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The detection and quantification of food components on stainless steel surfaces following use in an operational bakery

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9 Abstract

10 Food preparation areas in commercial bakeries present surfaces for continual organic fouling. The 11 detection of retained food components and microorganisms on stainless steel surfaces situated for 12 one month in the weighing in area, pastry and confectionary production areas of a bakery were 13 investigated using different methods. Scanning electron microscopy demonstrated the morphology 14 of the material on the surfaces from all three areas, with the weighing in area demonstrating a more 15 even coverage of material. Differential staining assays demonstrated a high percentage coverage 16 of organic material heterogeneously distributed across the surfaces. Differential staining also 17 demonstrated that the amount of organic material on the surface from the confectionary area was 18 significantly greater than from both the pastry and weighing in areas. Although, UV at 353 nm did 19 not detect residual surface fouling, performance of the UV detection was optimised and 20 demonstrated that the residual organic material on the weighing in area and the pastry samples was 21 best illuminated at 510 - 560 nm, and from the confectionary area of the bakery at 590 - 650 nm. 22 ATP bioluminescence revealed the confectionary production area contained the highest level of 23 biofouling. Contact plates determined that only low microbial counts (≤ 2) CFU/cm²) were 24 recovered from the surfaces. Changes in the physicochemistry (increased hydrophobicity) demonstrated that all the surfaces were fouled (ΔG_{iwi} -26.8 mJ/m² to -45.4 mJ/m²). Fourier 25 26 Transform Infra-Red Spectroscopy (FTIR) demonstrated that all the surfaces had retained fats,

- 27 carbohydrates and proteins. This work suggests that a range of methods may be needed to fully
- 28 detect organic and microbial fouling.
- 29 Keywords: Biofouling; food; conditioning film; detection; bakery; microorganisms

30 1. Introduction

31 Operational efficiency in food processing industries is affected by the fouling of preparation 32 surfaces, which is often due to the raw food materials. Fouling of food contact surfaces occurs as 33 a result of the transfer or buildup of adsorbed organic material on food processing surfaces. If such 34 fouling becomes problematic, it may impact the overall product quality and increase the operational 35 costs of food production due to frequent shut downs for cleaning (Barish and Goddard, 2013). 36 Biofouling leads to the production of a conditioning film on the food preparation surface, which 37 can influence subsequent microbial attachment and/or biofilm formation (Whitehead et al. 2008; 38 Whitehead et al. 2010).

39 In commercial bakeries, the ingredients include flour, yeast, salt, water, and oil/fat (Chen et al. 2005), which are all ideal for the formation of organic conditioning films on the surfaces. In 40 41 addition, they are a good source of nutrients for microorganisms. Therefore, food preparation 42 surfaces in bakeries present an interface for continual soiling as the food ingredients used in 43 production process are high in organic matter content, which easily disperse to form films on the 44 surfaces. It has been reported that the organic material involved in the fouling of surfaces influences 45 substratum properties and cell attachment and microbial retention with a subsequent impact on the 46 microbial fouling of surfaces (Whitehead et al. 2008).

Stainless steel is one of the most common materials used in bakery equipment (Chen et al. 2005).
The retention of organic material on industrial surfaces leads to such surfaces becoming a potential
nutrient source for microbial growth. These surfaces will then be new sources of microorganisms
transfer, which can lead to food contamination (Whitehead et al. 2010). Bacterial contamination of
food processing equipment is a central concern owing to the potential food spoilage and
transmission of foodborne pathogens.

53 There are currently a range of detection methods (e.g. Adenosine Triphosphate (ATP) 54 bioluminescence and Ultra Violet (UV)) for use in industry that allow rapid identification of surface 55 fouling. However, previous works have demonstrated that different methods have varying range in

the limit of detection for different types of organic fouling (Whitehead et al. 2008; 2009a; 2010; 2011). Some of these methods might be optimized, for example, using UV detection the wavelength can be optimised to detect residual organic material present on the surface (Adhikari and Tappel, 1975; Whitehead et al., 2010). Optimization of such methods is dependent on the molecular configuration of organic material allows some organic residues to fluoresce Another drawback of these methods is that they do not discriminate between the amount of organic fouling and microbial load (Everard et al. 2016: Salo et al. 2008; Verran and Whitehead 2006).

63 Scanning electron microscopy (SEM) has been utilized for a number of years to visualize both 64 organic material and cells retained on surfaces (Rajab et al. 2018; Whitehead et al. 2005; Zouaghi 65 et al. 2018) as has differential staining methods (Whitehead et al. 2009b). Fourier Transform Infra-66 Red Spectroscopy (FTIR) has been used to detect the fouling retained on surfaces (Whitehead et 67 al. 2011), however it can also be used to detect bacteria retained on surfaces (Schmitt and 68 Flemming, 1998). Although these and other methods have been carried out to detect organic fouling 69 and microbial contamination in vitro, to the authors knowledge, a comparison of such detection 70 methods have not been used to determine the amount of residual biofouling on surfaces from a 71 working bakery.

The aim of this work was to compare the detection and quantification of organic fouling on surfaces
 recovered from a bakery following one month exposure.

74 **2. Methods**

A range of methods were used to determine and compare the detection of residual fouling and microorganisms on stainless steel surfaces following their *in situ* placement for one month in a working bakery which produced a variety of confectionaries.

78 2.1 Sampling site and sample substrata

A bakery in the Northwest of England, UK, served as a host site where samples of stainless steel plates were placed on site prior to surface analysis investigations. The bakery used in this work produced both confectionary and pastry products. Stainless steel plates (2 mm thick, 340 2B finish) 82 were cut using an industrial hydraulic guillotine into 200 mm x 200 mm sample pieces and placed 83 in the bakery for one month. The stainless steel sample pieces were placed in three areas of the 84 industrial bakery; the weighing in area (where bags and boxes of food material are brought into the 85 bakery and dispensed), the pastry preparation area and the confectionary production area. The 86 stainless steel surfaces were exposed to the same routine food production and cleaning processes 87 as other work surfaces in the bakery. In the bakery's cleaning regime, workbenches surfaces were 88 cleaned each evening using a wipe method with the sanitizer Aphemclen® following the 89 manufacturer's instructions (Selden, UK). Following the in situ placement, samples were cut into 90 smaller substrata (20 x 20 mm) and used for subsequent surface analysis.

91 *2.2 Determination of surface fouling.*

92 2.2.1 ATP Bioluminescence

ATP bioluminescent measurements were carried out using an ATP bioluminescent device (Hygiena, UK), operated as per the manufacturer's instructions. However, although the manufacturer recommended swabbing a 100 x 100 mm area, in this study, both a 40 x 40 mm and 100 x 100 mm surface area were swabbed and the results were compared. Results were classified according to the manufacturers guidance whereby a result of < 10 relative light units (RLU) means that the surface could be considered clean, \geq 10 to \leq 29 RLU indicated that the surface was not adequately clean and >30 RLU the surface required cleaning (n = 3).

100 2.2.2 UV detection

101 A UV lamp (Labino trac-pack) with a lamp range of 353 nm was used for UV analysis of substrate 102 surface. The optimum wavelengths of UV to illuminate the different soils were then determined by 103 visualizing soil samples with different filters (330 nm - 380 nm, 510 nm - 560 nm and 590 nm -104 650 nm) using an Epifluorescence microscope (Nikon Eclipse E600, UK). For all UV illumination, 105 the unstained samples were place under the UV and the qualitative detection of biofouling on the 106 sample was recorded (n = 5).

107 2.2.3 Epifluorescence microscopy

The substrata were visualized using Epifluorescence microscopy (Nikon Eclipse E600, UK) with
an F-View II black and white digital camera (Soft Imaging System Ltd, UK) using a Cell F Image

110 Analysis package (Olympus, UK). The percentage coverage area of the stained material was

111 measured to indicate the surface coverage of the organic material (n = 60).

112 2.2.4 Scanning electron microscopy (SEM)

Samples of the stainless steel surfaces for observation by SEM were prepared. The surfaces were immersed in 4 % v/v gluteraldehyde (Agar Scientific, UK) for 24 h at 4 $^{\circ}$ C, washed in sterile distilled water, dried and stored at room temperature, in phosphorous pentoxide (Sigma Aldrich, UK) desiccators. The samples were sputter coated at a vacuum of 0.0921 mbar, for 3 min, at 2500 V, in argon gas at a power of 18-20 mA (Polaron E5100 SEM sputter coater, UK). Images of the control and fouled substrata were obtained (JEOL JSM 5600LV Scanning electron microscope, UK) (n = 3).

120 *2.2.5 Fourier transform infrared spectroscopy*

121 The stainless steel coupons (n=5) were analysed in reflection mode using a Nicolet Continuum 122 FTIR microscope (with liquid nitrogen cooled MCT detector) fitted to a Nicolet Nexus FTIR bench. 123 The apertures were set to maximum opening, 200 µm x 200 µm. Spectra were made up of 120 scans with resolution set to 4 cm⁻¹. All spectra were converted to absorbance and baseline corrected 124 125 using a spline fit. As the sampling area was fixed at 200 µm x 200 µm, it was possible to use the 126 combined areas (A_(combined)) of the OH and CH stretching (A_(OH&CH)), carbonyl (including amide) 127 $(A_{(C=O)})$ and C-O stretching peaks $(A_{(C-O)})$ as a measure of the relative amount of organic matter, 128 as summarized by equation 1.

129

130
$$A_{(combined)} = A_{(OH\&CH)} + A_{(C=O)} + A_{(C-O)}$$
 (1)

131

Peak areas were determined using the integration tool within the Nicolet Omnic software. Theintegration limits varied from spectrum to spectrum due to differing peak width, though

approximate integration limits were as follows: $A_{(OH\&CH)}$ 3710 – 2500 cm⁻¹; $A_{(C=O)}$ 1780 – 1490 cm⁻¹ and $A_{(C-O)}$ 1780 – 760 cm⁻¹s. The baseline limits were the same as the integration limits. The averages of five replicate $A_{(combined)}$ values were reported together with the standard deviation and the percentage ratio of the standard deviation to the average value.

138 2.2.6 Differential staining of organic material and microorganisms

139 Differential staining of the sample surfaces was carried out. Ten microliters of 4', 6-diamidino-2-140 phenylindole dye (DAPI, UK) was applied to the surface and spread across the sample surface 141 using a sterile plastic spreader, followed by the addition and spread of 10 μ L of rhodamine (Sigma, 142 UK). The samples were incubated for 10 min and then thoroughly rinsed with distilled water and 143 dried in a class 2 laminar flow hood in the dark for 1 h. The samples were visualized and analysed 144 using an epifluorescent microscope (Nikon Eclipse E600, UK).

145 *2.2.6 Contact plates*

146 The contact plate method for evaluating viable microbial counts from the surface of the substrata 147 was adapted from Eginton et al. (1995). The method involved using Plate Count Agar (PCA), de 148 Man, Rogosa, Sharpe agar ((MRS), for *Lactobacillus* detection), Tryptone soya agar (TSA), Brain 149 heart infusion agar (BHIA), Sabouraud agar ((SAB), for detection of moulds and yeast) and 150 MacConkey agar (for detection of Escherichia coli) all prepared according to the manufacturer's 151 instructions (Oxoid, UK). Stainless steel plates (20 mm x 20 mm) were placed onto the agar for 1 152 min, then removed. The agar was incubated for 24 h - 48 h at 30°C to allow the bacterial and fungal 153 cultures to grow. The microorganisms recovered from the plates were isolated onto the appropriate 154 agars to obtain pure cultures (n = 3).

155 *Physicochemistry of surfaces*

- 156 The physicochemistry of the surfaces was characterized according to Wilson-Nieuwenhuis et al.
- 157 (2017). Contact angles (θ) using HPLC grade water (BDH, UK), ethylene glycol or
- 158 dioodomethane (Alfa Aesar, USA) were measured with a MobileDrop goniometer (Krüss
- 159 GMBH, Germany). Both advancing and receding angles were determined, with five

160 measurements of each chemical on each sample taken (n = 10). Fresh coupons were used for each 161 solvent to ensure there was no cross contamination of solvents on the surfaces. The methods of 162 van Oss et al. (1988) and van Oss and Giese (1995) was used for calculating the surface energy 163 (γ_s^{SE}) of the films from these measurements, according to the following equation:

164
$$(1+\gamma_l)\cos\theta = 2\left(\sqrt{\gamma_s^{LW}\gamma_l^{LW}} + \sqrt{\gamma_s^A\gamma_l^B} + \sqrt{\gamma_s^B\gamma_l^A}\right)$$

where the subscripts *s* and *l* denote the surface energy of the solid and liquid respectively. The superscript *LW* denotes the Lifshitz-van der Waals components of the surface energy, and the superscripts *A* and *B* denote the Lewis acid and Lewis base parameters of the surface energy. The acid and base terms can be combined into the Lewis acid base (superscript *AB*) component of the surface energy:

171
$$\gamma_i^{AB} = 2 \sqrt{\gamma_i^A \gamma_i^B}$$

172

Subsequently the overall surface energy was calculated as the sum of the Lifshitz-van der Waalsand Lewis acid base components:

175 $\gamma_i = \gamma_i^{LW} + \gamma_i^{AB}$ 176 (4)

177 The components of the surface energy were then used to assess the hydrophobicity, or Gibbs free 178 energy of attraction between the surface and liquid (surface energies are denoted by subscript w), 179 and were calculated using the following²²:

180
$$\Delta G_{sw} = -2\left(\left(\sqrt{\gamma_s^{LW}} - \sqrt{\gamma_l^{LW}}\right)^2 + 2\left(\sqrt{\gamma_s^a \gamma_s^b} + \sqrt{\gamma_l^a \gamma_l^b} - \sqrt{\gamma_s^a \gamma_l^b} - \sqrt{\gamma_l^a \gamma_s^b}\right)\right)$$

181

182 2.2.7 Statistical analysis

(5)

(2)

(3)

183 The results were averaged and the standard deviation of the results calculated, and applied as error 184 bars. T-tests were carried out to determine the significant variance in the data whereby p < 0.05185 was considered significant.

186 **3. Results**

187 The ATP bioluminescence investigation revealed the presence of ATP on all the surfaces in all the 188 areas analyzed (Figure 1). There were no significant differences determined between the amount 189 of organic material demonstrated on the control and the weighing in area (40 mm x 40 mm control 190 = 5 RLU, weighing in area = 3.6 RLU; 100mm x 100 mm control = 7.7 RLU, weighing in area = 191 2.6 RLU). This suggests that both the control surfaces and the coupons deposited for one month in 192 the weighing in area can be considered as clean. The stainless steel samples from the pastry area 193 and confectionary area were significantly different. The pastry area demonstrated that the surfaces 194 were adequately clean (40 mm x 40 mm 13.3 RLU, 100mm x 100 mm = 13 RLU), whilst the 195 confectionary area demonstrated the greatest amount of fouling (40 mm x 40 mm = 57.3 RLU, 196 100mm x 100 mm = 59 RLU). The confectionary area demonstrated RLU levels 15 times that of 197 the weighing in area and 4 times that of the pastry indicating a highest organic matter presence. 198 Regardless of the area size, there was no significant difference between the bioluminescence results 199 of the 40 mm x 40 mm and the 100 mm x 100 mm areas investigated.

The investigations using the UV illumination carried out with the standard lamp at 353 nm revealed no differences observed between the control surfaces and the surfaces recovered from the three areas of the bakery (Figure 2a-d).

In order to optimize the UV illumination, the surfaces were exposed to different wavelengths of UV, so that the residual organic material on the surfaces were illuminated. The results demonstrated that residual organic material on the surfaces from the weighing in area and the pastry area were best illuminated by UV in the 510 - 560 nm.For the confectionary area of the bakery the best results were achieved in the 590 - 650 nm (Figure 3). And so? Do we have a percentage coverage to say

208 which surfaces were the most fouled, according to this detection method?

Scanning Electron Microscopy (SEM) was used to visualise the residual materials on the surfaces. SEM results demonstrated that the control surfaces were clean (Figure 4a), whereas the surfaces recovered from the weighing in area looked to have organic material distributed across the surface (Figure 4b). Images from the pastry and confectionary areas demonstrated the presence of particulate fouling (Figures 4 c and d). These results indicate that fouling occurred on the surfaces in the three areas of the bakery.

215 Following the differential staining assays, it was revealed that organic material was 216 heterogeneously dispersed across all surfaces used in the investigation (Figures 5b-d), when 217 compared to the control (Figure 5a). Indeed, samples taken from all areas of the bakery indicated presence of organic material in the grain boundaries (Figures 5b-d). However, the surfaces from 218 the pastry area showed the greatest amount of coverage (71.0 %) when compared to the weighing 219 220 in area (65.2 %) and confectionary areas (59.0 %) (Figure 5e). The organic material on surfaces 221 from the confectionary area was significantly different (p < 0.05) from both the pastry and 222 weighing in areas.

Following the contact plate assays onto the different types of agar, low numbers of bacteria, yeasts and moulds were isolated from the surfaces tested (< 2 CFU/cm²) (Figure 6). No *Lactobacillus* or *Escherichia coli* were detected from any of the samples taken from the bakery on the MRSA or MAC agar contact plates respectively.

227 The physicochemical results demonstrated that, following use in the bakery, all the surfaces became 228 more hydrophobic ($\Delta Giwi$ comprised between -26.8 mJ/m² and -45.4 mJ/m²) compared to the control ($\Delta Giwi = -18.0 \text{ mJ/m}^2$) (Table 1). The γs^+ decreased from 0.55 mJ/m² for the control 229 surfaces, to 0.2 mJ/m^2 for the other surfaces. The γs^{LW} values slightly increased from 34.4 mJ/m² 230 on the control surfaces to 35.6 mJ/m² – 39.4 mJ/m² on the used bakery surfaces. The ys^{AB} values 231 232 decreased from 6.2 mJ/m² on the control surfaces to 2.3-3.1 mJ/m², as did the ys⁻ values from 17.4 mJ/m^2 on the control surfaces to 8.2 $mJ/m^2 - 14.4 mJ/m^2$ on all the surfaces recovered from the 233 234 bakery.

Surface contamination was assessed using FTIR. Representative spectra from each of the three
sampling areas from within the bakery are shown in Figure 7. Surfaces from all three areas of the
bakery demonstrated significant common peaks that can be summarized as follows:

- i) hydrogen bonded N-H and O-H stretching $(3700 \text{ cm}^{-1} 2600 \text{ cm}^{-1})$ from adventitious water, sugars/carbohydrates (from flour) (O-H) and proteins (N-H),
- 240 ii) C-H stretching vibrations $(3000 \text{ cm}^{-1} 2800 \text{ cm}^{-1})$ mainly due to fatty alkyl chains
- 241 iii) Carbonyl stretching (1780 cm⁻¹ 1490 cm⁻¹) due to fatty esters and amide groups of
 242 proteins,
- iv) C-O bending 1170 cm⁻¹ 800 cm⁻¹, due to ether linkages in sugars / carbohydrates and
 esters in fats.

245 The absorption bands in the spectra obtained from the confectionary area indicated that there was 246 contamination, potentially from flour (C-O correlation), sugar (C-O correlation) and fat (ester 247 carbonyl correlation). There were also some proteins, indicated by amide I and II bands, present in 248 all the samples, which could be due to gelatin (setting agent) residues. The pastry area showed 249 contamination with diverse mixtures of fats and proteins. The weighing in area also showed a 250 diversity of contaminants with thick fat and flour contamination. The average combined OH/NH stretch, carbonyl, protein amide and C-O stretching peak areas ($\overline{A_{(combined)}}$) from the FTIR spectra 251 252 together with the standard deviation values for the data provide insight into the differences in levels 253 and uniformity of contamination within a particular area of the bakery (Table 2). The weighing in 254 area showed by far the highest level of fouling and also showed the most non-uniform fouling, as 255 indicated by the very high SD relative to the average value. The pastry and confectionary areas had 256 substantially lower levels of fouling; combined peak area values were broadly similar and the SD 257 values relative to the average indicated greater uniformity of fouling compared with the weighing 258 area.

- **4. Discussion**
- 260 Biofouling of industrial surfaces is of a major concern. When a food product comes into contact

261 with a stainless steel work surface, the contact surface will become contaminated with organic 262 material, and potentially microorganisms (Whitehead et al. 2011). Cleaning procedures aim to 263 remove organic material and microorganisms, but there is concern regarding organic material and 264 microorganisms that are retained on surfaces. There are numerous ways to detect organic and 265 microbial fouling on a surface. However, there is little information comparing the methods on 266 surfaces taken from a food industry. As a result, it is difficult to choose the best method to assess 267 the cleanliness of a surface in a food industry. In this study, the detection of both organic material 268 and microorganisms using different detection/quantification methods has been assessed on surfaces 269 recovered from a bakery, following one month exposure.

ATP bioluminescence has been widely used for the detection of microbial contamination and food residues in the food industry (Everard et al. 2016), providing a real time estimate of total surface cleanliness, including the presence of organic debris and microbial contamination. In this work, the ATP reading in the confectionary area was five times greater than that of the pastry area. It may be that on the confectionary surface, since sugar is water soluble and is easy to remove, oils are difficult to remove, and protein is very difficult to remove (Corrieu et al. 1981), that ATP was recovered from the surfaces with the most soluble organic component.

277 The use of UV has been shown to be a reliable, quick and cost effective method of accessing 278 hygiene after cleaning processes in both closed (Salo et al. 2008) and open surfaces (Verran and 279 Whitehead 2006). The need for rapid industrial detection methods such as UV is important in 280 determining the efficiency of cleaning and disinfection against organic soil and cell retention. UV 281 (353 nm) can be used for the detection of residual cells and soiling on industrial surfaces, although 282 (as with ATP) no distinction is made between the two components (Whitehead et al. 2009b). Using 283 the regular 353 nm UV lamp, the level of contamination was too low on the surfaces recovered 284 from the bakery to demonstrate residual fouling. However, using the UV with different wavelength 285 filters, it was demonstrated that the residual organic material were best illuminated in the 510 - 560 286 nm for the weighing in area and pastry areas, and in the 590 - 650 nm for the weighing in areas.

287 This occurs because the molecular configuration of organic material allows some organic residues 288 to fluoresce when illuminated by UV (Adhikari and Tappel, 1975). This is in agreement with work 289 carried out previously, whereby surfaces with residual fats were best illuminated using this method 290 (Whitehead et al. 2008). This result also corresponds to the surfaces in this work becoming more 291 hydrophobic, and non-polar as demonstrated from the physicochemistry assays. Furthermore, in 292 agreement with our work, Abban et al. (2014) demonstrated that one can change filters on the lamp 293 so that alternative wavelengths suitable for the various material surfaces are used to ensure that 294 residues on different material surfaces could be better visualized and thus the confidence in the 295 validation of the surface hygiene is increased.

296 SEM has been utilized for a number of years for the visualisation of cell:substratum interactions 297 (Rajab et al. 2018; Whitehead et al. 2005; Zouaghi et al. 2018). It was demonstrated that the control 298 surfaces were clean, whereas the surfaces recovered from the weighing in area looked to have 299 organic material distributed across the surface. Images from the pastry and confectionary areas 300 demonstrated the presence of particulate fouling, which is more reminiscent of protein and 301 carbohydrate fouling (Whitehead et al. 2010). However, the weighing in area demonstrated a 302 surface coverage which was more similar to the residual material left by fat deposits. This 303 corroborates with the results from the weighing in area surfaces, which was demonstrated to have 304 increased γ_s^{LW} measurements, and a more hydrophobic surface.

The development of novel staining systems has allowed quantitative and separate measurement of 305 306 cells and food soils retained on surfaces (Whitehead et al. 2009b). The differential staining results 307 demonstrated that the organic material was heterogeneously dispersed across all the surfaces used 308 in the investigation, and that it was particularly evident in the grain boundaries of the stainless steel. 309 Additionally, bacteria were not detected which correlates with the microbial counts obtained... 310 However, this could be because the areas analysed were much smaller than those covered using the 311 contact plate method. The pastry area showed the greatest amount of coverage when compared to 312 the weighing in area and confectionary areas.

313 The microbial investigation of the surfaces revealed an extremely low presence of microbial 314 contamination on the surfaces incubated in the bakery. The microbial data from the contact plates 315 was corroborated by the differential staining and SEM results, both of which indicated no incidence 316 of bacteria on the surfaces. This suggests that although organic material was retained on the 317 surfaces, this did not enhance microbial retention. The number of bacteria detected using the 318 manual counts was not detected using the differential stain, but this could be because the contact 319 plates used a much larger area than the differential staining method. Thus, the level and type or 320 organic material deposited on a surface and its effect on microbial retention merits further 321 investigation.

322 The physicochemical results demonstrated that following use in the bakery, all the surfaces became more hydrophobic compared to the control with increased ys^+ and ys^{LW} values. The ys^{AB} values 323 324 decreased as did the ys values on all the surfaces recovered from the bakery. This suggests that 325 following use, the surfaces became more electron accepting, and apolar. Previous work has 326 demonstrated that when compared to a pristine surface, in situ increases or decreases in the surface parameters γs^{AB} , γs^+ (the electron acceptor), and γs^- (the electron donor) suggest the presence of 327 retained surface soil. In agreement with these results changes in $\Delta Giwi$ and γs^{LW} suggest the 328 329 presence of certain oils and fats retained on the surface (Whitehead et al. 2009a) and this was 330 supported by the FTIR results. Further, Zouaghi et al. (2018) demonstrated that a lower surface 331 energy was shown to be an asset against whey protein deposits on stainless steel surfaces.

FTIR can be used to identify molecular species on a surface. FTIR has been used to detect milk traces in food (Cristina et al. 2016) and determine the impact of food treatments on the reconformation of allergens (Gomaa and Boyce, 2105). The FTIR spectroscopy of the sample areas revealed non-uniform contamination. Quantification of the amount of organic material on the surfaces, using the $\overline{A_{(combined)}}$ values (Table 2), revealed a different ranking of contamination level to the organic coverage and ATP measurements. The weighing area had the highest $\overline{A_{(combined)}}$ value, and hence showed the most prolific contamination. This result correlated with the physicochemistry (ΔG_{iwi} value) result and the SEM images.

340 Conclusions

The different microscopic investigations demonstrated particulate material on the surfaces from all three areas. UV detection was optimised and ATP bioluminescence revealed the confectionary production area contained the highest level of biofouling. Contact plates determined low microbial counts and demonstrated the advantages of screening larger surface areas for microbial contamination. All the fouled surfaces increased in hydrophobicity, which corresponded with the UV and FTIR results. This work suggests that a range of methods may be needed to detect organic and microbial fouling.

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| _ | ⊿Giwi | γs^{LW} | γs^{AB} | γs^+ | γs⁻ |
|------------------|-------|-----------------|-----------------|--------------|------|
| Control | -18.0 | 34.4 | 6.2 | 0.5 | 17.4 |
| Weighing in area | -45.4 | 38.2 | 2.3 | 0.2 | 8.2 |
| Pastry | -26.8 | 35.6 | 2.9 | 0.2 | 14.4 |
| Confectionary | -36.3 | 39.4 | 3.1 | 0.2 | 11.3 |

434 Table 1. Physicochemistry (mJ/m^2) results of the surfaces following removal from the bakery site.

- 437 Table 2. Average combined OH/NH stretch, carbonyl, protein amide and C-O stretching
- 438 absorbance peak areas ($\overline{A_{(combined)}}$) and respective standard deviation and SD as % on the
- 439 average values (n=5 unless otherwise stated)

| Sampling area in bakery | $\overline{A_{(combined)}}$ | SD | SD as % on the average |
|-------------------------|-----------------------------|--------|------------------------|
| Confectionary | 9.1 | 3.9 | 43 |
| Pastry | 13.5 | 12.9 | 96 |
| Weighing in area | 78.6* | 100.2* | 127* |



Figure 1. ATP bioluminescence measurements indicating microbial presence on the stainless steel surface of the three areas of the bakery using, (where routine cleaning pass mark = 30)



463 Figure 2. UV of stainless steel substrata obtained from the a) control surface b) weighing in area,





- 467 Figure 3. Organic matter coverage of stainless steel surfaces upon UV illumination from a) control,
- b) weighing in area, c) pastry area d) confectionary area of the bakery



472 Figure 4. SEM images of stainless steel surfaces obtained from the bakery a) control, b), weighing

473 in area c, pastry area and d) confectionary area (x 3500 magnification)







Figure 5 Images showing organic matter coverage of stainless steel surfaces from a) control, b) weighing in area, c) pastry area d) confectionary area of the bakery and e) percentage organic material coverage of stainless steel surfaces sampled from the industrial bakery









490 Figure 6. The microbial count obtained from Stainless steel substratum using contact agar plates a) 491 control b) weighing in area, c) pastry and d) confectionary areas PCA = Plate count agar, MRSA = 492 de Man, Rogosa, Sharpe agar, MAC = MacConkey agar, SAB = Sabouraud agar, TSA = Tryptone 493 soya agar and BHIA = Brain heart infusion agar.





497 Figure 7. FTIR spectra of three representative samples taken from the a) weighing in area, b)
498 confectionary and c) pastry areas. Note that the peak/baseline disturbance around 2340 cm⁻¹ is
499 due to atmospheric CO₂.