

Proceedings

1 Microbial succession and variation of physico-chemical charac-2 teristics and antioxidant capacity during spontaneous fermenta-3 tion of Mutchayan, a traditional fermented baobab derived 4 food * 5

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Abstract:

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17 Mutchayan is a traditionally fermented cooked cereal dough mixed to baobab pulp, consumed in Benin. The present study evaluated the physico-chemical and microbiological changes of this prod-18 uct during 0 to 120 hours of spontaneous fermentation. Analysis of the fermentation process re-19 vealed on one hand an increase of lactic acid and antioxidant capacity and a decrease of ascorbic 20 acid content. Dry matter content and Brix value decreased while pH did not significant change. 21 Microbiological analysis revealed the presence of molds at the beginning of the fermentation, which 22 growth was inhibited after 36 h, while the LAB and yeasts dominated the process. 23

Keywords: lactic acid bacteria; fermentation; yeast; antioxidant capacity; ascorbic acid

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1. Introduction

Fermentation is an old technic of food preservation, widely used in the world [1]. It 27 is a process based on biological activity of microorganisms for increasing food value 28 through the development of substances able to limit the growth of undesirable microor-29 ganisms in foods. It involves different categories of microorganisms depending on the 30 product and on the type of fermentation [2]. Materials rich in starch or glucose as cereals 31 and roots and tubers are usually submitted to acid and alcoholic fermentations. Acid fer-32 mentation improves the product flavor, the bioactive compounds content, minerals bio-33 availability, delays starch bioavailability [3] and provides probiotics for human health [4]. 34

Several products are derived from cereals acid fermentation: Mawe, [5], Ogi [6], de-35 rived from maize, Gowé [7] derived from sorghum, Mutchayan [8] derived from cereals 36 (maize or sorghum or millet). Mutchayan, a traditional food of "Otamari" socio-cultural 37 group of northern Benin, derives from the fermentation in a jar of cooked cereal dough, 38 mixed with diluted baobab (Adansonia digitata) fruit pulp, from 24 to 168 hours. It is char-39 acterized by a pH of about 4.2, lactic acid bacteria and yeast counts estimated at 7.6 log₁₀ 40 cfu/g and 7.2 log10 cfu/g respectively [8]. Mutchayan can be considered as promising, not 41 only because of the fermentation benefits, but also the functional properties reported for 42

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the baobab fruit pulp. Therefore, the present study aimed to evaluate the physico-chemical, microbiological and nutritional changes occurring during the fermentation process.

2. Materials and Methods

2.1. Materials

Red sorghum grains (Sorghum bicolor (L.) Moench) purchased on a local market of Abomey-Calavi (southern Benin), and baobab (Adansonia digitata) fruit pulp purchased in a local processing unit of baobab fruits, at Boukoumbé (northern Benin), were used for the production of Mutchayan.

2.2. Mutchayan production and sampling

Mutchayan was produced based on the traditional technology [8]. Around of 160 g of 10 sorghum grains flour was mixed in 1.5 L of water for making a porridge which was heated till boiling; then, 160 g of the flour was added to the porridge and mixed thoroughly. About 900 mL of diluted baobab fruit pulp (220 g in 760 mL of water) was immediately 13 added to the cooked dough. The mix was cooled, and let to spontaneously ferment at 14 room temperature during 120 hours, in a covered plastic bucket. Along this process, the 15 temperature was monitored with ibutton devices (temperature data logger), and samples 16 were collected at 0, 6, 12, 24, 36, 48, 72, 96 and 120 hours to assess the total viable count 17 (TVC), yeast count (YC), molds count (MC), lactic acid bacteria count (LABC), Enterobac-18 teriaceae count (EC), pH, Brix value (BV), total titratable acidity (TTA), dry matter (DM), 19 ascorbic acid content (AAC) and antioxidant capacity (VCEAC). 20

2.3. Nutritional and physico-chemical analyses

The extraction process for the antioxidant capacity was performed as follows accord-22 ing to Thaiponget al. [9], with modifications. Two grams (02 g) sample was mixed to 15 23 ml of 60% methanol solvent in falcon tubes (50 ml); preliminary works revealed 60% meth-24 anol solvent as the adequate mixture (methanol/water) for the maximal extraction of an-25 tioxidants from baobab fruit pulp. The antioxidant capacity was evaluated using DPPH 26 assay, according to Brand-Williamset al. [10], as described by Thaipong et al. [9], with 27 some modifications. The results were expressed as mg VCEAC (Vitamin C Equivalent of 28 Antioxidant Capacity) for 100 g of the product (dry weight). 29

The ascorbic acid content was determined by titrimetric method ISO 6557/2:1984; the results were expressed as mg/100mg dw (dry weight) of Mutchayan samples.

The pH of the collected samples was assessed with a pH meter, according to ISO 32 1842:1991. Total titratable acidity (TTA) was assessed according to ISO 750:1998, using the potentiometric method; 10 g of the sample was diluted in distilled water to make 100 ml of suspension which was titrated with 0.1 N hydroxide sodium (NaOH) solution. Dry 35 matter content was determined according to AOAC method 2.166 (1980). The Brix value 36 was determined using a digital refractometer, according to ISO 2173:2003. 37

2.4. Microbiological analyses

The general requirements and guidance for microbial examinations (ISO 7218:2007) 39 were used to prepare required dilutions and culture media. Total flora and lactic acid bac-40 teria (LAB) were enumerated on Plate Count Agar (PCA) and Man Rogosa and Sharpe 41 (MRS) media respectively; the Petri dishes were incubated at 30 °C for 72 h according to 42 the standard ISO 4833-1:2013 for total flora (TVC), and according to ISO 15214:1998 for 43 LAB. For yeasts and molds, 100 μ l of the considered dilution was surface inoculated on 44 solidified Sabouraud Dextrose Chloramphenicol Agar medium; incubation was per-45 formed at 25 °C for 72 to 120 h, according to ISO 21527-2:2008. Enterobacteriaceae count 46 was assessed by inoculating on Violet Red Bile Glucose (VRBG) medium; dishes were 47 incubated at 37 °C for 24 h, according to BS ISO 21528-2:2004. The expression of the results 48 was done according to ISO 7218:2007. 49

2.5. Statistical analyses

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The collected data were analyzed with R 4.0.5 software. An analysis of variance was 1 realized to assess the significance of the variability of physico-chemical parameters. Pois-2 son family models (Poisson and negative binomial models) were used for analyzing the3 variability of microbiological count data (Total viable count, yeast and molds, lactic acid4 bacteria and Enterobacteriaceae). Barplot test was performed as post-hoc test.5

3. Results

3.1. Changes of the physico-chemical characteristics of Mutchayan during fermentation

Table 1 shows that, from 0 h to 120 h, the titratable acidity increases (p<0.00) from</th>8 5.7 ± 0.1 to 6.8 ± 0.1 g of lactic acid for 100 g of the product (dry weight), while the pH did9not change. The dry matter contents and the Brix value decreased significantly (p<0.00)</td>10from 19.8±0.0 to 16.7±0.1 g/100g and from 7.6±0.11 °Bx to 5.0±0.0 °Bx. The vitamin C con-11tent and the antioxidant capacity also varied significantly (p<0.05) from 91.7±1.1 to</td>12 69.8 ± 4.0 mg/100 dw and 657.1 ± 1.2 to 802.9 ± 48.9 mg VCEAC /100g (dw) respectively.13

Table 1. Physico-chemical characteristics of sorghum flour, baobab pulp and Mutchayan14

	ıcts an- /zed	рН	°Brix	TTA (LA g/100g dw)	DM (g/100g)	Vitamin C (mg/100g dw)	Antioxidant ca- pacity (VCEAC mg/100g dw)
M. fer	0h	3.4±0.0 ª	7.3±0.3 ª	5.8±0.1 ^{e.f}	18.7±0.1 ^b	91.7±1.1 ^a	693.5±49.0 a.b
Mutchayan sa ferent lengths	6h	3.4±0.0 ª	7.6±0.1 ª	5.7 ± 0.1 f	19.8±0.0 ª	84.7 ± 3.8 a.b	699.2±51.7 ^{a.b}
hay ler	12h	3.3±0.0 ª	7.5±0.4 ª	6.0±0.0 d.e	18.6±0.2 ^b	79.1±5.0 ^{a.b}	677.7±0.7 ^{a.b}
an s ngtl	24h	3.3±0.0 ª	7.4±0.2 ª	6.1±0.0 d.e	18.3±0.0 ^c	69.8±4.0 ^b	657.1±1.2 ^b
san hs c	36h	3.4±0.0 ª	6.8±0.5 ª	6.3±0.0 ^{c.d}	18.1±0.0 ^c	78.8 ± 8.1 ^{a.b}	713.8±43.1 ^{a.b}
samples hs of fer	48h	3.4±0.0 ª	6.0 ± 0.4 a.b	6.5±0.1 ^{a.b.c}	17.5 ± 0.0 d	80.8 ± 1.1 ^{a.b}	802.9±48.9 a
	72h	3.3±0.0 ª	6.2±0.9 a.b	6.8±0.1 ^a	16.9±0.0 ^e	76.6±0.9 a.b	766.1±0.0 a.b
of d men	96h	3.3±0.0 ª	5.0±0.0 ^b	6.6±0.1 ^{a.b}	16.9±0.1 ^e	79.3±0.0 ^{a.b}	751.5±17.9 ^{a.b}
dif- enta-	120h	3.4±0.1 ª	5.1±0.4 ^b	6.3±0.1 b.c.d	16.7±0.1 ^e	82.6±3.9 a.b	798.2±23.2 ^a
	ghum our	6.1±0.0	8.2±0.3	0.6±0.0	91.3±0.1	0.0±0.0	-
Baoba	Baobab pulp		73.4±1.7	14.4 ± 0.1	87.3±0.5	299.3±4.4	3004.6±1.6

The mean values with different letters in column are significantly different at 5% threshold. TTA:15Total titratable acidity; DM: Dry matter;16

3.2. Microbial changes during Mutchayan fermentation

Different types of microorganisms were observed during the fermentation. Among 18 the lactic acid bacteria, two main types of colonies were observed: the colonies with a halo 19 (LABh) and the others without halo (LABwh) (Figure 1); the same tendance was observed 20 for yeasts: large yeast colonies (LY), and small yeast colonies (SY) which appeared on plate 21 whiter and lighter (Figure 1). 22

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LABh: Lactic acid bacteria with halo; LABwh: Lactic acid bacteria without halo; SY: Small yeast; LY: Large yeast; M: Molds

Figure 1. (a) Lactic acid bacteria in *Mutchayan;* (b) Yeast and mold in *Mutchayan.*

At the beginning of the fermentation (0 h), the total viable count was estimated at 2 about 3.5±3.4 log₁₀ cfu/g (Table 2), while the microbial activity of lactic acid bacteria, yeasts 3 and Enterobacteriaceae was observed at 12 hours of fermentation. Before 12 h, the dough 4 temperature was decreasing from 57.8±3.9 °C to 30.5±0.0, and these microorganisms' loads 5 were under 1 log₁₀ cfu/g. While these microorganisms were absent, molds were revealed 6 as dominant flora (at 0 h), with a population estimated at 2.7 log₁₀ cfu/g; after an increase 7 between 24 and 36 h, where their load reached a maximum estimated value of 6.9 log₁₀ 8 cfu/g, mold growth tended to be inhibited and the load decreasing below 5.6 log10 cfu/g at 9 the end of the fermentation. The total viable count increased significantly (p<0.05) from 10 12 to 36 h, reaching 7.8±7.8 log10 cfu/g and keep constant till 120 h, with a dough temper-11 ature ranging from 28.0 to 32.3 °C (Table 2). 12

Duration TVC YC SYC LYC MC LABC LABwhC LABhC EC 3.5±3.4 ° <1 <1 <1 0h <1 <1 2.7±0.0 <1 <1 6h <1 <1 <1 <1 <1 <1 <1 3.6±3.4 ° 3.0±0.0 12h 3.8±2.9 ° 3.1±0.0 d 3.1±0 d <1 2.7±0.0 2.7±0.0 d 2.7±0.0 ° <1 1.7±1.2 b 24h 6.9±7.0 b 7.0±5.9 ° 7.0±0.0 ° 6.1±0.0 ° <6.6 6.7±6.7 ° 6.5±6.5 b 6.2±6.3 b <1 7.8±7.8 a 7.3±6.7 b,c 7.4±7.1 b 7.2±6.9 a.b 2.1±0.9 a 36h 7.4±6.9 b 6.4±0.0 b,c 6.9±0.0 7.0±6.6 a 48h 7.7±6.8 a 7.9±7.4 ª 7.8±7.0 a,b 7.3±0.0 a <5.6 7.8±6.4 a.b 7.8±6.4 ª <5 <1 72h 8.0±7.6 a 8.0±7.7 a 7.9±7.7 a <6.6 7.8 ± 7.6 a.b 7.5±7.3 ª 7.5±7.3 ª 6.8±0.0 b,c 8.2±7.8 ^a 8.0±7.1 ª 96h 7.9±6.9 a 7.9±6.1 a,b 7.1±0.0 a,b 6.6±0.0 8.1±6.9 ^a 7.2±6.6 a 120h 8.2±7.4 ª 8.1±6.1 ª 8.0±7.4 ª 7.6±0.0 a <5.6 8.1±6.9 a 8.0±6.9 a 7.5±7.2 ª

Table 2. Microorganisms count during Mutchayan fermentation

The mean values with different letters in column are significantly different at 5% threshold. The values in bold represented the estimated count of microorganisms; TVC: Total viable count; YC: Yeast Count; SYC: Small Yeast count; LYC: Large Yeast Count; MC: Molds count; LABC: Lactic Acid Bacteria Count; LABwhC: Lactic acid bacteria without halo count ; LABhC:

Lactic acid bacteria with halo Count; EC: Enterobacteriaceae Count

Yeast count increased significantly (p<0.05) (Table 2) during the fermentation process, from less than $3.1\pm0.0 \log_{10}$ cfu/g after 12 hours to $7.9\pm7.4 \log_{10}$ cfu/g after 48 h of fermentation; only small yeasts colonies were detected at 12 h of fermentation. A significant increase of yeast population was observed from 12 to 24 h, for small yeast colonies and for large yeast colonies as well. Yeast population stayed statistically the same after 48 h of the fermentation; the same trends were observed for small yeasts count and large yeasts count which were respectively $7.8\pm7.0 \log_{10}$ cfu/g and $7.3\pm0.0 \log_{10}$ cfu/g.

Similarly to yeasts, lactic acid bacteria (LAB) count rapid increased between 12 and 21 36 h from 2.7±0.0 to 7.4±7.1 log₁₀ cfu/g; after 48 h, the lactic acid bacteria count did not 22

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change significantly and stayed at 7.8±6.4 log¹⁰ cfu/g (Table 2). Both lactic acid bacteria with and without halo presented similar growth tendencies.

The enterobacteriaceae count increased from <1 to $1.7\pm1.15 \log_{10}$ cfu/g between 6 and312 h, and decreased from 1.7 ± 1.15 to $<1 \log_{10}$ cfu/g between 12 and 24 h. After 36 h of4fermentation, similar observations were made; the enterobacteriaceae load increased from5<1 to $2.06\pm0.85 \log_{10}$ cfu/g, and later decreased to $<1 \log_{10}$ cfu/g. Enterobacteriaceae appear6to not play an important role during the fermentation of *Mutchayan*.7

4. Discussion

Yeasts and lactic acid bacteria are the main microorganisms counted during the fer-9 mentation of Mutchayan. During the first 12 hours of the fermentation, the microbial ac-10 tivity was limited and no significant change was observed on physico-chemical parame-11 ters till 24 h. After 24 h of fermentation, the total flora and the lactic acid bacteria load 12 became higher (about 6 log10 cfu/g), and that might allow rapid increase of lactate pro-13 duction from 36 to 120 h, leading to an increase of the titratable acidity content between 14 the beginning and the end of the fermentation. Despite the lactic acid production, the 15 quantity produced during the fermentation was not sufficient to decrease the already acid 16 pH of Mutchayan (pH=3.4 at 0 h of fermentation) as observed for the fermentation of ce-17 real-based products as Mawe [11] and Gowe [7], where the pH diminished from 6.1-6.2 to 18 3.5-3.6 [11], almost the pH of Mutchayan at 0 h of fermentation. 19

The absence of LAB and yeasts at the beginning of *Mutchayan* fermentation could be 20 linked to the cooking process applied and the relatively high temperature of the dough 21 during the first 4 hours (58 to 41 °C). But the presence of the molds at the same time is 22 probably related to their ability to survive in critical conditions; some mold species can 23 grow at low pH values (pH = 3.2) and low water activity [12]. The inhibition of molds 24 growth along the fermentation process could be linked to the LAB activity because it has 25 been proven that some strains of Lactobacillus sp. have a strong inhibitory capacity of fun-26 gal growth; that is the case of Lactobacillus plantarum strain ITEM 17215 which inhibit the 27 growth of mycotoxigenic fungi as Aspergillus flavus [13, 14]. 28

As the fermentation duration increased, yeasts and lactic acid bacteria counts became 29 higher (6-8 log¹⁰ cfu/g) than at the beginning, inducing a significant increase of antioxidant 30 capacity, associated to the activity of lactic acid bacteria without halo which should be 31 deeply studied for their relation with the increasing of antioxidant capacity. Association 32 of LAB and yeasts were found to be responsible for *Mutchayan* fermentation, which agrees 33 with the results of Chadare et al. [8] and other studies of indigenous cereal-based fermented foods, as Kenyan *Busaa, Kaffir* beer, Nigerian *Ogi, Pito, Sekete* and *Busa* [15]. 35

The occurrence of Enterobacteriaceae in *Mutchayan*, which was observed in other products, e.g. in *Gowé* [16], in *Calugi*, [17], could be linked to unmastered surrounding conditions and the fermenter (plastic bucket) which was not tightly covered. During *Mutchayan* fermentation, the decrease of Enterobacteriaceae load could be due to the low pH values or the increase of the lactic acid bacteria activity, since it was reported that LAB produce bacteriocins, effective to control their growth [18, 19].

5. Conclusion

The fermentation of *Mutchayan* is characterized by the domination of molds during 43 the 12 first hours. Their growth was inhibited by the development of yeast and lactic acid 44 bacteria which dominated the fermentation. Along this microbial changes, a decrease of 45 Brix value, dry matter, and an increase of lactic acid and antioxidant capacity were observed with the increasing of duration. Further investigations are needed for identifying 47 the different microbial species and their specific role in the fermentation of *Mutchayan*. 48

Conflicts of Interest

The authors declare there is no conflict of interest

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