INTRODUCTION

Fish and fishery products are an important food component for a large part of the world’s population, with an average consumption level of 20.1 kg per capita (FAO, 2016). In developing countries, fish is a relatively cheap and accessible protein source, suitable for complementing high carbohydrate-based diets of West African population (Adeyeye, Oyewole, Obadina, & Omemu, 2015; Ikutegbe & Sikoki, 2014). Among muscle food, fish is the most perishable and loses freshness after death due to autolytic and microbial spoilage (Dehghani, Hosseini, & Regenstein, 2018; Matak, Tahergorabi, & Jaczynski, 2015). In tropical regions, conservation of fresh fish remains a problem because of the lack of adequate infrastructures, and environmental and climatic conditions that contribute to its spoilage.
within few hours (Anihouvi, Kindossi, & Hounhouigan, 2012). To prevent fish spoilage and reduce postcatchature losses, various preservation methods including frying, fermentation, drying, salting, and smoking are used (Adeyeye et al., 2015; Ikutege & Sikoki, 2014). Smoking consists in submitting fish to direct or indirect action of smoke during the incomplete combustion of certain trees used as fuel. Smoking of foodstuffs improves food organoleptic characteristics, induces water loss, and reduces the microbial load, thanks to heat and the presence of aromatic and bactericidal substances (Chakroborty & Chakraborty, 2017; Yusuf et al., 2015).

In Benin, fish products are the most important source of animal proteins (Kpodékon et al., 2014). Traditional smoking, one of the main methods used for fish preservation in the country (Dégnon et al., 2013), generates two types of end products, smoked fish (SF) and smoked–dried fish (SDF), used for local consumption or exported to neighboring countries.

Fish and fish products are involved in 10%–20% of foodborne diseases (Pilet & Leroi, 2011), and the presence of pathogenic bacteria such as Staphylococcus aureus, Salmonella spp., pathotypes of Escherichia coli, and Listeria monocytogenes has been reported in SF (Adeyeye et al., 2015; Ayeloja, George, Jimoh, Shittu, & Abdulsalami, 2018; Ineyougha, Orutugo, & Izah, 2015; Likongwe, Kasapila, Katundu, & Mpeketula, 2018; Nunoo & Kombat, 2013; Udochukwu, Inetianbor, Akaba, & Omorotionmwan, 2016). Another concern is the contamination by fungi. In this respect, various studies have reported the occurrence of aflatoxigenic fungi in SDF (Ayeloja et al., 2015; Ikutegbe & Sikoki, 2014; Babalola, Odebode, Ojomo, Ogungbemile, & Jonathan, 2018; Job, Agina, & Dapiya, 2016; Wogu & Iyai, 2011), which under certain conditions can produce mycotoxins (Wogu & Iyai, 2011). It is therefore necessary to take action by improving the microbiological quality of SF and SDF. The present study aims to provide insights into the microbiological status of SF and SDF processed in Benin, and to identify associated contamination factors in order to formulate suitable corrective measures.

2 | MATERIAL AND METHODS

2.1 | Sample collection

A total of 66 fish samples including fresh fish and processed fish were collected. Samples of SF and SDF from six fish species (Scomber scombrus, Merluccius polli, Oreochromis niloticus, Cypselurus cyanopterus, Sphyraena baraccuda, and Ethmalosa fimbriata) were randomly purchased at different processing sites and markets at Aguégués, Cotonou, Abomey-Calavi, Comé, and Aplahoué municipalities (Benin). The samples were collected using individual sterile plastic bags and were transported under refrigeration to the laboratory.

2.2 | Follow-up of the manufacturing processing

The follow-up of the processing was performed on two fish species: S. scombrus (smoking) and C. cyanopterus (smoking–drying) identified as the most used by processors, according to a previous survey (data not shown). Twelve trials were performed with three experienced processors, each performing two smoking and two smoking–drying trials. Three types of samples were collected at sensitive steps: raw fresh fish, cleaned raw fresh fish, and SF (or SDF).

2.3 | Enumeration of the smoked and smoked–dried fish microflora

For each sample, 25 g was suspended in 225 ml of buffered peptone water (Bio-Rad, pH 7.0 ± 0.2) and homogenized (230 rpm for 2 min) using a stomacher (Lab Blender; Seward Medical) to obtain a 1/10 dilution. Decimal dilutions were prepared in BPW as described by ISO 6887-3 and inoculated in different media: (a) Plate Count Agar (Bio-Rad) for aerobic mesophilic bacteria (AMB), incubated at 30°C for 72 ± 3 hr (ISO 4833-1); (b) De Man–Rogosa–Sharpe Agar (Bio-Rad) for lactic acid bacteria (LAB), incubated at 30°C for 72 ± 3 hr (ISO 15124); (c) Violet Red Bile Glucose (Bio-Rad) for Enterobacteriaceae, incubated at 37°C for 24 ± 2 hr (ISO 21528-2); (d) TBX Agar (Bio-Rad) for E. coli incubated at 37°C for 21 ± 3 hr (ISO 16649-2); (e) Tryptose Sulfite Cycloserine Agar (Bio-Rad) supplemented with Perfringens Selective Supplement (SFP Oxoid) for Clostridium perfringens, incubated at 37°C for 24–48 hr (presumptive colonies confirmed with Lactose Sulfite (ISO 7937)); (f) Baird–Parker Agar (Bio-Rad) supplemented with Rabbit Plasma Fibrinogen (Bio-Rad) and incubated at 37°C for 24 hr (ISO 6888-2) for S. aureus; and (g) Mannitol Egg Yolk Polymyxin Mossel base (Biokar Diagnostics-Zac) with Egg Yolk (Biokar Diagnostics-Zac) incubated at 30°C for 18–24 hr (ISO 7932) for Bacillus cereus. Yeasts and molds were investigated on Yeast Glucose Chloramphenicol Agar (Bio-Rad) incubated at 25°C for 72–120 hr (ISO 7954). L. monocytogenes was sought on RapidL Mono Agar (Bio-Rad) incubated at 37°C for 24 ± 2 hr (BRD). Salmonella spp. were investigated on Rapid’Salmonella Agar (Bio-Rad) after enrichment in buffered peptone water with addition of active supplement (capsules Bio-Rad). Confirmation was performed using Salmonella Latex Kit (Bio-Rad) for agglutination test according to the validated method BRD 07/11-12/05.

2.4 | 16S rRNA sequencing

Sequencing of 16S rRNA gene was performed on fresh colonies. PCR mixture was prepared by using sterile bi-distilled water, MgCl₂ (25 mM), Universal primers Univ1 (10 μM) and Bact4 (10 μM), DNTP Mix (2 mM), GoTaQ Flexi Buffer and GoTaQ enzyme in adequate amount. Sequences of the universal primers Univ1 and Bact4 were 5’-ACTCTACCAGGAGGCAG-3’ and 5’-GGCGTGTGTCAAAGGCAGG-3’, respectively. One μl of the suspension of a fresh colony in Ringer was mixed with 49 μl PCR mixture. Amplification was performed in a C1000™ Thermal Cycler (Bio-Rad) as follows: 5 min at 95°C, 30 cycles (1 min at 95°C, 1 min at 50°C, 1 min at 72°C) and 10 min at 72°C. Five microlitre of products was run on 0.8% agarose gel (0.5% Tris-acetate–EDTA buffer) to which 5 μl/100 ml EtBr was added. DNA bands were visualized by UV light photography.
| Type of fish | Parameters          | Mean ± SD | Min. | Max. | Positive samples<sup>a</sup> | AL<sup>b</sup> | NCS |
|-------------|---------------------|-----------|------|------|-----------------------------|-------------|-----|
| SF (n = 18) | AMB                 | 7.4 ± 1.8<sup>c</sup> | 3.8  | 9.5  | 18 (100%)                  | <7          | 12  |
|             | LAB                 | 7.0 ± 1.7<sup>c</sup> | 3.7  | 9.2  | 18 (100%)                  | NA          | NA  |
|             | Enterobacteriaceae  | 4.4 ± 2.7<sup>c</sup> | <1   | 8.2  | 16 (88.9%)                  | 4           | 9   |
|             | *Escherichia coli*  | 2.8 ± 2.5<sup>c</sup> | <1   | 6.8  | 8 (44.4%)                   | 2           | 7   |
|             | *Bacillus cereus*   | 1.5 ± 1.5<sup>c</sup> | <1   | 5.1  | 5 (27.8%)                   | 5           | 2   |
|             | *Clostridium perfringens* | 1.1 ± 1.2<sup>c</sup> | <1   | 5.7  | 13 (72.2%)                  | 4           | 1   |
|             | Yeasts              | 3.5 ± 1.8<sup>c</sup> | <1   | 5.7  | 17 (94.5%)                  | NE          | NA  |
|             | Molds               | 3.3 ± 1.9<sup>c</sup> | <1   | 6.0  | 13 (72.2%)                  | NE          | NA  |
|             | *Staphylococcus aureus* | <1          | <1   | <1   | 0 (0%)                      | 2           | 0   |
|             | *Listeria monocytogenes* | <1         | <1   | <1   | 0 (0%)                      | 2           | 0   |
|             | *Salmonella spp.*   | Abs        | NA   | NA   | 0 (0%)                      | Abs         | 0   |
| SDF (n = 18)| AMB                 | 4.8 ± 1.7<sup>c</sup> | 2.9  | 7.8  | 18 (100%)                  | <7          | 4   |
|             | LAB                 | 4.1 ± 1.9<sup>c</sup> | <1   | 7.3  | 17 (94.4%)                  | NE          | NA  |
|             | Enterobacteriaceae  | 1.5 ± 1.4<sup>c</sup> | <1   | 5.1  | 7 (38.9%)                   | 4           | 1   |
|             | *E. coli*           | 0.8 ± 0.2<sup>c</sup> | <1   | 1.3  | 2 (11.1%)                   | 2           | 0   |
|             | *B. cereus*         | 2.8 ± 0.9<sup>c</sup> | <1   | 4.4  | 15 (83.3%)                  | 5           | 0   |
|             | *C. perfringens*    | 1.1 ± 0.5<sup>c</sup> | <1   | 2.1  | 8 (44.4%)                   | 4           | 0   |
|             | Yeasts              | 1.0 ± 0.6<sup>c</sup> | <1   | 2.7  | 5 (27.8%)                   | NE          | NA  |
|             | Molds               | 2.1 ± 1.1<sup>c</sup> | <1   | 4.1  | 15 (83.3%)                  | NE          | NA  |
|             | *S. aureus*         | <1          | <1   | <1   | 0 (0%)                      | 2           | 0   |
|             | *L. monocytogenes*  | <1          | <1   | <1   | 0 (0%)                      | 2           | 0   |
|             | *Salmonella spp.*   | Abs        | NA   | NA   | 0 (0%)                      | Abs         | 0   |

Abs, absence in 25 g; AMB, aerobic mesophilic bacteria; LAB, lactic acid bacteria; n, number of samples analyzed; NA, not applicable; NCS, noncompliant samples; NE, not established; SD, standard deviation; SF, smoked fish.

<sup>a</sup>Positive samples = samples in which the number of detected colonies is ≥ 1. <sup>b</sup>AL = acceptable limit according to Health Protection Agency (2009). <sup>c</sup>For each parameter, mean values followed by different letters indicate that they differ significantly between smoked versus SDF (p < 0.05).
PCR products were purified using purification kit (GenElute™
PCR Clean-Up; Sigma-Aldrich). Identification of the bacteria was
done by comparing their 16S rRNA sequence with those in data-
bases using www.ncbi.nlm.nih.gov/BLAST.

2.5  |  Data analysis

Statistical analyses were performed using STATISTICA 7.1. The
analysis of data was performed with Student’s t test, Mann–
Whitney U test, one-way analysis of variance (ANOVA), and
Kruskal–Wallis ANOVA. Significant difference was established at
p < 0.05, and means were separated using Student, Newman, and
Keuls range test.

3  |  RESULTS AND DISCUSSION

3.1  |  Microbiological characteristics of SF and SDF samples

Table 1 shows the microbial loads of SF and SDF samples. AMB and
LAB were the most frequent and dominant flora in both types of fish,
with AMB reaching concentrations up to 9.5 and 7.8 Log10 (CFU/g)
in the SF and SDF samples, respectively. Enterobacteriaceae, E. coli,
B. cereus, C. perfringens, yeasts, and molds were observed in number
of samples, while Salmonella spp., L. monocytogenes, and S. aureus
were not detected.

The minimum and maximum values recorded for each criterion
showed important variability within samples of each type of pro-
duct. This variation can be explained by the fact that the samples
were collected from various processors and sellers where the quality
of the raw material varied, as well as handling and hygiene practices.
Also, the density of AMB, Enterobacteriaceae, and E. coli was signifi-
cantly higher (p < 0.05) in SF than in SDF (Table 1). This is proba-
bly due to the fact that SF samples have a higher moisture content
(61 ± 11%) than the SDF ones (24 ± 11%). Also, in daily practices, SF
are often more handled than SDF both by processors and customers.

Furthermore, 66.7%, 50.0%, 38.9%, 11.1%, and 5.6% of SF were
not compliant with the stipulated limits for seafood products (Health
Protection Agency, 2009) regarding AMB, Enterobacteriaceae, E. coli,
B. cereus, and C. perfringens, respectively. Similarly, 22.2% and 5.6%
of SDF samples were not compliant with the acceptable limits for
AMB and Enterobacteriaceae, respectively. The high load of AMB
is likely due to a high contamination level of the raw material, and
these microorganisms were not fully eliminated during the smoking
treatment. The postprocess handling and storage conditions are also
potential sources of renewed pollution of the processed fish, as re-
ported by Kpodékon et al. (2014).

Thermosensitive bacteria like Enterobacteriaceae are used as indi-
cators of hygiene conditions and contamination of food after cooking
(Health Protection Agency, 2009). The detection of E. coli in ten sam-
ple (eight SF and two SDF) also suggested a contamination by fecal
matter from animal or human origin during postsmoking handling.

The presence of B. cereus and C. perfringens at levels above the
permitted limits may constitute a hazard to consumer’s health. B. ce-
reus and C. perfringens are foodborne pathogens mostly evoked in
gastrointestinal diseases in developed countries (Dierick et al., 2005;
EFSA & ECDC, 2016; Lindström, Heikinheimo, Lahti, & Korkeala,
2011). The presence of large numbers of molds, especially in SDF,
TABLE 4  Microbial load (Log$_{10}$ CFU/g) during the processing of smoked–dried fish (SDF) Cypselurus cyanopterus

| Parameters                  | Fresh fish (n = 3) | Cleaned fresh fish (n = 3) | SF (n = 6) | SDF (n = 6) |
|-----------------------------|--------------------|----------------------------|------------|-------------|
| AMB                         | 7.4 ± 0.8$^a$     | 6.5 ± 0.3$^b$             | 6.0 ± 1.2$^a$ | 5.1 ± 1.1$^b$ |
| Lactic acid bacteria        | 6.1 ± 2.0$^a$     | 4.4 ± 2.2$^b$             | 5.2 ± 1.1$^a$ | 3.9 ± 1.1$^b$ |
| Enterobacteriaceae          | 1.7 ± 1.0$^a$     | 2.1 ± 1.2$^a$             | 1.8 ± 1.8$^a$ | <1          |
| Escherichia coli            | <1                | <1                         | <1         | <1          |
| Bacillus cereus             | 1.2 ± 0.9$^a$     | <1                         | 1.0 ± 0.4$^a$ | 1.4 ± 0.8$^a$ |
| Clostridium perfringens     | <1                | <1                         | <1         | <1          |
| Yeasts                      | 1.8 ± 1.0$^a$     | 1.3 ± 0.7$^a$             | 1.9 ± 1.6$^a$ | 0.9 ± 0.5$^a$ |
| Molds                       | 2.3 ± 0.8$^a$     | 1.7 ± 1.3$^a$             | 2.5 ± 1.7$^a$ | 1.3 ± 0.9$^a$ |
| Staphylococcus aureus       | 1.0 ± 0.3         | <1                         | <1         | <1          |
| Listeria monocytogenes      | <1                | <1                         | <1         | <1          |
| Salmonella spp.             | Abs               | Abs                        | Abs        | Abs         |

Note. Legend as for Table 2.

also poses a risk, since they may produce mycotoxins during long-term storage (Job et al., 2016; Wogu & Iyayi, 2011).

As indicated in Table 2, no significant (p > 0.05) differences were observed between the microbial counts of samples from processing sites and those collected from markets regarding AMB, Enterobacteriaceae, B. cereus, C. perfringens, and yeasts in the case of SF. However, E. coli counts in SF samples collected from processing sites were significantly (p < 0.05) higher than those from markets. This could be explained by the additional smoking of leftover products intended to extend their shelf life. Likewise, mold counts in SF samples collected from markets were significantly higher (p < 0.05) than those from processing sites.

3.2 | Identification of MRS-associated bacteria

Bacteria grown on MRS constituted the dominant flora of both SF and SDF samples. Although MRS is a nonselective medium for LAB, it can also promote the growth of other microorganisms. 16S rRNA sequencing was performed on 12 CFU isolated on MRS. Five LAB species (Lactococcus garvieae, Pediococcus acidilactici, Weissella paramesenteroides, Enterococcus faecalis, and Enterococcus hirae) together with Klebsiella pneumoniae and Staphylococcus piscifermentans were found.

Klebsiella pneumoniae is known to possess histidine decarboxylase activity, enabling the bacterium to produce histamine in fish products (Visciano, Schirone, Tofalo, & Suzzi, 2012), which causes various health disorders to humans (Maintz & Novak, 2007). L. garvieae is found in aquatic environments (marine and freshwater aquaculture) and is a pathogen for fish (Vendrell et al., 2006). Wang et al. (2007) reported that it can be pathogenic for human with gastrointestinal disorder. E. faecalis has also been reported to cause endocarditis and diverse infections. Its transmission is nosocomial, but can also be done by food (Oprea & Zervos, 2007). Thus, beside the conventional microorganisms investigated for assessing the safety of ready-to-eat foods, SF and SDF samples also contained other potential pathogenic microorganisms exposing consumers to foodborne diseases. However, some of these bacteria can have positive effects. For instance, P. acidilactici produces a bacteriocin (Bacteriocin PA-1 or Pediocin AcH), which has an inhibitory effect on L. monocytogenes (Nieto-Lozano et al., 2010), and W. parameenteroides secretes a bacteriocin with a broad spectrum of inhibition of spoilage bacteria and food pathogens such as Salmonella typhimurium, Vibrio parahaemolyticus, or L. monocytogenes (Pal & Ramana, 2010).

3.3 | Changes in microbial loads during the processing of SF and SDF

Table 3 shows the evolution of the microbial loads during the processing of SF. AMB load in the raw frozen fish decreased significantly (p < 0.05) after the washing step (from 7.1 ± 0.5 to 6.2 ± 0.2), but remained stable after the smoking step (6.5 ± 0.4), close to the acceptable limit (<7 Log$_{10}$ CFU/g). This is surprising since a significant reduction in the microbial load of the fish was expected after the heat treatment. In addition, there were no significant (p > 0.05) changes in microbial loads for Enterobacteriaceae, B. cereus, yeasts, and molds during processing. Furthermore, potentially pathogenic organisms such as E. coli, C. perfringens, S. aureus, and Salmonella spp. were not detected in both fresh and SF samples.

Hot smoking as carried out in traditional processing units can induce a reduction in the microbial contamination comparable to pasteurization (Plahar, Nerquaye-Tetteh, & Annan, 1999). During the follow-up trials, temperature values recorded in the core of fish remained above 70°C during the last 30 min of the average duration of 90 min of smoking (data not shown). Since this temperature is expected to reduce the microbial load, the hypothesis of recontamination of the product during and after smoking is therefore likely. Indeed, the follow-up trials revealed many practices that could contribute to fish recontamination: (a) processors do not wear appropriate clothes during processing (no clean apron and charlotte) (b) the processing is performed in unhygienic conditions.
environment where the product is exposed to dust and flies, (c) processors use wastewater from raw fish washing to cool their hands, and (d) cement paper or frozen fish wrap is used to cover the end products.

Table 4 shows the evolution of the microbial loads during the processing of SDF. AMB and LAB counts decreased significantly (p < 0.05) at the end of the smoking–drying period (from 7.4 ± 0.8 to 5.1 ± 1.1 and 6.1 ± 2.0 to 3.9 ± 1.1 Log_{10} CFU/g, respectively). Moreover, Enterobacteriaceae were not detected at the end of the smoking–drying period. However, B. cereus, yeasts, and molds counts were not reduced significantly (p > 0.05), which could be explained by a recontamination of the product by these microorganisms during postprocess handling. As for SF, potential pathogenic bacteria such as L. monocytogenes were not detected at the end of the smoking–drying period. However, B. cereus, yeasts, and molds counts were not reduced significantly (p > 0.05), which could be explained by a recontamination of the product by these microorganisms during postprocess handling.

This study revealed that SF and SDF processed in Benin are not always of satisfactory microbiological status and represent potential sources of foodborne diseases. The quality of raw material, poor hygiene practices, and inappropriate handling practices during processing and selling is factors that contribute to the unsatisfactory microbiological quality of these products. Processors and sellers should be trained on good hygiene and handling practices in order to produce a safe and sound product for consumption.

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**CONFLICT OF INTEREST**

The authors have no conflicts of interest to declare.

**ETHICAL REVIEW**

This study does not involve any human or animal testing.

**INFORMED CONSENT**

Written informed consent was obtained from all study participants.

**ORCID**

Victor Anihouvi https://orcid.org/0000-0002-2609-3837

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Jacques Mahillon https://orcid.org/0000-0002-4463-1328

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