Yeasts from a traditional ferment “Rabilé”: Molecular and physiological characterization

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“Rabilé” is a popular traditional ferment in Burkina Faso, consisting mainly of yeasts. It is used as a food supplement or additive like Single Cell Protein (SCP). The present work focused on identifying yeast microbiota in local food, and studying their growth kinetic parameters. “Rabilé” sampling from the 13 regions of Burkina Faso was used to isolate yeast strains. Molecular methods, including PCR-RFLP, Sanger Sequencing, and Single Locus Analysis, were applied for strain identification. The kinetic parameters were determined in batch culture. The results show 390 isolates belonging to 12 species with a predominance of Saccharomyces cerevisiae, followed by Cutaneotrichosporon curvatus. Among the selected strains, S. cerevisiae OG22 and Kluyveromyces marxianus KY01 showed the highest maximum growth rate (0.566 and 0.568 h-1) concerning kinetic parameters. These results demonstrate that “Rabilé” is an important biotope of yeast strains, and could be a potential food supplement.

Key words: Yeast, Rabilé, traditional ferment, molecular identification, kinetic parameters.

INTRODUCTION

Yeasts have played an essential role in the human diet for millennia (Bekatorou et al., 2006; Gómez-Pastor et al., 2011; Steensels and Verstrepen, 2014). They are used in brewing, bread-making, winemaking, and producing many other fermented foods worldwide (Bekatorou et al., 2006; Carrau et al., 2015). Nowadays, advances in genetic engineering (recombinant protein) have made it possible to extend their application in many fields such as the environment, the food industry, and health (Feldmann, 2012; Fallah et al., 2016; de Souza Varize et al., 2019).

In West Africa, yeasts are mainly used to produce ethyl alcohol and fermented foods (Somda et al., 2011; Tankoano et al., 2017; Johansen et al., 2019). The production of fermented foods is an income-generating activity for local producers (Lyumugabe et al., 2012; Djêgui et al., 2015), who provide enriched and inexpensive food. Yeasts improve the nutritional and organoleptic quality of these foods by providing additional...
proteins, vitamins, minerals, and various metabolites (Suman et al., 2015; De Vuyst et al., 2016; Onofre et al., 2017; Johansen et al., 2019; Tofalo et al., 2019). Thus, they contribute to a balanced diet in West Africa, where undernourishment affects 15% of the population (FAO, 2020).

In Burkina Faso, “Rabilié” is a popular traditional ferment that mainly contains yeasts. It is obtained at the end of sorghum beer production after a semi-controlled fermentation and corresponds to the dried lees (Korlani et al., 1996a, 1996b). Specific pathogenic and opportunistic yeasts can probably proliferate, and produce compounds that are harmful to the body (Steensels and Verstrepen, 2014; Johansen et al., 2019; Tofalo et al., 2019). On the other hand, this semi-controlled fermentation denotes a great diversity of wild yeasts from “Rabilié” with unexplored potential. Empirically, the incorporation of “Rabilié” in sauces and ready meals is similar to Single Cell Protein (SCP) as a food supplement or additive. We, therefore, believe that “Rabilié” is an alternative for the treatment and prevention of malnutrition.

For all these reasons, we hypothesize that “Rabilié” resulting from a process of semi-controlled fermentation constitutes a source of diversified wild yeasts potentially usable for food enrichment.

Yeasts’ diversity analysis is mainly based on molecular genotyping. Esteve-Zarzoso et al. (1999) proposed a method of differentiating yeasts based on enzymatic restriction of the 5.8S-ITS region. This technique has been proven helpful in several diversity studies, particularly for yeasts in wine and beer (Granchi et al., 1999; Diaz et al., 2013; Corbett et al., 2019). However, to have a suitable level of discrimination for genetic analysis of populations, molecular techniques such as sequencing and microsatellites analysis are needed. This study aimed to identify the yeast microbiota in a traditional ferment, and to study their kinetic parameters.

**MATERIALS AND METHODS**

**Sampling of “Rabilié” (traditional ferment)**

Sampling was conducted in 13 sites representing the 13 administrative regions of Burkina Faso (Banfora, Bobo-dioulasso, Dédougou, Dori, Fada N’Gourma, Gaoua, Kaya, Kombissiri, Koudougou, Ouagadougou, Ouahigouya, Tenkodogo, Ziniaré). In each location, three local producers were randomly chosen from which one sample of “Rabilié” was collected (approximately 50 g). Samples collected were aseptically packaged in stomacher bag (Stomacher® 400 Classic Bags, Seward, UK), and transported under cold conditions (4°C).

**Isolation of yeasts from samples of traditional ferments**

To make serial decimal dilutions, 10 g of each sample were diluted in 90 mL of sterile physiological solution (0.9% NaCl). A volume of 100 µL of each dilution was inoculated on Sabouraud media with chloramphenicol. The set was incubated at 30°C for 48 to 72 h (Guimarães et al., 2006).

Well-individualized colonies were selected and purified by a series of three successive sub-culturing. A total of 30 isolates per site was selected, and stored at -20°C in Yeast Extract Peptone Dextrose Broth (YPD Broth) containing 20% glycerol for molecular and physiological characterization.

**Molecular characterization of yeast isolates**

**PCR-RFLP**

DNA was extracted from each yeast isolate using Cetyl Trimethylammonium Bromide (CTAB) buffer from a fresh culture (Kumar et al., 2014). The 5.8S-ITS region was amplified using the primer pair ITS1 (5’-TCCGTAGTGAACTGCGG-3’) and ITS4 (5’TCCCTCCTATTGATATGC-3’) as described by White et al. (1990). According to the manufacturer’s instructions, PCR products were digested separately using three restriction enzymes HhaI, HaelIII, and Hinfl (NEW ENGLAND Biolabs® Inc.). The PCR products and their restriction fragments were separated on a 2.5% agarose gel. After electrophoresis, gel was photographed under ultra-violet light of the transilluminator (UVP®) coupled to a computer. All fragments were analyzed with UVP Doc-It™ Ls Analysis software (Version 6.8.2) to determine their sizes. Isolates were then clustered according to their restriction profiles.

**Sequencing of the 5.8S-ITS region**

One strain was chosen from each restriction profile group for sequencing. The PCR products were purified (NEW ENGLAND Biolabs® Inc.) under a combined action of two hydrolytic enzymes (Exonuclease I and Shrimp Alkaline Phosphatase). The obtained products were re-amplified using the Genomelab-DC TS Quick Start Kit (Beckman Coulter, USA) following the cycle sequencing program of 30 cycles of denaturation (96°C, 20 s), annealing (50°C, 20 s), and extension (60°C, 4 min). Cycle sequencing products were purified by ethanol precipitation, and then separated using a CEQ™ 8000 Genetic Analyzer Sequencer. Finally, the isolates’ phylogenetic tree was constructed using the Geneious Prime software (Version 2020.0.3) from the sequences obtained using Tamura-Nei as Genetic Model Distance and Neighbor-Joining as Tree Build Method.

**Single locus analysis**

Three isolates of Saccharomyces cerevisiae were randomly selected per site to obtain 39 isolates for single-locus analysis. The ScAAT1 locus was amplified as described by Legras et al. (2005) using a pair of ScAAT1 primers represented by the ScAAT1 forward labeled (Cy5) and ScAAT1 reverse unlabeled. Using CEQ™ 8000 Genetic Analyzer Sequencer and DNA standard size 400 bp, the amplified products were separated by capillary electrophoresis. GenAlEx version 6.502 was used to determine diversity parameters.

**Study of kinetic parameters**

It consisted of monitoring the yeasts’ growth in a non-renewed medium and determining their kinetic parameters by following the steps that follow.
Table 1. Distribution of isolates according to the similarity of restriction profiles.

| Profile | Number of isolates | PCR product size (bp) | Restriction fragments size (bp) |
|---------|--------------------|-----------------------|--------------------------------|
|         |                    |                       | Hha | Haelli | HinfI |
| 1       | 322                | 874                   | 376+340+141 | 323+239+178+130 | 374+135 |
| 2       | 002                | 629                   | 305+230+100 | 414+218         | 344+215 |
| 3       | 002                | 500                   | 300+234   | 414+95          | 267+250 |
| 4       | 005                | 659                   | 224+200+157+105 | 331+187       | 333+188+136 |
| 5       | 026                | 516                   | 275+257   | 478             | 290+228 |
| 6       | 007                | 721                   | 287+191+165+89 | 608          | 239+177+103 |
| 7       | 005                | 362                   | 206+167   | 364             | 182+168 |
| 8       | 001                | 355                   | 206+167+143 | 371           | 177+97  |
| 9       | 001                | 390                   | 218+94    | 393             | 188+95  |
| 10      | 007                | 500                   | 218+183+77+62 | 400          | 218+150+136 |
| 11      | 007                | 350                   | 160+143+65 | 364            | 180+100+71 |
| 12      | 004                | 600                   | 296+257   | 400+120         | 311+280 |
| 13      | 001                | 410                   | 414        | 421            | 212+142 |

Strains were first grown on YPD agar for 24 h. Then pure colonies obtained were inoculated into YPD broth incubated under continuous agitation overnight at 30°C to constitute a stock culture. Pre-culture was performed by inoculating 6 µL of stock culture into fresh YPD Broth incubated under continuous agitation overnight at 30°C. Cells were harvested from pre-culture by centrifuging at 3000 g for 2 min to constitute an inoculum of 1.38 × 10⁷ cells/mL.

A volume of 10 µL (1.38×10⁷ cells) of the inoculum (1.38×10⁷ cells/mL) was transferred to the wells of a microplate (containing 290 µL of YPD Broth), except the three wells used as blank (control with no cells). The test was carried out in triplicate. The growth at 30°C was then monitored using a microplate reader spectrophotometer (Biotek ELx808, USA). The cellular biomass (Optical Density OD₆₀₀) was recorded every 20 min for 24 h. A growth curve (OD as a function of time) was performed to determine the kinetic parameters: maximum specific growth rate (μmax) and generation time (tg). Analysis of variance was applied to assess significant kinetic differences.

RESULTS

390 yeast isolates were obtained from “Rabilé” samples collected from the 13 regions of Burkina Faso (Table 1).

Molecular characteristics of yeasts from “Rabilé”

Based on the size of the PCR products and the restriction patterns obtained, 390 yeast isolates were clustered into 13 genetic groups corresponding to 13 distinct restriction profiles (Table 1).

Similarity of sequences

Homology search results revealed 12 species among the 13 genetic groups with a similarity of 98.1 to 100% (Table 2).

According to the sampling sites, we found seven species of yeast in samples from the Ziniaré site; five species in samples from Ouagadougou; four species in samples from Tenkodogo and Fada N’Gourma; three species in samples from Bobo-Dioulasso and Ouahigouya; two species in samples from Banfora, Dori, Gaoua, Kaya, Kombissiri, and Koudougou, and one species in samples from Dédougou (Figure 1). Figure 2 represents the Neighbor-Joining tree of these yeast species isolated from “Rabilé.”

A total of 38 Saccharomyces isolates were successfully analyzed using a single locus assay. We obtained 28 genotypes and 18 alleles with 0.902 as the Nei’s genetic diversity index for these isolates.

Kinetic parameters

The maximum specific growth rate is inversely proportional to the generation time, and is one of the most important parameters in modelling microbial growth (Barbera et al., 2019).

S. cerevisiae

The analysis of kinetic parameters showed that maximum specific growth rate (μmax) and generation time (tg) were significantly variable between isolates of S. cerevisiae. According to the classification of Oliveira et al. (2004), three of our isolates of S. cerevisiae (OG22, OG26, TK03) had a high μmax. Fifteen isolates reached a medium level. The isolate OG22 showed a high growth rate similar to that of the industrial strain used as a control (Table 3).
Table 2. Homology search for the sequences obtained.

| Isolate name | Accession number* | Sequence length (bp) | Quality (HQ %) | Similarity (%) | Query coverage | Species | Profile |
|--------------|-------------------|----------------------|---------------|----------------|----------------|---------|---------|
| BB15         | MW600341          | 104                  | 100           | 98.1           | 100            | Saccharomyces cerevisiae | 1 |
| BB21         | MW582617          | 413                  | 100           | 100            | 100            | Rhodotorula mucilaginosa | 2 |
| DR23         | MW582862          | 441                  | 99.5          | 100            | 100            | Candida parapsilosis     | 3 |
| FD01         | MW600340          | 215                  | 100           | 100            | 100            | Exophiala dermatitidis   | 4 |
| TK22         | MW583114          | 446                  | 100           | 100            | 100            | Cutaneotrichosporon curvatus | 5 |
| KY01         | MW592701          | 522                  | 100           | 99.8           | 100            | Kluyveromyces marxianus  | 6 |
| OG11         | MW592876          | 278                  | 100           | 100            | 100            | Yarrowia lipolytica      | 7 |
| OG12         | MW592844          | 158                  | 100           | 93.7/91.8      | 50/100         | Yarrowia lipolytica / Yarrowia phanggaensis | 8 |
| OY30         | MW582897          | 240                  | 100           | 100            | 100            | Clavispora lusitaniae    | 9 |
| TK23         | MW592836          | 342                  | 100           | 100            | 100            | Pichia kudriavzevi       | 10|
| ZN06         | MW592921          | 303                  | 100           | 99.7           | 100            | Yarrowia phanggaensis    | 11|
| ZN25         | MW592744          | 432                  | 100           | 99.3           | 100            | Meyerozyma cariblica     | 12|
| ZN26         | MW592740          | 322                  | 100           | 100            | 100            | Kodamaea ohmeri          | 13|

*Sequence accession number in Genbank NCBI.

Figure 1. Map of Burkina Faso illustrating the diversity of "Rabilé" yeasts according to the sampling site.

Non-saccharomyces species

Maximum specific growth rate and generation time were also significantly variable between species (Table 4). Kluyveromyces marxianus had a high maximum specific growth rate, while Yarrowia phanggaensis had the lowest.
DISCUSSION

This study highlights the “Rabilé” yeasts diversity from different production sites. It was hypothesized that “Rabilé” could contain a varied range of yeasts, as this starter results from uncontrolled alcoholic fermentation. It occurs in a non-sterile environment so that several microorganisms can intervene to enhance fermentation or as spoiling agents (Mogmenga et al., 2017). Nine yeast species identified in this study (S. cerevisiae, K. marxianus, Pichia kudriavzevii, Meyerozyma caribbica, Candida lusitaniae, Candida parapsilosis, Rhodotorula mucilaginosa, Yarrowia lipolytica, Kodameae ohmeri) are yeasts commonly encountered in fermented foods (N’guessan et al., 2011; Johansen et al., 2019). Some of these yeasts (Y. lipolytica, S. cerevisiae, and K. marxianus) belong to the list of microorganisms, and derived products approved by the Food and Drug Administration (FDA). It is reassuring that yeasts from traditional fermentations do not constitute a health risk for consumers in general (Steensels and Verstrepen, 2014).

Yeasts known to produce alcohol have been found in the ferment, including S. cerevisiae, K. marxianus, P. kudriavzevii, and M. caribbica (Djégui et al., 2015; Tolieng et al., 2018; Johansen et al., 2019). These strains may be potential candidates for standardized starter production. Some species involved in organoleptic and nutritional quality improvement have been isolated from

Figure 2. Phylogenetic tree of the 12 yeast species isolated from the “Rabilé”.

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Table 3. Kinetic parameters of *S. cerevisiae* isolates.

| ID    | $\mu_{\text{max}}$ (h$^{-1}$)* | $t_{\text{g}}$ (h) | Level  |
|-------|-------------------------------|-------------------|--------|
| SC1   | 0.593$^a$                     | 1.170             | High   |
| OG22  | 0.566$^a$                     | 1.226             | High   |
| OG26  | 0.557$^{ab}$                  | 1.244             | High   |
| TK03  | 0.555$^{abc}$                 | 1.248             | Medium |
| BF18  | 0.518$^{abcd}$                | 1.337             | Medium |
| KD05  | 0.518$^{bcd}$                 | 1.337             | Medium |
| TK05  | 0.513$^{bcd}$                 | 1.350             | Medium |
| ZN03  | 0.512$^{cdde}$                | 1.353             | Medium |
| DD29  | 0.510$^{de}$                  | 1.359             | Medium |
| KY24  | 0.500$^{def}$                 | 1.386             | Medium |
| BB19  | 0.499$^{def}$                 | 1.388             | Medium |
| KD06  | 0.498$^{def}$                 | 1.391             | Medium |
| FD25  | 0.487$^{defgh}$               | 1.423             | Medium |
| GA16  | 0.485$^{defgh}$               | 1.429             | Medium |
| GA13  | 0.482$^{defgh}$               | 1.437             | Medium |
| KD16  | 0.479$^{defgh}$               | 1.446             | Medium |
| DR06  | 0.469$^{efgh}$                | 1.479             | Medium |
| KB02  | 0.464$^{gh}$                  | 1.493             | Medium |
| OY24  | 0.460$^{ghi}$                 | 1.506             | Medium |
| BF20  | 0.446$^{ghij}$                | 1.553             | Low    |
| OY05  | 0.445$^{ghijk}$               | 1.559             | Low    |
| BB15  | 0.433$^{ij}$                  | 1.600             | Low    |
| KD04  | 0.432$^{ij}$                  | 1.604             | Low    |
| KB23  | 0.431$^{hjkm}$                | 1.607             | Low    |
| DR25  | 0.427$^{hjkm}$                | 1.622             | Low    |
| OY23  | 0.426$^{ijkl}$                | 1.649             | Low    |
| BF08  | 0.417$^{klm}$                 | 1.662             | Low    |
| DD09  | 0.412$^{klm}$                 | 1.684             | Low    |
| FD24  | 0.411$^{klm}$                 | 1.685             | Low    |
| DD17  | 0.405$^{klm}$                 | 1.712             | Low    |
| DR07  | 0.404$^{klm}$                 | 1.716             | Low    |
| KY06  | 0.401$^{klm}$                 | 1.729             | Low    |
| ZN24  | 0.395$^{lmno}$                | 1.736             | Low    |
| OG01  | 0.388$^{lmno}$                | 1.788             | Low    |
| KY21  | 0.368$^{lm}$                  | 1.883             | Very low |
| GA07  | 0.356$^a$                     | 1.948             | Very low |
| ZN10  | 0.307$^a$                     | 2.257             | Very low |
| BB07  | 0.303$^a$                     | 2.288             | Very low |
| TK25  | 0.218$^a$                     | 3.186             | Very low |
| KB13  | 0.203$^a$                     | 3.422             | Very low |

ANOVA Fisher (LSD) p-value $<0.0001$ -

*Values with the same letters are not significantly different.

the traditional ferment “Rabilé”. These yeasts (*K. marxianus*, *R. mucilaginosa*, and *Y. lipolytica*) would act with lactic acid bacteria to provide each production’s sensory characteristics and health benefits (Sørensen et al., 2011; Calabretti et al., 2012; Johansen et al., 2019; de Souza Varize et al., 2019).

In contrast, other species reported as opportunistic pathogens (*C. lusitaniae*, *C. parapsilosis*, *Exophiala*) have been detected. These yeasts are associated with producing harmful substances, fungal infections, and even respiratory diseases (Pires et al., 2016). The presence of these opportunistic pathogenic yeasts in the
“Rabilé” could constitute a health concern, especially since the resulting sorghum beer “Dolo” is consumed directly without pasteurization.

There is no systematic link between a yeast’s presence in a ferment, and its presence in the final product. N’guessan et al. (2011) pointed out that certain non-Saccharomyces yeast species appear sporadically, and may not appear in the final product. Nonetheless, non-Saccharomyces yeasts require safety tests beforehand (Steensels and Verstrepen 2014). Taxonomically, the data obtained indicate a predominance of hemi-ascomycetes. The oleaginous yeasts belonging to the group of hemi-ascomycetes appeared to be closer to the hemi-basidiomycetes present in “Rabilé” (Figure 2).

In contrast to previous studies, our findings prove that the yeast microbiota of “Rabilé” is more diverse with 11 additional species compared to two species (Konlani et al., 1996a, van der Aa Kühle et al., 2001). We are of the opinion that yeast microbiota from “Rabilé” could be more diverse in reality because the culture-dependent method used in this study for assessing yeast diversity relies on knowledge of the yeasts’ characteristics, and the relative abundance of each species (Lv et al., 2013). Further research at the subspecies level, using single-locus analysis, showed a high diversity of S. cerevisiae from the same production site, and within the same sample. These isolates showed a high allelic and genotypical richness.

Thus, this study determines these wild yeasts’ kinetic parameters for Saccharomyces and non-Saccharomyces species. There was a high kinetic variability depending on the strain or species of yeast used. Some of these yeasts had a high maximum specific growth rate close to an industrial strain used as a control in the study. These yeasts could constitute potential candidates for the improvement and the standardization of “Rabilé” production.

CONFLICT OF INTERESTS

The authors have not declared any conflict of interests.

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Table 4. Kinetic parameters of non-Saccharomyces yeasts.

| Species                     | ID   | \( \mu_{max} \) (\( h^{-1} \))^ \* | \( tg \) (h) | Level     |
|-----------------------------|------|-------------------------------------|------------|----------|
| Kluyveromyces marxianus     | KY01 | 0.568^a                           | 1.220      | High     |
| Clavispora lusitaniae      | OY30 | 0.527^a                           | 1.315      | Medium   |
| Pichia kudriavzevi         | TK23 | 0.400^b                           | 1.733      | Low      |
| Rhodotorula mucilaginosa   | BB21 | 0.346^c                           | 2.003      | Very low |
| Candida parapsilosis       | DR23 | 0.337^d                           | 2.057      | Very low |
| Meyerozyma caribbica       | ZN25 | 0.323^e,d                        | 2.146      | Very low |
| Kodamaea ohmeri            | ZN26 | 0.299^e                         | 2.318      | Very low |
| Yarrowia lipolytica        | OG11 | 0.296^e                          | 2.342      | Very low |
| Yarrowia sp                | OG12 | 0.284^f                          | 2.441      | Very low |
| Exophiala dermatitidis     | FD01 | 0.256^h                          | 2.708      | Very low |
| Cutaneotrichosporon curvatus | TK22 | 0.241^g                          | 2.876      | Very low |
| Yarrowia phanganaensis     | ZN06 | 0.116^i                          | 5.975      | Very low |

ANOVA Fisher (LSD) p-value <0.0001

*Values with the same letters are not significantly different.
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