EFFECT OF PRE-DRYING HANDLING ON QUALITY OF FINAL DRIED BOMBAY DUCK (HARPODON NEHERIUS)

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Key words: Bombay duck, drying, salting, non-protein nitrogenous substances, free fatty acids, peroxide value.

ABSTRACT

Proliferation of spoilage microorganisms during pre-processing delay, with associated deteriorative changes in non-protein nitrogenous substances and lipids, and its impact on the sensory characteristics of the final dried products were studied. Bombay duck was judged as unfit for human consumption beyond 8 hrs of exposure at 28°C. However, chilling the fish to 0°C or salting the fish after gutting at 28°C or drying at 45°C without delay effectively reduced the proliferation of spoilage microorganisms, and associated biochemical and sensory characteristics during drying. Higher levels of proteolytic and lipolytic bacterial load, free fatty acids, trimethylamine and total volatile bases due to predrying exposure at 28°C without salt or 10% salt solution resulted in dried fish with least sensorial characteristics. Predrying exposure at 0°C did not had any effect on the development of peroxides compared to 28°C, however exposure to saturated salt solution or dry salt enhanced the development of peroxides. Dressed Bombay duck, treated with 20% salt solution for 16 hours and artificially dried at 45°C was judged superior by sensory panel.

I. INTRODUCTION

Harpodon neherius, is a single species fishery of high magnitude along Gujarat and Maharashtra coast. It is popularly known as Bombay duck, and one of the largely produced and relished dried fish in the coastal and some interior parts of India. Due to very high moisture content of 90.98%, it is unsuitable to use either in fresh form or in frozen form, and almost entire catch of the Bombay duck is consumed in unsalted and sundried form [27]. Abundant catches of Bombay duck is transported from the fishing ground to the landing center without icing and depending on the distance from the shore to the fishing ground, it takes more than four hours to reach the shore. Heaps of the landed Bombay duck during odd hours during peak seasons has to wait for considerable length of time of their disposal, resulting in inevitable deterioration in quality. Fresh fish stored at ambient temperature becomes unsuitable for human consumption within 12 hours [28].

Such raw materials are dried on the shore without salting in sunshine that may takes more than three days depending on the weather conditions. Initial freshness of raw material affects the end product [27]. Main reasons for the availability of most of the low quality dried fish along Kerala, Tamilnadu and Maharashtra coast is due to the effect of the nature of the raw material, pre-drying delay, unhygienic handling, unpredictable weather condition, improper salting practices and unprotected storage on the quality of the dried fish and its keeping quality [3, 11, 19].

Non-protein nitrogenous compounds and lipids are involved in post mortem reactions in the flesh of fish. Accumulation of non protein nitrogenous substance such as trimethylamine, total volatile bases and free amino acids due to microbial and indigenous enzymes in the fish flesh results in unacceptability of the product for human consumption [5, 18, 30, 32, 33]. Deteriorative changes in lipids due to its susceptibility for oxidation or enzymatic reaction results in the formation of components that are associated with undesirable secondary reactions of the dried fish product [10, 12, 13, 26, 31]. Hence, changes in non protein nitrogenous substances and deteriorative changes in lipids in fish muscle serves as a measure of freshness or staleness of seafood, and helps in making a strategy to improve the existing traditional processing methods to produce the product of consumer acceptability.

Spoilage rate can be reduced to greater extent by chilling as the spoilage retarding effect is especially high close to freezing point of fish flesh, and on the other hand rate of spoilage and period of drying can be reduced, and sensory properties of the products can be improved by salting. Even after being an important fishery of Maharashtra and Gujarat coast not much attempt has been made to improve the traditional method of processing. Pre-drying spoilage contribute considerable nutritional loss in food chain and drying technique is often
considered to produce durable product of energy saving one compared to frozen products and little or no additional energy is required, and hence considerable scope exists for improving quality of dried fish produced in India to feed the poor people by improving the processing techniques. This is an attempt to study the impact of deteriorative changes taking place during pre-drying holding on the keeping quality of the final dried product, and we have made an effort to reduce the rate of deterioration in Bombay duck by chilling or salting after gutting, and artificial drying.

II. MATERIALS AND METHODS

1. Fish Samples

The Fresh Bombay Duck (FBD) samples were transported in an insulated container after adequately icing them in the proportion of 1:1 fish to ice, to the laboratory of Central Institute of Fisheries Education (ICAR). The time lapsed between catching at the fishing ground and processing may not exceed over four to six hours and temperature recorded during the catching and the processing did not exceed 4°C. The FBD samples belonging to size group of 21 to 23 cm long; weighing around 80 to 85 g were sorted out on a sanitized stainless steel working table were washed using chilled running water system maintained between 2-4°C. FBD procured in this manner was used in two lots for studying the effect of post harvest delay on the quality of the final dried product.

2. Effect of Preprocessing Delay on the Quality of the Freshly Caught Bombay Duck

To study the effect of preprocessing delay on the quality of the raw material, first lot freshly caught fish was divided in to four sub-lots, and the first three sub-lots were exposed to the temperature of 28°C for 4, 8, and 12 hours, and the last sub-lot was exposed to the temperature of 0°C for 12 hours in thermostatically controlled cooling incubator (Rotec, Cochin) in the polythene pouches (US-28C/4H, US-28C/8H, US-28C/12H and US-0C/12H respectively). Samples were drawn from each sub-lot at regular intervals of time for further analysis.

3. Effect of Gutting and Salting on the Deteriorative Changes of Bombay Duck

Combined effect of gutting and salting on pre-drying spoilage was studied on second lot of FBD after gutting by slit opening the belly without removing the head, fins or tails under sanitary conditions, washing under chilled running water system. Dressed Bombay duck (DBD) produced in this manner was divided in to four sub-lots, and the first three sub-lots were treated with 10, 20 and 36% (Saturated) salt solution in the ratio of 1:1.5 salt solution to fish proportion, and the fourth sub-lot was treated with dry salt in the ratio of 1:6 salt to fish proportion as per Bureau of Indian Standards specifications [16-17]. The common salt, having around 99% sodium chloride obtained locally from market with the size of 3-5 mm of coarse salt in 2/3 parts and the size of 1-0.5 mm fine salt in 1/3 parts were used. Salting was done as per the method explained by the Bureau of Indian Standards specifications [15-17]. Samples were drawn from each sub-lot at regular intervals of time for further analysis. The visceral organs were pulled out after split opening the belly. Washed and dressed Bombay duck is washed in 3.5% salt solutions, so as to remove blood, slimes, dirt etc. The material was drained prior to drying.

4. Artificial Drying of the Samples with Different Pre-drying Conditions

Samples held at 28 or 0°C with or without salts (DBD, US-28C/4H, US-28C/8H, US-28C/12H, US-0C/12H, SA10-DBD, SA20-DBD, SA36-DBD and SADS-DBD) were dried in Torry kiln (Torry Research Station, UK) at the temperature of 45°C with air velocity of 1.003 meter/second and relative humidity of 60±2%. All the samples were hung on one meter long iron rods after interlocking the jaws which were mounted on wooden frames of the trolley and were loaded in the drying chamber of the kiln with proper labeling. The temperature, air velocity and relative humidity was monitored regularly, recorded at regular intervals of time and maintained at constant rate. Samples were drawn from artificial dried samples at regular intervals of time for analysis.

5. Proximate Analysis

Samples drawn and analyzed at different intervals of processing were performed in quadruplicates. The wet fish samples were blended in a homogenizer (Rotec, Cochin) at 3,000 rpm for 10 minutes and dried fish samples were powdered in a Waring blender (Philips India, Bombay) at 22,000 rpm for 10 minutes. Moisture content of the samples were estimated as per FAO [9] and expressed as percentage of moisture. Salt content of samples were estimated as per FAO [8] and expressed as percentage of salt. The total lipid in the fish was extracted using chloroform methanol phase separation and peroxide value (POV) of the samples were estimated by the method described by Lima and others [24], and expressed as millimoles of oxygen/kg of fat. Free fatty acids (FFA) of the sample were estimated by the method described by IS: 5734 [15] and is expressed as percentage of oleic acid on lipid basis. Trichloro acetic acid extract was prepared as per FAO [9] and used for measuring nonprotein nitrogenous substances (NPNs) like trimethylamine nitrogen (TMAN), total volatile bases nitrogen (TVBN) and alpha amino nitrogen (AAN). TMAN and TVBN content of the sample was determined by the method described by Martin [25] and the values were expressed as mg/100 g of fish muscle. The AAN in the samples were estimated by the method described by Pope and Stevens [29] and value is expressed as mg/100 g fish muscle.

6. Microbiological Methods

Glassware and prepared media were sterilized using moist
heat at 121°C for 15 minutes. Petri dishes, homogenizers, pipettes were sterilized using dry heat at 180°C for 1 hour. Skimmed milk 10%, trybutyrin solution 10% were sterilized by tendylisation method, where solution was free steamed for 1 hour on first day and for 30 minutes on the next 2 successive days. Mesophilic bacterial count (MBC) was determined as per APHA [2] method. MBC was enumerated and expressed as mesophiles per gram of sample on dry weight basis. Proteolytic bacterial count (PBC) was determined using the method of Leo [23]. Proteolytic positive bacterial colony forms clear zone around the colony and was expressed as number of proteolytic bacteria per gram of sample on dry weight basis. Lipolytic bacterial count (LBC) was determined by the method explained by Collines [4] using Trybutyrin agar. Hydrolysis of trybutyrin results in clearing of medium and formation of a clear zone around the colony, which was enumerated and expressed as lipolytic count per gram of sample on dry weight basis. Total mould count (TMC) was determined according to the method described by Leo [23], which was enumerated and expressed as total mould count per gram of sample on dry weight basis.

7. Sensory Evaluation
Sensory attributes like appearance, colour, odor, texture, and flavour were conducted using a ten member panel of trained professionals. Samples were labeled in such a way that the panelist will not be able to identify them and were placed in separate booths. The panelists were provided with clean water to rinse their mouth after tasting each sample. The samples were evaluated using a nine point hedonic scale basis (9 = like extremely, 8 = like very much, 7 = like moderately, 6 = like slightly, 5 = neither like nor dislike, 4 = dislike slightly, 3 = dislike moderately, 2 = dislike very much and 1 = dislike extremely) [22].

8. Chemicals
All the chemicals used were of analytical grade and were obtained from Merck Limited (Mumbai, India).

9. Statistical Analysis
A completely randomized design was performed using quadruplicate samples through the study. One- and two-way ANOVA was performed using Statographies 2.1 (STSC Inc., Rockville, MD). The difference in means was analyzed using a Turkey HSD test ($p < 0.05$).

III. RESULTS AND DISCUSSIONS
1. Effect of Pre-drying Delay on the Quality of the Freshly Caught Bombay Duck
When FBD samples were held at 28°C for 12 hours (US-28C/12H), MBC was increased by 538.46 ± 0.25 folds, but as the temperature of the storage of the samples reduced to 0°C (US-0C/12H) the increase of MBC was reduced drastically to 7.89 ± 0.025 folds. US-28C/12H registered higher ($p < 0.05$) levels of MBC compared to US-0C/12H samples, all along the storage period. PBC increased by 79.00 ± 0.022 folds at 28°C, but only by 15.53 ± 0.021 folds at 0°C. Similarly during 12 hours of holding, LBC increased by 81.5 ± 0.025 folds at 28°C, but only by 17.20 ± 0.023 folds at 0°C. On the other hand, FFA in US-28C/12H samples increased by 1.60 ± 0.019 folds in 12 hours, but in US-0C/12H samples the increase was only by 1.30 ± 0.02 folds (Fig. 1).

Reduction in the release of FFA in chilled fish (US-0C/12H) compared to samples stored at ambient temperature (US-28C/12H) may be due to decreased lipolytic activity close to freezing point of fish flesh [20], as at ambient temperature autolytic or bacterial lipase catalyse the release of FFA from glycerides and phospholipids [12]. Accumulation of FFA in fish flesh is undesirable due to secondary reactions leading to quality deterioration [1]. Similarly, when FBD samples were stored at 28°C, TMAN, TVBN and AAN increased by 4.44 ± 0.021, 5.53 ± 0.022 and 2.40 ± 0.02 folds in 12 hours (US-28C/12H), but as the temperature of storage reduced to 0°C (US-0C/12H), rate of increase of these values remained, respectively at 1.37 ± 0.018, 1.35 ± 0.016 and 0.92 ± 0.019 folds only (Fig. 2).

TMAN, TVBN and AAN increased remarkably ($p < 0.05$) during the storage of FBD at 28°C, but there was a considerable reduction ($p < 0.05$) in the development of TMAN, TVBN and AAN was noticed when temperature reduced to 0°C. Trimethylamine oxidase produce by spoilage organisms reduces trimethylamine oxide of fish flesh to trimethylamine that is believed to react with fish fats to produce the typical spoilage odor that are associated with fish beyond their prime [32, 35]. FBD samples held at 28°C more than 4 hours (US-0C/8H and US-0C/12H) were found unacceptable by the sensory panel and scored less than 5 points on Hedonic scale.
FFA, TMAN and TVBN registered at this point of storage were 6.76 ± 0.02% of oleic acid, 11.83 mg% and 35.17 ± 0.15 mg% respectively. The trimethylamine level in fresh fish rejected by sensory panels varies between species but is around 10 to 15 mg% in aerobically stored fresh fish [6] and in the present study TMAN level increased more than this level beyond 8 hours of storage at ambient temperature (US-28C/8H and US-28C/12H). Total volatile bases is mostly formed by bacterial or tissue autolysis leading to a deteriorative odors and flavors and its limits well be 30-35 mg% in teleost fishes [5], and in the present study TVBN level increased more than this level in sample stored at ambient temperature beyond 8 hours (US-28C/8H and US-28C/12H). But POV of lipids increased by 2.00 ± 0.02 folds of the initial values in both samples stored at 28°C for 12 hours (US-28C/12 and US-0C/12) (Fig. 3). It is instructive to note here that even though accumulation of fatty acids reduced drastically ($p < 0.05$) as the storage temperature of the samples reduced from 28 to 0°C, it did not have significant effect ($p > 0.05$) on the release of peroxides. Since samples were exposed to atmospheric oxygen in both US-28C/12-FBD and US-0C/12-FBD samples, highly unsaturated fatty acids are susceptible for oxidation due to contact with oxygen [33]. Even though hydroperoxides are odorless and flavorless compounds, and not related directly to the actual sensorial objectionable rancification and discoloration as we observed in the final dried products might latter lead to objectionable secondary reactions [37].

FBD contained semitransparent body, silvery white abdomen and transparent fins, and during the exposure of the samples at the atmospheric temperature of 28°C turned grayish, translucent, and mushy by 12 hours, and was found unacceptable for human consumption by sensory panel. Since the samples were turned unfit beyond 8 hours of storage by sensory evaluation, we did not store the samples beyond 12 hours of storage. FFA, POV, TMAN, TVBN and AAN values registered at 8 hours of storage at 28°C was 8.19% oleic acid, 4.89 millimoles of oxygen/kg of fat, 29.61 ± 0.03 mg%, 67.34 ± 0.025 mg%, and 110.15 ± 0.02 mg% respectively with hedonic scale of 4.20 ± 0.1 points. Hence, if the huge catch of Bombay duck is waiting further processing, it is advised to chill the fish to reduce the rate of deterioration [35].

2. Effect of Gutting and Salting on the Deteriorative Changes of Bombay Duck

Second lot of freshly caught Bombay duck samples were degutted, washed and treated with 10, 20, 36% (Saturated) salt solution or dry salt in four different sub-lots (SA10-DBD, SA20-DBD, SA36-DBD and SADS-DBD, respectively). Since Bombay duck has a higher moisture content of 90.98% compared to other fishes, maximum period of salting in the present study was 16 hours. During pre-drying holding of DBD samples at different concentrations of salts at 28°C, MBC increased respectively by 1.40 ± 0.002 and 1.28 ± 0.003 folds in SA10-DBD and SA20-DBD samples, but decreased respectively by 0.32 ± 0.003 and 0.26 ± 0.002 folds in SA36-DBD and SADS-DBD samples. During these period PBC decreased by 0.52 ± 0.002, 0.46 ± 0.001, 0.4 ± 0.003, and 0.40 ± 0.002 folds, and LBC decreased by 0.31 ± 0.003, 0.28 ± 0.002, 0.23 ± 0.001, and 0.20 ± 0.002 folds of initial value in SA10-DBD, SA20-DBD, SA36-DBD and SADS-DBD samples,
Fig. 4. Changes in MBC, LBC and FFA in SA10-DBD, SA20-DBD, SA36-DBD and SADS-DBD during salting at 28°C.

respectively. On the other hand, FFA of samples were increased by $2.50 \pm 0.02$, $2.20 \pm 0.019$, $2.10 \pm 0.016$ and $1.90 \pm 0.019$ folds of its initial value in samples treated with 10, 20, 36% salt solution and dry salt, respectively by 16 hours (Fig. 4).

During salting period we have observed that the rate of increase of FFA decreased ($p < 0.05$) with the increase in the concentration of salt. Similarly TMAN increased by $1.60 \pm 0.02$, $1.50 \pm 0.021$, $1.30 \pm 0.02$ and $1.20 \pm 0.018$ folds (Fig. 5), and TVBN increased by $1.20 \pm 0.019$, $1.20 \pm 0.021$, $1.00 \pm 0.021$ and $1.00 \pm 0.016$ folds (Fig. 6) in samples treated with 10, 20, 36% salt solution and dry salt (SA10-DBD, SA20-DBD, SA36-DBD and SADS-DBD, respectively), respectively by 16 hours. Even though AAN levels fluctuated during salting period increase remained at a constant rate of $1.00 \pm 0.012$ fold (Fig. 7). TMAN and TVBN values increased ($p < 0.05$) gradually with salting period, but the rate of increase of these values decreased with the increase in salt concentration ($p < 0.05$). But it is interesting to note here that there was no significant ($p > 0.05$) difference in TMAN and TVBN values were observed in samples treated with saturated salt solution and dry salt (SA36-DBD and SADS-DBD). Similarly during salting of DBD samples, POV increased by $2.80 \pm 0.015$, $3.10 \pm 0.018$, $4.50 \pm 0.019$ and $4.80 \pm 0.015$ folds, respectively in SA10-DBD, SA20-DBD, SA36-DBD and SADS-DBD samples (Fig. 8). During this period contrary to FFA, rate of increase of POV increased ($p < 0.05$) with the increase in the concentration of salt, which may be due to the oxidation of highly unsaturated fatty acids in fish lipids by catalytic activity...
of common salt, prooxidation action of moisture and auto-oxidation by atmosphere oxygen [36]. At the end of the salting period moisture content was 86.19 ± 0.035, 81.29 ± 0.04, 70.56 ± 0.03, and 68.02 ± 0.03% in SA10-DBD, SA20-DBD, SA36-DBD and SADS-DBD samples (Fig. 8) and salt content was 10.89 ± 0.02, 13.01 ± 0.03, 16.66 ± 0.02 and 18.02 ± 0.02%, respectively on dry weight basis. Moisture and salt content in FBD was 89.92 ± 0.02% and 4.11 ± 0.1%, respectively. But increase in concentration of salt favors the formation of peroxides which in turn inversely correlates with the quality preference, even though washing salted Bombay duck with 3.5% salt solution prior to drying decreases salt crystal deposit over the product.

On the other hand, rate of release of fatty acids, and formation of non-protein nitrogenous substances decreased (p < 0.05) with the increase in the concentration of salt, may attributes to the inhibition of bacterial activity by sodium chloride [33]. Sensory panelist considered dressed Bombay duck treated with 20% salt solution for 16 hours (SA20-DBD) were best among the salted samples (SA10-DBD, SA36-DBD and SADS-DBD) and scored highest for flavors (Hedonic scale of 8.6 ± 1.4). Salting Bombay duck with either saturated sodium chloride solution or dry salt (SA36-DBD and SADS-DBD) was found unsuitable by sensory panelist due to the high salt content in the flesh, and samples turned opaque, faint straw color, with rough texture and longitudinal wrinkles, and samples scored 6.8 ± 0.8 and 6.5 ± 1.2 respectively on hedonic scale.

Samples treated with 10% sodium chloride solution were with translucent body, grayish abdomen, dark fins, and chalky surface, and samples scored 7.85 ± 0.17 on hedonic scale.

3. Effect of the Artificial Drying on the Quality of the Final Dried Products

Bombay duck samples with various pre-drying conditions (DBD, US-28C/4H, US-28C/8H, US-28C/12H, US-0C/12H, SA10-DBD, SA20-DBD, SA36-DBD and SADS-DBD), were dried artificially in Torry kiln under similar controlled conditions. At constant rate of 45°C, with the controlled air speed of 1.003 meters per second and relative humidity of 60%, it took around 40 hours to attain final moisture of 11.96 ± 0.02% in the unsalted samples (DBD, US-28C/4H, US-28C/8H, US-28C/12H or US-0C/12H), and took around 40, 36, 28 and 24 hours to attain final moisture content of 14.09 ± 0.2, 16.29 ± 0.2, 24.57 ± 0.03 and 27.36 ± 0.02%, respectively in samples treated with 10, 20, 36% salt solution, and dry salt (SA10-DBD, SA20-DBD, SA36-DBD and SADS-DBD) (Fig. 9). Hence preliminary salting shortens the overall drying period to the state of sufficient dryness.

During artificial drying of DBD, US-28C/4H, US-28C/8H, US-28C/12H, US-0C/12H, SA10-DBD, SA20-DBD, SA36-DBD and SADS-DBD samples, MBC increased by 79.49 ± 0.65, 58.67 ± 0.95, 25.00 ± 0.72, 28.33 ± 0.45, 150.00 ± 0.95, 272.00 ± 0.42, 100.00 ± 0.35, 44.0 ± 0.15 and 2.60 ± 0.45 folds and TMC increased by 23.13 ± 0.45, 87.88 ± 0.23, 51.43 ± 0.44, 44.07 ± 0.33, 138.46 ± 0.22, 33.87 ± 0.42, 11.66 ± 0.20, 1.20 ± 0.09 and 0.40 ± 0.09 folds from the initial value (Figs. 10 and 11). Rate of increase (p < 0.05) of MBC was more for
SA10-DBD, followed by US-0C/12H, SA20-DBD, DBD, US-28C/12H, US-28C/8H, US-28C/4H, SA36-DBD and SADS-DBD samples, however rate of increase ($p < 0.05$) of TMC was more for US-0C/12H, followed by SA10-DBD, SA20-DBD, DBD, US-28C/12H, US-28C/8H, US-28C/4H, SA36-DBD and SADS-DBD samples. It is interesting to note here that higher initial microbial load resulted in the dried fish products with higher microbial load ($p < 0.05$).

FFA values of the lipids in DBD, US-28C/4H, US-28C/8H, US-28C/12, US-0C/12, SA10-DBD, SA20-DBD, SA36-DBD and SADS-DBD samples at the initial period of drying was...
free fatty acids in the dried products. The rate of liberation of fatty acids and formation of peroxides is related to the formation of peroxide and moderately disliked by the entire panelist with very strong dry-fishy smell, very firm with longitudinal shrinkage, and scored 7.00 ° on hedonic scale. The samples held at 28°C for 8 and 12 hours (US-28C/8H, US-28C/12, US-0C/12, SA10-DBD, SA20-DBD, SA36-DBD and SADS-DBD samples). TMAN increased by 1.7 ± 0.02, 1.7 ± 0.01, 1.40 ± 0.03, 1.20 ± 0.01, 1.40 ± 0.02, 1.30 ± 0.01, 1.30 ± 0.03, 1.10 ± 0.01 and 1.00 ± 0.03 folds, TVBN increased by 1.20 ± 0.02, 1.40 ± 0.019, 1.20 ± 0.01, 1.10 ± 0.01, 1.20 ± 0.01, 1.20 ± 0.01, 1.10 ± 0.02 and 1.00 ± 0.09 folds (Fig. 14), and AAN increased by 1.10 ± 0.02, 1.40 ± 0.01, 1.40 ± 0.03, 1.20 ± 0.01, 1.20 ± 0.02, 1.20 ± 0.01, 1.10 ± 0.02 and 1.00 ± 0.02 folds during 40 hours of artificial drying (Fig. 15). In dried fish, spoilage was quite evident, when TMAN values reached 50 mg% [34], but in salted fish it was still quite low, [14]. Connell [5] suggested that TVBN of 100-200 mg% on dry weight basis as the limit beyond which salted or dried fish could be considered as spoiled, while, Koizumi et al. [21] observed that salted dried fish could be considered as stale when TVBN increased above 100 mg% dry weight basis. It is found out that if the TMAN and TVBN values are higher, prior to drying, the final dried product continues to exhibit higher TMAN and TVBN values [7].

Artificial dried DBD samples were grayish white, dry fish odor, very firm with longitudinal shrinkage, and scored 7.00 ± 1.4 on hedonic scale. The samples held at 28°C for 8 and 12 hours prior to drying (US-28C/8H, US-28C/12H) were unfit for human consumption and turned dark yellowish in color, with very strong dry-fishy smell, very firm with longitudinal shrinkage and moderately disliked by the entire panelist with

5.88 ± 0.04, 6.74 ± 0.03, 8.26 ± 0.02, 9.17 ± 0.025, 7.83 ± 0.03, 10.23 ± 0.06, 9.28 ± 0.04, 8.59 ± 0.06 and 7.87 ± 0.03% oleic acid (Fig. 12), while POV of the lipids were 2.59 ± 0.03, 3.56 ± 0.025, 4.96 ± 0.02, 5.25 ± 0.03, 5.18 ± 0.025, 9.21 ± 0.03, 10.13 ± 0.05, 14.41 ± 0.02 and 15.47 ± 0.04 millimoles of oxygen/kg of fat respectively (Fig. 13).

A POV of 10-15 mg/kg of lipids indicates rancidity [5]. During artificial drying of these samples (DBD, US-28C/4H, US-28C/8H, US-28C/12, US-0C/12, SA10-DBD, SA20-DBD, SA36-DBD and SADS-DBD), POV increased by 8.40 ± 0.02, 6.70 ± 0.012, 5.10 ± 0.011, 5.10 ± 0.022, 4.70 ± 0.021, 4.10 ± 0.02, 3.50 ± 0.021, 2.05 ± 0.019, and 1.50 ± 0.016 folds at end of respective drying period, but rate of development of FFA remained at a rate of 5.20 ± 0.001 folds in DBD, US-28C/4H, US-28C/8H, US-28C/12 and US-0C/12 samples, but 5.80 ± 0.02, 4.25 ± 0.018, 2.50 ± 0.019 and 1.50 ± 0.021 folds in SA10-DBD, SA20-DBD, SA36-DBD and SADS-DBD samples. Rate of liberation of fatty acids and formation of peroxides increased (p < 0.05) in the same order during artificial drying of the samples held at 28°C or 0°C (US-28C/4H, US-28C/8H, US-28C/12H or US-0C/12H). It is inferred here that the holding period at 28°C is related to the formation of peroxide and free fatty acids in the dried products.

Similarly, TMAN values were 10.57 ± 0.03, 11.85 ± 0.02, 29.44 ± 0.03, 47.51 ± 0.02, 13.96 ± 0.025, 16.82 ± 0.02, 15.52 ± 0.05, 12.68 ± 0.04 and 12.33 ± 0.02 mg%, TVBN values were 31.53 ± 0.02, 35.90 ± 0.03, 67.31 ± 0.03, 175.4 ± 0.02, 42.67 ± 0.025, 38.85 ± 0.02, 37.62 ± 0.02, 32.32 ± 0.02 and 32.46 ± 0.02 mg%, and AAN values were 79.44 ± 0.02, 99.32 ± 0.025, 10.76 ± 0.04, 189.90 ± 0.025, 72.60 ± 0.03, 76.43 ± 0.05, 77.17 ± 0.12, 78.66 ± 0.09, and 9.73 ± 0.02 mg%, respectively in DBD, US-28C/4H, US-28C/8H, US-28C/12, US-0C/12, SA10-DBD, SA20-DBD, SA36-DBD and SADS-DBD samples. TMAN increased by 1.7 ± 0.02, 1.7 ± 0.01, 1.40 ± 0.03, 1.20 ± 0.01, 1.40 ± 0.02, 1.30 ± 0.01, 1.30 ± 0.03, 1.10 ± 0.01 and 1.00 ± 0.03 folds, TVBN increased by 1.20 ± 0.02, 1.40 ± 0.019, 1.20 ± 0.01, 1.10 ± 0.01, 1.20 ± 0.01, 1.20 ± 0.01, 1.10 ± 0.02 and 1.00 ± 0.009 folds (Fig. 14), and AAN increased by 1.10 ± 0.02, 1.40 ± 0.01, 1.40 ± 0.03, 1.20 ± 0.01, 1.20 ± 0.02, 1.20 ± 0.01, 1.10 ± 0.02, 1.00 ± 0.01 and 1.00 ± 0.02 folds during 40 hours of artificial drying (Fig. 15).
the Hedonic scale of 3.23 ± 0.15 and 3.02 ± 0.13, respectively.

While samples held at 0°C for 12 hours (US-0C/12H) were
turned straw yellow color, with slight dry-fishy colors, very
firm with longitudinal shrinkage and scored 6.60 ± 1.5 and
those samples held at 28°C for 4 hours (US-28C/4H) were
turned straw yellow color, with slight dry-fishy odour, very
firm with longitudinal shrinkage and scored 6.00 ± 1.5, during
40 hours of artificial drying. While comparing salted and
artificial dried samples, artificial dried sample treated with
20% salt (SA20-28C/16H) was rated best (Hedonic score of
8.20 ± 1.4) with the better appealing color of grayish white and
firm texture. In general terms DBD samples scored 7.00 ± 1.4,
SA10-DBD samples scored 7.60 ± 1.6, SA20-DBD samples
8.20 ± 1.4, SA36-DBD samples scored 6.60 ± 0.09 and SADS-
DBD samples scored 6.20 ± 1.2 on hedonic scale. It is inter-
esting to note here that as the salt concentration increased,
colour of the samples turned from grayish white to increasing
yellowish ting with salt deposits on the surface, but became
softer and smoother. In fact direct comparison between salted
samples may not hold true as the moisture levels varied be-
tween 11.96 ± 0.02 and 27.36 ± 0.02% amongst the samples.
Hence the initial quality of the Bombay duck fish has a direct
impact on the quality of final dried products. At the end of drying period
dressed Bombay duck, treated with 20% salt solution for 16
hours and artificially dried at 45°C was judged superior by
sensory panel.

IV. CONCLUSION

Freshly caught Bombay duck becomes unfit for human
consumption beyond 8 hours of holding at ambient tempera-
ture as decided by the microbial, biochemical and sensory
characteristics. If Bombay duck caught are waiting for fur-
ther processing it is advisable to chill them just above the
freezing point to retard the spoilage, as indicated by lower
MBC, PBC, LBC, FFA, TMAN, TVBN and AAN. However,
this is inefficient in preventing the accumulation of peroxides.
On the other hand, gutting and salting the freshly caught
Bombay duck prior to artificial drying reduces the drying pe-
riod and reduces the deteriorative changes. Even though in-
crease in the concentration of salt decreases deteriorative
changes as indicated by higher microbial count and non-pro-
tein nitrogenous substances in Bombay duck, higher concen-
tration of salt facilitates the peroxide accumulation and de-
creases the quality preference due to discoloration and salt
deposits. The initial quality of the raw material used and
spoilage taking place during drying has a direct impact on the
quality of the final dried product. At the end of drying period
dressed Bombay duck, treated with 20% salt solution for 16
hours and artificially dried at 45°C was judged superior by
sensory panel.

REFERENCES

1. Aidos, I., Schelvis-Smit, R., Veldman, M., Luten, J. B., Van Der Padt, A.,
and Boom, R. M., “Chemical and sensory evaluation of crude oil ex-
tracted from herring byproducts from different processing operations,”
Journal of Agricultural and Food Chemistry, Vol. 51, No. 7, pp. 1897-
1903 (2003).
2. APHIA, Recommended Procedure for the Examination of Seawater and
12-13 (1982).
extension of fresh fish and shellfish,” Critical Reviews in Food Science
(1967).
fillets packed in vacuum or modified atmospheres,” International Journal
7. Dissaraphong, S., Benjakul, S., Visessanguan, W., and Kishimura, H.,
“The influence of storage conditions of tuna viscera before fermentation
on the chemical, physical and microbiological changes in fish sauce
during fermentation,” Bioresource Technology, Vol. 97, No. 16, pp. 2032-
8. FAO, “The prevention of losses in cured fish,” Fisheries Technical Paper,
9. FAO, “Support to regional aquaculture activities in Latin America and the
Caribbean,” In: Olvera-Novoa, M. A., Martinez Palacios, C. A., and Real
de Leon, E. (Eds.), Nutrition of Fish and Crustaceans- a Laboratory
Manual, AB479/E, Food and Agriculture Organization of the United
lems and their measurement in meat, poultry and fish product,” In:
Pearson, A. M. and Dutson, T. R. (Eds.), Quality Attributes and Their
Measurement in Meat, Poultry and Fish Products-Advance in Meat Re-


