1 2	Comparative Assessment of qPCR Enumeration Methods that Discriminate Between Live and Dead <i>Escherichia coli</i> O157:H7 on Beef
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20 Abstract

21 Quantitative Polymerase Chain Reaction (qPCR) is a molecular method commonly used to 22 detect and quantify bacterial DNA on food but is limited by its inability to distinguish between 23 live and dead cell DNA. To overcome this obstacle, propidium monoazide (PMA) alone or with 24 deoxycholate (DC) was used to prevent dead cell detection in qPCR. qPCR methods were used to 25 detect strains of *Escherichia coli* O157, which can cause infection in humans with an infectious dose of less than 10 cells. A 5 strain E. coli O157:H7 cocktail was inoculated onto beef steaks and 26 27 treated with interventions used in meat facilities (lactic acid (5%), peroxyacetic acid (200 ppm) or 28 hot water (80°C for 10 s)). Treatment of PMA or PMA + DC was applied to samples followed by 29 DNA extraction and quantification in qPCR. RNA was also quantified in addition to conventional 30 plating. For lactic acid intervention, qPCR DNA quantification of E. coli O157:H7 yielded 6.59± 0.21 and 6.30 ± 0.11 log gene copy #/cm² for control and lactic acid samples, respectively and after 31 32 treatment with PMA or PMA + DC this was further reduced to 6.31 ± 0.21 and 5.58 ± 0.38 , 33 respectively. This trend was also observed for peroxyacetic acid and hot water interventions. In 34 comparison, RNA quantification yielded 7.65 \pm 0.13 and 7.02 \pm 0.38 log reverse transcript/cm² 35 for rRNA control and lactic acid samples, respectively, and for plating (LB), 7.51 ± 0.06 and 6.86 $\pm 0.32 \log \text{CFU/cm}^2$, respectively. Our research determined that treatment of PMA + DC in 36 37 conjunction with qPCR prevented dead cell DNA detection. However, it also killed cells injured 38 from intervention that may have otherwise recovered. RNA quantification was more laborious and 39 results had higher variability. Overall, quantification with conventional plating proved to be the 40 most robust and reliable method for live EHEC detection on beef.

41 Keywords: Propidium Monoazide, Deoxycholate, qPCR, Detection, Beef, Escherichia coli

42 O157:H7

43 **1. Introduction**

44 Molecular detection methods have become a mainstay in assessment of microbiological 45 composition and activity within microbial ecosystems, including food safety assessment, as they 46 are faster and more selective than labor-intensive and time-consuming conventional plating 47 methods (Ge and Meng 2009). Quantitative polymerase chain reaction (qPCR) is one of the most 48 commonly used methods of gene quantification, and is a validated method for pathogen detection 49 in food (Health Canada 2018; Wong and Medrano 2005). However, qPCR is limited in that it does 50 not discriminate between DNA from live and dead cells in a food sample, which may lead to false 51 positive results and lead to unnecessary recalls of food products (Bae and Wuertz 2009). Current 52 methods for pathogen detection use an enrichment step to prevent detection of dead cells (Ge and 53 Meng, 2009). Enrichment of samples also reduces the detection limit from > 1000 cells / g (qPCR 54 only) to less than 1 cell per 400 g (qPCR after enrichment); however, an enrichment increases the 55 time required for detection and does not allow quantitative results to be obtained.

56 Monoazide DNA binding dyes can enter dead cells with compromised membranes, intercalate 57 with DNA bases and form covalent bonds upon exposure to light (Takahashi et al. 2017; Taskin 58 et al. 2011). This modification of DNA prevents its amplification in qPCR. Ethidium monoazide 59 (EMA) has previously been investigated for this purpose, but was found to cause significant DNA 60 loss from viable cells after extraction (Flekna et al. 2007; Nocker et al. 2006a). Propidium 61 monoazide (PMA) penetrates into cells with compromised membranes without affecting viable 62 cells with intact membranes (Pan and Breidt 2007). However, PMA is excluded from dead cells 63 with intact membranes and therefore PMA-qPCR may overestimate the number of viable cells 64 (Løvdal et al. 2011; Nocker et al. 2007; Pacholewicz et al. 2013). To better select for live cells in 65 qPCR, a sodium deoxycholate treatment prior to PMA application can enhance its penetration due

to the emulsifier's ability to permeabilize intact membranes of dead cells (Yang et al. 2011). While *E. coli* is highly tolerant to bile salts, the effect of deoxycholate on injured cells is unknown. An
alternative is quantification of RNA using qPCR as an indicator of cell viability as it degrades
more rapidly upon cell death than DNA, with mRNA degrading more rapidly than rRNA (Keer
and Birch 2003). However, the low stability of RNA also increases the risk of degradation of the
target molecule during sample preparation.

72 Ruminants including cattle are a major reservoir of enterohaemorrhagic E. coli (EHEC) 73 (Croxen et al., 2013); accordingly, foodborne illness caused by EHEC has often been linked to the 74 consumption of contaminated beef (Blagojevic et al. 2012; Rangel et al. 2005). Interventions 75 implemented in beef packing plants to control EHEC on carcasses include lactic acid and 76 peroxyacetic acid washes as well as hot water or steam application (Bacon et al. 2000). E. coli 77 O157, the prototype serogroup of EHEC, is regulated in Canada and the US and current methods 78 of detection of *E. coli* O157 in food include enrichment in conjunction with qPCR. However, 79 qPCR does not differentiate between live cells and cells killed the use of interventions, limiting its 80 use when both live and dead cells are present. This investigation aimed to compare four different 81 PCR-based detection methods to determine which method most accurately quantifies live cells of 82 E. coli O157, in comparison to viable plate counts as reference method for bacterial viability. 83 Reverse-transcription (RT-)qPCR of mRNA and rRNA as well as PMA or PMA and deoxycholate 84 treatment prior to DNA quantification in qPCR were evaluated. Quantification experiments were 85 performed on beef steaks inoculated with a cocktail of E. coli O157:H7 strains, and treated with 86 lactic acid, peroxyacetic acid or hot water as a decontamination treatments.

87 2. Material and Methods

88 **2.1. Preparation of Inoculum**

Five Shiga toxin (stx) negative *E. coli* O157:H7 (02-0628, 02-1840, 02-0304, 02-0627, 00-3581; Luciano et al. 2011) were grown individually overnight in 9 mL of Luria-Bertani (LB; Oxoid, Mississauga, Ontario, Canada) broth in 15 mL glass tubes at 37°C and 250 rpm. The next day, cells of each strains were harvested by centrifugation, cell pellets were washed twice with 9 mL of 0.1% (w/v) peptone water (Difco, Becton-Dickinson, Sparks, Maryland, USA) and then combined to create a 5 strain cocktail of *E. coli* O157:H7.

95 **2.2. Primers**

Quantification of DNA in qPCR was done with the species specific primers URL-301 (5'-TGT 96 97 TAC GTC CTG TAG AAA GCC C-3') and URR-432 (5'-AAA ACT GCC TGG CAC AGC AAT 98 T-3') that target *uidA* coding for β -glucuronidase activity of *E. coli* (Bej et al. 1991). For 99 quantification of rRNA in reverse transcription qPCR (RT-qPCR) primers targeting partial 16S 100 rRNA, HDAf (5'-ACT CCT ACG GGA GGC AGC AGT-3') and HDAr (5'-GTA TTA CCG CGG 101 CTG CTG GCA C-3') were used (Walter et al. 2000, Wang et al. 2013). For quantification of 102 mRNA in RT-qPCR, primers targeting glyceraldehyde-3-phosphate dehydrogenase were selected: 103 gapAf (5'-GTT GAC CTG ACC GTT CGT CT-3') and gapAr (5'-ACG TCA TCT TCG GTG 104 TAG CC-3'). While not all primers were species specific, all primers amplified the respective 105 target in E. coli and the beef samples were essentially free of other bacteria after decontamination 106 of the surface.

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2.3. Preparation of Beef Steaks for Intervention and Control Samples

For each trial, fresh beef (eye of round) was purchased from a local grocery store. It was then boiled in water for 2 min to decontaminate surface, the cooked outer layer (2 mm) was removed with a sterile knife and then the clean beef cut was sliced into 2 cm thick steaks with surface area of 88 cm². One mL of the 5 strain cocktail of *E. coli* O157:H7, corresponding to an inoculum of

 10^7 to 10^8 cfu/cm², was inoculated onto the top surface of a beef steak in a biosafety cabinet which 112 113 was immediately spread evenly across the surface of the steak with a sterile plastic spreader. Steaks 114 with inoculum were left to air-dry in the biosafety cabinet for 30 min to allow for attachment. 115 Three trials were performed per intervention and for each trial, eight steaks were inoculated with 116 the 5 strain *E. coli* O157:H7 cocktail: four for application of interventions (2 x 0 min and 2 x 30 117 min sampling) and four for inoculated control (no application of intervention; $2 \ge 0$ min and $2 \ge 0$ 118 30 min sampling). The sampling times were selected to allow for a comparison of the effect of 119 different interventions on cell viability, injury and recovery after immediate application and after 120 a 30 min holding time. One steak was left uninoculated as a control sample per trial to confirm 121 that the steak did not contain detectable bacteria or bacterial nucleic acids, where no intervention 122 was applied. The control steak was immediately placed in 99 mL of 100 mM PBS buffer (includes 123 buffering agents monopotassium phosphate and disodium phosphate) in a stomacher bag, 124 manually massaged for 2 min and suspension plated on LB and Violet Red Bile (VRB, Oxoid) 125 agars. Plates were incubated at 30°C for 24 h but all counts were below the detection limit of 100 126 CFU/g, corresponding to fewer than 2 colonies per plate. Samples of one mL were kept for DNA 127 qPCR analysis; similar to plating, quantification yielded negative result denoted by either 128 amplification at or above 35 Ct with species specific primers (*uidA*) for DNA qPCR or 10 Ct above 129 last sample's Ct value for non-specific primers (HDA, gapA) used in both RNA RT-qPCR 130 experiments.

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2.4. Lactic Acid or Peroxyacetic Acid Intervention

Prior to application of interventions, inoculated and control steaks were placed in separate
aluminum trays (23 x 15 x 1.5 cm). A 5% (w/v) lactic acid (Sigma-Aldrich, Mississauga, Ontario,
Canada) or 200 mg/L peroxyacetic acid (Inspexx, Ecolab Inc., St. Paul, Minnesota, USA) solution

was prepared and connected to the equipment used for application of antimicrobials; this equipment has been previously described in detail (Youssef et al. 2012). The aluminum trays containing steaks were placed on the stainless steel conveyor belt that carried trays under a spray nozzle, exposing steaks to a 5% lactic acid spray or 200 mg/L peroxyacetic acid spray at a volume of 0.4 mL of fluid per cm² steak surface. The steaks were sampled immediately (0 min) or after 30 min of holding at room temperature.

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2.5. Hot Water Intervention

To simulate spraying with hot water, which is used in beef processing as a pathogen intervention method, a 4 L beaker was filled with 2.5 L of sterile water and brought to 80°C on a hot plate. Once the target temperature was reached, steaks were submerged individually for 10 s using sterile stainless steel tongs and immediately removed and sampled at 0 or 30 min.

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2.6. Sampling of Steaks after Intervention

147 Each steak was placed in a sterile stomacher bag (VWR International, Radnor, Pennsylvania, 148 USA) with 100 mL of buffer. PBS was used to neutralize lactic acid or hot water intervention 149 samples and neutralizing buffer (includes buffering agent monopotassium phosphate and reducing 150 agent sodium thiosulfate) was used for peroxyacetic acid intervention-treated samples to neutralize 151 acidulant as well as quench oxidant. Steaks in buffer were then massaged manually in stomacher 152 bags for 2 min. From sample suspension, 15 x 1 mL aliquots were transferred to separate 2 mL 153 transparent Eppendorf tubes. Tubes were centrifuged at 10,000 x g for 5 min, supernatants 154 discarded, cells washed in 1 mL peptone water, centrifuged and resuspended in 300 μ L peptone 155 water. Each tube was designated to one of 5 different sampling groups for subsequent treatment 156 application, in triplicate: DNA extraction (control; no PMA or deoxycholate), PMA treatment and 157 DNA extraction, PMA with deoxycholate treatment and DNA extraction, RNA extraction, plating

158 (Figure 1). Samples for RNA extraction were further washed with RNAprotect Bacteria Reagent 159 (Qiagen, Mississauga, Ontario, Canada) according to the manufacturer's instructions to stabilize 160 and protect RNA from degradation, the resulting pellet was kept frozen (-20°C) until use.

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2.7. PMA and Deoxycholate Treatment

162 For PMA treatment of samples, a 20 mM stock solution of PMA (Biotium, Inc. Hayward, 163 California, USA) was stored in dark at 4°C until use. For each sample suspension (300 μ L) in 2 164 mL Eppendorf tubes, 1.5 µL of 20 mM PMA was added and vortexed thoroughly. Tubes were 165 placed on ice horizontally in the dark for 5 min followed by exposure to a 650-W halogen lamp 166 for 2 min, held 20 cm above the tubes, while slowly rocking samples back and forth to ensure uniform exposure. Samples were then removed from light, centrifuged at 8,000 x g for 3 min, the 167 168 supernatant was discarded, and the cells were washed with peptone water. The cell pellet was 169 stored at -20°C until further analysis.

170 For samples treated with both PMA and deoxycholate, deoxycholate was applied to samples 171 before PMA treatment using a protocol previously optimized by Yang et al. (2011). A 1% (w/v) 172 solution of deoxycholate (Sigma-Aldrich) was stored at -20°C until use. Sample suspensions (300 173 μ L) in 2 mL Eppendorf tubes were centrifuged at 8,000 x g for 3 min, and the supernatants were 174 discarded. The pellets were resuspended with 300 µL 1% deoxycholate and incubated in a rocking 175 water bath at 37°C for 30 min. After incubation, the suspension was treated with PMA as described 176 above.

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2.8. Enumeration of E. coli O157:H7 on Beef Steaks

178 Viable cell counts were enumerated by surface plating on the day of treatment. Ten-fold 179 dilutions were prepared in sterile 0.1% peptone water from original sample suspensions of beef

180 steaks in stomacher bags with buffer, and 100 μ L of dilutions estimated to provide 30-300 colonies 181 per plate was used to manually plate onto LB and VRB agar. Plates were incubated at 30°C for 24 182 h and enumerated next day. Uninjured cells (capable of reproduction without repair of cellular 183 injury) were enumerated on VRB agar, a medium which prevents the growth of injured cells with 184 compromised outer membrane due to the presence of bile salts (Gänzle and Vogel, 2001). All 185 viable (live) cells (capable of reproduction, including cells that require repair of cellular injury 186 prior to reproduction) were enumerated on LB agar, a non-selective medium (Scheusner et al. 187 1971).

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2.9. DNA Extraction and Quantification in qPCR

189 DNA was extracted from samples (DNeasy blood and tissue kit; Qiagen) following instructions 190 for Gram-negative bacteria with modifications. Cells suspended in buffer ATL (supplemented with 191 proteinase K) were incubated for 2 h, and DNA was eluted from mini columns twice with 100 µL 192 of butter AE. For quantification in qPCR, primers URL-301 and URR-432 were used to target 193 uidA gene (Sigma-Aldrich). qPCR master mix was prepared as follows per reaction: 12.5 μ L of 194 SYBR green (Qiagen), 2.5 µL each forward and reverse primer, and 2.5 µL nuclease free water, 195 with 5 μ L template DNA added to 20 μ L of master mix for a total volume of 25 μ L per reaction. 196 A standard curve was prepared from purified PCR product of E. coli O157:H7 (00-3581) DNA 197 (QIAquick PCR Purification Kit; Qiagen), which was diluted to 100 mg/L and then further diluted 198 in serial ten-fold dilutions. Six dilutions were used to generate a standard curve in the range of 199 DNA concentrations in the samples. Nuclease free water served as negative control for all PCR 200 reactions; uninoculated steaks served as negative control for any native E. coli that may be already 201 present on steak. A 7500 Fast Real-Time PCR system (Applied Biosystems) was used for 202 quantification of DNA by one initial denaturation of template DNA and activation of Taq DNA

polymerase at 95°C for 10 min, followed by 40 cycles of denaturation at 95°C for 30 s, primer
annealing at 60°C for 1 min, extension at 72°C for 1 min. Quantification data was acquired from
7500 Software (version 2.0.5, Applied Biosystems) and analyzed. Melting curves of final products
verified no formation of non-specific product or primer dimers. Each sample was quantified in
duplicate per qPCR plate.

208 2.10.

2.10. RNA Extraction and Quantification in RT-qPCR

209 RNA was extracted from samples (RNeasy Mini Kit; Qiagen) and reverse transcribed to cDNA 210 following instructions stated in the Kit's manual (QuantiTect Reverse Transcription Kit; Qiagen). 211 Two sets of primers were selected for RNA analysis, one targeting partial 16S rRNA (HDAf and 212 HDAr; Sigma-Aldrich) and the second targeting the glyceraldehyde-3-phosphate dehydrogenase 213 (GAPDH) (gapAf and gapAr; Sigma-Aldrich). Stocks were prepared for each primer set with 214 sterile nuclease free water. Master mix recipe per reaction was prepared as follows: 12.5 µL SYBR 215 green (Qiagen), 0.5 μ L each forward and reverse primers, 10.5 μ L nuclease free water, and 1 μ L 216 sample cDNA. A standard curve was developed by extracting genomic DNA from an overnight 217 culture of E. coli O157:H7 (00-3581) using Wizard Genomic DNA purification kit (Promega). 218 Extracted genomic DNA was run in qPCR with respective primers (HDA or gapA) and the PCR 219 product was purified according to purification kit instructions (QIAquick PCR Purification Kit, 220 Qiagen) and diluted to 100 mg/L. For each primer set, this concentration was further diluted ten-221 fold (up to 10⁻⁸) to generate a standard curve. Negative controls included nuclease free water, 222 samples after treatment with genomic DNA Wipeout but before reverse transcription, and isolation 223 of RNA from uninoculated steaks. The Reverse Transcription qPCR (RT-qPCR) method was one 224 initial denaturation of template DNA and activation of Taq DNA polymerase at 95°C for 5 min, 225 followed by 40 cycles of denaturation at 95°C for 30 s, primer annealing at 60°C for 30 s, extension at 72°C for 30 s. Data was acquired from 7500 software (version 2.0.5, Applied Biosystems) and
analyzed. Melting curves of final products were used to verify that non-specific products or primer
dimers did not form. Samples were run in triplicate reaction per plate.

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2.11. Statistical Analysis

Statistical Analysis Systems (SAS Institute, Inc., Cary, NC) was used to evaluate data by two way analysis of variance (ANOVA) followed by pair-wise comparison of means from different treatment groups with Bonferroni adjustment. A probability (p) of <0.05 was considered statistically significant. Three independent trials were performed for each intervention, with triplicate samples per trial. Linear regressions were calculated in SigmaPlot 12.5.

235 **3. Results**

3.1. Viable Plate Counts and Sublethal Injury on Treated and Untreated Beef.

Cell counts for untreated, inoculated beef were on average $7.27 \pm 0.15 \log \text{CFU/cm}^2$ (mean \pm 237 238 standard deviation); the difference of plate counts between LB and VRB, which represents the sublethally injured population, on average was $0.33 \pm 0.18 \log \text{CFU/cm}^2$. After intervention with lactic 239 240 acid and hot water, the difference in plate counts on LB and VRB increased, indicating that the 241 number of injured cells had increased. However, the average proportion of injured cells was not 242 increased by peroxyacetic acid treatment compared to the control (Table 1). Comparison of post-243 intervention counts to the controls indicated that the percentage of dead cells in samples was 68%, 244 45% and 90% for meat treated by lactic acid, peroxyacetic acid and hot water, respectively.

245 **3.2.** Quantification of *E. coli* O157:H7 on Beef by qPCR After Lactic Acid Intervention

To determine the method that most accurately quantifies viable *E. coli* O157:H7 on beef steak after lactic acid intervention, five different detection methods were compared. Figures 2A and B correlate viable plate counts of *E. coli* O157:H7 after lactic acid intervention to the gene

249 copy numbers obtained by quantification of microbial DNA or RNA. Control samples represent 250 steaks that were inoculated but not treated; control quantification represents DNA quantification 251 without PMA or PMA-DC. Lactic acid intervention reduced viable cell counts by about 1 log 252 CFU/cm² (Table 1), however, this reduction was not significant. Interestingly, significant 253 reduction in the log gene copy $\#/cm^2$ of E. coli O157:H7 was determined by qPCR only in samples 254 treated with PMA combined with deoxycholate. However, reductions observed at 0 and 30 min 255 indicate that the combination treatment not only prevented amplification of DNA from cells killed 256 by intervention, but it also prevented the amplification of DNA from sub-lethally injured cells 257 when compared to plate count data. Thich explains why reductions were significant as compared 258 to plating. Accordingly, the slope of the regression line correlating cell counts and gene copy 259 numbers was < 1.0 for qPCR and PMA-qPCR quantification but > 1 for PMA-deoxycholate qPCR. 260 Quantification with RT-qPCR targeting rRNA or mRNA provided comparable results (Figure 2B).

261 **3.3.** Quantification of *E. coli* O157:H7 on Beef by qPCR After Peroxyacetic Acid

262

2 Intervention

263 Similarly, after peroxyacetic acid intervention of steaks inoculated with E. coli O157:H7, five 264 different detection methods were compared to determine the most accurate method to quantify 265 viable cells. Figures 3 A and B correlate quantification of E. coli O157:H7 on beef after 266 peroxyacetic acid intervention. Enumeration on LB agar was compared to DNA or RNA 267 quantification in qPCR or RT-qPCR, respectively. After peroxyacetic acid intervention, plate 268 counts and the number of gene copies were not reduced at 0 or 30 min sampling times. The 269 reduction of cell counts or of the copy number of nucleic acids after combination treatment of 270 PMA with deoxycholate was also not significant. This is likely because this intervention did not 271 cause the same degree of sub-lethal injury as the other two interventions.

272 3.4. Quantification of E. coli O157:H7 on Beef by qPCR After Hot Water Intervention 273 The same five different detection methods were compared to determine the most accurate 274 method to quantify live cells after hot water intervention on beef steaks inoculated with E. coli 275 O157:H7. The correlation of quantification of E. coli O157:H7 on beef after hot water intervention 276 by means of enumeration on LB agar as compared to DNA or RNA quantification in qPCR or RT-277 qPCR is presented in Figures 4 A and B, respectively. Enumeration by plating, qPCR or RT-qPCR 278 determined that the number of viable E. coli O157:H7 on steaks at both 0 and 30 min were not 279 significantly reduced by hot water intervention. However, PMA with deoxycholate treatment and gPCR quantification reduced the log gene copy $\#/cm^2$ at 0 and 30 min. The significant reduction 280 281 is likely attributed the degree of sub-lethal injury that the intervention caused, and that this 282 population of cells was sensitive to the combination of PMA with deoxycholate treatment, which 283 then led to the prevention of amplification in qPCR. Accordingly, the slope of the regression line 284 correlating plate counts to gene copy numbers was 1.26 for quantification with PMA-deoxycholate 285 qPCR but only about 0.5 for quantification with qPCR and PMA-qPCR (Fig. 4A). Quantification 286 with RT-qPCR targeting rRNA or mRNA provided comparable results with a slope of 1.1 and 1.3, 287 respectively (Fig. 4B).

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3.5. E. coli O157:H7 Recovered on LB and VRB from Beef Steaks at 30 min After

Intervention and After Treatment of Cells with PMA or PMA with Deoxycholate

qPCR quantification indicated that the combination of PMA and deoxycholate treatment permeabilized viable *E. coli* cells to PMA after intervention when compared to control and cell count enumeration data by plating. To determine whether treatment with PMA and PMA-DC affects viability of *E. coli*, the organism was enumerated after intervention treatments on beef steaks and PMA or PMA-deoxycholate treatment (Table 1). PMA did not affect the viability of *E.*

coli (Table 1). Treatment with both PMA and deoxycholate reduced the viability of *E. coli*; this reduction was significant after treatment of *E. coli* that were previously exposed to hot water washes on beef. Taken together, these data suggest that PMA-deoxycholate may permeabilize and kill *E. coli* when cells were sublethally injured by intervention treatments.

4. Discussion

300 This study compared five different detection methods for the quantification of viable E. coli 301 O157 on beef, and after intervention treatments that are currently used in the meat industry. The 302 application of lactic acid, peroxyacetic acid and hot water interventions to beef steaks created a 303 heterogeneous population of viable, sub-lethally injured, and dead cells. These populations are 304 highly suitable for testing detection methods for the ability to discriminate between live and dead 305 bacterial cells. Lactic acid and hot water interventions were more lethal to E. coli O157:H7 on beef 306 than peroxyacetic acid, which is consistent with previous studies (Castillo et al. 1998; Ellebracht 307 et al. 2005; Gill and Bryant 2000; King et al. 2005; Youssef et al. 2012). Experiments reported in 308 this study were obtained with aseptic meat samples without background microbiota and primers 309 that were not specific to E. coli O157. Equivalent results can be expected if primer sets with more 310 specific targets are used to specifically detect *E. coli* O157 or STEC in commercial meat samples.

Overall, the cell counts obtained by plating on LB agar were similar to RT-qPCR based quantification that targeted rRNA; mRNA copy numbers and DNA copy numbers were generally lower. This discrepancy reflects that mRNA is less stable than rRNA and that *E. coli* cells contain several thousand ribosomes (Keer and Birch 2003). The decrease of viable cell counts after intervention treatments was reflected by a reduced copy number obtained by RT-qPCR. However, qPCR targeting DNA did not discriminate between viable and dead cells, though plating does (Sidari et al. 2011). Furthermore, rRNA provided a more robust estimation than mRNA, which is 318 present as a lower percentage of total RNA and is more susceptible to degradation as compared to319 rRNA (Kushner 2002).

320 Enumeration of viable cells by surface plating indicated that lactic acid was more lethal than 321 peroxyacetic acid; this difference was not observed by RT-qPCR, demonstrating that conventional 322 plating provided the most accurate estimate of viable cells. Hot water intervention resulted in a 323 greater reduction in viable cells from the plate count quantification method and both RNA RT-324 qPCR quantification methods. Hot water washes acted immediately while lactic acid acted more 325 slowly. The antimicrobial effect of lactic acid is based on penetration of cells and lowering 326 intracellular pH (Alakomi et al. 2000) while heat leads to immediate disruption of membranes and 327 irreversible denaturation of proteins (Lee and Kaletunc 2002).

328 PMA treatment does not affect viability of cells (Elizaquivel et al. 2012; Nocker et al. 2006b, 329 Yang et al. 2011; this study). PMA-qPCR, reduced the gene copy numbers detected in beef samples 330 but the difference between qPCR and PMA-qPCR was small and remained unchanged by 331 bactericidal intervention treatments (Figs. 2, 3, and 4). A high number of dead cells may inhibit 332 PMA crosslinking of DNA by light and thus reduce the correlation to viable plate counts (Løvdal 333 et al. 2011; Pan and Breidt 2007; Wagner et al. 2008; Yañez et al. 2011). To counter this, it has 334 been suggested that incubating samples for a longer period of time and at higher temperatures to 335 increase PMA penetration into dead cells (Nkuipou-Kenfack et al. 2013).

Several past studies compare PMA-qPCR to quantification of live cells of foodborne pathogens by plate counts. Løvdal et al. (2011) applied lethal heat treatment of 80°C for 15 min to *Listeria innocua* cells in broth and found large discrepancies between quantification methods: plating determined cells counts below the detection limit whereas PMA-qPCR quantified 8 log cells/mL

340 Enumeration of *Campylobacter jejuni* on raw chicken carcasses after different slaughterhouse 341 processing steps determined that PMA-qPCR consistently yielded up to a 1-1.5 log CFU/mL 342 higher cell count than plating (Pacholewicz et al. 2013). After severe heat treatment of C. jejuni in 343 chicken rinse at 95°C for 5 min, PMA-qPCR and plating both indicated that cells counts were 344 below the detection limit (Josefsen et al. 2010). Lastly, Ju et al. (2016) quantified live E. coli 345 O157:H7 on lettuce. Plating indicated that E. coli O157:H7 did not survive on leaf surfaces and 346 was reduced by 10⁷ cells within 96 h; however, RT-qPCR and PMA-qPCR indicated a reduction of 10^2 and 10^3 cells, respectively. Overall, a majority of studies indicate that PMA-qPCR can lead 347 348 to false positive results. This conforms to the observation that the structurally related propidium 349 iodide is excluded by a substantial proportion of dead cells after lethal interventions (Ulmer et al., 350 2000). Some studies defined PI or PMA excluding but non-culturable cells "viable but not 351 culturable"; however, sublethally injured cells of *E. coli* are readily recovered by cultivation media 352 containing pyruvate or catalase as scavengers for oxidative stress (Afari and Hunc, 2018; Mizunoe 353 et al., 1999). We used LB as reference medium for detection of viable cells and thus did not account 354 for those cells that require extended recovery times or addition of pyruvate or catalase to recover 355 from injury.

The use of deoxycholate prior to PMA treatment was proposed to increase cell membrane permeability to permeabilize all dead cells (Yang et al., 2014). In suspensions where *E. coli* cells were killed after heat treatment with no viable cells remaining, treatment with PMA prior to qPCR significantly decreases amplification of dead cells in qPCR as compared to PMA alone (Yang et al. 2014). *E. coli* cells that were killed by 72°C or less were permeable to PMA but the addition of the deoxycholate treatment rendered a large portion of these cells permeable to the dye (Yang et al. 2011). After lactic acid intervention, membranes of dead and injured *E. coli* cells that were 363 impermeable to PMA were rendered permeable to the dye after treatment with 1% deoxycholate 364 treatment (Wang et al. 2014). Injured cells were resuscitated in nutrient rich broth for 2 h prior to 365 deoxycholate treatment which restored injured cell membrane barrier properties. This could be a 366 possible solution to injured cell sensitivity to PMA with deoxycholate treatment. However, since 367 our study was performed in a more complex system rather than in broth or buffer, this could prove 368 to be more complicated and warrants further investigation. Zhou et al. (2016) optimized the 369 concentration of deoxycholate for use with *Cronobacter sakazakii* in powdered infant formula. 370 The authors found that heat injured cells were not affected by a concentration of 0.08% 371 deoxycholate and determined that at this concentration, the accuracy of viable cell detection when 372 combined with PMA and qPCR was increased. Sodium dodecyl sulfate has also been used to aid 373 in PMA penetration into dead cells and shows little impact on cell viability. While this could offer 374 an alternative emulsifier to use with PMA to increase the accuracy of live cell detection in qPCR, 375 it still requires further research in different food systems (Dong et al. 2018, Takahashi et al. 2017). 376 Our work indicates that combination treatment of PMA with deoxycholate prevents amplification 377 of DNA from sublethally injured cells, and could lead to false negative results.

378 The inability to discriminate between DNA in live and dead cells in qPCR is a problem that 379 extends beyond E. coli O157:H7 in meat, this combination treatment could be a solution in other 380 areas of research. This includes determination of viable probiotics in the gut or viable lactic acid 381 bacteria or bifidobacteria in fermented foods (Cocolin et al. 2011; Garcia-Cayuela et al. 2009; 382 Rantsiou and Cocolin 2006; Villarreal et al. 2013). However, it is important to note that 383 concentration of treatment should be adjusted to target microorganism, as Gram-positive 384 microorganisms have different resistance to deoxycholate than Gram-negative microorganisms 385 (Nkuipou-Kenfack et al. 2013).

386 **5.** Conclusions

387 This study compared five different methods to determine the most accurate quantification of viable E. coli O157:H7 on beef steaks after intervention. PMA treatment with qPCR did not 388 389 prevent DNA amplification from all dead cells in samples after intervention and therefore this 390 treatment does not resolve the issue of false positive result in qPCR. However, it does prevent 391 amplification of some dead cell DNA and therefore increases the accuracy of viable cell 392 quantification compared to qPCR alone. In contrast, treatment of cells with PMA with 393 deoxycholate prevented amplification of DNA from all dead cells but does so at the cost of sub-394 lethally injured cells since deoxycholate supported PMA penetration into injured cells which may 395 have otherwise recovered. This makes DNA inaccessible for qPCR quantification and would lead 396 to false negative result. Both rRNA and mRNA quantification in qPCR had more variability and 397 was not as sensitive for quantifying viable cells as plate counts. While PMA with deoxycholate 398 may not be suitable for pathogen detection in food where sublethal injury of cells is of concern, it 399 could be used in other areas of research that aim to exclude all dead cells at the expense of injured 400 cells. On the other hand, if research aims to amplify all viable cells including the injured cell 401 population at the cost of also amplifying some dead cell DNA, PMA or rRNA with qPCR detection 402 methods could be used. Overall, conventional plating provided the most sensitive and reliable 403 quantification of viable E. coli O157:H7 on beef.

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Table 1. Enumeration of *E. coli* O157:H7 (Log CFU/cm²) Recovered on LB and VRB agars from Beef Steaks held for 30 min after Interventions of Lactic Acid (LA), Peroxyacetic Acid (PA) and Hot Water (HW) and after Treatment of cells with PMA or PMA with Deoxycholate.

	Intervention					
Sample	Lactic Acid		Peroxyacetic Acid		Hot Water	
	LB	VRB	LB	VRB	LB	VRB
Control	7.24 ± 0.08	6.85 ± 0.27	7.00 ± 0.40	6.70 ± 0.70	7.35 ± 0.09^{a}	7.03 ± 0.18^{a}
Intervention	6.41 ± 0.14	5.27 ± 0.69	6.48 ± 0.13	6.08 ± 0.57	6.19 ± 0.54^{ab}	5.65 ± 0.62^{ab}
Control + PMA	6.65 ± 0.30	6.45 ± 0.17	6.98 ± 0.45	6.63 ± 0.79	$7.57\pm0.16^{\rm a}$	$7.11\pm0.46^{\rm a}$
Intervention + PMA	5.91 ± 0.54	5.44 ± 0.45	6.29 ± 0.61	5.75 ± 1.17	6.16 ± 0.64^{ab}	5.88 ± 0.77^{ab}
Control + PMA + DC	6.57 ± 0.09	6.15 ± 0.70	6.41 ± 0.16	6.22 ± 0.25	6.46 ± 0.22^{ab}	6.09 ± 0.48^{ab}
Intervention + PMA + DC	5.58 ± 0.47	4.85 ± 0.94	5.70 ± 0.41	5.62 ± 0.31	$5.20\pm0.72^{\text{b}}$	$4.85\pm0.68^{\text{b}}$

Values are means \pm standard deviation, n=3. Values with superscripts are significantly different among values in the same column; those that do not share a common superscript are significantly different (*p*<0.05).



Note: Leave 1 steak uninoculated for quality control purposes. Do not apply intervention; place in buffer, massage and only sample for control plating and control DNA/qPCR analysis.

Figure 1. Experimental flow chart per trial.



Figure 2. (**A**) Correlation of cell counts on LB agar & qPCR DNA quantification of *E. coli* 0157:H7 recovered from beef after lactic acid (LA) intervention & PMA or PMA + deoxycholate (DC) treatment. Circle, control; triangle, PMA; square, PMA + DC; black, control 0 min; white, LA 0 min; dark grey, control 30 min; light grey, LA 30 min. Treatments plotted at log CFU/cm² of respective control or LA sample at 0 or 30 min. Values are means of three trials with technical repeat (n=3), standard error of means of LB \leq 0.32; DNA \leq 0.72. †Significantly different log gene copy #/cm² from Control at 0 & 30 min (*p*<0.05) *Significantly different log gene copy #/cm² from Control + PMA, Control + PMA + DC and LA at both 0 & 30 min (*p*<0.05) Correlation coefficients r² of the linear regressions were 0.94 (control), 0.88 (PMA) and 0.94 (PMA + DC). (**B**) Correlation of cell counts on LB agar & RTqPCR RNA quantification of *E. coli* 0157:H7 recovered from beef after LA intervention. Diamond, rRNA; triangle, mRNA; black, control 0 min; white, LA 0 min; dark grey, control 30 min; light grey, LA 30 min. Treatments plotted at log CFU/cm² of respective control or LA sample at 0 or 30 min; light grey, LA 30 min.

three trials (n=3) with technical repeat, standard error of means of rRNA \leq 0.68; mRNA \leq 0.65. Correlation coefficients r² of the linear regressions were 0.49 (rRNA) and 0.55 (mRNA).



Figure 3. (**A**) Correlation of cell counts on LB agar & qPCR DNA quantification of *E. coli* 0157:H7 recovered from beef after peroxyacetic acid (PA) intervention and PMA or PMA with deoxycholate treatment. Circle, control; triangle, PMA; square, PMA with deoxycholate; black, control 0 min; white, PA 0 min; dark grey, control 30 min; light grey, PA 30 min. Treatments plotted at log CFU/cm² of respective control or PA sample at 0 or 30 min. Values are means of three trials with technical repeat (n=3), standard error of means of LB \leq 0.39; DNA \leq 0.68. Correlation coefficients r² of the linear regressions were 0.83 (control), 0.87 (PMA) and 0.94 (PMA + DC). (**B**) Correlation of cell counts on LB agar & RTqPCR RNA quantification of *E. coli* 0157:H7 recovered from beef after PA intervention. Diamond, rRNA; triangle, mRNA; black, control 0 min; white, PA 0 min; dark grey, control 30 min; light grey, PA 30 min. Treatments plotted at log CFU/cm² of respective control or PA sample at 0 or 30 min. Values are means of three trials with technical repeat (n=3), standard error of means of rRNA \leq 0.35; mRNA \leq 1.12. Correlation coefficients r² of the linear regressions were 0.87 (rRNA) and 1.00 (mRNA).



Figure 4. (A) Correlation of cell counts on LB agar & qPCR DNA quantification of *E. coli* 0157:H7 recovered from beef after hot water (HW) intervention & PMA or PMA with deoxycholate treatment. Circle, control; triangle, PMA; square, PMA with deoxycholate; black, control 0 min; white, HW 0 min; dark grey, control 30 min; light grey, HW 30 min. Treatments plotted at log CFU/cm² of respective control or HW sample. Values are means of three trials with technical repeat (n=3), standard error of means of LB \leq 0.54; DNA \leq 0.93. *Significantly different log gene copy #/cm² from Control, Control + PMA, Control + PMA + DC, HW, HW + PMA at 0 and 30 min (*p*<0.05). Correlation coefficients r² of the linear regressions were 0.99 (control), 0.93 (PMA) and 0.92 (PMA + DC). (B) Correlation of cell counts on LB agar & RTqPCR RNA quantification of *E. coli* 0157:H7 recovered from beef after HW intervention. Diamond, rRNA; triangle, mRNA; black, control 0 min; white, HW 0 min; dark grey, control 30 min; light grey, HW 30 min. Treatments plotted at log CFU/cm² of respective control or a present of the sample at 0 or 30 min. Values are means of three trials with technical repeat (n=3), standard error 0 min; white, HW 0 min; dark grey, control 30 min; light grey, HW 30 min. Treatments plotted at log CFU/cm² of respective control or HW sample at 0 or 30 min. Values are means of three trials with technical repeat (n=3), standard error of means of rRNA

 \leq 1.48; mRNA \leq 1.20. Correlation coefficients r² of the linear regressions were 0.66 (rRNA) and 0.75 (mRNA).