Importance of culture history on 17α-ethinylestradiol cometabolism by nitrifying sludge

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ABSTRACT

17α-ethinylestradiol (EE2), a synthetic estrogen which interfere the endocrine and reproductive function in living organisms, has been found extensively to be deposited into municipal wastewater treatment plants and the environment via human excretion. EE2 has long been known to be efficiently cometabolized by ammonia-oxidizing bacteria (AOB) during ammonia (NH3) oxidation. Current study aims to investigate the effect of culture history on the biotransformation of EE2 by nitrifying sludge which was enriched under different ammonia loading rates in continuous flow reactors. Result showed that past growth condition largely affected not only the metabolic rate of NH3 oxidation but also EE2 cometabolism. Sludge previously acclimated with higher NH3 loads as well as sludge dominated with AOB belong to high growth cluster (Nitrosomonas europaea-Nitrosococcus mobilis) showed higher rate of EE2 biotransformation than those one being acclimated with lower NH3 loads because of its ability to provide more reducing power from NH3 oxidation. Moreover, the correlation between the degradation rates of NH3 and EE2 was higher in sludge being acclimated with higher load of NH3 in comparison with other sludge. Implication of the findings emphasized the role of volumetric NH3 loading rate in determining EE2 removal in wastewater treatment system.

Keywords: Ammonia-oxidizing bacteria, Cometabolism, Culture history, Nitrifying sludge, 17α-ethinylestradiol

1. Introduction

Estrogens are essential group of hormones that regulate the growth and development of female sexual characteristics in humans and animals. Estrogens strongly interfere with endocrine and reproductive functions in living organisms including disturbance on immunological system and fertility, reproductive failure due to thinning of eggshells and reduction in egg production [1], feminization and masculinization, and altered sexual development [2]. 17α-ethinylestradiol (EE2), a synthetic estrogen, is generally used as the oral contraceptive pill to control the birth rate and hormone therapy. EE2 prompt to be released to the environment via human excretion which is later discharged into municipal wastewater treatment plants (WWTS) as hotspots for the release of estrogens in the environment [3].

Some estrogens are able to be degraded in the WWTS. The estrogens removal efficiencies in activated sludge process range from 61% to 98% for estrone (E1), 67% and 99% for estradiol (E2) and 85% to 90% for EE2 [4]. Natural estrogens (E1 and E2) are readily metabolized by heterotrophic microorganisms in WWTS, while the biodegradation of more recalcitrant hormone like EE2 is known mainly through cometabolism of ammonia monooxygenase enzyme (AMO) by ammonia-oxidizing bacteria (AOB) [5, 6]. Some essential factors like the availability of ammonia (NH3) as the growth substrate of AOB [5] have been investigated for its roles on EE2 cometabolism by AMO. However, microorganism generally produce enzyme and other macromolecules at a rate which is unique to their previous growth condition [7, 8], therefore, the culture history of AOB may be another factor determining not only the rate of NH3 utilization [9] but also the rate of EE2 cometabolism.
From Carl Erba Reagents (Val de Reuil, France). All chemicals were purchased from Sigma-Aldrich Corporation (St. Louis, MO, USA). The stock solution of EE2 was prepared to 500 mg/L in methanol. Other chemicals were purchased from Carlo Erba Reagents (Val de Reuil, France). All chemicals used in this study are analytical reagent grade.

2. Materials and Methods

2.1. Chemicals

EE2 (17α-ethinylestradiol) was purchased from Sigma-Aldrich Corporation (St. Louis, MO, USA). The stock solution of EE2 was prepared to 500 mg/L in methanol. Other chemicals were purchased from Carlo Erba Reagents (Val de Reuil, France). All chemicals used in this study are analytical reagent grade.

2.2. Enrichment Media for AOB

Enrichment media, which consist of synthetic wastewater plus NH3 at various concentrations, was used for enrichment of AOB. Synthetic wastewater A was prepared by dissolving NaCl (1 g), MgCl2•6H2O (0.4 g), CaCl2•2H2O (0.1 g), KH2PO4 (0.2 g), and KCl (0.5 g) into one liter of de-ionized (DI) water and then adding one milliliter (mL) of each mineral solution including the solution of non-chelated trace element, selenite-tungstate, vitamin mixture, thiamine, vitamin B12 [10]. Synthetic wastewater B was added with 0.68 and 2.02 g of (NH4)2SO4 per liter to make NH3 at various concentrations, was used for enrichment of AOB.

2.3. Enriched Reactors Setup and Operation

Three vessels with an effective volume of 6 liters were identically set up as continuous flow enriched reactors without cell recycle. For each reactor, air was supplied from air blower connecting with a small stone diffuser to provide oxygen and mix the mixed liquor completely. The concentration of dissolved oxygen (DO) in bulk solution was maintained between 7-8 mg O2/L by adjusting air flow rate with a small needle valve. pH in mixed liquor was controlled at 7.5 ± 0.2 by a pH controller (Liquitron DP 5000, LMI Milton Roy, USA) equipped with a pH probe (Orion 9156DWP, Thermo scientific, UK) as well as a solution of 0.5 N HCl and 0.5 N NaOH. Reactors were operated at room temperature between 23-25°C. Mixed liquor taken from full scale activated sludge process was used as inoculum to start up the reactors. Enriched reactors were operated differently in order to enrich AOB which belongs to different growth characteristics. Enrichment medium containing NH3 was continuously fed into each one of three reactors at different NH3 concentrations to enrich the AOB population which belongs to different rates of NH3 utilization. The reactors named ‘70NS’, ‘140NS’ and ‘420NS’ were fed with NH3 of 70, 140 and 420 mg N/L, and controlled at sludge age of 4, 3, and 4 d, respectively. For all tests, samples were periodically collected in a similar manner with that of EE2. Synthetic wastewater A was used particularly for tests with having initial NH3 levels of 28 and 70 mg N/L, while for tests belonging to the initial NH3 concentration of 140 and 420 mg N/L, wastewater B was used instead. Air and mixing was provided to the mixed liquor by shaking in an orbital shaker at 250 rpm. The reactor was incubated at room temperature. While running the tests, pH in bulk solution was maintained within a range of 8.0 ± 0.2 by adding HEPES buffer solution to a final concentration of 30 g/L.

2.4. Test procedure

Sludge containing the AOB population which belongs to different NH3 utilization rates was taken from each one of three enriched reactors during steady operation and then, brought into batch degradation tests. Various initial concentrations of NH3 were used in the tests in order to determine effect of growth substrate concentration on the rates of NH3 oxidation and EE2 degradation. Loss due to abiotic sorption and volatilization of NH3 and EE2 were observed through batch tests with heat-killed cell and without cell. EE2 biodegradability by heterotrophic microorganism in enriched sludge was investigated in the presence of 10 mg/L allylthiourea (ATU) for the suppression of the AMO usage by AOB [12]. Furthermore, 1,000 mg/L chloramphenicol was also added into the reactors in order to evaluate the NH3 and EE2 biodegradability remained in the harvested sludge under the suppression of AMO-induced de novo synthesis [13]. All tests were carried out in triplicate except the test with heat-treated cells and with no cells were done in duplicate. For all tests, samples were periodically collected and analyzed for the concentration of EE2 and inorganic nitrogen species in bulk solution for the whole two weeks of the test period.

2.5. Chemical Analysis

Mixed liquor samples were collected from enriched and batch reactors and then, filtered through 0.45 μm Polytetrafluoroethylene membrane filter. Filtrate were analyzed for EE2 and inorganic nitrogen species (NH3, NO2⁻, NO3⁻) while filtrated were measured for MLSS and mixed liquor volatile suspended solid by gravimetric method. EE2 was analyzed by a C18 reversed-phase high-performance liquid chromatography with ultraviolet detector (Shimadzu...
Series LC-10ADvp) using a 60:40 acetonitrile:H₂O as mobile phase. NH₃ was measured by gas-sensitive combination electrode (Cole Palmer, Co.) and salicylated-colorimetric method with UV visible spectrophotometer (Thermo Electron Corporation, Hexious α, Cambridge, UK) at 640 nm [14]. NO₂⁻ was colorimetrically analyzed with UV visible spectrophotometer at 543 nm while NO₃⁻ was analyzed by UV screening method at 220 and 275 nm [14]. pH, DO concentration and temperature in mixed liquor were monitored during sludge enrichment and degradation test. pH value was determined by pH probe (250A+, Thermo Orion Inc., Germany). The concentration of DO was determined by membrane sensing electrode (DO 083005D probe with 850A meter, Thermo Orion Inc., Germany). Temperature was measured by a laboratory glass thermometer.

2.6. Analysis of Ammonia-oxidizing Bacterial Communities

2.6.1. Sample preparation and DNA extraction

Sludge samples (2 mg dry sludge/L) from two enriched reactors including ‘70NS’ and ‘420NS’ were concentrated via centrifugation at 14,000 rpm for 10 min and stored at -20°C. Genomic DNA was extracted from each samples (2 mg dry sludge) using Fast-DNA SPIN kits for soil (QBiogenes, USA) following to the manufacturer’s instruction. The extracted DNA was verified using 1.5% agarose gel electrophoresis (Bio-Rad, Spain).

2.6.2. Polymerase chain reaction (PCR), cloning and sequencing

Primers amoA 1F and 2R [15] (forward 5-GGGGTTTCTACT GGTGGT-3; reverse 5-CCCTCKGSAAAGCCTTCTTC-3) were used for PCR amplification. Bacterial amoA gene fragments were amplified using a Takara polymerase (Takara Bio Inc., Japan) in a thermal cycler (Biorad Laboratories, USA) for 10 min at 95°C, followed by 35 cycles of 60 s at 95°C, 60 s at 56°C, and 30 s at 72°C, followed by 15 min of final extension at 78°C. Later, the amplified PCR products were purified by gel electrophoresis using a NucleoSpin Extract II Kit (Clontech Laboratories Inc., USA). Then, the purified PCR products were cloned using the pGEM-T Easy vector system (Promega, USA). Later, five and four clones of ‘70NS’ and ‘420NS’ samples were randomly chosen for sequencing at Macrogen Inc., Korea, respectively.

### Table 1. System Performance of Enriched Reactors

<table>
<thead>
<tr>
<th>Parameter</th>
<th>A (70 mg N/L NH₃-fed reactor)</th>
<th>B (140 mg N/L NH₃-fed reactor)</th>
<th>C (420 mg N/L NH₃-fed reactor)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Influent NH₃ (mg N/L)</td>
<td>69.14 ± 1.44</td>
<td>138.05 ± 3.79</td>
<td>409.37 ± 17.55</td>
</tr>
<tr>
<td>NH₃ loading rate (mg N/(L-d))</td>
<td>16.59 ± 0.35</td>
<td>33.13 ± 0.91</td>
<td>98.25 ± 4.21</td>
</tr>
<tr>
<td>Effluent NH₃ (mg N/L)</td>
<td>0.49 ± 0.15</td>
<td>1.95 ± 0.58</td>
<td>11.00 ± 10.76</td>
</tr>
<tr>
<td>NH₃ oxidation rate (mg N/(gVSS-day))</td>
<td>219.7</td>
<td>275.0</td>
<td>637.4</td>
</tr>
<tr>
<td>Effluent NO₂⁻ (mg N/L)</td>
<td>0.12 ± 0.08</td>
<td>0.84 ± 0.69</td>
<td>12.35 ± 13.16</td>
</tr>
<tr>
<td>Effluent NO₃⁻ (mg N/L)</td>
<td>66.47 ± 3.38</td>
<td>125.88 ± 3.20</td>
<td>353.55 ± 18.40</td>
</tr>
<tr>
<td>Total N</td>
<td>67.09 ± 3.01</td>
<td>128.67 ± 3.51</td>
<td>376.90 ± 27.03</td>
</tr>
</tbody>
</table>

Dominated AOB cluster: Nitrosomonas oligotropha cluster – Nitrosomonas europaeae-Nitrosococcus mobilis cluster

3. Results and Discussion

3.1. Enriched Reactors Performance

Table 1 and Fig. 1 showed long-term performance of all three enriched reactors. All reactors showed steady performance in terms of NH₃, NO₂⁻ and NO₃⁻ effluent concentrations after five weeks of operation. At steady state condition, NH₃ removal efficiencies...
for reactors fed with NH$_3$ of 70 mg N/L and 140 mg N/L were more than 99%. But for 420 mg N/L NH$_3$-fed reactor, a slight less efficiency for NH$_3$ removal (96.23%) was observed due to receiving higher NH$_3$ load compared to other reactors. In addition, an increase in NH$_3$ load led to stimulate specific NH$_3$ oxidation rate. No NO$_2^-$ accumulated in any reactors indicating NH$_3$ oxidation is the rate determining step for nitrification.

### 3.2. Ammonia-oxidizing Bacterial Communities in Enriched Sludge

Sludge ‘70NS’ and ‘420NS’ were brought to further analyze for ammonia-oxidizing bacterial communities using PCR-cloning-sequencing technique. Results showed that AOB belongs to *Nitrosomonas oligotropha* (N. oligotropha) cluster dominated (4 of 5 clones) in reactor fed with NH$_3$ of 70 mg N/L at which the NH$_3$ concentration were less than 1 mg N/L in effluent (Table 1). AOB belongs to *N. oligotropha* cluster is known as a high NH$_3$ affinity cluster as indicated by low Monod half-saturated coefficient (K$_s$: 0.32-3.16 mg N/L [16, 17]). However, reactor fed with 420 mg N/L NH$_3$, which its effluent contained 15.82 mg N/L NH$_3$, was dominated (all 4 clones) by AOB belongs to *Nitrosomonas europaea-Nitrosoccus mobilis* (N. europaea-N. mobilis) cluster. AOB belongs to *N. europaea-N. mobilis* cluster has much lower NH$_3$ affinity than that belongs to *N. oligotropha* cluster as indicated by much higher K$_s$ (1-100 mg N/L [16-19]). These results corresponded with the finding reported in Limpiyakorn et al. [20]. Limpiyakorn et al. [20] showed that *N. europaea-N. mobilis* cluster dominated in chemostats fed with > 140 mg N/L NH$_3$ while for the reactor fed with ≤ 140 mg N/L NH$_3$, the dominant AOB belonged to *N. oligotropha* cluster.

### 3.3. Batch NH$_3$ Oxidation

#### 3.3.1. Effect of culture history

In order to investigate effect of continuous culture history on batch substrate utilization, three enriched sludge being acclimated with various loads of the growth substrate (‘70NS’, ‘140NS’, and ‘420NS’) were brought to determine the rate of substrate utilization under batch condition. Results showed that continuous culture history of enriched sludge strongly influence on batch NH$_3$ oxidation rate (Table 2). For any given values of initial NH$_3$ concentration, higher NH$_3$ oxidation rate was found in sludge previously being acclimated with higher load of NH$_3$.

This is likely because continuous cultivation of mixed sludge at higher NH$_3$ load leading to the selection for a particular AOB having higher NH$_3$ utilization rate. Generally, the dominant bacterial population mainly determined the rate of substrate utilization and bacterial growth of mixed culture. Sludge dominated by AOB with *N. europaea-N. mobilis* cluster (420NS) showed higher batch NH$_3$ utilization rate as compared with that of dominated by *N. oligotropha* AOB cluster (70NS). For all results shown above, an intrinsic NH$_3$ oxidation rate clearly depended on the physiology of dominant AOB population in enriched nitrifying sludge.

#### 3.3.2. Effect of initial NH$_3$ concentration

As a result, NH$_3$ oxidation was limited by substrate availability. Larger specific NH$_3$ oxidation rate was observed at higher initial NH$_3$ (Fig. 2). However, maximum batch NH$_3$ oxidation rate of enriched sludge was attained under initial NH$_3$ concentration which had been previously used for feeding during sludge enrichment. Enriched sludge in reactors fed with 70 mg N/L NH$_3$ (70NS) achieved its maximum NH$_3$ oxidation rate at 70 mg N/L while at least 140 mg N/L was needed by sludge taken from the 140 and 420 mg N/L NH$_3$-fed reactors (140NS and 420NS). In other words, *N. oligotropha*-dominating sludge (70NS) achieved its maximum NH$_3$ oxidation rate at lower NH$_3$ concentration than that of sludge dominated with *N. europaea-N. mobilis* AOB cluster (420NS). Higher NH$_3$ affinity of the 70NS as compared to that of the 420NS because it was dominated by the *N. oligotropha* cluster which has higher NH$_3$ affinity than the *N. europaea-N. mobilis* cluster.

Moreover, at initial NH$_3$ concentration of 420 mg N/L, slight lower NH$_3$ was oxidized as a result of substrate inhibition. However, sludge dominated with *N. oligotropha* AOB cluster (70NS) were

### Table 2. Comparison of the Rates of EE2 Degradation by Nitrifying Activated Sludge and Enriched Nitrifying Sludge in Aerated Batch Reactor

<table>
<thead>
<tr>
<th>Culture</th>
<th>EE2 degradation rate (mg/gVSS-day))</th>
<th>NH$_3$ oxidation rate (mg N/gVSS-day)</th>
<th>Initial concentration of EE2 &amp; NH$_3$</th>
<th>Ref</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nitrifying activated sludge</td>
<td>0.0936</td>
<td>1,320</td>
<td>1.0 mg/L EE2; 100 mg N/L NH$_3$</td>
<td>[12]</td>
</tr>
<tr>
<td>AMO-containing extract from enriched nitrifying sludge</td>
<td>7.83-29.17</td>
<td>129.6-1,339.2</td>
<td>-</td>
<td>[23]</td>
</tr>
<tr>
<td>Enriched nitrifying sludge</td>
<td>0.096</td>
<td>1,260</td>
<td>0.75 mg/L EE2; 52.5 mg N/L NH$_3$</td>
<td>[26]</td>
</tr>
<tr>
<td>Nitrifying activated sludge</td>
<td>0.024</td>
<td>933.6</td>
<td>0.05 mg/L EE2; 50 mg N/L NH$_3$</td>
<td>[28]</td>
</tr>
<tr>
<td>Enriched nitrifying sludge</td>
<td>0.28-0.52</td>
<td>51.1-225.4</td>
<td>3.0 mg/L EE2; This study</td>
<td></td>
</tr>
<tr>
<td>70NS (N. oligotropha-dominating)</td>
<td>0.31-0.66</td>
<td>67.5-312.7</td>
<td>28-420 mg N/L NH$_3$</td>
<td></td>
</tr>
<tr>
<td>140NS</td>
<td>0.44-0.88</td>
<td>150.9-630.7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>420NS (N. europaea-dominating)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
apparently affected far less by the substrate inhibition than *N. europaea*-dominating sludge indicating higher tolerability to adverse condition in slow-growing AOB. This emphasized an importance of past cultivation condition of sludge on its rate of substrate utilization in batch reactor.

### 3.4. Batch EE2 Degradation

EE2 biodegradation by sludge enriched with different loads of NH3 was investigated to observe effect of culture history on the AMO cometabolism. As shown in Fig. 2 and 3, NH3 and EE2 degradation was observed only in the absence of ATU. Furthermore, there was not any EE2 degradation in batch with heat-treated cells or no cell (data not shown) indicating the loss of EE2 due to abiotic processes (i.e., sorption and volatilization) was negligible in our study. This indicates that AMO cometabolism by AOB is main degradation pathway of EE2 rather than heterotrophic and abiotic degradation that is correspondence to several studies [12, 21-25]. Moreover, EE2 and NH3 degradation in the presence of AMO-induced inhibitor was not found (data not shown) indicating residual AMO utilization by harvested cells in order to complete both substrate degradation was negligible.

EE2 biotransformation clearly depended on NH3 oxidation. During EE2 cometabolism, EE2 was degraded by using a reducing power from NH3 oxidation. Therefore, more NH3 was oxidized, higher EE2 was degraded via the cometabolism of AMO. Moreover, not only initial NH3 concentration largely affected on NH3 oxidation rate but on EE2 biotransformation rate as well. Higher available NH3 improved both the rates of NH3 and EE2 removals indicating a reductant-limiting condition for AMO metabolism and cometabolism. While a reductant-inhibiting condition decreased NH3 oxidation and EE2 degradation at initial NH3 of 420 mg N/L. The results shown here is in good agreement with previous studies [22-24] which found strong positive correlation between initial concentration of NH3 and batch EE2 biotransformation rate by nitrifying sludge.

In addition, after 5 d of sludge incubation with initial NH3 of 28-140 mg N/L, all NH3 were utilized, but substantial biotransformation of EE2 was still occurring even in the absence of NH3. This was consistent with the results obtained by a number of studies [22, 26-28] which revealed the capability of nitrifying sludge to cometabolize EE2 in NH3-free environment. This phenomenon is likely to be a result of utilizing AMO accumulated within the AOB cell for degrading EE2 during NH3-limiting condition.

Moreover, EE2 degradation of enriched sludge was affected by not only initial NH3 level but also the enrichment condition. Higher EE2 biotransformation rate was achieved by sludge previously being enriched with higher NH3 load (Fig. 4). As mentioned earlier, the enrichment of nitrifying bacteria at higher NH3 load may pro

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**Fig. 2.** Profiles of NH3 and NO3− concentration during incubation test with and without adding ATU at initial NH3 concentration of 28 mg N/L (a-c), 70 mg N/L (d-f), 140 mg N/L (g-i) and 420 mg N/L (j-l) using sludge 70NS (a,d,g,j), 140NS (b,e,h,k) and 420NS (c,f,i,l).
Fig. 3. EE2 degradation using different sludge at initial NH$_3$ concentration a) 28 mg N/L b) 70 mg N/L c) 140 mg N/L d) 420 mg N/L.

Fig. 4. The degradation rate of a) NH$_3$ and b) EE2 under different sludge and initial NH$_3$ concentrations.

note specific AOB having higher utilization rate of NH$_3$ over those having the lower one. The results show that AOB population in enriched nitrifying sludge affected on the EE2 degradation in the same manner as NH$_3$ oxidation. Higher EE2 degradation was found in sludge dominated by higher-growth-AOB than lower-growth- AOB-dominating sludge. EE2 cometabolic rate of sludge dominated with the N. europaea-N. mobilis AOB cluster were approximately 1.6-1.7 (420NS) times higher than that belongs to

Fig. 5. The relationship between NH$_3$ and EE2 degradation rates for a) 70NS b) 140NS c) 420NS.
the N. oligotropha AOB cluster (70NS). Greater EE2 cometabolic rate of higher-growth-AOB may be caused by more electron available for cometabolism.

Furthermore, the correlation between NH3 and EE2 degradation rates of sludge was found strongest in the sludge being previously cultivated with highest NH3 load (Fig. 5). It can be implied that more electron provided by NH3 oxidation is needed by sludge being previously cultivated with the highest load of NH3 to cometabolize a certain amount of EE2. Our finding firstly reveals the importance of past AOB growth condition regarding its determination on simultaneous N and EE2 removals. The rates of EE2 biotransformation found in this study were compared with those reported in previous studies (Table 2). Higher rates of EE2 degradation obtained in this study as compared to other studies is likely because much higher initial concentration of EE2 was used.

4. Conclusions

This research firstly determined the importance of culture history of nitrifying sludge on simultaneous NH3 and EE2 removals in wastewater. Higher available NH3 improved both the rates of NH3 and EE2 removals indicating a reductant-limiting condition for AMO metabolism and cometabolism. Sludge being acclimated with higher NH3 load as well as the one dominated with high-AMO metabolism and cometabolism. Sludge being acclimated with highest NH3 load (Fig. 5). It can be implied that more electron provided by NH3 oxidation is needed by sludge being previously cultivated with the highest load of NH3 to cometabolize a certain amount of EE2. Our finding firstly reveals the importance of past AOB growth condition regarding its determination on simultaneous N and EE2 removals. The rates of EE2 biotransformation found in this study were compared with those reported in previous studies (Table 2). Higher rates of EE2 degradation obtained in this study as compared to other studies is likely because much higher initial concentration of EE2 was used.

Acknowledgments

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