Efficacy of low-pressure foam cleaning compared to conventional cleaning methods in the removal of bacteria from surfaces associated with convenience food

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Abstract

Background: Food borne illnesses and food poisoning are cause for concern globally. The diseases are often caused by food contamination with pathogenic bacteria due largely to poor sanitary habits or storage conditions.

Objectives: Prevalence of some bacteria on cleaned and sanitised food contact surfaces from eight convenience food plants in Gauteng (South Africa) was investigated with the view to evaluate the efficacy of the cleaning methods used with such food contact surfaces.

Methods: The microbial load of eight convenience food manufacturing plants was determined by sampling stainless steel food contact surfaces after they had been cleaned and sanitised at the end of a day’s shift. Samples were analysed for Total Plate Count (TPC), *Escherichia coli*, *Salmonella* species, *Staphylococcus aureus* and *Listeria* species.

Results: Results showed that 59 % of the total areas sampled for TPC failed to comply with the legal requirements for surfaces, according to the Foodstuffs, Cosmetics and Disinfectants Act (< 100 cfu.cm⁻²). *S. aureus* and *Salmonella* were not detected, but *Listeria* was detected in 23 % and *E. coli* in 1.3 % of the samples. Fifty percent (50 %) of the plants applied conventional cleaning methods for cleaning and sanitation and 50 % used the low-pressure foam (LPF) method. There was significant difference (P ≤ 0.05) between the mean TPC values of the conventional cleaning method (14 358.82) compared to that of LPF method (2 386.51) but no significant difference (P > 0.05) in terms of *Listeria* species isolates obtained from both cleaning methods. The LPF method proved to be the superior cleaning option for lowering TPC counts.

Conclusion: Regardless of cleaning method used, pathogens continued to flourish on various surfaces, including dry stainless steel, posing a contamination hazard for a considerable period depending on the contamination level and type of pathogen. Intensive training for proper chemical usage and strict procedural compliance among workers for efficient cleaning procedures is recommended.

Key words: Disinfectants, food, *Listeria* spp., sanitation, Total plate count

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Introduction

Public health concerns with food safety and food poisoning emerged in Britain in the 1880s, following the first indication that acute gastric illness was caused by a specific organism. The word ‘sanitation’ is derived from the Latin word ‘sanitas’, which refers to health. In the food industry, this means the application of a regime to provide safe, wholesome food processed in a clean environment by healthy workers who pose a limited health threat to the end-consumer. The South African food industry is changing rapidly and ready-to-eat products (convenience foods) internationally are becoming more popular. According to Brand, “It is confirmed that there is a significant growth in the convenience food market.” This is due to the fact that the continuously fluctuating South African economy and the ever-increasing cost of living, has resulted in more people now working than ever before in order to survive as well as to sustain the average household income. Consequently, because of work pressure people seem to have less time to cook, thereby increasing the patronage of food convenience stores. Therefore, the need to assess the safety of food is increasingly being recognised. Unfortunately the attitude and/or knowledge required to practice effective hygiene control is inadequate or even lacking in some food businesses. The bacteria responsible for food poisoning can proliferate quickly in food, especially in warm and moist conditions. A single bacterial cell on an item of food left out of the fridge overnight could produce

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millions of bacteria by the morning – sufficient to cause foodborne illness.39

A recent study conducted in restaurants determined the incidence of a number of significant foodborne pathogens and the general hygiene status, as estimated by TPCs and total coliform counts (TCCs), on the interior surfaces of domestic refrigerators.25 Some of the microbial isolates were found to survive and grow while refrigerated or under mild temperature abuse conditions. Such pathogens (pschycrophiles) may transfer to food in domestic fridges and multiply until they reach clinically significant numbers.19 These risks are of particular concerns in relation to ‘ready-to-eat’ foods, which will not receive any further processing before consumption.25 A study by Chao et al.10 revealed that counts of Listeria were 13.4% higher on delicatessen foods than on cooked foods investigated during their study. Moreover, non-spore-forming bacteria might be able to withstand dry conditions on surfaces for an extensive period.31 Surveillance of bacteria has also become increasingly important due to the increase in international food trade.36 In addition, microbiological hazards could stem from the introduction of new techniques for mass production as well as the rapidly growing, widespread distribution of foodstuffs.1

Organisms such as Total Aerobic Mesophiles, E. coli, S. aureus, Listeria species and Salmonella species normally isolated from meat, dairy and vegetable products have been universally utilised as indicators to determine the level of contamination on contact surfaces after they have been cleaned and sanitised.5 Though the South African legal limit42 for TPC on food contact surfaces is < 100 cfu.cm², current legislation does not make provision for maximum counts related to E. coli, S. aureus, Listeria species or Salmonella species on food contact surfaces. E. coli is not considered a serious foodborne hazard in countries with high sanitary standards and practices. Water contaminated with human sewage may lead to contamination of foods, as can handling by infected food handlers resulting in the infrequent isolation of these organisms from such food products.27

S. aureus food poisoning usually occurs rapidly and is in many cases acute, depending on the individual's susceptibility to the toxin produced by this microbe, the amount of contaminated food eaten, the amount of toxin in the food ingested and the general health of the victim.27 Staphyloccoci exist in air, dust, sewage, water, milk and food or on food equipment, environmental surfaces, humans and animals. Humans and animals are the primary reservoirs of the bacteria,27 which are present in the nasal passages and throats and on the hair and skin of over 50% of healthy individuals. Although food handlers are usually the main source of food contamination in food poisoning outbreaks, equipment and environmental surfaces can also be sources of contamination.27

Salmonella food poisoning appears to be rising in the United States as well as in other industrialised nations.4 Salmonella enteritidis isolations from humans have risen dramatically in the past decade, particularly in the Northeast USA (sixfold or more), and the increase in human infections is spreading south and west, with sporadic outbreaks occurring in other regions.8 Salmonella inhabit the intestinal tracts of humans and other animals, including birds, and in any raw food of animal origin, such as meat, poultry, milk and dairy products, eggs, seafood and on some fruits and vegetables.27

The aim of this study was to identify whether selected organisms are present on cleaned and sanitised food contact surfaces from eight convenience food plants in Gauteng (South Africa), to relate the bacterial count to the legal limit and to compare and evaluate the cleaning methods used with such food contact surfaces.

Materials and methods

Sampling protocol

This study was conducted among a sample of convenience food manufacturers supplying convenience food products (ready-to-eat lunch foods) to retail outlets in the Gauteng area of South Africa. Eight outlets (amounting to 20% of the medium to large manufacturing plants that supply the retail industry in the region) were chosen because they mainly focus on preparing ready-to-eat lunch meals. Foods manufactured included ready-to-eat salads, sandwiches, fruit salads, filled pancakes or omelettes and cocktail burgers. The management staff at each of the manufacturing plants sampled, granted permission to conduct the survey and subsequent interviews. None of the premises were Hazard Analysis Critical Control Point (HACCP) certified. The various food manufacturers used different chemical suppliers and the chemical companies had different levels of cleaning technology, therefore, different levels of cleaning methods were applied.27 Fifty percent of the outlets used traditional methods such as manual cleaning (brush and bucket) and were supplied by local chemical manufacturers. The brush and bucket method refers to physical energy being carried out by people, using pressure and movement.
on surfaces. A brush with hot water and detergent are used for cleaning. Sufficient contact time is allowed to disinfect the surfaces with chemicals. Thereafter the chemical residue is removed by a final rinse and the surfaces are allowed to air dry. International companies supplied the remainder of the plants and they used more modern technologies (for instance, low-pressure foam cleaning systems). Low-pressure foam cleaning involves a mechanized system of low pressure in combination with specialized foam detergents and disinfectants in the following steps - pre-wash, application of foam detergent, exposure, rinsing, disinfecting and final finishing wash.

Stainless steel food contact surfaces at the manufacturing plants were sampled by means of swabs after they had been cleaned and sanitised at the end of each day’s shift. The samples were collected in accordance with local health legislation. To ensure that the usual level of cleaning applied to contact surfaces occurred in all of the manufacturing plants, workers were not informed of the planned sample collection. The sampling was performed on days that required no overtime work, as overtime would potentially decrease the time allocated for cleaning the contact surfaces. Thus, adequate time was available for cleaning and sanitising of all contact surfaces. All samples were analysed on the same day. A total of 477 microbiological samples (Table 1) were collected according to the SABS swab technique and all analyses were performed at least twice.

### Table 1. Total plate count (TPC) and bacterial content of samples collected from cleaned and sanitised convenience food contact surfaces at eight food manufacturing plants.

<table>
<thead>
<tr>
<th>Food Manufacturing Plants</th>
<th>^1TPC</th>
<th>^2Salmonella species</th>
<th>^3Listeria species</th>
<th>^4Staphylococcus aureus</th>
<th>^5Escherichia coli</th>
<th>Total/plant</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>25</td>
<td>3</td>
<td>17</td>
<td>3</td>
<td>10</td>
<td>58</td>
</tr>
<tr>
<td>2</td>
<td>41</td>
<td>5</td>
<td>27</td>
<td>5</td>
<td>8</td>
<td>91</td>
</tr>
<tr>
<td>3</td>
<td>21</td>
<td>3</td>
<td>14</td>
<td>3</td>
<td>8</td>
<td>49</td>
</tr>
<tr>
<td>4</td>
<td>30</td>
<td>4</td>
<td>20</td>
<td>4</td>
<td>12</td>
<td>70</td>
</tr>
<tr>
<td>5</td>
<td>14</td>
<td>3</td>
<td>10</td>
<td>3</td>
<td>7</td>
<td>37</td>
</tr>
<tr>
<td>6</td>
<td>23</td>
<td>3</td>
<td>16</td>
<td>3</td>
<td>9</td>
<td>54</td>
</tr>
<tr>
<td>7</td>
<td>32</td>
<td>4</td>
<td>21</td>
<td>4</td>
<td>11</td>
<td>72</td>
</tr>
<tr>
<td>8</td>
<td>19</td>
<td>2</td>
<td>14</td>
<td>2</td>
<td>9</td>
<td>46</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>205</strong></td>
<td><strong>27</strong></td>
<td><strong>139</strong></td>
<td><strong>27</strong></td>
<td><strong>79</strong></td>
<td><strong>477</strong></td>
</tr>
</tbody>
</table>

2. Method SWJM 42 (Swift Micro Laboratories)
5. ISO method 16649-2 (International Organisation for Standardization, 2001)

### Microbiological analysis

The TPC samples were analysed using the conventional pour plate technique specified in ISO Method 4833 (International Organisation for Standardization, 2003). To isolate other bacteria, the samples were first enriched using a non-selective enrichment broth (NSEB) and incubated at 35-37°C for 16-18 h. To isolate *E. coli*, solid growth media as stipulated in ISO Method 16649-2 was used. *S. aureus* was isolated using the spread plate technique by spread inoculating (0.1 ml) of the culture broth from NSEB onto the surface of a dried pre-poured Baird-Parker agar (BPA) medium in a Petri-dish and incubating at 37°C for 24 h. Coagulase production among *S. aureus* isolates was determined using ISO Method 6888-1. *Listeria* species were isolated culturing the NSEB broth culture onto solid media using the conventional technique described in ISO 11290-1. Similarly *Salmonella* species were also isolated by culturing the NSEB culture onto solid media using the Malthus’s method. All bacteria were characterized using morphological and biochemical characteristics before serological confirmation. Presumptive positive colonies were confirmed using latex agglutination kits as follows: Salmonella latex kit (SWJM 42) (Swift Micro Laboratories) for *Salmonella* spp., Latex agglutination
test kit (Hardy Diagnostics) for *E. coli*, Staph latex kit (Remel) for *S. aureus* and DR1126 Listeria test kit (Oxoid) for *Listeria* spp.

**Data analysis**
The results were analysed with assistance from Corrie Uys, statistician at the Cape Peninsula University of Technology’s Centre for Postgraduate Studies and were presented as frequencies and percentages in tables and graphs.

**Results and Discussion**
**Microbiological results for convenience food contact surfaces**

**Total Plate Count**
The sample size (205 samples) proved to be 95% accurate as a representative sample of the population when using the Confidence and Error method with a tolerance of 5%. Plant 1 showed the highest TPC values of $2.07 \times 10^5$ cfu.cm$^{-2}$. Although all plants sampled had areas where there was no bacterial growth, all exceeded the legal limit of $< 100$ cfu.cm$^{-2}$ average TPC (Figure 1).

![Figure 1. Comparison between the Total Plate Count (TPC) average and the standard deviation versus the legal limit of $< 100$ cfu.cm$^{-2}$ across manufacturing plants](image)

Results also showed that the average bacterial count and normal data distribution or standard deviation also considerably exceeded the legal limit. This may indicate insufficient cleaning and disinfection, since effective cleaning processes should yield a significantly reduced or zero TPC.

Table 2 presents a summary of the total samples taken and shows the compliance with the legal limit ($< 100$ cfu.cm$^{-2}$ for TPC). Overall, 84 of the 205 TPC samples (41%) complied with the legal requirement, whereas 121 of the 205 samples (59%) did not comply.

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Escherichia coli, Staphylococcus aureus, Salmonella species and Listeria species

Table 2 shows the total samples investigated for the presence or absence of E. coli, S. aureus, Salmonella species and Listeria species.

Table 2. Compliance with the legal limit (< 100 cfu.cm$^{-2}$ for TPC) and the prevalence of bacteria on the cleaned and sanitised convenience food contact surfaces in the eight food manufacturing plants

<table>
<thead>
<tr>
<th>Test</th>
<th>No. of samples (n)</th>
<th>COMPLIANCE</th>
<th>NON-COMPLIANCE</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Absent</td>
<td>0-100 cfu.cm$^{-2}$</td>
</tr>
<tr>
<td>TPC</td>
<td>205</td>
<td>84</td>
<td>41</td>
</tr>
<tr>
<td>Escherichia coli</td>
<td>80</td>
<td>79</td>
<td>98.7</td>
</tr>
<tr>
<td>Staphylococcus aureus</td>
<td>27</td>
<td>27</td>
<td>100</td>
</tr>
<tr>
<td>Salmonella species</td>
<td>27</td>
<td>27</td>
<td>100</td>
</tr>
<tr>
<td>Listeria species</td>
<td>139</td>
<td>107</td>
<td>77</td>
</tr>
</tbody>
</table>

N/O = not observed

Results showed that 1.3% of the samples investigated yielded E. coli isolates. Although S. aureus and Salmonella species were absent on these surfaces, 23% of all the samples yielded Listeria species. These pathogens pose a considerable threat to the safety of convenience food consumers.\(^{29}\) Listeria species especially L. monocytogenes are of great concern to retailers in South Africa. The presence of this organism immediately is a reason for concern and the retailer’s procurement divisions will act strongly against any supplier who supplies products that indicate the presence of Listeria species.\(^{3,32}\) The organism is associated with listeriosis, which can be life threatening and causes septicaemia, meningitis and even stillbirth in high-risk populations.\(^{29}\) Factory environments are not sterile and L. monocytogenes can be found anywhere in the natural environment.\(^{46}\) The bacteria are easily introduced into food production and processing facilities through many routes and may establish colonies on food processing equipment. They also have the properties needed to survive refrigeration temperatures and resist freezing.\(^{20,33}\) Based on recent estimates from the Centres for Disease Control and Prevention of the United States, the annual incidence of death caused by listeriosis is about eight times greater than the mortality due to E. coli O157:H7 infections.\(^{37}\)

Many commonly used disinfecting or sanitising agents, such as quaternary ammonium compounds, chlorine and iodophors have been shown to be effective against Listeria species in suspension, but organic material reduces the activity of disinfectants.\(^{46}\) Subsequently, food products may become contaminated during processing. L. monocytogenes may grow in biofilms that protect them against environmental stress and can be isolated from surfaces after they have been cleaned and disinfected. They can also adhere to all of the materials commonly used in the food industry. In many food processing environments, the slow flowing water rich in supplies of nutrient suspensions provide conditions that favour bacterial adherence.\(^{6}\) Therefore, several challenges - including their increased resistance to sanitisers and their ability to grow at the low temperatures found in...
ready-to-eat processing plants - exist in controlling the proliferation of *L. monocytogenes*.

Although there are limited specifications available on bacteria in food in South Africa, the norm is that all pathogens should be absent. Listeria species are very common and can be found almost anywhere in the environment. As such, with food processing and manufacturing, there is the potential to introduce the organism continuously. The challenge is to direct efforts to prevent the growth and establishment of *Listeria* within the plant through having appropriate controls such as sanitation, proper manufacturing practices and employee training.

*E. coli* was found on one sample in Plant 1 only, whereas the most positive *Listeria* samples were found in Plants 1 and 7. Plant 1 also showed the highest average bacterial count, followed by Plant 7. It appears that the overall hygiene standard of the plant influences the presence of *Listeria*.

Statistical comparisons of cleaning methods
Statistical analyses were used to determine which cleaning method (conventional cleaning methods or low-pressure foam cleaning) is most suitable for application in the convenience food industry. And to conclusively demonstrate the efficacies of the cleaning methods, SABS 1853 approved sanitisers that kill 99.9% of microorganisms were used in 7 out of the 8 plants. The expected outcome was that all samples should be close to zero or at least comply with the legal requirements of < 100 cfu.cm^-2. The samples were taken from identical surfaces to ensure uniform results. Results indicated that the LPF system is more effective, as proved by the lower mean of the TPC found on convenience food contact surfaces in which this method was employed for cleaning purposes (Table 3). A statistical significant difference (*P* ≤ 0.05) was found in the TPC means of the cleaning methods (Table 3). The LPF method consistently proved to be the better cleaning option for reducing TPCs. The presence of *Listeria* species on convenience food contact surfaces was statistically evaluated with no significant difference (*P* = 0.812) found between the cleaning methods used.

<table>
<thead>
<tr>
<th>Plants</th>
<th>N = 205</th>
<th>Mean TPC</th>
<th>Median TPC</th>
<th>Standard deviation</th>
<th>*P-value cleaning methods</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conventional method</td>
<td>1, 2, 3, 7</td>
<td>119</td>
<td>14 358.82</td>
<td>1 240.00</td>
<td>33 560.897</td>
</tr>
<tr>
<td>LPF method</td>
<td>4, 5, 6, 8</td>
<td>86</td>
<td>2 386.51</td>
<td>35.00</td>
<td>7201.980</td>
</tr>
</tbody>
</table>

*P*-values were calculated between the cleaning methods

**Conclusion**
The results highlighted the presence of high counts of bacteria, including *Listeria* spp that was detected on the sanitised or disinfected convenience food contact surfaces. Fifty-nine percent (59 %) of the TPC samples analysed exceeded the legal specification (< 100 cfu. cm^-2 for food contact surfaces) when measured against the requirements of the Foodstuffs, Cosmetics and Disinfectants Act. It is alarming that these plants use reputable chemical suppliers’ approved products but still exhibit a pathogen contamination as well as generally high bacterial counts on contact surfaces. The majority of positive samples for *Listeria* and TPC were found in Plants 1 and 7, with one sample in Plant 1 showing the presence of *E. coli*. Both of these plants made use of the conventional cleaning method. The LPF method was found to be significantly better (*P* ≤ 0.05) than the conventional cleaning method in the manufacturing plants utilising these methods, respectively. Result of this study also raises the question as to whether workers or cleaners have sufficient knowledge and/or insufficient training on how to apply the chemicals correctly to achieve the desired results. It is therefore recommended that the management of
the various plants consider the possibility of providing intensive training to the production workers and general cleaners. This study has further highlighted the fact that pathogens continue to flourish on various surfaces, including dry stainless steel, and present a contamination hazard for a considerable period, depending on the contamination level and type of pathogen.\textsuperscript{31}

References


