Glycerol-driven Denitratation: Process Kinetics, Microbial Ecology, and Operational Controls

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ABSTRACT: Denitratation, the selective reduction of nitrate to nitrite, is a novel process when coupled with anaerobic ammonium oxidation (anammox) could achieve resource-efficient biological nitrogen removal of ammonium- and nitrate-laden waste streams. Using a
fundamentally-based, first principles approach, this study optimized a stoichiometrically-limited, glycerol-driven denitratation process and characterized mechanisms supporting nitrite accumulation with results that aligned with expectations. Glycerol supported selective nitrate reduction to nitrite and near-complete nitrate conversion, indicating its viability in a denitratation system. Glycerol-supported specific rates of nitrate reduction (135.3 mg-N/g-VSS/h) were at least one order of magnitude greater than specific rates of nitrite reduction (14.9 mg-N/g-VSS/h), potentially resulting in transient nitrite accumulation and indicating glycerol’s superiority over other organic carbon sources in denitratation systems. pH and ORP inflection points in nitrogen transformation assays corresponded to maximum nitrite accumulation, indicating operational setpoints to prevent further nitrite reduction. Denitratation conditions supported enrichment of *Thauera* sp. as the dominant genus. Stoichiometric limitation of influent organic carbon, coupled with differential nitrate and nitrite reduction kinetics, optimized operational controls, and a distinctively enriched microbial ecology, was identified as causal in glycerol-driven denitratation.

KEYWORDS: partial denitrification; denitratation; glycerol; short-cut biological nitrogen removal; first-principles approach

1. Introduction

Conventional biological nitrogen removal (BNR), including energy and chemical-intensive nitrification and denitrification, is traditionally used to treat ammonium-laden \((\text{NH}_4^+)\) waste streams. The advent of engineered processes that achieve oxidation of \(\text{NH}_4^+\) to nitrite \((\text{NO}_2^-)\), termed nitritation, combined with denitrification (reduction of \(\text{NO}_2^-\) to nitrogen gas \((\text{N}_2)\))
or anaerobic ammonium oxidation (anammox) represent short-cut BNR alternatives to conventional BNR approaches. Such short-cut BNR processes can provide reductions in chemical (external carbon for denitrification and alkalinity for nitrification) and energy use (aeration for nitrification), driving the desire for NO$_2^-$ accumulation within these processes.

Alternatively, waste streams containing concomitantly high concentrations of NH$_4^+$ and nitrate (NO$_3^-$), such as those resulting from fertilizer$^1$ and explosives manufacturing,$^2,3$ provide similar energy and chemical reduction opportunities through distinct short-cut BNR processes. A particularly effective pathway for treating waste streams containing both NH$_4^+$ and NO$_3^-$ is through heterotrophic$^4-9$ or autotrophic$^{10}$ denitratation (selective reduction of NO$_3^-$ to NO$_2^-$) coupled with downstream anammox. A combined denitratation-anammox system used to treat waste streams containing equal concentrations of NH$_4^+$ and NO$_3^-$ would theoretically reduce aeration energy requirements by 100% and COD requirements by 80% compared to treatment of the same waste stream using conventional BNR. Recent studies$^4-9$ on heterotrophic denitratation have focused on performance in lab-scale sequencing batch reactors (SBRs) driven by acetate, methanol, glucose, and sludge fermentation liquid due to the lack of sufficient readily biodegradable chemical oxygen demand (COD) in typical waste streams. These studies have primarily been observational in nature, with particular emphasis placed on empirically identifying parameters and conditions that potentially contributed to NO$_2^-$ accumulation, such as influent COD:N ratios, pH, ORP, and loading rates. Stoichiometric limitation of influent COD:N ratios, specifically, has been shown to influence endpoint nitrogen speciation.$^{11}$ Various parameter combinations were optimized, denoted by the observation of stable NO$_3^-$-to-NO$_2^-$ conversion ratios as high as 90% during steady-state studies.$^6$
The selection of an external COD source to drive denitrification is critical when attempting to maximize NO$_2^-$ accumulation. Traditionally, methanol has been one of the most widely used external COD sources for denitrification due to its low cost and wide availability.$^{12}$ NO$_2^-$ accumulation has proven difficult with methanol due to methanol dehydrogenase’s direct delivery of electrons to cytochrome $c$ and proximal to NO$_2^-$ reductase as opposed to distribution solely through the ubiquinol pool to NO$_3^-$ reductase similar to other carbon sources.$^{13-15}$ The unique electron delivery locations during methanol oxidation within the respiratory denitrification chain potentially contribute to concomitant NO$_3^-$ and NO$_2^-$ reduction.

Several water resource recovery facilities are switching to glycerol due to the operational and safety risks associated with methanol.$^{12}$ Glycerol is similar in cost to methanol and less expensive than ethanol and acetate,$^{16-18}$ is available as a waste or byproduct,$^{19,20}$ and has no known inhibitory effects on the anammox process, unlike methanol.$^{21}$ NO$_2^-$ accumulation during glycerol supplementation was also anecdotally observed in full-scale treatment plants resulting in unintentional enrichment of anammox on the produced NO$_2^-$.$^{22}$ Nevertheless, to fully realize the operating benefits that a denitratation-anammox system could offer, it is imperative for the parameters and conditions leading to NO$_2^-$ accumulation in a glycerol-driven denitratation system to be systematically identified, defined, and addressed in relation to reactor operating strategies.

Accordingly, the overarching goals of this study were to use a fundamentally-based, first principles approach to characterize the process kinetics, nitrogen conversion efficiencies, and microbial ecology of a glycerol-fed denitratation process, and identify concomitant reactor operating strategies. The specific objectives were to (1) control selective conversion of NO$_3^-$ to NO$_2^-$ through stoichiometric limitation of influent glycerol dose, (2) quantify the rates of NO$_3^-$
reduction relative to rates of NO$_2^-$ reduction and understand their impact on the selective accumulation of NO$_2^-$; (3) elucidate the microbial community structure under varied carbon-loading levels in a functional glycerol-driven denitrification process, and (4) identify operational controls and reactor operating strategies to maximize denitrification rates and efficiencies.

2. Materials and Methods

2.1. Experimental Set-up and Reactor Operation

A lab-scale SBR with a working volume, V=12 L, was operated at room temperature (22±2°C) for a period of 232 d. The SBR was operated at a hydraulic retention time (HRT) of 1 d, utilizing 4 cycles per day with each cycle consisting of a 90-min anoxic feed and react period, a 180-min anoxic react period, a 50-min settling period, and a 40-min decant period. SBR feed contained 100.0 mg/L NO$_3^-$-N as the terminal electron acceptor to simulate the influent of a high NO$_3^-$-containing waste stream typical of a fertilizer$^1$ or explosives$^2$ manufacturing facility, 25.0 mg/L NH$_4^+$-N (to support assimilation), and macro and trace nutrients (Table S1). pH was controlled automatically at 7.50±0.05 using 0.5 M HCl and 1.0 M NaHCO$_3$ via a chemical dosing pump (Etatron D.S., Italy). Sludge wasting was controlled daily at the end of the anoxic feed and react period following COD exhaustion to maintain a solids retention time (SRT) of 3 d. Glycerol, diluted to a 15% solution by volume, served as the external COD source and was provided to meet influent COD:NO$_3^-$-N ratios from 2.4:1 to 5.0:1. Glycerol was fed at the end of the anoxic feed and react period so that examined influent COD:NO$_3^-$-N ratios were met during each cycle. Upon transitioning to each influent COD:NO$_3^-$-N ratio tested, a stabilization period of 4 x SRT was allowed prior to assessing performance relative to other conditions. Sequencing
and timing of SBR cycles and daily solids wasting was controlled and maintained by peristaltic
pumps (Masterflex, IL) using electronic timers (ChronTrol Corporation, CA).

2.2. Sample Collection and Wastewater Quality Analysis

All analytical procedures employed were in accordance with Standard Methods.\textsuperscript{23} Aqueous-phase samples were withdrawn during the decant period of the reactor cycle and concurrently from the influent for chemical species analysis after centrifugation (8,000 x G, 10 min, 4-8°C) to remove cells and cell debris. NO$_3^-$ and NH$_4^+$ were measured using ion selective electrodes (Thermo Fisher Scientific, MA). NO$_2^-$ concentration was measured via diazotization and colorimetry.\textsuperscript{23} The fraction of influent NO$_3^-$ lost to nitrogenous gases was determined via mass balance on nitrogen. Centrifuged aqueous-phase samples were filtered using 0.20 µm syringe filters (A Chemtek, MA) and stored at -20°C. Dionex ICS-2100 ion chromatography using a Dionex IonPac AS-18 IC column (Thermo Fisher Scientific, MA) was used to confirm ion selective electrode and colorimetric measurements of NO$_3^-$ and NO$_2^-$ concentrations, respectively. Similarly, a Dionex IonPac AS-14 IC column (Thermo Fisher Scientific, MA) was used to quantify volatile fatty acid production during unbuffered \textit{ex situ} batch kinetic assays. Separate aqueous-phase samples were extracted at the end of the anoxic react period and during the decant period of the reactor cycle to assess total biomass concentrations in the reactor and effluent, respectively, for SRT control. Aqueous-phase samples taken during the decant period were centrifuged (8,000 x G, 10 min, 4-8°C) and filtered using 0.45 µm syringe filters (A Chemtek, MA) to assess remaining soluble COD (sCOD) concentrations (Hach Chemical Company, CO). Biomass concentrations were approximated by subtracting sCOD measurements from total COD measurements to determine particulate COD (pCOD) (Hach Chemical
Company, CO). Additional aqueous-phase samples taken just prior to the end of the anoxic react period were centrifuged (8,000 x G, 10 min, 4-8°C), supernatant was discarded, and cell pellets were preserved at -80°C for subsequent DNA extraction and 16S rRNA gene sequencing.

2.3. Feeding Strategy Experiments

Two feeding strategies were tested to maximize NO$_2^-$ accumulation. A semi-continuous feeding strategy delivered NO$_3^-$-containing SBR feed and glycerol continuously for the first 75 and 72 min, respectively, of the anoxic feed and react period (Fig. S1). A pulse feeding strategy delivered a pulse of NO$_3^-$-containing SBR feed and glycerol every 45 min for the first 270 min of the SBR cycle (Fig. S1). Feeding rates were controlled to maintain equivalent mass loading rates of NO$_3^-$ and glycerol and influent COD:NO$_3^-$-N ratios for the two feeding strategies.

2.4. Batch kinetic assays

Batch assays, in situ (within the SBR) and ex situ, were conducted to measure extant process kinetics and optimize operational controls, including batch duration, pH, and ORP. In situ assays followed previously described sampling collection and chemical analysis procedures. Aqueous-phase samples were obtained from the primary SBR at steady-state over the course of a single 360-min reactor cycle. Ex situ assays were carried out in an anoxic, sealed, spinner flask batch vessel with a working volume, V=1 L, at room temperature (22±2°C). Mixed liquor was taken from the primary SBR at steady-state during the feed and react period, washed 4 times using SBR feed without NO$_3^-$, and supernatant was discarded. Prior to extant kinetic batch assays, the medium was buffered to pH 7.50 using 0.5 M HCl and 1.0 M NaHCO$_3$ and N$_2$ gas was sparged until dissolved oxygen (DO) levels were equal to 0.01 mg/L O$_2$, or the minimum
practical limit of the InPro 6850i polarographic DO sensor with M300 transmitter (Mettler-Toledo, OH). pH was maintained at pH 7.50±0.05 by manual control. pH optimization batch assays were conducted within normal pH operating ranges (see Supporting Information (SI)).

NO$_3^-$ and glycerol were dosed to meet the desired initial COD:NO$_3^-$-N ratio. NO$_3^-$ was dosed at the outset of the experiment (time=0 min) and the biomass was incubated for 30 min prior to the addition of glycerol to ensure that residual nitrogen species and glycerol from the primary SBR remaining in the washed mixed liquor were consumed prior to data collection. pH, ORP, and DO were measured and recorded continuously via an InPro 3253i/SG pH/ORP electrode and an InPro 6850i polarographic DO sensor, respectively, attached to an M300 transmitter (Mettler-Toledo, OH). Following extant kinetic batch assays, linear regression with $R^2 \geq 95\%$ of NO$_x$-N species from time points of maximum concentration to minimum concentration for each respective species was performed with pCOD concentrations taken just prior to glycerol input to determine true specific rates of NO$_3^-$ reduction (sDNaR) (Eqn. 1) and NO$_2^-$ reduction (sDNiR) (Eqn. 2). NO$_2^-$ production resulting from NO$_3^-$ reduction was not accounted for in the determination of specific rates of NO$_2^-$ reduction, yet this remains representative of a true reduction rate. During the time points assessed for each influent COD:NO$_3^-$-N ratio, NO$_3^-$ removal was complete or near-complete (<3% of initial dose) except at influent COD:NO$_3^-$-N=2.5:1 where NO$_3^-$ concentration measurements confirmed no continued NO$_3^-$ reduction. pCOD measurements were used to determine maximum specific substrate consumption rates (Eqns. 1-2).

$$sDNaR = \left( \frac{1}{\chi} \left( \frac{\Delta S_{NO_3^-}}{\Delta t} \right) \right)$$

Eqn. 1
\[ s_D N_i R = \left( \frac{1}{X} \right) \left( \frac{\Delta S_{NO_3^-}}{\Delta t} \right) \]  \hspace{1cm} \text{Eqn. 2}

Where:

- \( s_D N_a R \): maximum specific \( NO_3^- \) consumption rate (mg \( NO_3^- \)-N/g VSS/h)
- \( s_D N_i R \): maximum specific \( NO_2^- \) consumption rate (mg \( NO_2^- \)-N/g VSS/h)
- \( X \): volumetric biomass concentration approximated using pCOD measurements (g VSS/L)

\[ \frac{\Delta S_{NO_3^-}}{\Delta t} \]: volumetric substrate (\( NO_3^- \)) consumption rate (mg \( NO_3^- \)-N/L/h)

\[ \frac{\Delta S_{NO_2^-}}{\Delta t} \]: volumetric substrate (\( NO_2^- \)) consumption rate (mg \( NO_2^- \)-N/L/h)

2.5. DNA Extraction, Next-Generation Sequencing of Amplicon Library, and Bioinformatics

DNA was extracted from biomass samples and purified using a QIAamp DNA Mini Kit (Qiagen, Inc., MD). The quality and quantity of DNA were checked using a NanoDrop Lite spectrophotometer (Thermo Fisher Scientific, MA). Barcoded fusion primers with Ion Xpress™ sequencing adapters (Thermo Fisher Scientific, MA) and a 16S rRNA bacterial 1055F/1392R universal primer set were applied in each sample for multiplex sequencing. Amplification of genomic DNA targets was performed with \( \text{iQ}^\text{TM} \) SYBR® Green Supermix (Bio-Rad, CA) and purification via Agencourt AMPure XP Reagent (Beckman Coulter, CA). Library quantification was performed with an Agilent DNA 1000 Kit (Agilent, CA). Template preparation with the DNA library followed by Ion Spheres Particle (ISP) enrichment was performed using Ion OneTouch2 (Ion PGM Hi-Q View OT2 Kit). Enriched ISP was loaded onto an Ion Torrent 318 v2 BC chip and run on an Ion Torrent Personal Genome Machine (Ion PGM Hi-Q View Sequencing Kit). Ion Torrent Suite software was used for base calling, signal processing, and
quality filtering (Phred score of >15) of the raw sequences. The 1055F/1392R universal primer set targeted sequences of approximately 350 base pairs (bp). Mothur software was used to initially screen out likely incorrect amplicon sequences with bp lengths more than 50 bp different than the target sequence length.\textsuperscript{25} AfterQC software was utilized to further delete bad quality reads (Phred score of <20) and trim the tails of reads where quality dropped significantly.\textsuperscript{26} DADA2 programming via R Studio software was used to produce a table of non-chimeric amplicon sequence variants from the demultiplexed fastq files.\textsuperscript{27} QIIME2 software was applied in conjunction with the Silva version 132 reference taxonomy for further post-sequencing bioinformatic analysis.\textsuperscript{28}

2.6. Nitrogen Conversion Calculations

Reactor performance was normalized with respect to the influent characteristics. A NO\textsubscript{2}\textsuperscript{-} accumulation ratio (NAR) (Eqn. 3) was defined to relate the accumulation of NO\textsubscript{2}\textsuperscript{-} to the removal of NO\textsubscript{3}\textsuperscript{-}.\textsuperscript{29} A NAR equal to 100% indicated that all NO\textsubscript{3}\textsuperscript{-} removed accumulated as NO\textsubscript{2}\textsuperscript{-} compared to terminal reduction to N\textsubscript{2} gas, for which the NAR would be 0%.

\begin{equation}
NAR = \left(\frac{NO_{2\text{eff}} - NO_{2\text{inf}}}{NO_{3\text{inf}} - NO_{3\text{eff}}}\right) \times 100\%
\end{equation}

\textbf{Eqn. 3}

NO\textsubscript{3}\textsuperscript{-} reduction was also classified in terms of a NO\textsubscript{3}\textsuperscript{-} reduction ratio (NRR) (Eqn. 4), which normalized the conversion of NO\textsubscript{3}\textsuperscript{-} to the influent NO\textsubscript{3}\textsuperscript{-} concentration.\textsuperscript{9} A NRR equal to 100% would indicate conversion of all influent NO\textsubscript{3}\textsuperscript{-} to any reduced form, while a NRR of 0% would indicate no conversion.

\begin{equation}
NRR = \left(\frac{NO_{3\text{eff}} - NO_{3\text{inf}}}{NO_{3\text{inf}}}\right) \times 100\%
\end{equation}

\textbf{Eqn. 4}
\[ NRR = \left( \frac{(NO_{3}^{\text{inf}}-N) - (NO_{3}^{\text{eff}}-N)}{NO_{3}^{\text{inf}}-N} \right) \times 100\% \]  

Eqn. 4

3. Results and Discussion

3.1. Denitratation Reactor Performance

The influent COD:NO\textsubscript{3}^-\textsubscript{-N} ratio required for glycerol-driven denitrification (NO\textsubscript{3}^-\textsubscript{-N to N\textsubscript{2}} reduction) was thermodynamically\textsuperscript{30} determined to be 5.9:1 (see SI). This corresponded well with experimentally-determined operational ratios of 4.2:1 to 5.6:1,\textsuperscript{16,20,31} although the lowest reported ratio\textsuperscript{16} may not be fully representative as it was determined via \textit{ex situ} batch assays as opposed to steady-state continuous flow bioreactor or SBR operation. Stoichiometric analysis revealed that influent COD:NO\textsubscript{3}^-\textsubscript{-N}=2.4:1 (see SI) would provide only enough electrons via COD oxidation to reduce NO\textsubscript{3}^-\textsuperscript{'} to NO\textsubscript{2}^-\textsuperscript{'} on a theoretical electron equivalence basis as opposed to full denitrification. Therefore, influent COD:NO\textsubscript{3}^-\textsubscript{-N} ratios between 2.4:1 and 5.9:1 were referred to as stoichiometrically-limited for the purposes of this study. These calculations form the fundamentally-based foundation to the first principles approach used in this study to conduct and interpret the results of glycerol-driven denitratation presented herein.

The utilization of glycerol as the external COD source and electron donor resulted in significant NO\textsubscript{2}^-\textsuperscript{'} accumulation at stoichiometrically-limited influent COD:NO\textsubscript{3}^-\textsubscript{-N} ratios from 2.5:1 to 5.0:1, indicating that the use of glycerol was feasible to sustain a denitratation process. The highest degrees of NO\textsubscript{3}^-\textsuperscript{'} removal and NO\textsubscript{2}^-\textsuperscript{'} accumulation, as a function of influent COD:NO\textsubscript{3}^-\textsubscript{-N} ratio during steady-state SBR operation, occurred at influent COD:NO\textsubscript{3}^-\textsubscript{-N}=3.0:1 (Fig. 1). This resulted in an average NO\textsubscript{2}^-\textsuperscript{'} accumulation of 60.8±11.5 mg/L NO\textsubscript{2}^-\textsubscript{-N} (n=10) and NAR of 62\%, indicating that 62\% of the NO\textsubscript{3}^- reduced was converted to NO\textsubscript{2}^- rather than terminally reduced to N\textsubscript{2} gas. Additionally, the NRR was determined to be 96\%, indicating that...
a majority of the influent NO$_3^-$ was converted leaving only approximately 4% of influent NO$_3^-$ in the effluent (Table 1). Accumulation of NO$_2^-$ at influent COD:NO$_3^-$-N=2.8:1 compared to influent COD:NO$_3^-$-N=3.0:1 was not significantly different (p=0.49, α=0.05, n=10). Substantial NO$_3^-$ accumulation occurred at influent COD:NO$_3^-$-N=2.8:1 (31.7±11.4 mg/L NO$_3^-$-N, n=11), signifying that this ratio was less operationally optimal compared to influent COD:NO$_3^-$-N=3.0:1. The observed NO$_3^-$ accumulation at influent COD:NO$_3^-$-N=2.5:1 and 2.8:1 may be due to lower COD-supported biomass concentrations leading to reduced denitrification rates. However, effluent sCOD concentrations were negligible signifying that glycerol was nearly completely consumed (sCOD and biomass concentration data not shown). In situ performance profiles (Fig. 2) did not show significant endogenous denitrification, potentially indicating that COD uptake and storage was minimal. Rather, the observed NO$_3^-$ accumulation in these cases indicated that the influent COD:NO$_3^-$-N was not sufficient, potentially due to unrealized COD requirements for cell maintenance and synthesis or additional demand by fully denitrifying microorganisms remaining in the microbial community. Therefore, influent COD:NO$_3^-$-N=3.0:1 was selected as the optimal ratio due to the similar NO$_2^-$ accumulation to influent COD:NO$_3^-$-N=2.8:1 coupled to less than 4% of the influent NO$_3^-$ remaining in the effluent. The high sensitivity at influent COD:NO$_3^-$-N<3.0:1 highlighted significant implication for accurate system operation and control. A minimal reduction in influent COD:NO$_3^-$-N ratio from 3.0:1 to 2.8:1 yielded a sevenfold increase in effluent NO$_3^-$, signifying that strict control of the glycerol-driven denitratation system must be maintained. To this end, online dosing control based on appropriate signals of reactor performance seems necessary to maximize concomitant NO$_3^-$-N conversion selectively to NO$_2^-$ during partial denitratation.
Analysis of variance (ANOVA) across the influent COD:NO$_3^-$-N ratios identified a statistically significant difference in NAR ($p=4.8 \times 10^{-11}$, $\alpha=0.05$, $n=38$) with a decrease from 62% to 11% as the influent COD:NO$_3^-$-N ratio approached that for glycerol-driven denitrification (5.9:1; see SI). Further Holm-Sidak post-hoc multiple comparison analysis indicated that the significant difference in NAR was primarily caused by the expectedly lower NAR at influent COD:NO$_3^-$-N=5.0:1 ($p<9.7 \times 10^{-5}$ for all comparisons, $\alpha=0.05$; Table S2). The decrease in NAR from influent COD:NO$_3^-$-N=4.0:1 to 5.0:1 was most likely attributable to excess available COD.

Previous studies$^{4,6}$ observed that varying the influent COD:NO$_3^-$-N ratio had a negligible effect on the NAR determined at the point of maximum NO$_2^-$ accumulation during ex situ batch experiments, while a separate batch study$^{33}$ concluded that the COD source, as opposed to the influent COD:NO$_3^-$-N ratio, impacted the NAR more readily. In contrast, another separate batch study$^7$ concluded that NO$_2^-$ accumulation was influenced by both the COD source and COD dosing. While insightful, the utility of these results$^{4,6,7}$ to guide steady-state denitratation processes is limited as these studies failed to acclimate their batch experiment seed sludge to the conditions being investigated, which likely contributed to their discrepancy with the current study. Despite investigating the impact of various influent COD:NO$_3^-$-N ratios, Ge et al.$^7$ utilized a fully denitrifying inoculum, whereas Du et al.$^6$ inoculated batch experiments assessing various influent COD:NO$_3^-$-N ratios with a microbial community acclimated to a single stoichiometrically-limited influent COD:NO$_3^-$-N ratio. Both seed sludges likely contained phenotypes with NO$_2^-$ accumulation capabilities different than those expected following acclimation to the investigated conditions. Cao et al.$^4$ did not report conditions of their batch inoculum.
In an improvement over these previous efforts, our current study utilized a sludge stabilization and acclimation period of 4 x SRT following influent COD:NO$_3^-$-N ratio changes. This intentionally allowed the microbial community to adapt to the influent COD:NO$_3^-$-N ratio being investigated. In doing so, it was observed that the influent COD:NO$_3^-$-N ratio had similar impacts on NAR during both steady-state operation (Fig. 1) and ex situ batch assays, with NO$_2^-$ accumulation decreasing as influent COD:NO$_3^-$-N ratios increased (Fig. S2).

In comparison to other steady-state operation studies$^{6,9,34}$ using primarily sodium acetate as the external COD source, glycerol-driven NARs were at least 10% lower (Table 1). While most reported acetate-driven denitrification NARs were greater than 80%, glycerol-driven denitrification yielded NARs less than 70%. These respective acetate-driven steady-state studies$^{6,9,34}$ were deemed reasonable comparisons due to similar COD dosing regimens and results were reported for study periods sufficient in length to assume microbial community acclimation to and stabilization at the studied conditions. Despite this, the assessment of reactor performance based solely upon reported NARs can be misleading as the index does not account for complete or other conversion of influent NO$_3^-$, Thus, NAR=100% does not necessarily indicate that all influent NO$_3^-$ was converted. Several studies,$^{4-6,34}$ however, reported NRRs of nearly 100% that when coupled with a NAR approaching 100% indicated near-perfect denitrification performance (Table 1). It follows then that optimal performance in the current study occurred at influent COD:NO$_3^-$-N=3.0:1 with NAR=62% and NRR=96%. The inability of glycerol to achieve similar efficiency to acetate- or fermentate-driven denitrification is not currently understood. Possible explanations include a greater intracellular carbon and microbial energy storage mechanism during low substrate availability,$^{35,36}$ the COD-source supported enrichment of a microbial consortium with a greater abundance of true denitrifiers,$^{37}$ an
inefficient metabolism in support of denitratation due to a less direct assimilability of glycerol, or
the downstream delivery of electrons on the electron transport chain similar to methanol.\textsuperscript{14,15}

Effluent sCOD measurements, as an estimation of residual glycerol concentration, averaged 9.4±8.8 mg/L COD (n=29) across all influent COD:NO\textsubscript{3}⁻-N ratios assessed. The ca. 96% average decrease from influent to effluent sCOD indicates that nearly all of the glycerol was consumed, and that reactor cycle duration was adequate for COD consumption.

A likely contributing factor to the need for a higher than the theoretical influent COD:NO\textsubscript{3}⁻-N ratio (see SI) was an incomplete enrichment for a solely denitratating or progressive onset\textsuperscript{38} phenotype-dominated microbial community. The presence of microorganisms that express a complete denitrification metabolic pathway or those that exhibit a rapid, complete onset of denitrification genes\textsuperscript{38} would impose a competitive demand on influent COD, thus decreasing its availability for selective reduction of NO\textsubscript{3}⁻ to NO\textsubscript{2}⁻. This additional COD demand would result in a high NRR but low NAR, or significant gaseous-N products with limited NO\textsubscript{2}⁻ accumulation, which was supported by the results herein (Table 1).

3.2. Process Kinetics

Notably, extant kinetic analysis indicated that transient NO\textsubscript{2}⁻ accumulation at all influent COD:NO\textsubscript{3}⁻-N ratios assessed was potentially due to at least one order of magnitude greater specific rates of NO\textsubscript{3}⁻ reduction compared to the specific rates of NO\textsubscript{2}⁻ reduction driven by glycerol (Table 2).\textsuperscript{39} Observed performance at influent COD:NO\textsubscript{3}⁻-N>3.0:1 (Fig. S2) also supported this assertion as the maximum NO\textsubscript{2}⁻ accumulated never equaled the initial NO\textsubscript{3}⁻ concentration, indicating that there was concomitant reduction of NO\textsubscript{3}⁻ and NO\textsubscript{2}⁻. However, performance at influent COD:NO\textsubscript{3}⁻-N=3.0:1 resulted in near-complete selective reduction of
NO$_3^-$ to NO$_2^-$ prior to terminal reduction to N$_2$ gas (Fig. S2). It should be emphasized that the kinetic profiles in Fig. S2 were obtained from acclimated biomass from individual SBRs operated for at least 4 x SRT at each influent COD:NO$_3^-$-N ratio. In general, measured specific rates of NO$_3^-$ reduction and $\mu_{\text{max}}$ values were higher than those previously reported for glycerol-driven full denitrification studies (Table 2) and may be due to differences in the microbial community that was selected for by stoichiometric limitation during our current denitratation-specific study. Glycerol-driven specific rates of NO$_3^-$ reduction values were nearly double those reported for acetate-driven systems at similar influent COD:NO$_3^-$-N ratios, but slightly lower than those observed in an experiment utilizing a combination of external COD sources garnered from sodium acetate and endogenous carbon in a domestic wastewater stream (Table 2). The ratios of sDNaR:sDNiR achieved in this study with glycerol across different influent COD:NO$_3^-$-N values were also higher than previously reported with acetate (Table 2). This difference may be due to variations in the direct assimilability of each COD source with more assimilable COD sources such as glycerol or endogenous carbon in these cases supporting greater specific rates of NO$_3^-$ reduction, or the COD source-supported microbial community.

3.3. NO$_2^-$ Accumulation through the Management of Operational Controls

3.3.1. Denitratation Control via Batch Duration

Batch duration was identified as an effective process control parameter to maximize NO$_2^-$ accumulation. The duration of the anoxic feed and react period could be shortened to achieve comparable or improved performance. NO$_2^-$ concentrations decreased following peaks of NO$_2^-$ accumulation at higher influent COD:NO$_3^-$-N ratios (4.0:1, 5.0:1; Fig. 2). This decrease was not
observed at influent COD:NO$_3^-$-N=3.0:1, indicating that excess COD remained following completion of denitrification at higher ratios. Despite minimal NO$_2^-$ reduction following peak NO$_2^-$ accumulation at influent COD:NO$_3^-$-N=2.5:1, overall performance remained low, making this ratio less effective at achieving partial denitrification (Table 1; Fig. 2).

Results generally supported that influent COD:NO$_3^-$-N ratios have an inverse relationship with time to maximum NO$_2^-$ accumulation during the anoxic react period. Batch duration could be reduced to 150 minutes or less, or the time to maximum NO$_2^-$ accumulation (Fig. 2). Subtraction of the feed and react period of the SBR cycle from the reduced batch duration, by extension, would yield an optimal react time equivalent to a continuous flow system’s HRT (Fig. 2). The optimal react time is representative of when glycerol is available for NO$_3^-$ reduction in both systems. Therefore, the identified optimal react times in our SBR system would be equivalent to HRTs of approximately 30 minutes (COD:NO$_3^-$-N=4.0:1 and 5.0:1) to 60 minutes (COD:NO$_3^-$-N=2.5:1 and 3.0:1) in continuous flow systems operating at each respective influent COD:NO$_3^-$-N ratio.

3.3.2. Denitrification Control via pH and ORP

During unbuffered and non-carbon limited operation (influent COD:NO$_3^-$-N≥5.9:1), the denitrification-dominated phase of the denitrification profile exhibited a distinct decrease in the reactor’s pH and increase in the ORP until both reached inflection points after which pH increased and ORP decreased (Fig. 3). At this inflection point, NO$_3^-$ reduction decelerated due to the depletion of available NO$_3^-$ allowing for observable concomitant NO$_2^-$ reduction thus decreasing the NAR and negatively impacting the objective of maximizing NO$_2^-$ accumulation. Continuous monitoring of pH and ORP could provide an observable real-time control to maximize
denitratation. While feedforward online control of COD dosing tied to influent NO\textsubscript{x} loading has proven effective in controlling denitratation,\textsuperscript{17} this system requires online NO\textsubscript{x} sensors which may not be achievable at all plants due to potentially high capital\textsuperscript{40} and maintenance costs.\textsuperscript{41}

Rather, denitratation control via pH and ORP observation could provide a backup check or serve as a less costly alternative\textsuperscript{40} with widely available and utilized sensors.

pH and ORP were previously reported as control parameters for denitrification driven by acetate, methanol, endogenous carbon, soybean wastewater, and brewery wastewater.\textsuperscript{6,7,33,42,43} Contrary to the distinct glycerol-driven pH and ORP profile observed in the current study, Ge et al.\textsuperscript{7} and Du et al.\textsuperscript{6} described acetate-driven profiles exhibiting a general increase in pH whereby a “turning point” separated denitratation from denitritation. However, the observed pH profiles obtained experimentally in our study (Fig. 3) are in excellent concurrence with theoretically calculated net production of 0.43 equivalents of acidity per mole NO\textsubscript{3}-- reduced to NO\textsubscript{2}-- (Eqn. 5), which supported the observed pH fluctuation profiles.

\begin{equation}
\text{NO}_x^- + (0.14)C_3H_8O_3 \rightleftharpoons \text{NO}_2^- + (0.43)CO_2 + (0.57)H_2O
\end{equation} \text{Eqn. 5}

For completeness, stoichiometry (Eqn. 6) reveals that denitritation should result in a net consumption of 0.36 equivalents of acidity per mole NO\textsubscript{2}-- reduced to N\textsubscript{2} gas at pH 7.5.

\begin{equation}
\text{NO}_2^- + (0.21)C_3H_8O_3 + H^+ \rightleftharpoons (0.50)N_2 + (0.64)CO_2 + (1.36)H_2O
\end{equation} \text{Eqn. 6}
3.3.3. Denitrification Control via Feeding Strategy

The pulse feeding strategy resulted in a statistically significant improvement in denitrification performance ($\alpha=0.05$; $n=8$) over the semi-continuous feeding strategy in both NO$_2^-$ accumulation ($p=0.03$) and NO$_3^-$ reduction ($p=0.0003$), indicating that feeding methodology impacted the performance of the system (Table S3). As both feeding strategies maintained equivalent influent COD:NO$_3^-$-N ratio per substrate pulse or for the duration of the semi-continuous feeding period, this difference in system performance was thought to be influenced by the temporal distribution of substrate pulses. Those pulses occurring later in the anoxic feed and react period may have limited the time for the biotransformation of NO$_3^-$ to gaseous nitrogen thus allowing for greater NO$_2^-$ accumulation. This is counter to the semi-continuous feeding strategy, where fully denitrifying microorganisms within the microbial community had the full anoxic feed and react period to reduce influent NO$_3^-$-N. Therefore, in a continuous-flow BNR process, the spatial location of introducing glycerol could be another factor to promote partial denitrification if possible. Optimizing the dosing location of electron donors is quite widely practiced for increasing the efficiency of COD utilization even for full denitrification in step-feed BNR or Bardenpho configurations.44

3.4. Microbial Ecology

Proteobacteria was the most dominant phylum out of 14 identified at all influent COD:NO$_3^-$-N ratios (Fig. 4a). $\beta$-Proteobacteria made up at least 73% of the Proteobacteria phylum at all influent COD:NO$_3^-$-N ratios. In a survey of wastewater denitrifying bacterial 16S rDNA sequences retrieved from GenBank, Lu et al.45 found that approximately 72% of
prokaryotic microorganisms displaying denitrifying capabilities were taxonomically affiliated with *Proteobacteria*, while β sub-class affiliated microorganisms were typically abundant in denitrifying activated sludge,\textsuperscript{1,45,46} similar to the findings herein.

Within β*-Proteobacteria*, the *Rhodocyclaceae* and *Comamonadaceae* families were identified as those mainly involved in denitrification in activated sludge.\textsuperscript{46,47} Our findings supported this as *Thauera* sp., that belongs to the *Rhodocyclaceae* family within β*-Proteobacteria* was enriched as the most dominant genus with a relative abundance of nearly 80\% at influent COD:NO\textsubscript{3}−-N=3.0:1 (Fig. 4b). *Comamonadaceae* fam. was not found, indicating that their enrichment may not be favored under stoichiometrically-limited conditions imposed herein. Certain *Thauera* spp. strains were characterized according to two distinct regulatory phenotypes,\textsuperscript{48} including the immediate and simultaneous onset of all denitrification genes with no detectable NO\textsubscript{2}− accumulation, as well as the progressive and sequential onset of denitrification cascade genes with appreciable NO\textsubscript{2}− accumulation.\textsuperscript{38} Selective pressures were not identified for either, although the selection for progressive onset denitrifiers would be critical to facilitate denitratation. The coupling of a high relative abundance of *Thauera* sp. (Fig. 4b), high NRR, and high NAR (Table 1), with the ability to perform full denitrification when presented with sufficient COD (Fig. S2) indicated that the application of stoichiometric limitation in the influent COD:NO\textsubscript{3}−-N as a selective pressure may favor the progressive onset over rapid, complete onset phenotype. *Thauera* sp. may represent a key functional microorganism for denitratation systems indicated by its decreasing relative abundances away from the optimal influent COD:NO\textsubscript{3}−-N (Fig. 4b). Several recent denitratation-specific studies\textsuperscript{4,6,9,49} further supported this argument with reported *Thauera* sp. relative abundances from 55\% to 73\% under limited influent COD:NO\textsubscript{3}−-N ratios with acetate as the external COD source.
despite different seed sludges. In comparison, acetate-driven full denitrification studies reported no more than 12% relative abundance of *Thauera* sp. Therefore, the application of a stoichiometrically-limited influent COD: NO$_3^-$-N ratio as a selective pressure in a denitrification system may impart a stronger impact on the denitrifying community structure than previously recognized.

4. Conclusions

Denitrification, with downstream anammox processes, offers chemical and energy reductions through resource-efficient BNR of NH$_4^+$ and NO$_3^-$-laden waste streams. A fundamentally-based, first-principles approach was used to propose an influent COD:N ratio and other operating parameters that would promote denitrification and experimental results aligned with expectations. Glycerol supported the process kinetics and microbial ecology necessary to selectively convert NO$_3^-$ to NO$_2^-$ in denitrification systems. Process control strategies, including influent COD loading and pH, ORP, and batch duration operational setpoints were identified and used to further define reactor operating strategies that could maximize denitrification performance. Significant enrichment indicated *Thauera* sp. may represent a key functional microorganism in denitrification systems. This study implicated stoichiometric limitation of influent organic carbon, unique microbial community enrichment, and differential NO$_3^-$ and NO$_2^-$ reduction kinetics as determinant factors in glycerol-driven denitrification.

ADDITIONAL INFORMATION

E-supplementary data can be found in online version of the paper.
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Author Contributions: The manuscript was written through contributions of all authors. All authors gave approval to the final version of the manuscript. All authors contributed equally.

Notes: The authors declare no competing financial interests.

ACKNOWLEDGMENTS
This study was supported by Project Director, Joint Services, project USMA1740. Views and opinions expressed or implied herein are solely those of the authors and should not be construed as policy or carrying the official sanction of the Department of Defense, United States Army, United States Military Academy, or other agencies or departments of the U.S. Government.

REFERENCES


**Figure 1.** Steady-state denitratation performance and respective NAR and NRR assessed at each influent COD:NO$_3$-N ratio. *Effluent gaseous-N contributions were calculated via mass balance.
Table 1. Influence of external COD source and influent COD:NO$_3$-N ratios on denitrification performance.

<table>
<thead>
<tr>
<th>External COD Source</th>
<th>Influent COD:NO$_3$-N</th>
<th>NAR [%]</th>
<th>NRR [%]</th>
<th>Reactor Type</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sodium Acetate</td>
<td>3.0</td>
<td>51 – 73</td>
<td>~73 - 93</td>
<td>USB$^a$</td>
<td>34</td>
</tr>
<tr>
<td></td>
<td>3.0</td>
<td>80</td>
<td>~100</td>
<td></td>
<td>6</td>
</tr>
<tr>
<td></td>
<td>2.75</td>
<td>83</td>
<td>~100</td>
<td></td>
<td>34</td>
</tr>
<tr>
<td></td>
<td>2.5</td>
<td>87</td>
<td>85</td>
<td></td>
<td>9</td>
</tr>
<tr>
<td>Sodium Acetate / Domestic Wastewater</td>
<td>3.1$^b$</td>
<td>90</td>
<td>~100</td>
<td>SBR</td>
<td>4</td>
</tr>
<tr>
<td>Fermentation Effluent</td>
<td>3.0</td>
<td>80</td>
<td>~100</td>
<td></td>
<td>5</td>
</tr>
<tr>
<td>Glycerol</td>
<td>2.5</td>
<td>65</td>
<td>54</td>
<td></td>
<td>This study</td>
</tr>
<tr>
<td></td>
<td>2.8</td>
<td>69</td>
<td>73</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>3.0</td>
<td>62</td>
<td>96</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>4.0</td>
<td>57</td>
<td>97</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>5.0</td>
<td>10</td>
<td>99</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

$^a$ Upflow sludge blanket reactor (USB)

$^b$ Reported influent ratio includes COD associated both with the domestic wastewater and external COD source.
Table 2. Summary of process kinetic parameters for both full denitrification and denitrification studies with respect to external COD source and influent COD: NO\textsubscript{3}\textsuperscript{-}N ratio.

<table>
<thead>
<tr>
<th>COD Source</th>
<th>Inf. COD: NO\textsubscript{3}\textsuperscript{-}N [mg N/L]</th>
<th>Inf. NO\textsubscript{3}\textsuperscript{-}N [mg N/L]</th>
<th>( \mu_{\text{max}} ) [d\textsuperscript{-}1]</th>
<th>sDNA\textsubscript{R} \textsuperscript{b} [mg N/g VSS/h]</th>
<th>sDNi\textsubscript{R} \textsuperscript{c} [mg N/g VSS/h]</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sodium Acetate</td>
<td>1.22</td>
<td>2,700</td>
<td>--</td>
<td>23.0\textsuperscript{f}</td>
<td>19.0\textsuperscript{f}</td>
<td>51</td>
</tr>
<tr>
<td></td>
<td>5.0</td>
<td>150</td>
<td>--</td>
<td>82.3</td>
<td>32.0</td>
<td>6</td>
</tr>
<tr>
<td></td>
<td>1.0</td>
<td>--</td>
<td>--</td>
<td>52.0</td>
<td>--</td>
<td>7</td>
</tr>
<tr>
<td></td>
<td>6.0</td>
<td>--</td>
<td>--</td>
<td>280.0</td>
<td>--</td>
<td></td>
</tr>
<tr>
<td>Sodium Acetate / Domestic WW</td>
<td>3.4\textsuperscript{e}</td>
<td>1,000</td>
<td>--</td>
<td>190.0</td>
<td>--</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>5.0</td>
<td>100</td>
<td>--</td>
<td>6.5\textsuperscript{a,d}</td>
<td></td>
<td>20</td>
</tr>
<tr>
<td>Glycerol</td>
<td>26.0</td>
<td>22.5</td>
<td>3.4</td>
<td>1.7\textsuperscript{a,b}</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>26.0</td>
<td>22.5</td>
<td>2.0</td>
<td>1.35\textsuperscript{a,c}</td>
<td></td>
<td>16</td>
</tr>
<tr>
<td></td>
<td>2.5</td>
<td>100</td>
<td>--</td>
<td>112.3</td>
<td>1.8</td>
<td></td>
</tr>
<tr>
<td></td>
<td>3.0</td>
<td>100</td>
<td>--</td>
<td>135.3</td>
<td>14.9</td>
<td></td>
</tr>
<tr>
<td></td>
<td>5.0</td>
<td>100</td>
<td>--</td>
<td>147.1</td>
<td>40.9</td>
<td></td>
</tr>
<tr>
<td></td>
<td>20.0\textsuperscript{g}</td>
<td>100</td>
<td>6.2</td>
<td>--</td>
<td>--</td>
<td></td>
</tr>
</tbody>
</table>

\textsuperscript{a} Rates reported as mg NO\textsubscript{3}\textsuperscript{-}N/g VSS/hr based upon full denitrification studies.
\textsuperscript{b} Rate reported in study exhibiting no NO\textsubscript{2} accumulation.
\textsuperscript{c} Rate reported in study exhibiting NO\textsubscript{2} accumulation.
\textsuperscript{d} Suspended phase rates reported; biofilm rates not reported for comparison purposes to current study.
\textsuperscript{e} Reported influent ratio includes COD associated both with the domestic wastewater and external COD source.
\textsuperscript{f} Rates reported from original study for the pH utilized in current study.
\textsuperscript{g} Batch experiment used biomass acclimated to influent COD: NO\textsubscript{3}\textsuperscript{-}N=3.0.
\textsuperscript{h} Specific rate of NO\textsubscript{3} reduction (sDNA\textsubscript{R})
\textsuperscript{i} Specific rates of NO\textsubscript{2} reduction (sDNi\textsubscript{R})
Figure 2. Representative in situ NO$_2^-$-N profiles identified the optimal batch duration obtained during steady-state operation at each respective influent COD:NO$_3^-$-N ratio. Optimal batch durations corresponded to the points of maximum NO$_2^-$ accumulation at each respective influent COD:NO$_3^-$-N ratio. Decreases in NO$_2^-$ concentrations during the feed and react period were attributed to dilution.
Figure 3. NO$_x$, pH, and ORP profiles depicting the pH (a) and ORP (b) inflection points at the point of maximum NO$_2^-$ accumulation prior to which denitratation was dominant and after which denitrification became dominant (influent COD:NO$_3^-$-N=10.0:1; microbial ecology acclimated to influent COD:NO$_3^-$-N=3.0:1). Influent COD was provided in excess and beyond that at which biomass was acclimated in order to drive the process beyond denitrification and demonstrate the ability of pH and ORP to serve as denitrification process controls even under non-ideal influent COD:NO$_3^-$-N ratios.
Figure 4. Taxonomic analysis of the microbial consortium at the phylum (a) and genus (b) taxonomic levels under optimal operating conditions (influent COD: NO₃⁻-N=3.0:1, SRT=3 d). The grouping “Other” comprises OTUs with less than 1% total relative abundance (among all samples summed).