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Zhao, Nannan; Li, Xiaohu; Jin, Xiangdan; Angelidaki, Irini; Zhang, Yifeng

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Integrated electrochemical-biological process as an alternative mean for ammonia monitoring during anaerobic digestion of organic wastes

Nannan Zhao, Xiaohu Li, Xiangdan Jin, Irini Angelidaki, Yifeng Zhang*

Department of Environmental Engineering, Technical University of Denmark, DK-2800 Lyngby, Denmark

*Corresponding author, Tel.: +45 45251410; fax: +45 45932850.

E-mail address: yifz@env.dtu.dk; yifzmfc@gmail.com (Yifeng Zhang)
Abstract

Ammonia monitoring is important to control anaerobic digestion (AD) process due to inhibition effect. Here, an electrolysis cell (EC) was integrated with a complete nitrification reactor as an alternative approach for online monitoring of ammonia during AD processes. The AD effluent was pumped into nitrification reactor to convert ammonia to nitrate, followed by the introduction of nitrate-rich effluent to EC cathode. It was first evaluated with synthetic ammonia-rich digesters and was observed that the current at 5 min were linearly corresponding to the ammonia levels (from 0 to 7.5 mM NH$_4^+$-N, $R^2 = 0.9673$). The linear relationship was always observed regardless of different wastewater pH and external voltage. Pre-removal of other electron acceptors from digestate at cathode could eliminate their disturbances to sensor performance. Finally, the accuracy of biosensor was verified with real digestate test. The simple and reliable biosensor showed great promising for online ammonia monitoring of AD processes.

Keywords

Biomonitoring; Ammonia; Electrolysis cell; Nitrification; Anaerobic digestion; Digestate
1. Introduction

Anaerobic digestion (AD), as a sustainable technology to produce biogas from various wastes such as manure, solid waste and sludge, is becoming more and more attractive across the world (Chatterjee and Mazumder, 2016; Choong et al., 2016; Rayner et al., 2017; Senghor et al., 2017). However, ammonia coming from degradation of protein in AD substrate could be an inhibitor for AD process (Angelidaki and Ahring, 1994; Hansen et al., 1998; Rajagopal et al., 2013). As reported before, low free ammonia concentrations (e.g. 100 mg-N/L) could inhibit an unadapted AD process (Hansen et al., 1998). Therefore, ammonia monitoring is significantly important to maintain stability of AD. Currently, the common methods to test the ammonia in aquaculture include colorimeter procedure (Nessler, phenate, and salicylate methods), ion-selective electrode for NH$_4^+$, and test kits relying on the Nessler or salicylate methods (Krug et al., 1979; Le and Boyd, 2012; Zhou and Boyd, 2016). However, there are still some drawbacks among these methods. Firstly, the accuracy of colorimeter methods is easily disturbed by the sample color or turbidity. Secondly, the issue of disposal of hazardous waste, which increases the complexity and operation cost, becomes a sever concern in Nessler method. Thirdly, most of the colorimeter methods need extra sample preparations (e.g., membrane filtration) which are time-consuming and costly. Lastly, the ion-selective electrode is strongly affected by the sodium and potassium although with high precision and wide detection range. New conductive materials have also been developed to monitor dissolved ammonia in water, but the synthesis of materials and sensor fabrication are still complicated and the costs are relatively high (Tao et al., 2006; Huang et al., 2016). Thus, development of an alternative environmental-friendly, simple, rapid, online and reliable ammonia sensor is in urgent need for environmental monitoring and water quality control.

On the other hand, electrochemical system has been widely applied into ammonia sensor field because of their high selectivity and sensitivity (Ribeiro et al., 2012; Herzog, 2015; Zhybak et al., 2016; Ning et al., 2017). The electrochemical ammonia sensors, which are based on the ammonia ion transfer reactions across different electrolyte interfaces, mainly include potentiometric and amperometric methods (Bertocchi et al., 1996; Abass et al., 1998; Zolotov et al., 2014; Herzog, 2015). In Luo’s research (Luo and Do, 2006), two kinds of polyaniline-poly composite films were prepared and they exhibited a high sensitivity of NH$_4^+$-N up
to 100 mM. Ribeiro (Ribeiro et al., 2012) also developed an amperometric sensor to detect ammonia concentrations in the range from 4.2 to 66 uM. Most of the electrochemical ammonias sensors were focused on the development of new nanomaterials or fabricating electrodes. Although with high sensitivity and selectivity, these electrochemical sensors still require tedious and complex fabrication process and high consumption of nanomaterials.

Recently, nitrate as an alternative electron acceptor was removed at cathode of an EC reactor, which demonstrated the cathodic nitrate reduction in EC (Abdallah et al., 2014; Radjenovic and Sedlak, 2015; Rajmohan et al., 2016). In biological pathway, ammonia could be aerobically oxidized to nitrate through nitrification process (Zhang et al., 2009; Samarasinghe et al., 2016; Wang et al., 2016). Therefore, coupling nitrification stage with electrochemical nitrate reduction could be a possible way to monitor ammonium during biogas production. Compared to the conventional methods and current electrochemical ammonia sensors, it may offer a fast, environmentally friendly and simple potential approach for monitoring ammonia during AD. To the best of our knowledge, such integrated system has never been explored before.

The aim of present study is to develop an alternative and potentially sustainable approach for ammonia monitoring during AD process. To achieve this goal, an integrated EC-nitrification system was constructed. In this sensor, ammonia in the AD digester was firstly oxidized to nitrate in nitrification reactor. The effluent enriched with nitrate was introduced into cathode of EC reactor for further reduction. Firstly, the sensor performance was explored with synthetic wastewater, in terms of nitrification efficiency, ammonia detection range, response time and operational stability. The effects of external power supply and pH on sensor performance were also investigated. Besides, considering other possible electron acceptors in AD digester, the digester was pumped into cathode of EC first to eliminate effects. Finally, the new sensor was tested with real AD digesters to verify its reliability. The application of this new sensor may have the potential to provide an online and reliable approach to monitor ammonia, which could be helpful to maintain AD stability.
2. Material and methods

2.1 Reactor configuration and operation

An EC reactor, made of nonconductive polycarbonate plates, was a rectangular reactor composed of anode chamber (50 ml) and cathode chamber (100 ml) (Fig.1). The electrode materials were a titanium woven wire mesh (4×5 cm, 0.15 mm aperture, William Gregor Limited, London) coated with 0.5 mg/cm² Pt for cathode, and a Titanium mesh electrode coated with Ir MMO (dimensions: 4×5 cm; 1 mm thickness; specific surface area 1.0 m²/m², Magneto Special Anodes, The Netherlands) for anode. The anode chamber was filled with 50mM PBS (pH-7) and cathode chamber was filled with synthetic wastewater. The synthetic wastewater was prepared with ammonia chloride being dissolved into tap water at different ammonia levels. Two chambers were separated by a cation exchange membrane (CEM, CMI 7000, Membrane international, NJ). The rubber gaskets and screws were used to avoid leakage during the assembling. The membrane was put in 50 M NaCl solution for 24 hours, and then soaked in distilled water until use. A power supply (NEWARE Battery testing system 7.5.X, China) was used to provide the external voltage to the reactor. During the study of influence of other electron acceptors on sensor performance, an anion membrane (AEM) was used to avoid NH₄⁺ migration.

A lab-scale nitrification reactor was set up to complete the oxidation from ammonia to nitrate. A glass bottle (500 ml) was used for the reactor, and the effluent of a membrane bio-reactor (MBR) was used as the inoculum to set up this nitrification stage. The characteristics of inoculum were 2.5 g/L (total solid content) and pH 7.04. During the enrichment of nitrification bacteria, the medium, which contained 2.45 g/L NaH₂PO₄, 4.58 g/L Na₂HPO₄, 0.1 g/L KCl, 0.1 g/L MgCl₂•6H₂O, 0.1 g/L CaCl₂•2H₂O, 12.5 ml/L mineral solution and vitamin solution as describe before, was used. NaHCO₃ (2 g/L) was used as the carbon source and electron donor. The aeration was obtained directly from the air as it was open to air and the solution was fully mixed by magnetic stirrer.

In the investigation of linear relationship between current and ammonia concentration, the synthetic wastewater with different ammonia levels (0-7.5 mM NH₄⁺-N) was pumped into nitrification reactor first and then the effluent was purged with nitrogen and introduced into cathode chamber of EC. The current in circuit
was recorded by NEWARE Battery testing system 7.5.X. In one set of tests, the external voltage was set at 1.5 V, 1.8 V, 2 V and 2.3 V to elucidate the effect of external power on sensor performance. Then the effect of pH on sensor performance was studied at four different pH levels (pH 4, 6, 7 and 8). Subsequently, the pretreatment of sensor was evaluated by adding Fe$^{3+}$ and NO$_3^-$ to the synthetic wastewater. To verify the accuracy of sensor, it was tested with two real AD effluents. One was collected from a lab-scale AD reactor for biogas upgrading which was operated at 37 ºC (noted as AD effluent 1). The other one was taken from a thermophilic lab-scale AD reactor fed with manure (55 ºC) (noted as AD effluent 2). All experiments were carried out in duplicate at room temperature.

In the statistical analysis, the student’s t-test was applied and P-values<0.05 were considered to have significance effect on the response, while values > 0.1 indicate the variables are not significant.

Figure 1 Schematic diagram of the integrated EC-nitrification system. CEM: the cation exchange membrane.
2.2 Electrochemical analysis and calculations

The pH and conductivity of tested solution were measured by a PHM 210 pH meter (Radiometer) and a CDM 83 conductivity meter (Radiometer), respectively. The concentrations of ammonia, nitrate and Fe$^{3+}$ were determined by colorimetric test kits (Spectroquant 00683, 09713, Merck, Germany). The current in EC circuit was recorded by NEWARE Battery testing system 7.5.X every 1 min.

3. Results and discussion

3.1 Nitrification stage

Figure. 2 The correlation between initial ammonia level and accumulated nitrate concentration in nitrification reactor.

Fig.2 depicts the performance of lab-scale nitrification reactor. All the nitrifying process was completed within four hours. As shown, when ammonia level changed from 0 to 7.1 mM NH$_4^+$-N, the corresponding nitrate concentrations varied from 0 to 6.9 mM NO$_3^-$-N, with a stable conversion rate. The line slope is 0.9534, illustrating almost the same amount of ammonia to nitrate numerically. The high conversion efficiency (over 90%) was in agreement with literature (Tarre and Green, 2004). However, compared to other study (Zhang et al., 2009), the nitrification rate obtained here was a little lower (581.4 g/m$^3$·d) because of the possible different operational parameters such as the different attachment forms of nitrifying bacteria,
pH and dissolve oxygen concentrations. As demonstrated before, influent pH has a significant impact on the
performance of a membrane-aerated bioreactor with respect to nitrification rate (Shanahan and Semmens,
2015). Also, biofilm performs better than suspended-biomass in terms of nitrification rate (Tarre and Green,
2004). Nevertheless, the results above had already offered enough proof for the sensor concept. Considering
aerated nitrification process has already been developed commercially with a high rate, it could be replaced
with a commercialized nitrification reactor in the future industrial application. In a short conclusion, the high
and stable nitrification efficiency provided the insurance of the following investigation of relationship
between current and ammonia concentration.

3.2 The performance of EC: correlation between ammonia level and current

The effluent containing nitrate from nitrification process was firstly flushed with nitrogen and then
introduced into cathode chamber of EC. A fast response of current of EC to a stepwise increase from 0 to 7.1
mM NO₃⁻-N was observed within 5 minutes. Fig.3 (A) and (B) illustrate the current variation of EC versus
nitrate and ammonia concentration, respectively. It is clear from Fig.3 (A) and (B) that the current showed
positive slopes with nitrate and ammonia level increasing. The current linearly increased (5.329 ± 1.157 to
10.139 ± 1.233 mA, R² = 0.9673) when ammonia levels ranged from 0 to 7.5 mM NH₄⁺-N. The slopes for

![Figure. 3 The linear relationship between current and nitrate levels (A) and ammonia levels (B) at external
data-pow supply of 1.8 V and pH 6.](1.4 to 7mM NH₄⁺-N)
current and nitrate, ammonia levels were 0.6686 and 0.6389, respectively. The positive slopes meant with ammonia or nitrate concentrations increasing, the corresponding currents showed a gradual linear increase as well. The current, as a representative of the numbers of electron per second in the circuit, was closely connected with external voltage and amount of electron acceptors in cathode. Considering the stable external power of 1.8 V in this case, the different current response was mainly contributed by the different electron acceptor (i.e., nitrate) in cathode. In addition, the range of detection could reach up to 7.5 mM NH$_4^+$-N, which was much higher compared to other electrochemical ammonia sensor (0.3 mM) (Ribeiro et al., 2012). The above results demonstrated a good linear relationship between current generation and initial ammonia concentrations ranging from 0 to 7.5 mM. Contrast to the conventional testing approaches (e.g. testing kits and titration), the new biosensor developed here exhibited merits of rapid and real time monitoring properties. Compared to other fast electrochemical ammonia sensor, this new sensor also shows great advantages of relatively low cost and high sensitivity. Overall, it has been demonstrated that the new developed sensor has a good performance in monitoring ammonia and especially is most suitable for high ammonia concentration detection.

3.3 Monitoring performance under different operational conditions

![Graph showing current generation vs NH$_4^+$-N concentration under different operational conditions.](image)
Figure. 4 Effect of external applied voltage on the performance of ammonia sensor. The initial pH of the nitrification effluent is 6 in the cathode.

Since the current in circuit is highly related to external voltage, a series of tests were conducted at varied external applied voltages (1.5 V to 2.3 V). As shown in Fig.4, the currents always showed positive slopes with respect to ammonia levels regardless of the applied voltages. The slopes at 1.5, 1.8, 2 and 2.3 V were 0.7629, 0.6389, 1.2712 and 1.6488, respectively. The highest slope appeared at the external voltage of 2.3 V, while the slopes for 1.5 V and 1.8 V were relatively lower. The higher slope usually meant a higher current generation. For example, at 7.5 mM NH$_4^+$-N, the currents at 1.5, 1.8, 2 and 2.3 V were 8.6, 10.1, 11.2 and 15.6 mA, respectively. The higher slopes at 2 and 2.3 V were mainly contributed by the higher external voltage. The lowest currents appeared in 1.5 V could be explained by that the amount of electrons became the limiting factor of nitrate reduction when voltage was too low as 1.5 V. Additionally, the sensor exhibited high linearity more than 90% at all voltages, illustrating the good performance under different voltages.

![Figure 4](image.png)

Figure. 5 Effect of pH on the performance of ammonia monitoring at external applied voltage of 1.8 V.

Different pH of AD digesters are usually observed in various kinds of studies (Cook et al., 2017). Thus, it’s vital to investigate the influence of pH on sensor performance. In order to achieve this goal, different pH of synthetic wastewater were tested. The results are displayed in Fig.5. It was observed that at all pH conditions, the currents always increased linearly with ammonia levels. The slopes for pH 4, pH 6, pH 7 and pH 8 were
0.6746, 0.6389, 0.9395 and 0.862, respectively. The higher slopes tended to be achieved in neutral and weak alkaline condition. However, it should be noticed that higher currents were observed in weak acid environment (pH 6). For example, when with the initial ammonia concentration was 4.5 mM NH₄⁺-N, the currents at pH 6 and pH-8 were 8.8 and 6.3 mA, respectively. As shown in Fig.5, the current generation at weak acid condition (pH 6) actually exhibited a higher value compared to the neutral and weak alkaline condition. It has been reported that the weak acid condition may favour electrochemical nitrate reduction (Abdallah et al., 2014). Moreover, the correlation coefficients for pH 4, pH 6, pH 7 and pH 8 were 0.9825, 0.9652, 0.9291 and 0.9526, respectively. The sensor showed high linearity at all tested pH conditions.

3.4 Monitoring performance under the interference from other electron acceptors

![Figure. 6 The change of Fe³⁺ concentration (A) and current generation (B) over time in EC reactor.](image)

The operational parameters: external voltage of pretreatment is 4 V, initial pH is 6 and initial NH₄⁺-N of 6 mM in synthetic water.

In order to avoid the interferences from other possible electron acceptors, the wastewater was firstly introduced to the cathode of EC reactor before nitrification. The synthetic wastewater with 21.4 uM Fe³⁺ was used for the test. Fig.6 (A) shows the variations of Fe³⁺ concentration over time in the EC during pretreatment. It is clear from this figure that Fe³⁺ was removed totally within 90 minutes when external voltage was applied to 4 V. The removal mechanism was mainly electrochemical reduction in cathode. Fig.6 (B) displays the change of biosensor current over time with and without pre-treatment in the EC reactor. For
both conditions, the fast decrease of current within 20 minutes was observed. The stable current of biosensor with pre-treatment was 0.93 mA, while it was 0.67 mA for the biosensor without pre-treatment. The student’s test was conducted to evaluate the significant difference between two series of data. At 99% confidence level, there was a significant difference between the current generations in two cases. The results demonstrated that EC pretreatment before nitrification could be an efficiency method to eliminate the disturbance from other electron acceptors in the AD effluent.

![Figure 7](image)

Figure. 7 The change of NO$_3^-$-N concentration (A) and current generation (B) over time in EC reactor. The operational parameters: external voltage of pretreatment is 4 V, initial pH is 6 and initial NH$_4^+$-N of 80 mg/L in synthetic water.

Likewise, the nitrate, as another important electron acceptor in water, is also not negligible. To explore if the EC pretreatment also works for the removal of NO$_3^-$-N before nitrification, the synthetic wastewater enriched with 714.3 uM NO$_3^-$-N/L was used for the test. As displayed in Fig.7 (A), the nitrate showed a gradual decrease within 2 hours owing to cathodic electrochemical reduction when 4 V was applied. Fig.7 (B) shows the current change in the biosensor with and without EC pretreatment. Similarly, the results of student’s $t$-test showed a significant difference between two situations at 99% confidence level, showing the importance of EC pretreatment for the accurate sensing. To conclude, the effects of Fe$^{3+}$ and NO$_3^-$-N on sensor performance could be eliminated through the pre-treatment using EC, which could improve the stability
during ammonia monitoring with complex waste streams. But for AD process, electron acceptors such as Fe$^{3+}$ and NO$_3^-$ may not be existence, since they would be reduced by receiving the electrons from microbial oxidation of organics. Thus, pretreatment may not be necessary for ammonia monitoring of AD digestate. But the results offer insight into the extended application feasibility of the sensor for other water treatment processes.

3.5 Application in real AD effluent

Table. 1 Determination of NH$_4^+$-N in real waste streams by EC-nitrification based biosensor and ammonia testing kit.

<table>
<thead>
<tr>
<th>sample</th>
<th>NH$_4^+$-N$^a$</th>
<th>NH$_4^+$-N$^b$</th>
<th>pH</th>
<th>Conductivity (us/cm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>AD effluent 1</td>
<td>1542.30 ± 5.12</td>
<td>1647.8 ± 13.18</td>
<td>8.87 ± 0.10</td>
<td>1690 ± 2</td>
</tr>
<tr>
<td>AD effluent 2</td>
<td>917.24 ± 2.16</td>
<td>987.5 ± 20.04</td>
<td>8.20 ± 0.10</td>
<td>1329 ± 5</td>
</tr>
</tbody>
</table>

$^a$: tested by fast kits. $^b$: measured by sensor.

To verify the applicability for real digestate, the biosensor was further tested with two real AD effluents. The samples were stored at 4 ºC and fluxed with nitrogen before used. The ammonia levels obtained from fast kits and biosensor along with primary characteristics of two effluents were summarized in Table 1. Anova analysis was performed to investigate the difference between values tested with two methods. The results showed no significant difference between the results obtained from the two methods ($P > 0.05$) at 95% confidence level, which confirmed a good accuracy of the biosensor. According to the results, the sensor could monitor the AD samples with a wide range of ammonia levels.

4. Conclusions

The present study for the first time demonstrated the feasibility of integrated EC-nitrification process for ammonia monitoring during AD process. It was observed that the sensor responded immediately within 5 minutes to a stepwise increasing of ammonia level from 0 to 7.5 mM NH$_4^+$-N. Linear relationships between current and ammonia concentration were always observed regardless of voltage and pH. Additionally, EC could pre-remove potential electron acceptors before monitoring. Finally, the sensor showed high accuracy.
and sensitivity to real AD effluents. Though promising, more efforts are still needed to further improve the sensor by simplifying reactor configuration, shortening aerobic process etc.

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References


Graphical Abstract

Electrochemical system

Nitrification Reactor