodon hypophthalmus* [Sauvage, 1878]) cultured in pond in Vietnam. *Nature and Science* **4** 54-60.
- Zaman SB 2013: Characterization of bacterial flora of Koi (*Anabas testudineus*) fish harvested from different aquatic environments and its public health implications. MS Thesis, Department of Public Health and Food Hygiene, Faculty of Veterinary Science. Bangladesh Agricultural University (BAU), Bangladesh.