Abstract
Carbohydrate metabolism of bacterial pathogens conducts crucial roles in regulating pathogenesis but the molecular mechanisms by which metabolisms and virulence are modulated and coordinated remain to be illuminated. Here, we investigated in this regard Edwardsiella piscicida, a notorious zoonotic pathogen previously named E. tarda that could ferment very few PTS sugars including glucose, fructose, mannose, N-acetylglucosamine, and N-acetylglactosamine. We systematically characterized the roles of each of the predicted 23 components of phosphotransferase system (PTS) with the respective in-frame deletion mutants and defined medium containing specific PTS sugar. In addition, PtsH was identified as the crucial PTS component potentiating the utilization of all the tested PTS sugars. Intriguingly, we also found that PtsH while not Fpr was involved in T3SS gene expression and was essential for the pathogenesis of E. piscicida. To corroborate this, His15 and Ser46, the two established PtsH residues involved in phosphorylation cascade, showed redundant roles in regulating T3SS yields. Moreover, PtsH was shown to facilitate mannose uptake and transform it into mannose-6-phosphate, an allosteric substrate established to activate EvrA to augment bacterial virulence. Collectively, our observations provide new insights into the roles of PTS reciprocally regulating carbohydrate metabolism and virulence gene expression.

Key points
• PTS components’ roles for sugar uptake are systematically determined in Edwardsiella piscicida.
• PtsH is involved in saccharides uptake and in the regulation of E. piscicida’s T3SS expression.
• PtsH phosphorylation at His15 and Ser46 is essential for the T3SS expression and virulence.

Keywords Edwardsiella piscicida · Mannose · Phosphotransferase system · Type III secretion system (T3SS) · Virulence

Introduction
Bacteria show high metabolic versatility towards carbon sources for survival and adaptation in continuously changing environments. Various transportation systems are equipped in bacteria to cope with sugar specificities for saccharides utilizing ATP, ion gradients, and phosphoenolpyruvate as energy sources. Phosphoenolpyruvate carbohydrate phosphotransferase system (PTS) is a multi-functional system in a diverse array of bacteria and plays pivotal roles in saccharides uptake (Jeckelmann and Erni 2019, 2020; Galinier and Deutscher 2017). During the utilization of distinct sugars, the phosphorylation state of carbohydrates would be modified by cognate PTS components before being delivered into the cytoplasm, which is varied respectively to PTS sugars and the metabolic state of the cell.

Conventionally, PTS contains two general components, enzyme I (EI) and the histidine-containing phosphocarrier protein (HPr), as well as sugar-specific enzyme II (EII) complexes. EII family proteins consist of EII A/B/C/D subfamilies and EII D only appear in the PTS-mannose family (Stolz et al. 1993; Liu et al. 2019). In response to the availability of specific
carbohydrates, the phosphoenolpyruvate (PEP) functions as phosphoryl donor for the EI protein and the histidine-containing phosphate carrier (HPr) transmits the phosphate group to the downstream EII proteins, which phosphorylates imported sugars (Deutscher et al. 2006, 2014).

Until now, the notion that carbohydrate metabolism conducts a crucial role in regulating virulence in various bacterial pathogens is becoming an emerging theme. The defective expression of ptsI and crr, encoding enzyme I and enzyme IIαGlc, reduced the expression of PhoPQ regulon associated with quorum sensing and switched Salmonella from growth arrest to acute virulence through activation of virulence factor secretion (Maze et al. 2014; Lim et al. 2019). In Vibrio cholerae, PTS modulates virulence gene expression by regulating the expression of tcpH and aphAB to control the expression of toxT, the central activator of virulence gene expression (Wang et al. 2015). A promiscuous mannose-family PTS transporter of Streptococcus exerted influence on SLS-mediated hemolysis and the route of glucose uptake impacting their survivals in human blood (Sundar et al. 2017; 2018). In a mouse model, the impairments in PTS components in Bacillus anthracis and Borrelia burgdorferi attenuated in vivo virulence via affecting the expression of their master virulence regulators, respectively (Khajanchi et al. 2016; Bier et al. 2020). However, the mechanisms underlying that the specific phosphotransfer reactions mediated by individual sugar-specific PTS branches or components are integrated into a global virulence program to control bacterial pathogenesis remain elusive.

Edwardsiella piscicida is an important gram-negative zoonotic pathogen that infects a variety of farmed fish, mainly depending on its type III secretion system (T3SS) and type VI secretion system (T6SS) controlled by two-component system EsrA-EsrB (Leung et al. 2019). Given the sluggish growth in fermented sugars, E. piscicida was initially named E. tarda, implying that limited PTS sugars can be utilized by E. piscicida (Abott and Janda 2006; Wang et al. 2009; Shao et al. 2015). Recently, we revealed that EvrA is a mannose-6-phosphate (Man-6P) sensor that directly activates esrB expression to control T3SS production in E. piscicida (Wei et al. 2019). How mannose is transported and activated to fulfill virulence regulation is yet unknown.

Here, we identified 23 components predicted for sugar and nitrogen PTS systems in E. piscicida. We also systematically defined the distinct genes required for the 5 PTS sugars utilized by the bacterium. Specifically, PtsH not only participated in carbohydrate utilization but also controlled the expression of T3/T6SS genes in E. piscicida. Moreover, residues His15 and Ser46, the established sites for PtsH phosphorylation, showed redundancy in modulating virulence gene expression. In addition, PtsH while not Fpr played a role in modulating virulence gene expression. Finally, PtsH potentiated and transformed mannose to Man-6P to activate virulence regulator EvrA, augmenting virulence gene expression. Collectively, our findings provide new insights into the roles of PtsH in coordination between carbohydrate metabolism and the virulence gene expression of E. piscicida.

Materials and methods

Bacterial strains and culture conditions

The strains and plasmids used in this study are shown in Table S1. E. piscicida EIB202 and WTΔP, the wild-type strain cured of its endogenous Cm and Str resistance plasmid pEIB202 (Wang et al. 2009), were used as the parental strains for the construction of the in-frame deletion mutant strains (Supplemental Table S1). E. piscicida strains were grown in Luria–Bertani broth (LB) or on LB plates (Oxoid, England) and in Dulbecco’s modified Eagle’s medium (DMEM) (Invitrogen, USA) at 30 °C. DMEM was used to induce T3/T6SS production as well as to yield auto-aggregation phenotypes mimicking in vivo conditions (Zheng et al. 2005). When required, antibiotics were supplemented at the following concentrations: colistin (Col, 16.7 µg/mL), ampicillin (Amp, 100 µg/mL), and kanamycin (Km, 25 µg/mL).

Construction of deletion mutants and complemented strains

In-frame deletion mutants were generated using sacB-based allelic exchange (Yin et al. 2018). Overlap PCR was used to generate appropriate DNA fragments for creating in-frame deletions in each target gene. The fragments were inserted into the suicide vector pDM4 with Gibson assembly (Gu et al. 2016) and the resulting plasmids were introduced into Escherichia coli SM10 λpir for conjugation into E. piscicida EIB202 or WTΔP. Transconjugants were selected on LB plates containing Km and Col. Double-crossover events were subsequently selected on LB plates containing 12% sucrose. The fragments containing the corresponding ORF region and putative promoter region were cloned into plasmid pUTt to obtain complementation strains (Yin et al. 2018). All newly constructed plasmids were verified by sequencing. All primers used in this study are listed in Supplemental Table S2.

PTS transport analysis

The PTS-related metabolic pathways were predicted on Kyoto Encyclopedia of Genes and Genomes (KEGG) with default parameters (https://www.genome.jp/kegg/pathway.html). The PTS components were further identified using the BlastN algorithm (NCBI) based on their corresponding homologous sequences. Overnight cultures were grown
in LB medium at 30 °C for 24 h. Subsequently, 1% (v/v) inoculum of all *E. piscicida* cultures was inoculated in M9 medium with or without indicated carbohydrates. The M9 medium does not contain any carbohydrates, leading to the growth deficiency of *E. piscicida* strains. The initial pH of the M9 medium with 0.5% (w/v) various sugars was 7.27 ± 0.07. Bromothymol blue (0.1% w/v, Sigma-Aldrich) was used as a pH indicator of sugar fermentation.

**Determination of growth curve**

Growth curves were determined by Bioscreen C (Oy Growth Ab Ltd, Finland). All *E. piscicida* strains were inoculated into LB for overnight culture and then diluted into fresh LB or DMEM at an OD<sub>600</sub> of ~0.1. Two hundred microliter cultures were transferred into a 100-well plate and then statically cultured at 30 °C. The non-inoculated LB or DMEM were used as a control and all measurements (n = 6) were repeated at least three times in independent experiments.

**Total RNA extraction and qRT-PCR**

*E. piscicida* strains were statically cultured in DMEM. RNA samples were extracted with a commercial RNA isolation kit (Tiangen, China) and mRNA was reverse-transcribed into cDNA using the FastKing RT kit (Tiangen, China). qRT-PCR was performed with an Applied Biosystems 7500 cycler (Applied Biosystems, USA) with triplicate reactions for each sample. The comparative C<sub>T</sub> (2<sup>−ΔΔCT</sup>) method (Gu et al. 2016) was used to quantify the relative levels of each transcript with the specific primer pairs (Table S2) and *gyrB* gene was used as an internal control (Yin et al. 2018).

**RNA-seq**

For the preparation of mRNA for RNA-seq, the Ribo-Zero-rRNA kit (Epicentre, USA) was initially used to remove rRNA from the RNA samples. The final concentration of RNA samples was determined with a Qubit 2.0 Fluorometer (Thermo Fisher, USA). The VAHTS Stranded mRNA-seq Library Prep Kit for Illumina (Vazyme, China) was used to construct strand-specific RNA-seq libraries, and sequencing was conducted on an Illumina HiSeq 2500 platform, yielding the paired-end reads. Adapter sequences and low-quality bases (PHRED quality scores ≤ 5) were trimmed with the Trimmomatic package (Bolger et al. 2014) using the default parameters and reads smaller than 35 bp were discarded. The RNA-seq data processing procedures and statistical analysis were performed (Tjaden 2015).

**SDS-PAGE and western blotting analysis**

Whole cell proteins (WCPs) and extracellular proteins (ECPs) were extracted and concentrated (Yin et al. 2018). Overnight cultures of *E. piscicida* were sub-cultured into 50-mL fresh DMEM and statically incubated for 24 h to 72 h at 28 °C; bacteria were then harvested by centrifugation at 5,000×g for 10 min at 4 °C for WCPs. For ECPs, culture supernatants were filtered with 0.22-µm filters (Millipore, USA), and concentrated using 10 kDa cutoff centrifugal filter devices (Millipore, USA). Proteins were separated by 12% SDS-PAGE, followed by Coomassie blue staining. For western blots, separated proteins were wet transferred onto PVDF membranes (Millipore, USA) and incubated with a 1:1000 dilution of mouse anti-EseB (GL Biochem, China). HRP-conjugated anti-mouse IgG (Santa Cruz Biotechnology, CA) was used at a 1:2,000 dilution as a secondary antibody. Proteins were visualized with TMB substrate solution at 8,000×g for 2 min at 4 °C and washed three times with phosphate-buffered saline (PBS, pH 7.4). A total of 5.0×10<sup>5</sup> colony-forming units (CFUs) bacteria suspended in PBS were injected into each fish via the intraperitoneal (i.p.) route and PBS was used as a negative control. A total of 30 fish were injected with each strain and fish mortality was monitored daily. The infection experiments were performed at least three independent times.

**Animal survival and competitive index assays**

Turbot experiments were performed according to protocols approved by the Animal Care Committee of the East China University of Science and Technology (2,006,272) and the Experimental Animal Care and Use Guidelines from the Ministry of Science and Technology of China (MOST-2011–02). Healthy turbot weighing 30.0 ± 3.0 g (~2 months old and ~ 1:1 female to male) were obtained from a commercial farm (Yantai, China) and acclimatized with a continuous flow of seawater for 7 days at 16 °C. For fish survival assays, overnight cultures were harvested by centrifugation at 8,000×g for 2 min at 4 °C and washed three times with phosphate-buffered saline (PBS, pH 7.4). A total of 5.0×10<sup>5</sup> colony-forming units (CFUs) bacteria suspended in PBS were injected into each fish via the intraperitoneal (i.p.) route and PBS was used as a negative control. A total of 30 fish were injected with each strain and fish mortality was monitored daily. The infection experiments were performed at least three independent times.

Competitive index (CI) assays were performed between WT, or the indicated gene-deletion mutant strains, and WTΔp, the WT strain cured of its endogenous Cm and Str resistance plasmid pEIB202 (Wang et al. 2009). Inocula were prepared using fresh cultures of bacteria that were diluted and mixed at a 1:1 ratio. A total of ~10<sup>5</sup> CFUs bacteria in a 100 µL inoculum were i.p. injected into each fish. At 8-day post-injection (DPI), the livers from fish in each group (6 fish/group) were collected, homogenized, and plated on LB plates with or without the presence of Cm to distinguish WT Δp (Cm<sup>+</sup>) from other strains (Cm<sup>−</sup>) (Yang et al. 2017).
The ratios of the bacterial counts were used to determine CI values.

**Statistical analysis**

GraphPad Prism (version 8.0) was used for the statistical analyses. Data are presented as the mean ± SD of triplicate samples per experimental condition unless otherwise noted. Statistical analyses were performed using unpaired two-tailed Student’s t test for the metabolite level analysis, one-way ANOVA analyses followed by Bonferroni’s multiple-comparison post-test comparing the data of CI values, or Kaplan–Meier survival analysis with a log-rank test. Differences were considered significant at *P < 0.05, **P < 0.01, and ***P < 0.001.

**Results**

**Identification of the PTS components and cognate PTS sugars in E. piscicida**

*Edwardsiella* genus bacterium was initially given the name “tarda” to describe its weak ability of sugar fermentation and the genetic basis underlying this phenotype remains unknown. Bioinformatics analysis with KEGG indicated that E. *piscicida* EIB202 encodes 23 PTS components (Wang et al. 2009). Similarly, all the currently known Edwardsiella species encode very few PTS genes (24 for *E. ictaluri*, 29 for *E. anguillarum*, 28 for *E. tarda*, and 36 for *E. hoshinae*) as compared to other enteric pathogens, e.g., 58 for Salmonella Typhimurium LT2 and 52 for *E. coli* O157:H7 EDL933. KEGG entry etr02060 informed that *E. piscicida* encodes 20 genes for sugar PTS and 3 for nitrogen PTS (PTSNtr), respectively, including one protein for EI, 2 for HPs, and 7 for EIIIs (Fig. 1A and Table 1). Of them, PtsI was supposed to be PEP-related protein phosphotransferase EI and Hpr proteins were annotated to include Fpr for the utilization of fructose, and PtsH supporting the transport of the rest of sugars. A total of 7 saccharides were predicted as the cognate potential PTS sugars including glucose (Glc), fructose (Fru), mannose (Man), N-acetylglucosamine (GlcNAc), N-acetylgalactosamine (GalNAc), L-ascorbic acid (LAc), and β-glucoside (β-Glu).

**Functional analysis of PTS components respective PTS sugar transport in E. piscicida**

To verify the utilization of putative PTS sugars, the WT strain was inoculated into M9 medium with the corresponding saccharides (Fig. 2A). Ascorbyl glucoside (AGlu) was used as the β-glucoside carbon source substrate. The growth
of WT in M9 medium alone and M9 supplemented with AGlu and LAc was severely low, suggesting that β-glucoside and L-ascorbic acid could not be efficiently utilized by *E. piscicida*. Glucose, fructose, mannose, N-acetylglucosamine, and N-acetylgalactosamine enabled the growth of WT (Fig. 2A).

In order to further define the roles of PTS components in specific sugar transport, we set out to systematically construct the in-frame deletion mutants of them. In total, 16 in-frame deletion mutants of predicted sugar PTS system components were successfully constructed and yet we failed to obtain the deletion mutants of the essential genes *ptsG* (EII components) shown comparable to that of WT (Fig. 2B). The growth of Δ*ptsI* and Δ*ptsH* was severely impaired in all the rest of tested PTS sugars (Table 2), validating that *ptsG* operon corresponding PTS sugars were supplemented as the sole carbon sources (Table 2), validating that *ptsG* operon corresponding PTS sugars were supplemented as the sole carbon sources (Table 2), validating that *ptsG* operon corresponding PTS sugars were supplemented as the sole carbon sources (Table 2).

| Family | Saccharide | Component | Gene | Annotation |
|--------|------------|-----------|------|------------|
| EI     | Fructose   | EI(PtsI)  | ETAE_1131 ptsI | Phosphoenolpyruvate-protein phosphotransferase (PEP EI) |
| HPr    | Fructose   | FPPr      | ETAE_2303 ptsH | Phosphohistidine-protein-hexose phosphotransferase component |
| EII    | Fructose   | EII(BC^fru | ETAE_2301 fruA | Fructose-specific PTS system IIBC component |
| Glucose| Glucose    | EIAGlc    | ETAE_1130 crr | Glucose-specific PTS system enzyme IIA |
|        | L-Ascorbic acid | EIIAGlAc | ETAE_3026 sgaA | Putative phosphotransferase enzyme IIA component SgcA |
|        | Mannose    | EIIMan    | ETAE_1559 manX | Mannose-specific PTS system enzyme IIA component |
|        | N-Acetylgalactosamine | EIIGalNAc | ETAE_2532 agaF | N-Acetylglactosamine-specific PTS system enzyme IIA |
|        | N-Acetylglucosamine | EIICMan | ETAE_1558 manY | Mannose-specific PTS system enzyme IIBC component |
|        | β-Glucoside | EIICLAc 1 | ETAE_1557 manZ | Mannose-specific PTS system enzyme IIDD |
|        | N-Acetylglucosamine | EIICLAc 2 | ETAE_2533 agaE | N-Acetylglactosamine-specific PTS system enzyme IIDD |
|        | β-Glucoside | EIICLAc 1 | ETAE_1557 manZ | Mannose-specific PTS system enzyme IIDD |

We further measured the growth of all the 18 deletion mutants in glucose-supplemented DMEM mimicking intracellular conditions. Similarly, Δ*ptsI* growth was also severely impaired in this condition (Table 2). Δ*ptsH* and Δ*manZ* exhibited severe growth retardation in this glucose-containing DMEM (Fig. 2A–B, Table 2). The *ptsH*+ and *manZ*+ complement strains rescued their growth as that of WT (Fig. 3A–B). However, there were no growth defects in Δ*ptsH* and Δ*manZ* compared to that of WT in nutrient-rich conditions.
Collectively, PtsH and ManZ seem to play important roles for growth in intracellular conditions in E. piscicida. We further explored whether carbohydrate metabolism influences the pathogenesis of E. piscicida. Due to the appearance of EseB-mediated protein filament, T3SS activity was investigated. The results are shown as the mean ± S.D. (n = 6). ***P < 0.01, NS, not significant (P > 0.05), based on the student's t-test in comparison to WT. A: Growth of the wild-type EIB202 (WT) and PTS mutant strains in M9 medium supplemented with defined sugars. Growth of the WT and ΔptsH strains inoculated in M9 medium supplemented with 5 mg/mL of indicated PTS sugars, including glucose (Glc), fructose (Fru), mannose (Man), N-acetylglucosamine (GlcNAc), N-acetylgalactosamine (GalNAc), and ascorbyl glucoside (AGlu). The OD600 values of bacterial cultures were measured at 24 h post-inoculation. The results are shown as the mean ± S.D. (n = 6). B: Growth curves of 18 deletion mutants of PTS system components grown in M9 medium supplemented with 5 mg/mL glucose as the sole carbon source. The results are shown as the mean ± S.D. (n = 6).
production could be assessed by auto-aggregation phenotype in *E. piscicida* (Gao et al. 2015). Δ*ptsI* was not considered in the following experiments owing to its poor growth in DMEM. Intriguingly as Δ*agaE* and Δ*manZ* exhibited marked deficiency in auto-aggregation, whereas other mutants possessed similar auto-aggregation phenotypes as WT did (Fig. 2C). Consistent with the auto-aggregation phenotypes, the reduced amounts of extracellular T3SS proteins were observed in Δ*ptsH* and Δ*manZ* as confirmed by western blot analysis against EseB (Fig. 2D).

**PtsH modulates T3/T6SS gene expression as revealed by RNA-seq**

We further utilized RNA-seq to compare the transcriptomes of Δ*ptsH* and WT cells grown in DMEM. Differential transcriptomic analyses indicated 163 differentially expressed genes in Δ*ptsH* compared to those in WT ((|log2(FC)| > 1 and \(P < 0.05\)) (Fig. 4A, Supplemental Table S3). Specifically, the expressions of 36 genes were down-regulated and the expressions of 127 genes were up-regulated. With regard to virulence genes, almost all of the T3/T6SS genes \( (n = 49) \) had significantly lower transcript levels in Δ*ptsH* than those in WT (Fig. 4A–B). Consistent with the crucial role of PtsH in carbohydrate metabolism, the PtsH-regulated genes were mainly grouped into “energy production and conversion,” “amino acid transport and metabolism,” and “carbohydrate transport and metabolism” (Fig. 4C). Subsequently, we set out to detect transcript levels of established virulence regulatory genes (*esrA*, *esrB*, and *esrC*), T3SS genes (*eseB*, *eseC*, and *eseD*), and T6SS genes (*evpP*, *evpC*, and *evpI*) by qRT-PCR assays. Due to the absence of *ptsH*, T3/T6SS gene transcripts significantly declined compared to those in WT, whereas no apparent changes were found in the transcript levels of virulence regulators EsrA/B/C (Fig. 4D). These results demonstrated PtsH’s function in controlling not only the carbohydrate metabolism but also the expression of virulence determinants.

**Phosphorylation of PtsH is involved in the regulation of T3SS yields**

The complement of *ptsH* could fully restore the auto-aggregation and T3/T6SS production in Δ*ptsH* (Fig. 5A–B), demonstrating that PtsH modulates the expression of T3/T6SS genes. As a histidine-containing phosphocarrier protein, HPr is generally phosphorylated at the active site residues, i.e., histidine 15 (His15) and serine 46 (Ser46) (Waygood 1998; Marquez et al. 2002). In PtsH, His15~P and Ser46~P were previously established to be involved in PTS sugar utilization and bacterial virulence regulation, respectively (Deutscher et al. 2005;
Fig. 3 PtsH and ManZ are essential for the T3SS yields in *E. piscicida* cells grown in DMEM. **A**–**B** Growth curve of WT, ΔptsH, ΔmanZ, ptsH+, and manZ+ strains grown in DMEM. The OD$_{600}$ values of indicated cultures were measured at the indicated time point post-inoculation. Results were shown by the mean±SD (n=6). **C** Auto-aggregation in the deletion mutants of putative PTS component genes grown in DMEM supplemented with 5 mg/mL glucose. **D** Extracellular protein profiles (ECP, upper panel) and western blot (lower panel) analysis of EseB and EvpP yields in the indicated mutants. DnaK in the cell lysates (WCP) was used as the loading control. All images are representative of triplicated experiments.
To explore the crucial amino acid residue of PtsH contributing to the T3SS production, PtsH variants, i.e., H15A, S46A, H15A/S46A, D23A/D69A, H15A/D32A/D69A, and H15A/D32A/S46A/D69A, were expressed in the ΔptsH strain. The alanine substitutions of residues aspartic acid 32 as well as 69 (D23A/D69A) were used as negative controls (Fig. 5C). The variants containing both H15A and S46A, i.e., PtsH^{H15A/S46A} and PtsH^{H15A/D32A/S46A/D69A}, exhibited deficient T3/T6SS production similar to that of ΔptsH (Fig. 5B–C), while other PtsH variants including the single substitution of H15A or S46A possessed similar extracellular protein profiles comparable to that of WT. Additionally, we sought to determine the transcript levels of T3/T6SS genes (eseB, eseC, and eseD) and T3SS regulatory genes (esrA and esrB) in the indicated strains expressing PtsH variants. Congruent with the extracellular protein profiles, the significantly decreased T3SS gene transcripts were identified in strains harboring substitutions of both H15A and S46A, further indicative of the requirements of phosphorylation in both His15 and Ser46 for the modulation of T3SS expression (Fig. 5D).
Fig. 5 PtsH phosphorylation at His15 and Ser46 residues is redundant in the activation of T3SS gene expression. **A** Auto-aggregation of ΔptsH and the complement ptsH+ strains. The empty plasmid pUTt was introduced into WT or ΔptsH strains as a control. **B–C** ECP profiling (upper panel) and western blot (lower panel) analysis of EseB and EvpP yields in ΔptsH and ΔptsH strains expressing wt PtsH (ptsH+) (B), or PtsH variants with substitution in the residues that could not be phosphorylated (C). DnaK in the ECP and WCP was used as the loading control. **D** qRT-PCR analysis of the transcript levels of indicated T3SS genes in WT or ΔptsH cells expressing wt or variants PtsH grown in DMEM. The results are shown as mean±SEM (n = 3), ***P < 0.001. NS, not significant, based on the student’s t test. gyrB was used as the internal control.
PtsH coordinates mannose utilization and virulence gene expression

Mannose-6-phosphate (Man-6P) was established to bind to virulence regulator EvrA to activate esrB expression (Wei et al., 2019). We thus investigated the roles of mannose utilization in mediating virulence towards fish host in E. piscicida. Corroborating to our previous findings (Wei et al., 2019), the deletion of manZ resulted in impaired auto-aggregation and production of the extracellular T3SS proteins (Fig. 6A and B). The reintroduction of manZ fully rescued these defects in ΔmanZ. In line with these results, the transcript levels of T3SS genes (eseB, eseC, and eseD) and T6SS genes (evpP, evpC, and evpl) were all significantly declined in ΔmanZ cells grown in DMEM supplemented with mannose as compared to that of the WT (Fig. 6C). Moreover, the transcript levels of T3/T6SS genes were significantly augmented in E. piscicida cells grown in DMEM supplemented with mannose as compared to that with glucose (Fig. 6D). Mannose modulation of T3SS gene expression also appeared to be dependent on PtsH as the transcript levels of T3SS genes in ΔptsH cells showed no difference between DMEM containing mannose and glucose (Fig. 6D). It was also notable that ptsH expression seems not to be affected by mannose, and the expression of evrA was up-regulated in response to mannose in WT but not in ΔptsH cells (Fig. 6C–D). Taken together, these data indicated that PtsH and ManZ are essentially involved in mannose utilization, which facilitates virulence gene expression in E. piscicida.

A turbot infection model was used to explore PtsH role in controlling virulence in vivo. ΔptsH and ΔmanZ strains exhibited significantly attenuated virulence. Approximately 70% of fish survived after infection with ΔptsH and ΔmanZ, while the fish challenged with ptsH+ and manZ+ showed 100% mortalities at 12 DPI, similar to those infected with WT (Fig. 6E). Meanwhile, in vivo CI assays were carried out to explore whether PtsH is required for optimal E. piscicida fitness during infection (Fig. 6F) with the ΔptsH and ΔmanZ related strains and WT ΔP, the wild-type strain with pEIB202 cured, proficient in in vivo colonization (Fig. 6F) (Yang et al. 2017). The ΔptsH and ΔmanZ were significantly impaired in the in vivo competition with WT and the corresponding complement strains, ptsH+ and manZ+ resulted, showed WT level of colonization. Collectively, these data demonstrated that PtsH and ManZ are critical for E. piscicida optimal growth in vivo.

Discussion

E. piscicida has been increasingly recognized to be a huge threat to the world’s aquaculture industries. To elucidate the mechanism of the carbohydrate metabolism in E. piscicida and its potential relationship to pathogenesis, the PTS sugars utilized by E. piscicida the crucial PTS components facilitating the cognate sugar transport were determined in this study. Being the crucial component of PTS, PtsH supported E. piscicida to use mannose both as the carbon source for growth and as a specific host signal to trigger virulence program. Specifically, PtsH phosphorylation at His15 and Ser46 was revealed to be critical for extracellular T3SS yields. Additionally, the disruption in ptsH impaired the bacterial colonization and infection in fish host. These results further delineated that mannose represents as a crucial metabolic signal during infection (Wei et al. 2019), laying a good foundation for the development of novel therapeutics against bacterial infection.

In this work, we systematically defined the genetic basis for PTS sugar utilization in E. piscicida. As a versatile zoonotic pathogen threatening various piscine hosts as well as humans, it is a counterintuition that the bacterium encodes paucity of genetic resources for saccharides uptake. Comparative genomic analysis also discerned that E. piscicida significantly differs from other Enterobacteriaceae bacteria with the lowest ratio of gene contents related to carbohydrate transport and metabolism (G), reflecting that the organism may be well adapted to the aquatic ecosystems and intracellular niches, where may exist relatively mean carbohydrates (Wang et al. 2009; Yang et al. 2012; Leung et al. 2022). However, we might have underestimated the genetic flexibility of E. piscicida in dealing with carbohydrate utilization. An example come up with the observation that ΔptsH also showed marginal growth using glucose, mannose, N-acetylglucosamine, and N-acetylgalactosamine as the sole carbon source, respectively (Fig. 2A), suggesting compensatory pathways have been evolved for fermentation of these PTS sugars. Indeed, E. piscicida encodes ETAE_1144 and ETAE_2966 for a galactose permease (GalP) and a glucose kinase (Glik), respectively, that may support its growth in glucose and other hexoses in the absence of cognate PTS as in E. coli (Hernández-Montalvo et al. 2003). Reverse genetics technologies such as using the defined mutant library and transposon insertion sequencing technology (Wei et al. 2019; Yang et al. 2017) might be useful to facilitate the further annotation of genes for defined carbohydrates in E. piscicida and other bacteria.

Our data revealed the unexpectedly important role of PtsH in E. piscicida virulence. More importantly, His15 and Ser46 were identified as the crucial residues to coordinate T3SS activation (Fig. 4). Unlike the canonical phosphorylation of His15 controlled by PEP, the ATP-dependent phosphorylation of Ser46 requires a specific HPr kinase/phosphorylase (HprK/P), leading to the inhibition of the phosphorylation of His15 in other bacteria. For instance, the phosphorylation of Ser46 of B. subtilis retarded the phosphoryl transfer from EI to HPr at least 100-fold (Deutscher

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**A**

Comparative analysis of WSZ, WT, and manZ strains under DMEM+Man conditions. Bar graph showing relative fold change in gene expression.

**B**

Western blot analysis of protein expression levels under different conditions. MW (kDa) shown on the left.

**C**

Relative fold change in gene expression under DMEM+Man conditions. Bar graph with NS and *** indicating significance levels.

**D**

Relative fold change in gene expression under DMEM conditions. Bar graph with NS and *** indicating significance levels.

**E**

Percent survival (%) over DPI for WT, PBS, ΔptsH, ptsH+, ΔmanZ, and manZ+ strains. Graph showing survival rates.

**F**

CI (Combined Index) for different strain combinations. Scatter plot showing CI values with NS and *** indicating significance levels.
PtsH coordinates with ManZ to regulate bacterial virulence in a mannose-dependent manner. A Auto-aggregation of WT and mannose (Man) uptake-related mutant strains grown in DMEM supplemented with Man as the sole carbon source. B ECP profiling (upper panel) and western blot (lower panel) analysis of EseB and EvpP yields in of WT, ΔmanZ, and its complement strain ΔptsH. DnaK in the ECP and WCP was used as the loading controls. C qRT-PCR analysis of the transcript levels of indicated T3/T6SS genes in WT and ΔmanZ cells grown in DMEM supplemented with 5 mg/mL mannose. The results are shown as mean ± SEM (n = 3) and gyrB was used as the internal control. D qRT-PCR analysis of the transcript levels of indicated T3/T6SS genes in WT (red) and ΔptsH (blue) cells grown in DMEM supplemented with 5 mg/mL mannose compared with those with 5 mg/mL glucose. E Survival curves of turbot challenged with the ptsH and manZ related strains. Phosphate-buffered saline (PBS, pH 7.4) was used as a control. The bacterial strains were suspended in PBS and injected into each turbot at a dose of 5.0 × 10⁵ CFU/fish (n = 30 fish/group). Kaplan–Meier survival analysis with a log-rank test is shown. F Competition index (CI) assays of equally mixed ptsH or manZ strains with WTΔp, WT cured of plasmid pEIB202, in turbot. Data presented are the mean ± SD (n = 6), * p < 0.01; NS, not significant, P > 0.05, based on ANOVA followed by Bonferroni’s multiple-comparison post-test comparing the data to the values from the WTΔp / WT

et al. 2006). Moreover, PtsH/Ser ~ P was supposed to directly or indirectly interact with virulence regulators, thereby providing a link between carbon metabolism and pathogenesis (Deutscher et al. 2005). Our results here support a model that the PTS-mediated T3SS activation depends on both the level of PEP intermediates and the activities of HPr kinase (Fig. 7), both of which reflect the intracellular ATP level and other physiological states/signals of the cell. Further studies are required to unravel the detailed mechanisms of PtsH’s phosphorylation on the virulence of *E. piscicida*.

Since extracellular mannose augments *E. piscicida* virulence in an EvrA-dependent manner (Wei et al. 2019), the loss of ManZ attenuated virulence due to the dysfunction of transportation of extracellular mannose. Given the function of PtsH as the phosphate group carrier to deliver it to enzyme IIC to activate PTS-mannose, ΔptsH is blocked in the import of mannose, leading to the same attenuated virulence as that of ΔmanZ. The similar virulence phenotype of ΔptsH and ΔmanZ implied that PtsH and ManZ coordinate *E. piscicida* pathogenicity. Therefore, PtsH and ManZ are indispensable for the enhanced virulence by mannose, which facilitates pathogen sensing of the host environment and triggers virulence programs (Fig. 7). However, our data also support an alternative model in which PtsH might interact with some other factor(s) to facilitate the transfer of phosphoryl group to PTS-mannose complex, which warrants further investigation. Given the essential roles of PtsH in supporting the importation of mannose, the metabolic signal that *E. piscicida* encounters at specific points during infection, and in controlling virulence gene expression by reflecting ATP level and other physiological signals, it could be merited as a valuable gene target for construction of live attenuated vaccines. The same “one stone two bird” strategies have been adopted by using Fur and PhoP that play essential roles in metabolism and virulence regulator as the effective targets for the development of vaccines combating edwardsielliosis infection (Swain et al. 2020, 2022). Taken together, our work deepens our understanding of how bacterial pathogens utilize available sugar to control virulence programs and might be useful for live attenuated vaccine development.

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**Author contribution** Conceptualization: WQ, SS; data curation: MQQ, JIH; funding acquisition: WQ, SS; investigation: MQQ, JIH, WX, MY; project administration: WQ, SS; supervision: WQ, SS; validation: WQ, SS; visualization: WQ, SS; writing (original draft): SS, MQQ; writing (review and editing): WQ, SS.

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**Data availability** All data associated with this study are presented in the main text or supporting information. The raw data of RNA-seq analysis were deposited in NCBI Sequence Read Archive (SRA) database under accession number PRJNA765294.

**Declarations**

**Ethics approval** All animal protocols used in this study were approved by the Animal Care Committee of the East China University of Science and Technology (2006272). The Experimental Animal Care and Use Guidelines from Ministry of Science and Technology of China (MOST-2011–02) were strictly followed.

**Conflict of interest** The authors declare no competing interests.

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