Determination of the regulatory network and function of the lysR-type transcriptional regulator of Lactiplantibacillus plantarum, LpLttR

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Abstract
Background: Lactiplantibacillus plantarum has various healthcare functions including the regulation of immunity and inflammation, reduction of serum cholesterol levels, anti-tumor activity, and maintenance of the balance of intestinal flora. However, the underlying metabolic and regulatory mechanisms of these processes remain unclear. Our previous studies have shown that the LysR type transcriptional regulator of L. plantarum (LpLttR) regulates the biotransformation of conjugated linoleic acids (CLAs) through the transcriptional activation of cla-dh (coding gene for CLA short-chain dehydrogenase) and cla-dc (coding gene for CLA acetoacetate decarboxylase). However, the regulatory network and function of LpLttR have not yet been characterized in L. plantarum.

Results: In this study, the regulatory role of LpLttR in various cellular processes was assessed using transcriptome analysis. The deletion of LpLttR had no evident influence on the bacterial growth. The transcriptome data showed that the expression of nine genes were positively regulated by LpLttR, and the expression of only two genes were negatively regulated. Through binding motif analysis and molecular interaction, we demonstrated that the regulatory region of the directly regulated genes contained a highly conserved sequence, consisting of a 15-base long box and rich in AT.

Conclusion: This study revealed that LpLttR of L. plantarum did not play a global regulatory role similar to that of the other transcriptional regulators in this family. This study broadens our knowledge of LpLttR and provides a theoretical basis for the utilization of L. plantarum.

Keywords: Lactiplantibacillus plantarum, LysR type transcriptional regulator (LttR), Transcriptional regulation, Regulon

Background
The LysR type transcriptional regulatory factor (LttR) is a ubiquitous regulatory factor in prokaryotes. Many studies have shown that this family of regulators plays a regulatory role in many cellular processes including primary metabolism, secondary metabolism, stress response, cell division, toxicity, quorum sensing and protection, etc. [1, 2]. In Pseudomonas putida KT2440, CatR regulates the expression of catBCA, which is involved in the catechol gene cluster [3]. The CbnR in Ralstonia eutropha regulates the transcription of the catechol explanation related gene cluster cbnABCD [4]. The TfdR of R. eutropha JMP134 not only controls the expression of the o-phenylene diene gene cluster tfdDCEFB, but also regulates the transcription of tfdA to metabolize 2,4-dichlorophenoxyacetic acid isooctyl ester [5]. Moreover, LttRs regulate the genes involved in virulence, metabolism, quorum sensing and exercise [1]. As more target genes have been identified, LttRs have been identified as global regulators.

LttRs are structurally conserved, with most containing 276–324 amino acid residues. Their N-terminal is a
typical DNA binding domain named the HTH domain, and the C-terminal is the substrate or co-inducer binding domain, which is less conservative and is convenient for the recognition and binding of substrates [6]. LttRs are often induced or suppressed by small molecules, and usually form dimers or tetramers to activate or inhibit the expression of target genes [1].

*L. plantarum* is a common lactic acid bacteria (LAB) used in the production of several functional and fermented foods [7]. However, there have been few studies on its metabolism and regulatory mechanisms. Our previous studies have shown that *L. plantarum* ATCC BAA-793 can convert linoleic acid (LA) to conjugated linoleic acid (CLA), which plays an important role in reducing weight and regulating immunity. The LysR family transcriptional regulator, *Lp*LttR, activates the transcription of *cla-dh* and *cla-dc* during this process and promotes the biosynthesis of CLA [8]. In addition to *cla-dh* and *cla-dc*, it is still unknown which genes are regulated by *Lp*LttR and the biological processes in which it participates. With continuous study of prokaryotic transcriptional regulation, more target genes of LttRs have been identified. However, the regulatory mechanisms in different species remain to be clarified, especially in *L. plantarum*, which has a wide range of application prospects in the food industry.

Here, we studied the regulon of *Lp*LttR of *L. plantarum* ATCC BAA-793. The sequence analysis of *Lp*LttR was performed to predict its function. We then performed transcriptome sequencing of the WT and *Lp*lttR knockout mutant strains (Δ*Lp*lttR) to identify the differentially expressed genes. Moreover, the presumed binding sites of the target genes were predicted and verified using molecular interactions based on interferometric techniques. The findings of this study revealed the regulatory network of *Lp*LttR in *L. plantarum* and provided new insights into the functions of LttRs.

**Results**

*Lp*LttR conservative analysis

The primary structure of *Lp*LttR in *L. plantarum* ATCC BAA-793 was analyzed in this study. It was encoded by *LP_RS00230* and composed of 295 amino acids. Pfam domain analysis showed that *Lp*LttR contained an HTH domain and a LysR substrate binding domain at the C and N terminals at residues 3–64 and 85–293, respectively (Fig. 1A). The amino acid sequence of the HTH domain of LttRs was aligned with other LttRs which have been extensively studied and reported as CatR in *P. putida* KT2440 [4], ClcR in *P. putida* [9], TcbR in *Pseudomonas* sp. strain P51 [10], CbnR in *Ralstonia eutropha* [11], TfdR in *R. eutropha* JMP134 [12], CatR in *P. putida* PaW85 [13],

![Lp*LttR conservative analysis. A](image)

Fig. 1 *Lp*LttR conservative analysis. A The illustration of the domain organization of *Lp*LttR. *Lp*LttR was composed of two domains: one HTH domain (position 3–64) at the N-terminal and one LysR substrate-binding domain (position 85–293) at the C-terminal. B Sequence conservation analysis of the HTH domain of LttRs including CatR (*Pseudomonas putida* KT2440), ClcR (*P. putida*), CbnR (*R. eutropha*), TfdR (*R. eutropha* JMP134), CatR (*P. putida*), CatM (*Acinetobacter*), PcaQ (*Agrobacterium*), BenM (*Acinetobacter*), TfdT (*Burkholderia*), NtdR (*Acidovorax*), LinR (*Sphingomonas*), and *Lp*LttR (*Lactobacillus plantarum*)
CatR in P. putida PRS1 [4], TfdR in Ralstonia [12], TfdS in Ralstonia [5], CatM in Acinetobacter [14], PcaQ in Agrobacterium [15], BenM in Acinetobacter [16], TfdT in Burkholderia [12], NtdR in Acidovorax [17], and LinR in Sphingomonas [18]. The sequence alignment showed a high sequence conservation between the HTH domain of LpLttR and other LttRs, especially for the first 50 amino acid residues (Fig. 1B). This suggests that LpLttR might also directly bind to DNA regulatory regions and play a regulatory role in a variety of cellular processes, similar to other LttRs.

**LpLttR knockout strain construction**

To study the function of LpLttR, we constructed the knockout strain (ΔLplttR) based on CRISPR-Cas9 gene editing technology. As shown in Fig. 2A, the upstream and downstream 1000 bp of LplttR were selected as the upstream and downstream homologous arms (Ha-1 and Ha-2), respectively. The homologous arms and sgRNA were inserted into the pLCNICK knockout plasmid. Verification primers were designed on both sides of the homologous arms to verify the transformers using colony PCR. As shown in Fig. 2B, the amplified product of the putative knockout strain (ΔLplttR) was approximately 1000 bp smaller than that of the WT, suggesting that the LplttR gene was deleted successfully. To study the effect of LplttR on bacterial growth, the growth curves of WT and ΔLplttR strains were investigated. As shown in Fig. 2C, during the first 24 h, the ΔLplttR strain grew slower than the WT but not thereafter.

**Identification of the LpLttR regulon**

As the main family of transcriptional regulators in prokaryotes, LttRs regulate a variety of genes and even play a global regulatory role in P. aeruginosa, S. thermophilus, and Salmonella enterica [19–21]. To identify the regulons of LpLttR in L. plantarum, we analyzed the mRNA expression levels of WT and ΔLplttR. As shown in Fig. 3A, in ΔLplttR, there were 11 mRNAs with a fold difference of more than two-fold, of which the expression of nine genes were down-regulated. Among the differentially expressed genes, LP_RS15475 and LP_RS15480 expressions were down regulated most substantially (Fig. 3B), and were described as replication proteins and hypothetical proteins, respectively. The expression of LP_RS14775 and LP_RS14610, separately annotated as

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**Fig. 2** Construction of the LplttR knockout strain and its effects on bacterial growth. **A** Schematic diagram of the LplttR knockout principle. Ha-1 and Ha-2 represent the upstream and downstream homologous arms respectively. The black arrows indicate the location of colony PCR primers used for verification. **B** Identification of LplttR knockout by colony PCR, **C** Effect of the LpLttR mutation on bacterial growth. Error bars indicate the standard deviations of three independent experiments.
SLC45 family MFS transporter and IS1182 family transposase, were upregulated. The transcriptional levels of these 11 genes were analyzed using RT-qPCR. Although the genes differed with respect to fold-change, the trend of up-regulation and down-regulation was similar, thus, confirming the reliability of the transcriptome data. All of the differentially expressed genes and their descriptions are listed in Table 1. Base on the number and function of the differential genes, we inferred that unlike other LttRs, LpLttR did not play a global regulatory role in L. plantarum.

To further analyze the regulatory role of LpLttR in L. plantarum, we also analyzed the differentially expressed genes with a change of more than 1.5-fold. As shown in Table 2, there were 70 genes with a fold difference of more than 1.5-fold, with a P-value < 0.05. LpLttR mainly acted as a transcriptional activator, since 49 of the 70 differential genes were down-regulated, whereas

| Gene name     | Gene description                          | FC(ΔLttR/WT) |
|---------------|------------------------------------------|--------------|
| LP_RS00230    | LysR family transcriptional regulator    | 0.291        |
| LP_RS15475    | replication protein                      | 0            |
| LP_RS15480    | hypothetical protein                     | 0            |
| LP_RS14775    | SLC45 family MFS transporter             | 2.198        |
| LP_RS14610    | IS1182 family transposase                | 100.747      |
| LP_RS13240    | cell wall hydrolase/muramidase           | 0.46         |
| LP_RS11700    | MFS transporter                          | 0.433        |
| LP_RS13245    | SH3 domain-containing protein            | 0.47         |
| LP_RS07440    | ABC transporter permease                 | 0.44         |
| LP_RS12305    | LysM peptidoglycan-binding domain-containing protein | 0.423 |
| LP_RS14610    | APC family permease                      | 0.436        |

Fig. 3 Analysis of the target genes with more than twofold difference in expression. A Scatter plot of the differentially expressed genes. The X-axis and Y-axis represent the expression of genes in the WT and LpLttR knockout strains. The values are all log-transformed. Each dot in the image represents a specific gene. The red dots indicate the up-regulated genes and green dots indicate the down-regulated genes. B Heatmap of the differentially expressed genes. Each column represents different strains and each row represents a gene. The depth of the color in the heatmap indicates the amount of the gene expression. For the specific amount of the expression, please see the digital label on the color bar.
| Gene name   | Gene description                                      | Type     | Regulate | FC(ΔlttR/WT) |
|------------|-------------------------------------------------------|----------|----------|--------------|
| LP_RS14610 | IS1182 family transposase                            | mRNA     | Up       | 100.747      |
| LP_RS14775 | SLC45 family MFS transporter                         | mRNA     | Up       | 2.198        |
| LP_RS15415 | DUF916 and DUF3324 domain-containing protein          | mRNA     | Up       | 1.935        |
| LP_RS01010 | DUF871 domain-containing protein                      | mRNA     | Up       | 1.931        |
| LP_RS11220 | GntR family transcriptional regulator                 | mRNA     | Up       | 1.702        |
| LP_RS12660 | Glycoside hydrolase family 1 protein                  | mRNA     | Up       | 1.7          |
| LP_RS01000 | PTS sugar transporter subunit IIC                    | mRNA     | Up       | 1.696        |
| LP_RS13760 | Hemolysin III family protein                         | mRNA     | Up       | 1.613        |
| treR       | Trehalose operon repressor                           | mRNA     | Up       | 1.611        |
| LP_RS01020 | DgaE family pyridoxal phosphate-dependent ammonia lyase| mRNA     | Up       | 1.572        |
| LP_RS02110 | Serine transporter                                    | mRNA     | Up       | 1.57         |
| spx        | Transcriptional regulator Spx                         | mRNA     | Up       | 1.541        |
| LP_RS10520 | Hypothetical protein                                  | mRNA     | Up       | 1.535        |
| LP_RS14565 | Galactose mutarotase                                  | mRNA     | Up       | 1.532        |
| LP_RS05750 | Aspartate-semialdehyde dehydrogenase                  | mRNA     | Up       | 1.5          |
| LP_RS06040 | Hypothetical protein                                  | mRNA     | Up       | 1.512        |
| LP_RS15420 | WxL domain-containing protein                         | mRNA     | Up       | 1.512        |
| LP_RS11200 | PTS sugar transporter subunit IIA                    | mRNA     | Up       | 1.506        |
| LP_RS12525 | DUF916 and DUF3324 domain-containing protein          | mRNA     | Up       | 1.503        |
| LP_RS05765 | NADH-dependent flavin oxidoreductase                  | mRNA     | Up       | 1.503        |
| LP_RS00610 | Helix-turn-helix transcriptional regulator             | mRNA     | Up       | 1.5          |
| LP_RS05365 | Peptide ABC transporter substrate-binding protein      | mRNA     | Down     | 0.668        |
| LP_RS12020 | DufM peptidoglycan-binding domain-containing protein  | mRNA     | Down     | 0.668        |
| LP_RS13890 | HIT family protein                                    | mRNA     | Down     | 0.663        |
| LP_RS01675 | AraC family transcriptional regulator                 | mRNA     | Down     | 0.659        |
| LP_RS00140 | Hypothetical protein                                  | mRNA     | Down     | 0.656        |
| LP_RS00290 | ASCH domain-containing protein                        | mRNA     | Down     | 0.655        |
| LP_RS06330 | Response regulator transcription factor               | mRNA     | Down     | 0.654        |
| LP_RS13055 | ABC transporter ATP-binding protein                   | mRNA     | Down     | 0.646        |
| LP_RS06150 | DUF916 and DUF3324 domain-containing protein          | mRNA     | Down     | 0.643        |
| LP_RS06145 | LPXTG cell wall anchoring protein                     | mRNA     | Down     | 0.643        |
| LP_RS05380 | ABC transporter ATP-binding protein                   | mRNA     | Down     | 0.642        |
| mvk        | Mevalonate kinase                                     | mRNA     | Down     | 0.642        |
| LP_RS07140 | ACP S-malonyltransferase                              | mRNA     | Down     | 0.637        |
| LP_RS07130 | Ketolactyl-ACP synthase III                           | mRNA     | Down     | 0.636        |
| LP_RS05375 | ABC transporter permease                              | mRNA     | Down     | 0.633        |
Table 2 (continued)

| Gene name     | Gene description                                      | Type         | Regulate | FC(ΔlttR/WT) |
|---------------|-------------------------------------------------------|--------------|----------|--------------|
| LP_RS11365    | LysR family transcriptional regulator                | mRNA         | Down     | 0.629        |
| LP_RS07625    | GIY-YIG nuclease family protein                       | mRNA         | Down     | 0.625        |
| fabZ          | 3-hydroxyacyl-ACP dehydratase FabZ                    | mRNA         | Down     | 0.623        |
| LP_RS05800    | Membrane protein                                     | mRNA         | Down     | 0.619        |
| LP_RS12295    | Helix-turn-helix transcriptional regulator            | mRNA         | Down     | 0.615        |
| LP_RS04070    | MucBP domain-containing protein                       | mRNA         | Down     | 0.611        |
| LP_RS04675    | AraC family transcriptional regulator                | mRNA         | Down     | 0.609        |
| LP_RS08765    | Hypothetical protein                                 | mRNA         | Down     | 0.609        |
| LP_RS13480    | Transporter substrate-binding domain-containing protein | mRNA       | Down     | 0.606        |
| LP_RS14300    | C40 family peptidase                                 | mRNA         | Down     | 0.603        |
| LP_RS07350    | Ribonuclease Hf family protein                        | mRNA         | Down     | 0.601        |
| LP_RS12415    | Zinc ribbon domain-containing protein                | mRNA         | Down     | 0.597        |
| LP_RS12410    | Zinc ribbon domain-containing protein                | mRNA         | Down     | 0.59         |
| LP_RS11005    | Virophilin family protein                            | mRNA         | Down     | 0.589        |
| LP_RS11130    | Hypothetical protein                                 | mRNA         | Down     | 0.583        |
| LP_RS11350    | Aminotransferase class I/II-fold pyridoxal phosphate-dependent enzyme | mRNA       | Down     | 0.576        |
| LP_RS12030    | Methylated-DNA-[protein]-cysteine S-methyltransferase | mRNA         | Down     | 0.571        |
| LP_RS15375    | 2-keto-4-pentenoate hydratase                        | mRNA         | Down     | 0.558        |
| LP_RS02570    | AEC family transporter                               | mRNA         | Down     | 0.544        |
| LP_RS08575    | ISL3 family transposase                              | mRNA         | Down     | 0.532        |
| LP_RS13185    | YxeA family protein                                  | mRNA         | Down     | 0.528        |
| LP_RS13235    | Ldh family oxidoreductase                            | mRNA         | Down     | 0.524        |
| LP_RS12675    | LysM peptidoglycan-binding domain-containing protein | mRNA         | Down     | 0.521        |
| LP_RS08270    | GNAT family N-acetyltransferase                      | mRNA         | Down     | 0.511        |
| LP_RS00810    | Peptide ABC transporter substrate-binding protein     | mRNA         | Down     | 0.51         |
| LP_RS13245    | SH3 domain-containing protein                        | mRNA         | Down     | 0.47         |
| LP_RS13240    | Cell wall hydrolase/muramidase                       | mRNA         | Down     | 0.46         |
| LP_RS11700    | MFS transporter                                      | mRNA         | Down     | 0.443        |
| LP_RS07440    | ABC transporter permease                             | mRNA         | Down     | 0.44         |
| LP_RS12305    | APC family permease                                  | mRNA         | Down     | 0.436        |
| LP_RS01255    | LysM peptidoglycan-binding domain-containing protein | mRNA         | Down     | 0.423        |
| LP_RS00230    | LysR family transcriptional regulator                | mRNA         | Down     | 0.291        |
| LP_RS15475    | Replication protein                                 | mRNA         | Down     | 0        |
| LP_RS15480    | Hypothetical protein                                 | mRNA         | Down     | 0        |
the remaining 21 genes were up-regulated in ΔLplttR (Fig. 4A). The 70 differentially expressed genes were assigned to different groups based on KEGG pathway analysis. As shown in Fig. 4B, 12 genes participated in metabolism, nine in environmental information processing, three in genetic information processing, and three in human disease. Notably, eight of the environmental information processing genes were carbon transport and sensing related proteins, that were closely correlated with carbohydrate metabolism, especially the fatty acid metabolism (Table 3). The PTS sugar transporters (agaF and celB), galactose mutarotase (galM), glycoside hydrolase (bglA), fatty acid metabolism-related enzymes (fabD, fabZ, fabH), and several ABC transporter proteins (oppA,
agaF, fliY, oppC, metI, and oppD) were under the control of LpLttR, suggesting that LpLttR might play an important role in carbohydrate metabolism in L. plantarum.

### Preliminary exploration of the regulatory mechanism of LpLttR

To better understand the sequence characteristics of the LpLttR binding box, a MEME search was performed to analyze the conserved motifs. The binding motif of LpLttR was predicted using MEME (https://meme-suite.org/meme/) according to the binding sites reported in previous studies, including the LplttR and clc operon regulated by LpLttR in L. plantarum [8], catBCA regulated by CatR in P. putida [4], clcABD operon on plasmid pAC27 regulated by Cler in Pseudomonas sp. strain P51 [9], tcbCDEF on plasmid pP51 controlled by TcbR of Pseudomonas sp. strain P51 [10], cbnABCD controlled by CbnR in R. eutropha [22], tfdA regulated by TfdR/S in R. eutropha JMP134 [5], catBCI-JFD regulated by CatM in Acinetobacter sp. benABCDE controlled by BenM in Acinetobacter, and linE-linD regulated by LinR in Sphingomonas [23]. Default settings were used in the site distribution (zero or one occurrence per sequence (zoops)). As shown in Fig. 5A, the predicted motif of LpLttR possessed a conserved binding motif 5′-(A/T)TAC-n7-(G/A)(T/A)-3′.

Next, the regulatory regions of the differentially expressed genes that increased or decreased more than twice were analyzed using bioinformatics. As shown in Fig. 5B, the regulatory regions of LP_RS00230, LP_RS13240 (located in the same transcription unit with LP_RS13245), LP_RS11700, LP_RS07440, LP_RS01255, and LP_RS12305 contained the potential LttR binding motif, suggesting that the promoter of these genes might be directly bound to LpLttR. LP_RS15475 and LP_RS15480

### Table 3 Differential genes involved in carbohydrate metabolism, lipid metabolism, and environmental information processing in KEGG pathway analysis

| Gene ID   | Description                                                                 | KO Name | KO Description                                                                 |
|-----------|------------------------------------------------------------------------------|---------|--------------------------------------------------------------------------------|
| Carbohydrate metabolism |                                                                             |         |                                                                                 |
| LP_RS11200 | PTS sugar transporter subunit IIa                                             | agaF    | PTS system, N-acetylgalactosamine-specific IIa component [EC:2.7.1.-]           |
| LP_RS14565 | galactose mutarotase                                                         | galM    | aldose 1-epimerase [EC:5.1.3.3]                                                |
| LP_RS12660 | glycoside hydrolase family 1 protein                                         | bgIA    | 6-phospho-beta-glucosidase [EC:3.2.1.86]                                        |
| LP_RS01000 | PTS sugar transporter subunit IIC                                            | celB    | PTS system, cellobiose-specific IIa component [EC:2.7.1.-]                      |
| LP_RS07140 | ACP S-malonyltransferase                                                     | fabD    | [acyl-carrier-protein] S-malonyltransferase [EC:2.3.1.39]                       |
| fabZ       | 3-hydroxyacyl-ACP dehydratase FabZ                                            | fabZ    | 3-hydroxyacyl-[acyl-carrier-protein] dehydratase [EC:4.2.1.59]                  |
| LP_RS07130 | ketoacyl-ACP synthase III                                                     | fabH    | 3-oxacyl-[acyl-carrier-protein] synthase III [EC:2.3.1.180]                     |
| Lipid metabolism |                                                                             |         |                                                                                 |
| LP_RS05365 | peptide ABC transporter substrate-binding protein                            | oppA    | oligopeptide transport system substrate-binding protein                           |
| LP_RS13055 | ABC transporter ATP-binding protein                                          |         | iron complex transport system ATP-binding protein [EC:3.6.3.54]                 |
| LP_RS11200 | PTS sugar transporter subunit IIa                                             | agaF    | PTS system, N-acetylgalactosamine-specific IIa component [EC:2.7.1.-]           |
| LP_RS13480 | transporter substrate-binding domain-containing protein                      | fliY    | L-cystine transport system substrate-binding protein                             |
| LP_RS05375 | ABC transporter permease                                                      | oppC    | oligopeptide transport system permease protein                                   |
| LP_RS07440 | ABC transporter permease                                                      | metI    | D-methionine transport system permease protein                                   |
| LP_RS05380 | ABC transporter ATP-binding protein                                          | oppD    | oligopeptide transport system ATP-binding protein                                |
| LP_RS01000 | PTS sugar transporter subunit IIC                                            | celB    | PTS system, cellobiose-specific IIa component [EC:2.7.1.-]                      |
| LP_RS06330 | response regulator transcription factor                                       | nreC    | two-component system, NarL family, response regulator NreC                      |

Environmental information processing |                                                                             |         |                                                                                 |
| LP_RS05365 | peptide ABC transporter substrate-binding protein                            | oppA    | oligopeptide transport system substrate-binding protein                           |
| LP_RS13055 | ABC transporter ATP-binding protein                                          |         | iron complex transport system ATP-binding protein [EC:3.6.3.54]                 |
| LP_RS11200 | PTS sugar transporter subunit IIa                                             | agaF    | PTS system, N-acetylgalactosamine-specific IIa component [EC:2.7.1.-]           |
| LP_RS13480 | transporter substrate-binding domain-containing protein                      | fliY    | L-cystine transport system substrate-binding protein                             |
| LP_RS05375 | ABC transporter permease                                                      | oppC    | oligopeptide transport system permease protein                                   |
| LP_RS07440 | ABC transporter permease                                                      | metI    | D-methionine transport system permease protein                                   |
| LP_RS05380 | ABC transporter ATP-binding protein                                          | oppD    | oligopeptide transport system ATP-binding protein                                |
| LP_RS01000 | PTS sugar transporter subunit IIC                                            | celB    | PTS system, cellobiose-specific IIa component [EC:2.7.1.-]                      |
| LP_RS06330 | response regulator transcription factor                                       | nreC    | two-component system, NarL family, response regulator NreC                      |
Fig. 5  LpLttR binding sites analysis. A Detecting the conserved binding motif of LpLttR using the MEME online tool. The LttrS used as the MEME input included the LpLttR and cla operon regulated by LpLttR in Lactobacillus plantarum, catBCA regulated by CatR in Pseudomonas putida, clcABD operon on plasmid pAC27 regulated by ClcR, tcbCDEF on plasmid pP51 controlled by TcbR of Pseudomonas sp. strain P51, cbnABCD controlled by CbnR in Ralstonia eutropha, tfdA regulated by TfdR/S in R. eutropha JMP134, catBCJFD regulated by CatM in Acinetobacter sp. benABCDE controlled by BenM in Acinetobacter, and linE-linD regulated by LinR in Sphingomonas. The motif count setting was searching for one motif. Motif width was between 6 and 50. B The predicted binding sites of LpLttR on the target gene promoters. The operons of the differential genes were predicted by the website (http://www.microbesonline.org/operons/gnc220668.html). C Molecular interaction of LpLttR to the regulatory region of the target genes. Both the correspondence of the gene names annotated in KEGG database and that used in the transcriptome sequencing are listed. The interaction mainly contains two processes: association and dissociation. During the association process, the spectral interference shift increased, while the wavelength shift decreased during dissociation. Different colors represent different target gene promoters.
were located on plasmids p0203 and p0203, respectively. The cis-element analysis of these two genes was not performed.

To further verify whether LpLttR regulates the expression of these genes directly, we used the purified LpLttR-His protein and the regulatory region DNA of the target genes to carry out molecular interaction experiments. LP_RS14775, with no predicted LttR-binding motif, was used as the negative control. As shown in Fig. 5C, LpLttR had the strongest affinity to the promoter of LP_RS12305 (KD = 0.104 μM) and different degrees of binding with LP_RS00230, LP_RS13240, LP_RS11700, LP_RS07440, and LP_RS01255, but not with LP_RS14775, which was consistent with the results of the binding site analysis. The specific binding between LpLttR and its own promoter region suggested that LpLttR in Lactiplantibacillus plantarum was self-regulated, similar to other LttRs. These results suggested that LpLttR in this species remained uncharacterized. In this study, we identified the regulon of LpLttR by transcriptomic analysis of the WT and LplttR knockout strains. LpLttR was highly conserved in sequence, and its knockout caused the transcriptional difference of 70 genes to be more than 1.5-fold, and 11 genes to be more than twofold. Many of the differentially expressed genes were mapped to the perception, metabolism, and transportation of carbon sources, revealing that LttRs might perform important functions in carbon metabolism. Through bioinformatics analysis and molecular interactions, we further verified that LpLttR directly regulated the expression of LP_RS00230, LP_RS13240, LP_RS11700, LP_RS07440, LP_RS01255, and LP_RS12305 by binding to the promoter regions. The binding motif consisted of a highly conserved consensus sequence: 5′- (A/T) TAC-N7- (G/A)/(T/ A) a(T/ A) -3′.

Our previous study showed that LplttR responded to LA and activated the transcription of cla-dh and cla-dc, promoting the biotransformation of CLA [8]. However, the LplttR regulon identified in this study does not contain cla-dh and cla-dc. This may be due to the difference in the culture media and conditions. The medium used in this study did not contain LA, under which condition the transcription of cla-dh and cla-dc was suppressed.

### Discussion

*L. plantarum* was one of the best studied lactobacilli; hundreds of studies and dozens of reviews have described its metabolism and regulation of metabolism. In our previous study, we found that the CLA biotransformation was transcriptionally regulated by LpLttR. However, the regulon of LpLttR in this species remained uncharacterized. In this study, we identified the regulon of LpLttR by transcriptomic analysis of the WT and LplttR knockout strains. LpLttR was highly conserved in sequence, and its knockout caused the transcriptional difference of 70 genes to be more than 1.5-fold, and 11 genes to be more than twofold. Many of the differentially expressed genes were mapped to the perception, metabolism, and transportation of carbon sources, revealing that LplttR might perform important functions in carbon metabolism. Through bioinformatics analysis and molecular interactions, we further verified that LpLttR directly regulated the expression of LP_RS00230, LP_RS13240, LP_RS11700, LP_RS07440, LP_RS01255, and LP_RS12305 by binding to the promoter regions. The binding motif consisted of a highly conserved consensus sequence: 5′- (A/T) TAC-N7- (G/A)/(T/ A) a(T/ A) -3′.

Our previous study showed that LplttR responded to LA and activated the transcription of cla-dh and cla-dc, promoting the biotransformation of CLA [8]. However, the LplttR regulon identified in this study does not contain cla-dh and cla-dc. This may be due to the difference in the culture media and conditions. The medium used in this study did not contain LA, under which condition the transcription of cla-dh and cla-dc was suppressed.

### Table 4 Strains and plasmids used in this study

| Strain or plasmid | Characteristic | Source or reference |
|-------------------|---------------|---------------------|
| Strains           |               |                     |
| Lactiplantibacillus plantarum ATCC BAA-793 | Wild type | [25] |
| LP_RS00230, LP_RS13240, LP_RS11700, LP_RS07440, and LP_RS01255 | LplttR deletion mutant strain | [8] |
| LP_RS15480, LP_RS13240, LP_RS11700, LP_RS07440, and LP_RS12305 | LttR overexpression strain, carrying pIB184-LplttR | [8] |
| E.coli BL21(DE3) Expression strain | Novagen |
| Plasmid | pET28a-LplttR | pET28a derivative carrying LplttR |
| pIB184-LplttR | pIB 184 carrying LplttR for gene overexpression |
| pLCNICK-LplttR | Used for LplttR deletion | [8] |

According to previous studies, LttRs are often induced or suppressed by environmental or metabolic co-inducers, thus activating or inhibiting the expression of target genes. For example, the metabolic intermediates of aromatic compounds generally act as inducers of LttR in the regulation of aromatic compound metabolism-related genes. It has even been shown that BenM could bind to different co-inducers leading to an altered protein conformation [1]. Therefore, the cellular processes in which LpLttR participates and its regulon may vary under different culture conditions.

### Conclusions

In summary, we investigated the function and regulation of LplttR in *L. plantarum*. The sequence of LpLttR was highly conserved with that of other transcriptional regulators in this family. However, the knockout of LpLttR showed no significant effect on the bacterial growth. Coinciding with the growth, only 11 genes exhibited a more than twofold transcriptional increase. LP_RS00230, LP_RS13240, LP_RS11700, LP_RS07440, LP_RS01255, and LP_RS12305 were directly regulated by LpLttR. LpLttR regulated the transcription of these genes by binding to the conserved LttR box (5′- (A/T) TAC-N7- (G/A)/(T/ A) a(T/ A) -3′). In this study, we systematically investigated the regulation of LplttR in *L. plantarum* and revealed that LpLttR regulated different genes and performed different functions in different species. These findings deepened our understanding of the regulatory mechanisms of LttRs and provided a theoretical basis.
for the metabolism and regulation mechanisms of *L. plantarum*.

**Methods**

**Strains**
The strains and plasmids used in the present study were listed in Table 4. *L. plantarum* ATCC BAA-793 and the derivate strains were cultured in MRS medium at 37 °C under aerobic conditions. *E. coli* BL21(DE3) was used for the expression and purification of LpLttR. It was cultured in LB medium at 37 °C, 200 rpm. The process of protein expression and purification has been described in a previous study [8].

**RNA extraction**
The WT and ΔLplttR strains were collected during the exponential growth period with three biological replicates. The bacteria were sent to Majorbio Bio-Pharm Technology Co., Ltd (Shanghai, China) for subsequent transcriptome sequencing. TRIzol Reagent (Invitrogen) was used to extract the total RNA according to the manufacturer's instructions. Agilent 2000 was used for the RNA quality determination, and Nanodrop2000 (NanoDrop Technologies) was used for the quantification. The integrity of RNA was assessed by agarose gel electrophoresis. Only high-quality RNA samples (OD260/280 ≈ 1.8 ~ 2.0, OD260/230 ≥ 2.0, RIN ≥ 6.5, 23S:16S ≥ 1.0, Concentration ≥ 100 ng/μl, and total amount of RNA ≥ 2 μg) were used for subsequent library construction.

**Library construction and transcriptome sequencing**
The TruSeq™ Stranded Total RNA Library Prep Kit from Illumina (San Diego, CA, USA) was used to construct the library for the experiment. After removing rRNA and adding fragmentation buffer, mRNA was randomly broken into small fragments of approximately 200 nt. Under the action of reverse transcriptase, one-strand cDNA was synthesized using random primers and mRNA as templates. For the second strand synthesis, dUTP was used instead of dTTP to form the base of the second strand of cDNA containing dTTP. Before PCR amplification, the second strand of cDNA was digested with the UNG enzyme so that only the first strand of cDNA was included in the library. Finally, Illumina HiSeq × 10 (2 × 150 bp read length) was used for sequencing. Processing of the original images to sequences, base-calling, and quality value calculations were performed using the Illumina GA Pipeline (version 1.6), in which 150 bp paired-end reads were obtained.

**Bioinformatics analysis**
The data generated from the Illumina platform were used for bioinformatics analysis. All the analyses were carried out using the I-Sanger cloud platform (www.i-sanger.com) from Majorbio Bio-Pharm Technology Co., Ltd (Shanghai, China). Sequencing
reads were compared to those in the Rfm database. The accession number of the reference genome was GCF_000203855.3. The transcriptome sequencing raw data in fastq format were deposited in the National Center for Biotechnology Information Sequence Read Archive (NCBI SRA) database (accession number: PRJNA751435).

Protein-DNA interaction
The regulatory regions of the target genes were amplified by PCR using the primers listed in Table 5. To label the DNA with biotin, a second PCR reaction was performed using a universal biotinylated primer (5′-biotin-AGC CAG TGG CGA TAAG-3′). The PCR products were purified using a PCR purification kit (Shanghai Generay Biotech). The quality and concentration of the biotin-labelled DNA probe was analyzed using 1% agarose gel electrophoresis and a Nanodrop 2000 spectrophotometer (Thermo Fisher Scientific), respectively.

The specificity of binding and affinity constant of LpLltR to the promoter of target genes were determined using the Octet system (Octet, ForteBio, USA) based on bio-membrane interference technology, according to the previous study [24]. The samples were added to the 96-well plates in a total volume of 200 μL. The reactions were performed at 37 °C with shaking at 1000 rpm. After loading with the biotinylated DNA, the streptavidin biosensors were transferred to His-LplltR solutions to associate, and then moved to running buffer to dissociate. The obtained data were processed by Octet Data Analysis version 7.0 using a 1:1 binding model.

RNA preparation and RT-PCR
L. plantarum and the derivative strains in the exponential growth period were collected by centrifugation at 4 °C. The total RNA was prepared and analyzed by qPCR as previously described [8]. The primers used for RT-PCR were listed in Table 6. Each PCR condition was performed in triplicate on the LightCycler 96 qRT-PCR system (Roche Diagnostics, Switzerland). The PCR procedure was as follows: 95 °C for 5 s, followed by 40 cycles of denaturation at 95 °C for 5 s and annealing at 60 °C for 30 s. 16S rRNA was used as the internal control. The obtained data was analyzed using analytical LightCycler 96 system, and the 2−ΔΔct method was used to calculate the transcriptional fold changes.

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Authors’ contributions
X‑XL designed the study, analyzed the data, and wrote the manuscript. LL performed the research. X‑S, G‑QW, Z‑QX, and Y‑JX contributed new methods. L‑ZA conceived the study. All authors read and approved the final manuscript.

Data availability
All data analyzed in this study are included in this published article.

Code availability
Not applicable.

Declarations

Ethics approval and consent to participate
This article did not contain research involving humans or animals performed by any of the authors.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no conflict of interest.

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Table 6 Primers used in qRT-PCR

| Primers   | Sequence (5′–3′)               |
|-----------|-------------------------------|
| 0057-qPCR-s | ACTTTGGTCGCCAGAAAGG           |
| 0057-qPCR-a | GACGACGATGATAGGCC             |
| P0202-qPCR-s | TGGTTCGATTTGGATTG            |
| P0202-qPCR-a | GGCATACCTGTGCCTTCC           |
| P0203-qPCR-s | GAGCCTTTATAGTGGTC            |
| P0203-qPCR-a | TTACCAACGGCGTCATC            |
| 3533-qPCR-s | ACGGCGATGATTTGGTTC           |
| 3533-qPCR-a | CTGGGATGACACCTGGAGA          |
| 3151-qPCR-s | ACAAGGGAAAGCTGATC            |
| 3151-qPCR-a | AGCAGCTTTCGCCGTA             |
| 2768-qPCR-s | CTGCCTTCTGCTTGTCC            |
| 2768-qPCR-a | GTTGCGCGAAATAATGGTAT         |
| 3153-qPCR-s | GCCATTCAGATTTACGA            |
| 3153-qPCR-a | TGATAGTGGACAGATAAGG          |
| 1745-qPCR-s | CGGCTGTTGACGAGATAA           |
| 1745-qPCR-a | GCACCGAAGCAGAGAA             |
| 0302-qPCR-s | ATTAACGCAATGCGATCAC          |
| 0302-qPCR-a | AAACCATACGGACACAGA           |
| 2920-qPCR-s | CACGTCTTTCGCGGTAT            |
| 2920-qPCR-a | ATGGGAAAGTTGCTGGTAT          |
| qPCR-16s-F  | CACATTGGGACTGAGACACGG        |
| qPCR-16s-R  | CGATGCACCTCTTGCCTGGTGG        |

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