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Characteristics and Complete Genome Analysis of a Pathogenic *Aeromonas Veronii* **SJ4 from Diseased** *Siniperca Chuatsi*

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Abstract

As an opportunistic pathogen, *Aeromonas veronii* can cause hemorrhagic septicemia of various aquatic animals. In our present study, a dominant strain SJ4, isolated from naturally infected mandarin fsh (*Siniperca chuatsi*), was identifed as *A. veronii* according to the morphological, physiological, and biochemical features, as well as molecular identifcation. Intraperitoneal injection of *A. veronii* SJ4 into *S. chuatsi* revealed clinical signs similar to the natural infection, and the median lethal dosage (LD_{50}) of the SJ4 to *S. chuatsi* in a week was 3.8×10^5 CFU/mL. Histopathological analysis revealed that the isolate SJ4 could cause cell enlargement, obvious hemorrhage, and infammatory responses in *S. chuatsi*. Detection of virulence genes showed the isolate SJ4 carried *act*, *fm*, *fgM*, *ompA*, *lip*, *hly*, *aer*, and *eprCAL*, and the isolate SJ4 also produce caseinase, dnase, gelatinase, and hemolysin. In addition, the complete genome of *A. veronii* SJ4 was sequenced, and the size of the genome of *A. veronii* SJ4 was 4,562,694 bp, within a G+C content of 58.95%, containing 4079 coding genes. Nine hundred ten genes encoding for several virulence factors, such as type III and VI secretion systems, fagella, motility, etc*.*, were determined based on the VFDB database. Besides, 148 antibiotic resistance-related genes in 27 categories related to tetracyclines, fuoroquinolones, aminoglycosides, macrolides, chloramphenicol, and cephalosporins were also annotated. The present results suggested that *A. veronii* was etiological agent causing the bacterial septicemia of *S. chuatsi* in this time, as well as provided a valuable base for revealing pathogenesis and resistance mechanism of *A. veronii*.

Keywords *Aeromonas veronii* · *Siniperca chuatsi* · Pathogenesis · Resistance mechanism · Complete genome

Introduction

Siniperca chuatsi is one of the most economically important freshwater fsh species in China and has a high market value due to fast growth, high nutrition, and low fat (Shen et al. [2021\)](#page-15-0). The scale expansion of the mandarin fsh industry has resulted in a high annual production of 373.9 thousand tons according to China Fishery Statistical Yearbook in 2021. Recently, aquaculture has become highly sensitive to frequent pathogenic infections because of the high-density culture and the deterioration of water environment, which seriously threaten the steadiness and development of the fishery industry, and the bacterial infections percentage has risen to about 25% in freshwater fsh disease outbreaks (Tyagi et al. [2022](#page-15-1)). In recent years, various bacterial pathogens were frequently increasing and causing highly troublesome losses to *S. chuatsi* industry, including *Flavobacterium cloumnare*, *Aeromonas hydrophila*, *Gytophaga columnaris*, *Streptococcus uberis*, and *Aeromonas salmonicida* (Chen et al*.* [1995](#page-14-0); Zhou et al. [2015;](#page-15-2) Luo et al. [2017](#page-15-3); Chen et al. [2018;](#page-14-1) Lin et al. [2020](#page-15-4)). In this study, mass mortalities of *S. chuatsi* with bacterial septicemia occurred in Yangzhou, Jiangsu Province China, and *A. veronii* was identifed as the primary pathogen for the hemorrhagic septicemia in *S. chuatsi*.

A. veronii, a Gram-negative, rod-shaped, facultative anaerobic bacterium, has a strong adaptability to the environment and is widely distributed, especially freshwater areas and estuaries (Tekedar et al. [2019\)](#page-15-5). Among

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various bacterial pathogens, mesophilic *Aeromonas* species (*A. veronii*, *A. hydrophila*, *Aeromonas sobria*, etc.), causing motile *Aeromonas* septicemia (hemorrhagic septicemia), have often been reported as the major cause of fish mortalities around the globe (Janda and Abbott [2010](#page-14-2); Mishra et al. [2017](#page-15-6)). In recent years, *A. veronii* has been recognized as an aquatic pathogen to various fish species, such as *Ictalurus punctatus*, *Oreochromis niloticus*, *Dicentrachus labrax*, *Misgurnus anguillicaudatus*, *Carassius auratus*, *Labeo rohita*, *Odontobutis potamophila*, *Silurus asotus*, and *Astronotus ocellatus* (Huang et al. [2010;](#page-14-3) Sreedharan et al. [2011](#page-15-7); Kim et al. [2013;](#page-15-8) Dong et al. [2017;](#page-14-4) Smyrli et al. [2017](#page-15-9); Kaur et al. [2020;](#page-15-10) Shameena et al. [2020](#page-15-11); Zhang et al. [2020](#page-15-12); Liu et al. [2022a,](#page-15-13) [b\)](#page-15-14). It is visible that the cases of economic loss in aquaculture caused by the spread of *A. veronii* were rapidly increasing. Therefore, more attention should be attracted to the widespread infections and virulence mechanisms of *A. veronii* in aquatic animals.

As is well-known, the bacterial pathogenicity was based on many virulence-related factors, such as colonization, invasiveness, exotoxins, endotoxins, and extracellular enzymes (Tyagi et al. [2022](#page-15-1)). These virulence-related factors help the pathogen to colonize, invade, and attack the host, immune escape, proliferate, and cause disease (Sharma et al. [2017](#page-15-15)). Relative to single action, multiple virulence factors work synergistically, which is more beneficial for bacterial invasion (Haslinger et al*.* [2005\)](#page-14-5). Thus, the genes encoding for virulence factors have often been used to determine the potential pathogenicity of bacterial pathogens (Nailis et al*.* [2010](#page-15-16)). Now, with the update of next-generation sequencing technology (NGS) technologies and the development of utility tools, it has become more feasible to investigate whole genomes of various bacterial pathogens with high speed and accuracy. Thus, the whole genome sequencing (WGS) technology would play a crucial role into species identification, gene function annotation, and genome structure difference analysis of various pathogens (Donkor [2013\)](#page-14-6).

In the present study, *A. veronii* SJ4 was isolated from the kidneys of diseased *S. chuatsi*, and the pathogenicity of SJ4 was confirmedly determined by experimental infection, histopathology changes, and virulence gene examination. Additionally, the complete genome of *A. veronii* SJ4 was sequenced, and the virulence-related genes and antibiotic resistance genes were analyzed for revealing pathogenesis and resistance mechanism of *A. veronii* SJ4. The present results suggested that *A. veronii* were etiological agent causing the bacterial septicemia in *S. chuatsi* aquaculture of this time, and the whole genome sequencing analysis provided prediction for the pathogenesis and resistance mechanism of *A. veronii*.

Materials and Methods

Fish Disease and Bacterial Isolation

The diseased moribund *S. chuatsi* were collected from the aquaculture farm in Yangzhou, Jiangsu Province, China. The samples were examined for external clinical signs and aseptically dissected after sterilization of body surface with 75% ethanol. The liver, gill, kidney, and spleen samples were inoculated and streaked separately onto Luria–Bertani (LB) agar plates, which were cultured for 12 h at 28 °C. The dominant single colony was re-streaking on the LB agar slant, and the bacteria were stored in LB medium with 30% glycerol at−80 °C for further study.

Morphology Observation

The representative dominant strain SJ4 was selected and incubated in LB broth at microbiological incubator (180 rpm, 28 °C) for 18 h. The pure cultured SJ4 cells were collected by centrifuge at 6000 g, 4 °C for 10 min and washed thrice with sterile PBS. Then the bacteria cells were cooled at 4° C with glutaraldehyde (0.25 g/L), postfxed with osmium tetroxide in 0.1 M cacodylate bufer, and adhered onto poly-l-lysine coated coverslips. Then the graded ethanol was used for dehydration, and the gold palladium alloy was coated on EMS (Electron Microscopy Science). Finally, the Zeiss EM10 was used for the observation of the fagella cells, the types and sizes of which would be analyzed.

Identifcation of Bacterial Isolates

Thirty-three kinds of the biochemical reaction tubes (Hangzhou Binhe Microorganism Reagent Co., Ltd., China) were selected for the biochemical tests, including Gram staining, oxidase, voges–proskauer, indole production, sucrose, maltose, rafnose, lactose, melibiose, cellobiose, galactose, glucose, and mannitol, arabitol. The results of the tests would be compared with the description in Bergey's Manual of Systematic Bacteriology (Brenner et al. [2021\)](#page-14-7).

The *16S rRNA* and *gyrB* genes of SJ4 were PCR amplifed with primers as per previously described protocols of Zhang et al. ([2014](#page-15-17)). After sequencing in Shanghai Sangon Biotech Co., Ltd., the homology of the *16S rRNA* and *gyrB* sequences of isolate SJ4 were analyzed using the Basic Local Alignment Search Tool (BLAST) in NCBI. The maximum likelihood method was adopted to construct the phylogenetic trees using MEGA 7.0 (Kumar et al. [2016](#page-15-18)).

Bacterial Virulence Assay

After incubated in LB broth at microbiological incubator (180 rpm, 28 \degree C) for 18 h, the pure cultured SJ4 fluid was suspended with sterile PBS (pH 7.4) to the concentrations from 2.4×10^8 to 2.4×10^5 CFU/mL. Twenty *S. chuatsi* (100–120 g) confrmed to be free of pathogen infection in each tank were injected intraperitoneally with 100 µL bacteria within diferent concentrations $(2.4 \times 10^8, 2.4 \times 10^7, 2.4 \times 10^6, 2.4 \times 10^5 \text{ CFU/}$ mL) per fsh, respectively. In the control group, sterile PBS was injected intraperitoneally in each fsh. The survival situation of each fsh was monitored daily for 14 days (d), and the cumulative mortalities of experimental fsh were recorded to calculate the LD_{50} of *A. veronii* to *S. chuatsi* by using methods of Behreans and Karber [\(1953](#page-14-8)).

Histopathology

The tissue samples of the livers, spleens, kidneys, and gills were fxed in 10% neutral bufered formalin. The formalinfxed tissues were dehydrated in ethanol and embedded in paraffin wax blocks, which sectioned at 6 μ m and H&Estained for optical microscope observation (Pei et al. [2021](#page-15-19)).

Determination of Extracellular Enzymes and Hemolysin

The extracellular enzymatic activities including phospholipase, lipase, amylase, hemolysin, urease, caseinase, dnase, and gelatinase were screened by the methods described by Gao et al. ([2020](#page-14-9)). LB agar plates were supplemented with 10% egg yolk, 1% Tween-80, 2% starch, 7% rabbit erythrocytes, 2%

urea, 10% skim milk, 0.05% DNA, and 1% gelatin, respectively. Five microliters bacteria suspension was spot-inoculated onto the center of plates, and these plates were cultured at 28 °C for 18 h with triplicates. If the isolate SJ4 possesses the corresponding extracellular enzyme activities, the colonies would surround a circle of lytic halo.

Detection of Virulence‑Related Genes

The polymerase chain reaction was performed using Easy Taq PCR Super® Mix (Tolo Biotech Co., Ltd) to detect the presence of virulence-related genes in the isolate SJ4, including *act*, *fm*, *fgM*, *ompA*, *Lip*, *hly*, *aer*, and *EprCAI*. The specifc primers were designed according to the genome data (Table [1\)](#page-2-0). The PCR products were detected by 1% Agarose gel electrophoresis.

Whole‑Genome Sequencing and Assembly

For next-generation sequencing (NGS), SJ4 bacterial genomic DNA isolation kit (TransGen Biotech Co., Beijing) was used to extract the genomic DNA from the overnight grown culture of *A. veronii* SJ4. Agarose gel electrophoresis was also used to check the quality of genomic DNA. The whole genome of SJ4 was sequenced by the third-generation PacBio RS II combined with the second-generation Illumina HiSeq2000 sequencing method, which was completed by Shanghai OE Biotech Co., Ltd. Qualifed DNA samples were sequenced using PacBio RS II singlemolecule real-time (SMRT) sequencing technology (Pacifc Biosciences, Menlo Park, CA, USA) to have their entire genome sequenced. Then the hierarchical genome-assembly process (HGAP) was used to assemble randomly interrupted sequence

read segments to obtain very long sequence read segments. Next, the second-generation sequencing technology Illumina HiSeq 2000 was used for sequencing, and Bowtie2 (v2.3.0) was used to compare the Illumina gene fragments with the assembled SMRT-contigs, so as to obtain the complete circular whole genome sequence information. Glimmer 3.02 was used for gene prediction and annotation; tRNAscan-SE was used for prediction and screening of tRNA genes; RNAmmer software was used for prediction and analysis of rRNA; and Rfam (v10.0) was used for prediction and screening of sRNA. Subsequently, RepeatMasker (v4.0.7) software was used to predict the tandem repeat and interspersed repeat.

Genome Annotation and Comparative Genomics

In terms of genome annotation, several database annotations are commonly used, including NR annotation, COG/KOG functional annotation, GO classifcation, Swissprot, eggNOG, KEGG, and Pfam. DIAMOND software was employed for comparison of NR, COG/KOG, GO, Swissprot, eggNOG, and KEGG database annotations, and annotations with $E < 1E-5$ were obtained by selecting proteins that possess the highest sequence similarity. The Pfam database comprises a compilation of protein families, each represented by multiple sequence alignments and hidden Markov models. HMMER software was utilized to compare protein family models, and the family with the highest score was chosen. Moreover, the Blast function of DIAMOND software was used to compare with the CARD resistance gene database, and annotations with $E < 1E$ -10 were chosen. The VFDB database is specifcally designed for studying the pathogenic factors of pathogenic bacteria, chlamydia, and mycoplasma. Annotations with $E < 1E-10$ were selected via comparison with the VFDB database using DIAMOND software.

Results

Pathological Clinical Signs of Diseased Fish and Bacterial Isolation

According to the epidemiological investigation, the diseased *S. chuatsi* exhibited multiple symptoms including gill pallor, muscle bleeding and necrosis, fn erosion, as well as abdominal enlargement, redness and swelling (Fig. [1\)](#page-3-0). Anatomical examination revealed that internal organ damage caused the accumulation of yellow or red ascites in the abdominal cavity and led to organ enlargement, specifcally in the liver, spleen, and kidney. Abundant pure bacteria isolated from the livers, spleens, kidneys, and gills of the diseased *S. chuatsi*, and these colonies exhibited the characteristics of white, translucency, circle, convex, glaze surface, and intact edge. A representative strain from these was selected for this study, which was tentatively named SJ4.

Fig. 1 External clinical symptoms of experimentally infected *S. chuatsi*. GR gill rot, HS hemorrhage spot, H hemorrhage

Electron Microscopic Observation of the Isolate

The isolate SJ4 exhibited single polar fagellum and a shape of rod-shaped with round-ends, about 1.4–2.1 µm wide and 2.8–5.3 μ m long, upon the electron microscopic observation (Fig. [2](#page-4-0)).

Physiological and Biochemical Characterization

By physiological and biochemical observation, the isolate SJ4 showed similar characteristics of *A. veronii* bv *veronii* as the description in Bergey's Manual of Systematic Bacteriology. As shown in Table [2](#page-5-0), oxidase, voges–proskauer,

 $30. t i f$ $10:02:42$ a $05-05-17$ TEM Mode: Imaging

 500 nm $HV=80.0kV$ Direct Mag: 6000x AMT Camera System

Fig. 2 Electron micrograph of SJ4, bar = 0.5μ m.

 "+", positive; "−", negative; D, 11 89% positive with incubation at 35 ℃ for 7 days except for *A. veronii*, which were incubated at 25 ℃. "*" the data of *A. veronii* come from Bergey's Manual of Systematic Bacteriology

indole production, fructose, cellobiose, and β-galactosidase were produced, but not rafnose, lactose, xylose, arabitol, tartrate, and acetate. The sorbitol, amygdalin, and rhamnose of the isolate SJ4 showed positive activity, which exhibits different characteristics with *A. veronii* bv *veronii* described in Bergey's Manual of Systematic Bacteriology.

Molecular Identifcation

The *16S rRNA* gene and gyrase submit B gene of SJ4 were sequenced by Sengon Biotech and assembled by Seqman after PCR. The *16S rRNA* sequences of isolate SJ4 showed 99% sequence identities with *A. veronii* isolates in the NCBI Reference RNA Sequences (refseq rna) database (accession number: OQ826708), and the phylogenetic tree showed the isolate SJ4 belong to *A. veronii* (Fig. [3A](#page-6-0)). Besides, the gyrB sequences showed 98% sequence identities with *A. veronii* isolates in the NCBI Reference RNA Sequences (refseq rna) database (accession number: OQ845908), and the phylogenetic tree also showed the isolate SJ4 belong to *A. veronii* (Fig. [3](#page-6-0)B).

Virulence of the Isolate

The results of bacterial virulence assay were shown in Fig. [4.](#page-6-1) The *S. chuatsi* of experimental group started death from first day. The 2.4×10^8 , 2.4×10^7 , 2.4×10^6 , 2.4×10^5 , and 2.4×10^4 CFU/mL of SJ4 caused 100%, 100%, 70%, 50%, and 10% mortalities after 14 dpi, respectively, while all fsh of control group were still alive. The calculated LD_{50} of SJ4 to *S. chuatsi* was 3.8×10^5 CFU/mL. Furthermore, the isolate SJ4 was re-isolated from the infected *S. chuatsi*, confrming that this experiment satisfes Koch's postulates.

Histological Observation

Compared to the control group, the liver tissues of the experimental group presented with swelling, hemorrhage, necrosis, and interstitial hyperplasia among liver cells (Fig. [5B](#page-8-0)). Similarly, the respiratory epithelial cells of the secondary gill plate exhibited clear signs of necrosis with the bended branchial lamellae, while the epithelial cells exhibit proliferate and necrosis (Fig. [5](#page-8-0)D). Furthermore, the spleen tissues of the experimental group showed distinct manifestations of telangiectasia, hyperemia, hemolysis, and blood spot formation, with severe regional rupture (Fig. [5F](#page-8-0)). The focal area of the kidney displayed symptoms of loss of brush border, glomerular necrosis, and necrotic interrenal tissue cells with chapped morphology (Fig. [5H](#page-8-0)).

Virulence Factors of the Pathogenic Isolate

The cytolytic enterotoxin (*act*), fagellin (*fm*, *fgM*), outer membrane protein A (*ompA*), lipid A acyltransferase (*Lip*), hemolysin (*hly*), aerolysin (*aer*), and epsilon proteobacterial calcium-binding autotransporter-like protein (*eprCAL*) were detected by PCR detection. The extracellular enzymes activities of SJ4 were shown in Fig. [6](#page-8-1). The strain SJ4 could produce caseinase, dnase, gelatinase, and hemolysin activity.

Genome Structure and General Features of *A. Veronii* **SJ4 Genome**

The genome of *A. veronii* strain SJ4 was sequenced using Nanopore, and the genome characteristics are also

Fig. 3 (**A**) Phylogenetic tree of Aeromonas species based on 16S rRNA sequences. (**B**) Phylogenetic tree of Aeromonas species based on gyrB sequences.

summarized. The genome size of SJ4 was determined to be 4,562,694 bp with a GC content of 58.95%. A total of 4079 coding genes were identifed, with a combined length of 3,973,236 bp. In addition, 229 simple repetitive sequences were found, totaling 10,430 bp and representing 0.23% of the genome size. as shown in Fig. [7,](#page-9-0) the genome also contained 124 tRNA genes, 43 sRNA genes, and 31 rRNA genes.

COG Analysis

The protein-coding genes with biological functions in the genome of experimental strain SJ4 were annotated by

comparing them with the COG database. Based on the COG classifcation standards, these genes were categorized into 25 classes represented by uppercase English letters (A–Z), and a homologous gene annotation was performed. A total of 3271 protein-coding genes were annotated. As shown in Fig. [8,](#page-10-0) the annotation results for the basic function prediction genes were the most abundant, with 456 genes accounting for 13.94% of the annotated genes. The next most abundant category was amino acid transport and metabolism, with 371 genes accounting for 11.34% of the annotated genes. Signal transduction mechanisms (287 genes; 8.77%), transcription (284 genes; 8.68%), and energy production and conversion

Fig. 4 The survival rates of largemouth bass challenged by SJ4

Fig. 5 Histological changes of S. chuatsi infected by the isolate SJ4. ◂ CV represents Swollen Central Vein; LN represents liver nucleus; CL represents cytoplasmic lysis; IH represents interstitial hyperplasia; BF represents branchial flament; SG represents the secondary gill lamellae; SN represents spleen nucleus; MC represents melanomacrophage center; RT represents rupture; GR represents renal tubules; CT represents connective tissue; FN represents focal necrosis

(240 genes; 7.34%) were also well-annotated. In addition, 299 genes (9.14%) with unknown functions require further investigation in future studies.

GO Analysis

The protein sequences of the predicted genes were compared with the GO database by BLASTp to obtain GO annotation information. The DIAMOND software was used for comparison, and proteins with the highest sequence similarity were selected for GO functional clustering analysis. A total of 2945 protein-coding genes were successfully annotated with GO functions. The GO annotation includes three aspects: biological processes, cellular components, and molecular functions. As shown in Fig. [9](#page-10-1), in terms of biological processes, the annotated genes were mainly related to cellular processes, metabolic processes, and localization. In terms of cellular components, the annotated genes were mainly related to cells, cell parts, and membranes. In terms of molecular functions, the annotated genes were mainly related to catalytic activity, binding, and transporter activity.

KEGG Analysis

In this study, we compared the genome of *Aeromonas* sp. SJ4 with the KEGG database and found that 2461 genes could be mapped to 165 metabolic pathways, accounting for 60.33% of all genes. As shown in Fig. [10](#page-11-0), the metabolic pathway-related genes could be classifed into 29 categories and 5 major classes, namely, cellular processes, environmental information processing, genetic information processing, metabolism, and organismal systems. KEGG enrichment analysis revealed that the strain was mostly involved in metabolism pathways, followed by environmental information processing.

Prediction of Virulence Genes of the Strains SJ4

By comparison using the virulence factor database (VFDB), it was found that there are 910 coding sequences that may be virulence genes in the whole genome of the pre-test strain SJ4. Among them, most of the genes are related to secretion systems, toxins, two-component systems, superoxide dismutase, etc. (Table [3\)](#page-11-1). The experimental strain SJ4 has type III and type VI secretion systems, and the efector factors *lpg* (gene0212, gene1207, gene1209, gene1211) of the Dot/Icm type IV secretion system, the efector factor *LirB* (gene3402) of the type IV secretion system, the ABC transporter protein gene *Pmen_2312* (gene1763), and other genes are involved in the formation and regulation of the type IV secretion system, while

Fig. 6 The extracellular enzyme test results of *A. veronii* SJ4

Fig. 8 The Clusters of Orthologous Genes (COG) functional annotation in the whole genome of *A. veronii* SJ4

the efector factor *VPA0450* (gene1674), the ATPase *hrcN* (gene2447), *ssaN* (gene2488), and other genes are involved in the formation and regulation of the type III secretion system. Genes such as the exfoliative toxin gene *eta* (gene0151), the toxin co-regulated pilus biogenesis protein gene *tcpI* (gene0280, gene0886, gene1105, gene2909), the structural toxin protein gene *rtxA* (gene0808), the aerosol toxin gene *aerA* (gene2830), and the enterotoxin gene *act* (gene2830) are

Fig. 9 Gene Oncology (GO) functional annotation in the whole genome of *A. veronii* SJ4

Fig. 10 Gene distribution based on the Kyoto Encyclopedia of Genes and Genomes (KEGG) classifcation of *A. veronii* SJ4

related to the virulence of bacteria. The encoded proteins are important proteins involved in carbohydrate and amino acid metabolism and also have a certain impact on the expression of bacterial fagella and pili.

Analysis of Antibiotic Resistance

To further investigate the drug-resistant characteristics of strain SJ4, the main drug-resistant genes of SJ4 were predicted (Table [4](#page-12-0)). The results showed that the SJ4 genome contains drug-resistant genes related to tetracyclines, fuoroquinolones, aminoglycosides, macrolides, chloramphenicol, and cephalosporins (Fig. [11\)](#page-12-1). These genes mainly confer resistance to antibiotics through mechanisms such as antibiotic target alteration, antibiotic target replacement, reduced permeability to antibiotics, antibiotic inactivation, and antibiotic efflux.

Discussion

As a common opportunistic pathogen in freshwater fsh in China, *A. veronii* has been recognized as one of the major pathogens in the aquaculture industry (Ran et al. [2018](#page-15-22)). Recently, it has been rapidly increasing in the cases of

Table 3 The annotation of virulence factors of the strain SJ4 in VFDB databases

Table 4 Physiological and biochemical characteristics of strain SJ4

economic fsh death, exhibiting pathological signs such as ulceration, organ hemorrhage, and severe ascites, as well as abdominal enlargement and redness (Zhu et al. [2022](#page-16-0)). In this study, *A. veronii* SJ4 was isolated from diseased *S. chuatsi*, and the major pathological symptoms exhibit gill pallor, muscle bleeding and necrosis, fn erosion, as well as abdominal enlargement, redness and swelling. Experimental infection in vivo showed that the LD₅₀ of *A. veronii* SJ4 to *S. chuatsi* was 3.8×10^5 CFU/mL and the *S. chuatsi*

infected by *A. veronii* developed similar clinical signs to the naturally infected fsh, suggesting that the isolate SJ4 has the high virulence to *S. chuatsi*. Previous studies show that *A. veronii* has been confrmed to be the pathogen of ulcerative disease. Wang et al. ([2021\)](#page-15-23) reported that the *A. veronii* strain 18BJ181 which was frstly isolated from diseased *Lateolabrax maculatus*, caused acute death in aquaculture, and the LD₅₀ of the 18BJ181 to *L. maculatus* in 10 days was 8.5×10^5 and 4.2×10^5 CFU/g under the

Fig. 11 The CARD database annotation in the whole genome of *A. veronii* SJ4. Note: The diferent sizes of fan-shaped sections in various colors indicate the proportion of genes annotated to the corresponding term, while the numbers following the legend located in the upper left corner represent the total number of genes annotated to the corresponding term

Functional Categories

conditions of intraperitoneal injection and intramuscular injection, respectively. Hassan et al. [\(2017\)](#page-14-10) reported the *A. veronii* to be the most predominant pathogen of *O. niloticus*. Zhu et al. ([2022\)](#page-16-0) reported that the *A. veronii* strain GJL1 could cause the mass mortalities of *M. salmoides*, and the LD_{50} of the GJL1 to *M. salmoides* in 10 days was 3.6×10^5 CFU/mL.

It is well known of *Aeromonas* spp. to possess several virulence factors, such as adhesion factor, fagellum, exotoxins, endotoxins, and extracellular enzymes (Castro-Escarpulli et al*.* [2003](#page-14-11)). Previous studies have demonstrated the crucial role of extracellular products (ECPs) in the pathogenicmechanism of bacteria pathogens (Sharma et al. [2017](#page-15-15)). ECPs produced by *Aeromonas* genus bacteria, including *A. veronii* and *A. hydrophila* have been proved to cause huge damage to organisms (González-Serrano et al*.* [2002\)](#page-14-12). The haemolysin and aerolysin proteins promote the process of pathogen invasion in host, while lipases catalyze the hydrolysis of membrane lipids, leading to intestinal damage (Sughra et al. [2022](#page-15-24)). Through these virulence factors, the bacteria pathogen adheres to the surface of host tissue cells, disrupts the immune system, and hence starts colonization. In this study, it was observed that *A. veronii* SJ4 exhibited a dense coverage of long fmbriae upon microscopic observation. This fnding implies that *A. veronii* SJ4 may possess a high level of both adhesion and virulence. *A. veronii* SJ4 has demonstrated various enzymatic activities such as caseinase, Dnase, gelatinase, and hemolysin, which may facilitate the bacterium's ability of invading the host. Moreover, the presence of virulence genes including *act*, *fm*, *fgM*, *ompA*, *lip*, *hly*, *aer*, and *eprCAL* plays a crucial role in protein code and toxin secretion, which contribute to the pathogenicity of *A. veronii* SJ4. The various toxins and enzymes expressed by these genes have the potential to infict damage on host cells, leading to their deterioration.

Whole genome sequencing technology plays a critical role in molecular epidemiology research of pathogenic bacteria, as it reveals the genomic characteristics, infers transmission routes and epidemiological features, and aids in infection control and prevention (De Maio et al*.* [2019\)](#page-14-13). Zhou et al. [\(2022\)](#page-15-25) used whole-genome sequencing to reveal the virulence and resistance mechanisms of non-O1 Vibrio cholerae, the pathogen of shrimp disease. Pang et al. ([2015\)](#page-15-26) conducted wholegenome sequencing analysis of *A. hydrophila*, identifying three metabolic pathways unique to the epidemic strain. Kang et al. [\(2016](#page-15-27)) sequenced the whole genome of the highly virulent *A. veronii* strain TH0426 and also determined the genome framework of the weakly virulent strain AV161 and the nonpathogenic strain CL8155, revealing the diferences among *A. veronii* strains with strong, weak, and no virulence at the genome level through comparative analysis. In present study, the complete genome nucleotide sequence of *A. veronii* SJ4 was determined with a total of 4079 predicted genes, among a GC content of 58.95% and 124 tRNA genes, 43 sRNA genes, and 31 rRNA genes. A considerable quantity of genes were identifed and annotated within the NR, COG/KOG, GO, Swissprot, eggNOG, KEGG, and Pfam pathways. Notably, these pathways have been previously linked to various human diseases. Additionally, earlier investigations have demonstrated that *A. veronii* can induce numerous maladies, including toxemia, diarrhea, and endocarditis, among others. This evidence underscores the potential risk that *A. veronii* strain SJ4 may pose to human health (Liu et al. [2022a](#page-15-13), [b\)](#page-15-14).

The pathogenesis of *Aeromonas* infection is a complex process involving multiple factors. The initial stages of bacterial pathogenicity entail movement, invasiveness, and colonization. Specifcally, the single polar fagellum allows the *Aeromonas* genus bacteria to swim in liquid environments, and the extra lateral fagellar system contributed to the collective motions and bioflm formation of 60% *Aeromonas* genus bacteria (Fernández-Bravo and Figueras [2020](#page-14-14)). In the present study, a total of about 128 genes encoding for fagellar component protein were determined in the *A. veronii* SJ4 genome. Besides several fagellar related genes, many virulence-related gene encoding for extracellular toxins and enzymes have a potential virulence property to the *Aeromonas* genus. The genes encoding for caseinase, dnase, gelatinase, and hemolysin were also determined in *A. veronii* SJ4 genome, which is consistent with the phenotypic results of extracellular enzyme activity. Hemolysin and aerolysin play a critical role in host tissue damage, bacterial invasiveness and colonization, as well as the suppression of immune system, which contributed to the phenotypic symptom of hemorrhagic septicemia. Additionally, the presence rate of hemolysin, aerolysin, lipases, and serine proteinase is remarkably high among *Aeromonas* isolates derived from diseased fsh (Tyagi et al. [2022\)](#page-15-1). *A. veronii* SJ4 genome also carried 15, 15, 6, 5, 7 genes involved in T2SS T3SS T4SS T6SS T7SS biosynthesis, respectively. Bacteria use T2SS to release cytoplasmic proteins, including virulence factors like hemolysin, aerolysin, caseinase, and dnase in *Aeromonas* (Matys et al. [2020\)](#page-15-28). The bacterial effector proteins could directly invade into the host cell with a needle-like structure created by T3SS, which disrupt signaling pathways and cytoskeleton, as well as induce apoptosis (Zeb et al. [2019](#page-15-29)). T4SS transports DNA, proteins, and other molecules through multiple proteins, while T6SS indirectly interacts with host cells through peptidoglycan-like efectors, such as Vibrio's T6SS uses toxic efectors with an actin cross-linking domain to directly act on eukaryotic cells (Wang et al. [2016\)](#page-15-30).

Multiple antibiotic resistance genes related to antibiotic target alteration, antibiotic target replacement, reduced permeability to antibiotics, antibiotic inactivation, and antibiotic efflux were also determined in the *A. veronii* SJ4 genome. However, it is extremely complicated of resistance mechanisms in pathogenic bacteria that the bacteria could control the antibiotic content in

the cells through complex biochemical pathways, or change/ degrade the antibiotics through enzymatic action. Moreover, the bacteria gene mutation may also infect the efectiveness of the antibiotics (Miller et al*.* [2014](#page-15-31)). Thus, it is not inevitably that antibiotic resistance genes could be transformed into antibiotic resistance phenotype. In the present study, despite that many antibiotic-related genes (fuoroquinolone, penicillin tetracycline, etc*.*) were determined in the genome of *A. veronii* SJ4, it was observed that no resistance against some tested antibiotics (forfenicol, ofoxacin, enrofoxacin, etc*.*) in antibiotics susceptibility testing (the supplementary table 1). The observed phenomenon may be attributed to either non-functionality of the efflux pump in *A. veronii* SJ4, the absence of antibiotic gene expression, or the dysfunction of the enzyme itself (Martínez and Baquero [2002\)](#page-15-32).

Conclusion

In summary, the *A. veronii* SJ4 was identifed as a high virulence pathogen of causing the bacterial septicemia and mass mortalities of *S. chuatsi* in this time. After whole genome sequencing, many virulence-related genes and antibiotic resistance genes are annotated in the database. The present study provides theoretical basis to clarify characteristics, pathogenicity, antibiotic resistance, and virulence factors of highly pathogenic *A. veronii* at the molecular level, and the whole genome sequencing analysis provided prediction for the pathogenesis and resistance mechanism of *A. veronii*.

Author Contribution X.H. Zhu fnished writing—original draft, methodology, conceptualization, and data curation. L.J. Qin fnished methodology and data curation. Q.Q. Qian fnished methodology, data curation. Y.J. Zhu finished methodology, data curation. X.J. Gao finished writingreview. Qun Jiang fnished methodology. Jun Wang fnished data curation. Guoxing Liu supported funding acquisition. Xiaojun Zhang fnished the writing—review, conceptualization and funding acquisition. All authors reviewed the manuscript.

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Data Availability The datasets generated during and/or analyzed during the current study are available from the corresponding author on reasonable request.

Declarations

Ethics Approval All treatments involving animals were carried out under the strict guidelines of Animal Experiment Ethics Committee of Yangzhou University. (Permit number: 201802003).

Competing Interests The authors declare no competing interests.

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