

Original Research Article

Screening and Characterization of Putative Probiotic *Lactobacillus* Strains from Honey Bee Gut (*Apis mellifera*)

ABSTRACT

The objective of this work was to isolate, identify and characterize lactobacilli strains from the intestinal tract of honey bees as putative probiotics. We obtained eighty-five isolates. At the end of screening based on physiological properties, 17 isolates were pre-selected and their resistance to gastrointestinal stress was evaluated. Twelve (12) with good resistance after 3 h exposure to low pH values (pH2, pH3) were subjected to determination of their *in vitro* BSH activity. The research of the *bsh-A*, *bsh-B*, *Bsh1* and *Bsh-Lp1* genes encoding the BSH enzyme was also conducted. Four isolates (H46, H82, H21 and H28) were resistant, seven others tolerant (H6, H15, H47, H24, H67, H44, H80) and only one was sensitive (H63) to oxgall bile salt. Determination of BSH activity revealed that all strains hydrolyze bile salts, with a preference for oxgall as opposed to Taurodeoxycholate. H15 and H47 isolates showed the highest BSH activities, which were 103.82 ± 12.93 U/mg and 98.53 ± 2.86 U/mg, respectively, with no significant difference ($p > 0.05$). Only the *bsh-Lp1* gene was amplified in isolate H24 and H28. None of the strains showed the *bsh-1*, *bsh-A* or *bsh-B* genes. After sequencing *bsh-Lp1* gene of H24 and H28, the BSH proteins deduced from the complete ORF showed high similarity with those of GenBank database. Antimicrobial activity revealed the inhibition zone against pathogenic and food spoilage bacteria. Isolates were identified based on the sequencing of 16S rRNA encoding gene as *Lactobacillus plantarum* (75%) and *Lactobacillus paraplantarum* (25%).

Keywords: Honey bees; Lactobacilli; Bile salt hydrolase; Antimicrobial Activity; 16S rRNA; Gene sequencing.

1. INTRODUCTION

Over the last few decades, probiotic lactic acid bacteria (LAB) have become increasingly popular in fighting diseases that impair human health [1]. Factors contributing to this enthusiasm include the emergence of scientific and clinical evidence demonstrating the efficacy of certain probiotic strains and the increasing consumer demand for natural drug substitutes. Probiotics are "live microorganisms which, when administered in sufficient quantities, confer benefits to the host's health" [2].

Various studies indicated that probiotics might improve lactose intolerance, have a positive influence on the intestinal flora of the host, stimulate/modulate mucosal immunity, reduce inflammatory or allergic reactions, reduce diarrhea, constipation, candidiasis, blood cholesterol and competitively exclude pathogens [3].

Cholesterol is a vital substance in the human body. Long-standing high blood cholesterol levels may lead to atherosclerosis and therefore, may cause a major risk of developing cardiovascular diseases (CVDs). By the year 2030, CVDs will affect approximately 23.3 million people around the world [4]. Although a drug is used, it is often suboptimal, expensive and can cause adverse side effects [5]. These

55 pharmaceuticals are mostly based on the interruption of the enterohepatic circulation (EHC) of bile salts
56 [6]. In recent years, LAB identified as probiotics are increasingly popular in challenging these diseases.

57
58 Bile salt metabolism and cholesterol metabolism are closely linked. Bile is a digestive secretion that plays
59 a key role in lipid solubilization, as it behaves like biological detergents [6]. The deconjugation, one of the
60 mechanisms that the lactobacilli use to counteract the action of bile salts, is catalyzed by bile salt
61 hydrolases (BSH) which release glycines/taurines from the steroid nucleus, thus reducing the solubility of
62 bile at low pH and reduces its detergent activity [7]. The hydrolysis of bile salts by LAB breaks the
63 enterohepatic cycle of bile salts and may contribute to lower the blood cholesterol level (cholesterol-
64 lowering effect). Oral consumption of probiotic LAB has been shown to considerably decrease cholesterol
65 levels by as much as 22 to 33%. This cholesterol-lowering property can be in part attributed to BSH
66 activity [8,9].

67
68 But, once ingested, probiotic LAB come into contact with the stressful conditions of the gastrointestinal
69 tract. They must survive and remain metabolically active under these circumstances. Gastric acidity and
70 the disaggregation properties of bacterial membranes by bile salts are the primary challenges. It is also
71 important for these LABs to exhibit resistance against the autochthonous microflora to improve the ability
72 to colonize the digestive tract and express their probiotic functions [10]. This resistance to the microflora
73 can occur particularly by the production and secretion of antimicrobial compounds. Some strains of lactic
74 bacteria can synthesize bactericidal/bacteriostatic molecules such as organic acids, hydrogen peroxide,
75 carbon dioxide, diacetyl and especially bacteriocins [11]. The selection of LAB strains that are able to
76 withstand the stressful conditions into the gastrointestinal tract of humans and improve their probiotics
77 benefits is a challenge.

78
79 Several microorganisms live in symbiosis with insects that have special diets. The bees are an example.
80 Their stomach is filled with nutrients and nectar, and hence constitutes a micro-aerobic environment
81 which, at an optimal temperature of 35°C in the hive serves as a useful ecological niche for LAB [12]. In
82 fact, LAB and predominantly Lactobacilli has been found in the gastrointestinal tract of bees [13].
83 Previous studies have reported the isolation from honey bee of LAB with probiotic potential and their
84 applicability in controlling infections in bees. However, none to the best of our knowledge have addressed
85 the selection of potential probiotic LAB from honey bees for their use in Human. Moreover, the microbiota
86 of the honey bees in Cameroon has not yet been explored, whereas it may possess LAB that can provide
87 beneficial effects in humans. The selection of probiotic LAB has been based on *in vitro* physiological tests
88 to different stress factors such as low pH, and bile salts [14]. In addition to these physiological tests, the
89 use of molecular markers is an approach that would lead to improved screening in order to obtain the
90 strains presenting the most wanted potentials.

91
92 In the present study, LAB isolated from honey bee digestive tract were screened for properties such as
93 pH and bile salts tolerances, bile salts hydrolysis as well as antimicrobial activity using phenotypic criteria
94 as well as molecular markers.

95 96 **2. MATERIAL AND METHODS**

97 98 **2.1 Isolation and purification of lactic acid bacteria**

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100 100 to 125 honey bees (*Apis mellifera*) were collected from hives and honey vendors in five localities of
101 the Menoua Division (West-Cameroon): Fossong Wentcheng (5°24'N; 9°56'E), Penka-Michel centre
102 (5°27'N; 10°18'E), Dschang ("Marché B") (5°27'N; 10°02'E), Bamendou "Qt Nguim" (5°26'N; 10°12'E),
103 Balessing ("King Place") (5°30'N; 10°15'E). The samples were collected by trapping in sterilized bottles.
104 Once in the laboratory, the bottles were stored at + 4°C for 3-5 min to stop or decrease the mobility of the
105 bees. Using the method described by Mahesh et al. [15], the stomach contents of the bees were collected
106 and introduced into 5 ml of MRS broth supplemented with 5% (w/v) L-cysteine-HCl for 48 h activation at
107 37°C. Each culture was subsequently streaked onto MRS Agar medium supplemented with 5% (w/v) L-
108 cysteine-HCl and incubated at 37°C for 48 h. At the end of the incubation period, colonies of different
109 appearance were isolated and cultured MRS broth medium. The purity of isolates was assessed by re-
110 streaking on a fresh MRS agar medium. Gram staining was carried out, and Gram-positive rod-shaped

111 bacteria were selected and evaluated for their physiological parameters such as catalase activity, CO₂
112 production from glucose, growth at 10°C and 45°C.

113

114 **2.2 Phenotypic and genotypic tests related to acid and bile salts tolerance**

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116 **2.2.1 Evaluation of the ability to tolerate acidity**

117

118 The ability of the LAB isolates to tolerate acidity was assessed using the method of Verdenelli et al. [16].
119 Resting cell suspensions were prepared by harvesting (10,000 g, 10 min at 4 °C) exponentially grown
120 (16-18 h) lactobacilli cultures. Resting cell suspension (10⁸ CFU/ml) was introduced in different citrate
121 buffers (pH 2, pH 3, and pH 6.5) for 3 h. The suspensions were then centrifuged at 5,000 g for 5 min at 4
122 °C twice and washed in sterile saline solution to eliminate the citrate buffer. Cell pellets were suspended
123 in physiological solution, and a series of tenfold dilution (10⁻² to 10⁻¹⁰) were prepared. 50 µl of each
124 dilution was plated on to MRS-Cys-HCl agar and incubated at 37°C for 24-48 h. Percentage of viable
125 bacteria was expressed as the ratio between the counts after 3 h and at 0 h incubation time.

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127 **2.2.2 Evaluation of the ability to survive in the presence of bile salts**

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129 The capacity to grow in the presence of bile (bile salt tolerance) was also evaluated following the method
130 of Verdenelli et al. [16]. with slight modifications. MRS broth containing 0, 0.3, 0.5 or 1% w/v oxgall (a
131 mixture of conjugated and unconjugated bile salts, a natural dried bovine bile component; DIFCO) were
132 used. The absorbance at 560 nm (A_{560nm}) was measured at hour intervals up to 8 h. The results were
133 expressed as the time difference of growth in the control (MRS without oxgall) and the test media (MRS
134 containing 0.3, 0.5 or 1% oxgall) measured by a 0.3 unit increase in A_{560nm} as described by Gilliland et al.
135 [17].

136 The difference between the time required to increase the A_{560nm} of 0.3 units for a given bile salt
137 concentration and that needed for the control is the stunted growth retardation. This time was calculated
138 at 0.3% of bile salts and the isolates classified according to their sensitivity to bile salts based on their
139 growth retardation (d) according to the criteria described by Château et al. [18]. All experiments were
140 carried out in triplicate.

141

142 **2.2.3 Screening for the presence of genes involved in resistance to acidity and bile salts.**

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144 The genes involved in pH and bile salt tolerances that were screened are shown in Table 1. The
145 genomes of the LAB isolates were screened by direct colony PCR. The primers used for each PCR
146 reaction were designed based on the literature (references in Table 1).The following conditions were used
147 for PCR: initial denaturation at 95°C for 5 min, then 40 cycles of the denaturation set at 95°C for 1 min,
148 hybridization (at the annealing temperature of each gene) for 1 min, polymerization at 72°C for 1 min and
149 a final step of additional elongation at 72°C for 10 min. Then, 10 µl of PCR product was analyzed on 1%
150 agarose gel with GoldView™ for DNA staining in Tris-acetate-EDTA buffer 0.5X (TAE, pH 8.5) for 20 min
151 at 130 V and the reading done by UV trans-illumination.

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166 **Table 1. List of primers used to screen genes involved in acid and bile salts tolerances in the**
 167 **collection of bacteria**
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General function	Gene	Predicted Function	Primer (5' to 3' sequence)	An.T (°C)	Size (bp)	References
Survival to acidity	<i>Hdc</i>	Histidine decarboxylase	Fd-AGATGGTATTGTTTCTTATG	52.0	367	[19]
			R-AGACCATACACCATAACCTT			
pH and bile salt survival	<i>gtf</i>	Glucan synthase	Fd-ACACGCAGGGCGTTATTTTG	58.0	374	[20]
			R-GCCACCTTCAACGCTTCGTA			
survival	<i>clpL</i>	ATPase	Fd-GCTGCCTTYAAAACATCATCTGG	56.0	158	[21]
			R-AATACAATTTTGAARAACGCAGCTT			

169 *An. T = Annealing temperature, Size = Expected amplicon size, bp = base pair*

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 171 **2.3 Phenotypic and genotypic tests related to Bile Salt Hydrolase activity**
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173 **2.3.1 In vitro evaluation of Bile Salt Hydrolase activity**
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175 The BSH activity was measured by determining the concentration of amino acids released from
 176 conjugated bile salts (oxgall and taurodeoxycholate, DIFCO) as described by [9]. One unit of BSH activity
 177 (U/mg) was defined as the amount of enzyme that releases 1 µmol of amino acids from the substrate per
 178 minute.
 179

180 **2.3.2 Screening for the presence of genes involved in the Bile Salt Hydrolase activity**
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182 The primers used are shown in Table 2 and the PCR reaction was carried out in a reaction mixture
 183 consisting of 25 µl of 2 x Master Mix, 2 µl of primer (1µM) and a bacterial colony of the pure isolate in a
 184 final volume of 50 µl. A heating step was performed at 94 °C for 2 min, and the PCR program consisted of
 185 30 cycles composed of 3 steps as follows: denaturation at 94 °C for 1 min, hybridization at 58°C for 20 s,
 186 elongation at 72°C for 2 min. After these 30 cycles, a final extension step at 72°C for 10 min was
 187 performed. Then, the amplicons were analyzed as previously described.
 188

189 **Table 2 List of primers used to amplify the genes responsible for the expression of the BSH**
 190 **enzyme**
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Genes	Primer (5' to 3' sequence)	An.T	Reference
<i>bsh-A</i> (<i>Lb acidophilus</i> BshA)	F: TACAACACTATTCATTTAGACGCAATATCC R: CACTCTGCCAACACTCCATAACG	58°C	[22]
<i>bsh-B</i> (<i>Lb acidophilus</i> BshB)	F: CAAAAGCCATTTATTCCGACTGA R: CATAATTTATTACTTCTTTGTTAGACAGC		
<i>bsh-Lp1</i> (<i>Lb plantarum</i> Bsh1)	F: TGTATTTTAGTAGGTATTTCAAGCATCTC R: CAATGAAATGGTTACGATTACGC		
<i>bsh-1</i> (<i>Lb casei</i> Bsh)	F: GCCATTAAGCAATTCGGGTTATA R: CCAATGATTGGTCTCTCGTTCA		

192 *An. T = Annealing temperature*

193
 194 **2.3.3 Sequence Analysis of bsh**
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196 The purified amplicons of the *bsh* gene of the isolates were sequenced by an automated DNA sequencer
 197 using the services of a commercial company (<http://www.ruibiotech.com>). The sequences were aligned
 198 with similar sequences present in the National Center for Biotechnology Information (NCBI) gene
 199 collection (<http://blast.ncbi.nlm.nih.gov/Blast.cgi>). The BLAST2 program from the NCBI was used for
 200 nucleotide sequence analysis and amino acid sequence deduction. Protein sequences were aligned
 201 using ClustalW software package. The nucleotide sequences were deposited in the GenBank database to
 202 obtain Accession Number.
 203

204 **2.4 Antimicrobial activity**

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206 The direct antimicrobial activity of the LAB strains was evaluated as described by [23]. Indicator bacteria
207 were selected based on their involvement in gastrointestinal infections and food spoilage: *Listeria innocua*
208 ATCC 33090, *Staphylococcus aureus* ATCC 25923, *Streptococcus mutans* DSM 20523, *Bacillus cereus*
209 11778, *Proteus mirabilis* (Clinical isolate), *Escherichia coli* ATCC 13706, *Salmonella enterica* serovar
210 Typhi ATCC 6539, *Pseudomonas aeruginosa* ATCC 20027. *Lactobacillus plantarum* 5S is a bacteriocin's
211 sensitive strain obtained from our laboratory collection and used as positive control.

212 **2.5 Molecular Identification of selected LAB isolates by 16S rRNA gene sequencing**

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214 The primers (Forward: 5'-AGAGTTTGATCCTGGCTCAG-3' and Reverse: 5'-
215 CTACGGCTACCTTGTTACGA-3') previously designed by Weisburg et al. [24] were used to amplify the
216 nearly completed 16S rRNA encoding gene. Direct colony PCR reaction was carried out in a reaction
217 mixture consisting of 25 µl of 2X Master Mix, 4 µl of primers (1µM) and a bacterial colony of the pure
218 isolate in a final volume of 50 µl. A step of heating was carried out at 94°C for 2 min. The PCR program of
219 30 cycles consisting of 3 steps was done: denaturation at 94°C for 1 min, hybridization at 42°C for 20 s,
220 elongation at 72°C for 2 min. After these 30 cycles, a final extension step at 72°C for 10 min was
221 performed. The amplicons were analyzed as described earlier. After amplification of the 16S rRNA
222 encoding gene, DNA fragments of about 1400bp were observed in the agarose gel. The amplicons were
223 then purified and sent to a commercial facility for sequencing (<http://www.ruibiotech.com>, China, Beijing).
224 After sequencing, the chimeras within the sequences were identified and trimmed using ChromasPro
225 1.7.7 software. The sequences were aligned with similar sequences retrieved from the NCBI GenBank
226 (<http://blast.ncbi.nlm.nih.gov/Blast.cgi>). CLUSTAL multiple sequence alignment was performed and the
227 16S rRNA gene sequences of the strains were deposited in the NCBI GenBank to obtain their nucleotide
228 sequences accession numbers.

229 **2.6 Statistical analysis**

230
231 The results were expressed as the mean \pm standard deviation and then analyzed by the analysis of
232 variance (ANOVA) using the Graph Pad InStat software (GraphPad Software Inc., www.graphpad.com,
233 V3). When differences existed, means were compared between them by the Student-Newmann-Keuls
234 test at the probability threshold 0.05.

235
236 Principal Component Analysis (PCA) was applied to acid survival (pH2 and pH3), 1% bile salt survival
237 rate, time to increase absorbance (ΔDO_{600nm}) to 0.3 units at 1% bile salt, as well as the BSH activity on
238 the Oxgall. The XLSTAT 2007.8.04 software (Addinsoft, Paris, France, <http://www.slstat.com>) was used
239 and a normalized Pearson (n) PCA was applied.

240 **3. RESULTS**

241 **3.1 Isolation of LAB**

242
243 A total of eighty-five pure and Gram-positive isolates were obtained. Microscopic observation revealed
244 that they were rod-shaped and arranged in pairs or chains of varying length. Among these isolates, 17 do
245 not produce CO₂ from glucose and were preselected for future testing.

246 **3.2 Tolerance to acidity by the pure isolates**

247
248 The 17 preselected isolates were tested for resistance to low pH and the results are presented in Table 3.
249 In general, there was a significant decrease ($p < 0.05$) in the survival percentage of the isolates when the
250 pH decreases from 6.5 to 3 and then to 2. At pH2, 8 isolates out of 17 (47.06%), namely H21, H32, H45,
251 H48, H51, H51, H55, and H63 showed survival percentages below 50%, while the significantly ($p < 0.05$)
252 highest survival percentage was $93.00 \pm 1.73\%$ with isolate H15. At pH3, only the isolate H55 has a
253 survival percentage below 50%, while the other isolates had survival percentages greater than or equal to
254

259 65%. The significantly ($p < 0.05$) highest values were those of the isolates H15 and H47, respectively
 260 $95.67 \pm 2.08\%$ and $96.17 \pm 1.26\%$. The 12 isolates with survival percentages greater than 45% at pH 2
 261 have been selected for the further assays.

262
 263 **Table 3. Percentage of survival of the isolates after incubation for 5 h in citrate buffer at different**
 264 **pH values**

Isolates	Survival (%)		
	pH2	pH3	pH6.5
H6	61.67±2.89 ^{Ab}	81.67±1.53 ^{Bbejk}	96.00±2.00 ^{Cbdeghi}
H15	93.00±1.73 ^{Ac}	95.67±2.08 ^{Bci}	98.00±2.00 ^{Cbeghi}
H21	45.67±2.08 ^{Ad}	73.00±2.65 ^{Bdg}	98.67±1.15 ^{Cbegh}
H24	62.67±2.52 ^{Ab}	83.33±2.89 ^{Bekj}	95.00±0.00 ^{Ca}
H28	75.67±2.08 ^{Ae}	91.00±3.61 ^{Bcthi}	98.67±1.15 ^{Cbet}
H32	30.00±5.00 ^{Af}	70.67±3.06 ^{Bdg}	90.33±2.52 ^{Ca}
H44	50.00±5.00 ^{Ad}	72.33±2.52 ^{Bdg}	91.33±1.15 ^{Cad}
H45	13.67±3.21 ^{Aa}	77.33±6.43 ^{Bbg}	94.67±4.16 ^{Cae}
H46	72.33±2.52 ^{Ag}	92.33±2.52 ^{Bhic}	97.67±2.08 ^{Cbeghi}
H47	88.67±1.15 ^{Ah}	96.17±1.26 ^{Bi}	98.33±0.58 ^{Cbeghi}
H48	8.33±2.89 ^{Aa}	65.00±5.00 ^{Bd}	90.67±1.15 ^{Cac}
H51	8.33±2.89 ^{Aa}	87.67±2.52 ^{Bthjk}	97.17±1.04 ^{Cbeghi}
H55	11.67±2.89 ^{Aa}	13.33±2.89 ^{Ba}	95.50±1.80 ^{Calh}
H63	48.33±7.64 ^{Ad}	70.00±5.00 ^{Bdg}	97.00±1.73 ^{Cbeghi}
H67	58.33±2.89 ^{Ab}	87.67±2.52 ^{Bik}	95.67±2.08 ^{Cbcdeghi}
H80	63.33±2.89 ^{Ab}	72.67±2.52 ^{Bd}	95.00±3.00 ^{Calg}
H82	61.67±2.89 ^{Ab}	71.67±2.89 ^{Bdg}	95.50±1.80 ^{Calh}

266 ^{A,B,C}: On the same row, values with identical letters do not differ significantly ($p > 0.05$) compared to the MRS-Cys
 267 control.
 268 ^{a, b, c, d, e, f, g, h, i, j, k}: On the same column, values with identical letters do not differ significantly ($p > 0.05$). Values represent
 269 the mean±SD of three trials ($n=3$).
 270

271 **3.3 Tolerance to bile salts**

272
 273 The survival percentages of isolates at different oxgall concentrations (0.3, 0.5 and 1%), vary between
 274 79.64 ± 0.78 and 93.71 ± 0.92 after exposure to 0.3% oxgall (Table 4). At 0.5% oxgall, only isolate H44
 275 had a survival rate lesser than 72%, i.e., 59.23 ± 3.99 . The isolate H46 showed a survival rate greater
 276 than 91% for any concentration. This isolate has a higher survival ($p < 0.05$) at 1% oxgall compared to all
 277 other isolates. However, only the isolate H44 has a survival of less than 50% with 1% of bile salts.
 278

279 At all oxgall concentrations, the time (min) required to increase the absorbance by 0.3 units for each
 280 isolate (Table 4) didn't increase significantly for isolates H46, H82, H6, H15, H47, H21, and H24
 281 compared to the control (MRS-Cys). For the isolate H67, this time was significantly different from the
 282 control ($p < 0.05$) at 0.5% oxgall. It differed significantly ($p < 0.05$) from the control at 1% oxgall with the
 283 isolates H44, H80 and H63.
 284

285 The difference between the time required to increase the OD of 0.3 units for a given bile salt
 286 concentration and that required for the control represents the accrued growth delay (in minutes). This
 287 value was calculated (Table 4). This growth delay varies between 10 ± 0.0 min and 50 ± 17.32 min for the
 288 concentration of 0.3% of bile salts.
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296 **Table 4. Parameters indicating the behavior of the isolates in Bile Salt**
 297

Isolates	Parameters							
	Survival rate after 24 h			Time (min) required to increase the absorbance by 0.3 unit				GD *
	0.3% BS	0.5% BS	1% BS	MRS-Cys	0.3% BS	0.5% BS	1% BS	
H6	92.48±0.98 ^{Abcd}	84.28±1.30 ^{Bbcehi}	82.06±1.80 ^{Bchi}	140±17.32 ^{Aac}	170±17.32 ^{Aac}	170±17.32 ^{Aac}	180±00.00 ^{Aa}	II
H15	92.20±0.41 ^{Abcde}	75.48±0.87 ^{Bdfkl}	71.19±2.39 ^{Cbde}	130±17.32 ^{Aac}	160±17.32 ^{Aac}	160±17.32 ^{Aac}	160±17.32 ^{Aa}	II
H21	89.22±1.25 ^{Abcdefgh}	75.52±1.06 ^{Bfkl}	67.08±1.28 ^{Cbdefgk}	160±17.32 ^{Aac}	170±17.32 ^{Aac}	170±17.32 ^{Aac}	170±17.32 ^{Aa}	I
H24	81.47±1.06 ^{Aa}	72.19±0.30 ^{Bdfgk}	69.60±0.45 ^{Cbdeg}	120±00.00 ^{Aa}	140±17.32 ^{Aac}	140±17.32 ^{Aa}	140±17.32 ^{Aa}	II
H28	93.57±1.95 ^{Abc}	87.65±1.82 ^{Bh}	85.69±1.47 ^{Bh}	140±17.32 ^{Aac}	170±17.32 ^{Aac}	170±17.32 ^{Aac}	180±00.00 ^{Ba}	I
H44	83.10±4.38 ^{Aa}	59.23±3.99 ^{Bj}	48.67±1.76 ^{Cj}	170±17.32 ^{Abc}	200±34.64 ^{Abc}	200±34.64 ^{Aac}	260±34.64 ^{Bb}	II
H46	93.71±0.92 ^{Ab}	92.10±1.08 ^{Aa}	91.84±1.15 ^{Aa}	130±17.00 ^{Aac}	140±17.00 ^{Aac}	140±17.00 ^{Aa}	140±17.00 ^{Aa}	I
H47	81.52±0.08 ^{Aah}	84.82±0.49 ^{Bbeh}	74.25±1.09 ^{Cbcei}	130±17.32 ^{Aac}	160±17.32 ^{Aac}	160±17.32 ^{Aac}	160±17.32 ^{Aa}	II
H63	79.64±0.78 ^{Aa}	77.06±0.54 ^{Al}	62.59±13.56 ^{Adfkg}	170±17.32 ^{Abc}	220±34.64 ^{Abc}	220±34.64 ^{Abc}	260±34.64 ^{Bb}	III
H67	90.12±0.87 ^{Abcdefg}	84.96±1.35 ^{Bbhi}	82.96±0.72 ^{Bhi}	120±00.00 ^{Aa}	130±17.32 ^{Aa}	160±17.32 ^{Bac}	170±17.32 ^{Ba}	II
H80	84.28±2.14 ^{Aa}	76.15±1.49 ^{Bkl}	69.46±1.70 ^{Cbdegk}	170±17.32 ^{Abc}	200±34.64 ^{Abc}	200±34.64 ^{Aac}	260±34.64 ^{Bb}	II
H82	90.52±3.89 ^{Abcdef}	85.66±4.30 ^{Abh}	75.53±3.53 ^{Bbci}	130±17.32 ^{Aac}	140±17.32 ^{Aac}	140±17.32 ^{Aa}	140±17.32 ^{Aa}	I

298 ^{A,B,C}: On the same row, values with identical letters do not differ significantly ($p>0.05$) compared to the MRS-Cys
 299 control. ^{a,b,c,d,e,f,g,h,i,j,k}: On the same column, values with identical letters do not differ significantly ($p>0.05$). Values
 300 represent the mean±SD of three trials ($n=3$). BS: Bile Salts (oxgall); *GD=Growth delay: Distribution of isolates
 301 according to the growth delay and classification designed by Château et al. [18]. I= Resistant ($d\leq 15min$), II= Tolerant
 302 ($15<d\leq 60 min$), III= Poorly tolerant ($40<d\leq 60 min$), IV= Sensitive ($d>60min$).
 303

304 **3.4 Genes involved in acid and bile salts resistance**

305
 306 Genes have been sought to provide an explanation for the mechanism used by the isolates to tolerate
 307 acid and bile salts. The results showed that only the *clpL* gene (encoding ATPase) was found in the
 308 genome of all the 12 isolates tested (Table 5).
 309

310 **Table 5. Genes responsible for acid and bile salts survival in different isolates**
 311

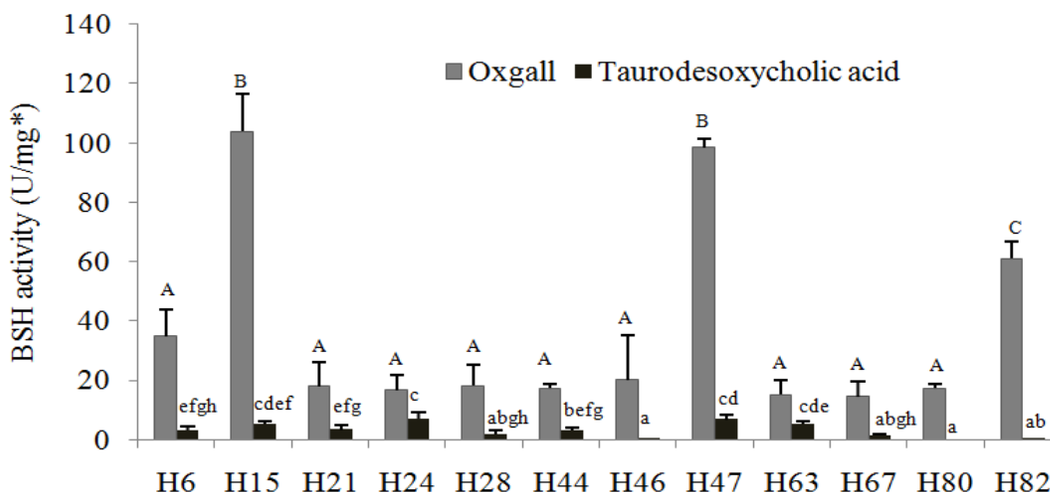
Isolates	Genes responsible for acid and bile salts survival			Bile salt hydrolase encoding genes			
	Acid resistance	Acid and bile salts resistance		<i>Bsh-Lp1</i>	<i>Bsh-1</i>	<i>Bsh-A</i>	<i>Bsh-B</i>
	<i>Hdc</i>	<i>clpL</i>	<i>gtf</i>				
H6	-	+	-	-	-	-	-
H15	-	+	-	-	-	-	-
H21	-	+	-	-	-	-	-
H24	-	+	-	+	-	-	-
H28	-	+	-	+	-	-	-
H44	-	+	-	-	-	-	-
H46	-	+	-	-	-	-	-
H47	-	+	-	-	-	-	-
H63	-	+	-	-	-	-	-
H67	-	+	-	-	-	-	-
H80	-	+	-	-	-	-	-
H82	-	+	-	-	-	-	-

+ = Presence of the gene, - = Absence of the gene

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 314 **3.5 In vitro activity of Bile Salt Hydrolase of the isolates**
 315

316 Fig. 1 shows the BSH activity of the isolates in the presence of 0.3% each of oxgall and
 317 taurodeoxycholate. This experiment showed that isolates exhibited a different level of hydrolysis activity
 318 on oxgall and taurodeoxycholate. As we can notice, the BSH activity of the isolates is higher in the

319 presence of oxgall than Taurodeoxycholate. Isolates H47 and H15 showed the highest activity on oxgall
 320 (98.53 ± 2.86 U/mg and 103.82 ± 12.93 U/mg respectively), whilst the lower value was observed with
 321 isolate H63 (15.10 ± 4.74 U/mg). On the other hand, BSH activity in the presence of taurodeoxycholate,
 322 was relatively low but still detectable. Compared to oxgall, the high value was 7.1 ± 1.9 U/mg (isolate
 323 H24) and the lower value at 0.03 ± 0.028 U/mg (isolate H24).



339 **Fig. 1. Bile Salts Hydrolase activity of different isolates**

340 *A,B,C;a,b,c,d,e,f,g,h*: For the same type of bile salt, values with identical letters do not differ significantly ($p>0.05$). Values
 341 represent the mean \pm SD ($n=3$), Error bars represent standard deviation. * 1 U/mg represents the amount of enzyme
 342 which releases 1 μ mol of amino acid from the substrate per minute. TDC: sodium Taurodeoxycholate.

344 **3.6 Genes involved in the Bile salts hydrolysis**

346 For specific genes responsible for the hydrolysis of the bile salts, the *bsh-Lp1* (*L. plantarum* Bsh1) gene
 347 was amplified only in isolate H24 and H28. While, none of the *bsh-1*, *bsh-A* and *bsh-B* genes were
 348 amplified on any of the isolates (Table 5).

350 After sequencing the *bsh-Lp1*, DNA sequences of BSH were obtained. They were designated Bsh_H24
 351 and Bsh_H28 respectively for the two isolates H24 and H28. The fragment contained single ORF 705
 352 nucleotides encoding a 234 amino acids protein with Bsh_H24, and ORF 726 nucleotides encoded 241
 353 amino acids protein with Bsh_H28 (Fig. 2). Both nucleotides are flanked by an alanine start codon (GCT)
 354 and a translational termination codon (TAA). The complete sequence has been deposited in GenBank
 355 database under the accession number of MF098540 and MF098541 respectively for the Bsh_H24 and
 356 Bsh_H28. Using the ClustalW program, these BSH sequences were aligned with other from GeneBank
 357 database. In general, the deduced amino acid sequence of the Bsh_H24 and Bsh_H28 display 100%
 358 identity with BSH-related proteins from *Lactobacillus plantarum* subsp. *Plantarum* P-8 (Accession
 359 number: AGL65610.2). They also exhibit 99% identity with BSH-related proteins from *Lactobacillus* sp.
 360 DPP8 (Accession number: ALT14558.1) and *Lactobacillus plantarum* (Accession number: ACA49878.1).

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BSH_LP1_H24 GENE ORF1 CDS translation	BSH_LP1_H28 GENE ORF1 CDS translation
1 GCTGATAAAGTTAATATCACACCATTGAAATTAATCTTGGTTA A D K V N I T P F E L I P W L	1 GCTGATTAAAAAATATGATGCTGATAAAGTTTATATCACACCA A D Y K K Y D A D K V Y I T P
46 TTGGGACAATTTCAAGTGTAGAGAAGTGA AAAAGACATACAA L G Q F S S V R E V K K N I Q	46 TTTGAATTAATTCCTGGTTATGGGACAATTTCAAGTGTAGA F E L I P W L L G Q F S S V R
91 AAACAAACTTGGTAAATTAATTTAGTGAACAATTACCATT K L N L V N I N F S E Q L P L	91 GAAGTGAAAAAAGAACATTCAAAAACAACTTGGTAAATTAAT E V K K N I Q K L N L V N I N
136 TCACCGTACATTGGTTGGTGTGATAAACAGGAATCGATAGTT S P L H W L V A D K Q E S I V	136 TTTAGTGAACAATTACCATTATCACCGCTACATTGGTTGGTGT F S E Q L P L S P L H W L V A
181 ATGAAAGTGTAAAGAAAGGACTAAAAATTCAGACAATCCAGTA I E S V K E G L K I Y D N P V	181 GATAAACAGGAATCGATAGTTATTGAAAGTGTAAAGAAAGGACTA D K Q E S I V I E S V K E G L
226 GGTGTGTTAACAAAACATCCTAATTTGACTACCAATTAATTAAT G V L T N N P N F D Y Q L F H	226 AAAATTTACGACAATCCAGTAGGTGTGTTAACAAAACATCCTAAT K I Y D N P V G V L T N N P N
271 TTGAACAACATATCGTGCTTATCAATAGCACACCTCAAAATAGT L N N Y R A L S N S T P Q N S	271 TTTGACTACCAATTAATTTGAAACAATATCGTGCTTATCA F D Y Q L F N L N N Y R A L S
316 TTTTGGAAAAAGTGGATTTAGATAGTTATAGTAGAGGAATGGGC F S E K V D L D S Y S R G M G	316 AATAGCACACCTCAAAATAGTTTTTCGGAAAAAGTGGATTTAGAT N S T P Q N S F S E K V D L D
361 GGACTAGGATTACCTGGAGACTTGTCTCAATGCTAGATTTGTC G L G L P G D L S S M S R F V	361 AGTTATAGTAGAGGAATGGCGGACTAGGATTACCTGGAGACTTG S Y S R G M G G L G L P G D L
406 AGAGCCGCTTTTAACTAAATTAACCTGTTGCGGATCGAGACAGAG R A A F T K L N S L P M Q T E	406 TCCTCAATGCTAGATTTGTCAGAGCCGCTTTTAACTAAATTAAC S M S R F V R A A F T K L N
451 AGTGGCAGTGTAGTCAGTTTTTCCATATACAGGAGTCTGTAGAA S G S V S Q F F H I L G S V E	451 TCGTTGCCGATGACAGACAGAGAGTGGCAGTGTAGTCAGTTTTTC S L P M Q T E S G S V S Q F F
496 CAACAAAAAGGGCTATGGAAGTTACTGACGGAAGTACGAATAT Q Q K G L C E V T D G K Y E Y	496 CATATACTAGGGTCTGTAGAACAACAAAAAGGGCTATGTGAAGTT H I L G S V E Q Q K G L C E V
541 ACAATCTATTCTTCTTGTGTATGGACAAGGGAGTTTATTAC T I Y S S C C D M D K G V Y Y	541 ACTGACGGAAGTACGAATATACAATCTATTCTTCTTGTGTGAT T D G K Y E Y T I Y S S C C D
586 TATAGAACTTATGACAATAGTCAAAATTAACAGTGTCAATTTAAAC Y R T Y D N S Q I N S V N L N	586 ATGGACAAGGGAGTTTATTACTATAGAACTTATGACAATAGTCAA M D K G V Y Y Y R T Y D N S Q
631 CATGAGCACTTGGATACGACTGAATTAATTTCTATCCATTACGA H E H L D T T E L I S Y P L R	631 ATTAACAGTGTCAATTTAAACCATGAGCACTTGGATACGACTGAA I N S V N L N H E H L D T T E
676 TCAGAAGCACAACTATGACAGTTAACTAA S E A Q Y Y A V N *	676 TTAATTTCTTATCCATTACGATCAGAACACAATACTATGCAGTT L I S Y P L R S E A Q Y Y A V
	721 AACTAA H *

Fig. 2. Nucleotide sequence and deduced amino acid sequence of the BSH_H24 and BSH_H28 gene of the respectively H24 and H28 Isolate (further identified as *Lactobacillus plantarum* sp.)
On the left, *Bsh_H24* gene (accession number MF098540) and on the right *Bsh_H28* gene (accession number MF098541) deposited in NCBI GenBank data base (<http://blast.ncbi.nlm.nih.gov/Blast.cgi>).

3.7 Principal Component Analysis (PCA)

With PCA, we noticed that the first axis (F1) makes it possible to explain 60.54% of the total variance, and separates the isolates into two groups: those that survive acidity and bile salts (right) and those presenting excellent growth time performance (low time required to increase the absorbance by 0.3 unit and growth delay) (left). The second axis (F2) that opposes survival to the acidity (top) and survival to the bile salts (bottom) explains 23.93% at its level (correlation Biplot, Fig. 3). The variables BSH activities on oxgall and survival are almost orthogonal represented, indicating that they are significantly uncorrelated. The isolates well represented on the F1 axis are H44, H63 and H47, while on the second Principal Component F2 has a high contribution of isolates H15 and H47.

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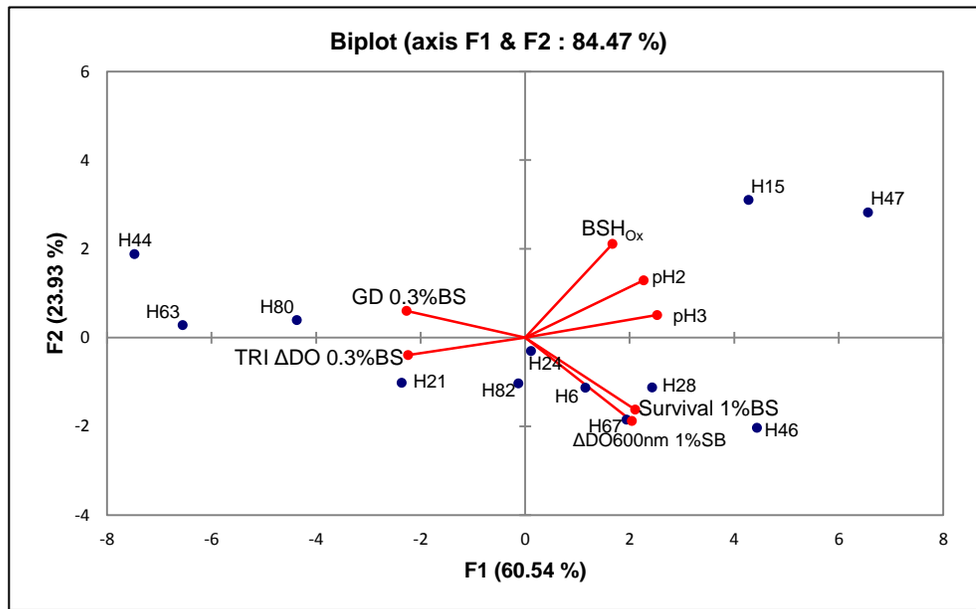


Fig. 3. Graphical representation of the correlation Biplot from the Principal Component Analysis
 pH3, pH2: Survival rate to pH2 and pH3. ΔDO0.3%BS: Variation of absorbance A600nm after 24h at 0.3% bile salt.
 TRI ΔDO 0.3%BS: Time Required to Increase A600nm by 0.3 units (min). Survival 1%BS: Survival rate in 1% Bile
 Salt after 24h. GD 0.3%BS: Delay of growth during the exponential phase at 0.3% bile salt (min). BSH_{ox}: Bile Salt
 Hydrolase Activity on the Oxgall (U/mg).

3.8 Antimicrobial activity

Table 6 indicates results of preliminary antimicrobial activity against a range of indicator bacteria such as *L. plantarum* 5S (bacteriocin's sensitive strain), food spoilage or pathogenic bacteria on the agar medium by the spot technique. It appears that all our isolates exhibit the antimicrobial effect against *L. plantarum* 5S strain. Furthermore, isolates showed antibacterial activities against the indicator bacteria with different levels. The isolates H15 and H24 showed higher antagonistic activity.

Table 6. Inhibitory activity of the isolates against *L. plantarum* 5S and the indicator spoilage or pathogenic bacteria

Strains	H6	H15	H21	H24	H28	H44	H46	H47	H63	H67	H80	H82
<i>Lb.p</i> 5S	+++	+++	+++	+++	+++	+++	+++	+++	+++	+++	+++	+++
SA	-	++++	-	+++	+++	-	+	-	-	-	+++	-
ST	-	+++	++	++++	-	-	-	-	++	++	-	-
BC	-	+++	+	+++	-	-	-	-	+++	+	+	-
EC	-	+++	-	+++	-	+	-	-	-	-	-	+
SM	-	++	-	+	-	-	+	-	-	-	-	-
LI	-	++	-	-	-	-	-	-	++	++	++	-
PA	-	-	-	+	+++	+++	+++	-	+	-	+	-
PM	+++	++++	-	++++	++	++	-	+++	-	+	-	++

472 – no inhibition; + 1.0–3.0 mm (weak); ++ 3.1–6.0 mm (good); +++ 6.1–14.0 mm (very good); ++++ >14.0 mm (strong).
 473 The diameter of inhibition was calculated as the difference between the total of inhibition zone and the diameter of
 474 growth spot of selected strains (n=3). *Lb.p* 5S: *L. plantarum* 5S. *LI*: *Listeria innocua* ATCC 33090. *SA*:
 475 *Staphylococcus aureus* ATCC 25923. *SM*: *Streptococcus mutans* DSM 20523. *BC*: *Bacillus cereus* 11778. *PM*:
 476 *Proteus mirabilis* (Clinical isolate). *EC*: *Escherichia coli* ATCC 13706. *ST*: *Salmonella enterica* serovarTyphi ATCC
 477 6539. *PA*: *Pseudomonas aeruginosa* ATCC 20027.
 478

479 **3.9 Molecular Identification of Lactic Bacteria**

480
 481 A step of characterization of the isolates based on the PCR amplification and the sequencing of the gene
 482 encoding the 16S rRNA was carried out to identify the 12 pre-selected isolates and the results are
 483 presented in Table 7. The gene targeted in all these isolates has been amplified. Therein, we can find the
 484 code of each isolate and its origin, the accession number provided by NCBI (from KU886166 to
 485 KU886177), the genus and species name of the corresponding lactic bacteria strain. All the 12 isolates
 486 were categorized as the genus *Lactobacillus* which showing more than 99% identity to *L. plantarum* and
 487 *L. paraplantarum* already present in the NCBI GenBank.
 488

489 **Table 7. The 16S rRNA gene sequencing identification of LAB isolates**

490

Strains	Origin	16S rRNA sequencing identification	Sequence length (bp)	% Query coverage	% Max identity	Accession number
H6	Fossong Wentcheng	<i>Lactobacillus plantarum</i>	816	100	99	KU886167
H15	Penka-Michel centre	<i>Lactobacillus plantarum</i>	720	100	100	KU886171
H21	Penka-Michel centre	<i>Lactobacillus plantarum</i>	748	100	100	KU886168
H24	Penka-Michel centre	<i>Lactobacillus plantarum</i>	777	100	99	KU886174
H28	Penka-Michel centre	<i>Lactobacillus plantarum</i>	708	100	99	KU886169
H44	Bamendou, <i>QtNguim</i>	<i>Lactobacillus plantarum</i>	597	100	99	KU886176
H46	Bamendou, <i>QtNguim</i>	<i>Lactobacillus plantarum</i>	708	100	99	KU886170
H47	Bamendou, <i>QtNguim</i>	<i>Lactobacillus plantarum</i>	939	100	100	KU886166
H63	Balessing, <i>King Place</i>	<i>Lactobacillus paraplantarum</i>	729	100	100	KU886177
H67	Balessing, <i>King Place</i>	<i>Lactobacillus paraplantarum</i>	726	100	99	KU886173
H80	Balessing, <i>King Place</i>	<i>Lactobacillus paraplantarum</i>	1147	100	99	KU886175
H82	Balessing, <i>King Place</i>	<i>Lactobacillus plantarum</i>	588	100	100	KU886172

491
 492 **4. DISCUSSION**

493
 494 The sensitivity of the isolates to low pH was carried out to predict their behavior during gastrointestinal
 495 transit in human. We found that 52.94% and 94.11% could survive respectively to pH 2 and pH3. It has
 496 been shown that species of the genus *Lactobacillus* are tolerant to gastric acid conditions [25]. Our
 497 results are in agreement with those of Prasad et al. [26] who obtained acid-tolerant strains from 200
 498 isolates, following their survival of nearly 80% after exposure to pH3 for 5 h. Several mechanisms have
 499 been elucidated to explain the resistance of lactobacilli to acid stress. Among them are the protomotor
 500 force F1F0 ATPase, DNA repair mechanisms, modification of the composition, architecture and stability of
 501 the plasma membrane, production of alkaline compounds by the action of urease or Arginine deiminase
 502 (ADI), and the management of denatured proteins [27].
 503

504 Bile salts are the second important factor faced by probiotic LAB in the digestive tract. The growth delay
 505 at 0.3% bile salt obtained with our isolates is significantly better than that of strains of commercial lactic
 506 acid bacteria isolated from faeces by Mirolohi et al. [28]. They showed that strain *L. plantarum* A7 exhibited
 507 growth delay greater than 1 hour at 0.3% bile salts. For the use as a probiotic in humans, LAB must

508 survive at a concentration of 0.3% bile salts [29]. According to the classification of Château et al. [18],
509 none of our isolates was classified as sensitive to bile salts. Thus, all the isolates with survival of more
510 than 79%, at 0.3% bile salt concentration, can probably overcome the bile stress in the intestine if subject
511 to *in vivo* assay.

512
513 Known mechanisms can explain this resistance. We can mention the extrusion of the bile, achieved by
514 efflux systems including the multidrug resistance (MDR) family [30]. Another mechanism is the
515 deconjugation of bile acids. It is catalyzed by bile salts hydrolases (BSH), enzymes that release
516 glycines/taurins from the steroid nucleus, which lowers the solubility of bile at low pH and reduces their
517 detergent activity on bacterial membranes [7].

518
519 All the 12 isolates tested were found to possess the *clpL* gene coding for ATPase. This gene could have
520 played an important role in the resistance to acid and biliary stress found during *in vitro* phenotypic tests.
521 Turpin et al. [31] reported the presence of *clpL* gene allowing low pH resistance in 91% to 100% of the
522 isolates from their collection. In *Lactobacillus reuteri* ATCC 55730, studies have shown that inactivation of
523 *clpL* has resulted in a significant decrease in bacterial survival after incubation at pH 2.7 [32] or medium
524 containing 0.3% bile salts [33].

525
526 Genes have been sought to provide an explanation for the mechanism used by these isolates tolerate the
527 gastrointestinal stress conditions. None amplification of *gtf* and *hdc* genes can be explained by their
528 phylogenetic distribution. Indeed, the primers used would be restricted to a set of species or subspecies
529 absent from the collection of lactic bacteria that we have isolated; or it is possible that the tolerance to
530 acidity and resistance to bile salts found *in vitro* is due to a mechanism controlled by other genes. Similar
531 results were obtained by Turpin et al. [31]. They showed that none out of 38 isolates tested harbored the
532 genes *gtf* and *hdc*. Non-expression of the *hdc* gene is an advantage because; ingestion of a large amount
533 of histamine can cause serious problems into the body. This biogenic amine is formed after
534 decarboxylation of histidine by the enzyme histidine decarboxylase encoded by the *hdc* gene.

535
536 In this study, we also evaluated the ability of isolates to perform BSH activity. We observed that they
537 possessed this activity. According to Tanaka et al. [34], in an analogous manner, all the lactobacilli
538 isolated from the gastrointestinal media possess the BSH activity. In all of our isolates, BSH activity was
539 higher with oxgall than with Taurodeoxycholate. Therefore, they have a substrate preference for oxgall. It
540 has been demonstrated that most strains of LAB exhibit high activity with conjugated bile salt mixtures
541 than with a particular type of conjugated bile acid [9]. Also, Kumar et al. [35] showed that the majority of
542 lactobacilli tested had more valuable BSH activity on glycocholate than on taurocholate or
543 taurodeoxycholate. However, since glycocholate is the most abundant of the bile salts found in humans, it
544 would be advantageous to have isolates exhibiting a preference for glycocholate. According to Brashears
545 et al. [36], these isolates could be candidate for the *in vivo* reduction of serum cholesterol levels. Lately, it
546 has been revealed that the BSH take part in a most important role in cholesterol metabolism, thus
547 influencing the serum cholesterol levels [35]. It has also been suggested that BSH activity must be
548 essential in the choice of probiotic organisms with cholesterol-lowering properties, given that
549 microorganisms that do not deconjugate bile salts cannot reduce cholesterol from a medium to a
550 significant level [9, 35].

551
552 Several authors have suggested that the resistance of lactobacilli to the toxicity of bile salts in the
553 duodenum could be attributed to the activity of the BSH enzyme [37]. Although BSH activity is widespread
554 in lactobacilli, there is not always a direct relationship with their ability to resist bile [38]. This situation is
555 supported by the numerous functional studies carried out on lactobacilli strains whose genome is
556 sequenced but for which the deletion of a gene coding for BSH does not necessarily have any
557 consequence on the strain survival in the presence of bile salts [39,40]. This enzyme is encoded by the
558 *bsh* gene. Our results showed only the *bsh*-Lp1 gene (*L. plantarum* Bsh1) in H24 and H28. Studies have
559 shown that the presence and genetic organization of *bsh* genes in lactobacilli are very variable [41].
560 There are four alleles of the *bsh* (*bsh*-Lp1, *bsh*-Lp 2, *bsh*-Lp 3 and *bsh*-Lp 4) in *L. plantarum*, but the
561 highest BSH activity is correlated with the *bsh*-Lp 1 [39].

562

563 Some of our isolates have very high percentages of survival at high bile salt concentrations without
564 having such a high BSH activity. The case of the isolate H46 which exhibited a very low BSH activity of
565 20.20 ± 14.83 U/mg compared to that of the H15 isolate (103.82 ± 12.93) on oxgall. The same
566 observation was also noted by the principal component analysis (PCA), where the variables such as the
567 activity of BSH on oxgall and bile salt survival were shown to be significantly uncorrelated. Recent studies
568 have shown that the resistance of lactobacilli can not necessarily be associated with the presence of BSH
569 [42]. According to the PCA, it can be concluded that the strains H15 and H47, based on their
570 representation, are strongly resistant to acidity; the strains H28, H67 and H46 are more tolerant to bile
571 salts.

572
573 The satisfying probiotics are supposed to exhibit their antimicrobial activities generally against pathogens
574 in the gastrointestinal system [43]. In this study, we used pathogenic bacteria (such as *L. innocua*, *S.*
575 *aureus*, *S. Typhi*, *E. coli* and *B. cereus*) because they are often found as food-borne pathogens that might
576 cause gastroenteritis. The presence of the inhibition zones indicated the antibacterial activity of our
577 isolates on the indicator bacteria. In fact, LABs are recognized for their production of various antimicrobial
578 substances (organic acid, hydrogen peroxide, diacetyl, reuterin and bacteriocins) [44]. The production of
579 these substances responsible for the antagonistic phenomenon by our isolates is important to their
580 antimicrobial property, and they could more expect to be used as probiotic.

581
582 According to the joint FAO/WHO [2] expert report on the presence of probiotics in food, it is necessary to
583 know the genus and species of a probiotic strain. All the 12 isolates were found to belong to the genus
584 *Lactobacillus*. Indeed, as reported by Tannock [25], the genus *Lactobacillus* occurs in a variety of habitats
585 including plants, the gastrointestinal tract of animals such as bees, and is the most dominant LAB found
586 in the intestinal tract of bees [45]. The high presence of the *plantarum* species was noted by [46] who
587 found that the *L. plantarum* strain was the most abundant (51.02%) of the 5 *Lactobacillus* phylotypes
588 identified in the honey bee *Apis dorsata* in Malaysia. The high presence of *plantarum* and *paraplantarum*
589 species in this collection could be explained by the origin of the type of bee collected during sampling. We
590 collected foraging bees, which are frequently in contact with plants (including pollen and nectar of the
591 flowers); those are habitats of *plantarum* species. This species may well be found in the intestinal tract of
592 bees because most lactic acid bacteria that exist in the intestinal tract of bees are also isolated from
593 pollen [47].

594 595 **5. CONCLUSION**

596
597 The results obtained in this study show that Lactobacilli isolated from honeybee gut in the Menoua
598 division (West-Cameroon) can survive low pH and bile salts. Their BSH activity may contribute to lower
599 the blood cholesterol levels. They also possess antibacterial activity on pathogenic bacteria. Thus, the
600 cultures obtained in this work could be presumed as potential probiotic bacteria. Complementary
601 investigation of certain strain will attest their safety, probiotics properties and more.

602 603 604 **CONFLICT OF INTEREST**

605
606 The authors declare no competing interests
607

608 609 **REFERENCES**

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