Nitrogen Removal Performance and Metabolic Pathways Analysis of a Novel Aerobic Denitrifying Halotolerant *Pseudomonas balearica* strain RAD-17

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**Abstract:** An aerobic denitrification strain, *Pseudomonas balearica* RAD-17, was identified and showed efficient inorganic nitrogen removal ability. The average NO$_3$–N, NO$_2$–N, and total ammonium nitrogen (TAN) removal rate (>95% removal efficiency) in a batch test was 6.22 mg/(L·h), 6.30 mg/(L·h), and 1.56 mg/(L·h), respectively. Meanwhile, optimal incubate conditions were obtained through single factor experiments. For nitrogen removal pathways, the transcriptional results proved that respiratory nitrate reductases encoded by *napA*, which was primarily performed in aerobic denitrification and cell assimilation, were conducted by *gluS* and *gluD* genes for ammonium metabolism. In addition, adding the strain RAD-17 into actual wastewater showed obvious higher denitrification performance than in the no inoculum group (84.22% vs. 22.54%), and the maximum cell abundance achieved 28.5 ± 4.5% in a ratio of total cell numbers. Overall, the efficient nitrogen removal performance plus strong environmental fitness makes the strain RAD-17 a potential alternative for RAS (recirculating aquaculture system) effluent treatment.

**Keywords:** aerobic denitrification; *Pseudomonas balearica* RAD-17; nitrogen removal; metabolic pathways; bioaugmentation

1. Introduction

Biological nitrogen removal is crucial for wastewater treatment, and the heterotrophic denitrification is the most selective method due to its high efficiency and flexibility [1]. During the process, organic carbons are necessary to supply the electrons to the nitrate. Substances such as methanol, ethanol, etc., were commonly used [2]. However, precise liquid organic was difficult to measure, and an overdose can add organic loading to the bio-filter, which would have negative effects on the aquaculture system stability [3]. In wastewater with low C/N ratio characteristics, solid-phase denitrification that use biodegradable polymer as a simultaneous organic carbon source, as...
well as biofilm carriers, are considered more appropriate in specific fields [4]. For example, in recirculating aquaculture system (RAS) effluent, PBS (poly-butylene succinate) or PHBV (poly-3-hydroxybutyrate-co-3-hydroxyvalerate)-based denitrifying reactors demonstrated efficient nitrogen removal performance [3,5]. Therefore, using biodegradable polymer as slow-release organic will also have additional benefits to increase the fish survival rate [3,6].

The drawback of the biodegradable polymer-based denitrification is the relative high cost in media when compared with liquid carbon [4]. On the other hand, residual DOC (dissolved organic carbon) in effluent commonly appeared in solid-phase denitrifying reactors [3,7,8], which indicated that the entirety of the carbon source was not optimally used since part of degrading bacteria have an incapable ability in denitrification [9]. Moreover, the residual organic substances were considered to support DNRA (dissimilatory nitrate reduction to ammonium) over denitrification [10], as well as the SRB (sulfate reduced to sulfide) process [3], especially in marine environment conditions. Therefore, enhancement of the denitrification performance is crucial for this technology in practice.

An interesting alternative to promote the denitrification performance is the supplement of functional bacteria through bioaugmentation technology [11]. In a previous study, adding the Diaphorobacter polyhydroxybutyricum strain SL-205 enabled rapid reactor startup and improved nitrate removal performance when compared with active sludge inoculation in an anoxic solid-phase denitrification reactor [12]. This indicated that the initial microbial regulation can support relative bacterial community in reactors. Furthermore, to suppress the DNRA and SRB pathways in a marine PBS denitrification reactor, the alternant aerobic/anoxic operations instead of continuous anoxic were demonstrated to be feasible in our previous study [6]. Therefore, the bacteria screening for bioaugmentation need strong fitness in such salinity and aerobic conditions.

Traditional denitrification or the similar reduction process only appeared under anoxic conditions [13,14], as the narG gene encoded for the nitrate reductase is sensitive to oxygen presence, which could block the sequential energy and electron transfer under aerobic condition [15]. Recently, many aerobic denitrifying groups were found to support a potential pathway for a biological nitrogen removal process [16]. The main characteristics of these strains are that they have a gene cluster of the napFDAGHBC family while napA was responsible for synthesis of the catalytic subunit for electron delivery from NADH+ to nitrate aerobically [17]. Under aerobic or alternate aerobic/anoxic conditions, the nitrate reductase encoded by the napA gene, which is located in the periplasm, was primarily infiltrated by nitrate and oxygen as compared to the narG gene located in plasma membrane [18]. Therefore, the high activity of the napA gene make those aerobic denitrifying strains use nitrate over oxygen preferentially [16]. Potentially, this wide niche can reduce technological requirements in current biological nutrient treatment processes, which always cause temporal (Anoxic/Oxic, A/O process) or spatial (sequencing batch reactor, SBR process) division for different microbota [19]. Until now, many aerobic denitrifiers were reported in plenty of genera including Pseudomonas stutzeri YZN-001 [20], Acinetobacter sp. HA2 [21], Pseudomonas stutzeri T13 [22], Marinobacter hydrocarbonoclasticus RAD-2 [23], Pseudomonas stutzeri C3 [24], etc. In addition, several strains were also found to have aerobic ammonium removal ability, which show potential through heterotrophic nitrification or assimilation pathways [22,24–26].

In this study, a novel aerobic denitrifying halotolerant strain, Pseudomonas balearica RAD-17, was isolated for a long-term PBS-supported denitrification reactor for RAS effluent treatment, demonstrated in a previous study [6]. The 16S rDNA gene was amplified to identify the phylogenetic relationship for the isolated strain, while API 20NE (analytical profile index of Gram-negative with non-Enterobacteriaceae) was used for its physiological feature. Meanwhile, the inorganic nitrogen removal performance was also evaluated by different nitrogen sources. Moreover, the aerobic nitrogen metabolic pathways were investigated by quantifying the key denitrifying genes (napA, nirS, norB, and nosZ) and glutamic biosynthesis genes (glaD and gluS) that are potentially related with ammonium assimilation. In addition, the strain’s bioaugmentation performance was also evaluated by adding it into actual RAS effluent. To the best of our knowledge, this is the first report of a functional strain with efficient aerobic nitrogen removal ability in the Pseudomonas haloterrica species.
Overall, the results might provide new insight in aerobic denitrifying microbial resources and potential alternatives for enhancing nitrate-removal performance for RAS practice.

2. Materials and Methods

2.1. Cultured Media

The culture media used in this study were according to our previous study [23]. The LB (Luria-Bertani) media was prepared by using 5.0 g/L yeast extract, 10.0 g/L peptone, 25.0 g/L NaCl, and 1.5% (v/v) agar. The DM (denitrification media) was prepared in the ratio of 2.0 g/L sodium acetate, 2.0 g/L KNO₃ (or NaNO₃), 0.2 g/L MgSO₄⋅7H₂O, 1.0 g/L K₂HPO₄, and 1.0% (v/v) trace-element solution for aerobic denitrification performance evaluation. The HNM (heterotrophic nitrification media) was prepared as follows: 2.0 g/L sodium acetate, 0.3 g/L NH₄Cl, 0.2 g/L MgSO₄⋅7H₂O, 6.7 g/L NaHPO₄, 1.0 g/L KH₂PO₄, and 1.0% (v/v) trace-element solution for ammonium-nitrogen removal evaluation. The trace-element solution contained 50.0 g/L EDTA, 2.2 g/L ZnSO₄, 5.5 g/L CaCl₂, 5.06 g/L MnCl₂⋅4H₂O, 5.0 g/L FeSO₄⋅7H₂O, 1.1 g/L (NH₄)₆Mo₇O₂₄⋅4H₂O, 1.57 g/L CuSO₄⋅5H₂O, and 1.61 g/L CoCl₂⋅6H₂O. In addition, the amounts of nitrogen and carbon in DM or HNM can also change according to the experimental setting. The initial pH of all media was set to 7.2 and then autoclaved for 20 min at 121 °C.

2.2. Bacteria Isolation and Identification

The RAD-17 strain was screened from a long-term PBS based denitrifying reactor, which operated under alternant aerobic/anoxic conditions in our previous study [6]. The reactor influent contained around 10 mg/L NH₄-N and 150 mg/L NO₃-N and showed average TAN and nitrate removal rates of 47.35 ± 15.62 g NH₄-N m⁻³ d⁻¹ and 0.64 ± 0.14 kg NO₃-N m⁻³ d⁻¹ with no obvious nitrite accumulation [6]. For screening, a 15 mL mixture of mature PBS and solutions were transferred into a 150 mL flask aseptically with 100 mL LB media for 10 days of preculture. The temperature and revolution were set at 30 °C and 150 rpm (revolutions per minute), respectively. Afterwards, the homogenized suspensions were serially diluted and plated using a DM media and incubated at 30 °C for 72 h. Then, a single colony with a pale-yellow circle was dilution-streaked onto a DM agar plate for further purification. Finally, a strain of the *Pseudomonas baharia*., named RAD-17, was isolated. The genomic DNA of the RAD-17 strain was extracted using a DNA extraction kit (TaKaRa Biotechnology Co. Ltd, Beijing, China). The 16S rRNA amplified product was sequenced by the Zhejiang Institute of Microbiology (Hangzhou, China). Phylogenetic relationships of the strain RAD-17 with other denitrifying bacteria were constructed using the molecular evolutionary genetics analysis software (MEGA 5, The Biodesign Institute, Tempe, USA). In addition, the purified strain RAD-17 was also stored in a 30% glycerol solution at -80 °C for following experiments.

2.3. Nitrogen Removal Performance

The inorganic nitrogen removal performance of the strain RAD-17 was evaluated on DM or HNM media. For aerobic denitrification capacity, a sole nitrogen source of NO₃–N (around 300 mg/L) or NO₂–N (around 300 mg/L) was tested in DM media, which contained KNO₃ or NaNO₃, respectively. For heterotrophic ammonium removal, sole nitrogen source of NH₄Cl (around TAN 80 mg/L) was carried out in similar operation in HNM media. For process, 3% (v/v) seed suspension was inoculated in 250 mL Erlemeyer flasks and cultured for 48 h under aerobic condition at 25 °C and 150 rpm, respectively. Meanwhile, the cell growth (O₆₅₀ value) and nitrogen concentrations were measured every 4 h.

2.4. Single-Factor Experiments

Single-factor experiments were also carried out to evaluate the effect of various conditions on the aerobic denitrification performance of the strain RAD-17, for optimized incubated conditions. The basal condition was determined as follows: NO₃–N concentration of 300 mg/L, C/N ratio 10, NaCl
concentration 25‰, temperature 25 °C, rotation 150 rpm, and 3% inoculation (v/v). On C/N ratio test, the C/N ratios were set to 2, 5, 10, 15, and 20. On salinity test, the NaCl concentrations were set at 0‰, 2.5‰, 5‰, 15‰, and 25‰. On carbon sources, fructose, sodium acetate, lactin, glucose, and sodium citrate were tested. On revolution test, the speeds were set to 0 rpm, 50 rpm, 100 rpm, 150 rpm, and 200 rpm. On temperature test, 5 °C, 15 °C, 25 °C, and 40 °C were used. All tests were conducted in triplicate and non-seeded samples were used as blank control.

2.5. qRT-PCR Analysis

The transcripational level gene expression intensity of the strain RAD-17 on nitrogen removal processes was investigated to reveal the metabolic pathways. In this study, real-time quantitative PCR was conducted to amplify the denitrifying genes napA, nirS, norB, nosZ, and the ammonium incorporation genes gluD (NADP-specific glutamate dehydrogenase) and gluS (glutamate synthase) with RNA samples in 48-h experiments. All primers were designed by the genome sequence of the strain RAD-17 and are listed in Table 1. The amplification specificities of these primer pairs were verified through agarose gel electrophoresis (Figure S2). The housekeeping gene 16S ribosomal RNA was used as an internal control to normalize the data. Total RNA extraction and cDNA synthesis were performed by using an RNAprep Bacteria Kit and a FastQuant RT Kit (Tian Gen BiotechCo. Ltd, Beijing, China), respectively. PCR amplification was performed with the following protocol: Initial denaturation at 95 °C for 2 min followed by 40 cycles of denaturation at 95 °C for 15 s, annealing at 55 °C for 15 s, and synthesis at 72 °C for 15 s; a melting curve was generated by linear heating from 70 °C to 95 °C over 25 min [27]. All quantitative amplifications were conducted in triplicate using the SYBR Green Real-Time PCR Kit (Novland, Shanghai, China) and respective primers on a StepOne PCR instrument (Applied Biosystems, Forest City, CA, USA).

Table 1. All the primers used in this study for Pseudomonas balearica strain RAD-17.

| Gene | Primer sequences (5’-3’) | References |
|------|--------------------------|------------|
| 16S rRNA | F: CCTACGGGAGGCAGCAG | This study |
| | R: ATTACCGCGGCTGCTGG | |
| gluD | F: GCTATGGCACAGAGAAC | This study |
| | R: CATCACTTCGTTGTCGCTC | |
| gluS | F: CGCAACATCTTCTCCAACC | This study |
| | R: TICTCTCACCCCATTCGAC | |
| napA | F: TTCATGGCCTGCTGTACCTG | This study |
| | R: TCATCCTGGCAATCGAAC | |
| nirS | F: TGGAAAGCCAGATGCAGCAC | This study |
| | R: ACGCTCCTTGACGAAGTGGATG | |
| norB | F: TTCTACAACCCGGAGAACC | This study |
| | R: GCCATGATGCTACAGGCC | |
| nosZ | F: CAACATCGCCAGATCGAAG | This study |
| | R: TGCACTAGTACCAGTGCAG | |

2.6. Bioaugmentation Performance Evaluation

The strain RAD-17 was added into actual RAS effluent to evaluate its bioaugmentation performance. The experimental tanks had a total volume of 10 L. A total of 5 L was used for raw RAS wastewater. The NO₃–N concentration and the C/N ratio were adjusted at approximately 100 mg/L and 15 by adding KNO₃ and sodium acetate, respectively. Then, 500 µL of the strain RAD-17 (OD₆₀₀ = 1.0) solution was added into the experimental tanks and aerobically cultured for 140 h. Another group without inoculation operated as control. Air-pumps (ACO-003, 120 W, Sengseng Co., Ltd., Taipei, Taiwan) were used for aerobic condition and temperature was set at 25 ± 1 °C in a thermostatic chamber. All treatments were carried out in triplicate.
During the experimental phase, 2 mL solutions were cultured on DM medium to evaluate the potential denitrification strains by the most probable number (MPN) standard method. Here, the amount of CFU can partly reveal the potential denitrifying ability as DM is a specific media for the denitrifier [28]. Meanwhile, the growth rate of the strain RAD-17 was also detected through monitoring the copies ratio of the napA gene by the strain RAD-17 that is relative to the total 16S rDNA genes using absolute qPCR. The qPCR primers are listed in Table 1. In the absolute qPCR assay, the standard curves were constructed using serial dilution of purified target DNA from PCR amplification. The amount of the template DNA was determined by the NanoDrop ND-2000 ultraviolet absorption assay. There are different gene copies between the napA gene and the 16S rDNA gene. The average number of 16S rDNA copies per bacterial cell is 4.2, and the napA gene carries a single copy in the strain RAD-17 cell [19]. The different copy numbers of two genes were used to normalize the ratio of qPCR data.

2.7. Analytical Methods

The solution samples were filtered through a 0.45 µm filter membrane before analyzing. The TAN, NO3−:N, and NO2−:N concentrations were analyzed according to standard methods [29]. Bacteria biomass was measured by ODo value using a spectrophotometer at 600 nm (Agilent Technologies Cary 60 UV-vis, Santa Clara, USA). The morphology of the strain RAD-17 was observed by a scanning electron microscope (SEM) (SU8010, Hitachi High-Technologies Corporation, Tokyo Japan). The pH value was measured using a portable pH meter (58, Mettler Toledo, Zurich, Switzerland). DO was measured using a DO meter (SC9-FK2, Mettler Toledo, Zurich, Switzerland).

Physiological and biochemical characteristics were tested using API 20NE kits (BioMérieux Shanghai Co. Limited, Shanghai, China), and test strips were checked after incubation for 24 h [23]. Nitrogen balance analysis was done according to the previous study. For process, 3% (v/v) seed suspension was inoculated in 250 mL Erlenmeyer flasks and cultured for 20 h under aerobic condition at 25 °C and 150 rpm, respectively. The incubated nitrate was set at around 30 mg/L. Then, the nitrogen balance can be calculated on the initial and final nitrogen concentration [30].

3. Results and Discussion

3.1. Bacteria Characteristics and Identification

For strain screening, more than 20 pure isolates were obtained from the DM medium, while one named RAD-17 showed the highest aerobic denitrification performance. The colonies of the RAD-17 strain were pale yellow, salient, semitransparent, circular in shape, and presented a moist surface on the LB medium. The RAD-17 strain was a gram-negative strain with a bacilliform sharp in size of around 0.3-0.4 µm in diameter and 0.8-1.6 µm in length, respectively (Figure S1). The 16S rDNA gene sequence was submitted to the NCBI database with the accession number MK881511, and the highest similarity of the RAD-17 strain was found with Pseudomonas balearica DSM 6083. The phylogenetic analysis (threshold 100%) further confirmed the identification of the RAD-17 strain as Pseudomonas balearica (Figure 1). For nitrogen removal, many strains in the genus Pseudomonas were demonstrated to have aerobic denitrification ability, such as Pseudomonas stutzeri C3 [24], Pseudomonas stutzeri T13 [25], Pseudomonas stutzeri YZN-001 [20], Pseudomonas stutzeri PCN-1 [31], Pseudomonas tolai Y-11 [32], etc. In addition, several groups also have ammonium removal capacity under aerobic conditions [22]. However, in Pseudomonas balearica, though strain DSM6083 presented genome information (ASM81801v1), no study on the denitrifying function was reported in this sub-lineage.
Figure 1. Neighbor-joining phylogenetic tree based on 16S rRNA gene sequences showing the position of the RAD-17 strain and closely related strains. Bootstrap values based on 1000 replicates are presented in branch nodes.

The API 20NE tests were carried out for further identification of the physiological and biochemical characteristics of the strain RAD-17 (Table 2). The RAD-17 strain was positive for nitrate reduction, but was negative for urease, β-glucosidase, protease, and β-galactosidase. On cell biosynthesis, it could use glucose, maltose, gluconate, capric acid, etc., while arabinose, mannose, mannitol, and N-acetyl-glucosamine could not be assimilated.

Table 2. of the RAD-17 strain determined by analytical profile index of Gram-negative with non-Enterobacteriaceae (API 20NE) tests.

| API 20NE Results | Strain RAD-17 |
|------------------|---------------|
| Oxidase test     | +             |
| Nitrate reduction| +             |
| Arginine dihydrolase | --         |
| Urease           | --            |
| β-Glucosidase    | --            |
| Protease         | --            |
| β-Galactosidase  | --            |
| Assimilation of  |               |
| Glucose          | +             |
| Arabinose        | --            |
| Mannose          | --            |
| Mannitol         | --            |
| N-acetyl-glucosamine | --          |
| Maltose          | +             |
| Gluconate        | +             |
| Capric acid      | +             |
3.2. Nitrogen Removal Performance

3.2.1. Nitrogen Removal Ability

The aerobic inorganic nitrogen removal capacity of the strain RAD-17 is shown in Figure 2. In general, the strain RAD-17 can use three typical nitrogen forms, such as nitrate, nitrite, and ammonia. For aerobic denitrification, nitrate as sole nitrogen showed a quick start with the NO₃⁻-N concentration decreasing from the initial 301.43 ± 1.06 mg/L to a final 2.94 ± 1.66 mg/L, which indicated a 99.02% removal efficiency (Figure 2A). The lag phase was only observed during the initial 8 h, and the logarithmic growth phase occurred during the following 8 h. Nitrite accumulation was also presented and the peak concentration of 69.67 ± 6.64 mg/L was found in 16 h but disappearing rapidly after 20 h. The strain biomass reached its maximum OD₆₀₀ value with 1.61 ± 0.08 in 24 h but then slightly decreased. The reason might be that the nitrogen substance was consumed in this period. Additionally, when using nitrite as sole nitrogen, a backward lag phase was observed in 0–24 h, though the NO₂⁻-N concentration also decreased from 302.27 ± 1.11 mg/L to 0.56 ± 0.37 mg/L rapidly (Figure 2B). The final strain biomass was around 0.74 ± 0.04, which was lower than using nitrate as the nitrogen source that potentially indicated that nitrite added toxicity to the cell.

For the aerobic ammonium removal ability of the strain RAD-17 is illustrated in Figure 2C. When using ammonium as nitrogen source, no obvious lag phase occurred, indicating that ammonium should be an accessible element for cell growth. The TAN concentration decreased from the initial 77.73 ± 2.35 mg/L to the final 2.94 ± 1.66 mg/L, resulting in 1.56 mg TAN·L⁻¹·h⁻¹ removal rate. This phenomenon could also be supported by the higher OD₆₀₀ value of 2.24 ± 0.05, which revealed more efficient strain yield when compared with nitrate or nitrite as substance. It was interesting to note that a slight nitrate concentration of maximum 8.79 ± 0.80 mg/L was accumulated in 28–48 h. Under the current condition, ammonium was the sole nitrogen source, so the mechanism and transfer pathway of ammonium to nitrate and nitrogen gas still needs further study.
3.2.2. Nitrogen Balance Analysis

The result of the nitrogen balance analysis is shown in Table 3. Under aerobic denitrification, the strain RAD-17 gained 87.76% nitrogen loss, which indicated that the nitrate substance prioritized transfer to gaseous products rather than biomass synthesis. Similar reports which showed 12.6% cell assimilation from nitrate were presented in Paracoccus versutus KS293 [35] and 19.8% in Pseudomonas stutzeri ZF31 [30]. On the other hand, using ammonium as sole nitrogen source, increased cell yield revealed by higher OD<sub>600</sub> value was obtained (Figure 2C), which were also consisted with nearly 50% ammonium-nitrogen assimilated in Acinetobacter sp. HA2 and Paracoccus versutus LYM, respectively [21,26].

Table 3. Balance analysis for the strain RAD-17 under aerobic denitrification.

| Substance | Initial TN (mg/L) | NO<sub>3</sub>−-N | NO<sub>2</sub>−-N | NH<sub>4</sub>+-N | Organic-N | Intracellular N | N loss (%) |
|-----------|------------------|-----------------|--------------|-------------|-------------|----------------|-----------|
| Nitrate   | 30.56 ± 0.02     | 0.14 ± 0.03     | 0.05 ± 0.02  | 0.17 ± 0.06 | 0.15 ± 0.04 | 3.23 ± 0.34    | 87.76     |

Note: N loss = (Initial TN−Final N−Intracellular N)/Initial TN * 100%.
3.2.3. Single Factor Experiments

The aerobic denitrification performance under different single factor conditions is shown in Table 4. In general, the C/N ratio and the kinds of carbon source should strongly associate with nitrate removal efficiency, due to this server as electron donor and energy support [13,15]. For carbon sources, the strain RAD-17 was found unable to metabolize lactin while fructose also obtained low efficiency. In contrast, more than 90% nitrate removal performance was presented in glucose, sodium acetate, and sodium citrate, respectively. In general, these substances were thought to be easily utilized as raw material for a TCA (tricarbosylic acid) cycle for maximum energy efficiency and ATP (adenosine triphosphate) synthesis under aerobic metabolism [37]. For organic amounts, a C/N ratio with a range of 5–15 was found to have optimal denitrification performance, which gained nearly complete nitrate removal. For a C/N ratio of 2, low nitrate removal performance with residual nitrate indicated electron donor deficiency. In addition, a slight decrease in nitrate removal efficiency was presented in a C/N ratio of 20, which revealed that excess organic substances also had a negative effect on denitrification. Seral similar reports were also found, optimum C/N ratio 5-15 in the Marinobacter hydrocarbonoclasticus RAD-2 strain [23], C/N ratio 15 in Marinobacter sp. Fe [38], C/N ratio 7-9 in Bacillus methylolotrophicus L7 [39], and C/N ratio 6-10 in Pseudomonas stutzeri YG-24 [33].

For salinity, the strain RAD-17 gained commendable denitrification performance in a NaCl concentration range of 0-25%. The similar ODout value and nitrate removal performance were presented in a NaCl amount of 0% and 25%, which indicated that salinity has no effect on growth and the denitrifying activity of the strain RAD-17. An inferential mechanism was that Na+ ion supported from sodium acetate can offset the NaCl deficiency. This was also demonstrated in Marinobacter hydrocarbonoclasticus, as Na+ ion was absolutely required, no matter whether the K+ ion and Cl- ion existed [40], and a minimal amount for its growth is a Na+ ion beyond 0.08 molarity concentration [41]. Therefore, the current phenomenon revealed that the strain RAD-17 might have wide ecological niche fitness in practice.

For temperature, a typical mesophilic characteristic of the strain RAD-17 was present when more than 90% nitrate removal efficiency was found in 15-40 °C. No denitrification occurred under a temperature of 5 °C, which indicated that low a temperature might have a significant negative effect on denitrifying enzyme activity. This result was consistent with previous studies that used mostly mesophilic aerobic denitrifiers [23,42,43].

In this study, the effect of rotation speeds that revealed DO concentrations on denitrification performance were also evaluated. It should be noted that the strain RAD-17 gained both ideal nitrate removal efficiency in anoxic (0 rpm, DO 0.2 ± 0.1 mg/L) and aerobic (150 or 200 rpm, DO 3.3 ± 0.6 mg/L or 4.7 ± 0.9 mg/L) conditions, respectively. However, a decreased denitrification performance was observed in oxygen-limited conditions (50 or 100 rpm, DO 0.9 ± 0.4 mg/L or 1.7 ± 0.8 mg/L). In a previous study, several denitrifying strains, especially in the genus Paracoccus versatius, have both aerobic and anoxic nitrate removal ability. For example, Paracoccus versatius KS293 exhibited 82% and 85% total nitrogen removal under anoxic and aerobic conditions, respectively [35]. Since denitrification is a respiratory process, the regulation of the denitrification respirome in Paracoccus denitrificans is related to transcription factors fnrP, narR, and marR to adopt the oxygen, nitric oxide, and nitrate shift conditions [44,45]. However, whether the Pseudomonas family shares similar pathways or not is still unclear. Therefore, further studies need to reveal the oxygen triggering mechanism for the strain RAD-17 denitrification in future.

Table 4. Varied single factors on the aerobic denitrification performance of strain RAD-17 after 48 h incubation.

| Factor | Variations | Initial Nitrate (mg/L) | Final Nitrate (mg/L) | Final Nitrite (mg/L) | Final TAN (mg/L) | Growth (ODout) |
|--------|------------|------------------------|----------------------|---------------------|-----------------|---------------|
| C/N ratios | 2 | 295.49 ± 0.60 | 149.00 ± 2.87 | 50.21 ± 1.52 | 8.61 ± 1.19 | 0.86 ± 0.16 |
|           | 5 | 298.42 ± 0.26 | 0.92 ± 0.80 | 0.36 ± 0.02 | 1.88 ± 0.04 | 1.17 ± 0.10 |
Temperature

Pseudomaonas balearica in assimilation is still unclear. Therefore, the difference of response and remodeled rules by the respiratory nitrate reductases under transcriptional results, the nitrate distribution were conducted together through a responsible norB reductases related genes were significantly upregulated by nitrate inducing, including 3.3.1.

3.3.1. Microorganisms

Rotation

Carbon

NaCl (%)

Temperature

Carbon source

Fructose

Lactate

Glucose

NaAc

10 299.59 ± 0.37 8.17 ± 1.82 0.34 ± 0.01 1.19 ± 0.03 1.43 ± 0.03
15 301.51 ± 0.71 8.35 ± 3.10 0.30 ± 0.00 1.02 ± 0.03 1.04 ± 0.25
20 300.09 ± 0.71 23.41 ± 9.78 0.64 ± 0.02 0.95 ± 0.00 1.09 ± 0.14
0 302.47 ± 0.19 3.14 ± 2.00 0.34 ± 0.01 0.26 ± 0.08 1.73 ± 0.08
2.5 299.78 ± 0.56 6.52 ± 1.37 0.42 ± 0.03 0.30 ± 0.05 1.30 ± 0.06
5 301.29 ± 0.24 9.37 ± 5.57 0.27 ± 0.03 1.04 ± 0.08 1.30 ± 0.15
15 300.36 ± 0.17 8.11 ± 4.29 0.23 ± 0.07 1.05 ± 0.02 1.40 ± 0.35
25 301.46 ± 0.33 9.04 ± 4.11 0.25 ± 0.04 1.14 ± 0.06 1.66 ± 0.24

Fructose 299.61 ± 0.22 205.59 ± 8.50 5.71 ± 4.06 29.13 ± 12.13 0.27 ± 0.03

NarA 303.14 ± 0.11 2.59 ± 1.11 0.87 ± 0.02 3.34 ± 0.33 1.99 ± 0.11

NorB 298.45 ± 0.27 296.90 ± 2.59 0.36 ± 0.03 – 0.69 ± 0.15

NorB 301.09 ± 0.14 0.57 ± 0.27 0.51 ± 0.04 0.10 ± 0.09 1.86 ± 0.11

NorB 304.56 ± 0.15 19.16 ± 6.26 0.35 ± 0.11 1.01 ± 0.40 1.37 ± 0.06

NorB 302.42 ± 0.31 36.57 ± 4.53 1.07 ± 0.25 1.68 ± 0.57 0.76 ± 0.05

NorB 301.77 ± 0.17 116.52 ± 9.91 13.74 ± 0.20 0.90 ± 0.05 1.07 ± 0.11

NorB 299.46 ± 0.20 8.99 ± 1.33 0.54 ± 0.04 1.71 ± 0.29 2.00 ± 0.07

NorB 301.26 ± 0.05 2.59 ± 0.94 0.53 ± 0.16 1.86 ± 0.93 2.19 ± 0.06

NorB 299.76 ± 0.16 298.57 ± 1.50 – 0.45 ± 0.29 0.74 ± 0.06

NorB 298.31 ± 0.09 7.79 ± 0.91 0.62 ± 0.06 1.63 ± 1.42 1.89 ± 0.18

NorB 300.55 ± 0.11 2.28 ± 2.21 0.41 ± 0.07 2.49 ± 0.10 2.03 ± 0.25

NorB 300.98 ± 0.17 18.51 ± 5.32 0.60 ± 0.18 2.50 ± 1.20 1.70 ± 0.15

3.3. Nitrogen Metabolism Pathways Analysis

3.3.1. Aerobic Denitrification Pathway

The transcriptional expression levels of the denitrification genes under aerobic condition of the RAD-17 strain are shown in Figure 3. The results reveal that the expression of four respiratory nitrate reductases related genes were significantly upregulated by nitrate inducing, including narA, nirS, norB, and nosZ. The narA firstly showed a quicker response to nitrate than nirS, norB, and nosZ, which is consistent with the fact that this process is a sequence of electrons transfer [13,18]. However, though the nosZ and nirS were present, which was consistent with the nitrite peak concentration after 16 h (Figure 2A). It was interesting to note that the enhanced expression of norB, which lasts 16 hours incubation, presented a maximum intensity earlier than nirS and nosZ. In Pseudomonas stutzeri PCN-1, coincident peak expression was found in nirS, norB, and nosZ [31]. Therefore, these results implied that the environmental signals NO− and NO might both be the indirect inducer for norB [13]. In a previous study, the transcriptional activators, ferP, narR, and narK, were thought to be responsible as primary effectors to oxygen, NO, and NO2/NO2: [13,44,45]. Furthermore, when using nitrate as sole nitrogen source, both respiratory nitrate reductases and assimilatory nitrate reductases were conducted together through a corporate chaperone encoded by narA in the Paracoccus denitrification strain [46]. However, though this study proved the aerobic nitrate removal was caused by the respiratory nitrate reductases under transcriptional results, the nitrate distribution mechanism in assimilation is still unclear. Therefore, the difference of response and remodeled rules in Pseudomonas balearica with other aerobic denitrifiers need to be further studied.
3.3.2. Heterotrophic Ammonium Removal Pathway

The transcriptional expression levels of the ammonium assimilation genes under aerobic condition of the strain RAD-17 are shown in Figure 4. In general, ammonium assimilation into different amino acids was the start for glutamate. Based on the KEGG nitrogen metabolism pathways, there are two major biosynthesis pathways of ammonium into L-glutamate, which involve glutamate dehydrogenase (1.4.1.2, 1.4.1.3, 1.4.1.4) as well as the glutamine synthetase (6.3.1.2) and glutamate synthase (1.4.1.13, 1.4.1.14, 1.4.7.1) [6]. In this study, an obvious up-regulation of gltS and gltD genes occurred during 4–16 h, which were approximately 420 and 3100-folds compared with the control sample, respectively. The ammonium concentrations also showed apparent consistency in this phase with a sharp decrease (Figure 2C). A similar phenomenon was also reported in other aerobic denitrificans, like *Pseudomonas stutzeri* T13 [22], *Acinetobacter* sp. HA2 [21], *Paracoccus versatius* LYM [36], and *Klebsiella* sp [47]. Therefore, this indicated that a certain assimilation pathway is performed in the strain RAD-17 when using ammonium as the sole nitrogen source.

On the other hand, the respiratory nitrate reductases-related genes of napA, nirS, norB, and nosZ did not show obvious enhanced expression during 0–24 hours, which indicated that ammonium should not be a direct inducer for aerobic denitrification. It should be noted that a slight nitrate accumulation occurred between 24–48 hours (Figure 2C), which also caused inconspicuous increase in the expression of nitrate reductases genes (Figure 4). In a previous study, ammonium translated into nitrate in aerobic denitrifications through hydroxylamine related genes [47]. However, we searched the whole genome of the strain RAD-17 (data not shown), but no hydroxylamine genes were found, which indicated that another potential pathway existed for the nitrate production and further denitrification. Thus, the results indicated that a novel pathway was existed for ammonia change to nitrogen-gas that none study was reported before. Hence, a hypothesis was proposed that the strain RAD-17 was inclined to reserve nitrate under ammonium feast condition. Basically, ammonium is a more available nutrient for microbes, while nitrate is a selective substance. Therefore, this might be a characteristic of the strain RAD-17 to fit a wider niche, but the mechanism needs further study.
3.4. Bioaugmentation Performance Evaluation

The bioaugmentation performance by adding the strain RAD-17 into actual RAS effluent is shown in Figure 5. In general, the inoculated groups have higher nitrate removal efficiency compared with control groups (84.22% vs. 22.54%). The nitrate concentrations decreased from initially 99.27 ± 0.43 mg/L to 15.66 ± 3.85 mg/L or 76.89 ± 5.79 mg/L, respectively. In addition, the inoculated groups also gained relative higher pH values and lower DO values (Figure 5A, C), since heterotrophic denitrification was an alkalinity produced process [48]. Meanwhile, the strain RAD-17 also showed obvious preponderant cell abundance in wastewater, which revealed its potentially strong fitness in environment. The maximum denitrifying strains of 4.9 × 10^{7} ± 2.0 × 10^{6} cells/mL was found after 36 h of incubation by inoculated the strain RAD-17, while only 5.1 × 10^{5} ± 2.0 × 10^{6} cells/mL was gained in control groups (Figure 5B). It should be also noted that after 50 h, the relative abundance of strain RAD-17 was decrease, and the reasons might be related to substance insufficient or other strain competitive. Based on the qPCR detection, the strain RAD-17 occupied a peak ratio of 28.5 ± 4.5% in total strain cell numbers (Figure 5D). No significant amplification of the napA gene was detected in control groups by gel electrophoresis analysis (Figure S3). Bioaugmentation was a convinced technology to improve bioremediation system performance [49]. Therefore, our results provided clear evidence that the strain RAD-17 can function as efficient nitrate removal in RAS effluent treatment.
Figure 5. The bioaugmentation performance of the RAD-17 strain in actual recirculating aquaculture system (RAS) effluent. (A) pH and DO values; (B) maximum potential denitrifying strains numbers;
(C) inorganic nitrogen concentrations; (D) strain RAD-17 abundance. Data shown are means ± SD (error bars) from three replicates.

3.5. Research Prospective

A proposed model for aerobic nitrogen removal mechanisms of the strain RAD-17 is shown in Figure 6, which indicated the convinced substance utilization and electron transportation pathways. The nitrate reductases that encoded at least five denitrification relative genes, napA, napB, napC, napD, and napE, were found to orderly arrange in clusters by genome annotation of the strain RAD-17, which should support the high nitrate removal efficiency. Respiratory nitrate reductases were carried out by napA primarily for aerobic denitrification, as well as the cell assimilation were a predominant approach started from gluS and gluD genes for ammonium metabolism. Taking into consideration the halotolerant characteristic and bioaugmentation performance, the above abilities support that the strain RAD-17 owns width ecological niche, and thus might have stronger fitness in the application.

However, the current phenomenon also indicated that deeper insightful research should be done for characterizing the strain RAD-17. The express mechanism of respiratory nitrate reductases and assimilatory nitrate reductases should be clearer under aerobic or alternate aerobic/anoxic conditions. The DO shifts commonly existed in temporal or spatial difference in practice to reveal potential chaperone, and transcriptional activators were in favor of setting unequivocal operated parameters in wastewater treatment project. Furthermore, the novel pathways responsible for the ammonia translated to nitrogen-gas should be illuminated clearly.

Finally, the practical purpose is to use the strain RAD-17 for bioaugmentation to improve denitrification performance in RAS effluent. The whole cell immobilization and integrated with packing carrier should be considered. In addition, the function and ecological fitness of this individual species with other microbes that relate with QS (quorum sensing) regulation should be further clarified, and the relevant control methodologies still need further study.

Figure 6. Proposed model for aerobic nitrogen removal mechanisms of the Pseudomonas balearica strain RAD-17. Red arrow means gene up-regulation; green arrow means inconspicuous gene regulation.
4. Conclusions

An aerobic denitrification Pseudomonas balsiria strain RAD-17 showed efficient nitrogen removal performance with average NO$_3$-N, NO$_2$-N, and TAN removal rates of 6.22 mg·L$^{-1}·$h$^{-1}$, 6.30 mg·L$^{-1}·$h$^{-1}$, and 1.56 mg·L$^{-1}·$h$^{-1}$, respectively. The transcriptional results proved that aerobic nitrogen metabolic pathways were performed in respiratory nitrate reduction genes (napA, nirS, norB, and nosZ) for nitrate removal, or cell assimilation (glnD and glnT) for ammonium utilization. In addition, the bioaugmentation performance by the strain RAD-17 achieved maximum cell abundance of 28.5 ± 4.5% in total environmental cell numbers, as well as obvious higher denitrification performance than in the no inoculum group (84.22% vs. 22.54%).

Supplementary Materials: The following are available online at www.mdpi.com/2076-2607/8/1/72/s1. Figure S1: Scanning electron microscope micrograph of Pseudomonas balsiria RAD-17; Figure S2: The specificity evaluation of PCR amplification assay. The specific DNA bands were detected by agarose gel electrophoresis, lane 1-7 represent the amplified products of 16S rDNA, gldD, gldS, napA, nirS, norB and nosZ, respectively; Figure S3: The agarose gel electrophoresis of the qRT-PCR amplification product of napA gene (A) and 16S rDNA genes (B) in one of the no-inoculum treatments. Lane 1-8 represent the time point of eight samples from 0 hour to 72 hours (including 0, 6, 12, 24, 30, 48, 72 hours); M, marker; CK, positive control.

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