Adaptive mechanism of *Lactobacillus amylolyticus* L6 in soymilk environment based on metabolism of nutrients and related gene-expression profiles

Yongtao Fei1,2,3 | Li Huang1 | Hong Wang1,2 | Jinglong Liang1,3 | Gongliang Liu1,2,3 | Weidong Bai1,2,3

1Guangdong Provincial Key Laboratory of Lingnan Specialty Food Science and Technology, Zhongkai University of Agriculture and Engineering, Guangzhou, China
2College of Light Industry and Food Science, Zhongkai University of Agriculture and Engineering, Guangzhou, China
3Academy of Contemporary Agricultural Engineering Innovations, Zhongkai University of Agriculture and Engineering, Guangzhou, China

**Correspondence**
Gongliang Liu and Weidong Bai, Guangdong Provincial Key Laboratory of Lingnan Specialty Food Science and Technology, Zhongkai University of Agriculture and Engineering, Guangzhou, China.

**Emails:** gongliangliu@126.com (G.L.) and whitebai2001@163.com (W.B.)

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**Abstract**
*Lactobacillus amylolyticus* L6 isolated from naturally fermented tofu-whey was characterized as potential probiotics. To give insight into the adaptive mechanism of *L. amylolyticus* L6 in soymilk, the gene-expression profiles of this strain and changes of chemical components in fermented soymilk were investigated. The viable counts of *L. amylolyticus* L6 in soymilk reached $10^{12}$ CFU/mL in the stationary phase (10 hr). The main sugars reduced gradually while the acidity value significantly increased from 45.33° to 95.88° during fermentation. About 50 genes involved in sugar metabolism and lactic acid production were highly induced during soymilk fermentation. The concentration of total amino acid increased to 668.38 mg/L in the logarithmic phase, and 45 differentially expressed genes (DEGs) in terms of nitrogen metabolism and biosynthesis of amino acid were detected. Other genes related to lipid metabolism, inorganic ion transport, and stress response were also highly induced. Besides, the concentration of isoflavone aglycones with high bioactivity increased from 14.51 mg/L to 36.09 mg/L during the fermentation, and the expression of 6-phospho-β-glucosidase gene was also synchronously induced. This study revealed the adaptive mechanism of *L. amylolyticus* L6 in the soymilk-based ecosystem, which gives the theoretical guidance for the application of this strain in other soybean-derived products.

**KEYWORDS**
chemical components, *Lactobacillus amylolyticus* L6, soymilk, transcriptome
1 | INTRODUCTION

Soybean and its derived food products are important part of the Asian diet. These foods are rich in various nutrients, such as protein, oligosaccharides (raffinose and stachyose), grease, vitamins, and insoluble dietary fiber (Lokuruka, 2011). Meanwhile, they were reported to have many beneficial functions to consumers, including the prevention of cardiovascular disease, osteoporosis, hormone-related cancers, and modulation of immunity and intestinal flora (Ko, 2014; Yan et al., 2017). As consumers become increasingly interested in functional foods, they have higher requirements for the varieties of functional soybean products. In recent years, fermentation of soymilk by probiotics has become one of the research hotspots because of the function-promoting effects brought about by these microorganisms and soymilk (Marazza et al., 2013; Wang et al., 2012; Wei et al., 2007).

Probiotics were able to attach to the surface of intestinal mucosa and colonize in the intestinal tract, which could allow them to bring beneficial effects to human health (Bron et al., 2012). For example, the probiotics could competitively exclude and inhibit pathogens in the intestinal tract (Kholy et al., 2014), enhance intestinal flora (Gerritsen et al., 2011), augment both cellular and humoral immunity (Yan & Polk, 2011), and relieve inflammation and food allergy (Majamaa & Isolauri, 1997). Except for the above functional characteristics, the probiotics used as soymilk starter were required to adapt to a complex nutritional environment of soymilk. In general, the minimum number of living probiotics in the final product of soybean yoghurt should reach $10^8$ CFU/ml (Shah, 2000). Meanwhile, the pH for coagulating soymilk was 4.5–5.0 (Qiao & Li, 2018), which required the strong acid-producing ability of probiotics in soymilk. Although the stachyose and raffinose in soymilk have been regarded as prebiotics, excessive intake by human body would cause gastric bloating or diarrhea, requiring probiotics to own ability of hydrolyzing soybean oligosaccharides in soymilk with α-galactosidase (Donkor et al., 2007). In addition, soymilk is rich in low-absorptive isoflavone glycosides (occupying approximately 90% of isoflavone content) (Izumi et al., 2000), and the probiotic strain with the ability of converting isoflavone glycosides into high-absorptive aglycones by β-glucosidase were the best choice (Donkor et al., 2007). On the other hand, stachyose and raffinose in soymilk could promote the proliferation of fermentation probiotic strains (Kim et al., 2010; Sarina et al., 2017). In addition, the soymilk could be used as food vehicles of probiotics, protecting bacterial cells from adverse environment such as low pH of gastric acid, bile salt, and various digestive enzymes in the gastrointestinal tract (Zhuang et al., 2009). Therefore, the selection of probiotic strains suitable for soymilk environment is very important for the production of soybean yoghurt.

Lactobacillus amylyticus L6 was isolated from naturally fermented tofu-whey (Fei et al., 2018) and its safety, potential probiotic characteristics, and fermentation properties in tofu-whey have been extensively studied (Fei et al., 2020; Fei, Li, et al., 2017; Fei, Liu, et al., 2017). Since L. amylyticus L6 was one of the dominant bacteria in naturally fermented tofu-whey for a long time, it has evolved the adaptability to nutritional environment in soybean products, which makes L. amylyticus L6 one of the best candidate probiotic strains for fermenting soymilk. In this study, the changes of nutrient and functional substances in soymilk and gene-expression profiles of L. amylyticus L6 during fermentation were investigated to reveal the molecular mechanisms of synergistic effect between soymilk and L. amylyticus L6.

2 | MATERIALS AND METHODS

2.1 | Strains and cultivation

Lactobacillus amylyticus L6 (CGMCC NO.9090) was isolated from naturally fermented tofu-whey (Fei et al., 2018). This strain was preserved in 15% glycerol at −80°C and cultivated in De Man, Rogosa and Sharpe (MRS) (Guangdong Huankai Microbiology Biotech Inc., Guangzhou, China) plate at 37°C for 36 hr before use. A single colony was then picked and inoculated into 10 ml of MRS broth and incubated for 24 hr.

2.2 | Preparation of fermented soymilk

Soymilk was prepared according to the method described by Salma et al. (Elghali et al., 2014) with slight modification. Soybean (100 g) was washed and then soaked in 600 ml of drinking water with 0.5% NaHCO$_3$ at 26°C for 14 hr. The soaked soybean was ground and heated with 800 ml of drinking water in a soymilk maker (DJ12B-DEF4, Midea, China). The slurry was filtered through a double-layered cotton cloth and then mixed with drinking water in a ratio of 8:2. Glucose (Sigma Chemical Co., Ltd, Guangzhou, China) with a concentration of 1.5% (w/v) was added to make soymilk. Soymilk was heated at 85°C for 15 min for sterilization and then cooled to 37°C. Subsequently, the soymilk was inoculated with 10% (w/v) L. amylyticus L6 and incubated at 37°C for 24 hr. The growth curve was plotted according to the viable counts determined at 0 hr, 2 hr, 4 hr, 6 hr, 8 hr, 10 hr, 12 hr, 14 hr, and 16 hr during fermentation (Tang et al., 2007). All analyses were performed in triplicate.

2.3 | Transcriptomic analysis

The fermented soymilk was sampled at the fermentation time of 4 hr, 7 hr, and 10 hr corresponding to the lag phase, logarithmic phase, and stationary phase, respectively. Three parallel samples were obtained in each sampling point. The quality and integrity of total RNA were assessed by 1% agarose gels and RNA 6000 Nano Assay Kit of the Bioanalyzer 2100 system. Probes were used to purify mRNA from the total RNA of prokaryotic samples. Fragmentation was carried out using divalent cations under hyperthermal temperature in first strand synthesis reaction buffer (5X). Synthesis of first strand complementary DNA (cDNA) was performed with random hexamer primer and...
Moloney murine leukemia virus (M-MuLV) reverse transcriptase. The second strand was synthesized by DNA Polymerase I and M-MuLV reverse transcriptase. The 3’ ends of DNA fragments were adenylated and then ligated to the adaptor with hairpin loop structure for hybridization. The cDNA library fragments with 350–400 bp were selected and purified with AMPure XP system. Polymerase chain reaction (PCR) was carried out with Phusion High-Fidelity DNA polymerase and the PCR products were purified with AMPure XP system. Finally, library quality was evaluated with Agilent 2100 Bioanalyzer system (Cheng et al., 2019). Gene descriptions and annotations were performed in the Genome Database of L. amylolyticus strain L6 in National Center for Biotechnology Information (NCBI) (https://www.ncbi.nlm.nih.gov) with GenBank Accession Number of CP020457.1. The annotated genes were then used to predict biochemical pathways. Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways and gene ontology (GO) terms were retrieved from the KEGG database (http://www.kegg.jp/kegg) and gene ontology (GO) database (http://geneontology.org), respectively.

Real-time quantitative PCR (RT–qPCR) was applied to verify the accuracy of transcriptomic results. Primers were designed and synthesized according to gene sequences on NCBI (Table S1). The gene-expression levels were calculated via the 2ΔΔCt method (Xu et al., 2015), which was used to compare with the sequencing results of the transcriptome.

2.4 Sugars and organic acid

The determination of sugars, including sucrose, glucose, fructose, galactose, raffinose, and stachyose in fermented soymilk, was performed by high-performance liquid chromatography (HPLC) according to the National Standard of China GB/T 22,221–2008. The samples with a volume of 5 ml were collected at 0 hr, 4 hr (lag phase), 7 hr (log phase), and 10 hr (stationary phase) and then centrifuged at 10,000 r/min for 10 min at 4°C. The supernatant (4 ml) was transferred to a 10-mL volumetric flask, diluted with methanol to the constant volume, and extracted with sonication for 1 hr. The resulting extracts were filtered through a 0.22-µm membrane into HPLC vials for HPLC testing.

The content of amino acid in fermented soymilk was determined by HPLC according to Agilent AdvanceBio AAA method (www.agilent.com/chem/advancedbioaa). Briefly, the pretreated samples were derivatized with o-phthalaldehyde (OPA), and the specific operations were carried out according to the method provided by Agilent. Analysis of amino acid by HPLC was carried out on Agilent 1100 equipped with an Agilent AdvanceBio AAA amino acid column (4.6 mm×250 mm, 5 µm) under isocratic elution. Na2HPO4 with a concentration of 0.01 mol/L and acetonitrile–methanol solution (acetonitrile:methanol:water 45:45:10) were used as mobile phase A and mobile phase B, respectively, with a flow rate of 1.5 ml/min. The detection wavelength was 338 nm, and the column temperature was set at 35°C. The content of sucrose, glucose, fructose, galactose, raffinose, and stachyose (Sigma Chemical Co., Ltd, Guangzhou, China) was dissolved in deionized water and transferred to a 10-mL volumetric flask for gradient dilution. The equation parameters of standard curve were used to determine the concentration of organic acid in fermented soymilk.

2.5 Analysis of isoflavones and amino acids by HPLC

The content of lactic acid and acetic acid in fermented soymilk was detected by HPLC according to GB/T 5009.157–2003. The samples with a volume of 5 ml were collected at 0 hr, 4 hr (lag phase), 7 hr (log phase), and 10 hr (stationary phase) and then centrifuged at 10,000 r/min for 10 min. The supernatant was filtered through a 0.22-µm syringe membrane into HPLC vials for testing. HPLC was performed on Agilent 1100 equipped with Luna C18(2) 100A column (4.6 mm×250 mm, 5 µm) and VWD 3100 ultraviolet detector. KH2PO4 (95%) with a concentration of 10 mmol/L-methanol (5%) was used as mobile phase with a flow rate of 0.5 ml/min. The detection wavelength was 210 nm, and the column temperature was set at 25°C. The standard of lactic acid and acetic acid (Sigma Chemical Co., Ltd, Guangzhou, China) was dissolved in deionized water and transferred to a 10-mL volumetric flask for gradient dilution. The equation parameters of standard curve were used to determine the concentration of organic acid in fermented soymilk.

2.6 Statistical analysis

Analyses were performed using SPSS (SPSS Inc., Chicago, IL, USA, V23.0.0). One-way analysis of variance (ANOVA) was performed
That might be because tofu-whey isolated L. amylolyticus L6 were identified. There were 260 SDEGs in logarithmic phase versus a total of 313 significantly differentially expressed genes (SDEGs) transcriptomic analysis of different growth phases next to each other. (RNA-seq). Our research mainly focused on the comparative transcriptomic analysis between L. amylolyticus L6 and Lactobacillus casei Zhang grown from the lag phase into the logarithmic phase at a time of 3 hr and reached stationary phase at 14 hr with a cell concentration of 10^9 CFU/mL. It was reported that Lactobacillus casei Zhang grew from the lag phase into the logarithmic phase at a time of 3 hr and reached stationary phase at 14 hr with a cell concentration of 10^12 CFU/mL (Wang et al., 2012). L. amylolyticus L6 need less time than L. casei Zhang to grow into stationary phase, while L6 could produce more bacterial cells in soymilk than L. casei Zhang. That might be because tofu-whey isolated L. amylolyticus L6 is more adaptable in the soymilk-based ecosystem than koumiss-isolated L. casei Zhang (Fei, Liu, et al., 2017).

### RESULTS AND DISCUSSION

#### 3.1 Growth characteristics of L. amylolyticus L6 in soymilk

The growth curve of L. amylolyticus L6 in soymilk was plotted according to viable counts (Figure 1). L. amylolyticus L6 started to grow after 2 hr inoculation in soymilk. It needed approximately 4 hr for bacteria to grow from lag phase into the logarithmic phase. Bacteria grew into the stationary phase at the time of 10 hr with a cell concentration of 10^12 CFU/mL. It was reported that Lactobacillus casei Zhang grew from the lag phase into the logarithmic phase at a time of 3 hr and reached stationary phase at 14 hr with a cell concentration of 10^9 CFU/mL (Wang et al., 2012). L. amylolyticus L6 need less time than L. casei Zhang to grow into stationary phase, while L6 could produce more bacterial cells in soymilk than L. casei Zhang. That might be because tofu-whey isolated L. amylolyticus L6 is more adaptable in the soymilk-based ecosystem than koumiss-isolated L. casei Zhang (Fei, Liu, et al., 2017).

#### 3.2 Gene-expression profiles of L. amylolyticus L6 during fermentation in soymilk

The gene-expression profiles of L. amylolyticus L6 during growth in the soymilk ecosystem were investigated by the RNA-sequencing (RNA-seq). Our research mainly focused on the comparative transcriptomic analysis of different growth phases next to each other. A total of 313 significantly differentially expressed genes (SDEGs) were identified. There were 260 SDEGs in logarithmic phase versus lag phase and 171 SDEGs identified in the stationary phase versus logarithmic phase (Figure 5). The SDEGs of L. amylolyticus L6 in logarithmic phase versus lag phase were functionally categorized, indicating that SDEGs mainly enriched in biological process (transmembrane transport, oxidation and reduction, and translation), cellular component (ribosome and membrane), and molecular function (structural constituent of ribosome, nucleotide binding, and catalytic activity) (Figure 2a). In the stationary phase compared to the logarithmic phase, SDEGs mainly enriched in molecular function, such as structural constituent of ribosome, nucleotide binding, and catalytic activity (Figure 2b). The pathway enrichment for different growth phases is shown in Figure 3 according to the KEGG pathway database. As for logarithmic phase versus Lag phase group, the pathway of SDEGs enriched in starch and sucrose metabolism, fatty acid degradation, ribosome, biosynthesis of secondary metabolites, and folate biosynthesis. The pathway of SDEGs in stationary versus logarithmic phase group enriched in ribosome, starch, and sucrose metabolism, adenosine triphosphate (ATP)-binding cassette (ABC) transporter, phosphotransferase system (PTS), and amino acid metabolism.

In order to determine the reliability of transcriptomic results, expression changes of nine target genes (B1704_03855, B1704_02440, B1704_01765, B1704_01760, B1704_00925, B1704_00910, and B1704_06165) between lag phase and stationary phase were selected for detection. The result indicated a high consistency between platform of RNA-seq and real-time quantitative polymerase chain reaction (RT-qPCR), proving the validity of RNA-seq data (Figure 4).

#### 3.3 Carbon metabolism of L. amylolyticus L6

It has been reported that Lactobacillus amylolyticus could metabolize various carbohydrates such as dextrin, fructose, galactose, glucose, maltose, mannose, sucrose, melibiose, and raffinose, and in some cases salicine, esculin, amygdalin, and starch (Bohak et al., 1998; Fei et al., 2018). Soymilk mainly contained four different kinds of sugars, including glucose, sucrose, raffinose, and stachyose (Table 1). To provide a guide for industrial applications of L. amylolyticus L6 in fermenting soymilk, 1.5% (w/v) of glucose was added to provide enough carbon source for the growth of L6. The metabolism of carbohydrate to produce organic acid by L. amylolyticus L6 during its fermentation in soymilk is shown in Figure 5. The results indicated that four kinds of sugars reduced significantly (p < .05) during the fermentation and the main carbon sources used for the growth of L. amylolyticus L6 were sucrose and glucose (Table 1 and Figure 5). Many genes related to glucose metabolism were significantly up-regulated in logarithmic phase, such as genes coding for PTS β-glucoside transporter (B1745_01765), 6-phospho-alpha-glucosidase (B1745_05130), gluconate kinase (B1745_01565), and glucose-6-phosphate dehydrogenase (B1745_01805) (Table 2). However, several genes involved in sucrose transportation, especially sugar ABC transporters (B1745_06760, B1745_06745, and B1745_06750), and galactose metabolism (B1745_05485 and B1745_05490) were
significantly down-regulated in logarithmic phase. Microbes intend to utilize easily metabolizable carbohydrate and inhibit the metabolism of the other carbohydrate by down-regulating the expression of related genes (Luesink et al., 1998), the phenomenon of which, called carbon catabolite repression (CCR), has been widely found in lactic acid bacteria (LAB) (Görke & Stülke, 2008; Wang et al., 2012). In the stationary phase, few genes related to glucose metabolism were induced while many genes involved in sucrose (B1745_04485, B1745_04615, B1745_06775), raffinose, and stachyose utilization were found to be significantly up-regulated (Table 3). Among these sugars, only the content of galactose increased slightly (Table 1), which was due to the partial hydrolysis of raffinose and stachyose by α-galactosidase (B1745_RS08070) (Table 2). This phenomenon has also been reported in several researches of soybean products fermented by LAB (Battistini et al., 2018; Elghali et al., 2014; Xia et al., 2019). The production of energy for *L. amylolyticus* L6 is mainly through the Embden–Meyerhof–Parnas pathway (EMP). The gene (B1745_01805) of glucose 6-phosphate dehydrogenase that is the key regulatory enzyme of the Hexose Monophosphate Pathway (HMP) was highly expressed in the log phase, indicating that HMP was also indispensable in the glycometabolism of *L. amylolyticus* L6. Besides, two genes (B1745_05365 and B1745_06945) relevant to ATP production were also significantly up-regulated. During the fermentation, the acidity of soymilk increased significantly from 45.33° to 95.88° (Table 1). The acidity increment was mainly derived from lactic acid with its content increased from 2.62g/L to 4.65g/L (stationary phase) (p < .05). In addition, the content of acetic acid also increased slightly. Organic acid production was produced by *L. amylolyticus* L6 through the consumption of sugars in the soymilk (Wang et al., 2012; Xia et al., 2019). The production

**FIGURE 2** Significantly differentially expressed genes (SDGEs) between different growth phase based on gene ontology (GO) analysis. Logarithmic phase versus Lag phase (a), Stable phase versus Logarithmic phase(b); BP, CC, and FF refer to biological process, cellular component, and molecular function, respectively

**FIGURE 3** Scatter plot of Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway enrichment analysis for different growth phases. Rich factor is the ratio of the number of differentially expressed genes (DEGs) annotated to the Pathway Term to the number of genes annotated to the Pathway entry
of acetic acid indicated that this strain was a facultatively heterofermentative bacterium. After glucose is converted into pyruvate by glycolysis, it can generate lactic acid through the action of lactate dehydrogenase under anaerobic conditions (Figure 5). RNA-Seq results showed that the expression of LDH gene (B1745_03165) encoding L-lactate dehydrogenase was significantly increased in the log phase (Table 2). In addition, it was also observed that adhE (B1745_05695) encoding acetaldehyde hydrogenase related to acetic acid production was up-regulated in the log phase. Therefore, high expression of LDH and adhE genes promotes the production of lactic acid and acetic acid which are important for coagulating soymilk.

3.4 | Nitrogen metabolism and biosynthesis

Due to the lack of various biosynthetic pathways, especially amino acid synthesis pathways, LAB generally need various nutritional ingredients and therefore they are usually found in nutrient-rich environments, such as vegetables, meat, and milk (Fernández & Zúñiga, 2006). Amino acids as an important nitrogen resource for LAB played important roles in physiological functions such as intracellular pH maintenance, stress resistance, and energy generation (Lei et al., 2018; Slonczewski et al., 2009). As a consequence, the proteolytic enzyme system serves a key role for LAB to grow in protein-rich soymilk. A total of 17 kinds of amino acids were detected in the fermented soymilk, including seven kinds of essential amino acids (EAAs) and 10 kinds of nonessential amino acids (NEAAs) (Table 4). The content of EAAs decreased gradually along with the fermentation and reached 177.26 mg/L in the stationary phase. But the content of total amino acids and NEAAs increased significantly and reached the highest 668.38 mg/L and 187.40 mg/L in the logarithmic phase respectively. Although the content of total amino acids and NEAAs decreased slightly in the stationary phase, it was still higher than that of unfermented soymilk. The increase of free amino acid content in the soymilk fermented by different lactobacilli and their mixes has been widely

### Table 1: Changes of sugar and organic acid in soymilk fermented with L. amylophilus L6

|                          | Unfermented | Lag phase | Logarithmic phase | Stationary phase |
|--------------------------|-------------|-----------|-------------------|------------------|
| Sugars (g/L)             |             |           |                   |                  |
| Sucrose                  | 3.75 ± 0.09a| 3.36 ± 0.09b| 2.89 ± 0.10c | 2.61 ± 0.05c    |
| Glucose                  | 8.01 ± 0.15a| 7.65 ± 0.12b| 7.36 ± 0.10b | 7.06 ± 0.14c    |
| Fructose                 | 0.13 ± 0.01a| 0.11 ± 0.00b| 0.09 ± 0.00b | 0.09 ± 0.00b    |
| Raffinose                | 0.59 ± 0.00a| 0.59 ± 0.01b| 0.57 ± 0.01b | 0.53 ± 0.05c    |
| Stachyose                | 2.89 ± 0.02a| 2.88 ± 0.02a| 2.88 ± 0.01a | 2.76 ± 0.01b    |
| Galactose                | 0.23 ± 0.01c| 0.35 ± 0.01b| 0.41 ± 0.01a | 0.44 ± 0.02a    |
| Organic acids (g/L)      |             |           |                   |                  |
| Lactic acid              | 2.62 ± 0.04d| 3.38 ± 0.04c| 4.12 ± 0.04b | 4.65 ± 0.06a    |
| Acetic acid              | 0.36 ± 0.02c| 0.43 ± 0.03b| 0.57 ± 0.04a | 0.62 ± 0.03c    |
| Acidity (°)              | 45.33 ± 1.14c| 47.64 ± 1.86c| 88.42 ± 2.31b | 95.88 ± 2.65a   |
| pH                       | 6.7 ± 0.1a  | 6.6 ± 0.1a  | 5.5 ± 0.2b    | 4.3 ± 0.2b      |

Note: Data are the mean ± standard deviation (n = 3). Means in the same column with different superscript letters (a-d) are significantly different (p < .05)
reported (Ceh et al., 2020; Song et al., 2008). Besides, the content of glutamate and arginine was higher than those of other amino acids in unfermented and fermented soymilk, accounting for approximately 40% content of total amino acid. This phenomenon has been found in soy powder yoghurt fermented by L. brevis WCP02 and L. plantarum P120 that content of arginine is the highest (reached 380 mg/g), and accounted for almost 50% content of the total amino acid in soy powder yoghurt (Ceh et al., 2020). Therefore, fermentation of soymilk by L. amylolyticus L6 could promote the hydrolysis of protein into amino acid, improving the nutritional quality and digestibility of soymilk.

The proteolytic system of LAB generally consisted of protease, transport systems of amino acid or peptides, and peptidases (Wang et al., 2012). The protein in soymilk was first hydrolyzed by protease into amino acids and peptides, which were then transported to cytoplasm by transport systems. Finally, the translocated peptides were degraded by peptidases (Savijoki et al., 2006). Transcriptomic results indicated that the expression of clpP (B1745_04695), clpC (B1745_01265), and clpE (B1745_04960) genes encoding subunits of caseinolytic protease (Clp) previously identified in Lactobacillus plantarum IIA-1AS (Mega et al., 2020) was highly induced in the logarithmic and stationary phase. Meanwhile, two genes (B1745_00550 and B1745_06165) coding for metalloprotease were also found to be up-regulated in the logarithmic and stationary phase. Besides, the gene of another intramembrane metalloprotease showed higher induction levels in the stationary phase than in the logarithmic phase. These highly expressed protease genes indicated the strong proteolytic ability of L. amylolyticus L6 in the fermentation of soymilk.

The transport of peptides into the cell is an essential step for LAB multiplying in soymilk (Hagting, 1995). Transcriptomic data showed that genes involved in the transport and hydrolysis of peptide in the cytoplasm also exhibited high expression levels (Tables 2 and 3). The gene cluster oppDFBCA encoding the oligopeptide transport system (Opp) and PepC encoding the aminopeptidase, which have been identified in an operon of Lactococcus lactis (Tynkkynen et al., 1993), were found to be up-regulated in soymilk-grown L. amylolyticus L6 cells. Five genes in Opp operon were highly expressed, including B1745_00920 (OppA coding for substrate-binding proteins), B1745_00925 and B1745_00940 (OppB and OppC coding for membrane proteins), and B1745_00945 (OppD and OppF coding for ATP-binding proteins). Meanwhile, pepC coding for aminopeptidase that could hydrolyze oligopeptide into amino acid also exhibited high expression levels. Additionally, another five peptidase genes were highly induced in the logarithmic phase, which include two peptidases (pepO, B1745_02860; pepT, B1745_06070), two aminopeptidases (pepC, B1745_00910 and B1745_00955), and a dipeptidyl-peptidase (pepX, B1745_02545). Dipeptidyl-peptidase (pepX, B1745_02545) identified in Lactobacillus helveticus (Ojennus et al., 2019) and Lactococcus lactis (Nurizzo et al., 2002) could hydrolyze peptide bonds from the N-terminus of substrates when the penultimate amino acid residue is a proline. The highly induced dipeptidyl-peptidase might suggest the high content of oligopeptides
| Function group and ORF | Gene              | Description                                                | Value of log₂ variable expression |
|------------------------|-------------------|------------------------------------------------------------|-----------------------------------|
| Genes up-regulated     |                   |                                                            |                                   |
| Carbohydrate transport and metabolism |                   |                                                            |                                   |
| B1745_05615           | nagZ              | beta-N-acetylhexosaminidase                                | 3.54                              |
| B1745_05130           | glvA              | 6-phospho-alpha-glucosidase                                | 3.26                              |
| B1745_03165           | ldh               | L-lactate dehydrogenase                                    | 3.18                              |
| B1745_05695           | adhE              | acetaldehyde dehydrogenase/alcohol dehydrogenase          | 2.67                              |
| B1745_RS08070          | galA              | alpha-galactosidase                                        | 2.32                              |
| B1745_01765           | scrA              | PTS beta-glucoside transporter subunit EII BCA             | 1.98                              |
| B1745_04615           | spp               | HAD family hydrolase                                       | 1.93                              |
| B1745_03820           | gyoR              | hypothetical protein                                        | 1.80                              |
| B1745_03825           | spp               | sugar-phosphatase                                          | 1.69                              |
| B1745_01805           | zwf               | glucose-6-phosphate dehydrogenase                          | 1.62                              |
| B1745_06170           | gpmB              | histidine phosphatase family protein                       | 1.58                              |
| B1745_07215           |                  | transcriptional regulator                                   | 1.44                              |
| B1745_02045           | docC              | D-alanyl-D-alanine carboxypeptidase                         | 1.44                              |
| B1745_02005           | bcrC              | phospholipid phosphatase                                   | 1.44                              |
| B1745_03140           | fumC              | class II fumarate hydratase                                | 1.43                              |
| B1745_04590           | murB              | UDP-N-acetylenolpyruvoylglucosamine reductase              | 1.38                              |
| B1745_04460           | glgC              | YebC/PmpR family DNA-binding transcriptional regulator    | 1.36                              |
| B1745_04525           | spp               | HAD family hydrolase                                       | 1.33                              |
| B1745_03145           | frdA              | flavocytochrome c                                          | 1.32                              |
| B1745_02465           | galE              | UDP-glucose-4-epimerase                                    | 1.30                              |
| B1745_07245           | poxL              | pyruvate oxidase                                           | 1.29                              |
| B1745_04860           | pgl               | 3-carboxymuconate cyclase                                  | 1.29                              |
| B1745_06310           | rpe               | ribulose phosphate epimerase                               | 1.22                              |
| B1745_00870           |                  | aldose 1-epimerase                                         | 1.19                              |
| B1745_07130           | bdhAB             | aldo/keto reductase                                        | 1.19                              |
| B1745_03235           | lysozyme          | lysis                                                      | 1.12                              |
| B1745_01565           | gntK, idnK        | gluconate kinase                                           | 1.09                              |
| B1745_04605           | pta               | phosphate acetyltransferase                                | 1.09                              |
| B1745_05160           | rpiA              | ribose 5-phosphate isomerase A                             | 1.03                              |
| B1745_04420           | ackA              | acetate kinase                                             | 1.00                              |
| B1745_03855           | bglA              | 6-phospho-beta-glucosidase                                 | 1.00                              |
| B1745_05365           | zntA              | copper-translocating P-type ATPase                         | 1.09                              |
| B1745_06945           |                  | cadmium-translocating P-type ATPase                       | 2.16                              |
| Amino acids transport and metabolism |                   |                                                            |                                   |
| B1745_06855           | asnA              | aspartate–ammonia ligase, asparagine biosynthetic process  | 4.16                              |
| B1745_02860           | pepO              | peptidase M13                                              | 2.44                              |
| B1745_00965           | hsp20             | heat-shock protein Hsp20                                   | 2.39                              |
| B1745_04695           | clpP              | ATP-dependent Clp protease proteolytic subunit             | 2.35                              |
| B1745_04960           | clpE              | Clp protease ClpE                                          | 2.25                              |
| B1745_05185           | glnP              | glutamine ABC transporter permease                         | 2.16                              |
| B1745_03105           | att               | amino acid permease                                        | 2.14                              |
| B1745_00925           | oppB              | peptide ABC transporter substrate-binding protein           | 2.07                              |
| Function group and ORF | Gene     | Description                                      | Value of log<sub>2</sub> variable expression |
|------------------------|----------|--------------------------------------------------|---------------------------------------------|
|                        | htx      | zinc metalloprotease Htx                          | 1.97                                        |
| B1745_00550            | pepT     | peptidase T                                      | 1.89                                        |
| B1745_09050            | oppF     | ABC transporter ATP-binding protein              | 1.83                                        |
| B1745_03200            | prmA     | ribosomal protein L11 methyltransferase          | 1.81                                        |
| B1745_00910            | pepC     | aminopeptidase                                   | 1.67                                        |
| B1745_01265            | clpC     | ATP-dependent Clp protease ATP-binding protein   | 1.63                                        |
| B1745_04890            | -        | transcriptional activator, Rgg/GadR/MutR family domain-containing protein | 1.60                                        |
|                        | glnM     | glutamine ABC transporter permease               | 1.60                                        |
|                        | pepX     | dipeptidyl-peptidase                             | 1.56                                        |
|                        | dacD     | D-alanyl-D-alanine carboxypeptidase              | 1.44                                        |
|                        | pepC     | aminopeptidase                                   | 1.44                                        |
|                        | oppD     | peptide ABC transporter ATP-binding protein      | 1.43                                        |
|                        | lysC     | aspartate kinase                                 | 1.41                                        |
|                        | att      | amino acid permease                              | 1.40                                        |
|                        | atpA     | haloacid dehalogenase                            | 1.40                                        |
|                        | pepC     | aminopeptidase                                   | 1.38                                        |
|                        | glnH     | glutamine ABC transporter substrate-binding protein | 1.32                                        |
|                        | groEL    | chaperonin GroEL                                 | 1.23                                        |
|                        | lysC     | metalloprotease                                  | 1.18                                        |
|                        | GlnP     | glutamine ABC transporter permease               | 1.08                                        |
|                        | oppA     | peptide ABC transporter substrate-binding protein | 1.07                                        |
|                        | cth      | aluminum resistance protein                      | 1.05                                        |
|                        | uvrB     | excinuclease ABC subunit B                       | 1.02                                        |
|                        | oppC     | peptide ABC transporter permease                 | 1.01                                        |
|                        | gadC     | glutamate:gamma-aminobutyrate antiporter        | 1.79                                        |

**Lipid metabolism, inorganic ion transport and stress response**

| Gene     | Description                                      | Value of log<sub>2</sub> variable expression |
|----------|--------------------------------------------------|---------------------------------------------|
| B1745_05970 | esterase                                        | 1.41                                        |
| B1745_00245 | esterase                                        | 1.03                                        |
| B1745_02830 | acyl-CoA thioesterase                           | 1.40                                        |
| B1745_05670 | biotin carboxylase                              | 1.58                                        |
| B1745_00100 | mgtC putative Mg<sup>2+</sup> transporter family protein | 1.26                                        |
| B1745_00845 | pot potassium transporter                       | 1.36                                        |
| B1745_05305 | amt ammonium transporter                        | 1.97                                        |
| B1745_06945 | cadmium-translocating P-type ATPase             | 2.16                                        |

**Genes down-regulated**

**Carbohydrate transport and metabolism**

| Gene     | Description                                      | Value of log<sub>2</sub> variable expression |
|----------|--------------------------------------------------|---------------------------------------------|
| B1745_06195 | hypothetical protein                            | -1.03                                       |
| B1745_05125 | glvB PTS alpha-glucoside transporter subunit IICB | -1.06                                       |
| B1745_05025 | atoB 3-ketoacyl-CoA thiolase                     | -1.08                                       |
| B1745_00695 | ddl D-alanine-D-alanine ligase A                 | -1.27                                       |
| B1745_05485 | tagA galactosyltransferase                       | -1.33                                       |
| B1745_05490 | Malk sugar ABC transporter ATP-binding protein   | -1.56                                       |
| B1745_06760 | acyP acylphosphatase                            | -1.50                                       |
with penultimate proline residue in the fermented soymilk. In the stationary phase, there were only two aminopeptidase genes (pepC, B1745_00910 and B1745_01515) that were significantly up-regulated, which might be due to the stagnation of cell growth and proliferation, reducing the requirement for peptide and amino acid.

Genome analysis of L. amylolyticus L6 with KEGG pathways revealed that this strain was able to synthesize nine kinds of amino acids, including valine (Val), leucine (Leu), isoleucine (Ile), phenylalanine (Phe), tryptophan (Trp), tyrosine (Tyr), aspartate (Asp), and arginine (Arg), alanine (Asn), and arginine (Arg). The overexpression of the glutamate transporter operon has also been reported in L. casei Zhang under soymilk environment (Wang et al., 2012). Meanwhile, many uncharacterized amino acid permease genes (B1745_03105, B1745_06875, B1745_06870, and B1745_06860) were up-regulated, while two amino acid permease genes (B1745_04680 and B1745_03815) were down-regulated in the logarithmic and stationary phase. Interestingly, two genes livB and brnQ coding for branched-chain amino acid transport system II carrier protein and branched-chain amino acid ABC transporter permease were significantly down-regulated in the stationary phase. That’s because L. amylolyticus L6 could synthesize branched-chain amino acids (leucine, isoleucine, and valine) and did not require the help of their transporters, therefore repressing the expression of corresponding genes.

### 3.5 Lipid metabolism, inorganic ion transport, and stress response

There are 14 genes involved in fatty acid biosynthesis which were identified in the genome of L. amylolyticus L6, which includes accA (acetyl-CoA (coenzyme A) carboxylase), accB (acetyl-CoA carboxylase biotin carboxyl carrier protein), accC (acyl-CoA carboxylase, biotin carboxylase subunit), accD (acyl-CoA carboxylase carboxyl transferase subunit beta), fabD (acyl-carrier-protein S-malonyltransferase), fabF (3-oxoacyl-[acyl-carrier-protein] synthase II), fabG (3-oxoacyl-acyl-carrier protein reductase), fabH (3-oxoacyl-3-oxoacyl-acyl-carrier protein synthase III), fabI (enoyl-[acyl-carrier protein] reductase I), fabZ (3-hydroxyacyl-[acyl-carrier-protein] dehydratase), and mch (medium-chain acyl-[acyl-carrier-protein] hydrolase). Only accC gene (B1745_05670) was found to be up-regulated in the logarithmic phase during the growth of L. amylolyticus L6 in soymilk. It was reported that soymilk contained 2.64% grease, and the relative

| Function group and ORF | Gene | Description | Value of log2 variable expression |
|------------------------|------|-------------|----------------------------------|
| B1745_06765 | pgmB | beta-phosphoglucosamide | -1.61 |
| B1745_06730 | gmpB | hypothetical protein | -2.38 |
| B1745_06745 | ganQ | sugar ABC transporter permease | -2.58 |
| B1745_06750 | ganP | sugar ABC transporter permease | -2.79 |

### Amino acids transport and metabolism

- **Table 2**

| Gene | Description |
|------|-------------|
| B1745_01435 | rpoA | DNA-directed RNA polymerase subunit alpha |
| B1745_02350 | yidC | insertase |
| B1745_04680 | - | amino acid permease |
| B1745_05775 | secG | preprotein translocase subunit SecG |
| B1745_06880 | lepB | S26 family signal peptidase |
| B1745_04160 | valS | valine-tRNA ligase |
| B1745_01405 | secY | preprotein translocase subunit SecY |
| B1745_02435 | cth | aluminum resistance protein |
| B1745_03815 | - | amino acid permease |

| Gene | Description |
|------|-------------|
| B1745_05195 | glnH | | -1.02 |
| B1745_05185 | glnP | | -1.07 |
| B1745_05775 | glnQ | | -1.14 |
| B1745_06740 | glnR | | -1.17 |
| B1745_06750 | glnS | | -1.19 |
| B1745_06760 | glnT | | -1.22 |
| B1745_06770 | glnU | | -1.38 |
| B1745_06780 | glnV | | -1.74 |
| B1745_06790 | glnW | | -2.08 |
| Function group and ORF                | Gene      | Description                                                                 | Expression ratio |
|--------------------------------------|-----------|-----------------------------------------------------------------------------|------------------|
| Genes up-regulated                   |           |                                                                             |                  |
| **Carbohydrate transport and metabolism** |           |                                                                             |                  |
| B1745_04550                          | glmS      | glutamine-fructose-6-phosphate transaminase (isomerizing)                    | 2.34             |
| B1745_04615                          | spp       | HAD family hydrolase                                                        | 1.91             |
| B1745_04860                          | pgl       | 3-carboxymuconate cyclase                                                   | 1.30             |
| B1745_06170                          | gpmB      | histidine phosphatase family protein                                        | 1.26             |
| B1745_04485                          | spp       | sugar-phosphatase                                                           | 1.20             |
| B1745_06775                          | nplT      | alpha-glycosidase                                                           | 1.14             |
| **Amino acid transport and metabolism** |           |                                                                             |                  |
| B1745_04550                          | glmS      | glutamine-fructose-6-phosphate transaminase (isomerizing)                    | 2.34             |
| B1745_00615                          | adaB      | cysteine methyltransferase                                                   | 2.00             |
| B1745_01515                          | pepC      | aminopeptidase                                                              | 1.55             |
| B1745_06860                          |           | amino acid permease                                                         | 1.54             |
| B1745_00965                          | hsp20     | heat-shock protein Hsp20                                                     | 1.52             |
| B1745_03805                          | uvrC      | excinuclease ABC subunit C                                                   | 1.45             |
| B1745_06885                          | clpE      | ATP-dependent Clp protease ATP-binding subunit                               | 1.40             |
| B1745_04960                          | clpE      | Clp protease ClpE                                                           | 1.33             |
| B1745_04860                          | pgl       | 3-carboxymuconate cyclase                                                   | 1.30             |
| B1745_01540                          | cysS      | cysteine-tRNA ligase                                                         | 1.29             |
| B1745_04890                          |           | transcriptional activator, Rgg/GadR/MutR family domain-containing protein    | 1.23             |
| B1745_03010                          | grpE      | nucleotide exchange factor GrpE                                             | 1.23             |
| B1745_00550                          | htxP      | zinc metalloprotease HtxP                                                    | 1.19             |
| B1745_04855                          | atpA      | haloacid dehalogenase                                                       | 1.18             |
| B1745_01260                          |           | histidine kinase                                                            | 1.16             |
| B1745_00985                          |           | CPBP family intramembrane metalloprotease                                   | 1.15             |
| B1745_04695                          | clpP      | ATP-dependent Clp protease proteolytic subunit                               | 1.14             |
| B1745_07110                          | pcp       | pyroglutamyl-peptidase I                                                    | 1.06             |
| B1745_06165                          |           | metalloprotease                                                             | 1.05             |
| B1745_00910                          | pepC      | aminopeptidase                                                              | 1.03             |
| **Lipid metabolism, inorganic ion transport, and stress response** |           |                                                                             |                  |
| B1745_02830                          |           | acyl-CoA thioesterase                                                       | 2.30             |
| B1745_01775                          | groEL     | chaperone                                                                    | 1.22             |
| B1745_03010                          | grpE      | nucleotide exchange factor GrpE                                             | 1.23             |
| B1745_03015                          | dnaK      | molecular chaperone                                                         | 1.10             |
| B1745_00745                          | uspA      | universal stress protein                                                     | 1.14             |
| B1745_01850                          | trxA      | thioredoxin                                                                  | 1.20             |
| **Genes down-regulated**             |           |                                                                             |                  |
| **Carbohydrate transport and metabolism** |           |                                                                             |                  |
| B1745_01165                          | murF      | UDP-N-acetyl-D-muramoyl-tripeptide--D-alanyl-D-alanine ligase                 | -1.02            |
| B1745_05730                          | manY      | PTS mannose/fructose/sorbose transporter subunit IIC                         | -1.04            |
| B1745_05730                          | manY      | PTS mannose/fructose/sorbose transporter subunit IIC                         | -1.04            |
| B1745_05735                          | manX      | PTS mannose transporter subunit IIAB                                         | -1.15            |
| B1745_05130                          | glvA      | 6-phospho-alpha-glucosidase                                                  | -1.33            |
| B1745_07135                          | nagB      | glucosamine-6-phosphate deaminase                                           | -2.70            |
content of unsaturated fatty acid is more than 80% (Xiangnan et al., 2019), which inhibited the expression of genes related to fatty acid biosynthesis in *L. amylolyticus* L6. Meanwhile, acyl-CoA thioesterase gene (B1745_02830) that catalyzes the hydrolysis of acyl-CoAs to the free fatty acid and regulates intracellular levels of free fatty acids and acyl-CoAs (Tillander et al., 2017) was highly induced during its growth in soymilk. Besides, several genes coding for esterase (B1745_05970 and B1745_00245) were also up-regulated in logarithmic phase to utilize the grease in soymilk.

Inorganic ions, especially metal ions, are important for LAB to maintain normal functions in the metabolism (Mrvčić et al., 2012). Generally, membrane transporters play a crucial role in regulating

| Function group and ORF | Gene | Description | Expression ratio |
|------------------------|------|-------------|------------------|
| Amino acid transport and metabolism | B1745_04680 | - amino acid permease | -1.01 |
| | B1745_05320 | livB branched-chain amino acid transport system II carrier protein | -1.02 |
| | B1745_05735 | manX PTS mannose transporter subunit IIAB | -1.15 |
| | B1745_02915 | tsf translation elongation factor Ts | -1.17 |
| | B1745_03935 | - peptide-binding protein | -1.24 |
| | B1745_02235 | thrS threonine-tRNA ligase | -1.41 |
| | B1745_00785 | asnB asparagine synthase (glutamine-hydrolyzing) | -1.47 |
| | B1745_00270 | brnQ branched-chain amino acid ABC transporter permease | -1.69 |

**TABLE 3** (Continued)

| Function group and ORF | Gene | Description | Expression ratio |
|------------------------|------|-------------|------------------|
| Lipid metabolism, inorganic ion transport, and stress response | B1745_00715 | - acetylesterase | -1.10 |

**TABLE 4** Changes of amino acids in soymilk fermented with *L. amylolyticus* L6

| Free amino acids (mg/L) | Period | Unfermented | Lag phase | Log phase | Stationary phase |
|-------------------------|--------|-------------|-----------|-----------|------------------|
| Essential amino acids   |        |             |           |           |                  |
| Lysine                  |        | 39.68 ± 1.72 | 36.60 ± 1.21 | 35.13 ± 0.28 | 33.06 ± 0.99 |
| Phenylalanine           |        | 29.62 ± 0.79 | 30.39 ± 0.86 | 28.87 ± 2.03 | 26.79 ± 1.39 |
| Methionine              |        | 47.04 ± 1.50 | 43.05 ± 1.79 | 43.72 ± 1.09 | 42.30 ± 1.85 |
| Threonine               |        | 8.59 ± 0.21  | 9.31 ± 0.25  | 9.59 ± 0.25  | 9.01 ± 0.22 |
| Isoleucine              |        | 20.32 ± 1.33 | 21.28 ± 0.92 | 20.80 ± 0.53 | 19.57 ± 2.50 |
| Leucine                 |        | 50.91 ± 1.98 | 48.33 ± 1.76 | 46.65 ± 1.20 | 43.40 ± 2.11 |
| Valine                  |        | 5.74 ± 0.33  | 2.19 ± 0.22  | 2.63 ± 0.08  | 3.13 ± 0.08 |
| Total of EAA            |        | 201.90 ± 7.95 | 191.14 ± 8.18 | 187.40 ± 4.82 | 177.26 ± 9.56 |

| Nonessential amino acids |        |             |           |           |                  |
| Asparagine              |        | 21.70 ± 0.76 | 14.10 ± 1.14 | 10.64 ± 1.02 | 5.48 ± 0.52 |
| Glutamate               |        | 102.04 ± 3.21 | 133.14 ± 2.90 | 133.30 ± 3.15 | 129.07 ± 0.99 |
| Serine                  |        | 23.27 ± 0.82 | 26.16 ± 1.63 | 27.98 ± 0.92 | 27.53 ± 0.66 |
| Histidine               |        | 4.65 ± 0.41  | 7.03 ± 0.23  | 6.82 ± 0.13  | 6.15 ± 0.25 |
| Glycine                 |        | 48.10 ± 1.61 | 61.88 ± 1.93 | 76.37 ± 1.00 | 83.97 ± 1.12 |
| Arginine                |        | 125.25 ± 2.85 | 121.10 ± 2.56 | 120.45 ± 2.65 | 115.86 ± 1.73 |
| Alanine                 |        | 60.34 ± 1.87 | 51.74 ± 2.00 | 53.91 ± 1.00 | 52.86 ± 0.98 |
| Tyrosine                |        | 11.42 ± 0.19 | 16.97 ± 0.24 | 16.23 ± 0.51 | 15.09 ± 0.41 |
| Cysteine                |        | 0.79 ± 0.06 | 1.21 ± 0.04 | 1.92 ± 0.09 | 3.18 ± 0.05 |
| Proline                 |        | 32.66 ± 1.55 | 32.83 ± 1.84 | 33.35 ± 0.67 | 32.69 ± 1.05 |
| Total of NEAAs          |        | 430.23 ± 13.34 | 466.16 ± 14.53 | 480.99 ± 11.14 | 471.86 ± 7.77 |
| Total amino acids       |        | 632.12 ± 21.28 | 657.30 ± 22.70 | 668.38 ± 15.97 | 649.12 ± 17.33 |

Note: Data are the mean ± standard deviation (n = 3). Means in the same row with different superscript letters (a–d) are significantly different (p <.05).
the intracellular concentrations of metal ions (Boyaval, 1989). The expression of five genes, such as mgtC (B1745_00100) coding for Mg$^{2+}$ transporter, pot (B1745_00845) coding for potassium transporter, amt (B1745_05305) coding for ammonium transporter, and cadmium-translocating P-type ATPase gene (B1745_06945), was significantly induced in the logarithmic phase, indicating the importance of inorganic ions in regulating physiological functions of L. amylolyticus L6, such as ion homeostasis, coenzyme factor, and electron transport system.

During the fermentation, the pH values and acidity of soymilk in the stationary phase could reach 4.0 and 95.88°, respectively, which would induce the expression of genes in responding to acidity stress. Molecular chaperones have been regarded as a ubiquitous feature of cells, including LAB, in which these proteins cope with stress-induced denaturation of other proteins (Feder & Hofmann, 1999). Chaperone proteins GroL, DnaK, and GrpE participate actively in the response to stress conditions by preventing the aggregation of stress-denatured proteins (Lemos et al., 2007). Transcriptomic analysis indicated that the expression of genes groEL (B1745_01775), dnaK (B1745_03015), and grpE (B1745_03010) coding for chaperone proteins was highly up-regulated in the stationary phase, while these two genes were not significantly induced in the logarithmic phase. The difference was mainly due to a relatively higher pH value in logarithmic phase that is not enough to cause acid stress to L. amylolyticus L6 (Table 1). The increased expression level of a universal stress protein (B1745_00745) in the stationary phase that was required for resistance to DNA damage also engaged in acid tolerance of L. amylolyticus L6. In addition, the high transcript level of thioredoxin (trxA, B1745_01850) in the stationary phase that acts as an antioxidant by promoting the reduction of other proteins through the cysteine thiol–disulfide bond exchange was related to stress adaptation in L. amylolyticus L6. The gene highly expressed in logarithmic phase was glutamate:γ-aminobutyrate antiporter (gadC, B1745_00320) that exchanges the intracellular γ-aminobutyric acid (GABA) with extracellular Glu to expel protons in the cytoplasm (Dan et al., 2012).

## 3.6 | Change of isoflavones in fermented soymilk

Soymilk was rich in isoflavones in the form of isoflavone aglycones (10%) and their corresponding glucosidic conjugates (90%) (Rodriguez-Roque et al., 2013). Isoflavones’ glucosidic conjugates could be converted into highly bioactive aglycones by β-glucosidase in lacticobacilli (Tang et al., 2007; Wei et al., 2007; Xia et al., 2019). As shown in Table 5, most of the isoflavones in unfermented soymilk occurred in the form of glucosides with the concentration of 285.77 mg/L and the content of aglycones was only 14.51 mg/L. During the fermentation, the total concentration of isoflavone aglycones increased from 14.51 mg/L to 36.09 mg/L, and three forms of aglycones’ (daizein, glycine, and genistin) concentration also increased significantly. However, the content of glucosidic isoflavones changed irregularly during the fermentation. Compared with the unfermented phase, the glucosidic isoflavones (daizaid, glyctin, and genistin) exhibited a decreasing tendency in the lag phase (2h) and then the content of glucosidic isoflavones increased gradually in the logarithmic and stationary phase. A similar phenomenon has been reported in the soymilk beverage fermented by Kombucha rich in LAB (Xia et al., 2019). It is presumed that the fermentation of L. amylolyticus L6 could promote the release of free flavonoids from binding forms with soluble fibers in the soymilk. Transcriptomic data indicated that the expression of bg1A gene coding for 6-phospho-β-glucosidase increased significantly in logarithmic phase, which was consistent with the increasing concentrations of isoflavone aglycones. 6-phospho-β-glucosidase that could convert isoflavone glucosides into aglycones has been reported in our previous study (Fei, Liu, et al., 2017).

### 4 | CONCLUSION

This study revealed the chemical component changes and transcriptomic changes of L. amylolyticus L6 in fermented soymilk.

| Isoflavones (mg/L) | Period           | Unfermented | Lag phase  | Log phase | Stationary phase |
|-------------------|------------------|-------------|------------|-----------|------------------|
| **Glycosides**    |                  |             |            |           |                  |
| Daidzin           | 216.65 ± 3.80a   | 154.96 ± 1.92c | 188.45 ± 1.83b | 217.22 ± 2.01a |
| Glyctin           | 46.02 ± 1.77b    | 35.17 ± 0.47d | 43.60 ± 0.19c | 50.73 ± 0.29a  |
| Genistin          | 23.09 ± 0.12a    | 11.87 ± 0.63b | 12.10 ± 0.06b | 12.62 ± 0.77b  |
| **Total**         | 285.77 ± 5.50a   | 202.01 ± 1.80c | 244.15 ± 1.87b | 280.57 ± 1.41a |

| **Aglycones**     |                  |             |            |           |                  |
| Daizein           | 10.03 ± 0.49c    | 9.17 ± 0.48d | 11.89 ± 0.08b | 16.63 ± 0.14a |
| Glycine           | ND               | ND          | ND         | 5.61 ± 0.07a  |
| Genistein         | 4.48 ± 1.16b     | 5.65 ± 1.87b | 12.11 ± 0.99a | 13.85 ± 1.27a |
| **Total**         | 14.51 ± 1.65 c   | 14.82 ± 0.62 c | 24.00 ± 1.07 b | 36.09 ± 1.48 a |

**Note:** Data are the mean ± standard deviation (n = 3). Means in the same column with different superscript letters (a–d) are significantly different (p < .05). ND means not detected.

### Table 5 | Concentration of isoflavones (mg/L) in soymilk fermented with L. amylolyticus L6

| Isoflavones (mg/L) | Period         | Unfermented | Lag phase | Log phase | Stationary phase |
|-------------------|----------------|-------------|-----------|-----------|------------------|
| **Glycosides**    |                |             |            |           |                  |
| Daidzin           | 216.65 ± 3.80a | 154.96 ± 1.92c | 188.45 ± 1.83b | 217.22 ± 2.01a |
| Glyctin           | 46.02 ± 1.77b | 35.17 ± 0.47d | 43.60 ± 0.19c | 50.73 ± 0.29a  |
| Genistin          | 23.09 ± 0.12a | 11.87 ± 0.63b | 12.10 ± 0.06b | 12.62 ± 0.77b  |
| **Total**         | 285.77 ± 5.50a | 202.01 ± 1.80c | 244.15 ± 1.87b | 280.57 ± 1.41a |

| **Aglycones**     |                |             |            |           |                  |
| Daizein           | 10.03 ± 0.49c | 9.17 ± 0.48d | 11.89 ± 0.08b | 16.63 ± 0.14a |
| Glycine           | ND             | ND          | ND         | 5.61 ± 0.07a  |
| Genistein         | 4.48 ± 1.16b | 5.65 ± 1.87b | 12.11 ± 0.99a | 13.85 ± 1.27a |
| **Total**         | 14.51 ± 1.65 c | 14.82 ± 0.62 c | 24.00 ± 1.07 b | 36.09 ± 1.48 a |
Large amount of genes related to carbon metabolism in *L. amylolyticus* L6 were significantly up-regulated in the logarithmic phase and stationary phase, which allowed this strain to metabolize various sugars in soymilk. Highly expressed α-galactosidase gene could help to reduce the content of raffinose and stachyose that caused flatulence of human body. Meanwhile, the concentration of total amino acid increased significantly in the logarithmic phase for highly induced genes involved in the proteolysis, hydrolysis, and transport of peptide, transport and biosynthesis of amino acid. Highly efficient utilization of carbon and nitrogen sources significantly raised the viable counts of *L. amylolyticus* L6 in soymilk. High expression of β-glucosidase promoted the conversion of isoflavone glycoside into highly bioactive aglycones. Besides, other genes related to lipid metabolism, inorganic ion transport, and stress response were also up-regulated. Further study should be conducted in terms of applying this strain into developing soymilk products and vitro digestion simulation test to testify its production performance. In conclusion, this study reveals that *L. amylolyticus* L6 isolated from the soybean-derived environment exhibited excellent adaptability in a soymilk-based ecosystem, which is expected to become the specific probiotic strain used for the fermentation of soybean products.

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**CONFLICT OF INTEREST**

None declared.

**ETHICAL APPROVAL**

The authors declare that they have no conflict of interest. This article does not contain any studies involving animal’s trials performed by any of the authors. Furthermore, this article does not contain any studies involving human participants performed by any of the authors.

**DATA AVAILABILITY STATEMENT**

The original contributions presented in the study are included in the article/Supplementary Material, and further inquiries can be directed to the corresponding authors.

**ORCID**

Yongtao Fei © https://orcid.org/0000-0001-7175-4406

**REFERENCES**

Battistini, C., Gullon, B., Ichimura, E. S., Gomes, A. M. P., Ribeiro, E. P., Kunigk, L., Moreira, J. U. V., & Jurkiewicz, C. (2018). Development and characterization of an innovative sybiotic fermented beverage based on vegetable soybean. *Brazilian Journal of Microbiology*, 49, 303–309. https://doi.org/10.1016/j.bjm.2017.08.006

Bohak, I., Back, W., Richter, L., Ehrmann, M., Ludwig, W., & Schleifer, K. H. (1998). *Lactobacillus amylolyticus* sp. nov., isolated from beer malt and beer wort. *Systematic & Applied Microbiology*, 21, 360–366. https://doi.org/10.1078/0723-2020(98)00453-3

Boyaval, P. (1989). Lactic acid bacteria and metal ions. *Le Lait*, 69, 87–113. https://doi.org/10.1051/laith:198927

Bron, P. A., Van Baarlen, P., & Kleerebezem, M. (2012). Emerging molecular insights into the interaction between probiotics and the host intestinal mucosa. *Nature Reviews Microbiology*, 10, 66–78. https://doi.org/10.1038/nrmicro2690

Ceh, A., Su, C., Du, H., Hyl, A., Hks, B., Kmc, A., & Jin, H. (2020). Enhancement of isoflavone aglycone, amino acid, and CLA contents in fermented soybean yogurts using different strains: Screening of antioxidant and digestive enzyme inhibition properties. *Food Chemistry*, 340, 128–199. https://doi.org/10.1016/j.foodchem.2020.128199

Cheng, L., Zhang, X., Zheng, X., Wu, Z., & Weng, P. (2019). RNA-seq transcriptomic analysis of green tea polyphenols regulation of differently expressed genes in Saccharomyces cerevisiae under ethanol stress. *World Journal Microbiology Biotechnology*, 35, 59–69. https://doi.org/10.1007/s11274-019-2639-4

Donkor, O. N., Henriksen, A., Vasiljevic, T., & Shah, N. P. (2007). α-Galactosidase and proteolytic activities of selected probiotic and dairy cultures in fermented soymilk. *Food Chemistry*, 104, 10–20. https://doi.org/10.1016/j.foodchem.2006.10.065

Elghali, S., Mustafa, S., Amid, M., & Manap, M. Y. A. (2014). Variations in soy milk components during fermentation by Lactobacillus and Bifidobacterium strains. *Journal of Food, Agriculture & Environment*, 12, 1–5.

Feder, M. E., & Hofmann, G. E. (1999). Heat-shock proteins, molecular chaperones, and the stress response: Evolutionary and ecological physiology. *Annual Review of Physiology*, 61, 243–282. https://doi.org/10.1146/annurev.physiol.61.1.243

Fei, Y., Jiao, W., Wang, Y., Li, G., & Lu, L. (2020). Cloning and expression of a novel alpha-galactosidase from *Lactobacillus amylo-lyticus* L6 with hydrolytic and transgalactosyl properties. *PLoS One*, 15, e0235687. https://doi.org/10.1371/journal.pone.0235687

Fei, Y., Li, L., Chen, L., Zheng, Y., & Yu, B. (2018). High-throughput sequencing and culture-based approaches to analyze microbial diversity associated with chemical changes in naturally fermented tofu whey, a traditional Chinese tofu-coagulant. *Food Microbiology*, 76, 69–77. https://doi.org/10.1016/j.fm.2018.04.004

Fei, Y., Li, L., Zheng, Y., Liu, D., Zhou, Q., & Fu, L. (2017). Characterization of *Lactobacillus amylolyticus* L6 as potential probiotics based on genome sequence and corresponding phenotypes. *LWT - Food Science and Technology*, 90, 460–468. https://doi.org/10.1016/j.lwt.2017.12.028

Fei, Y., Liu, L., Liu, D., Chen, L., Tan, B., Fu, L., & Li, L. (2017). Investigation on the safety of *Lactobacillus amylolyticus* L6 and its fermentation properties of tofu whey. *LWT - Food Science and Technology*, 84, 314–322. https://doi.org/10.1016/j.lwt.2017.05.072

Fernández, M., & Zúñiga, M. (2006). Amino acid catabolic pathways of lactic acid bacteria. *CRC Critical Reviews in Microbiology*, 32, 155–183. https://doi.org/10.1080/10408410600880643

Gerritsen, J., Smidt, H., Rijkers, G. T., & Vos, W. (2011). Intestinal microbiota in human health and disease: The impact of probiotics. *Genes & Nutrition*, 6, 209–240. https://doi.org/10.1007/s12263-011-0229-7
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