**N-acylhomoserine lactonase-based hybrid nanoflowers: a novel and practical strategy to control plant bacterial diseases**

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**Abstract**

**Background:** The disease caused by plant pathogenic bacteria in the production, transportation, and storage of many crops has brought huge losses to agricultural production. N-acylhomoserine lactonases (AHLases) can quench quorum-sensing (QS) by hydrolyzing acylhomoserine lactones (AHLs), which makes them the promising candidates for controlling infections of QS-dependent pathogenic bacteria. Although many AHLases have been isolated and considered as a potentially effective preventive and therapeutic agents for bacterial diseases, the intrinsically poor ambient stability has seriously restricted its application.

**Results:** Herein, we showed that a spheroid enzyme-based hybrid nanoflower (EHNF), AhlX@Ni3(PO4)2, can be easily synthesized, and it exhibited 10 times AHL (3OC8-HSL) degradation activity than that with free AhlX (a thermostable AHL lactonase). In addition, it showed intriguing stability even at the working concentration, and retained ~100% activity after incubation at room temperature (25 °C) for 40 days and approximately 80% activity after incubation at 60 °C for 48 h. Furthermore, it exhibited better organic solvent tolerance and long-term stability in a complicated ecological environment than that of AhlX. To reduce the cost and streamline production processes, CSA@Ni3(PO4)2, which was assembled from the crude supernatants of AhlX and Ni3(PO4)2, was synthesized. Both AhlX@Ni3(PO4)2 and CSA@Ni3(PO4)2 efficiently attenuated pathogenic bacterial infection.

**Conclusions:** In this study, we have developed N-acylhomoserine lactonase-based hybrid nanoflowers as a novel and efficient biocontrol reagent with significant control effect, outstanding environmental adaptability and tolerance. It was expected to overcome the bottlenecks of poor stability and limited environmental tolerance that have existed for over two decades and pioneered the practical application of EHNFs in the field of biological control.

**Keywords:** Biocontrol, Quorum sensing and quenching, AHL lactonases, Enzyme-based hybrid nanoflower, *Erwinia carotovora*, *Burkholderia glumae*, Robustness
Introduction

Plant bacterial diseases severely decrease the global crop yield each year. Treating plant bacterial diseases with chemical bactericides is effective but causes many environmental problems [1, 2]. The pathogenicity of many Gram-negative bacteria is coordinated by the quorum sensing (QS) system and is based on diffusible N-acylhomoserine lactones (AHLs) [3]. Soft rot from crops, such as potato, cabbage, radish, garlic, onion, cucumber, carrot, eggplant, squash, and tomato, is caused by Erwinia carotovora due to the production of exoenzymes (protease, cellulases and pectinases) in response to 3-oxohexanoyl-L-homoserine lactone [4, 5]. Seedling and grain rot from rice caused by Burkholderia glumae result from toxoflavin biosynthesis, lipase production and secretion, as well as motility, in response to N-octanoyl homoserine lactone [6–8]. Quorum quenching (QQ) enzymes, which can interfere with the infection process (which is modulated by the QS system of bacterial pathogens) by degrading AHLs, have become promising candidates for controlling plant bacterial disease in the past 20 years [9–11].

According to the different mechanisms of action on AHLs, QQ enzymes are conventionally divided into lactonases, acylases, and oxidoreductases [12, 13]. Among them, AHL lactonases (AHLases) have received increasing attention due to their widespread presence [14–18]. A large group of AHLases with a broad substrate spectrum...
and excellent catalytic performance have been isolated and characterized. However, most of these AHLases are easily destroyed by harsh environmental elements, including proteases, high temperatures, and high salt levels, which strongly limits their application in controlling plant bacterial diseases. To address stability issues, some thermostable AHLases, such as GKL (an orthologous phoshoetriesterase-like lactonase from the thermophile Geobacillus kaustophilus) [19], AiiTA196 (a thermostable N-acetylhomoserine lactonase from Bacillus sp. strain AI96) [20], Aii20J (a wide-spectrum thermostable N-acetylhomoserine lactonase from the marine bacterium Tenacibaculum sp. 20 J) [21], AiiIT (a thermostable N-acetylhomoserine lactonase from the thermophilic bacterium Thermaerobacter marianensis) [22] and AidB (a thermostable N-acetylhomoserine lactonase from the bacterium Bosea sp.) [23], were successively mined, and the thermostability of PPH (a phosphotriesterase-like lactonase) was improved by directed evolution [24]. Despite these efforts, the long-term stability of these AHLases under complicated ecological environments is not satisfactory. In the previous work, we identified a typical AHLase (called AhlX) from a marine bacterium, Salincola salinaria MCCCA1A01339 [25]. AhlX shows highly efficient AHL degradation activity, thermostability, and salt tolerance, and can attenuate Erwinia carotovora infection. Nevertheless, the activity of AHLase is still destroyed when it is exposed to the natural environment. It cannot be applied practically due to its inherent vulnerability in natural environments. Therefore, further improvement of AHLases stability remains quite important for their practical use.

Different from the inherent poor stability of free enzymes, enzyme immobilization based on physical, covalent, and affinity interactions with or without carrier is an effective strategy to improve enzyme stability and recyclability [26–32]. Despite this, there still exist some deficiencies in many conventional immobilization ways such as complicated preparation process, excessive cost, the use of toxic and harmful reagents, loss of activity, inferior reproducibility, and mass transfer limitations [33, 34]. Enzyme-based hybrid nanoflowers (EHNFs) are composites with a large specific surface area and porous and uniformly controllable flower-like morphology that are formed by enzymes and inorganic metal phosphates through self-assembly [35, 36]. Prepared EHNFs can effectively improve the stability, reusability and enantioselectivity of an enzyme, reduce mass transfer limitations without losing enzyme activity, and even increase the enzyme activity in many cases [37–44]. In 2012, Ge et al. accidentally discovered that enzymes can spontaneously form nanoflower-like complexes with Cu3(PO4)2 crystals and increase their enzyme activity by 650% [45]. Since then, an increasing number of enzymes, peptides, amino acids, nucleic acids, bioextracts, organic molecules, and metal ions have been attempted in the preparation of hybrid nanoflowers [32, 46–48]. The production process of EHNFs is convenient, fast, and does not use toxic, harmful, and expensive chemical reagents. For these reasons, EHNFs have become a highly important enzyme immobilization method and have been used in the fields of biosensors, biocatalysts, biomedicine, and wastewater treatment [39, 45, 49–52].

In this study, we developed an efficient strategy to increase the activity and robustness of AHLases for breaking their limitation in practical application. AhlX-based EHNFs were synthesized and applied to quench AHL-dependent quorum sensing of phytopathogenic bacteria. As shown in Fig. 1, we prepared and characterized a highly stable and highly active AHLase EHNF, AhlX@Ni3(PO4)2. AhlX was first incubated with Ni2+ and PO43− and then self-assembled into an EHNF, AhlX@Ni3(PO4)2. AhlX@Ni3(PO4)2 showed an excellent thermal stability, storage stability, organic solvent tolerance and high catalytic activity. It also displayed excellent stability in a natural water environment and exhibited long-term inhibition of plant bacterial infection. The AhlX@Ni3(PO4)2-dependent QQ strategy should largely benefit the practical control of plant bacterial diseases.

**Materials and methods**

**Bacterial strains and chemicals**

*Escherichia coli* BL21 (DE3) was used to achieve a heterologous expression of the AhlX protein in the pET28a (+) vector. The Agrobacterium tumefaciens NT1 strain expressing AHL-regulated LacZ was used to evaluate AHL activity with the bioassay method described by Dong et al. [9]. *Erwinia carotovora* SCG1 was provided by Prof. Ziduo Liu (HuaZhong Agricultural University, Wuhan, China) and was used to test the quenching activity of AhlX@Ni3(PO4)2 for potato soft rot. *Burkholderia glumae* CCTCC AB 2,016,346 was purchased from the China Center for Type Culture Collection (CCTCC) and was used to test the quenching activity of AhlX@Ni3(PO4)2 for rice seedling rot. N-(3-oxoctanoyl)-l-homoserine lactone (3OC8-HSL) was purchased from Sigma Aldrich (Missouri, USA). Other chemicals, if not specified, were purchased from Sinopharm Group Co. Ltd. (Shanghai, China).

**Expression and purification of AhlX**

The expression and purification of AhlX were performed as described in a previous study [25]. To assess the purity of the purified AhlX, 15% (v/v) sodium dodecyl sulfate-denatured polyacrylamide gel electrophoresis (SDS–PAGE) analysis was performed. Protein concentrations were measured with the Modified Bradford Protein Assay Kit (Sangon Biotech).
Synthesis of Ahlx-EHNFs

According to the promoting effect of divalent metal ions on free Ahlx activity, nickel sulfate hexahydrate, zinc sulphate heptahydrate, manganese sulfate monohydrate, magnesium sulfate, and cobalt chloride hexahydrate were chosen to synthesize Ahlx-EHNFs. The general synthetic process is as follows. Eighty microliters of aqueous divalent metal ion solution (200 mM) were added to 1 mL of 10 mM phosphate buffer (PB) with different pH values containing Ahlx at different concentrations, followed by incubation at 4 °C or 25 °C for different times. For the divalent metal ion selection experiments, the pH value of PB was 8.0, the final Ahlx concentration was 3.2 μM, and the incubation time was 36 h. For the pH preparation experiments, the pH range tested was from pH 5 to pH 9.4, the final Ahlx concentration was 3.2 μM, and the incubation time was 36 h. For the experiments preparing the Ahlx concentrations, the pH value of PB was 8.0, the final Ahlx concentration ranged from 3.2 μM to 25.6 μM, and the incubation time was 36 h. For the incubation time experiments, the pH value of PB was 8.0, the final Ahlx concentration was 3.2 μM, and the incubation time ranged from 0.5 h to 10 h. After that, the mixtures were centrifuged for 2 min at 12,000 rpm, the supernatants were decanted, and the precipitates were washed three times with ultrapure water. Protein concentrations in all supernatants were determined by a Modified Bradford Protein Assay Kit (Sangon Biotech). Amount of immobilized Ahlx is calculated by deducting total Ahlx used for immobilization from the amount of un-immobilized

Fig. 1 Schematic illustration of the preparation and application of Ahlx@Ni₃(PO₄)₂
AhlX. And the encapsulation efficiency was defined as the ratio of AhlX immobilized to the total amount of AhlX used. The final precipitates were resuspended in 10 mM PB (pH 8.0) to a final AhlX concentration of 3.2 μM.

Activity and kinetics assay of AhlX and AhlX-EHNFs
The activity and kinetics assay of free AhlX and AhlX-EHNFs was performed as described in a previous study [25]. Briefly, 5 μL of free AhlX or AhlX-EHNFs was added to 200 μL of 10 mM phosphate buffer (PB, pH 8.0) containing 1.6 mM 3OC8-HSL substrate and was incubated at 30 °C for 10 min. The reaction was terminated with SDS solution at a final concentration of 5%, and the remaining 3OC8-HSL was detected by high-performance liquid chromatography (HPLC) on a Diamonsil®C18 column (4.6 × 250 mm, 5 μm, Dikma, Beijing, China) with a mobile phase of C2H3N:HCOOH:H2O (50: 0.2: 49.8) at 1 mL min⁻¹. The detection wavelength was set to 201 nm. The concentration of 3OC8-HSL was set as 0.25–10 μM during kinetic detection. Kinetic parameters were obtained by nonlinear regression analysis of the Michaelis–Menten equation with origin 2017 (Origin Laboratory). All samples were analyzed in triplicate.

Characterization of AhlX@Ni₃(PO₄)₂ EHNF
The centrifuged AhlX@Ni₃(PO₄)₂ pellet was washed three times with ultrapure water, filtered on a 0.22 μm MCE filter membrane (Millipore), and dried at room temperature for subsequent tests. The morphology and elemental analysis of AhlX@Ni₃(PO₄)₂ were observed by a field emission scanning electron microscopy (SEM) S-4700 (Hitachi). The samples were coated with a thin layer of gold before they were placed in the SEM chamber. Fourier transform infrared spectroscopy (FTIR) spectra were collected on a Nicolet iS10 FTIR spectrometer (Nicolet). The conventional KBr pressed pellet method was used to prepare the samples. Each sample was scanned from 4000 to 400 cm⁻¹. X-ray photoelectron spectroscopy (XPS) was performed on an Axis Ultra DLD with a monochromic Al X-ray source (Kratos). Circular dichroism (CD) spectroscopy was performed with a J-815 CD spectropolarimeter (Jasco).

Biochemical characterization of free AhlX and AhlX@Ni₃(PO₄)₂
The optimum temperature and pH for free AhlX and AhlX@Ni₃(PO₄)₂ activity toward 3-OC8-HSL was assessed at different temperatures (10 to 60 °C) and different pH values (PB for pH 5.0–9.4, the activity of free AhlX at pH 8.0 was defined as 100%) under standard conditions. To test the organic solvent tolerance of free AhlX and AhlX@Ni₃(PO₄)₂, 5% of different organic solvents (ethanol, acetonitrile, isopropanol, n-butanol and isooamyl alcohol) were added to the reaction system. The activity of free AhlX in the absence of organic solvent was defined as 100%. The temperature stability of free AhlX and AhlX@Ni₃(PO₄)₂ was measured by a preprocessing treatment at 0 to 100 °C for 30 min. In addition, to measure the time-dependent thermal stability of free AhlX and AhlX@Ni₃(PO₄)₂, their residual activities were tested after incubation at 25°C for 0–40 days and 60°C for 0–48 h. All samples were analyzed in triplicate.

To detection the free AhlX and AhlX@Ni₃(PO₄)₂ stabilities in the ecological environment, they were added to sterilized and unsterilized river water (in the Zhejiang University of Technology sections of the Beijing-Hangzhou Grand Canal) to a final AhlX concentration of 3.2 μM and were incubated at room temperature for 0–30 days. Mixtures were taken at day 0, day 4, day 8, day 12, day 20, and day 30 for the activity assay with the agar strip diffusion method and 15% (v/v) SDS–PAGE analysis. For proteinase K treatment, 0.3 U of proteinase K was added to 20 μL of 3.2 μM free AhlX or AhlX-EHNF solution (10 mM PB, pH 8.0) and incubated at 37 °C for 0–60 min. Next, the activity was analyzed by the agar strip diffusion method, and the remaining AhlX was detected with 15% (v/v) SDS–PAGE analysis. Before 15% (v/v) SDS–PAGE analysis, 5 μL of 0.1 M HCl solution was added into 20 μL of AhlX@Ni₃(PO₄)₂ samples to release AhlX from it. All samples were analyzed in triplicate.

The agar strip diffusion method was performed in the following manner. Five microliters of the mixtures and 4 μL of 20 μM 3OC8-HSL were added to 11 μL of 10 mM PB (pH 8.0) and incubated at 30 °C for 10 min. The reaction was terminated with SDS solution at a final concentration of 5%. Ten microliters of reaction solution was dot coated on the anterior end of agar strips surface coated with Agrobacterium tumefaciens NT1. Cultures were grown overnight (20 h) at 30 °C, and the activity was assessed by the length of blue plaque.

Quenching the E. carotovora and B. glumae infection by AhlX@Ni₃(PO₄)₂ and CAS@Ni₃(PO₄)₂
To assess the in vitro biocontrol effect of free AhlX and AhlX@Ni₃(PO₄)₂ on the soft rot caused by E. carotovora, 1 μL of E. carotovora overnight cultures (3.0 × 10⁷ CFU mL⁻¹) and 10 μL of free AhlX or AhlX@Ni₃(PO₄)₂ (AhlX concentration was 3.2 μM) were mixed well and inoculated in the center of the potato slices. As a control, 10 μL of ddH₂O, 10 μL of AhlX, 10 μL of AhlX@Ni₃(PO₄)₂, and 1 μL of E. carotovora were inoculated in the center of the potato slices. All samples were analyzed in triplicate. The progression of lesion development was photographed every 2 days after incubation at 30 °C for
0–10 days, and the lesion area was quantified using Photoshop software (Adobe).

To investigate the in vitro suppressive efficacy of CSA@Ni3(PO4)2 in E. carotovora infection, surface-sterilized potato, marrow squash, radish, carrot, pepper, Chinese cabbage, and eggplant were inoculated separately with PB (pH = 8.0, as a control), E. carotovora, E. carotovora and CSA, E. carotovora and CSA@Ni3(PO4)2. Potato, marrow squash, and pepper slices were inoculated with 11 µL of PB (pH 8.0), a mixture of 1 µL of E. carotovora (3.0 × 10⁷ CFU·mL⁻¹) and 10 µL of PB, a mixture of 1 µL of E. carotovora (3.0 × 10⁷ CFU·mL⁻¹) and 10 µL of CSA (3.2 µM), and a mixture of 1 µL of E. carotovora (3.0 × 10⁷ CFU·mL⁻¹) and 10 µL of CSA@Ni3(PO4)2 (3.2 µM of CSA) and incubated at 30 °C for 5, 5, and 3 days, respectively. Radish, carrot, and Chinese cabbage slices were inoculated with 11 µL of PB, a mixture of 1 µL of E. carotovora (1.5 × 10⁸ CFU·mL⁻¹) and 10 µL of PB, a mixture of 1 µL of E. carotovora (1.5 × 10⁸ CFU·mL⁻¹) and 10 µL of CSA (3.2 µM), and a mixture of 1 µL of E. carotovora (1.5 × 10⁸ CFU·mL⁻¹) and 10 µL of CSA@Ni3(PO4)2 (3.2 µM of CSA) and incubated at 30 °C for 3, 3, and 2 days, respectively. Eggplant slices were inoculated with 11 µL of PB, a mixture of 1 µL of E. carotovora (1.5 × 10⁸ CFU·mL⁻¹) and 10 µL of PB, a mixture of 1 µL of E. carotovora (1.5 × 10⁸ CFU·mL⁻¹) and 10 µL of CSA (3.2 µM), and a mixture of 1 µL of E. carotovora (1.5 × 10⁸ CFU·mL⁻¹) and 10 µL of CSA@Ni3(PO4)2 (3.2 µM of CSA) and incubated at 30 °C for 2 days. To evaluate the in vivo inhibition efficacy of CSA@Ni3(PO4)2 in B. glumae infection, rice seedlings (two weeks old) were cut to the bottom 2 cm aboveground and drenched with tap water (as a control), B. glumae (1.0 × 10⁹ CFU·mL⁻¹), B. glumae and CSA (3.2 µM), B. glumae and CSA@Ni3(PO4)2 (3.2 µM CSA). The infection degree was evaluated 2 weeks after treatment.

Results and discussion
Preparation of AhlX-EHNFs
The formation of EHNFs goes through three major steps [31, 46, 53]: (a) during nucleation, divalent metal ions react with phosphate ions (PO₄³⁻) to form primary metallic phosphate crystals, (b) in step of growth, primary crystals combine with enzymes via coordination interaction between metal ions and carboxyl, amine and imidazole groups of enzymes to create agglomerates, and (c) in the final step, flower-like structures are eventually formed by anisotropic growth. In this study, AhlX-EHNFs were produced through the self-assembly of AhlX and metal phosphates. The influence of the metal ion kinds, temperature, pH value, enzyme concentration, and preparation time of AhlX-EHNFs was investigated.

The active center of the AhlX monomer contains a conserved Zn²⁺-binding domain, HXHXDH [23]. This site binds two Zn²⁺ ions and plays an important role in the catalytic ability of this type of enzyme [54–56]. In previous experiments, we found that Cu²⁺, Fe³⁺, Ca²⁺ possessed varying degrees of inhibition on the catalytic ability of AhlX, while five divalent metal ions, including Ni²⁺, Zn²⁺, Mg²⁺, Mn²⁺, and Co²⁺, have different promoting effects (between 1.5 and 3.5 times) on it [25]. Therefore, these five divalent metal ions were selected as the metal ion donors for the preparation of AhlX-EHNFs. The results showed that, except for the Mg²⁺ group, the other 4 groups all formed precipitation of AhlX-EHNFs (Fig. 2a). Zn²⁺ was also discarded in the subsequent experiments due to its weak influence in the activity of AhlX-EHNF. AhlX-EHNFs prepared by the ions of Co²⁺, Mn²⁺, Ni²⁺ show 2-, 3.5- and 6.5-times catalytic activity to free enzyme, respectively. No significant difference on activity of AhlX-EHNFs prepared at 4 °C and 25 °C had been found. The preparation temperature at about 25 °C was more energy-efficient and convenient.

As the protein concentration increased from 3.2 to 25.6 µM, the encapsulation efficiency gradually decreased (Fig. 2c). The AhlX@Ni₃(PO₄)₂ prepared at the protein concentration of 6.4 µM showed the best catalytic activity. The influence of preparation time from 0.5 to 10 h on the activity of AhlX@Ni₃(PO₄)₂ was studied. Different from EHNFs of the most enzymes, AhlX@Ni₃(PO₄)₂ with total catalytic activity could be prepared within only 0.5 h (Fig. 2d).

Finally, an optimized AhlX@Ni₃(PO₄)₂, in which the activity was 10 times higher than that of free AhlX, was successfully prepared in 10 mM PB (pH = 9.4) containing 6.4 µM AhlX and 15 mM NISO₄ at room temperature (RT, 25 °C) for 0.5 h (Fig. 2).

Structure characterization of AhlX@Ni₃(PO₄)₂
To explore the morphological and structural characteristics, AhlX@Ni₃(PO₄)₂ was analyzed by SEM, XPS, FTIR, and CD. It was found that AhlX@Ni₃(PO₄)₂
has a spherical structure with a porous surface (average size, ~2 μm) and high surface-to-volume ratios, while Ni₃(PO₄)₂ crystals exhibit small balls (average size, ~0.1 μm) and cannot form a three-dimensional stacked structure (Fig. 3a). In the presence of AhlX, AhlX can be used as a “glue” to bond small Ni₃(PO₄)₂ spherical particles into a three-dimensional structure [57]. The sponge-like surface and interior portion of AhlX@Ni₃(PO₄)₂ allow AHLs to penetrate freely and be degraded rapidly. Such a structure may be one of the reasons for the improvement in activity.

The full XPS spectra of Ni₃(PO₄)₂ and AhlX@Ni₃(PO₄)₂ revealed that they both had C, O, Ni and P elements on their surfaces (Fig. 3b). C on the surface of Ni₃(PO₄)₂ particles was attributed to CO₂ contamination from the air [58]. In contrast to Ni₃(PO₄)₂, the peak of N 1s only appeared in AhlX@Ni₃(PO₄)₂, and the peaks of C 1s and C KLL were significantly enhanced. XPS results demonstrated the successful construction of AhlX@Ni₃(PO₄)₂.

FTIR provides direct evidence of the interaction associated with the fabrication of AhlX@Ni₃(PO₄)₂ (Fig. 3c). AhlX@Ni₃(PO₄)₂ not only had a strong P-O stretching vibration absorption peak at 1050 cm⁻¹ but also had bending vibrations of bridging phosphorous such as O=P–O at 573 cm⁻¹ [59], which indicated the presence of phosphoric acid groups in AhlX@Ni₃(PO₄)₂. In addition, the FTIR image of AhlX@Ni₃(PO₄)₂ at 1650 cm⁻¹, 1542 cm⁻¹ and 1237 cm⁻¹ also demonstrated the existence of the following protein secondary structure bands: an amide I band (1660 ~ 1640 cm⁻¹, C=O stretching vibration), amide II band (1550 ~ 1530 cm⁻¹, overlap of N–H bending vibration and C-N stretching vibration) and amide III band (1340 ~ 1220 cm⁻¹) [60–62]. The peak at 1452 cm⁻¹ was assigned to the in-plane bending vibration of the C-H and O–H bonds of the protein [62]. The peaks between 2800 and 3000 cm⁻¹ correspond to the C-H bond stretching vibration peaks of -CH₂ and -CH₃ in AhlX [63]. Compared with free AhlX, no obvious peak shifts were found at the amide I, amide II and amide III bands, and no new absorption peaks appeared. The above results indicated that AhlX@Ni₃(PO₄)₂ is formed by the self-assembly of AhlX and Ni₃(PO₄)₂ through intermolecular forces and without the formation of covalent bonds.

Although no new chemical bond forms, the self-assembly of AhlX and Ni₃(PO₄)₂ still changed the
secondary structure of the protein. The shape of the CD spectra curve provides detailed information about the secondary structure of AhlX (Fig. 3d). The structure of the α-helix presents a positive peak at 192 nm and 'w'-shaped negative peaks at approximately 208 and 222 nm, while a positive peak at 196 nm and a negative peak near 218 nm are indicative of β-sheets [64]. The CD spectra of AhlX exhibited a positive peak of α-helices at 192 nm and a negative peak of β-sheets at 216 nm. Because the Ni₃(PO₄)₂ structure lacked chirality, there were no distinct peaks in the 190 ~ 240 nm UV region. After immobilization with Ni₃(PO₄)₂, the peak intensity of AhlX was dramatically reduced in the CD spectra, and the positive peak at 192 nm was redshifted and appeared at 194 nm, indicating that the proportion of α-helices and β-sheets decreased and increased, respectively.

Enzymatic properties of free AhlX and AhlX@Ni₃(PO₄)₂
The operating temperature range of AhlX@Ni₃(PO₄)₂ is wider than that of free AhlX. Compared with free AhlX, AhlX@Ni₃(PO₄)₂ was more resistant to high temperatures and could maintain the same activity at 30 °C and 60 °C, while free AhlX lost 60% of its activity at the higher temperature (Fig. 4a). At the same time, AhlX@Ni₃(PO₄)₂ had improved acid and alkali tolerance (Fig. 4b). The activity of free AhlX was completely lost at pH 5.0, while AhlX@Ni₃(PO₄)₂ maintained 19.26% of its activity. There was no decreasing trend for free AhlX when the pH was greater than 9. Moreover, AhlX@Ni₃(PO₄)₂ has good reusability. It is well known that free enzymes cannot be recovered after use. After immobilization, a free enzyme gains an insoluble solid-state, which allows it to be recovered after the reaction, and its usability and economy are improved. After 8 repeated uses, the remaining enzyme activity was still as high as 73% (Fig. 4c). After four cycles of use and recovery, this recovery rate was higher than 60% for the AiiA (a N-acylhomoserine lactonase from Bacillus sp. 240B1) that was immobilized onto magnetic nanoparticles [65]. As reusability was tested in a small reaction volume, inevitable physical loss occurred with each recycling process, which increased the loss in activity. Thus, the activity was better preserved with AhlX@Ni₃(PO₄)₂.

The kinetic constants $k_{cat}$ and $K_M$ of AhlX@Ni₃(PO₄)₂ and free AhlX were determined (Table 1). No obvious change was found in the $K_M$ value between free AhlX and AhlX@Ni₃(PO₄)₂, indicating that the affinity of AhlX to 3OC8-HSL remained unchanged. The increase

![Fig. 3 Structural characterization of the biomineralized enzyme AhlX@Ni₃(PO₄)₂. a SEM image of the biomineralized enzyme AhlX@Ni₃(PO₄)₂. b XPS spectra of Ni₃(PO₄)₂ and AhlX@Ni₃(PO₄)₂. c FTIR spectra of Ni₃(PO₄)₂, AhlX, and AhlX@Ni₃(PO₄)₂. d CD spectra of Ni₃(PO₄)₂, AhlX, and AhlX@Ni₃(PO₄)₂.](image-url)
in catalytic efficiency was due to the increase in the conversion number ($k_{cat}$) value. After immobilization, the $k_{cat}$ value had a tenfold enhancement from 21.57 s$^{-1}$ to 214.80 s$^{-1}$, and the $k_{cat}/K_M$ value increased from $6.05 \times 10^3$ M$^{-1}$·s$^{-1}$ to $6.12 \times 10^4$ M$^{-1}$·s$^{-1}$. This result was consistent with the prior activity improvement results. The increased activity of EHNFs may be derived from the following factors \[31, 45, 66\]: (a) reduced mass-transfer limitations due to high surface area, (b) enzyme favorable conformation in EHNFs, and (c) synergistic effects between the enzyme molecules and the enzyme-metal ions. According to the CD results, there was an increase in the proportion of β-sheets and a decrease in the proportion of α-helices in the AhlX structure after immobilization, indicating that the hydrophobicity of AhlX increases \[67, 68\]. According to the study of Permyakov et al. \[55\], the AHLase AiiA, which belongs to MLL AHLases, degrades the Y conformation C8-C12AHL in the active state when only one water molecule is connected to Zn$^{2+}$ in the active center. The long distance between the two Zn$^{2+}$ allows more water molecules to approach the active center, which shifts the substrate binding mode, which is critical to the catalytic ability of AiiA. The water repellency of the active center is important for the continuous catalytic ability of AiiA. Therefore, it is not difficult to speculate that the increased hydrophobicity of AhlX@Ni$_3$(PO$_4$)$_2$ improves the water repellency of the active center, eventually improving its catalytic ability for the Y conformation of 3OC8-HSL. In a nutshell, the increased activity of AhlX@Ni$_3$(PO$_4$)$_2$ is mainly due to the favorable conformation of AhlX.

**Stability of Free AhlX and AhlX@Ni$_3$(PO$_4$)$_2$**

The stability of AhlX is crucial in practical applications. To investigate the stability of free AhlX and AhlX@Ni$_3$(PO$_4$)$_2$, both were heat-treated at different temperatures for 30 min, and then the remaining activity was detected (Fig. 5a). Unlike free AhlX, which lost its total activity after a heat treatment for 30 min at 80 °C, AhlX@Ni$_3$(PO$_4$)$_2$ retained 20.00% of its enzymatic activity after the same treatment, and the activity even remained at 5.00% after a heating treatment at 100 °C for 30 min. The RT stability of AhlX@Ni$_3$(PO$_4$)$_2$ with a 3.2 µM AhlX concentration is better than that of free AhlX. After storage for 40 days, AhlX@Ni$_3$(PO$_4$)$_2$ maintained full activity, whereas free AhlX retained only 26.52% of the activity (Fig. 5b). Consistent with this, AhlX@Ni$_3$(PO$_4$)$_2$ had a better stability at 60 °C (Fig. 5c). AhlX@Ni$_3$(PO$_4$)$_2$ maintained 80.26% of its activity after 48 h of heating at 60 °C, whereas the activity of free AhlX was completely lost after 24 h of heating at 60 °C. Overall, AhlX@Ni$_3$(PO$_4$)$_2$ possesses better thermal stability than that of free AhlX. In a previous study \[25\], we found that AhlX displayed higher stability at a high concentration (0.6 mM) at 25 °C. However, the activity of AhlX showed a constantly declining trend when it was diluted to a working concentration of 3.2 µM (Fig. 5b). For practical applications, the low concentration of enzymes will contribute to a lower cost. Hence, AhlX@Ni$_3$(PO$_4$)$_2$ shows better applicability as it has outstanding stability at lower concentrations. Meanwhile, compared with free AhlX, AhlX@Ni$_3$(PO$_4$)$_2$ exhibited a better tolerance to organic solvents (Additional file 1: Figure S1). Free AhlX maintained an activity higher than 35% only when it was under 5% ethanol and acetonitrile.

![Fig. 4 Biochemical characterization and reusability of free AhlX and AhlX@Ni$_3$(PO$_4$)$_2$.](image)
However, AhlX@Ni$_3$(PO$_4$)$_2$ retained the same proportion of activity under 5% ethanol, acetonitrile, isopropanol, and $n$-butanol.

We speculated that the improvement in the stability of AhlX@Ni$_3$(PO$_4$)$_2$ to temperature and organic solvent environments is due to the packaging of the enzymes being good. The stability of inorganic Ni$_3$(PO$_4$)$_2$ under heat and acid–base conditions provides a relatively stable microenvironment for the AhlX wrapped in it, making it more resistant than free AhlX to changes in the external environment. To further validate our hypothesis, the samples were diluted with sterilized and unsterilized river water for different time spans. The activity was detected by the agar strip method (the longer the blue agar strip, the lower the activity), and the remaining protein was detected by SDS–PAGE (Fig. 5d, e). The activity of free AhlX treated with unsterilized river water on day 0 was significantly lower than that of the other three groups and basically lost all activity after 8 days. SDS–PAGE results showed that there was no obvious degradation of AhlX at the beginning. It is likely that there were some substances in the unsterilized river water that inhibited the activity of free AhlX. Considering both the activity test and SDS–PAGE results, it could be observed that a small residual amount of free AhlX in sterilized river water that was incubated for 8 days still showed great activity, but eventually it was completely degraded and lost activity after 30 days. It seemed that bioactive factors such as proteases or microorganisms could degrade the AhlX protein. At the same time, some substances can inhibit the activity of AhlX but do not induce degradation. In contrast, the activity of AhlX@Ni$_3$(PO$_4$)$_2$ was maintained well when it was treated in sterilized or unsterilized river water for 30 days. AhlX of AhlX@Ni$_3$(PO$_4$)$_2$ was barely degraded after a treatment in sterilized or unsterilized river water for 30 days.

To further evaluate the anti-proteinase ability of AhlX@Ni$_3$(PO$_4$)$_2$, free AhlX and AhlX@Ni$_3$(PO$_4$)$_2$ were digested with proteinase K for 0–60 min before activity and SDS–PAGE assays. The results showed that AhlX@Ni$_3$(PO$_4$)$_2$ maintained full activity after proteinase K digestion for 60 min, while free AhlX completely lost its
activity (Fig. 5f). This difference was derived from the anti-proteinase K effect of AhlX@Ni$_3$(PO$_4$)$_2$ (Fig. 5g). AhlX@Ni$_3$(PO$_4$)$_2$ can protect AhlX from proteinase K digestion. In summary, AhlX@Ni$_3$(PO$_4$)$_2$ was embedded in a safe three-dimensional structure. This structure provided a good protective barrier and structural support for AhlX, and it avoids inhibition and degradation, which is caused by many factors, and enables AhlX to keep a stable activity for a long time. Due to its structural rigidity and stability in the natural environment, AhlX@Ni$_3$(PO$_4$)$_2$ has superb potential in the control of bacterial infections.

The biological control effect of AhlX@Ni$_3$(PO$_4$)$_2$ on bacterial infections

Many gram-negative plant pathogenic bacteria regulate virulence with AHL-based QS systems. With the goal of targeting QS systems, many biological control methods, including treatment with wild or transgenic QQ bacteria and transgenic plants expressing QQ enzymes, have been attempted against plant pathogens [11]. Compared with these methods, the directed use of highly stable enzymes in ecological environments has little effect on microecology and no gene contamination. In the in vitro biological control experiment, potato slices were treated with free AhlX and AhlX@Ni$_3$(PO$_4$)$_2$, and these were all mixed with _E. carotovora_ SCG1 (Fig. 6). The free AhlX treatment group showed that the potato soft rot caused by _E. carotovora_ was effectively alleviated. In addition, AhlX@Ni$_3$(PO$_4$)$_2$ completely inhibited potato soft rot. There were no soft rot symptoms on potato slices within 10 days of culture, and the inhibitory effect was as high as 100%. Compared with free AhlX, AhlX@Ni$_3$(PO$_4$)$_2$ had an effective and lasting effect on the biological control of _E. carotovora_ infection. The excellent biocontrol effect of AhlX@Ni$_3$(PO$_4$)$_2$ is due to its efficient activity toward AHL molecules and strong tolerance to complex environmental factors.

For practical application, it is important not only to have a good biological control effect but also to produce agents in a convenient and inexpensive way. David et al. encapsulated an AHLase, PPH, in nanospherical capsules composed of tertbutoxycarbonyl-Phe-Phe-OH peptide [24]. It showed significantly improved thermal resistance and effectively attenuated _E. amylovora_ infection in plants in the field compared to that of previous AHLases. However, the use of purified AHLases and expensive peptides still restrict its practical application due to the complex process and high cost. Using purified AhlX to produce EHNFs is also costly in scale-up production. To reduce the cost and streamline production processes, after cell lysis, the heat-treated crude supernatant of AhlX (CSA) was used to synthesize CSA@Ni$_3$(PO$_4$)$_2$. We tested the in vitro inhibitory effect of CSA@Ni$_3$(PO$_4$)$_2$ against _E. carotovora_ infections. As shown in Fig. 7a, CSA effectively inhibited the _E. carotovora_-induced infectious symptoms of soft rot on potato, marrow squash, radish, pepper, Chinese cabbage and eggplant slices after incubation at 30 °C for different times, while full inhibition was observed on almost all of the plants inoculated with _E. carotovora_ and CSA@Ni$_3$(PO$_4$)$_2$ except for the marrow squash.
group, in which minimal soft rot was found. In addition, we validated the in vivo inhibitory effect of CSA@Ni$_3$(PO$_4$)$_2$ against rice seedling rot that was caused by B. glumae. Inoculation with B. glumae alone resulted in serious seedling rot symptoms, whereas no seedling rot was observed in the rice seedlings treated with B. glumae and CSA (Fig. 7b). These data together illustrate that AhlX@Ni$_3$(PO$_4$)$_2$ confers a great utility to AhlX and hold immense potential to become a next-generation antimicrobial agent.

Conclusions

In this work, we prepared and characterized an extremely stable and highly active AHLase EHNF, AhlX@Ni$_3$(PO$_4$)$_2$. AhlX@Ni$_3$(PO$_4$)$_2$ maintained highly effective and durable activity in a complicated ecological environment. The efficient control of E. carotovora and B. glumae by AhlX@Ni$_3$(PO$_4$)$_2$ in vitro and in vivo indicates its significant potential for further applications in controlling bacterial diseases.

EHNFs show apparent advantages when compared with some other biocontrol methods. First, in contrast to transgenic plants, there is no rigorous approval process [69]. Second, unlike transgenic biocontrol bacteria, there is no gene contamination with the environment [70]. Third, the materials used in the process of synthesizing EHNFs are all natural. Unlike enzymes immobilized with resin, AhlX@Ni$_3$(PO$_4$)$_2$ can be easily degraded into natural substances. Finally, since the EHNF prepared from CSA possessed outstanding biocontrol capabilities, the preparation procedure is straightforward and can be easily scaled up at a low cost.

This is the first time that EHNF was used in the biological control of plant diseases. Practically, AHLases can also be applied in other fields. Many AHLases (AiiA, AiiM, SsoPox, MomL and BpiB09) could effectively decrease the production of motility, virulence factors and biofilm formation of a common human and animal pathogen, Pseudomonas aeruginosa, in various in vitro or in vivo models, Caenorhabditis elegans, Drosophila melanogaster and rodents [13]. AiiA reduce the pathogenic of Vibrio sp. in brine shrimp and manila clam [71]. A membrane bio-reactor with QQ bacteria entrapping beads successfully achieved the control of membrane biofouling by suppressing the AHLs responsible for motility, the secretion of extracellular polymeric substances and biofilm attachment to the membrane surface [72, 73]. Due to the high activity and stability of AhlX@Ni$_3$(PO$_4$)$_2$, it has great promise for application in human and animal pathogens and biofouling control, and this hypothesis needs further experimental validation. In addition, although many AHLases have been isolated and studied in the laboratory, they cannot be applied practically due to their vulnerability in real environments. This work provides a feasible and universal strategy to improve the biological robustness of AHLases. Hence, AHLase-EHNFs are expected to solve the bottlenecks of poor stability and limited environmental tolerance, which have existed over two decades since AHLases were discovered, in practical applications.
Supplementary Information

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Additional file 1: Figure S1. The organic solvent tolerance of free AhIX and AhIX@Ni3(PO4)2. Figure S2. Detection of the remaining AhIX by SDS–PAGE after the sterilized and unsterilized river water treatments for 0 day. Figure S3. Detection of the remaining AhIX by SDS–PAGE after the sterilized and unsterilized river water treatments for 4 days. Figure S4. Detection of the remaining AhIX by SDS–PAGE after the sterilized and unsterilized river water treatments for 8 days. Figure S5. Detection of the remaining AhIX by SDS–PAGE after the sterilized and unsterilized river water treatments for 30 days. Figure S6. Detection of the remaining AhIX by SDS–PAGE after the proteinase K treatment. Standard marker protein (lane M).

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Author contributions

YC and PL contributed equally to this work. JW and XC obtained supporting funding. BY, YC, and SX contributed equally to this work. JW, YC, and PL developed the experimental protocols. WY, YC, and SX contributed equally to this work. All authors read and approved the final manuscript.

Availability of data and materials

All data generated or analyzed during this are included in this published article.

Declarations

Ethics approval and consent to participate

Not applicable.

Consent for publication

All authors agreed to submit this manuscript.

Competing interests

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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