Biochemical and functional properties of lactic acid bacteria isolated from Ivorian cocoa fermenting beans

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ABSTRACT

Objective: Fermentation is an important step in the post-harvest processing of cocoa beans. The quality of beans and chocolate depends strongly on the type and characteristics of microbial strains involved in the fermentation. This study investigates some functional properties of lactic acid bacteria (LAB), involved in Côte d’Ivoire cocoa fermentation.

Methodology and results: Bacterial growth was monitored during spontaneous heap fermentation using plate count and decimal dilution methods. LAB was present throughout the fermentative process with a load comprised between 6.0 log (UFC/g of bean) at the beginning to 6.60 log (UFC/g of bean) at the end. The maximum load of LAB corresponding to 7.22 log (UFC/g of bean) was reached after 24 h of fermentation in the condition of 42 °C of temperature and pH round 3.55. Among the 150 strains isolated, 99 were lactobacilli and 51 lactococci. Almost all these LAB showed ability to ferment glucose, fructose and sucrose with an important acidification capacity while only 8 strains belonging to lactococci proved to be able to assimilate citrate. Moreover, a weak proportion (7.33 %) of LAB strains isolated was heterofermentative type. These heterofermentative LAB strains were particularly characterized by a great fermentative capacity especially with glucose and sucrose with a gas production up to 5 mL.

Conclusion and application of results: The citrate metabolism and the strong fermentative capacity of certain LAB strains, indicate interesting technological traits potentially useful for their utilization as starters in cocoa fermentation.

Key words: lactic acid bacteria, fermentative power, cacao fermentation, acidification, citrate metabolism.

INTRODUCTION

The processing of cocoa beans consists of two major steps namely fermentation and drying (Hii et al., 2009; Guehi et al., 2010). Fermentation as the first stage in the preparation of chocolate begins immediately after the beans embedded in mucilaginous pulp are removed from the pods. Cocoa fermentation is a key step in the technological transformation of cocoa into chocolate, because the highly bitter, astringent unfermented cocoa beans lack the full chocolate flavour. During fermentation, cocoa pulp is hydrolyzed and this aids the drying process by allowing the pulp to be drained. Most importantly, fermentation triggers an array of chemical changes
within the cocoa bean that are vital to the development of the complex, beloved flavour of "chocolate" (Pereira et al., 2012). The fermentation of cocoa beans occurs at two levels. The first involves reactions that take place in the pulp, in the outer part of the beans, and the second involves several hydrolytic reactions that occur within the cotyledons (Lehrian and Patterson, 1983). The microbial activity in the cocoa pulp is a well-defined microbial succession in which numerous microbial species, mainly yeasts and bacteria including *Bacillus*, acetic acid bacteria (AAB) and lactic acid bacteria (LAB) are involved (Schwan et al., 2004; Ouattara et al., 2008). At the onset of cocoa beans fermentation, yeasts liquefy the pulp through depectinization, which reduces pulp viscosity, and produce ethanol from sugars (and citric acid) under anaerobic conditions in an acid (pH < 4.0), carbohydrate-rich environment. As pulp is drained away, ethanol formation proceeds and both temperature and pH increase, which creates ideal conditions for the growth of LAB and AAB. LAB converts sugars and organic acids mainly into lactic acid but others species such as heterofermentative LAB product in addition acetic acid, gas (CO\(_2\)) and ethanol. As more air comes in, AAB start to grow that, oxidize the ethanol, initially produced by the yeasts, to acetic acid. Ethanol and acetic acid diffuse into the beans, and this, in combination with the heat produced by this exothermic bioconversion, causes the death of the seed embryo as well as the end of fermentation. In addition, this initiates biochemical changes in the beans, leading to the formation of precursor molecules for the development of a characteristic flavour and colour of the beans (Hansen et al., 1998, Thompson et al., 2001). Therefore, the activities of LAB, AAB and yeast in this process are essential for the production of high-quality cocoa. On the other hand, fermentation is a natural microbial process, which remains difficult to control. In this context, many studies suggest the use of starter microbial culture as a best approach to improve fermentation process (Carr et al., 1980; Passos et al., 1984; Hansen et al., 1998; Hashim et al., 1998; Nielsen et al., 2005; Papalexandratou et al., 2011; Koen et al., 2012). Thus, several microbial cultures, including lactic acid bacteria have been assayed to analyze their potential as starters. These assays are mainly to analyze the effect of chosen strains, on the quality of fermented product. However, before using as potential starter, the functioning and role of this strain during cocoa bean fermentation, particularly their kinetics of growth, substrate consumption and metabolite production should be studied in more detail. Several studies concerned the identification of microflora from cocoa fermenting bean have been reported in other cocoa producing country such as Ghana, Brazil, Malaysia, Trinidad (Ardhana and Fleet, 2003; Nielsen 2006; Nielsen et al., 2007; Schwan, 1998; Jespersen et al., 2005). Although Côte d’Ivoire stands as the first cocoa producer in the world with up to 37 % of global production, the microbial flora responsible for its cocoa fermentation is still poorly characterized and unknown. In the present study, we investigate the characteristic and functional properties of lactic acid bacteria, one of the major microfloras involved in Côte d’Ivoire cocoa fermentation.

**MATERIAL AND METHODS**

**Fermentation condition and sampling:** Cocoa pods were harvested from Agboville (geographic coordinates 5°59’ north 4°28’West), situated at 79 km from Abidjan (Côte d’Ivoire). Beans were removed from pods and fermented traditionally by heap fermentation during six days. The fermenting mass about 100 kg, set on banana leaves and covered with banana leaves was constituted of mixed genotypes (Forastero, Trinitario and Criollo cultivars). Banana leaves are the main material used for traditional cocoa fermentation in Côte d’Ivoire. Samples of fermenting cocoa bean were taken according to a fixed time schedule, notably at the start of the fermentation (0 h) and after 24, 48, 72, 96 h, 120 and 144 h of fermentation. Each sample consisted of 50 g of beans that was transferred into sterile plastic bags. The pH and temperature were also regularly recorded directly at 15 cm depth on the fermenting heap, with portable pH-meter and thermometer.
Lactic acid bacteria isolation and numeration: The culture dependent approach was performed immediately after sampling. Therefore, 225 ml of 0.1 % (wt/v) peptone water (Oxoid, Basingstoke, United Kingdom) was added to 25 g of beans in a sterile Stomacher bag that was vigorously shaken for 5 min in a Stomacher 400 (Seward, Worthington, United Kingdom) to obtain a uniform homogenate. Samples (1.0 ml) of the homogenate were serially diluted 10-fold in Trypton salt buffer, from which aliquots (0.1 ml) were plated on different selective agar media; MRS (Man Rogossaa and Sharpe) agar, MSE (Mayeux, Sandine and Elliker, 1962) agar, and M17 agar (all from OXOID, Basingstoke, Hampshire, UK); to obtain the widest possible species variety of EAB associated with fermenting cocoa and capable of growing on these media. Plates were incubated at 37 °C for 48-72 h under anaerobic conditions to enable colony count enumeration (expressed as CFU per gram cocoa pulp-bean mass), as described previously (Kostinek et al., 2008). The three medium were supplemented with 50 µg/mL of nystatin to inhibit fungal growth. Eactic acid bacteria were identified as Gram positive, oxidase and catalase negative. The strains isolated were stored at -80 °C in MRS buffer medium supplemented with 20 % (v/v) glycerol in Eppendorf tubes, for further studies.

Analysis of carbon metabolism of strains isolated: The EAB isolates were analyzed for their ability to produce acid by catabolizing glucose, fructose, sucrose and citrate, which are the main carbohydrates, contained in the cocoa pulp (Afoakwa et al., 2013). The study of sugar metabolism was performed in a modified MRS medium containing the appropriate carbohydrate at 2 % as sole carbon source, 1.7 % agar and supplemented with 0.005 % of bromocresol purple. A volume of 7 mL of medium was put in a 20 mL tube and then sterilized for 15 min at 121 °C. Each strain was cultivated by central sting in the medium and then incubated at 30 °C for 72 hour in anaerobic conditions. A negative control was prepared in the same conditions and not inoculated with the microbial culture. The capacity of strains to metabolize the carbon source is assessed by the presence of colony in the tube and the change of medium colour due to pH lowering, comparatively to the negative control.

Evaluation of acidification capacity and fermentative type of LAB: Fermentative type and acidification capacity of bacterial strains was evaluated as previously described by Dicks and Van Vuuren (1987) with slight modification. Each strain was cultivated and incubated as described above, in the same MRS medium. The carbohydrates tested were glucose, fructose and sucrose that are known to be the sugar contained in the cocoa pulp (Afoakwa et al., 2013). Acid production was monitored by formation of yellow area in the tube and acidification capacity was analyzed by evaluating in a visual scale, the spread of the yellow area. The fermentative type was determined by ability of strains to produce gas from carbon source. Indeed, the presence of gas at the bottom of the tube accompanied by yellow zone indicates heterofermentative LAB type.

Evaluation of fermentative capacity of heterofermentative LAB: The quantity of gas produced by heterofermentative LAB is related to the intensity of their carbon metabolism according to the following equation (Raimbiault 1995).

\[
V = R^2 \times \pi \times h
\]

where \( V \) is the volume of gas produced in the tube after incubation, \( R \) is the ray of tube, \( h \) is the height occupied by the gas produced.

RESULT

Fermentation conditions and bacterial growth: The evolution of fermentation parameters (pH and temperature) of the cocoa beans during the process are shown in figure 1.
The pH value inside the cocoa fermenting mass was in the range 3.9-4.45 during the initial stage of the fermentation (0 to 48 h). After 48 h, the pH increased to a value of 6.07 at 96 h and reached a maximum of 7.9 at the end of the fermentative process (Figure 1). Concerning the cocoa bean temperature, the value was 29 °C at the start of the process and progressively rose to reach a peak of 45 °C within 48-72 h (Figure 1). Then the remaining time, a gradual decrease of temperature was observed, dropping at 36 °C at the end of fermentation. The dynamic of lactic acid bacteria population obtained from numeration during fermentation shows a biphasic mode of growth (Figure 2). The first phase concerned a rapid increase of LAB with a change in microbial load from 6.0 log (UFC/g of bean) at the beginning to 7.22 log (UFC/g of bean) at 24 h. This was followed by a decrease of LAB population in the timing 24-48 h. The second phase concerned a slow increase of bacterial growth that reaches a peak of at 96 h of fermentation. At the end, the process the LAB remained at relatively high load 6.60 log (UFC/g of bean).
Samples of fermenting bean were harvested at 24 h interval, and microbial numeration was performed by decimal dilution method on MRS, MSE and M17 media as described by Kostinek et al. (2008).

**LAB metabolism capacity:** A total of 150 LAB strains characterized as Gram positive, oxidase negative, catalase negative and cocci or rods shaped were selected from agar plates of MRS, M17 and MSE during the cocoa bean fermentation process. Among these strains, 99 were lactobacilli and 51 were lactococci. The LAB isolates were screened for their ability to produce acid by catabolizing glucose, fructose, sucrose and citrate. All the strains were naturally able to utilize glucose regarding the change of medium colour due to pH lowering comparatively to the control. It was also observed that, all the 99 lactobacilli strains were able to ferment fructose and sucrose (Table 1). However, among the 51 lactococci strains, 4 strains were not able to ferment fructose while 6 were unable to metabolise sucrose. In contrast, a wide proportion of LAB was not capable to ferment citrate, only 8 strains all belonging to lactococci showed the property to ferment citrate (Table 1).

**Table 1:** Distribution of lactic acid bacteria (LAB) strains isolates according to the sugar metabolism

<table>
<thead>
<tr>
<th>Strain</th>
<th>Tested strains</th>
<th>Glucose</th>
<th>Fructose</th>
<th>Sucrose</th>
<th>Citrate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lactobacilli</td>
<td>99</td>
<td>99</td>
<td>99</td>
<td>99</td>
<td>0</td>
</tr>
<tr>
<td>Lactococci</td>
<td>51</td>
<td>51</td>
<td>47</td>
<td>45</td>
<td>8</td>
</tr>
</tbody>
</table>

Strains were grown in modified MRS agar medium containing one of these carbohydrates and incubated for 72 hours at 30 °C.

**Acidification capacity:** When more acidity is produced, the medium is fully acidified and the yellow zone spread widely in the tube. In contrast, a weak acidification leads to a partial spread of yellow area in the medium. This allowed classifying the strain according to their acidification power (Figure 3). Basing on this principle, strains were classified as fully, moderately or weakly acidifying. All the 150 strains showed naturally, acidification capacity when glucose is used as sole carbon source. Of them, 141 strains proved to be fully acidifying while 8 strains were moderately acidifying and one showed low acidification (Table 2). Moreover, the acidification capacity of strain depended on the sugar metabolized. When fructose is used as sole carbon source, 146 strains presented full acidifying capacity while 4 strains showed an absence of acidification capacity. Additionally, 2 strains among these 146 showed full acidifying capacity with this sugar while they presented a low acid production.
capacity when glucose is use as sole carbon source. The use of sucrose as sole carbon source permit to determine 138 LAB as fully acidifying strains whereas 6 strains presented a low acid production capacity and 5 strains were not able to produce acid with this sugar (Table 2).

![Figure 3: Medium acidified at different levels.](image)

Fully acidified, moderately acidified and weakly acidified. Strains were cultivated in MRS medium by central sting touching the bottom of the tube and incubated for 72 h at 30°C.

<table>
<thead>
<tr>
<th>Table 2: Distribution of lactic acid bacteria (LAB) strains isolates according to the fermentative type</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Heterofermentatives LAB</strong></td>
</tr>
<tr>
<td>Glucose</td>
</tr>
<tr>
<td>Fructose</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Sucrose</td>
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<td></td>
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</tr>
</tbody>
</table>

**LAB group isolates and fermentative capacity:** The ability of LAB isolated, to produce gas from glucose metabolism was investigated in order to identify homofermentative and heterofermentative strains. Among the 150 isolates, only 11 (7.33 %) were heterofermentatives LAB which produced gas while 124 strains were homofermentatives (Figure 4). Among the heterofermentative LAB, 4 were lactococci and 7 lactobacilli. Additionally, it was observed that some strains producing gas with glucose failed to produce gas with fructose or sucrose. Hence, for a given strain, the same profile regarding gas production was not necessarily observed when the sugar used was different. We further estimated the fermentative power of strains by measuring the volume of gas produced from glucose metabolism. Figure 4 shows that heterofermentative strains produced different volume of gas although they were cultivated in the same conditions. Subsequently, these strains present different fermentative power according to the intensity of metabolism. Similarly, to acidification property, the table 3 shows that gas production also varied with the sugar used for a given strain. For instance, the most important gas producer strain BL 25 produced 2-folds more gas with glucose than with fructose. It appears that glucose offers the most efficient fermentative power with higher gas production in general (Table 3). Furthermore, the best gas producers BL25, BL40 and...
BL08 were also fully acidifying, with the ability to ferment all the sugars studied except citrate and belonged to lactococci. These strains were classified as having a strong fermentative capacity. It was also observed that 2 strains (BL 87 and BL 122) demonstrated the most efficient fermentative power with fructose than glucose and sucrose. We further noted that the 8 strains, which were able to catabolizing citrate, showed a fully acidifying capacity with glucose, fructose and sucrose but without gas production capacity. Moreover, these strains presented a strong fermentative capacity with gas volume ranged 3-4 mL such as strains BL 08 and BL 40.

**Figure 4:** Medium with different volume of gas.

Strong gas production, middle gas production and low gas production. Strains were cultivated in MRS medium by central sting touching the bottom of the tube and incubated for 48 h at 30 °C.

**TABLE 3:** Quantity of gas produced by heterofermentative lactic acid bacteria (LAB) strains

<table>
<thead>
<tr>
<th>tested strains</th>
<th>Gas volume (mL) produced with glucose</th>
<th>Gas volume (mL) produced with fructose</th>
<th>Gas volume (mL) produced with sucrose</th>
</tr>
</thead>
<tbody>
<tr>
<td>BL 07</td>
<td>&lt; 0.5</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>BL 08</td>
<td>3.534</td>
<td>&lt; 0.5</td>
<td>2.8272</td>
</tr>
<tr>
<td>BL 25</td>
<td>5.301</td>
<td>2.6505</td>
<td>3.534</td>
</tr>
<tr>
<td>BL 40</td>
<td>3.004</td>
<td>2.1204</td>
<td>2.6505</td>
</tr>
<tr>
<td>BL 55</td>
<td>2.1204</td>
<td>&lt; 0.5</td>
<td>2.4738</td>
</tr>
<tr>
<td>BL 56</td>
<td>2.297</td>
<td>1.414</td>
<td>1.0602</td>
</tr>
<tr>
<td>BL 61</td>
<td>2.474</td>
<td>1.5903</td>
<td>0</td>
</tr>
<tr>
<td>BL 87</td>
<td>&lt; 0.5</td>
<td>1.5903</td>
<td>&lt; 0.5</td>
</tr>
<tr>
<td>BL 121</td>
<td>&lt; 0.5</td>
<td>0</td>
<td>&lt; 0.5</td>
</tr>
<tr>
<td>BL 122</td>
<td>&lt; 0.5</td>
<td>1.8835</td>
<td>1.2369</td>
</tr>
<tr>
<td>BL 139</td>
<td>2.1204</td>
<td>&lt; 0.5</td>
<td>4.7709</td>
</tr>
</tbody>
</table>
DISCUSSION
The quality of cocoa beans and chocolate depends strongly on the type and characteristics of microbial strains involved in the fermentation. On this basis, the use of starter microbial strains presenting a high fermentative ability is believed to be favourable for cocoa fermentation improvement and producing high quality. In this study, LAB involved in fermentation was analyzed in order to know the properties and the characteristics of LAB involved in Côte d'Ivoire cocoa fermentation. Analysis of heap spontaneous fermentation condition indicates that the pH of fermentation heap ranged from 3.9 at the beginning to 7.9 at the end of the fermentation process, while the temperature ranged from 29 to 36 °C with a peak at 45 °C within 48 -72 h. The same profile of temperature and pH variation has been regularly recorded in cocoa fermenting mass in many countries (Schwan and Wheals, 2004, Papalexandratou et al., 2011; Guehi et al., 2010) indicating that the increase of both parameters constitutes an inherent property of cocoa fermentation worldwide. Moreover, pH continuously increasing during the fermentation became alkaline at the end of the process. Guehi et al. (2010) also reported an alkaline pH (8.5) at the end of spontaneous cocoa fermentation in Côte d'Ivoire. This result is particularly surprising since, it is very often observed that pH of cocoa fermentation is maximum toward the end, but remains in acidic range (Schwan and Wheals, 2004, Schwan 1998; Guehi et al., 2010). We could not explain the reason for why, sometimes the pH became alkaline in cocoa fermentation, but this seems to be a particularity of Côte d'Ivoire cocoa fermentation, since alkaline pH has not been yet reported in other country. On the other hand, the variation of temperature and pH of cocoa fermenting mass influence the microbial growth (Schwan and Wheals, 2004). The dynamic of lactic acid bacteria population obtained during fermentation shows that LAB are present throughout the fermentative process although a variation in bacterial load was observed, notably the rapid increase of LAB during the first stage and their slow decrease toward the end of fermentation. The same pattern of LAB growth was observed during small scale fermentation in Mexico (Pereira et al., 2012). In contrast, an initial increase of LAB load was followed by a drastic decrease from 80 h until the microorganisms were not detectable during spontaneous wooden box cocoa fermentation in Brazil (Schwan 1998). Hence, fermentation may sometime imply stress conditions susceptible to limit the microbial growth and subsequently to counterbalance their favourable technological properties. Regarding their growth kinetic during fermentation, the results suggest that LAB studied should be competitive and capable to survive under certain stress conditions and eventually to express their fermentative potential during the fermentation process. Biochemical and morphological identification of strains isolated, revealed as expected the presence of Lactococci (34 %) and Lactobacilli (66 %) in Ivorian spontaneous cocoa fermentation. Lactobacilli were also found to be dominant in Ghanaian and Nigerian cocoa fermentation (Camu et al., 2007; Kostinek et al., 2008). At date, the possible impact of the balance between both types of LAB, on the quality of bean is not elucidated. However, the results also showed that homofermentative strains are widely dominant among the LAB isolated as it is reported in Ghana (Camu et al., 2007) and Nigeria (Kostinek et al., 2008). Homofermentative LAB strains are known to convert sugars almost exclusively into lactic acid and then produce more lactic acid than heterofermentative strains (Raimbiault 1995). Although acid production during cocoa fermentation is desirable for development of precursors chocolate aroma (Biehl et al., 1993; Jinap 1994), a high production of less volatile lactic acid is liable to confer to fermented and dried beans a final acidity that deteriorate their quality (Schwan and Wheals, 2004). To this point of view heterofermentative strains producing other acid more volatile such as acetic acid, may be more interesting and desirable, but this type of strain remains in minor proportions in spontaneous cocoa fermentation as also showed by our results. Heterofermentative metabolism is characterized by gas (carbon dioxide) and acid other than lactic acid production (Raimbiault 1995). This study showed different level of gas production from sugar metabolism suggesting different fermentative power. Strains producing more gas were also observed to be fully acidifying suggesting that both properties are linked. The acidification of bean, allow activation of hydrolytic enzymes deep into the cotyledons (Jinap 1994); these enzymes notably aspartic endopeptidase and serine carboxypeptidase catalyse the production of precursor of chocolate flavour from globular proteins (Biehl et al., 1993; Voigt et al., 1994; Serra and Ventura, 1997) while gas production reflect the fermentative power, and then the ability of strain to speed the fermentation. One the other hand, some
homofermentative LAB isolates all belonging to lactococci presented the capacity to catabolizing citric acid. These results are consistent with the observation according to which lactococci are more susceptible to catabolize citrate than lactobacilli (Drinan et al., 1976). It was also reported that in experimental conditions, homofermentative lactococci produce more acetoin and diacetyl amount from citrate degradation than heterofermentative Lactobacilli (Drinan et al., 1976). Furthermore, citrate metabolism constitutes an important and particular property, since LAB is not usually able to utilize citric acid as carbon source. To our knowledge, no report exists on bacterial strain from cocoa fermentation metabolizing citrate as carbon source. Citric acid is the compound responsible for the initial pH of cocoa pulp before fermentation processing (Petitpier 1986; Thompson et al., 2001; Ardhana and Fleet, 2003, Lefeber et al., 2011). The degradation of this acid in the first stage of fermentation allows the raise of the pH favourable for the development of many bacterial groups (Schwan and wheals, 2004). Accordingly, possessing citrate metabolism constitutes a valuable trait for cocoa fermentation, since this could allow enhancing microbial metabolism, strengthening fermentation and subsequently have a possible contribution to cocoa and chocolate quality.

In conclusion, LAB flora studied were composed of lactobacilli and lactococci, presenting either homofermentative or heterofermentative metabolism. These bacteria were able to grow throughout the fermentation with capacity to metabolize glucose, fructose and sucrose. Furthermore, some homofermentative lactococci strains presented the capacity to metabolize citrate as carbon source, a valuable property for cocoa fermentation. All together, these results show that strains behave differently and indicate that some of them present interesting technological traits potentially useful for their utilization as starters in cocoa fermentation.

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