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Ouoba, L. Irène I., Vouidibio Mbozo, Alain Brice, Anyogu, Amarachukwu ORCID: https://orcid.org/0000-0001-9652-7728, Obioha, Promiselynda I., Lingani-Sawadogo, Hagrétou, Sutherland, Jane P., Jespersen, Lene and Ghoddusi, Hamid B. (2019) Environmental heterogeneity of Staphylococcus species from alkaline fermented foods and associated toxins and antimicrobial resistance genetic elements. International Journal of Food Microbiology, 311. p. 108356. ISSN 0168-1605

http://dx.doi.org/10.1016/j.ijfoodmicro.2019.108356

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Environmental heterogeneity of *Staphylococcus* species from alkaline
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20 Abstract

Different samples of three products including Bikalga and Soumbala from 21 Burkina Faso (West Africa) and Ntoba Mbodi from Congo-Brazzaville (Central Africa) 22 were evaluated. The bacteria (400) were phenotyped and genotypically characterized 23 by Rep-PCR, PFGE, 16S rRNA and *rpoB* gene sequencing and *spa* typing. Their 24 PFGE profiles were compared with those of 12,000 isolates in the Center for Disease 25 Control (CDC, USA) database. They were screened for the production of enterotoxins, 26 susceptibility to 19 antimicrobials, presence of 12 staphylococcal toxin and 38 AMR 27 28 genes and the ability to transfer erythromycin and tetracycline resistance genes to Enterococcus faecalis JH2-2. 29

Fifteen coagulase negative (CoNS) and positive (CoPS) species characterised 30 by 25 Rep-PCR/PFGE clusters were identified: Staphylococcus arlettae, S. aureus, S. 31 cohnii, S. epidermidis, S. gallinarum, S. haemolyticus, S. hominis, S. pasteuri, S. 32 condimenti, S. piscifermentans, S. saprophyticus, S. sciuri, S. simulans, S. warneri 33 and Macrococcus caseolyticus. Five species were specific to Soumbala, four to 34 Bikalga and four to Ntoba Mbodi. Two clusters of S. gallinarum and three of S. sciuri 35 were particular to Burkina Faso. The *S. aureus* isolates exhibited a *spa* type t355 and 36 their PFGE profiles did not match any in the CDC database. Bacteria from the same 37 cluster displayed similar AMR and toxin phenotypes and genotypes, whereas clusters 38 39 peculiar to a product or a location generated distinct profiles. The toxin genes screened were not detected and the bacteria did not produce the staphylococcal 40 enterotoxins A, B, C and D. AMR genes including *blazA*, *cat*501, *dfr*(A), *dfr*(G), *mecA*, 41 mecA1, msr(A) and tet(K) were identified in CoNS and CoPS. Conjugation 42 experiments produced JH2-2 isolates that acquired resistance to erythromycin and 43 tetracycline, but no gene transfer was revealed by PCR. 44

The investigation of the heterogeneity of *Staphylococcus* species from alkaline fermented foods, their relationship with clinical and environmental isolates and their safety in relation to antimicrobial resistance (AMR) and toxin production is anticipated to contribute to determining the importance of staphylococci in alkaline fermented foods, especially in relation to the safety of the consumers.

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52 **Key words**: Alkaline fermented foods; *Staphylococcus* spp.; diversity; distribution;

53 safety; toxins; AMR

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55 **1. Introduction**

Alkaline fermented foods are products that are generally rich in proteins (30-56 50%) and their production involves a fermentation process associated with an increase 57 of pH to values as high as 10.0 (Parkouda et al., 2009; Vouidibio Mbozo et al., 2017). 58 They are mainly known in Africa and Asia, where they constitute low cost sources of 59 protein foods and play an important part in the diet of people (Parkouda et al., 2009). 60 It is generally recognized that the pH rise is the result of accrued degradation of raw 61 material proteins that leads to the release of peptides, essential amino acids as well 62 63 as alkalinizing substances such as ammonia (Kiers et al., 2000). In some cases, alkalitreatments during the production are responsible for the pH augmentation (Parkouda 64 et al., 2008). 65

Bikalga, Soumbala, and Ntoba Mbodi are alkaline fermented foods which are 66 produced through the traditional fermentation of roselle seeds (Hibiscus sabdariffa), 67 African locust bean seeds (Parkia biglobosa) and cassava leaves (Manihot esculenta 68 Crantz) respectively. Production of Bikalga and Soumbala comprises several 69 processing stages of the seeds: cleaning, boiling (about 95°C for 24-40 h), dehulling 70 (Soumbala), fermentation (48-72 h), steaming (Bikalga) and drying (Ouoba et al., 71 2008a; Parkouda et al., 2008; Parkouda et al., 2009). They are the most popular 72 traditional fermented foods used as condiments in Burkina Faso (West Africa) and are 73 74 also produced in other African countries. Ntoba Mbodi, which is popular in the Republic of the Congo (Central Africa), is made of cassava leaves and the production process 75 does not involve heating steps. After being cleaned and cut, the leaves are distributed 76 into small portions in larger leaves, wrapped and left to ferment for 2-4 days (Vouidibio 77 Mbozo et al., 2017). The fermented cassava leaves are then used to prepare sauces, 78 which are eaten with starch-based foods. In Bikalga, Soumbala and Ntoba Mbodi, pH 79

values up to 8.5, 9 and 10 have been respectively reported (Ouoba, 2015; Ouoba,
2017; Parkouda et al., 2009; Vouidibio Mbozo et al., 2017).

The key microorganisms involved in alkaline fermentations are *Bacillus* species 82 such as B. subtilis, B. licheniformis, B. amyloliquefaciens, B. pumulis, and B. 83 xylanilyticus that can degrade proteins, lipids, and carbohydrates of the raw materials 84 and bring about the nutritional guality and safety of the product (Parkouda et al., 2009). 85 However, secondary microorganisms such as lactic acid bacteria, Staphylococcus 86 spp. and various Gram-positive and Gram-negative bacteria are often isolated with no 87 88 clear indication of their role in the fermentation. The production techniques of most alkaline fermented foods are traditional, leading to variability of microbial population, 89 nutritional quality, product stability and potential safety issues. It is believed that some 90 of the secondary microorganisms are in fact contaminants from the producers, 91 processing materials and environments that have poor hygienic quality in most 92 traditional fermentation settings. Some secondary microorganisms are part of the 93 original microbiota of the raw materials that are suppressed by products released 94 during the fermentation. For example, those that are alkali sensitive will be eliminated 95 or reduced during the fermentation. 96

Staphylococci are Gram-positive non-motile cocci ubiquitously spread in 97 nature, and are commonly isolated from humans, animals, food and environmental 98 99 sources. More than 70 species and subspecies of the Staphylococcus genus have been characterized so far (LPSN, 2018) with Staphylococcus aureus being one of the 100 most studied species due to its clinical importance. Staphylococcus species are 101 102 usually divided into two groups i.e. coagulase-positive (CoPS) and coagulasenegative (CoNS) according to their ability to coagulate rabbit plasma. Various 103 Staphylococcus species have been recurrently isolated in many alkaline and non-104

alkaline vegetable based fermented foods, but their role in such fermentation is 105 unknown. However, their importance in the fermentation of meat-based products has 106 been well documented. They occur in the production of various types of sausages 107 where they constitute a part of the autochthonous microbiota or are added as starter 108 cultures in combination with lactic acid bacteria (Fonseca et al., 2013; Milicevic et al., 109 2014). Staphylococcus equorum, S. saprophyticus S. xylosus and S. carnosus are 110 the most common species found in fermented sausages, but other species such as S. 111 warneri, S. vitulinus, S. pasteuri, S. epidermidis, S. lentus, S. haemolyticus, S. 112 113 intermedius, and S. hominis can also be occasionally isolated (Milicevic et al., 2014). They have been reported to be involved in biochemical activities during meat 114 fermentation such as the degradation of proteins, lipids and peroxide as well as the 115 reduction of nitrite (Casaburi et al., 2008; Milicevic et al., 2014). These activities 116 generate compounds such as peptides, amino acids, carbonyls and volatile 117 compounds that contribute to the nutritional quality, specific organoleptic properties 118 and safety of the fermented meat products (Milicevic et al., 2014). 119

Although beneficial properties of *Staphylococcus* species in food have been 120 documented, many species are most known worldwide for causing food poising and 121 other types of clinical infections in humans and animals including e.g. bacteraemia, 122 wound infections, pyogenic lesions, and mastitis. Moreover, they present detrimental 123 124 antimicrobial resistance (AMR) profiles that make some of their infections hard to treat. With regard to toxin production in food, staphylococcal food poisoning is one of the 125 most common food-borne diseases. It results from the ingestion of heat stable 126 staphylococcal enterotoxins (SEs) produced in food by enterotoxigenic strains of 127 mainly S. aureus (Hennekinne et al., 2012). Symptoms related to ingestion of such 128 toxins are fast-acting, occurring generally within 30 min to 8 h and lasting for about 24 129

hours. Individuals generally experience vomiting, nausea, stomach cramps, and 130 diarrhoea. Severe illnesses related to staphylococcal food poisoning are rare but 131 occasionally, it can be deadly, especially in children, elderly and patients weakened 132 by a long-term illness. Staphylococcus aureus is considered one of the world's leading 133 causes of disease outbreaks associated with food consumption. Other CoPS such as 134 S. intermedius and CoNS such as S. epidermidis, S. sciuri, S. simulans, S. xylosus, 135 S. equorum, S. lentus and S. capitis have been reported to harbour genes coding for 136 SEs and to produce some SEs (Casaes Nunes et al., 2015; Crass and Bergdoll, 1986). 137 138 However, information on their real implication in staphylococcal food poising outbreaks is scarce (Breckinridge and Bergdoll, 1971; Podkowik et al., 2013). Significantly, the 139 incidence of AMR has continually increased in staphylococci, as well as in other types 140 of bacteria. This is attributed to factors such as the frequent antibiotic administration 141 to livestock, or misuse and overuse of antimicrobials. Therefore, the food chain 142 constitutes one possible route of AMR transfer (Angulo et al., 2004). Both CoPS and 143 CoNS from food origin exhibit a wide range of AMR genetic elements coding for 144 resistance to different antimicrobials such as methicillin, tetracycline, penicillin, and 145 vancomycin (Gundogan et al., 2005; Simeoni et al., 2008). The most frequently 146 encountered resistant staphylococcus is methicillin-resistant Staphylococcus aureus 147 (MRSA), which is resistant to multiple antimicrobials and is responsible for hard-to-148 149 treat infections. Thus, it constitutes a serious public health concern, as it causes thousands of deaths per year worldwide and constitutes a global economic burden. 150

The study aimed to screen the environmental heterogeneity of *Staphylococcus* species from alkaline fermented foods, their relationship with clinical isolates worldwide and their safety with regards to genes encoding antimicrobial resistance

(AMR) and toxin production. This is anticipated to elucidate their distribution andimportance in such foods, especially in relation to the safety of the consumers.

156 **2. Materials and Methods**

157 2.1 Identification of the bacteria

158 2.1.1 Phenotypic characterisation

The bacteria (400) were isolated from different samples of Bikalga (9), Ntoba Mbodi 159 (8) and Soumbala (10) collected at different markets and production sites in Burkina 160 Faso (Bikalga and Soumbala) and the Republic of the Congo (Ntoba Mbodi), 161 from areas constituting major consumers of these food products. Each sample (10 g) 162 was homogenised for 1 min in 90 ml of maximum recovery diluent (MRD, Oxoid) and 163 ten-fold dilutions (10⁻² - 10¹⁰) prepared. The dilutions (0.1 ml) were spread on nutrient 164 agar (NA; Oxoid CM0003, Basingstoke, UK) for the enumeration of total aerobic 165 mesophilic bacteria. The plates were incubated aerobically at 37°C and the count 166 For enumeration and isolation of the presumptive 167 recorded after 2 days. Staphylococcus population, the dilutions were spread on Baird Parker (BPA; Oxoid) 168 and Mannitol Salt (MSA; Oxoid) agars and the plates incubated aerobically at 37°C for 169 2 days. Bacteria isolated from the two the latter media were purified by repeated 170 streaking on NA, maintained in nutrient broth (NB; Oxoid CM001) containing 20% (v/v) 171 glycerol (Sigma G8773, Gillingham, UK) as well as on cryobeads and stored at -20°C 172 for further analyses. 173

For the phenotypic characterization, the isolates were streaked on NA and examined for colony and cell morphology, as well as tested for Gram, catalase, oxidase and coagulase reactions. The Gram reaction was evaluated using the KOH method (Gregersen, 1978) and the oxidase test using *Pseudomas fragi* as positive control and *Brochothrix thermosphacta* as negative control. The coagulase test was performed using the Staphylase test kit (Oxoid DR0595) and following the manufacturer's instructions. The cell morphology was determined by light microscopy (Nikon Model Eclipse, E400, Japan). The phenotypic screening resulted in the elimination from the study of 310 non-staphylococcal isolates which were able to grow on BPA and MSA and whose colony morphologies initially suggested they were potential *Staphylococcus* spp. Thus, 90 isolates were retained for further characterization.

186 2.1.2 Molecular typing

187 2.1.2.1 DNA extraction

The bacteria were grown on Brain Heart Infusion Agar (BHI-A) prepared by mixing BHI (Oxoid CM1135) and technical agar (Oxoid LP0013, 15g/L). After incubation at 37°C for 48 h, the DNA was extracted from purified cultures following the manufacturer's instructions using InstaGene Matrix (Bio-Rad 732-6030, Hemel Hempstead, UK).

193 2.1.2.2 Repetitive element palindromic-polymerase chain reaction (rep-PCR)

The isolates were characterized by rep-PCR using the GTG5 primer as described by Ouoba et al. (2008a). Bacteria showing the same DNA profile were clustered in the same group by visual screening and cluster analysis using the Bionumerics system (2.50, Dice's Coefficient of similarity, Unweighted pair group method with arithmetic: UPGMA; Applied Maths, Saint-Martens-Latem, Belgium).

199 2.1.2.3 Pulse Field Gel Electrophoresis (PFGE)

In order to assess the worldwide distribution of the types of *S. aureus* found in the fermented foods, PFGE of Smal-digested genomic DNA was performed according to the method proposed by the Center for Disease Control and Prevention/Pulse Net (CDC, USA) as described by Van Balen et al. (2013). The Salmonella serotype

Branderup strain H9812 digested with Xbal was used as a molecular size marker. The PFGE profiles of the *Staphylococcus* isolates were compared with those in a CDC database that contains worldwide profiles of about 12,000 *S. aureus* strains recovered from outbreaks, surveillance, reference and the environment, and submitted to the database.

209 2.1.2.4 Sequencing of the 16S rRNA and rpoB genes

The bacteria were identified by the amplification and sequencing of the 16S 210 rRNA and rpoB genes. For the 16S rRNA gene amplification, the method and the pA 211 and pE primers described by Ouoba et al. (2008a) were used. For the amplification 212 of the *rpoB* gene, primers 2491F and 3554R described by Drancourt and Raoult (2002) 213 were used with the following PCR mixture: 5 µl of 10 X PCR buffer containing 15 mM 214 of MqCl₂ (Applied Biosystems N8080160), 0.5 µl of dNTP (1.25 mM), 0.5 µl of each 215 primer (21 pmol/µl), 0.2 µl of AmpliTaq polymerase (5 U; Applied Biosystems N808-216 0160) and 41.3 µl of sterile high purity water. The PCR conditions were as follows: 217 initial denaturation at 94°C for 2 min, 35 cycles of 94°C for 30 s, 52°C for 30 s, 72°C 218 for 1 min and a final extension step at 72°C for 5 min. 219

After the amplification of both genes, positive amplicons were checked by electrophoresis and purified using the QIAquick PCR purification kit (Qiagen, Crawley, UK). The 16S rRNA and *rpoB* genes were sequenced (SourceBioscience, Cambridge, UK) using primers pD (3.2 pmol/µl) and 2491F/ 3554R (3.2 pmol/µl) respectively. The 16S rRNA gene sequences were analysed in the ezbiocloud server (Yoon et al., 2017) whereas those of the *rpoB* gene were screened in the GenBank/EMBL/DDBJ database (National Center for Biotechnology, MD, USA).

227 2.1.2.4 *Spa* typing

A multiplex PCR method and primers recommended by the European 228 Reference Laboratory for antimicrobial resistance (EURL-AR, 2012), and that allows 229 at the same time the detection of the spa gene and the S. aureus methicillin resistance 230 genes mecA, mecC was used. The method includes the use of a PCR mixture 231 containing 2.5 µl of 2xGreen PCR Master Mix (Thermo Scientific K1081), 2 µl (10 µM) 232 of a mixture of the forward primers, 2 µl (10 µM) of a mixture of the reverse primers 233 234 and 6.5 µl of high purity water. The PCR conditions described by Stegger et al. (2012) were applied. 235

236 2.2 Screening haemolysis and toxin production potential of the bacteria

237 2.2.1 Haemolytic activity on blood agar

The *Staphylococcus* isolates were screened for their haemolytic activity on blood agar as follows: Columbia agar base (Oxoid CM003) was autoclaved at 121 °C for 15 min and sheep blood (5%; Oxoid SR0051B) added after cooling to 50 °C, before distribution into Petri dishes. The isolates were then streaked on the agar and incubated at 37 °C for 48 h. Haemolysis was identified by the appearance of a zone of clearing around the colonies.

244 2.2.2 Production of staphylococcal enterotoxins A, B, C and D

The ability of the bacteria to produce the staphylococcal enterotoxin A, B, C 245 and D was investigated using the staphylococcal enterotoxin reversed passive latex 246 agglutination (SET-RPLA) toxin detection kit (Oxoid TD0900). Each isolate was grown 247 at 37°C for 24h in 10 ml of Tryptone Soya broth (TSB; Oxoid CM0129) and 1ml of the 248 249 culture centrifuged using a benchtop centrifuge at 3000 rpm for 20 min. The presence of toxins was screened in the supernatant according to the manufacturer's instructions. 250 251 2.2.3 Detection of staphylococcal enterotoxin, exfoliative and toxic shock syndrome toxin genes 252

The isolates were tested for the presence of the staphylococcal enterotoxin 253 genes sea, seb, sec, sed, see, seg, seh, sei, sej, exfoliative toxin genes eta and etb, 254 and toxic shock syndrome toxin gene tsst-1. Primers described by Johnson et al. 255 (1991), Omoe et al. (2002) and Nashev et al. (2004) were used. The PCR mixture was 256 that used by Ouoba et al. (2008b). The PCR conditions were as follows: 94°C for 2 257 min followed by 35 cycles of denaturation at 94°C for 1 min, annealing at the 258 corresponding temperature for 1 min and elongation at 72°C for 1 min. The PCR ended 259 with a final extension at 72°C for 5 min. 260

261 2.3 Screening the AMR phenotypic and genotypic profiles of the bacteria

262 2.3.1 Inoculum preparation

The isolates were grown on BHI-A and pure colonies re-suspended in 1 ml of Maximum Recovery Diluent (MRD; Oxoid CM0733). A quantity of the latter suspension was transferred to 10 ml of MRD to obtain a turbidity of 0.5 (10⁷-10⁸ CFU/ml) on the McFarland scale using a nephelometer (TREK Diagnostic systems, West Sussex, UK).

268 2.3.2 Determination of the susceptibility of the bacteria to antimicrobials

The Minimal Inhibitory Concentration (MIC) for 19 antimicrobials (Table 1) was 269 determined using a 96 well sensititre[™] plate (EUST, TREK Diagnostic Systems Ltd, 270 East Grinstead, UK) containing varying amounts of each antimicrobial. The method 271 recommended by EURL-AR (DTU, Denmark) was applied (EURL-AR, 2009). Briefly, 272 50 µl of the inoculum was mixed with 10 ml Mueller Hinton broth (MHB; Oxoid 273 CM0405) and 50 µl of this mixture were dispensed in each well. Incubation was carried 274 out aerobically at 37°C for 24 h and visible growth recorded. The susceptibility of the 275 isolates to each antimicrobial was established using the cut-off values recorded in 276 Table 1. 277

278 2.3.3 Detection of AMR genes

The presence of 38 genes encoding resistance to chloramphenicol, erythromycin, gentamycin, methicillin, penicillin, quinupristin/dalfopristin, tetracycline, trimethoprim, streptomycin and vancomycin were investigated (Table 1).

For the detection of the genes, the PCR mixtures described by Naghizadeh 282 Raeisi et al. (2018) were used. However, for the detection of the S. aureus methicillin 283 284 resistance genes *mec*A and *mec*C, the multiplex PCR (described in 2.1.2.4) and primers recommended by EURL-AR (2012) were used. For the mecA1 gene pertaining 285 286 to S. sciuri, the same mixture (except the primers) as that used for the detection of erythromycin (Naghizadeh Raeisi et al., 2018) was applied. The references of the 287 primers are depicted in Table 1. Positive control isolates (EURL-AR) for most genes 288 289 screened were included. Moreover, all positive amplicons were sequenced to confirm the identity of the genes. 290

All PCR amplifications were performed in a thermocycler (GeneAmp PCR 2700 system) using the following program: initial denaturation at 94°C for 3 min, followed by 25 - 35 cycles of 94°C for 1 min, 45-68°C for 1 min depending on the annealing temperature for the individual set of primers, 72 °C for 1 min and a final extension at 72 °C for 10 min. After purification, the presumptive positive amplicons were sequenced using the amplification primers at a concentration of 3.2 pmol/µl and the similarity of the sequence tested against published sequences in GenBank.

298 2.3.4 Screening the transferability of AMR

The ability of *S. warneri* 105 to transfer the *msr*(A) gene to *Enterococcus faecalis* JH2-2 and *S. saprophyticus* A17M105 to transfer the *tet*(K) gene to the same recipient was investigated by conjugation experiments according to Ouoba *et al.* (2008c) and Naghizadeh Raeisi et al. (2018). The donors are resistant to erythromycin

(S. warneri) and tetracycline (S. saprophyticus), but sensitive to rifampicin and fusidic
 acid. The recipient *E. faecalis* JH2-2 is resistant to rifampicin and fusidic acid and
 sensitive erythromycin and tetracycline.

Inocula were prepared as described above and 10 ml of the donor suspension 306 and 1 ml of that of the recipient mixed and filtered through a sterile membrane filter 307 (0.45 µm) (Whatman Laboratory Division, Maidstone, UK) using a vacuum pump 308 (Welch Thomas 2522C-02, Skokie, Illinois, USA). The filters were incubated 309 aerobically on BHI-A at 37 °C for 48h. The growth was washed off the filter with 3 ml 310 311 of MRD, diluted (up to 10⁻⁵) and 100 µl of each dilution spread on agar plates containing different antimicrobials. For the *msr*(A) gene, the inoculation was made on 312 BHI-RFE [rifampicin (25µg/ml), fusidic acid (25 µg/ml) and erythromycin (16 µg/ml)], 313 BHI-RF [rifampicin (25 µg/ml) and fusidic acid (25 µg/ml)], BHI-E [Erythromycin (16 314 µg/ml)] and BHI agar plates. For the tet(K) gene transfer, the mating mixtures were 315 spread on BHI-RFT [rifampicin (25µg/ml), fusidic acid (25 µg/ml) and tetracycline (10 316 µg/ml)], BHI-RF [rifampicin (25 µg/ml) and fusidic acid (25 µg/ml)], BHI-T [(tetracycline 317 (10 µg/ml)] and BHI agar plates. The plates were incubated at 37 °C for 7-21 days. 318 Cells of the donors and recipient that had not undergone a conjugation procedure were 319 all spread on the same type of media as those used to inoculate the mating mixtures. 320 Only recipients that had acquired resistance to erythromycin and tetracycline 321 322 are able to grow on BHI-RFE and BHI-RFT agar plates, respectively. Potential E. faecalis JH2-2 transconjugant (PET) DNA fingerprints were compared with those of 323 the recipients and donors by rep-PCR using the GTG5 primer (Ouoba et al., 2008a). 324 In addition, their MICs for erythromycin and tetracycline were determined. 325

2.3.5 Determination of the genetic background of acquired erythromycin and
 tetracycline resistance in the PETs

328 2.3.5.1 Extraction of DNA

Total DNA was extracted as described above. Plasmid DNA was extracted as follows: the donors, recipient and PETs were cultured for 48 h on BHI-A. A pure colony was then transferred into 10 ml BHI and incubated in a shaking water bath at 37°C for 12 h. Plasmid DNA was extracted using the QIAGEN Plasmid miniprep Kit (QIAGEN GmbH, Hilden, Germany) according to the manufacturer's instructions.

2.3.5.2 Amplification of the AMR genes in the PETs

PCR using total and plasmid DNA were performed to detect the presence of the *msr*(A) and *tet*(K) genes in the PETs. Since the *S. warneri* isolate contained also the *cat*501, *blaz*A and *dfr*(A) genes, these were also screened in the PETs. The use of plasmid DNA aimed to determine the location of genes in the bacteria and to increase the chance to detect a potential transfer in the PETs.

340 **3. Results**

341 3.1 Identification of the bacteria

The pH values of the products were 5-6 in the unfermented samples and 8-9 in 342 the fermented products. The bacterial numbers recovered (CFU/g) for each product 343 were as follows: Soumbala: 10⁶-10¹³ CFU/g on NA, 10³-10¹⁰ on MSA, 10¹-10⁸ on BP; 344 Bikalga: 10⁸-10¹¹ on NA, 10⁷-10⁸ on MSA, 10⁴-10⁸ on BP; Ntoba Mbodi: 10⁹-10¹¹ on 345 NA, 10⁸-10¹⁰ on MSA, 10⁵-10⁸ on BP. Out of 27 samples analysed, *Staphylococcus* 346 species were recovered from 18 samples. The products where Staphylococcus 347 isolates were not detected included mainly samples from the production steps of 348 Bikalga and Soumbala that had undergone heat treatment, but also one sample of 349 350 each of the three foods obtained from different markets. As mentioned earlier, many bacteria recovered on MSA and BP did not possess typical staphylococcal 351 characteristics and were not included in the advanced screening. Most of those 352

bacteria were Gram-positive (many of which were spore-forming) and Gram-negative rods that were able to grow on the selective media (BPA and MSA). The 90 isolates that were further screened showed variable colony and cell morphology. All were cocci, arranged differently (single, pairs, clusters) according to the isolate, catalase and Gram positive and oxidase negative. Two isolates from Bikalga were coagulase positive and the remaining 88 bacteria were coagulase negative.

359 The microbial profile was variable according to the fermented product, the production site and country (Table 2, Table 3). Genotyping by rep-PCR and PFGE 360 361 both generated 25 clusters representing 15 species (Table 2, Fig 1). The comparison of the PFGE profiles of the S. aureus isolates did not generate a similarity with any of 362 the 12,000 profiles in the CDC database. By sequencing the 16S rRNA and rpoB 363 genes, the bacteria of eight rep-PCR/PFGE clusters were identified as S. sciuri (45.6 364 % of the total isolates), three as *S. gallinarum* (7.8 %), two as *S. cohnii* (4.4%), two as 365 S. warneri (5.6%) and one cluster each as S. arlettae (2.2%), S. aureus (2.2%), S. 366 epidermidis (1.1%), S. haemolyticus (1.1%), S. hominis (1.1%), S. pasteuri (1.1%), S. 367 condiment (1.1%), S. piscifermentans (1.1%), S. saprophyticus (1.1%), S. simulans 368 (18.9%), and Macrococcus caseolyticus (previously designated as S. caseolyticus, 369 5.5%). In general, S. sciuri was the most frequently isolated species and was dominant 370 in the samples of Bikalga (42.9%) and particularly Ntoba Mbodi (71.1%) where it 371 occurred from the beginning of the fermentation and persisted throughout the process. 372 In Soumbala, S. simulans (54.8%) was the dominant species. The isolate of S. 373 haemolyticus was clearly identified by 16S rRNA gene sequencing/EZtaxon analysis 374 with 100 % similarity, but the *rpoB* gene sequencing generated a 96.5% (30 base pairs 375 difference) similarity with S. hominis and 93.5% (30 base pairs difference) with S. 376 haemolyticus. The spa type of the S. aureus isolate was identified as t355. 377

The species of S. sciuri and S. gallinarum were common to the two countries. 378 One cluster (5) of S. sciuri was detected in all three types of food although this 379 particular cluster was not present in all samples (Table 3). Clusters 4 and 6 of the 380 same species were detected in samples from both countries but not in all three 381 products. Staphylococcus gallinarum was also recovered from both countries but not 382 all products, with one cluster (15) observed in Soumbala and Ntoba Mbodi. Nine 383 384 species were specific to Burkina Faso including five isolated solely from Soumbala (S. epidermidis, S. pasteuri, S. condimenti, S. piscifermentans, S. simulans) and four from 385 386 Bikalga (S. aureus, S. cohnii, S. haemolyticus, M. caseolyticus). On the other hand, four species were peculiar to Congo/ Ntoba Mbodi (S. arlettae, S. hominis, S. 387 saprophyticus, S. warneri). Moreover, two clusters of S. gallinarum (1, 23) and two of 388 S. sciuri (2, 7) were peculiar to Soumbala while one cluster of S. sciuri was particular 389 to Bikalga. 390

391 3.2 Haemolysis on blood agar and toxin production

Haemolytic activity of the isolates on sheep blood agar was variable as seen in Table 4. Positive haemolysis was observed mainly in isolates of *S. aureus* and *S. simulans. Staphylococcus epidermidis*, *S. haemolyticus*, *S. pasteuri* and *S. warneri* exhibited partial haemolysis, whereas the rest of the species did not show haemolytic activity within the 48 h of incubation. The different species clusters showed similar haemolysis potential.

None of the bacteria screened produced the staphylococcal enterotoxins A, B, C and D. Furthermore, the genes coding for the pre-cited enterotoxins as well as the other eight genes were not detected in the isolates.

401 3.3 Antimicrobial resistance

402 All bacteria were susceptible to gentamycin, kanamycin, streptomycin, and vancomycin. For the rest of the antimicrobials, susceptibility was variable according to 403 the species, the cluster, and the antimicrobial screened (Table 4). Most of the 404 staphylococci were resistant to trimethoprim with 87.8 % of the isolates, representing 405 11 species, exhibiting phenotypic resistance to that antimicrobial. Both S. cohnii 406 clusters exhibited the highest phenotypic resistance profiles involving 11 407 antimicrobials. The bacteria with the least phenotypic resistance were S. condimenti 408 and S. pasteuri, which showed reduced susceptibility toward sulfamethoxazole only. 409 410 Within the same species, the different clusters exhibited different resistance patterns, but all isolates from the same cluster showed a similar pattern irrespective of the origin. 411 For example, isolates of cluster 1 of S. gallinarum were resistant to cefoxitin, 412 clindamycin, quinupristin/dalfopristin and tiamulin, whereas those of cluster 15 were 413 resistant to clindamycin, fusidate, penicillin, quinupristin/dalfopristin, tiamulin, 414 sulfamethoxazole, and trimethoprim, resulting in phenotypic resistance profiles which 415 differ by five antimicrobials (Table 4). 416

None of the 38 AMR genes screened was detected in the isolates of S. 417 condimenti, S. pasteuri, S. piscifermentans, S. simulans, and M. caseolyticus (Table 418 4). For the other species, different genes encoding resistance to multiple 419 antimicrobials including penicillin: *blazA*, chloramphenicol: *cat*501, trimethoprim: 420 421 dfr(A), dfr(G), erythromycin: *msr*(A), methicillin: *mec*A, *mecA*1 and tetracyline: *tet* (K) were detected, with a degree of AMR genotypic profile variability observed between 422 some species and clusters of the same species (Table 4). The mecA1 gene was 423 detected in 45.6% of the isolates, mecA in 20%, tet (K) in 11.1%, blazA in 10%, dfr(A) 424 in 10%, *msr*(A) in 6.7%, *cat*501 in 1.1%, and *dfr*(G) in 1.1%. Isolates of all six clusters 425 of S. sciuri showed mecA1, but bacteria in only three clusters displayed mecA. None 426

of the *S. aureus* isolates contained *mecA*, *mecA1* or *mecC*. The isolates of *S. epidermidis* and *S. warneri* showed the highest numbers (4) of AMR genes, including *blazA*, *dfr*(A), *dfr*(G) and *tet*(K) for *S. epidermidis* and *blazA*, *Cat501*, *dfr*(A) and *msr*(A)
for cluster 13 of *S. warneri*. The *blazA*, *mecA*, *mecA1* and *tet* (K) genes were found in
isolates from all products and all countries, *cat501* in Ntoba Mbodi /Congo, *dfr*(A) in
Ntoba Mbodi/Congo and Soumbala/Burkina Faso, *dfr*(G) in Soumbala/Burkina Faso,
and *msr*(A) in Ntoba Mbodi /Congo and Bikalga/Burkina Faso

Conjugation experiments produced E. faecalis JH2-2 PETs with increased 434 435 resistance to erythromycin and tetracycline. The MIC of some PETs increased up to 6 and 4 fold for erythromycin and tetracycline respectively (Table 5). All PETs showed 436 the same rep-PCR profiles as that of the recipient E. faecalis JH2-2 (results not 437 shown). No bacteria were recovered on the antimicrobial selective agar plates 438 inoculated with cells which did not undergo a conjugation process. Positive amplicons 439 were obtained for the genes screened for transfer using both total and plasmid DNA 440 samples from the donors. However, no positive amplicon was observed with DNA 441 samples from the recipient and the PETs under the experimental conditions used. 442

443 **4. Discussion**

The specificity in the bacterial profile of the products can be attributed to 444 differences in the raw materials and production processes. Also, environmental factors 445 can affect bacterial populations in a given geographical area and its associated plants 446 and thus can affect the bacterial population of the products derived from such plants. 447 However, since a similar type of fermentation occurs during production of the 448 fermented foods, similarities can be anticipated. High bacterial counts as observed in 449 our study have been reported in other studies describing the microbial composition of 450 alkaline fermented products (Ahaotu et al., 2013; Azokpota et al., 2006; Dakwa et al., 451

2005; Ouoba et al., 2010). Furthermore, the increase of pH constitutes a selection of 452 analogous alkali resistant microorganisms in the foods. The presence of the same 453 clusters of S. sciuri and S. gallinarum in both countries suggests a wide distribution of 454 those strains in the environment. A peculiarity of the S. sciuri isolates recovered from 455 the three products is that they were all oxidase negative. This seems unusual because 456 the S. sciuri group of species including S. sciuri, S. fleuritti, S. lentus, S. stepanovicii, 457 and S. vitulinus are known as the only oxidase positive species of the Staphylococcus 458 genus (Becker et al., 2014). The discrepancy could be related to the origin specificity 459 460 and to a lesser extent, the screening methods. The fact that the S. aureus PFGE profile did not match that of any clinical and environmental isolates in the CDC database 461 suggests that the cluster is uncommon and not globally distributed. 462

The species of S. sciuri, S. gallinarum, S. epidermidis, S. aureus and M. 463 caseolyticus have been reported in other African alkaline fermented products such 464 maari, dawadawa, and soydawadawa (Parkouda et al., 2009). However, most of the 465 other species, as well as the high species and strain diversity in the *Staphylococcus* 466 profiles observed in the current study, seem to have not been reported before. The 467 reason could be that *Staphylococcus* populations in such foods are usually screened 468 as part of the general microbial population and are not addressed specifically and 469 thoroughly using both phenotypic and genotypic methods. The species of S. 470 471 condimenti and S. piscifermentans have been reported to be typically associated with fermented foods such as fermented fish, sausages and their starter cultures (Becker 472 et al., 2014). It is believed that Staphylococcus spp. from fermented foods are 473 contaminants from the raw materials, equipment, environment and possibly the 474 producers. To produce Ntoba Mbodi, no heating step is included in the process, thus 475 the original Staphylococcus isolates, derived from the cassava and wrapping leaves, 476

that can withstand alkaline conditions probably constitute the major part of the 477 staphylococcal population. However, production of Soumbala and Bikalga includes 478 long cooking steps (24-48h), so it can be assumed that there is a recontamination of 479 the products from the processing and storage materials, the producers and the 480 environment, especially during the drying process. The limited recovery of 481 Staphylococcus isolates (90/400 presumptive isolates) supports their status as a minor 482 483 component of the microflora of the foods. They have often been isolated from alkaline fermented foods as secondary microorganisms, but a clear indication of their role in 484 485 the fermentation is not established. The main microorganisms responsible for alkaline fermentation, and which bring about desired changes to the foods, have been 486 demonstrated to be Bacillus species (Odunfa and Adewuyi, 1985; Ouoba et al., 2017). 487 Fermentation trials conducted by Odunfa and Adewuyi (1985) showed that S. 488 saprophyticus was unable to ferment African locust bean seeds to generate a product 489 with the typical organoleptic and nutritional properties. Nevertheless, it can be 490 suggested that the presence of Staphylococcus species contributes to some 491 biochemical activities such as those related to the degradation of proteins and lipids 492 that some species are able to perform during meat fermentation to produce sausages 493 (Casaburi et al., 2008; Milicevic et al., 2014). 494

495 *Staphylococcus* spp., especially CoPS, are known to cause various illnesses 496 including food intoxication due to their capacity to produce heat resistant enterotoxins. 497 Interestingly, none of the isolates investigated, including the two CoP *S. aureus,* 498 exhibited a potential toxin-producing phenotype and genotype. It cannot be ruled out 499 that they may contain other toxin genes that were not screened, but since the most 500 common genes were not detected, it can be assumed that these bacteria do not 501 represent a serious threat for consumers. In Burkina Faso and the Republic of the

Congo, there are no official reports of foodborne illnesses resulting from the 502 consumption of alkaline traditional fermented foods, although this may be related to 503 non-availability of data and lack of statistics. If unrecorded cases had occurred, these 504 would probably have been attributed to toxins elaborated by the fermenting *B. cereus* 505 strains that commonly occur in the products (Ouoba et al., 2008 b; Vouidibio-Mbozo 506 et al., 2017). In the study by Fowoyo and Ogunbanwo (2017a), the sea, seb, sec and 507 508 sed, and tsst-1 genes were detected in CoNS, including S. epidermidis, S. simulans, S. xylosus, S. kloosii and S. caprae, isolated from six Nigerian traditional fermented 509 510 foods including iru, an alkaline fermented African locust bean product that is similar to Soumbala. The positive amplicons were not sequenced to confirm the identity of the 511 gene, but the ability of some isolates to produce the staphylococcal toxins A, B, C, and 512 D suggest that these bacteria may pose a potential threat to consumers. 513

Antimicrobial resistances that are not associated with the presence of a gene 514 are of concern, but those involving a gene represent an enhanced threat, due to the 515 potential for transfer to other bacteria by mechanisms such as horizontal transfer, and 516 therefore can be spread. The tet (K), blazA, dfr(A), msr(A), cat501, and dfr(G) genes 517 detected can be associated with transferable mobile genetic elements such as 518 plasmids and transposons, and can therefore potentially be transferred to other 519 bacteria including pathogenic microorganisms. In the current research, tet (K), blazA, 520 521 *msr*(A), *cat*501 and *dfr*(A) genes were amplified using plasmid DNA, suggesting that they are located on plasmids, at least in the isolates screened. The AMR threat is 522 greater with bacteria with multi-drug genotypic profiles, such as one of the isolates of 523 S. warneri which exhibited AMR genes for four antimicrobials: erythromycin, 524 chloramphenicol, penicillin and trimethoprim, and those of S. epidermidis and S. 525

526 *hominis* which showed respectively four and three genes encoding resistance to 527 penicillin, tetracycline and chloramphenicol.

Methicillin resistance is an important factor when screening AMR in 528 Staphylococcus. This is because MRSA is responsible for a serious and persistent 529 problem in treatment of infections caused by such bacteria. The most virulent MRSA 530 are also resistant to other beta-lactam antimicrobials. Methicillin resistance is usually 531 532 caused by the presence of the *mecA* gene coding for a penicillin-binding protein (PBP2a) associated with a significantly lower affinity for beta-lactams. Thus, cell-wall 533 534 biosynthesis by the bacteria continues, even when a typically inhibitory concentration of antibiotic is administered (Peacok et al., 2015). Variants of *mecA* include genes 535 such as mecA and mecC (mecALGA251) originally found in S. aureus isolates, mecA1, 536 mecA2 and mecB (mecAm) initially detected in the S. sciuri group, S. vitulinus and M. 537 caseolyticus respectively (Ito et al., 2012). Isolates of three clusters of S. sciuri from 538 both countries exhibited both mecA1 and mecA, whereas the isolates of the three 539 other clusters also found in both countries exhibited only mecA1. The presence of the 540 *mec*A1 gene in *S. sciuri* is a common trait of this species and does not generally imply 541 methicillin resistance in the bacteria (Couto et al., 2000). However, the presence of 542 mecA in the isolates implicates methicillin resistance and represents a threat, although 543 a phenotypic resistance was not observed for cefoxitin. Couto et al. (2000) showed 544 that only S. sciuri isolates that possess a copy of mecA in addition to mecA1 exhibit 545 significant levels of resistance to beta-lactams. In our study, such isolates displayed 546 phenotypic resistance to penicillin. Interestingly, the S. aureus did not display any of 547 the methicillin resistance genes screened and were susceptible to cefoxitin. However, 548 the strains were resistant to penicillin and contained the *blaZ* gene. 549

The prevalence of AMR genes in various CoNS and CoPS from processed 550 and unprocessed foods such as meat, fish and fermented dairy and meat products 551 from different origins has been widely documented (Argudín et al., 2011; Kürekci, 552 2016; Perreten et al., 1998; Zdolec et al., 2012). However, such information on CoNS 553 and CoPS from alkaline fermented products and especially those from Africa are, to 554 the best of our knowledge, non-existent. This can be due to the limited importance 555 attributed to these bacteria in alkaline fermentation. The study by Fowoyo and 556 Ogunbanwo (2017b) on CoNS from different Nigerian fermented foods, reported 557 558 phenotypic resistance to antimicrobials such as trimethoprim-sulfamethoxazole, ciprofloxacin, erythromycin, gentamicin, and cefotaxime, but no AMR gene was 559 reported, probably because only *mec*A was screened in the isolates with no positive 560 561 results.

The significant increase of the MIC above the cut-off values in the PETs for 562 erythromycin and tetracycline indicates that they have acquired resistance to the 563 antimicrobials. Since the genes screened were not detected in the PETs, it can be 564 assumed that they were not transferred, but that the isolates acquired resistance 565 through other mechanisms, such as mutations. The occurrence of resistances was 566 triggered by the presence of the Staphylococcus donors because PETs were not 567 recovered from recipient cells which did not undergo a conjugation process. Mutations 568 can occur as a result of a mechanism by which the *Staphylococcus* spp. stimulate an 569 evolution of *E. faecalis* JH2-2 by increasing, for instance, the expression of error-prone 570 DNA repair mechanisms (Naghizadeh Raeisi, 2018). Alternatively, the growth of the 571 recipient could be stimulated by the donor leading to a larger recipient population in 572 which mutations can arise (Naghizadeh Raeisi, 2018). However, it is worth noting that 573 when a low number of plasmid copies which carry a gene is transferred, it may not be 574

detected using conventional PCR. Moreover, other genes coding for resistance to the 575 antimicrobials and which were not screened, may be present in the donors and 576 transferred to the recipients. Transfer potential varies according to different factors 577 related to the environment, type of donor and recipient, and the location of the AMR 578 genes. Thus, the genes screened may have not been transferred into E. faecalis JH2-579 2 and therefore the threat with regards to the potential of the donors to spread AMR is 580 581 limited. However, the possibility of gene transfer to other types of bacteria cannot be ruled out since the genes are located on plasmids which are mobile genetic elements 582 583 that can be transferred in appropriate conditions.

The current study demonstrates ecological similarities and differences in the 584 Staphylococcus population of different types of alkaline fermented foods. The main 585 microorganisms identified are primarily CoNS with a very limited number of CoPS, as 586 would be expected in non-clinical samples. The bacteria did not suggest a safety issue 587 with regards to the potential to produce toxins responsible for foodborne diseases. 588 Due to the presence of multiple AMR genes in some isolates, they are reservoirs of 589 AMR determinants which may be spread under particular conditions. However, in 590 general, the results of the study herein reported suggest little safety importance of the 591 staphylococcus isolates in the fermented products screened. 592

593

594 Acknowledgements

595 The authors are grateful to the following people for their technical and scientific support 596 in the realization of the research project: Professor Simon C. Kobawila, and Professor 597 Delphin Louembe (Marien N'Gouabi University, Brazzaville, Congo), Dr Brigitte 598 Awamaria (London Metropolitan University, London, UK), Dr Rene S. Hendriksen, 599 Susanne Karlsmose Pedersen, Ana Rita Rebelo, and Dr. Lars Bøgo Jensen (Technical University of Denmark, Lyngby, Denmark), Janne M. Benjaminsen
(University of Copenhagen, Copenhagen, Denmark), Dr Eva Møller Nielsen, and Dr
Jesper Larsen (Statens Serum Institut, Copenhagen, Denmark), Dr Molly Freeman,
Dr Jeniffer Concepcion Acevedo, and Valerie S. Albrecht (Centers for Disease Control
and Prevention, Atlanta, GA, USA)

605

606 **Declaration of interest:** None

607 **Funding:** This research did not receive any specific grant from funding agencies in

608 the public, commercial, or not-for-profit sectors

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789 Figure caption

Figure 1: Cluster analysis (Bionumerics, Dice's Coefficient of similarity, Unweighted pair group method with arithmetic: UPGMA) of the different rep-PCR and PFGE fingerprints of *Staphylococcus species* isolated from Bikalga, Soumbala and Ntoba Mbodi

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810 Table 1: Antimicrobials studied and their effect on the *Staphylococcus* isolates

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Antimicrobials	Concentration	Cut off	Genes screened ^e	Staphylococcus	isolates	
	Range screened (mg/L)	values(mg/L)		^f MIC Range (mg/L)	Resistant (%)	Susceptible
Cefoxitin	0.5 - 16	4 ^{a,b}	<i>mec</i> A, <i>mec</i> A1 and <i>mec</i> C for methicillin resistance	1 - 8	8.9	91.1
Chloramphenicol	4 - 64	8 ^c	cat501. catA1. cm/A	<4 - 64	52.2	47.8
Ciprofloxacin	0.25 - 8	1 ^a	,,.	<0.25 - 2	1.1	98.9
Clindamycin	0.12 - 4	0.25 ª		<0.12 - 4	41.1	58.9
Erythromycin	0.25 - 8	1 ^a	erm(A), erm(B), erm(C), erA, erB, msrA, mphA, mefA/mefE	<0.25 - >8	11.1	88.9
Fusidate	0.5 - 4	1 ^c		<0.5 - >4	43.3	56.7
Gentamicin	1 - 16	2 ª	aac(6′)aph(2″), aac(3″)II, aac(3″)IV, ant(2″)-I	<1	0	100
Kanamycin	4 - 64	16 ^d		<4	0	100
Linezolid	1 - 8	4 ^a		<1 - 8R	1.1	98.9
Mupirocin,	0.5 - 256	1 ^a		<0.5 - > 256	1.1	98.9
Penicillin	0.12 - 2	0.125 ^b	BlaZ	<0.12 - >2	43.3	56.7
Quinupristin- dalfopristin	0.5 - 4	1 ^a	VatD, VatE	<0.5 - 4	64.4	35.6
Rifampicin	0.016 - 0.5	0.5 ^c		<0.016 - >0.5	4.4	95.6
Streptomycin	4 - 32	16 ^b	StrA, StrB, aadA, aadE	<4 - 16	0	100
Sulfamethoxazole	64 - 512	128 ª		<64 - >512	61.1	38.9
Tetracycline	0.5 - 16	1ª	tet(M), tet(L), tet(S), tet(Q), tet(K), tet(O), tet(W)	<0.5 - >16	6.7	93.76
Tiamulin	0.5 - 4	2ª		<0.5 - > 4	48.9	51.1
Trimethoprim	2 - 32	2 a	dfr(A), dfr(G), dfr(K)	<2 - >32	87.8	12.2
	1 16	2 a	Von A Von P Von V	_ 0_	0	100

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813 Cut-off values according to a) EFSA (2012), b) *EURL-AR (2017), c)* EUCAST (2018) and d) Virdis et 814 al. (2010)

e) Most primer sequences are from EURL-AR (2013) and Ouoba et al. (2008), except for mecA1

816 (Tsubakishita et al., 2010), erA, erB, msrA, mphA, mefA/mefE (Sutcliffe et al., 1996), dfr(G), dfr(K

817 (Argudín et al., 2011)

818	f) MIC: Minimal Inhibitory Concentration

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824	Table 2: Identity	[,] and origin	of the Staphvl	lococcus species	recovered from Soumbala.

Bikalga, and Ntoba Mbodi

Samples/Origin	Fermentation time/	Isolates	^b Rep-PCR/	Identification by 16S RNA and
Soumbala	Sampling Site			Ipob gene sequencing
ooumbulu				
Burkina Faso				
S3	Site 1/ 0h	S35M (47)	14	Staphylococcus epidermidis
S5	Site 1/ 48h	S51B (33)	23	Staphylococcus gallinarum
-		S54a M (34)	23	Staphylococcus gallinarum
		S54b M (35)	23	Staphylococcus gallinarum
S6	Site 1/ Dried	S63aM (41)	3	Staphylococcus simulans
SF	Site 2/ ≥72h	SF1M (Ì) Ú	1	Staphylococcus gallinarum
		SF3M (2)	3	Staphylococcus simulans
		SF2M (3)	5	Staphylococcus sciuri
		SF2bB (6)	3	Staphylococcus simulans
		SF2aB (7)	3	Staphylococcus simulans
		SF1B (8)	3	Staphylococcus simulans
		SF8M (21)	2	Staphylococcus sciuri
		SF7M (22)	3	Staphylococcus simulans
		SF6M (23)	7	Staphylococcus sciuri
SP	Site 3/ ≥72h	SP6M (26)	10	Staphylococcus piscifermentans
		SP5M (27)	3	Staphylococcus simulans
		SP3M (28)	3	Staphylococcus simulans
		SP1M (29)	11	Staphylococcus pasteuri
		SP4B (30)	3	Staphylococcus simulans
		SP2B (31)	3	Staphylococcus simulans
		SP1B (32)	3	Staphylococcus simulans
SZ	Site 4/ ≥72h	SZ1B (10)	3	Staphylococcus simulans
		SZ2B (11)	3	Staphylococcus simulans
		SZ6B (13)	3	Staphylococcus simulans
		SZ1M (14)	5	Staphylococcus sciuri
		SZ3M (15)	3	Staphylococcus simulans
		SZ4M (16)	5	Staphylococcus sciuri
	XV	SZ5M (17)	9	Staphylococcus condimenti
		SZ6M (18)	15	Staphylococcus gallinarum
		SZ7M (24)	3	Staphylococcus simulans
		SZ9M (25)	3	Staphylococcus simulans
Bikalga				
Durk Frank				
Burkina Faso				
22	Sita 5/ 19h	P21M (50)	6	Stanbylaggagua gajuri
DZ	Sile 5/ 4011	DZ IIVI (30) DOM (51)	0	Staphylococcus sciuri
		B2ZIVI (31) B27aM (53)	4	Staphylococcus sciuli Staphylococcus cobnii
		DZ7 alvi (33) D276M (54)	10	Staphylococcus connii
		DZTDIVI(34) D25M(56)	10	Staphylococcus connii
		DZOIVI (00)	0	Staphylococcus comm
		D23D(37)	0	Staphylococcus sciuri
D2	Sita 5/70h	D24D (00)	24 5	Staphylococcus comm
DJ	Sile 5/ 7211	DJOIVI (00)	5 5	Staphylococcus sciuri
		B34M (62)	5	Staphylococcus sciuri
		D341VI (02)	5	Staphylococcus sciuri
		B31M (65)	5	Staphylococcus sciuri
		B31B (66)	0 8	Staphylococcus sciuri

^aSite 1: Ouagadougou (Samandin production site), Site 2: Fada market, Site 3: Ouagadougou (Patte
 d'Oie market), Site 4: Ouagadougou (Zogona market), Site 5: Ouagadougou (Zone Pilote production

828 site) ^b Rep-PCR : Repetitive element palindromic-polymerase chain reaction ; PFGE : Pulse Field Gel

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829 **Electorphoresis**

830 Table 2 *continued*: Identity of the *Staphylococcus* species recovered from Soumbala,

831 Bikalga, and Ntoba Mbodi

Samples/Origin	Fermentation time/	Isolates	<mark>♭</mark> Rep-PCR/	Identification 16S RNA and
	sampling site ^a		PFGE clusters	<i>rpoB</i> gene sequencing
B5	Site 5/ Dried	B57B (69)	18	Staphylococcus aureus
		B51B (70)	18	Staphylococcus aureus
BP	Site 6/ Dried	BP2B (73)	19	Macrococcus caseolyticus
		BP2M (74)	19	Macrococcus caseolyticus
		BP3B (75)	19	Macrococcus caseolyticus
		BP6B (76)	19	Macrococcus caseolvticus
		BP5B (77)	19	Macrococcus caseolvticus
BB	Site 7/ Dried	BB3B (84)	20	S. haemolvticus
Ntoba Mbodi		(
Congo				
	Site 8/ 0h	A04M (93)	25	Stanbylococcus sciuri
A00		$\Delta 05M(93)$	12	Staphylococcus warneri
		AOSM(34)	12	Staphylococcus warneri
		A00101 (95)	12	Staphylococcus warnen
		AU/IVI (103)	12	Staphylococcus warnen
A 1 1 4	04-0/04-	AU6B (193)		Staphylococcus warnen
AU1	Site 8/ 24h	A12B (96)	6	Staphylococcus sciuri
		A15M (104)	25	Staphylococcus sciuri
		A1/M (105)	13	Staphylococcus warneri
		A18M (106)	17	Staphylococcus arlettae
		A15M (176)	4	Staphylococcus sciuri
		A16M (177)	6	Staphylococcus sciuri
		A111B (186)	5	Staphylococcus sciuri
		A110B (194)	5	Staphylococcus sciuri
AU2	Site 8/ 48h	A25B (111)	17	Staphylococcus arlettae
		A215B (113)	15	Staphylococcus gallinarum
		A21M (115)	4	Staphylococcus sciuri
		A27B (178)	5	Staphylococcus sciuri
		A29B (179)	6	Staphylococcus sciuri
		A26M (180)	6	Staphylococcus sciuri
		A23M (191)	4	Staphylococcus sciuri
		A24M (192)	4	Staphylococcus sciuri
AU3	Site 8/ 72h	A37bB (123)	6	Staphylococcus sciuri
		A39B (124)	21	Staphylococcus hominis
		A34M (128)	6	Staphylococcus sciuri
		A31M (188)	6	Staphylococcus sciuri
		Δ33M (187)	6	Staphylococcus sciuri
100	Site 9/0h	103B (131)	6	Staphylococcus sciuri
300		104B (122)	6	Staphylococcus sciuri
		J04D (13Z)	0	Staphylococcus sciuri
		JUID (109)	4	
	0:1.0/70	JUZB (190)	0	Staphylococcus sciun
103	Site 9/ / 2h	JUJIVI (138)		Stapnylococcus saprophyticus
		J311B (148)	15	Stapnylococcus gallinarum
		J33M (150)	6	Staphylococcus sciuri
		J34M (151)	6	Staphylococcus sciuri
		J31M (181)	6	Staphylococcus sciuri
		J32M (182)	6	Staphylococcus sciuri
MB	Site 10/ 72-96h	MB1M (156)	5	Staphylococcus sciuri
		MB11M (184)	5	Staphylococcus sciuri

aSite 5: Ouagadougou (Zone Pilote production site), Site 6: Ouagadougou (Patte d'Oie market), Site

833 7: Bobo Dioulasso (market), Site 8: Brazzaville (Production site A), Site 9: Brazzaville (Production site

B), Site 10: Brazzaville (market)

835 836	^b Rep-PCR : Repetitive element palindromic-polymerase chain reaction ; PFGE : Pulse Field Gel Electorphoresis
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857	Table 3 Geographical distribution of the Staphylococcus species and associated clusters
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Species	Rep-PCR/	Burkina F	aso/ Soumb	ala		Burkina	Faso/ Bikalo	ja	Congo/ I	Ntoba Mbod	i
	PFGE Clusters	Site 1 ^b	Site 2	Site 3	Site 4	Site 5	Site 6	Site 7	Site 8	Site 9	Site 10
S. arlettae	17	_c	-	-	-	-	-	-	+	-	-
S. aureus	18	-	-	-	-	+	-		-	-	-
S. cohnii	16	-	-	-	-	-	+		_	-	-
	24	-	-	-	-	-	+		-	-	-
S. epidermidis	14	+	-	-	-	-	-		-	-	-
S. gallinarum	1	+	-	-	-	-	-	_	-	-	-
-	15	-	-	-	+	-	-	-	+	+	-
	23	+	-	-	-	-	-	-	-	-	-
S. haemolyticus	20	-	-	-	-	-		+	-	-	-
S. hominis	21	-	-	-	-	-	-	-	+	-	-
S. pasteuri	11	-	-	+	-		-	-	-	-	-
S. condimenti	9	-	-	-	+	-	-	-	-	-	-
S. piscifermentans	10	-	+	-	-	-	-	-	-	-	-
S. saprophyticus	22	-	-	-	-		-	-	-	+	-
S. sciuri	2	-	+	-	-	-	-	-	-	-	-
	4	-	-	-	-	+	-	-	+	+	-
	5	-	+	-	+	+	-	-	-	-	+
	6	-	-	-	-	+	-	-	+	+	-
	7	-	+		-	-	-	-	-	-	-
	8	-	-		-	+	-	-	-	-	-
	25	-	-	-	-	-	-	-	+	-	-
S. simulans	3	+	+	+	+	-	-	-	-	-	-
S. warneri	12	-	-	-	-	-	-	-	+	-	-
	13	-		-	-	-	-	-	+	-	-
M. caseolyticus	19	-		-	-	-	+	-	-	-	-

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860 ^a Rep-PCR : Repetitive element palindromic-polymerase chain reaction ; PFGE : Pulse Field Gel Electrophoresis

861 ^bSite 1: Ouagadougou (Samandin production site), Site 2: Fada market, Site 3: Ouagadougou (Patte d'Oie market), Site 4: Ouagadougou (Zogona market),

862 Site 5: Ouagadougou (Zone Pilote production site), Site 6: Ouagadougou (Patte d'Oie market), Site 7: Bobo Dioulasso (market), Site 8: Brazzaville

863 (Production site A), Site 9: Brazzaville (Production site B), Site 10: Brazzaville (market)

864 ^c - : not detected, + : detected

Table 4 Profiles of antimicrobial resistance (AMR) and toxin production potential of the species and clusters of *Staphylococcus* from
 Bikalga, Soumbala, and Ntoba Mbodi

Origin ^a	Rep-PCR/ PFGE Cluster	Species	Number of Isolates	AMR phenotype ^b	AMR genes	Toxin genes/ Toxin A, B, C, D	Haemolysis on blood agar
Ntoba Mbodi (Co)	17	S. arlettae	2	CHL, CLI, FUS, SYN, SMX, TIA, TMP	dfr(A)	_ c	-
Bikalga (BF)	18	S. aureus	2	CHL, ERY, PEN, SYN, SUL	blazÁ	-	+
Bikalga (BF)	16	S. cohnii	3	FOX, CHL, CLI, ERY, FUS, PEN, RIF, SMX, SYN, TIA, TMP	msr(A)	-	-
Bikalga (BF)	24	S. cohnii	1	FOX, CLI, ERY, FUS, PEN, RIF, SMX., SYN, TET, TIA, TMP	<i>msr</i> (A)	-	-
Soumbala (BF)	14	S. epidermidis	1	CHL, PEN, SMX, TET, TMP	blazA, dfr(A), dfr(G) tet(K)	-	±
Soumbala (BF)	1	S. gallinarum	1	FOX, CLI, SYN, TIA	<i>tet</i> (K)	-	-
Soumbala, Ntoba Mbodi (BF, Co)	15	S. gallinarum	3	CLI, FUS, PEN, SYN, TIA, SMX, TMP	<i>tet</i> (K)	-	-
Soumbala (BF)	23	S. gallinarum	3	FOX, CLI, FUS, PEN, SYN, TIA, TMP	<i>tet</i> (K)	-	-
Bikalga (BF)	20	S. haemolyticus	1	CLI, ERY, PEN, SYN, TIA, TMP	tet(K), <i>msr</i> (A)	-	±
Ntoba Mbodi (Co)	21	S. hominis	1	CHL, PEN, SMX, TET, TMP	blazA, <i>te</i> t(K), dfr(A)	-	-
Soumbala (BF)	11	S. pasteuri	1	SMX		-	±
Soumbala (BF)	10	S. piscifermentans	1	CHL, ERY,SMX, TIA		-	-
Soumbala (BF)	9	S. condimenti	1	SMX		-	-
Ntoba Mbodi (Co)	22	S. saprophyticus	1	CHL, PEN, TET, TMP	<i>tet</i> (K)	-	-
Soumbala (BF)	2	S. sciuri	1	CLI, FUS, SMX, TIA, TMP	mecA, mecA1	-	
Bikalga, Ntoba Mbodi (BF, Co)	4	S. sciuri	6	CLI, FUS, TIA, TMP	mecA1	-	-
Bikalga, Soumbala, Ntoba Mbodi (BF, Co)	5	S. sciuri	13	CLI, FUS, TMP	mecA1	-	-
Bikalga, Ntoba Mbodi (BF, Co)	6	S. sciuri	16	CHL, PEN, SYN, SMX, TIA, TMP	mecA, mecA1	-	-
Soumbala (BF)	7	S. sciuri	1	CLI, ERY, FUS, SYN, TIA, TMP	mecA, mecA1	-	-
Bikalga (BÈ)	8	S. sciuri	2	CHL, PEN, SYN, TET, TIA, TMP	mecA1	-	-
Ntoba Mbodi (Co)	25	S. sciuri	2	CLI, FUS, SMX, SYN, TIA, TMP	mecA1		
Soumbala (BF)	3	S. simulans	17	CHL, LZD, SYN, SMX, TMP		-	+
Ntoba Mbodi (Co)	12	S. warneri	4	FUS, PEN, SYN, SMX, TMP	blazA, dfr(A)	-	±
Ntoba Mbodi (Co)	13	S. warneri	1	CHL, ERY, PEN, SMX, TMP	blazA, Cat501 dfr(A), <i>msr</i> (A)	-	-
Bikalga (BF)	19	M. caseolyticus	5	CIP, MUP		-	-

⁸⁶⁷ ^a BF: Burkina Faso, Co: Congo

^b The profile represents the antimicrobials to which the isolates were resistant to; Cefoxitin: FOX, Chloramphenicol: CHL, Ciprofloxacin: CIP,

869 Clindamycin: CLI, Erythromycin: ERY, Fusidate: FUS, Mupirocin: MUP, Quinupristin/Dalfopristin (Synercid): SYN, Linezolid: LZD, Penicillin:

870 PEN, Rifampicin: RIF, Sulfamethoxazole: SMX, Tiamulin: TIA, Trimethoprim: TMP ° -: negative, ±: partial haemolysis, +: positive haemolysis

>32 <1 16 <1	Donor S. <i>saprophyticus</i> Recipient <i>E. faecalis</i> PET-T1	64 1 8
<1 16 <1	Recipient <i>E. faecalis</i> PET-T1	1 8
16 <1	PET-T1	8
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<i>></i> 3∠	PET-T3	16
<1	PET-T4	8
<1	PET-T5	8
4	PET-T6	16
<1	PET-T7	16
<1	PET-T8	4
>32	PET-T9	16
<1	PET-T10	4
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<1		
>32		
<1		
	<1 4 <1 >32 <1 32 <1 >32 <1 >32 <1	<1 PET-T5 4 PET-T6 <1 PET-T7 <1 PET-T8 >32 PET-T9 <1 PET-T10 32 <1 >32 <1 >32 <1

Table 5: Minimal Inhibitory Concentration (MIC) for donors, recipient and potential *E. faecalis* JH2-2 transconjugants (PET)



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