Quorum quenching bacteria *Bacillus* sp. QSI-1 protect zebrafish (Danio rerio) from *Aeromonas hydrophila* infection

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Quorum Sensing (QS) is a bacterial regulatory mechanism, which is responsible for controlling the expression of various biological macromolecules such as the virulence factors in a cell density-dependent manner. Disruption of the QS system of pathogens has been proposed as a new anti-infective strategy. Biodegradation of AHLs proves to be an efficient way to interrupt QS, since AHLs are the main family of QS autoinducers used in Gram negative bacteria. In this study, the effect of *Bacillus* sp. QSI-1 as an efficient quorum quencher on virulence factors production and biofilm formation of fish pathogen *Aeromonas hydrophila* was investigated. QSI-1 reduced the accumulation of AHLs but did not affect the growth of *A. hydrophila* YJ-1 when cocultured. In the result, the supernatant of QSI-1 showed significant inhibition of protease production (83.9%), hemolytic activity (77.6%) and biofilm formation (77.3%) in YJ-1. In biocontrol experiment, QSI-1 significantly reduced the pathogenicity of *A. hydrophila* strain YJ-1 in zebrafish (Danio rerio). The fish fed with QSI-1 was observed to have a relative percentage survival of 80.8%. Our results indicate that AHLs degrading bacteria should be considered as an alternative for antibiotics in aquaculture for the biocontrol of bacterial fish diseases.

Antibiotics are critical in preventing, controlling, and treating disease in farm animals. By preventing essential processes such as cell wall synthesis, DNA proliferation, and protein synthesis, antibiotics become an integral part in keeping our food supplies safe from infectious bacteria. Unfortunately, the widespread use of antibiotics eventually lead to the selection of bacteria which is resistant to the antibiotic. These antibiotic-resistant bacteria population render the antibiotics ineffective, and new antibiotics are needed. In order to combat pathogenic bacteria without fear of developing an antibiotic resistant strain, alternative methods need to be investigated.

Quorum sensing (QS) is a mechanism in which bacteria coordinates gene expressions in response to their population density by producing, releasing, and recognizing small signal molecules called autoinducers. QS regulates various phenotypes such as biofilm formation, bioluminescence, virulence factors, and swarming, which have been shown it’s contribute to bacterial pathogenesis. Since pathogenicity traits of bacteria are controlled by QS, disruption of quorum sensing QS has been suggested as a new anti-infective strategy to control pathogenic bacteria, without interfering with their growth, in the field of aquaculture and animal husbandry. Quorum quenching (QQ), the disruption of QS, can be performed by small molecule antagonists or signal degrading enzymes. Many microorganisms can produce enzymes which can degrade N-Acyl-homoserine lactones (AHLs). The QQ enzymes produced by microorganisms were classified into three major types according to their enzymatic mechanisms: AHL lactonase (lactone hydrolisis), AHL acylase (amidohydrolisis) and AHL oxidase and reductase (oxido/lipoxygenation). These enzymes can degrade AHLs, and as a consequence, prevent pathogenic bacteria from producing virulence factors, forming biofilms, and reducing virulence. So the QQ microorganisms can be used as potential quenchers of quorum-sensing-regulated functions in pathogenic bacteria. Therefore it could act as an alternative to antibiotics.

In the present study, we investigate the previously isolated bacterium with quorum quenching activity, *Bacillus* sp. QSI-1, at degrading AHL signals of *A. hydrophila* and interrupting of its QS functions. Furthermore, we investigate the effect of QSI-1 quorum quenching activity on the pathogenicity of *A. hydrophila* in zebrafish. The results demonstrate its potential as an environmentally friendly alternative to antibiotics in aquaculture.
**Results**

**AHL inactivation assay.** Inactivation of C6-HSL was confirmed with the loss of purple pigment using the resting cells of QSI-1 at different time intervals at (0, 6, 12, 24 hours) and at different concentrations of (4 µl, 6 µl, 8 µl, 10 µl) by using *C. violaceum* CV026 as bio-reporter strain. Our results suggested that QSI-1 showed high AHL degrading ability even at 0.1 µg/µl C6-HSL within 6 hrs and no AHL molecules were found after further incubation to 24 h (Fig. 1). Disappearance of purple pigmentation production was also observed with the quorum quenching activities of QSI-1 supernatant on CV026 lawn. Nearly all C6-HSL was degraded after 8 hours of incubation. Our results showed that QSI-1 can degrade AHLS and at the same time, the metabolites of QSI-1 also can degrade AHL.

*Bacillus* sp. QSI-1 blocked AHL signal in *A. hydrophila* accumulation but did not affect its growth. From antagonistic assay, the *Bacillus* sp. QSI-1 did not have any negative effect on YJ-1, since YJ-1 grew properly in presence of QSI-1 filtered supernatant (Fig. 2). To test the effect of *Bacillus* sp. QSI-1 on AHL accumulation and growth of *A. hydrophila* YJ-1, YJ-1 was cocultured with or without *Bacillus* sp. strain QSI-1. AHL autoinducer was evaluated on LB agar, again using CV026 as a biosensor. In single culture, AHL produced by strain YJ-1 was detectable 3 h after inoculation, and a rapid increase was observed between 8 to 10 h after incubation, parrelling to the exponential proliferation stage of the bacterial cells (Table 1). However, no AHL was detected in the culture supernatant of YJ-1 co-cultured with QSI-1, which produce AHL-lactonase. In the competition assay, when *A. hydrophila* was grown with or without *Bacillus* sp. QSI-1, the obtained results showed no considerable variation in values of bacterial growth curve of both QSI-1-treated and untreated bacterial cultures of *A. hydrophila* YJ-1. This indicates that, *Bacillus* sp. had no effect on the growth of YJ-1 (Fig. 3).

**Discussion**

It has been demonstrated that the QQ enzymes could reduce the virulence of pathogenic bacteria. Zhang, et. al., expressed the aiiO-AIO6 in *Escherichia coli*. They also studied the effects of purified enzyme solution on the expression of virulence factors. Their results showed that AiiO-AIO6 could inhibit *A. hydrophila* virulence factors.
expression29. Cao et al., supplied the expressed AiiAA196 into fish feed and found that the oral administration of AiiAA196 significantly decreased the mortality of infected zebrafish29. In the present investigation, we showed the quorum quenching strain Bacillus sp. QSI-1 which originally isolated from the gut of health Carassius auratus gibelio, can inhibit the AHL-dependent violacein production in C. violaceum and virulence factors such as protease, hemolysins production and biofilm formation in A. hydrophila YJ-1. When QSI-1 cocultured with YJ-1, there is no effect on the growth of YJ-1, but the production of quorum sensing molecules was decreased compared with the mono-culture of YJ-1. Similar to Zhang et al.’s results, the total proteolytic, hemolytic activity and biofilm formation of YJ-1 were significantly decreased when added with the supernatant of QSI-1. In our protection ability experiments, QSI-1 was supplemented into experimental diets to control A. hydrophila infection. The results showed that the oral administration of QSI-1 significantly decreased the mortality of infected zebrafish. This result may provide a new explanation of modes of action of probiotics.

In summary, in this study we demonstrated yet another pharmacologically important property of bacterial resources against bacterial QS systems and a new way to combat bacterial diseases instead of antibiotics in aquaculture. The results of the present study clearly show a significant decrease in the QS-mediated phenotypic expression of the target bacterial pathogens without any reduction in its growth. And fish experiments also showed that the QQ bacteria can increase the survival rate. Owing to the importance of QS during bacterial pathogenesis, isolated compounds from Bacillus sp. QSI-1 will be a promising tool to human health as they prevent the onset of bacterial diseases as anti-pathogenic drugs, which is not subject to antibiotic resistance.

Methods

Bacterial strains and culture conditions. The quorum quenching strain Bacillus sp. QSI-1 which was isolated from the intestine gut of health Carassius auratus gibelio was cultivated in Luria-Bertani (LB) broth consisting of 1% tryptone, 0.5% yeast extract and 0.5% NaCl (pH 7.2 ± 0.2), and incubated for 24 h at 30°C. The cell-free supernatant from QSI-1 was obtained by centrifugation at 12,000 rpm, the supernatant filtered through 0.22 μm membrane filter. The filtered supernatant was then extracted with an equal volume of ethyl acetate; the solvent layer was separated and evaporated to dryness. The dried extract was weighed and used for anti-QS bioassays. The target pathogen A. hydrophila YJ-1 used in this study was isolated from a diseased fish. It was cultivated in Luria-Bertani (LB) broth (pH 7.2 ± 0.2) at 150 rpm in a shaker overnight at 28°C. CV026, a mini-Tn5 mutant derived from a Chromobacterium violaceum strain30 was used as an AHL-reporter to detect the residual AHL concentration in the rearing water. This mutant cannot produce AHL, but it can detect and respond to a range of AHL molecules (with acyl chain of four to eight carbons), by synthesis of the purple pigment violacein. CV026 was grown in LB at 30°C. This medium was supplemented with 20 mg l−1 of kanamycin.

AHL inactivation assay. AHL inactivation assays was conducted as described by Chan et al.31. QSI-1 was grown overnight at 30°C with shaking (220 rpm) in LB medium to approximately 108 cfu/ml, cells (100 μl) were collected by centrifugation, washed and resuspended in 10 ml of PBS (100 mM, pH 6.5) equalized to an OD of 1.0 at 600 nm and used directly as a source of resting cells for in vitro AHL inactivation assays. Aliquots of C6-HSL (10 μg/μl) (Sigma-Aldrich, St. Louis, Missouri, USA) in absolute ethanol were dispensed into sterile tubes and the solvent evaporated to dryness under sterile condition. The resting cells were used to rehydrate the C6-HSL to a final concentration of 0.1 μg/μl. The mixtures were incubated at 37°C for 4.5 h with gentle shaking in a hybridization oven. Heat-denatured suspension (10 μl) was inoculated onto LB agar seeded with the bio-reporter CV026 and incubated at 28°C. E. coli strain DH5α served as negative controls. Disappearance of C6-HSL from the mixture was assessed at different time using CV026 bioreporter. Degradation of C6-HSL is evident by loss of purple pigmentation shown by C. violaceum and the results were digitally recorded.

For AHLs activity in the supernatant produced by A. hydrophila, the well-diffusion assay was used. Ten milliliters of warm molten soft top LB agar (0.6%) agar was seeded with 100 μl of an overnight CV026 culture. This was gently mixed and poured immediately over the surface of a solidified LBA plate as an overlay. Wells of 5 mm in diameter were made on each plate after the overlay had solidified. Each well was filled with 50 μl of filter sterilized culture extract.

The plates were incubated for 48 h at 28°C and the diameter of pigment production by the monitor strain measured.

In vitro competition assay between Bacillus sp. QSI-1 and A. hydrophila. This experiment was done in two different steps; First the antagonistic activity of Bacillus sp. QSI-1 against A. hydrophila YJ-1 was evaluated. LB broth cultures were prepared freshly from the A. hydrophila agar slopes (stored at 4°C) and were incubated on LB agar plates (diameter, 98 mm) separately using a lawn culture technique32. Then, a 6-mm-diameter well was made in each plate with the help of a cork borer. A 100-μl aliquot of cell-free supernatant was charged in the well and incubated at 37°C for 24 h. The zones of inhibition around the well was recorded (in millimetres) after the incubation. Control plates were simultaneously maintained with sterile phosphate buffer saline (pH 7.2) poured into respective well prepared as mentioned above. Triplicate plates were maintained along with the PBS control for the antagonistic activity study.

In the second step in order to investigate the effect of QSI-1 on the growth of A. hydrophila YJ-1 was evaluated in mixed liquid cultures. Erlenmeyer flasks containing...
100 ml of LB were inoculated with 100 μl of independent overnight cultures of the Bacillus sp. strain QSI-1 and A. hydrophila YJ-1. The turbidity of the overnight cultures was measured at 600 nm and adjusted with sterile LB medium to obtain inoculums with the same cell ratio. The inoculated flasks were incubated at 30°C with shaking at 150 rpms in an orbital incubator. Samples were collected after 0, 3, 5, 8, 10 and 24 hours. For cells detection, 500 μl of culture were removed and after proper dilutions in PBS, all cultures were plated on LB agar plates. After 16–18 h cultured at 30°C, A. hydrophila and Bacillus sp. strain QSI-1 colonies were easily distinguishable based on their unique colony morphologies (Fig. 7). The experiment was repeated four times.

Effect of supernatant of QSI-1 on virulence factors production and biofilm formation in A. hydrophila. A. hydrophila YJ-1 cultures was grown for 24 hrs to O.D600 of 2.0 (for the protease, hemolytic activity and biofilm formation assays) treated with 200 μg/ml (w/v), Bacillus sp. QSI-1 supernatant in 2 ml fresh LB medium. Cells were harvested by centrifugation and the supernatant were filter sterilized. The supernatant were either used immediately or stored at −20°C.

Proteolytic activity was determined by azocasein assay by following the method of Swift et al.32 with slight modifications. Briefly, 150 μl of both treated and untreated YJ-1 culture supernatants was added to 1 ml of 0.3% azocasein (Sigma, St. Louis, USA) in 0.05 M Tris-HCl and 0.5 mM CaCl₂ (pH 7.5), and incubated at 37°C for 15 min respectively. The reaction was stopped by the addition of trichloroacetic acid (10%, 0.5 ml) followed by centrifugation, and the absorbance of clear supernatant was measured at 400 nm in a UV–visible spectrophotometer.

Blood agar, for detecting secreted hemolytic activity, was prepared by adding 5% washed erythrocytes to LBA33. Hemolysis activity in liquid assay was determined as described previously34. Briefly, serial dilutions of hemolytic samples were prepared with phosphate-buffered saline (PBS).100 ml aliquots of each dilution were incubated with 1 ml of erythrocyte suspension for 15 min at 25°C and then centrifuged for 1 min. The fluorescence absorbance at 405 nm (A405) of released hemoglobin was measured. Hemolytic activities are presented as the percentage of the total erythrocytes lysed (percent hemolysis). The percent hemolysis is defined as [(A405 for the
sample with hemolysin = A_{405} (for the control without hemolysin) \times 100/(A_{405} for the total lysis caused by sodium dodecylsulfate = A_{405} for the control).

Standard Tube Method (STM) was used for quantitative analysis of biofilm production. Briefly, overnight bacterial cultures were inoculated into 5 ml LB medium with 1:1000 dilution and incubated in the boricolite glass tubes without shaking at 28°C for 48 h. After the bacterial cultures were poured out, the tubes were washed extensively with water, then fixed with 2.5% glutaraldehyde, followed by washing alternatively with water, and staining with 0.4% crystal violet solution. After solubilization of the crystal violet with ethanol-acetone (80:20, vol/vol), the absorbance at 590 nm was determined. The experiment was performed in triplicate and repeated three times.

Fishextperiment. Wild-type zebrafish were brought from an ornamental fish farm in Nanjing, Jiangsu Province, and are allowed to acclimate for at least 3 days before use. Basal diet was prepared as mentioned elsewhere. For 50% lethal dose (LD50) determinations, seven groups of 10 Wild-type zebrafish (4 months of age, with an average weight of \sim 300 mg and an average length of \sim 2.5 cm) intraperitoneally injected with 0.02 ml of serially tenfold diluted bacterial suspensions containing 10¹–10⁸ CFU. A control group was injected intraperitoneally with 0.02 ml sterile PBS only. The fish were monitored at 28°C for mortality for 7 days. During this period, activity and behavior of fish were recorded daily, and LD50 were calculated by the method of Reed and Muench.

For QSI-1 contained diet, the QSI-1 was cultured for 24 h at room temperature (27 ± 2°C) in LB medium, and cells were harvested by centrifugation at 10,000 rpm 4°C for 20 min. The pellets were resuspended in PBS at the concentration of 10¹⁰ cfu/ml and mixed with feed to contain the final count of 10¹⁰ cfu/g of feed and sun dried. Prior to the experiment, the amount of diet consumed per fish was monitored for 7 days, and based on the data, the QSI-1 amount in the experimental feed was adjusted approximately 4.0 × 10⁶ cells per 5 g fish/day. Control fish were fed with the dry diet without QSI-1 bacterial culture.

Before challenge, all fish were clinically healthy and there was no evidence of A. hydrophila infection, nor was there any death due to A. hydrophila infection. Control and QSI-1 infected fish (30 per group) were challenged via intramuscular injection with 0.1 ml PBS or 6.0 × 10⁶ cells (5 × LD₅₀ of A. hydrophila) Y-1 in 0.1 ml of PBS. The fish were observed daily for 2 weeks after the challenge. Fish were observed to determine survival rate, and formation of ulcers and hemorrhagic lesions on the skin. Observation was made for a period of 2 weeks and data on mortality were calculated according to Amend.22 Bacterial isolation was performed by cultivation from hemolysin and ulcerous lesions, and from visceral organs of dead fish. This was also done in all surviving fish. Animal experiments were carried out in compliance with the National Institutes of Health Guide for the Care and Use of Laboratory Animals and approved by IACUC (Institutional Animal Care and Use Committee of China Pharmaceutical University).

Data analysis. Data are presented as mean ± SD of the number of fish per group. Data were statistically processed for one-way analysis of variance (ANOVA) to find out any significant differences among the experimental groups and the comparison between two treatments were done. Significant differences were determined at P < 0.05.

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**Additional information**

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