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3 **Screening and Identification of Lactic Acid Bacteria**

4 **with D-tagatose Production Capability**

5

6 **ABSTRACT**

D-tagatose is a natural ketohexose which can be used as a functional sweetener in foods, dairy and beverages products. Isolation of new bacterial strains having the ability to produce D-tagatose is a continuously trending topic of research. In this study, 4 strains of lactic acid bacteria (LAB) were isolated from kimchi sample. The isolates were identified as *Lactobacillus* spp. (*Lactobacillus plantarum*, *Lactobacillus fermentum* and *Lactobacillus salivarius*) on the basis of morphological, physicochemical characteristics and analysis of 16S rDNA gene sequence. Because of the novelty, strain designated as *L. salivarius* UJS 003 was considered for D-tagatose yield. Fermentation of D-tagatose was carried out using galactose as substrate for 48 hr at 37 °C, and HPLC method was used to determine the yield. The experimental results exhibited a D-tagatose yield of 3.134 g/L by *L. salivarius* UJS 003. The strain UJS 003 represented as a potent D-tagatose producer and could be useful in a variety of biotechnological and industrial processes, particularly food and beverage industries.

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8 *Keywords: D-tagatose; identification; L. salivarius; strain- screening; biotransformation*

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10 **1. INTRODUCTION**

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12 The rare sugar monosaccharide D-tagatose, which has the advantage of having low calorific content, accompanied by more than 90% sweetness compared to sucrose. As D-tagatose has been classified by the US Food and Drug Administration (FDA) to be generally recognized as safe (GRAS), it is considered as a promising sweetener that can be utilized in various applications in food industries [1-3]. Due to its relative low glycemic index, it can be considered as an alternative to glucose by playing an important role in mitigating the effect on hyperglycemia, type-2 diabetes, probiotic function, and antioxidant activity [3-5]. Being ketohexose and isomer of D-galactose, the production D-tagatose via chemical isomerization through the calcium catalyst was found to be inapplicable due to the lengthy and complex purification steps as well as the environmental unfriendly wastes. Therefore, the production of D-tagatose by biological approaches is found to be more desired as safer regards to both health and environment. In recent years, many studies demonstrated the production of D-tagatose via the enzymatic catalysis by employing L-arabinose isomerase which is capable of directly converting D-galactose to D-tagatose [6-9]. However, the key determinative steps are to screen the strains with high L-arabinose enzyme activity suitable for large-scale production, to construct the enzyme expression system and to explore the appropriate conversion process. The use of lactic acid bacteria (LAB) strains in food industry is a major field as the final fermentation products are readily available for human consumption [10]. To the above context, the current study aims to screen and identify novel D-tagatose producing lactic acid bacterium strain from kimchi liquid sample.

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31 **2. MATERIALS AND METHODS**

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33 **2.1 Samples, Chemical and Reagents**

34 Kimchi liquid sample, the source of isolated strains, was obtained from in local markets in Zhenjiang city
35 (Jiangsu, China). All chemical and reagents used are of analytical grade quality.
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37 **2.2 Enrichment and Isolation of D-tagatose Producing Bacteria**

38 A potential D-tagatose producer was isolated by applying enrichment culture technique using De Man
39 Rogosa and Sharpe (MRS) medium. The composition of liquid medium was as following (g/L): Glucose
40 (4.0), Beef extract (2.0), Peptone (1.0), Yeast extract (1.0), CH₃COONa (1.0), C₆H₁₇N₃O₇ (0.4),
41 Na₂HPO₄ (0.4), MgSO₄ (0.24), MnSO₄ (0.06) and C₆H₁₂O₅ (2.0). The pH of the medium was adjusted
42 to 6.2 and sterilized by autoclaving at 121 °C for 20 min. 1 mL of kimchi liquid sample as inoculated in the
43 MRS medium and incubated at 37 °C on the rotary shaker at 150 rpm for 24 hr. Following the incubation
44 period, 3 mL of culture was transferred to fresh medium containing and re-incubated for another 24 hr.
45 This step was repeated several times, and the growth was considered when the natural purple color of
46 the culture medium becomes yellowish. Then, samples from appropriate multiple dilutions were
47 inoculated on MRS agar plates containing (g/L): CaCO₃ (20), bromcresol purple (16) and Agar (15). After
48 the incubation period of 24 hr, single colonies exhibiting a yellowish color were selected and serially
49 cultured on MSR agar plates. Further, the isolated strains were stored as frozen stock cultures at -70 °C
50 in 25% (v/v) glycerol.
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52 **2.3 Screening for D-tagatose Producers**

53 The isolates were inoculated in MRS broth for 12 hr at 37 °C and activated for 2 generations. Later, 1 mL
54 of static inoculum culture was used to access the test tube containing 30 mL fermentation medium. The
55 fermentation broth was centrifuged for 20 min (5000 rpm, 4 °C), washed 2 times with phosphate buffer,
56 and the supernatant was discarded. Later, the bacterial cells possessing the characteristics of yellowish
57 colouration and the CaCO₃-dissolving capacity were selected and inoculated on MRS agar plate at 37 °C
58 for 48 hr. Ketoses production ability of strains were tested by using the cysteine carbazole sulfuric-acid
59 (CCSA) method, and by measuring absorbance at 560 nm [8, 11].
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61 **2.4 Strain Identification**

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63 **2.4.1 Phenotypical Characterization of Isolated Strains**

64 The bacterial cell morphology was observed under microscope after Gram staining. Additionally,
65 gluconate test, carbohydrate fermentation test, and mannitol test were conducted for physiological and
66 biochemical identification of the strain according to Bergey's manual of systematic bacteriology.

67 **2.4.2 Molecular Identification of Isolated Strains**

68 The isolates were identified by sequencing the 16S ribosomal DNA gene. For 16S rDNA sequencing, total
69 genomic DNA was extracted from the isolate and amplified by using the universal primers 27F [5-
70 AGAGTTTGATCCTGGCTCAG-3] and 1492R [5- GGTTACCTTGTTACGCTT-3]. The PCR conditions
71 were as follows; an initial denaturation at 94 °C for 5 min, followed by 35 cycles of denaturing at 94 °C for
72 30 s, annealing at 53 °C for 30 s, extension at 72 °C for 90 s and a final extension step at 72 °C for 10 min.
73 The resulted PCR products were loaded on 2.0% agarose gel and analyzed under UV transilluminator
74 after staining with ethidium bromide solution. The PCR amplified DNA was purified using Tiangel Midi
75 Purification Kit (TIANGEN, Beijing, PR China). DNA sequencing was performed by Suzhou Hongxun
76 Biotech Co., Ltd., (Suzhou, China). All the 16S rDNA gene sequences were cross-checked manually,
77 edited for phylogenetic analysis and the similarities were determined by using BLASTN multiple sequence
78 alignment.
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80 **2.5 D-tagatose Production**

81 D-tagatose production was carried out in the fermentation medium with the following composition (g/L);
82 Galactose (20), Peptone (5.0), Yeast extract (10), K₂HPO₄ (0.2), MgSO₄ (0.2), MnSO₄ (0.1) and NaCl
83 (0.01). The pH of the medium was adjusted to 6.2 and sterilized by autoclaving at 121 °C for 20 min. An

84 aliquot of 2 mL pre-cultured isolate from the MRS culture broth was inoculated in the flask containing 200
 85 mL of fermentation medium and incubated for 48 hr at 37 °C. After the incubation period, the fermentation
 86 broth was centrifuged for 20 min (5000 rpm, 4 °C), washed 2 times with phosphate buffer, and then finally
 87 the supernatant was discarded. D-tagatose production ability of strains was determined by High
 88 Performance liquid chromatography (HPLC) equipped with Xtimate Sugar-Ca, 5 µm, 7.8 × 30 mm (Welch,
 89 Shanghai, China) column and RID-20A refractive index detector (Shimadzu, Japan). The mobile phase
 90 was pure water running at a flowrate of 0.5 mL/min. A sample volume of 10 µL was injected to the column
 91 temperature (80°C) [1].
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93 3. RESULTS AND DISCUSSION

94 3.1 Bacterial Strains Screening

95 Due to diversity of microorganisms and different niches in which they inhabit, there is a need of efficient
 96 isolation and screening methods. Apart from direct isolation of strains by diluting and plating, enrichment
 97 cultures very promising for the isolation of D-tagatose producing microbes. The initial screening of all
 98 isolates obtained from kimchi solution revealed a total of 15 potential D-tagatose producing strains which
 99 were identified as Gram-positive bacteria based on the enrichment and isolation strategy employed in this
 100 study. These selected strains obtained were inoculated into the fermentation medium, and 4 strains with
 101 the ability of D-tagatose production was identified by the cysteine carbazole sulfuric-acid method. The
 102 strains were thereafter designated as UJS001, UJS002, UJS003 and UJS004 respectively.
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104 3.1.1 Morphology Physiology and Biochemistry Characteristics

105 The selected strains were cultured on the MRS solid medium for 24 hr at 37 °C, and the morphological
 106 characteristics of the colonies were observed. The results of morphological, physiological and
 107 biochemical characteristics of the selected isolates designated as UJS 001, UJS 002, UJS 003 and UJS
 108 004 are presented in Table 1. Colony shapes varied between irregular circle and circular, with white or
 109 milky white color, whereas cells were Gram-positive, rod-shaped, and capable of producing aeris via
 110 gluconate. The morphological and biochemical characteristics of these 4 strains matched with the
 111 description about genus *Lactobacillus* which is available in the literature [13-15].
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113 **Table1 . Morphological, physiological and biochemical characteristics of the isolated strains**

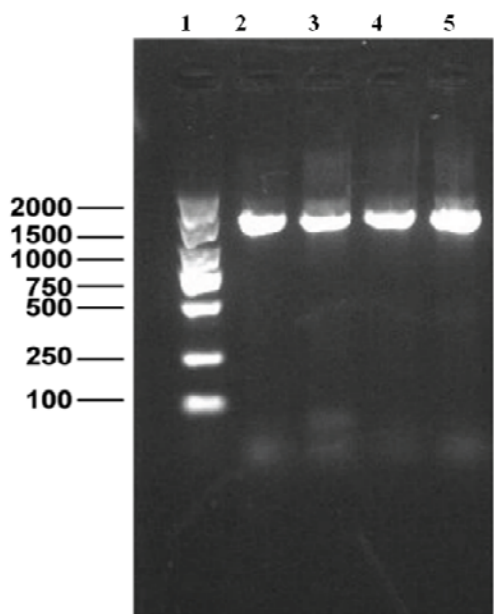
Strain No.	Colour	Colony shape	Pellucidity	Mycelial morphology	Gluconate	Carbohydrate fermentation	Mannitol
UJS001	White	Irregular circle	Lucency	Short rod	-ve	-ve	+ve
UJS002	Milky white	Irregular circle	Translucency	Short rod	-ve	-ve	+ve
UJS003	Milky white	Circular	Lucency	Short rod	-ve	-ve	+ve
UJS004	White	Circular	Lucency	Short rod	-ve	-ve	+ve

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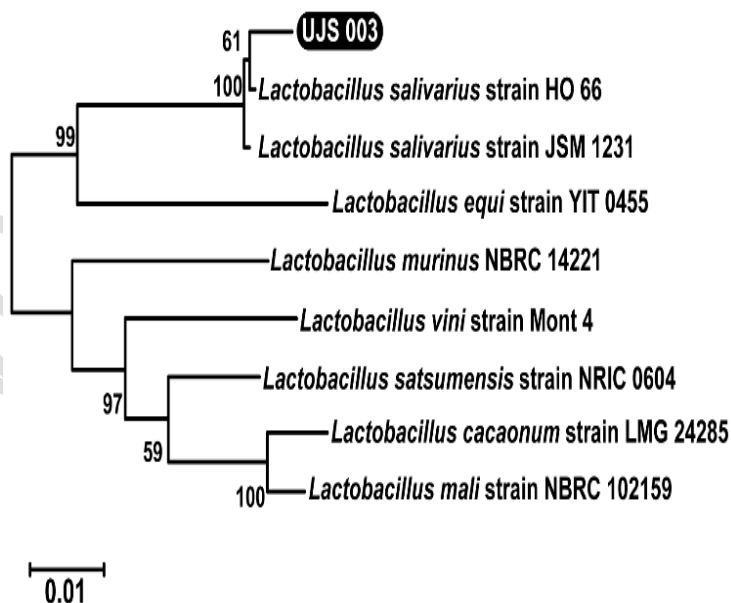
116 3.2 Molecular Identification of Bacterial Strain

117 Fig.1. represents the 16S rDNA gel electrophoresis exhibiting four bands corresponding to each of the
 118 four screened strains. Subsequent phylogenetic analysis confirmed that the strain UJS 001, UJS 002,
 119 UJS 003 and UJS 004, belongs to genus *Lactobacillus*. The relationship of these strains with the nearest
 120 phylogenetic relatives are described in Fig.2. According to the similarity calculations following the
 121 phylogenetic analysis, the closest relatives of UJS 001, UJS 002, UJS 003 and UJS 004, were *L.*
 122 *plantarum* (99 %, UJS 001/002), *L. salivarius* (100%) and *L. fermentum* (99%), respectively. Since there
 123 are many studies on *L. plantarum* and *L. fermentum* that can produce D-tagatose, the authors have
 124 selected UJS 003 for further research. The strain UJS 003 is being designated as *L. salivarius* UJS 003.

125 The similarity between the UJS003 and the JCM 1231 of *L. salivarius* was 99%. However, *L. spp.* are
126 well documented for D-tagatose production except for *L. salivarius* [7, 9, 12-14]. Henceforth, authors have
127 considered to *L. salivarius* UJS 003 as a model strain to determine the D-tagatose producing capability.
128



129 **Fig.1. Agarose gel electrophoresis showing partial rDNA sequence by genomic PCR**
130 Lane 1: 2000 bp DNA marker, Lane 2, 3, 4 and 5 represents the DNA isolated from UJS 001, UJS 002, UJS 003 and
131 UJS 004, respectively.
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134 **Fig.2. Phylogenetic tree of *L. salivarius* UJS 003**
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138 3.3 Quantitative Analysis of D-tagatose

139 HPLC was employed for the detection and quantification of tagatose production, after analysing, the yield
140 is 3.134g/L. Several studies demonstrated the use of *L.* strains for the biosynthesis of D-tagatose
141 including *L. sakei* [7], *L. fermentum* [12], *L. reuteri* [13] *L. plantarum* [9] and *L. lactis* [14]. This study
142 reports for the first time, the use of the probiotic bacterial strain *L. salivarius* for the production of D-
143 tagatose [15]. However, there are currently few studies about *L. salivarius*, so we have considered study
144 the capability of producing D-tagatose.
145

146 4. CONCLUSION

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148 A D-tagatose producing *L.salivarius* UJS 003, isolated from kimchi sample is reported. At present, the use
149 of enzymes to convert D-galactose into D-tagatose has a lot of reports, but there is a dirt report on the
150 conversion of *L. salivarius*. We need to further study the transformation mechanism to improve the
151 conversion rate. This is the first report on D-tagatose production by *L. salivarius* which may show potential
152 application in some industries especially for food industries. And therefore, merit for further research.
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