Microbiological Control of Internal Surfaces of Appliances in the Butcher’s Cold Chain in Lubumbashi

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Abstract

Introduction: Food contamination with microorganisms can occur at any stage of the process, from food production to consumption, and can be the result of environmental contamination. This study was carried out in order to identify the microorganisms forming the flora and to determine the level of microbial contamination of the internal surfaces of devices in the cold chain of butcher shops. Method: This is a comparative cross-sectional descriptive study of 30 internal surfaces of ten devices in the chain of two butchers. Results: It seems that the sampling points of Butchery 1 have the highest level of contamination at 87% compared to Butchery 2 (46.67%) with a p = 0.02508 (<0.05). The organisms isolated were *S. aureus* (54%) and *albus* (23%), *fusarium* and *A. niger* respectively with 8%, and *A. fumigatus* (7%); against 47% (Boucherie II) with a, The isolated microorganisms were *S. aureus* (43%), 29% for *K. planticola* and 14% for *A. fumigatus* and *fusarium spp*. The level of surface contamination for all isolated organisms was >1000 CFU/cm². The operating temperature range of the devices varied between −2˚C and 6˚C, with an average of −3.2˚C in Butchery 1 versus −4.6˚C and 6˚C, with an average of −4˚C, 7˚C for the Butcher 2. Conclusion: Food pathogens and opportunists can survive on the surfaces of cold chain equipment in butcher shops and therefore pose a risk of cross-contamination.

Subject Areas

Microbiology
1. Introduction

Cross-contamination of food with pathogens in the retail environment is a significant public health problem that contributes to an increased risk of foodborne illness [1]. The World Health Organization estimates that 1800 million episodes of diarrhea and 3 million deaths of children under 5 occur each year worldwide, mainly due to contaminated food. Foodborne pathogens cause both acute and chronic illnesses [2]. Food contamination with microorganisms can occur at any stage of the process, from food production to consumption, and can be the result of environmental contamination.

The internal faces of cold chain devices can be contaminated by pathogenic microorganisms of food origin [3]. Although the low temperature delays food spoilage, but even sub-freezing at a temperature of around 7˚C does not prevent the multiplication of all microorganisms, therefore refrigerated food is subject to spoilage by molds, yeasts and bacteria.

Existing global food safety standards help improve and standardize manufacturing practices, good hygienic practices, risk analysis, critical control point, for safer processed foods [4], are not necessarily implemented in the food industry around the world. In addition, there is insufficient information regarding the assessment of food safety practices, foodborne illnesses and the microbial load on food contact surfaces, including internal surfaces of appliances. The butcher’s cold chain is on a regular basis.

Hence the interest to deepen the knowledge on the current state of butchers, includes bacteriological analysis of food samples and environmental risk assessment and training of staff, in order to improve the quality microbiological of internal surfaces of cold chain devices. We therefore sought to identify the microorganisms forming the flora and to determine the level of microbial contamination of the internal surfaces of devices in the cold chain of butcher shops.

2. Methodology

We carried out a comparative cross-sectional descriptive study in two butchers in the city of Lubumbashi. Haut-Katanga province in the Democratic Republic of Congo (DRC) is over a period from June 29 to July 11, 2020, i.e. two weeks. Thirty (30) samples were taken from the internal surfaces of 10 devices in the cold chain of butcher shops accessible with the consent of the promoter. The sample was taken under aseptic conditions. Samples were taken from the bottom and from two surfaces using the dry sterile swabs and sent to the laboratory for analysis within one hour of collection. Different culture media were prepared according to the recommendations of the manufacturers. To obtain the isolated bacte-
ria the samples were inoculated on CLED Agar, mannitol salt agar and Mac Conkey Agar. **Microbiological analysis:** To obtain the fungal isolates, the samples were inoculated on Sabouraud Dextrose Agar (SDA). The initial identification of the isolated bacteria was based on their cultural and morphological characteristics. Further identification was by biochemical characteristics using standard procedures. **GRAM coloring:** A thin smear from a 24-hour culture was made on a fat-free, heat-fixed slide by passing the slide quickly over a Bunsen burner flame after air drying. The prepared smear was then flooded with crystal violet solution for about one minute, rinse with tap water; followed by mordanting with Lugol (iodine-iodine solution); spread the Lugol and leave to act for 1 minute, rinse with tap water; after d colorless with alcohol (+acetone); pour the alcohol or an alcohol-acetone mixture drop by drop onto the slanted slide, and monitor the discoloration (30 seconds). Rinse with tap water; the smear was counterstained with Safranin for 1 minute, washed with tap water; Air dry the slide; Observe with a drop of objective 100 (×1000) immersion oil. Color purple signifies a gram positive organism while pink to red signifies gram negative. **Catalase test:** Place on a glass slide one or two drops of hydrogen peroxide at 10 volumes. Using the taper of a Pasteur pipette, take a fragment of a colony and dissociate the culture in hydrogen peroxide. The release of bubbles immediately indicated a positive test while it was negative when no bubbles were formed.

**Interpretations**

**Statistical analysis:** The microbiological results obtained after incubation were entered using MS Excel 2016 software; processed and analyzed with Excel 2016 and Epi info software version 7.2.2.6. For the analysis and interpretation of the data, the following calculations and statistical tests were used: frequency, mean and standard deviation, the Fischer Exact test and the Mann-Whitney/Wilcoxon test.

**3. Results**

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**3.1. Level of Contamination of Surfaces and Isolated Microorganisms**

**Table 1** shows that the points of sampling butchery 1 have a highest contamination level to 87% relative compared to the butchery 2 which has a frequency of 46.67% of contaminated areas. There is a significant difference between the contamination of the internal surfaces of the cold chain devices between butchery I and II 0.02508 (<0.05). Concerning the microorganisms isolated from the internal surfaces of the cold chain devices, we observed a predominance of bacteria at 75%, and molds were in second position at 25%. The bacterial contamination and
fungal were respectively distributed in the following manner depending on whether the slaughter 1 and 2: 77% and 23% against 71% and 29% (Table 2). The distribution of bacteria and molds is greater than 1000 CFU/cm² or 1 Log_{10} CFU/cm² on all surfaces sampled. The level of hygiene/cleaning of the internal surfaces of refrigerators and refrigerated cabinets was respectively unacceptable and distributed as follows depending on whether it was butchery 1 and 2: 89% and 33% against 83% and 56% (because > 1 Log_{10} CFU/cm²), and require effective cleaning/disinfection. In terms of percentage (Table 3), *Staphylococcus aureus* represent 54% of the germs counted and against 7% of *Aspergillus fumigatus*. (Butcher I) and *Staphylococcus aureus* represent 43% of the counted flora, *Klebsiella planticola* (29%) and 14% respectively for *Aspergillus fumagitus* and *Fusarium spp.*. The average counts of the microorganisms detected on the internal surfaces of the cold chain devices are presented in Table 4, as follows depending on whether

**Table 1.** Distribution of the overall contamination of surfaces by cleaning assessment.

| Structure       | Contamination of surfaces |       |       |
|-----------------|----------------------------|-------|-------|
|                 |                           | No    | Yes   |
|                 | Frequency | Percentage (%) | Frequency | Percentage (%) |
| Butcher I       | 2         | 20            | 13      | 87            |
| Butcher II      | 8         | 53            | 7       | 47            |

*p = 0.02508.

**Table 2.** Distribution of microorganisms isolated from internal surfaces of cold chain devices.

| Isolated microorganisms | Butcher I    |       | Butcher II   |       |
|-------------------------|--------------|-------|--------------|-------|
|                         | Frequency | Percentage (%) | Frequency | Percentage (%) |
| Bacteria                | 10        | 77            | 5         | 71            |
| Molds                   | 3         | 23            | 2         | 29            |

**Table 3.** Global distribution of microbial strains isolated from internal surfaces of cold chain devices.

| Isolated microorganisms | Percentage (%) |       |       |
|-------------------------|----------------|-------|-------|
| *Staphylococcus*        |                |       |       |
| *Staphylococcus aureus* | 54             | 43    |       |
| *Staphylococcus albus*  | 23             | 0     |       |
| *Klebsiella planticola* | 0              | 29    |       |
| *Molds*                 |                |       |       |
| *Aspergillus fumagitus* | 7              | 14    |       |
| *Aspergillus niger*     | 8              | 0     |       |
| *Fusarium spp.*         | 8              | 14    |       |
Table 4. The average of the microbiological analyzes compared to the sampling points.

| Flora                | Average ± Etype | Butcher I | Butcher II |
|----------------------|-----------------|-----------|------------|
| *Staphylococcus*     | 3.47 ± 0.5      | 0.80 ± 2.21 |            |
| *Klebsiella planticola* | -              | 0.51 ± 1.57 |            |
| *Fusarium spp.*      | 0.27 ± 1.04     | 0.26 ± 0.97 |            |
| *Aspergillus*        | 0.53 ± 1.68     | 0.27 ± 1.04 |            |

*p = 0.7490.

it was butchery 1 st 2 (in log10 CFU/cm²): *S. aureus and albus* 3.47 and 0.80, *aspergillus* fumagitus and niger 0.53 and 0.27 for fumagitus; *fusarium* 0.27 and 0.26. *Klebsiella planticola* 0.51. The difference between the averages of microbial contamination of the internal surfaces of devices in the cold chain of butcheries is not significant. With a p = 0.7490 (>0.05).

3.2. Operating Temperature of Cold Chain Devices

The temperature of the devices of the observed Butcher I ranged from −2°C to 6°C, with an average temperature of −3.2°C. The temperature ranges of the cold chain devices observed (Butcher 2) varied between −4.6°C and 6°C, with an average temperature of −4.7°C.

3.3. Cleaning/Disinfection Procedure

The answers of the people questioned reveal that the cleaning of the internal surfaces of the cold chain devices is carried out every morning before the start of activities and the opening for the sale of food in the two butcheries, and at the end of each. Day to restore the surfaces to a clean state in the butcher’s shop I; The sponge (cloths), squeegee and buckets are the cleaning tools most often used in both structures. The cleaning/disinfection products used contained acetic acid and bleach as active ingredients.

4. Discussion

4.1. Surface Contamination Level

The sampling points of Butchery 1 have the highest level of contamination at 87% compared to Butchery 2 which has a frequency of 46.67% of contaminated areas. There is a significant difference between the contamination of the internal surfaces of the cold chain devices between butchery I and II 0.02508 (<0.05). Otubassey *et al*., 2017 found that 100% of refrigerators inspected show bacterial contamination, 32% fungal contamination and 8% contamination by parasitic organisms.

According to the indications of the service note DGAL/SDSSA/N2007-8275 France on the microbiological criteria applicable to carcasses of slaughter animals and poultry, and of the guidelines relating to the surface controls of the material
in slaughterhouses and in cutting plants slaughter animals and poultry. Our results are unsatisfactory because they are greater 1000 CFU/cm² or 1 Log_{10} CFU/cm² on all surfaces sampled [5]. The level of hygiene/cleaning the internal surfaces of refrigerators and refrigerated cabinets was not acceptable, respectively, and distributed as follows depending on whether it was the slaughterhouse 1 and 2: 89% et 33% against 83% et 56%, and require effective cleaning/disinfection. This result is similar to one of Zickrick K. et al., who found that 17.2% of domestic refrigerators in Germany contained > 1000 CFU/m³ [6]. And are not consistent with those found by Altunatmaz et al. who found psychrotrophic bacteria in all air sampled to be <200 CFU/m³ [7].

At the butcher I, *S. aureus* account for 54% of enumerated bacteria, followed by *S. albus* (23%) of *A. Niger and Fusarium spp* (8%), *A. fumagitus* (7%); on the other hand, at the level of butchery II, *S. aureus* represent 43% of the counted flora, *K. planticola* (29%) and respectively 14% for *asp. fumagitus and fusarium spp*. Macias-Rodriguez et al., In 2013; Kampmann et al., In 2008; Otu-Bassey et al., In 2017 reported that the bacterial genera isolated in decreasing order of frequency were: *Staphylococcus aureus* 27.3%, *Escherichia coli* 20.2%, *Shigella spp*. 13.0%, *Pseudomonas aeruginosa* 11.9%, *Aeromonas hydrophilia* 8.3%, *Salmonella typhi* 5.9%, *Klebsiella pneumonia* 5%, *Streptococcus pyogenes* 4.7% and *Proteus mirabilis* 2.3%. The fungal organisms isolated were *C. andida albicans* 54%, *penicilium spp* 43.2% and a *spergilus flavus* 2.7% while the parasites detected were *Entamoeba histolytica/dispar* 50% and *Ascaris lumbricoaides* 50% [6] [7] [8].

The averages of the count of microorganisms detected on the internal surfaces of the cold chain devices are presented as follows depending on whether it was butchery 1 or 2 (in log_{10} CFU/cm²): *Staphylococcus aureus and albus* 3, 47 and 0.80, *aspergillus fumagitus and niger* 0.53 and 0.27 for fumagitus; *fusarium* 0.27 and 0.26. *Klebsiella planticola* 0.51. The difference between the averages of microbial contamination of the internal surfaces of devices in the cold chain of butcheries is not significant. With a p = 0.7490 (>0.05). Otu-Bassey et al., In 2017; Oluwafemi et al., in 2015. Have isolated other species of bacteria and fungi in home refrigerators, *Staphylococcus aureus*, *Escherichia coli*, *Bacillus subtilis*, *Enterobacter spp.*, *Klebsiella spp.* and *Shigella spp.*, *Aspergillus flavus*, *A. niger*, *A. fumigatus*, *Saccharomyces cerevisae* and *Rhizopus spp.* [9] [10] [11].

These unwanted organisms may have entered cold chain devices from unwashed raw foods, poorly packaged foods (meats, eggs, and milk), dirty hands, through an open refrigerator door, fluctuating temperatures, and surfaces of unsuitable containers placed in the refrigerator.

### 4.2. The Operating Temperature of Cold Chain Devices

Temperature is one of the main factors in controlling food quality and safety due to its influence on microbial growth rates. Despite the fact that the low temperature can reduce the growth rate of many species of microorganisms, it has been reported that psychrotrophic microorganisms can thrive under normal refriger-
ration temperatures [12]. For Butcher I, the temperature range varied between 
−2˚C and 6˚C, with an average temperature of −3.2˚C. For Butcher II, the tem-
perature range varied between −4.6˚C and 6˚C, with an average temperature of 
−4.7˚C.

The WHO [13] recommends the temperature at which food is stored at a 
maximum of 5˚C in the refrigerator [14], reported that a drop in temperature is 
important to retard the growth of pathogens and other microorganisms during 
the shelf life of the product, especially after the product is purchased and under 
the responsibility of producers or consumers of food.

4.3. Cleaning/Disinfection Process

To eliminate a maximum of microorganisms, it is advisable to carry out a clean-
ing with a detergent, in order to eliminate the major part of the soils present on 
the surfaces, then to disinfect.

The antimicrobial properties of acetic acid are attributed to their acetic acid 
and citric acid content, respectively [15]. It is believed that these organic acids 
cross the cell membrane of bacteria and that the release of protons (H +) in the 
cells results in their death [16]. Bleach, a bactericide, a fungicide tested in the 
laboratory according to the standard EN 13697 after a contact time of 15 min-
utes. Used at a high concentration, bleach may attack the surface to which it 
has been applied and thus cause crevices favorable to the implantation of micro-
organisms.

The responses on the frequency of cleaning/disinfection obtained in our sur-
vey are similar to those obtained in the INCA survey carried out by [17] [18] in 
the survey on consumer practices and hygiene recommendations. In fact, 56% 
and 64% of respondents said they cleaned their refrigerator at least once a quarter.

The sponge (cloths), squeegee and buckets are the cleaning tools most often 
used in both structures. Using the same sponge to clean both cold chain devices 
and other surfaces promotes the circulation of microorganisms in the structure. 
In addition, a frequently used and thus almost always wet sponge is therefore very 
loaded with microorganisms [19]. Sponge washing can therefore constitute a mode 
of contamination of cold chain devices [20] [21]. The use of squeegees and a hand 
brush, which are aggressive for surfaces, should be avoided as they may cause 
scratches and thus promote the retention of dirt and micro-organisms.

5. Conclusion

Food pathogens and opportunists can survive on the surfaces of cold chain equip-
ment in butcher shops and therefore pose a risk of cross-contamination. Through 
our results, we have established that butcher’s cold chain devices for food storage 
are not as sterile. Enterobacteriaceae as well as molds constitute the resident fl-
ora of the internal surfaces of the devices of the cold chain in our environment, 
hence the importance of the control of the operating temperature, regular effec-
tive cleaning/disinfection regimes which must be communicated butcher shops
in order to ensure the quality and safety of the surfaces made available to the production function and the sale of foodstuffs.

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Conflicts of Interest

The authors declare no conflicts of interest.

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