

Isolation and selection of nitrifying bacteria with high biofilm formation for treatment of ammonium polluted aquaculture water

Phân lập và tuyển chọn vi khuẩn nitrate hóa hình thành màng sinh học để xử lý nước nuôi trồng thủy sản bị ô nhiễm ammonium

Research article

Hoang, Phuong Ha^{*1}; Nguyen, Hong Thu¹; Tran, Trung Thanh¹; Tran, Thanh Tung²; Do, Lan Phuong³; Le, Thi Nhi Cong¹

¹Institute of Biotechnology, Vietnam Academy of Science and Technology, ²University of Science and Technology of Hanoi, ³Vietnam Academy of Traditional Medicine

A biofilm is any group of microorganisms in which cells stick to each other and adhere to a surface by excreting a matrix of extracellular polymeric substances (EPS). The chemoautotrophic nitrifying bacteria hardly form biofilms due to their extremely low growth rate; however, biofilm formation of nitrifying bacteria trends to attach in carrier by extracellular polysaccharides that facilitate mutual adhesion, the forming biofilm is also beneficial in nitrogen removal in biological filter systems, especially in aquaculture water treatment systems. The microbial activity within bio-carrier is a key factor in the performance of biofilm reactor. Selection the nitrifier bacteria that biofilm formation and immobilization on the carrier for application in ammonium polluted water treatment technologies, especially in aquaculture is our research objective. Therefore, in this study, ten and six strains of ammonia oxidizing bacteria (AOB) and nitrite oxidizing bacteria (NOB) respectively were isolated from six different aquaculture water samples collected from Quang Ninh and Soc Trang. Basing on their high nitrification activity and biofilm forming capacity, six bacterial strains have been selected to take photo by scanning electron microscope (SEM) and carry out in 2 – liter tanks with and without carriers. As the results, the system with carriers (30% of total volume) increased nitrogen compounds elimination efficiency from 1.2 times to 2 times in comparison with the system without carrier. Two representatives of ammonia oxidizing bacterial group (B1.1; G2-1.2) were classification based on characteristics and they were classified as *Nitrosomonas* sp. and *Nitrosococcus* sp.

Màng sinh học được hình thành từ vi sinh vật nhờ các tế bào tiết ra các chất cao phân tử ngoại bào (EPS) và dính vào nhau đồng thời được gắn lên một bề mặt vật thể lỏng hoặc rắn. Vi khuẩn nitrate hóa tự dưỡng có thể tạo ra màng sinh học nhưng khá khó khăn do tỷ lệ sinh trưởng rất chậm của chúng. Tuy nhiên vi khuẩn nitrate hóa tạo màng sinh học thường có xu thể bám lên giá thể nhờ sự gắn kết của các polisaccarit ngoại bào. Sự hình thành màng sinh học cũng là lợi thể để loại bỏ các hợp chất nitơ trong các hệ thống lọc sinh học, đặc biệt là trong các hệ thống xử lý nước nuôi trồng thủy sản. Hoat tính vi sinh vật cùng với giá thể sinh học là một yếu tố quan trong để thực hiện trong các bể phản ứng màng sinh học. Trong nghiên cứu này, mục tiêu của chúng tôi là lựa chọn được các vi khuẩn nitrate hóa có khả năng tạo màng sinh học và cổ định chúng lên giá thể để ứng dụng trong các công nghệ xử lý nước bị ô nhiễm ammonia đặc biệt là trong nuôi trồng thủy sản. Kết quả cho thầy, từ sáu mẫu nước nuôi trồng thủy sản khác nhau từ Quảng Ninh và Sóc Trăng, 10 chủng vi khuẩn oxy hóa ammonia (AOB) và 6 chủng vi khuẩn oxy hóa nitrite (NOB) đã được phân lập. Dựa vào hoạt tính nitrate hóa và khả năng tạo màng sinh học của các chủng vi khuẩn phân lập được 6 chủng điển hình đã được lựa chọn để chụp ảnh kính hiến vi điện tử quét và được ứng dụng trong hai bể sinh học với dung tích 2 lít có chứa và không chứa chất mang (giá thể). Sau 7 ngày, hệ thống sinh học chứa giá thể (chiếm 30% thể tích) có hiệu suất loại bỏ các hợp chất nitơ tăng hơn từ 1,2 đến 2 lần so với bể sinh học không chứa chất mang. Hai đại diện của nhóm vi khuẩn oxy hóa ammonia (B-1.1 và G2-1.2) đã được phân loại sơ bộ dựa vào một số đặc điểm sinh học và chúng đã được xác định thuộc chi Nitrosomonas và chi Nitrosococcus.

Keywords: aquaculture, biofilm, carrier, chemoautotrophic, nitrifier bacteria

1. Introduction

Brackish water shrimp farming has developed with an increasing intenseve farming, productivity and output are constantly being raised, many processing factories to export the shrimp and shrimp export is become the largest exportable commodity in Vietnam, the annual average of over \$ 2 billion. However, frequent outbreaks of microbial diseases over the last few years and high risk. One of the main causes of disease that is causing environmental pollution increases, caused by excessive food and animal excretion discharged into the environment has formed the organic compounds and waste as fertilizer, including nitrogen pollution accounts for 30-40%. Nitrifying bacteria were involved nitrification process to convert ammonia to nitrate via nitrite, then they are applied in ammonia polluted water treatment technology especially in aquaculture. Nitrification is widely used to remove ammonia from wastewater by biological oxidation. Wastewaters containing high concentrations of ammonia create environmental problems because ammonia may be toxic to aquatic organisms and can cause fertilization of lakes and reservoirs which leads to algal growth and eutrophication (Forgie 1988, Welander 1998). The overall biochemical process of oxidation of NH_4^+ to NO_2^- , then finally to NO_3^- is known as nitrification. Nitrification is performed by the group of bacteria known as nitrifiers. The nitrifying process takes place in two steps and each step is carried out by a specific group of nitrifying organisms. The two microbes involved have been identified in many studies and are the aerobic autotrophic main genera Nitrosomonas and Nitrobacter (Reynolds, 1996). The reactions are as follows:

 $2NH_4^+ + 3O_2 \rightarrow 2NO_2^- + 4H^+ + 2H_2O + Nitrosomonas$ (AOB)

 $2NO_2^- + O_2 \rightarrow 2NO_3^- + Nitrobacter$ (NOB)

AOB performs the first step by oxidizing ammonium to nitrite. NOB completes the oxidation by converting the nitrite to nitrate.

Total ammonia nitrogen is the critical factor in aquaculture systems as it accumulates in the system as a by-product of feed and animal excretion and it has direct effect on the health of aquatic animals. In young shrimp larvae, concentrations of non-ionic ammonia (NH₃) as low as 0.01mg/L can result in mortalities besides causing pathological disturbances as well as depressing growth rates (Ostrensky and Wasielesky Jr., 1995; L-iu and Chen, 2004). Nitrite is also harmful to larvae as it causes reduction of hemolymph oxyhemocyanin (in Penaeus monodon) with concomitant increase in the partial pressure of oxygen (pO2) in hemolymph and reduced oxygen affinity (p50) (Cheng and Chen, 1995). However, it is less toxic than ammonia (Alcaraz et al., 1999) and, only under conditions of long term exposure the toxicity is found manifested (Wheaton et al., 1991) in the reared animals. Likewise, ammonia and nitrite toxicity in Macrobrachium rosenbergii adults and larvae have been investigated by various researchers (Chen and Lee, 1997; Cavalli et al., 2000; Naqvi et al., 2007). Nitrate is relatively harmless to the cultured aquatic organisms (Tomasso, 1994) and it has not been a limiting factor for Macrobrachium larval culture (Mallasen et al., 2004).

In biological ammonia removal systems nitrifying activity of bacteria suspended in seawater has been reported to be extremely low due to their slow growth rate and inhibition of nitrification by free ammonia and nitrite (Furukawa et al., 1993). However, immobilization techniques have been useful to overcome the situation and accordingly (Sung-Koo et al., 2000), fixed film nitrification biofilters are commonly used for ammonia removal in recirculation aquaculture systems (Wheaton et al., 1994; Seo et al., 2001; Shnel et al., 2002). In such installations attached growth as biofilm offer several advantages over suspended culture based systems, such as handling convenience, increased process stability to shock loading and prevention of the bacterial population from being washed off (Fitch et al., 1998; Nogueira et al., 1998). Conventional fixed-film biofilters applied in aquaculture systems include: fluidized bed reactor, biological rotating contactor, trickling filter, submerged filter, and floating packed-bed reactor (Jewell and Cumming, 1990; Nijhof and Bonverdeur, 1990; Yang, Lin-sen and Shieh, 2001, Valenti and Daniels 2000; New 2002). New biofilter types being recently introduced to RAS include: moving bed reactor, three-phase fluidized filter, and hybrid biofilter. The significance of nitrifying biofilters in aquaculture has been reviewed well (Eding et al., 2006; Colt, 2006; Gutierrez-wing and Malone, 2006) and rating standards for these systems (Colt et al., 2006; Malone and Pfeiffer, 2006) are developed recently. Many of the submerged biofilters use crushed oyster shell, dolomite or coral (5 mm particles) as the filter media, but calcareous media contain an inexhaustible source of buffer material which slowly dissolves into the water. Therefore, plastic filter media which have no buffering capacity are often used in biological filters (Seo et al., 2001; Sandu et al., 2002). Biofilter selection influences capital and operating costs of recirculating aquaculture systems, their water quality, and even the consistency of water treatment (Summerfelt, 2006). A perfect biofilter would remove all of the ammonia entering the unit, produce no nitrite, and support dense microbial growth on an inexpensive support material that does not capture solids, require little or no water pressure or maintenance, and require a small footprint. However, no biofilter type commercially available has been found to meet all those objectives, nevertheless each one has its own advantages and limitations. A great deal of research has been conducted on the design and operation of major biofilters in aquaculture systems. In addition to the general reviews on characteristic advantages and simple engineering methods for commonly used biofilters (Wheaten et al., 1994; Timmons et al., 2001), researchers have provided valuable information in terms of system design, operation and performance evaluations on fluidized bed reactors (Sandu et al., 2002; Summerfelt, 2006), floating bead filters (Golz et al., 1999; Malone-and Beecher, 2000), trickling filters (Kamstra et al., 1998), and moving bed filters (Greiner and Timmons, 1998; Yossi Tal et al., 2003) for their applications in aquacultural systems. However, information on process mechanism and kinetics relative to nitrification biofilters applied to aquaculture systems is still insufficient. Simply employing data from traditional wastewater treatment processes to the design of aquaculture biofilters is not appropriate as nitrification conditions in aquaculture systems differs from domestic and industrial wastewater. Compared with domestic wastewater (Metcalf and Eddy Inc., 1991), aquaculture wastewater has a relatively low concentration of pollutants

(Piedrahita, 2003). The nitrification rate is influenced by substrate concentration, dissolved oxygen (DO), mixing regime, pH, temperature, salinity and the level of organics (Chen et al., 2006).

Therefore, our tasks are to detect autotrophic nitrifying bacteria group involved in converting nitrogen compounds in the environment, to study the biofilm formation of nitrifying bacteria and the effect of attached biofilm on the carrier in ammonium polluted water treatment system. This will lead to the preparation of the probiotics to thoroughly treat aquaculture water ammonium contaminated in Vietnam.

2. Material and method

Samples: Six water samples were collected from aquaculture ponds in QuangNinh and SocTrang city, namely Bun, NuocVao, G1, G2, G3 and STII.

Microorganism culture media: Winogradsky (improve) I for AOB; II for NOB with inorganic carbon is NaHCO₃ (2g/L) exchange CaCO₃ (pH = 7.5) (Atlas, 1995). Carrier: foam in size of (1x1x1) cm was utilized as carriers for immobilizing nitrifiers.

Enrichment of nitrifying bacteria: Microorganisms were enriched before isolation by continuous pump of oxygen and supplements of Winogradsky I medium (100mg/l N-NH₄⁺) without agarose in 7 - 10 days before isolation.

Isolation of nitrifying bacteria: After enrichment, water samples were serially diluted to 10^{-1} , 10^{-2} , 10^{-3} , 10^{-4} and 10^{-5} in penicillin vials. Subsequently, water sample from each vial was cultured on petri dishes of Winogradsky I and Winogradsky II medium by spread plating technique. Then, bacteria cultures were incubated at $30 \pm 2^{\circ}$ C in 5 – 7 days. After bacterial colonies could be observed by naked eyes, single colonies were sub – cultured on respective Winogradsky medium by streak plating technique. Subcultures were then incubated at the same conditions for 4 – 6 days before further experiments.

Evaluation of nitrogen removing capability of bacteria isolates

Nitrifying bacteria activity test protocols

Nitrifying bacteria were grown in 20ml of liquid Winogradsky medium in incubator shaker (N – Biotek) at 200 rpm, 30°C. Starting concentration of nitrogenous compounds was 5mg/l N – NH₄⁺ or 5mg/l N – NO₂⁻. After 5 days, culture media were withdrawn for activity test. All specimens were tested under the same condition.

Activity test protocol

Culture media were centrifuged at 10000 rpm for 5 minutes. After centrifugation, cell pellets were removed to obtain supernatant. AOB activity was determined by the amount of $N - NH_4^+$ lost and $N - NO_2^-$ formed by Nessler

and Griss methods. NOB activity was determined by the amount of $N - NO_2^-$ lost and $N - NO_3^-$ formed by Griss and Salicylate methods.

Chemistry analysis methods: Determinate the ammonium concentration by Nessler, nitrite by Griss and nitrate by Brucine (Franson, 1995).

Biofilm forming analysis

Bacterial isolates were cultured in Winogradsky I or Winogradsky II liquid medium in a shaker incubator (200rpm, 30°C) for 3 days, 1ml of culture media was transferred into aseptic Eppendorf tubes and cultured for 3 days (200rpm, 30°C). Subsequently, all culture media were propagated into another set of Eppendorf tubes for bacterial growth determination (measure absorbance at OD₆₀₀). The original tubes were stained with 1ml crystal violet 1% for 20 minutes at room temperature. After staining, sample tubes were washed gently with distilled water, 1ml of ethanol 70% was added to dissolve adhered crystal violet. Absorbance was measured at OD₅₇₀.

SEM photographing: SEM photographing was performed at Vietnam National Institute of Hygiene and Epidemiology, department of Cell Metaphysics.

Large – scale culture system of AOB and NOB together

Bacteria isolates on agarose cultures were transferred to 2 systems of liquid media for evaluation of growth and activity in practice. One system did not contain carriers while the other contain foam (30% of total volume). Culture systems were maintained for 5 days with continuous supply of oxygen before any experiment.

3. Results and Discussion

3.1. Isolation of nitrifying bacteria

After being enriched the samples, that isolated and cultured on petri dishes and selected 10 single colonies on Winogradski I and 6 single colonies on Winogradski II different the phenotype, activity tests were performed for AOB and NOB.

Nitrifying capacity of bacterial isolates

After 5 days in shaker incubator bacterial, nitrifying activities were measured and demonstrated as the Table 1 following.

Table 1 shows that, all 16 isolated bacteria strains have nitrifying activity. In the ammonia-oxidizing group, the B-1.1, B-1.3, B-2.2, G2-1.2 strains has higher ammonia oxidizing activity, another nitrite oxidizing group, the ST-SII2, ST-SII3 and ST-SII4 strains has higher oxidizing nitrite activity, which is potential for further study.

Ammonia Oxidizing Bacteria			Nitrite Oxidizing Bacteria			
Isolate	NH4 ⁺ -N remain- ing (mg/l)	NO ₂ ⁻ -N formed (mg/l)	Isolate	NO ₂ ⁻ -N remain- ing(mg/l)	NO ₃ ⁻ -N formed (mg/l)	
Negative	4.92	Free	Negative	5	Free	
B- 1.1	Free	3.766	ST-SII 2	Free	4.826	
B- 1.3	Free	4.064	ST-SII 3	Free	3.987	
B- 2.1	1.262	3.427	ST-SII 4	Free	4.923	
B- 2.2	Free	4.297	G1-2.1	3.21	1.527	
G2 -1.1	2.657	2.153	G2- 2.2	2.543	2.131	
G2 -1.2	Free	4.510	G3 -2.1	3.016	1.73	
NV -1.2	1.261	3.342				
NV- 1.1	1.115	3.597				
G1- 1.1	2.53	2.641				
G3 -1.2	3.114	1.877				

Table 1. Activities of isolated bacteria

3.2. Biofilm formation of bacterial isolates

Aside from determining activities of isolates, their biofilm formation was also studied. Based on the result of activity test, only six AOB strains and three NOB strains were tested for growth in liquid medium and biofilm forming capacity (Table 2).

Although all achieved bacteria could form biofilm, some isolates were not clearly visible. Thus, further testing and research are required. From Table 2, ST-SII 4 has the best biofilm formation, B-1.1, ST-SII2, G2-1.2, ST-SII3 and B-1.3 had weaker biofilm forming capacity, however, they are still prominent for further study in large – scale culture system. By looking at Figure 1, the results from measuring absorbance correlate with naked eye observation. ST-SII 4 sample had the strongest crystal violet adhesion on Eppendorf tube while B-1.1, ST-SII2, G2-1.2, ST-SII3 and B-1.3 had less. Practical analysis showed that biofilm formation of isolated bacteria was reduced after 6 - 7 days in Eppendorf tubes. Further testing should be conducted to obtain specific data.

Table 2. biofilm forming capacity of bacterial isolates

3.3. SEM photographing of representative bacteria strain

B-1.1, B-1.3, B-2.2, G2-1.2, ST-SII 2 and ST-SII4 bacterial strains were selected to photograph by Scanning Electron Microscope at Vietnam National Institute of Hygiene and Epidemiology (NIHE). Examples are shown in Figure 2-Figure 7.

Six bacteria strains different complete the phenotype (Figure 2-7), five out of six bacteria strains has rod or pear shape, the only exception cell of G2-1.2 strain is spherical shape. All of them appeared to have rough surface while surfaces of ST-SII4 were rough and slot. For rod – shaped bacteria, cell length varied from 1.15μ m to 3.4μ m while width was approximately 0.5 to 0.8μ m, for G2-1.2 strain, cell diameter ranged from 0.7 - 0.8 nm, the characteristics of isolated bacteria strains shown thatAll of isolated bacteria strains has phenotype similarity to the bacteria of *Nitrosomonas* genus (follow Martin D et al., 2006) (Table 3).

Isolate	OD 600	OD 570
Negative control	0.008	0.087
B-1.1	0.276	0.591
B -1.3	0.113	0.389
B -2.1	0.124	0.302
B- 2.2	0.185	0.366
NV- 1.1	0.085	0.279
G2-1.2	0.289	0.411
ST-SII 2	0.236	0.475
ST-SII 3	0.212	0.408
ST-SII 4	0.418	0.710

Table 3. Characteristics of isolated bacteria strains

Characteristics Cell shap	B-1.1 Long slender	B-1.3 Long straight	B2.2 Pear shap	G2-1.2 spherical shap	ST-SII2 Slender pear	ST-SII4 Pear shap
	curved rods	rods			shap	
Cell size (µm)	0.5 x 2.7	0.5-2,5x 1.2-	0.5x1.3	0.7x0.8	0.4x1.15	0.8-1.5x 1-2
		3.4				
Surface of	Less rough	Less rough	rough surface	rough surface	rough surface	rough and slot
cells	surface	surface				surface
Growth	cell dichot-	cell dichot-	cell dichot-	cell dichot-	cell dichot-	cell dichot-
	omy	omy	omy	omy	omy	omy
Habitat	Brackish wa-	Brackish wa-	Brackish wa-	Brackish wa-	Brackish wa-	Brackish wa-
	ter	ter	ter	ter	ter	ter

 Table 4. Characteristics and preferred habitats of the Nitrosococcus, Nitrosomonas genus and G2-1.2, B-1.1 species (Hans-Peter Koops et al., 2006)

 Characteristic

Characteristis	Genus				
	Nitrosococcus	G2-1.2	Nitrosomonas	B-1.1	
Cell shape	Spherical to ellipsoidal	Spherical shap	Straight rods	Slender rods	
Cell size (µm)	$1.5 - 1.8 \times 1.7 - 2.5$	0.7x0.8	$0.7 - 1.5 \times 1.0 - 2.4$	0.5 x 2.7	
Salt requirement	obligate salt require-				
Maximum salt toler-	100-1800	600	150-900	600	
ance (in mM)	T G . C G 11.	N. (. h	Dalasta	Dalanta	
motile cells	i unt of flagella	Not observed	subpolar	subpolar	
Motion	Motile cells possess a tuft of flagella	Not observed	Some cells motile; some cells has not observed	Not observed	
Arrangement of	Central stacks	Central stacks	Peripheral	Peripheral	
intracytoplasmic mem- branes	of vesicles	of vesicles	flattened vesicles	flattened vesi- cles	
Maximum ammonia tolerane NH ₄ Cl (in mN; pH 8.0)	500-1000	600	50-600	600	
Preferred habitats	Sea water, brackish water and industrial wastewater treatment Plant.	Brackish water	Sewage disposal plants; eutrophica- tion; freshwaters; fertilized soils. or Brackish water	Brackish water	



B 1.1 B 1.3 B 2.1 B 2.2 NV 1.1 G2 1.2 STSII2 STSII3 STSII4 Figure 1. Examples of bacteria biofilm formation



Figure. 2. B- 1.1

Figure 3. B-1.3

Figure 4. B-2.2



Figure 5. G2-1.2

Figure 6. ST-SII2

Figure 7. ST-SII4

3.4. Liter culture of bacterial isolates

Six isolates selected (B-1.1, B-1.3, B-2.2, G2-1.2, ST-SII 2 and ST-SII4) were co – cultured in 2-liter tanks with and without carriers for comparison and determination of bacterial growth under natural conditions. Culture system was maintained for 5 days before add the ammonium concentration to be at 5mg/l. The results are shown in Figure 8.



Figure 8. Activity of nitrifying bacteria with and without carriers

In both systems with and without carriers, concentration of ammonium remaining reduced significantly. However, with carriers, bacteria were able to eliminate NH_4^+ totally from liquid medium with higher efficiency. For instance, at 2 hour mark, ammonium elimination by bacteria on carriers was enhanced by 1.2 times, at 4 hour was 1.37 times and at 6 hour was 2.1 times. To observe less nitrite concentration in medium because it was converted in to nitrate concentration by nitrite oxidizing bacteria, so highest nitrate concentration in medium on the 8th day.

3.5. General characteristics of the genera and species of AOB

The taxonomic framework of the AOB stems from early investigations of Winogradsky. The isolates originally were categorized into genera on the basis of the shape of cells and the arrangement of their intracytoplasmic membranes. Using these criteria, the genera *Nitrosomonas*, *Nitrosococcus*, *Nitrosospira*, *Nitrosolobus* and *Nitrosovibrio* have been established (Hans-Peter Koops *et al.*, 2006). In the following, a brief listing of the distinguishing morphological features of the two recognized genera of AOB are *Nitrosomonas* and *Nitrococcus* compare with B-1.1 and G2-1.2 species. Differential characteristics of the genera of AOB and of B-1.1 and G2-1.2 species are presented in Tables 4. The result shown that, characteristics of G2-1.2 strain nearly with *Nitrosococcus* genus and B-1.1 strain nearly with *Nitrobacter* genus in preliminary classification based on characteristics and preferred habitats.

4. Conclusion

Isolated 16 bacterial strains that could grow on Winogradsky medium. In which, 10 strains oxidize ammonium to nitrite while 6 oxidize nitrite to nitrate. Six strains from both groups (B1.1, B1.3, B2.2, G2-1.2, ST-SII 2, ST-SII4) that show superior activity and biofilm forming capacity were selected for further studies.

Selected bacteria were co - cultured in 2 - liter tanks with and without carriers. The system with carriers (30% of total volume) increased nitrogen elimination efficiency range from 1.2 to 2 times after 2 hours, 4 hours and 6 hours.

Two representatives of ammonia oxidizing bacterial group (B1.1; G2-1.2) were classification based on characteristics and they were classified as *Nitrosomonas* sp. and *Nitrosococcus* sp.

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