Study on nitrogen removal performance and microbial community structure succession under the condition of anaerobic sludge and low carbon source

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Abstract. Under anaerobic low carbon source, the reactor was started-up by inoculating anaerobic sludge. The nitrogen removal efficiency and nitrogen conversion pathway during the start-up process were analyzed. At the same time, the structure succession of bacteria in the process of start-up was analyzed by PCR-DGGE technology from molecular biology level. The results showed that inoculated anaerobic sludge had a good effect on nitrogen removal under the condition of anaerobic low carbon source.

1 Introduction

In recent years, a large amount of nitrogen containing substances produced in the process of production and living have entered the water, causing serious pollution to the water environment due to the development of industry and agriculture and the improvement of the people's living standard[1,2]. The traditional biological nitrogen removal process is to remove nitrogen from the water by ammonification, nitrification and denitrification[3,4]. However, nitrification requires adequate dissolved oxygen, and denitrification requires carbon source, which increased the cost of the wastewater denitrification process[5,6,7]. Therefore, it has attracted wide attention from researchers that how to remove nitrogen economically and efficiently. In this paper, anaerobic sludge was inoculated in a ASBR reactor, and its nitrogen removal characteristics and microbial community structure succession were studied under anaerobic and low carbon source conditions.

2 Experimental materials and methods

2.1 Experimental device method

The experiment was carried out in a ASBR reactor. The reactor was made of plexiglass. It is 700mm high, and the inner diameter is 80mm, which has the air outlet, the air intake, the water outlet, the water intake. The water quality of the reactor is shown in Table 2.1. In order to remove dissolved oxygen, the inlet is blown to 15min by nitrogen. The operation period of the reactor is 48 hours, the settling time is 1h, and the 3L of each cycle needs to be changed. The reactor was stirred by nitrogen circulation and stirred 15min per hour. The reactor avoids light and maintains a constant temperature at 30 degrees by using the temperature controller.

2.2 Experimental materials

2.2.1 Inoculating sludge

In this experiment, anaerobic sludge from a water treatment plant in Jin'nan was used as inoculate sludge. The sludge concentration in the reactor was controlled at 3000±50 mg/L.

2.2.2 Experimental water

Simulated wastewater is used for experimental water, which is composed of NH₄Cl, NaNO₂, KH₂PO₄, MgSO₄, 7H₂O, NaHCO₃ and CaCl₂ 2H₂O. Trace elements are supplied from garden soil leach solution. Water quality is shown in Table 2.1

| Index  | Range     |
|--------|-----------|
| NH⁺⁻⁻⁻⁻N | 20-40 mg/L |
| NO₂⁻⁻⁻⁻N | 10-30 mg/L |
| TN      | 60-130 mg/L|
| PH      | 7.5-7.9 mg/L|

2.3 Experimental method

2.3.1 Conventional index analysis method

During the experiment, the influent and effluent samples were collected 1 times. The concentrations of ammonia nitrogen, nitrate nitrogen, nitrite nitrogen and total

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nitrogen were analyzed. At the same time, morphological changes of microorganisms at different stages were observed by scanning electron microscope (SEM, Hitachi S-2500C). All of the above test methods were carried out according to the national standard method[8].

2.3.2 Molecular biological analysis method

In order to clarify the principle of Microbiology in the process of start-up systematically, the microbial population structure and succession law of the sludge were carried out by PCR-DGGE technology for 2 months and 4 months. The sampling sludge was centrifuged for 10 minutes under the condition of 4000rpm, and then the microorganism DNA in the sludge was extracted with the OMEGA DNA Extraction Kit (E.Z.N.A.TM Soil DNA Kit). The DNA was amplified by nested PCR reaction. The first template was diluted 5 times, and the second template was diluted 50 times. The reaction conditions of PCR were as follows: pre-denaturing 5min at 94℃, denaturing 30s at 94℃, annealing 45s at 57℃, refolding 90s at 72℃, 32 cycles, and finally extending 7min at 72℃.

PCR products were analyzed in DGGE by Bio-Rad DcodeTM Universal Mutation Detection System. DGGE adhesive with 16mL concentration 35%-55% system. DGGE electrophoresis first runs 5min under 120V conditions, and then runs 12h under 80V conditions. After the electrophoresis, Using SDNA-Nucleic Acids Stain Dye (BIO BASIC INC., Canada) to dye 30 min, the strip was cut, and the frozen thawing was repeated for 3 times, and then the DNA was dissolved. The above products were amplified by primers R518 and F357 for PCR, and the purified samples were commissioned in Beijing Huada gene company to be sequenced. The sequencing results were compared on NCBI to determine the microbial population, and Quantity one software was used to analyze the similarity and diversity of the results, and the phylogenetic tree was constructed by Mega5.0 software.

3 Analysis and discussion of the results

3.1 Study on Denitrification performances of anaerobic sludge with low carbon source

After the ASBR reactor was inoculated with anaerobic sludge, the operation of controlled anaerobic low carbon source has been running for 120 days, and the results are as follows.
provide carbon source for denitrifying bacteria, and do not contain dissolved oxygen in the simulated wastewater, and the ammonification effect is inhibited. The ammonia nitrogen concentration in the effluent is high, and therefore the ammonia nitrogen concentration in the effluent of the reactor is high, and the nitrate nitrogen and nitrite nitrogen concentration in the effluent are low and the total nitrogen removal rate is high. As the reaction goes on, the organic matter in the reactor was degraded gradually. The denitrifying bacteria decreased gradually because of the lack of carbon source. The removal of nitrite and nitrate decreased gradually and the removal rate of total nitrogen was reduced. And at this time, the removal rate of ammonia nitrogen has been increasing, because the new bacterial community that has the effect of removing ammonia nitrogen is enriched in the reactor. In the late reaction, the denitrifying bacteria were gradually replaced by some anaerobes, and the removal rate of ammonia nitrogen, nitrite nitrogen and total nitrogen increased again, reaching stable, while the concentration of nitrate nitrogen in the reactor increased. We can speculate that the dominant population in the late stage of the reactor is an anaerobes that can use ammonia nitrogen and nitrite nitrogen to produce nitrate nitrogen with no additional carbon source.

3.2 Microorganism population structure and Succession Law

3.2.1 Analysis of DGGE Atlas

The DGGE Atlas of different stages of the reactor is shown in Fig 3.5.

![DGGE Atlas](image)

Fig 3.5 DGGE atlas at different stages of the reactor

As can be seen from Fig 3.5, there are obvious differences in the biological population of the reactor at different stages, and the strip of the initial inoculation sludge is obviously more than that of the samples that run for two months and four months. In the DGGE map, the initial dominant population showed more performance. With the reaction goes by, the bands of 3, 4, 14, 15, 17, 18, 20, 21, 22, 23, 25, 28 were not detected in the follow-up samples. This may be due to the lack of organic matter in the water, resulting in the partial elimination of some heterotrophic microbial populations. At the same time, bands 12 and 16 appeared in all stages of the reactor, indicating that the biological population could survive under anaerobic low carbon environment. Bands 2, 26 and 13 are new bands, which are newly domesticated populations in anaerobic environment. Bands 5, 11, 19, and 24 disappear gradually throughout the operation of the biological system. With the operation of the reactor, these biological populations are gradually eliminated.

3.2.2 Cluster and diversity analysis in reactor

Fig. 3.6 is a cluster analysis map of microbial population in different stages of the reactor.

![Cluster Analysis](image)

Fig 3.6 Reactor cluster analysis

According to the cluster analysis of microbial population in different stages of the reactor in Fig. 3.6, it is clear that the reactor has obvious biological population succession during operation. The inoculated sludge runs in a completely anaerobic environment. After a long period of operation, some of the sludge's organisms are gradually eliminated, resulting a decline in the number of populations in the system and the similarity index decreased from 0.57 to 0.43 with the increase of running time.

Table 3.1 Diversity analysis of DGGE Atlas of sludge

| Number | 1  | 2  | 3  |
|--------|----|----|----|
| Shannon| 3.26| 2.68| 2.62|

Table 3.1 showed that the diversity index of the reactor decreased significantly after two months of operation, and the Shannon index decreased from 3.26 to 2.68. However, As it continues to run , the diversity index of the biological population changed little. The results showed that some aerobic and heterotrophic bacteria were gradually eliminated during the two months' operation, and the diversity of the biological population in the reactor remained stable.

3.2.3 Phylogenetic tree analysis of biological population

Phylogenetic tree were constructed by Mega5.0.

According to the comparison of the results of sequencing on NCBI, we can see that the microbial population in the reactor system is mainly composed of Proteobacteria, and there are a few other microbes. In combination with the DDGE map, some organic and aerobic bacteria are eliminated, such as Fibrobacteres (strip 3), Propionivibrio (strip 4), and Lysobacter (strip 20) gradually eliminated. At the same time, Nitrosomonas (strip 12) and Accumulibacter (strip 16) can be detected at various stages of the reactor.
According to the comparative literature, it was found that Nitrosomonas belongs to ammoniation bacteria, and the study showed that Nitrosomonas can show anammox activity under anaerobic conditions\(^{(9)}\). Accumulibacter belongs to the anaerobic oxidizing energy bacteria, which exists in the whole stage of biological system operation, indicating that this kind of bacteria has strong impact resistance\(^{(10)}\). Bradyrhizobium (band 13) and Panthomonas (strip 26) are newly acclimated microorganism, in which Bradyrhizobium can use ammonia salt and nitrate as nitrogen source and is another group of nitrogen removal bacteria within the system. Ralstonia (band 5), Rhodospirillum (strip 11), Chitinophagaceae (strip 18) and Conchiformibius (strip 21) keep the advantage for a long time, but they are gradually eliminated with the increase of running time.

![Fig 3.7 Phylogenetic tree of reactor](image)

The main biological systems are β- deformable bacteria (Betaproteobacteria) and α- deformia (Alphaproteobacteria), which belong to the deformable bacteria gate (Proteobacteria). With the operation of the reactor, the anaerobic population is the main group, and aerobic and heterotrophic bacteria are gradually eliminated.

4 Conclusions

1. The reactor reached stability after running 83d under anaerobic low carbon source condition. During the stable operation, the removal rate of ammonia nitrogen and nitrite nitrogen reached 95%, and the total nitrogen removal rate was more than 60%.
2. The denitrifying bacteria in the reactor were mainly Bradyrhizobium and Nitrosomonas This work was supported by the Higher Education Science and Technology Plan of Shandong Province (J17KA198).

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