Biochar-immobilized *Sphingomonas* sp. and *Acinetobacter* sp. isolates to enhance nutrient removal: potential application in crab aquaculture

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ABSTRACT: The frequency of water exchange and reducing the risk of eutrophication to surrounding water bodies have always been water-quality control issues in recirculating aquaculture systems. In this study, maize straw biochar prepared through pyrolysis showed great potential for both bacterial immobilization and pollutant adsorption. Heterotrophic bacterial strains of *Sphingomonas* sp. PDD-57b-25 and *Acinetobacter towneri* were isolated in situ from wastewater for pollutant remediation through a 16S rDNA-based method, which has been rarely reported to date. The selected strains had higher ammonia nitrogen (NH₄⁺-N, 63%), nitrite nitrogen (NO₂⁻-N, 38%), nitrate nitrogen (NO₃⁻-N, 25%) and total phosphorus (TP, 35%) assimilation capacities than those of other widely applied bacteria under similar medium conditions. In addition, more NH₄⁺-N (+16%), NO₂⁻-N (+14%), NO₃⁻-N (+17%) and TP (+19%) was removed by biochar-immobilized isolated strains than dissociated strains, suggesting their use may provide a means of improving water-quality control in recirculating aquaculture. With specific additions (4 g l⁻¹) of biochar-immobilized *Sphingomonas* sp. PDD-57b-25 and *A. towneri*, the dissolved inorganic nitrogen (approximately 0.45 mg l⁻¹) and TP (approximately 0.09 mg l⁻¹) levels were maintained below the clean water threshold for recirculating aquaculture of crab *Eriocheir sinensis*. Furthermore, the added strains exhibited high bio-safety and were capable of improving the yield and quality of crabs. Results indicate the potential applicability of biochar-immobilized *Sphingomonas* sp. PDD-57b-25 and *A. towneri* in agricultural sewage treatments. Further, the experimental methodology developed here may be used for the exploration of new strains for practical aquaculture.

KEY WORDS: *Sphingomonas* · *Acinetobacter* · *Eriocheir sinensis* aquaculture · Inorganic nitrogen · Phosphorus

1. INTRODUCTION

Crab *Eriocheir sinensis* aquaculture is one of the most important agricultural industries in Jiangsu Province, China, and plays a vital role in the promotion of local economies and human living standards. According to a survey in 2018, the total area of crab aquaculture in China was approximately 2 million ha, crab yield exceeded 40 million t, and annual production reached US$ 30 billion (J. Yu et al. 2019). Due to the rapid expansion of intensive crab aquaculture, the excessive nutrients in aquaculture water have resulted in significant environmental pollution, which has increased the remediation input costs. With increasing awareness of environmental protection and sustainable development, water-quality control has become a necessary step in aquaculture to decrease agricultural non-point source pollution (Gelfand et al. 2003).

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Levels of dissolved inorganic nitrogen (DIN) (ammonium nitrogen \([\text{NH}_4^+\text{-N}]\), nitrite nitrogen \([\text{NO}_2^-\text{-N}]\) and nitrate nitrogen \([\text{NO}_3^-\text{-N}]\) and total phosphorus (TP) are 2 important parameters in crab aquaculture, as these nutrients can be toxic at high concentrations and hazardous to crab health (N. Yu et al. 2019). Generally, the abovementioned nutrients are derived from the metabolic waste of aquatic animals and microbial decomposition of the remaining fish bait. Lowering their concentrations to within environmentally friendly ranges is a key issue in the successful implementation of crab aquaculture. Water replacement is the most widely applied approach to maintain DIN and TP at desirable levels in conventional crab cultivation (Qin et al. 2019). However, with this strategy, discharged nutrients can result in eutrophication of surrounding water bodies. Moreover, any included pathogens may be dispersed, which could be hazardous to plants and animals.

Recently, the recirculating aquaculture system (RAS), which uses microbes and water-processing equipment, has become a popular approach to water-quality maintenance. Nevertheless, the comparatively low recirculating efficiency and high cost restrict its large-scale application, and improving RAS performance in terms of denitrification and phosphorus removal is key to overcoming these bottlenecks. Currently, removal of DIN is typically achieved through nitrification and denitrification processes (Liu et al. 2016, S. Li et al. 2019). These processes involve several biochemical reactions and require alternating between anaerobic and aerobic conditions to ensure suitable performance. In this way, nitrifying and denitrifying bacterial activity imbalances can occur, which may result in the accumulation of \(\text{NO}_2^-\) and \(\text{NO}_3^-\) (Liang et al. 2014). In addition, RAS nitrogen removal is conducted with autophytic bacteria in a process that requires extra equipment, such as biofilters or biocarriers, to enhance bacterial propagation and function; however, notable enhancement is still lacking (Zhu et al. 2019). Unlike nitrogen, the phosphorus in aquaculture sewage is generally removed through chemical precipitation by using agents such as iron or aluminum salts (Wu et al. 2019). However, such chemical remediation approaches have the potential to threaten aquatic animal health and increase secondary pollution. Even though the biological phosphorus removal method using algae has been developed to overcome the aforementioned shortcomings, studies on the mechanisms of phosphorus assimilation and optimal conditions for algal growth are rare and therefore this technique still lacks experimental support (Filipe et al. 2001, Acevedo et al. 2015, Zhou et al. 2016).

Because of these bottlenecks, wastewater-borne bacteria have received much attention in RAS water-quality control. These organisms are heterotrophic bacteria that possess nitrogen and phosphorus assimilation capabilities and can convert nutrients into cellular biomass. Due to their rapid adaptability and preferable ammonium removal and phosphorus accumulation potential in aquatic environments, these bacteria may be suitable alternatives to other widely-used bacteria for water remediation (Sun et al. 2015). It is critical to examine strains of wastewater-borne bacteria more closely and to improve their nutrient removal efficiency; only a few suitable strains have been isolated and identified in aquaculture sewage thus far. Moreover, most of these strains are efficient at ammonium removal and phosphorus accumulation under certain medium conditions, but their application in high-content natural aquaculture wastewaters has been rarely reported (Zhu et al. 2018). To enhance the performance of selected ammonia nitrogen removal strains (ANRSSs) and phosphate removal strains (PRSSs), biochar possessing a large specific surface area (SSA) and complex pore structure could be used in conjunction with bacteria. Due to attachment effects, biochar can serve as a habitat to simultaneously promote bacterial propagation and reduce nitrogen and polyphosphate \((\text{PO}_4^{3-}, \text{HPO}_4^{2-}, \text{H}_2\text{PO}_4^-)\) concentrations in chicken farm effluents (Agyarko-Mintah et al. 2017). However, research on the enhanced performance of biochar-based probiotics on nutrient removal in RAS is still lacking.

The objectives of this study were as follows: (1) to characterize the physico-chemical properties of prepared biochar and assess its bacterial immobilization and nutrient adsorption potential, (2) to isolate and identify wastewater-borne strains of bacteria that possess optimal nitrogen and phosphorus assimilation capacities, (3) to evaluate the enhancement of biochar-immobilized bacteria on nitrogen and phosphorus removal and (4) to elucidate the water-quality control and bio-safety effects of biochar-immobilized bacteria on recirculating \textit{Eriocheir sinensis} aquaculture.

## 2. MATERIALS AND METHODS

### 2.1. Experimental design

The experiments were implemented in 3 steps. First, maize straw biochar was prepared through pyrolysis for bacterial adsorption, and optimal ANRSSs and PRSSs were selected and identified through
a 16S rDNA method (Fergola et al. 2007, Fitzgerald et al. 2015). Second, the performance of the biochar-immobilized isolated strains on sewage remediation was measured in laboratory-scale experiments. Relative nutrient-removal enhancement was estimated by comparing the efficiencies of the ANRS and PRS among different treatments. Third, the biochar-immobilized isolated strains were applied in practical crab aquaculture. Dynamic variations of nutrients were monitored, and the survival rate of crabs was measured to evaluate biosafety and application potential of the ANRS and PRS. All treatments and relevant analyses were conducted in triplicate.

2.2. Sewage collection and analysis

The sewage samples were collected from *Eriocheir sinensis* aquaculture ponds in Guchen Lake (31° 14’ N, 118° 53’ E), Nanjing city, Jiangsu Province, China. The samples were stored in cool conditions (4°C) before use; the water-quality parameters and determination methods of the collected aquaculture sewage are summarized in Table 1. All analyses were conducted in triplicate. The concentrations of heavy metal(loids) were below the safety threshold of the Chinese Water Quality Standard for Fisheries (serial number GB 11607-89). The NH4⁺-N, NO2⁻-N, NO3⁻-N, total nitrogen (TN) and TP concentrations were 8.6, 1.4, 0.7, 12.7 and 2.5 mg l⁻¹, respectively, thereby reaching a severe eutrophication level according to the classification criteria of the Environmental Quality Standard for Surface Water (GB 3838-2002).

| Parameter | Method | Unit | Value       |
|-----------|--------|------|-------------|
| pH        | Glass electrode method (GB 11901) | 7.2 ± 0.4 |
| NH4⁺-N    | Nessler’s reagent photometry (GB 7479) | 8.6 ± 0.7 |
| NO2⁻-N    | Phenol disulfonic acid method (GB/T 7493) | 1.4 ± 0.2 |
| NO3⁻-N    | Ultraviolet spectroscopy (GB HJ/T 346) | 0.7 ± 0.1 |
| TN        | GB HJ/T 346 | mg l⁻¹ | 12.7 ± 0.8 |
| TP        | Ammonium molybdate spectrophotometry (GB 11893-89) | mg l⁻¹ | 2.5 ± 0.3 |
| DO        | Iodometry (GB 7489) | mg l⁻¹ | 6.2 ± 0.3 |
| CODMn     | Potassium permanganate method (GB 7488) | mg l⁻¹ | 14.2 ± 0.6 |

| Heavy metals | Method                                      | Unit  | Value       |
|---------------|---------------------------------------------|-------|-------------|
| Cd            | Atomic absorption spectrometry (GB 7475)    | mg l⁻¹ | 0.002 ± 0.0003 |
| Pb            | GB 7475                                     | mg l⁻¹ | 0.033 ± 0.008 |
| Cr            | GB 7475                                     | mg l⁻¹ | 0.045 ± 0.01  |
| Hg            | GB 7475                                     | mg l⁻¹ | 0.0003 ± 0.0001 |
| As            | GB 7475                                     | mg l⁻¹ | 0.021 ± 0.004 |

Table 1. Mean (±SD) water-quality parameters, measurement methods used (GB number: relevant Chinese national standard method), and measured values of the collected aquaculture sewage. TN: total nitrogen; TP: total phosphorus; DO: dissolved oxygen; CODMn: chemical oxygen demand determined by acid potassium permanganate oxidation.

2.3. Preparation of the biochar-immobilized bacteria

2.3.1. Biochar production and characterization

Maize straw of the variety Suyu No. 9 (Nanjing Lianghua Ecological Agriculture Technology Development Co.) was selected to produce biochar for the immobilization of bacteria. Maize straw (20 g) was first weighed and broken into small pieces (5 mm length). The straw was then placed in a 1000 ml porcelain crucible and pyrolysed in a muffle furnace at 400°C for 2 h. The pyrolysis was maintained in anoxic conditions at a heating rate of 10°C min⁻¹. Subsequently, the prepared maize straw biochar was washed with deionized water and stored in a plastic bag for characterization after grinding (particle diameter: <0.5 mm).

The measured physico-chemical properties of the biochar are summarized in Table 2. The carbon yield (η) was calculated through the following equation:

\[ \eta = \frac{M_{\text{Biochar}}}{M_{\text{Biomass}}} \times 100\% \]  \hspace{1cm} (1)

where \( M_{\text{Biochar}} \) (g) represents the mass of biochar and \( M_{\text{Biomass}} \) represents the mass of maize straw before pyrolysis.

The measurement of ash content was compared to the national standard (GB/T 12496); 1 g of maize straw was put into a crucible and pyrolysed in a muffle furnace at 800°C to a constant weight. The ash content (%) was determined by:

\[ \text{Ash content} = \frac{M_2 - M_1}{M} \times 100\% \]  \hspace{1cm} (2)

where \( M \) is the mass of maize straw before ashing, \( M_1 \) is the mass of the crucible after drying and \( M_2 \) is the mass of the maize straw biochar and crucible after ashing.
The pH and electrical conductivity of biochar were measured using a pH meter (GB/T 12496) and conductivity meter (GB/T 24525), respectively. Dissolved organic carbon was determined using a total organic carbon analyzer (GB/T 3257). The mass ratios of biochar carbon and hydrogen were measured with an elemental analyzer (GB/T 15460); the mass ratio of oxygen was achieved by deducting the ash quality based on the mass balance. The surface structure of the biochar was observed via scanning electron microscopy (SEM). Moreover, the SSA, pore volume and pore diameter of the biochar were measured with a SSA and aperture analyzer (GB/T 19587): first, N2 adsorption−desorption isotherm tests were carried out (−196.15°C) with a 2 h pre-degassing process of biochar at 200°C; then SSA was calculated using the Brunauer-Emmett-Teller (BET) method, and the total pore volume was determined under specific relative pressure (P/P0 = 0.95); subsequently, the t-plot method was applied to determine the micropore volume and the Barrett-Joyner-Halenda (BJH) method was utilized to calculate the pore size distribution.

2.3.2. Bacterial isolation and screening

A synthetic medium consisting of NaAc (3 g l−1), Na2HPO4 (30 mg l−1), NH4Cl (60 mg l−1), MgSO4 (130 mg l−1), K2SO4 (25 mg l−1) and CaCl2·2H2O (18 mg l−1) was prepared for bacterial isolation. Additional (NH4)2SO4 (200 mg l−1) and K2HPO4 (100 mg l−1) were added individually to the base synthetic medium for ANRS and PRS screening, respectively. Moreover, beef extract medium, which contained beef extract (5 g l−1), peptone (10 g l−1), NaCl (15 g l−1) and agar (20 g l−1), was applied for further enrichment of the selected bacteria. The pH levels of the abovementioned mediums were adjusted to 7.0 before use.

Aquaculture sewage was added to the synthetic medium at a volume ratio of 9:1 (total volume: 250 ml) for bacterial incubation at 25°C in a shaker. After 48 h, an aliquot of bacterial solution was extracted on Luria-Bertani agar plates and incubated overnight at 25°C. Then, based on the colony morphologies, potential target bacterial colonies were selected and transferred to the beef extract medium for enrichment (de-Bashan et al. 2008). Thereafter, the enriched bacteria were streaked twice on the synthetic medium for purification, and specific strains (10 each) of the potential NH4+-N and TP removal bacteria were obtained. The selected strains were cultured overnight, and the bacterial cell densities were adjusted to 1 × 108 cfu ml−1. Subsequently, the selected strains (1 ml each) were incubated in the synthetic medium (1 l) for 48 h at 25°C. The initial and final NH4+-N and TP contents in the medium were measured to determine the most suitable strains for nutrient removal. The selected strains were stored (−20°C) for genomic identification and subsequent experiments.

2.3.3. Identification of the selected strains

The bacterial strains that possessed optimal NH4+-N and TP removal abilities were identified through a 16S rDNA-based method. The genomic DNA of the strains was extracted using a DNA extraction kit (TaKaRa). The primers 27f (5'−AGR GTT GAT CMT GGC TCA G-3') and 1492r (5'−GYT ACC TTG TTA CGA CTT-3') were further applied for PCR amplification of the 16S rDNA gene sequences (Liu et al. 2014, Chen et al. 2016). The obtained sequences were then compared with known sequences in the GenBank database using BLAST, and the genomic homologies between the isolated and GenBank strains were analyzed.

2.3.4. Adsorption of isolated bacterial strains on the prepared biochar

For bacterial adsorption, 1 g of prepared biochar was mixed with 10 ml isolated ANRS or PRS (cell density: 1 × 108 cfu ml−1). The mixtures were added into 250 ml conical flasks, and sterile water was added to adjust the total volume to 100 ml. Then the mixtures were incubated in a shaker (160 rpm) for 6 h at 25°C. Afterwards, the bacterial cell densities in the supernatant liquid were measured using the plate count method. In this way, we further determined the total ANRS and PRS amounts that had been adsorbed.

| Parameter                  | Unit   | Value     |
|----------------------------|--------|-----------|
| Carbon yield               | %      | 47.6 ± 2.7|
| Ash content                | %      | 38.2 ± 3.5|
| pH                         |        | 8.6 ± 0.4 |
| Electrical conductivity    | μS cm−1| 2032 ± 236|
| Dissolved organic carbon   | mg g−1 | 11.2 ± 1.3|
| H/C                       |        | 0.83 ± 0.05|
| O/C                       |        | 0.46 ± 0.03|

Table 2. Mean (±SD) physico-chemical properties of the maize straw biochar prepared through a 2 h pyrolysis at 400°C
onto the biochar surface, which were approximately $6.3 \times 10^7$ and $4.5 \times 10^7$ cfu g$^{-1}$, respectively.

### 2.4. Biochar-immobilized isolated bacteria for enhanced nutrient removal

Batch experiments were conducted at laboratory-scale to evaluate the performance and enhancement of nitrogen and phosphorus removal by biochar-immobilized isolated bacteria. In this step, 5 treatments were defined: a control group without the addition of agent (T1); sewage remediated with effective microorganisms (EMs) (T2); sewage remediated with prepared maize straw biochar (T3); sewage remediating with unfixed ANRSs and PRSs (T4); and sewage remediating with biochar-immobilized ANRSs and PRSs (T5). The EMs were purchased from EMRO-CHINA and are currently among the most widely applied water purification agents in crab aquaculture. To simulate the practical crab aquaculture environment, the pH, light intensity, temperature, air humidity and dissolved oxygen (DO) levels were controlled at approximately 7.0, 100 μmol m$^{-2}$ s$^{-1}$, 25°C, 40% and 10 mg l$^{-1}$, respectively. The NH$_4^+$-N, NO$_2^-$-N, NO$_3^-$-N and TP contents were measured every 12 h, and the nutrient removal efficiencies among the different treatments were compared after 120 h.

### 2.5. Application potential of biochar-immobilized isolated bacteria in crab aquaculture

In this step, the RAS, which included a polypropylene aquaculture pond, drum micro-filter, aerator, programmable temperature chamber, ozonator and sewage pump, was applied to *Eriocheir sinensis* aquaculture (n = 3 ponds per treatment; untreated control vs. biochar-treated). A dosage of the biochar-immobilized isolated bacteria was homogeneously added to the polypropylene pond without water exchange during the aquaculture. Environmental conditions were monitored daily and strictly controlled within the desired range as mentioned in Section 2.4: pH was measured by the glass electrode method (GB 11901), light intensity was detected with an underwater irradiance meter (GB 7000), temperature and DO levels were monitored by a water-quality analyzer (GB T31962), and air humidity was determined using a capacitive digital hygrometer (GB T18204). In addition, the dynamic variations in DIN (NH$_4^+$-N, NO$_2^-$-N and NO$_3^-$-N) and TP contents were measured every 5 d during the aquaculture process (90 d) to assess the water remediation efficiencies of the biochar-immobilized isolated bacteria in the RAS. The methods and relevant standards for nutrient measurements are summarized in Table 1. At the end of the experiment, random sampling (50 samples) of *E. sinensis* from both sets of polypropylene aquaculture ponds was conducted. Indexes of survival rate, weight, stocking density, biomass, crude protein, crude fat and water content (GB T19957) were determined to evaluate the bio-safety and potential application of the biochar-immobilized isolated bacteria for aquaculture within a RAS.

### 2.6. Statistical analysis

The data were processed and averages calculated in Microsoft Excel. Relevant standard deviations were statistically analyzed in SPSS v.11.5 software for illustrating the degree of dispersion in the data. The calculations and analysis in BET, t-plot and BJH methods were conducted in Origin v.8.5. In addition, Duncan’s multiple range tests in MATLAB v.7.0 were applied for analysis of nitrogen and phosphorus removal rates with utilization of the biochar-immobilized isolated bacteria ($p < 0.05$).

### 3. RESULTS AND DISCUSSION

#### 3.1. Characterization of maize straw biochar

The SEM results of the maize straw biochar prepared through 2 h of pyrolysis (400°C) are presented in Fig. 1. The biochar surface observed under 1000× SEM magnification was rough, with a complex pore structure (Fig. 1a). The surface pores that were irregularly distributed across the cellular structure were much clearer under 2500× magnification (Fig. 1b). Generally, the surface pore volume and diameter of the biochar strongly determine its SSA, which is closely related to its bacterial adsorption potential (Fu et al. 2019). Thus, N$_2$ adsorption–desorption isotherm analysis of the biochar was conducted to reveal its possible surface pore distribution.

As shown in Fig. 2a, under high temperature (77 K), the N$_2$ adsorption–desorption volumes increased from approximately 88 to 163 cm$^3$ g$^{-1}$ as an increase of relative pressure ($P/P_0$). It should be noted that the N$_2$ adsorption–desorption volume increased slowly at low $P/P_0$ (<0.9) but increased rapidly when $P/P_0 > 0.9$. This phenomenon indicates that the variations in N$_2$ adsorption–desorption of the prepared biochar in this study were consistent with Type IV isotherms according to
the classification criteria reported by the International Union of Pure and Applied Chemistry. No significant difference was observed between the N₂ adsorption and desorption curves, which suggests that there were no large pores (pore size: >50 nm) on the surface of the prepared biochar (J. Li et al. 2019, Wang & Wang 2019). Moreover, a hysteresis loop between the N₂ adsorption and desorption curves appeared when $P/P_0$ ranged from 0.4−0.65. This indicates that the pores distributed on the biochar surface mainly consisted of micropores and mesopores (2 nm < pore size < 50 nm), as depicted in Fig. 2b.

The biochar SSA, total pore volume, micropore volume and average pore size (APS) were further measured using the BET, $t$-plot and BJH methods, respectively (Table 3). The high micropore ratio (79%) and small APS (2.84 nm) resulted in a relatively large SSA (376.4 m² g⁻¹) of the prepared biochar. This may enable a suitable adsorption capacity for both bacteria and nutrients due to the abundant binding sites on its surface.

3.2. Nutrient removal potential and identification of selected strains

The NH₄⁺-N and TP removal capacities of the isolated ANRSs and PRSs are presented in Fig. 3. Clearly, dif-
Different strains possessed different NH$_4^+$-N and TP removal efficiencies. After 48 h of cultivation, the highest NH$_4^+$-N and TP removal rates were 58 and 34%, respectively, which were obtained in synthetic media containing the ANRS5 and PRS4 strains. Thus, ANRS5 and PRS4 were good candidates for genetic identification.

The 16S rDNA gene sequences of ANRS5 and PRS4 were further compared with known sequences in the GenBank database with BLAST. The analysis of their genomic homologies are summarized in Table 4. The results indicated that the selected ANRS5 and PRS4 are most likely members of the genera *Sphingomonas* and *Acinetobacter*, respectively, due to their high similarity (>98%). Moreover, through repeated comparisons, ANRS5 and PRS4 were found to have 100% proximity with *Sphingomonas* sp. PDD-57b-25 and *A. towneri*, respectively, of which the known strains in GenBank have accession numbers KR922120 and AF589023. Therefore, we speculated that the wastewater-borne ANRS5 and PRS4 in this study were *Sphingomonas* sp. PDD-57b-25 and *A. towneri*, respectively.

### 3.3. Enhanced nitrogen and phosphorus removal by biochar-immobilized isolated bacteria

The NH$_4^+$-N, NO$_2^-$-N, NO$_3^-$-N and TP removal efficiencies in batch experiments are shown in Fig. 4. Since side effects can result in slight intensity variations, the temperature, air conditions and DO levels were well simulated based on previous practical aquaculture data. The results of the batch experiments accurately reflected the pollutant removal performances of the different treatments and revealed possible treatment enhancement using biochar-immobilized ANRS5 and PRS4 (BIAP). BIAP (T5) was added to aquaculture sewage at a ratio of 2 g:1 l. The biochar amount and bacterial cell quantity applied in the other treatments (T2–T4) were in accordance with this ratio.

After 120 h, the NH$_4^+$-N concentration was reduced from 8.6 to 1.8 mg l$^{-1}$, which represents a removal rate of approximately 79% with biochar-immobilized ANRS5. In addition, decreases in NO$_2^-$-N and NO$_3^-$-N were observed, with levels decreasing from 1.3 to 0.67 mg l$^{-1}$ and from 0.67 to 0.37 mg l$^{-1}$, respectively. The genus *Sphingomonas* was first reported to have a satisfactory NH$_4^+$-N degradation ability by Yabuuchi et al. (1990). It was characterized as a beneficial strain for NH$_4^+$-N oxidation in circulating aquaculture systems or bioreactors. In many cases, NH$_4^+$-N was reported to be gradually converted into NO$_2^-$-N and subsequently nitrified into NO$_3^-$-N through

### Table 3. Mean (±SD) specific surface area, total pore volume, micropore volume and average pore size of the maize straw biochar

| Indexes         | Unit   | Value     |
|-----------------|--------|-----------|
| Specific surface area | m$^2$ g$^{-1}$ | 376.4 ± 12.3 |
| Total pore volume   | cm$^3$ g$^{-1}$ | 0.212 ± 0.008 |
| Micropore volume    | cm$^3$ g$^{-1}$ | 0.168 ± 0.007 |
| Average pore size   | nm     | 2.84 ± 0.04 |

### Table 4. The results indicated that the selected ANRS5 and PRS4 are most likely members of the genera *Sphingomonas* and *Acinetobacter*, respectively.
effective microorganisms (T2), prepared maize straw biochar (T3), unfixed ammonia nitrogen and phosphate removal strains (ANRS and PRS) (T4) and biochar-immobilized ANRS and PRS (T5) in laboratory experiments. Error bars: SD

Table 4. Genomic proximities between the isolated and GenBank strains. ANRS: ammonia nitrogen removal strain; PRS: phosphate removal strain

| Strain/rRNA length (bp) | GenBank strain (accession no.) | Proximity (%) |
|-------------------------|--------------------------------|---------------|
| ANRS5/1456              | Sphingomonas sp. PDD-57b-25 (KR922120) | 100           |
|                         | Sphingomonas sp. aurantiaca (AJ429236) | 98.63         |
|                         | Sphingomonas sp. PDD-60B-30 (KR922201) | 99.71         |
|                         | Sphingomonas sp. NMC17 (GU321356) | 99.28         |
|                         | Sphingomonas sp. PDD-63b-46 (KR922143) | 98.85         |
|                         | Sphingomonas sp. PDD-63b-1 (KR922270) | 98.74         |
|                         | Sphingomonas sp. PDD-57b-28 (KR922123) | 98.36         |
|                         | Sphingomonas sp. Ze13 (KR088450) | 98.18         |
| PRS4/1447               | Acinetobacter towneri (AF589923) | 100           |
|                         | Acinetobacter baumannii (AM410709) | 99.97         |
|                         | Acinetobacter haemolyticus (Z93437) | 98.26         |
|                         | Acinetobacter calcoaceticus (X81661) | 98.56         |
|                         | Acinetobacter lwoffii (Z93441) | 98.36         |
|                         | Acinetobacter junii (AM410704) | 99.14         |
|                         | Acinetobacter genomic (Z93439) | 98.74         |
|                         | Acinetobacter baumannii (Z93435) | 99.88         |

the nitrification processes, which may result in increases of NO$_2^-$-N and NO$_3^-$-N in certain periods (Kirimura et al. 1999, Su et al. 2019). However, in the batch experiments with the application of ANRS5, NH$_4^+$-N was largely removed within 70 h without increases of NO$_2^-$-N and NO$_3^-$-N. These results indicate a higher nitrification efficiency of the selected ANRS5 compared to that of EMs. Furthermore, as heterotrophic bacteria, ANRS5 assimilate NH$_4^+$-N, NO$_2^-$-N and NO$_3^-$-N as cell components, as evidenced by the continuous decreases in NO$_2^-$-N and NO$_3^-$-N during the process (Yun et al. 2019).

The immobilized ANRS5 (T5) was able to remove more NH$_4^+$-N (16%), NO$_2^-$-N (14%) and NO$_3^-$-N (17%) compared to the dissociated ANRS5 (T4). These enhancements were attributed to the maize straw biochar since its large SSA and complex pore structure contributed to nitrogen adsorption, which was observed in T3. Furthermore, the attachment sites on the biochar surface provided a habitat for ANRS5, which was beneficial for its growth and prevented bacterial diffusion caused by water flow fluctuations, thus allowing the high nitrogen assimilation activities of ANRS5 (Dalgaard et al. 2011, Rezaei Rashti et al. 2019). The NH$_4^+$-N removal efficiency in T4 was approximately 28% higher than that in T2 with application of EMs. This result demonstrated the feasibility of using ANRS5 as an alternative to EMs in controlling water quality. The use of EMs is currently the most widely applied water purification method due to the multiple functions induced by the included pro-
Biotic bacteria, such as photosynthetic bacteria, bacillus and actinomycetes (Peng et al. 2018). Nevertheless, with the addition of the same cell quantity, the NH4-N removal performance of the EMs was not as good as that of ANRS5. Moreover, the NO3-N content in T2 significantly increased from 0.92 to 3.05 mg l\(^{-1}\), which suggests that EMs might not be able to assimilate NH4-N.

Similar enhancements were also obtained in TP removal; the relevant removal rates were ranked as T5 (54.3 %) > T4 (35.4 %) > T3 (13.2 %) > T2 (10.4 %). Based on these results, we speculate that *Acinetobacter* spp. (PRS4) utilized the aquaculture sewage phosphorus. Heterotrophic bacteria such as PRS4 are thought to be able to store polyphosphate in their cells under aerobic or anoxic conditions, which would enable the simultaneous assimilation of phosphorus during energy uptake from external carbon sources (Burut-Archanai et al. 2013, Liu et al. 2019). This assumption was verified by the existing literature, confirming the phosphorus removal potential of *Acinetobacter* spp. The same mechanisms mentioned above could explain the TP removal enhancement by biochar-immobilized PRS4. In addition, the phosphorus accumulation capacity of PRS4 was much higher (25 %) than that of the EMs in this study, which also indicates that PRS4 is more suitable than EMs for TP removal.

### 3.4. Assessment of water quality and bio-safety with utilization of biochar-immobilized isolated bacteria for practical crab aquaculture

Aquaculture water quality is one of the main factors influencing the environmental ecology and health of aquatic animals. In our study, dynamic variations in DIN (NH4-N, NO3-N and NO2-N) and TP levels were monitored during 90 d of *Eriochir sinensis* cultivation. The results can guide optimization of the applied BIAP dosage, thus reducing operating expenses for water purification. A total of 3 treatments with BIAP dosages of 2 g l\(^{-1}\) (T7), 4 g l\(^{-1}\) (T8) and 6 g l\(^{-1}\) (T9) were conducted; the results are illustrated in Fig. 5.

Clearly, in the control group (T6) without the addition of BIAP, the DIN and TP contents increased from 0.8–10.1 and 0.2–2.6 mg l\(^{-1}\), respectively, with the continuous addition of fish bait. With the addition of BIAP, these concentrations were maintained within a certain range with only small fluctuations. It should be noted that with the application of BIAP at a level of 4 g l\(^{-1}\), the level of nutrients fluctuated below the clean water threshold for aquaculture (DIN: ≤1 mg l\(^{-1}\); TP: ≤0.2 mg l\(^{-1}\)) as defined in the environmental quality standard for surface water (GB 3838-2002). By further increasing the dose (6 g l\(^{-1}\)) of BIAP, more substantial water purification effects were obtained, with DIN and TP levels fluctuating around approximately 0.45 and 0.09 mg l\(^{-1}\), respectively. These results suggest that, in practical applications, immobilized ANRS5 and PRS4 are capable of transforming different forms of nitrogen and phosphorus into their respective components in several biochemical steps (Cai et al. 2007, Guedes et al. 2019, Qin et al. 2019). These immobilized strains likely experience rapid adaptation and growth in aquatic environments, and play an important role in nitrogen and phosphorus cycles. Furthermore, at 4 g l\(^{-1}\) BIAP, the cost of this application (including raw materials, production and labor) was calculated to be approximately US$10 per 1000 m\(^{2}\), which is quite acceptable.

The bio-safety of the isolated strains to aquatic animals in farming environments has not received great attention. In our study, this issue was related to the survival rate, weight, stocking density, biomass, crude protein, crude fat and water content of *E. sinensis* with BIAP application (4 g l\(^{-1}\)); the results are shown in Fig. 6 and Table 5. There was no significant difference in survival rate between the control and experimental treatments during crab cultivation. In addition, the cumulative survival rate of *E. sinensis* in the experimental group (88 %) was slightly higher than that in the control group (86 %) at harvest. With respect to yield and quality, crab weight, stocking density, biomass, crude protein, crude fat and water content in the experimental group were 231 g ind.\(^{-1}\), 12.3 kg m\(^{-2}\), 0.62 g cm\(^{-3}\), 18.1 %, 62.3 % and 10.8 %, respectively, which were improvements compared to those in the control group. Considering the very large cell quantity (approximately 21.6 × 10\(^{8}\) cfu l\(^{-1}\)) that was added to the aquatic environment, the results indicate that BIAP presents no bio-safety concerns for *E. sinensis* and might represent a potential economic value for aquaculture operations.

### 4. CONCLUSIONS

The biochar prepared from maize straw possessed a large SSA and complex pore structure, which was beneficial for immobilization of the isolated bacteria. The satisfactory surface physico-chemical properties of the prepared biochar also enhanced the
adsorption of pollutants in the aquaculture sewage. The isolated *Sphingomonas* sp. PDD-57b-25 and *Acinetobacter towneri* were more efficient in assimilating DIN and polyphosphate compared to currently used bacterial agents. With the application of a specific amount of BIAP, the water quality in a recirculating *Eriocheir sinensis* aquaculture system was maintained in a healthy range according to the relevant water quality standard. In addition, BIAP presented no bio-safety risks for *E. sinensis*, which demonstrates its potential economic value in sewage remediation and aquaculture applications.

In summary, the results in the present study are important in the context of the existing literature and for the practical application of biochar-immobilized bacteria as a decontamination technology for agricultural sewage. Moreover, we believe that the experimental and evaluation methodology developed here may be applied as a useful tool for practical intensive aquaculture in other RASs.
Table 5. Mean (±SD) yield and quality of Eriocheir sinensis crabs with practical application of biochar-immobilized ammonia nitrogen and phosphate removal strains (ANRS and PRS; 4 g l⁻¹). Control group was given no biochar-immobilized ANRS and PRS; experimental group was challenged by biochar-immobilized ANRS and PRS. Crabs sampled per treatment = 50; all measured crab parameters were significantly different between treatments at p < 0.01

| Indexes          | Unit       | Control group | Experimental group |
|------------------|------------|---------------|--------------------|
| Weight           | g ind⁻¹    | 182 ± 4.5     | 231 ± 6.3          |
| Stacking intensity | kg m⁻²     | 9.7 ± 0.8     | 12.3 ± 1.4         |
| Biomass          | g cm⁻³     | 0.48 ± 0.05   | 0.62 ± 0.07        |
| Crude protein    | %          | 16.3 ± 0.3    | 18.1 ± 0.4         |
| Crude fat        | %          | 58.6 ± 2.7    | 62.3 ± 2.3         |
| Water content    | %          | 9.4 ± 0.7     | 10.8 ± 0.9         |

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