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Removal of antibiotic resistance genes in an anaerobic membrane bioreactor treating primary clarifier effluent at 20 °C

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Anaerobic membrane bioreactors (AnMBR) play a key role in future plans for sustainable wastewater treatment and resource recovery because they have no energy-intensive oxygen transfer requirements and can produce biomethane for renewable energy. Recent research results show that they can meet relatively stringent discharge limits with respect to BOD₅ and TSS when treating municipal wastewater primary effluent. Sustainable used water recovery plans should also consider removal of unregulated pollutants. Antibiotic resistance genes (ARGs) represent an important emerging contaminant due to public health concerns surrounding the spread of infections resistant to common antibiotics. Conventional activated sludge processes have demonstrated mixed results regarding ARG removal. The objective of this research was to determine the impact of an AnMBR on ARG removal when treating municipal primary clarifier effluent at 20 °C. AnMBR treatment resulted in 3.3 to 3.6 log reduction of ARG and the horizontal gene transfer determinate, *int1*, copies in filtrate. Membrane treatment significantly decreased the total biomass as indicated by a decrease in 16S rRNA gene concentration. Microbial community analysis via Illumina sequencing revealed that the relative abundance of putative pathogens was higher in membrane filtrate compared to primary effluent although the overall bacterial 16S rRNA gene concentrations was lower in filtrate. Membrane treatment also substantially reduced microbial diversity in filtrate compared to anaerobic reactor contents.

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Water impact

Mainstream anaerobic wastewater treatment is a promising technology for sustainable water resource recovery. Anaerobic membrane bioreactors (AnMBRs) can meet relatively stringent BOD₅ and TSS regulatory standards (BOD₅ <10 mg L⁻¹, TSS <10 mg L⁻¹) for municipal wastewater at temperatures as low as 10 °C. The research reported herein demonstrated that AnMBR technology can also significantly reduce antibiotic resistance gene copies.

Introduction

Treating wastewater to only meet discharge limits may be unsustainable due to increasing population, urbanization, water stress, resource consumption and water reuse plans.¹ These factors also increase the demand on food and energy. Collectively, these challenges may be better overcome by con-

sidering not just wastewater treatment, but the holistic food–energy–water nexus.² The practice of employing used water reclamation is being more widely embraced as an opportunity to yield value-added products while reducing energy demands and yielding clean water at water resource reclamation facilities (WRRFs).^{1,3} WRRFs provide a centralized opportunity to recover nutrients (particularly phosphorus), reduce energy demands or produce energy, and provide water for reuse.^{4–6} To implement resource recovery, WRRFs may need to move beyond conventional activated sludge treatment that requires energy for oxygen transfer and biosolids disposal with limited opportunities for nutrient recovery.^{3,7}

Since anaerobic treatment is often a key technology for sustainable water resource recovery management scenarios of the future, anaerobic membrane bioreactors (AnMBRs) have garnered attention for multiple reasons.^{3,7} AnMBRs produce less solids than aerobic systems and do not require energy

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for aeration. Additionally, they produce methane that could be recovered for energy.^{8–10} Recent research results show that AnMBRs can yield effluent BOD₅ less than 10 mg L⁻¹ at temperatures as low as 10 °C.^{11,12} Furthermore, AnMBR filtrate can be treated using processes such as ion-exchange to concentrate nutrients and recover valuable products such as struvite.^{5,6}

In addition to producing valuable products and recovering energy, sustainable water resource recovery must produce water having appropriate quality for reuse. Chemical micropollutants are prevalent in wastewater and pose a variety of ecological risks including fish feminization and the spread of antibiotic resistance.^{13–18} Results vary in the few existing publications regarding chemical micropollutant removal through AnMBRs.^{19,20} Additional research is warranted to help determine if pre- or post-treatment is needed to achieve chemical micropollutant removal.²¹ Removal of biological micropollutants, including antibiotic resistance genes (ARGs), is also important to consider when evaluating treatment options.

Over 23 000 deaths a year in the US are attributed to infections caused by antibiotic resistant bacteria.²² Health costs associated with antibiotic resistant infections are approximately \$1 billion per year.²² The World Health Organization has deemed antibiotic resistance to be a major threat to modern medicine and noted that the role the environment plays in increasing antibiotic resistance is a serious concern.²³ Appropriate technical solutions could be applied at hotspots, such as WRRFs,²⁴ to limit ARG dissemination and slow the spread of antibiotic resistance.²⁵ Indeed, ARGs can even be enriched during activated sludge treatment.^{26,27} Thus, alternative treatment processes to conventional activated sludge could be beneficial to limit dissemination of ARGs in the environment.

Understanding the impact of AnMBRs on the fate of ARGs is important to evaluate sustainable water resource recovery options to remove emerging pollutants while minimizing energy costs and generating value-added products. Membrane treatment is a plausible option to increase ARGs removal. Membrane (100 kDa) filtration of wastewater spiked with plasmid-associated ARGs resulted in 1- to 2-log reduction of ARGs.²⁸ Therefore, a membrane bioreactor could result in ARGs removal due to physical filtration alone. During the biological process, ARGs increased in an aerobic bioreactor in one full-scale study, but were significantly lower in MBR filtrate compared to the wastewater treated.²⁹ The ARGs decrease was correlated to decreased 16S rRNA gene copies.²⁹ In general, aerobic MBR treatment results in higher removal of ARGs than conventional activated sludge treatment,³⁰ but little research is available that describes how AnMBR processes impact ARG fate.

The objective of this research was to determine changes in ARG abundance due to AnMBR treatment of primary clarifier effluent. The most effective mechanism for ARG removal was hypothesized to be membrane filtration rather than biological treatment. To test this hypothesis, a laboratory-scale an-

aerobic fluidized bed bioreactor followed by an external crossflow membrane was fed primary clarifier effluent from a full-scale water reclamation facility. The abundance of individual ARGs, *sul1*, *erm(B)*, and *tet(O)*, as well as, the integrase from class 1 integrons (*intI1*) and 16S rRNA genes were quantified at different points within a lab-scale AnMBR. Furthermore, the microbial communities were analyzed in the membrane bioreactor effluent and the membrane filtrate.

Materials and methods

AnMBR setup, operation, and sampling

An AnMBR comprised of a fluidized bed bioreactor (FBR) followed by an external, ceramic tubular crossflow membrane with a nominal 0.05 µm pore size (Type1/16, Atech Innovations, Gladbeck, Germany) was operated at 20 °C (Fig. 1) as described by Seib *et al.*^{9,12} Operational, performance, and energy demand details can be found elsewhere.^{9,12} Briefly, the FBR hydraulic residence time (HRT) was 6 h and the nominal solids retention time (SRT) was 49 d. The FBR was initially fed synthetic primary effluent (235 ± 35 mg L⁻¹ BOD₅; 115 ± 40 mg L⁻¹ volatile suspended solids [VSS]) for 320 d and then fed actual primary clarifier effluent (160 ± 60 mg L⁻¹ BOD₅; 77 ± 25 mg L⁻¹ volatile suspended solids [VSS]) from a full-

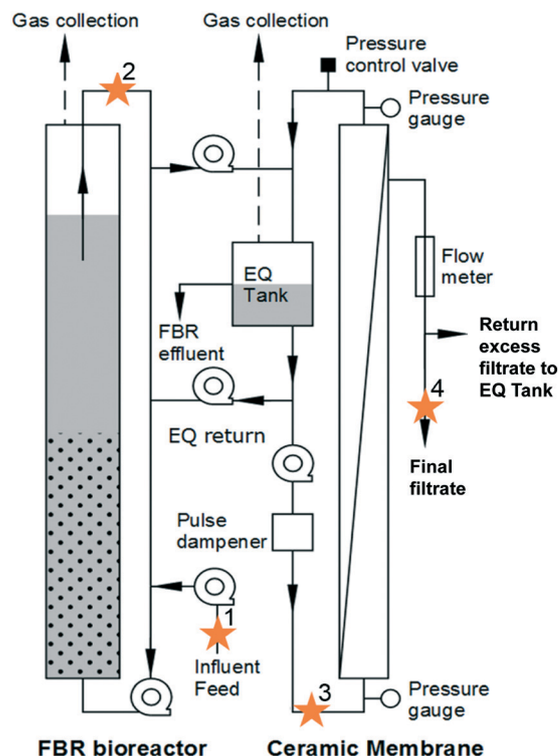


Fig. 1 Schematic of anaerobic membrane bioreactor (AnMBR) system with sampling points labelled with stars. The system consisted of a fluidized bed reactor (FBR bioreactor) that contained biomass and granular activated carbon media. Primary influent feed (1) entered the FBR reactor. FBR effluent (2) was conveyed to an equalization tank that served as the influent to the ceramic membrane (3). Final filtrate (4) after the membrane was also collected for analysis.



scale facility (South Shore WRRF, Oak Creek, WI) for 45 d.⁹ The FBR with external crossflow membrane achieved greater than 90% BOD₅ and chemical oxygen demand (COD) removal, with filtrate BOD₅ less than 10 mg L⁻¹.

Samples were taken on days 348, 349, 352, and 353. Four locations were sampled (Fig. 1): (1) primary influent feed, (2) FBR effluent, (3) liquid in the equalization tank connected to the membrane that served as influent to the membrane (membrane feed), and (4) final filtrate from the membrane. For sample locations 1 through 3, 50 mL were collected and centrifuged at 15 000 × *g* for 15 minutes. The sample pellet was resuspended in 1.0 mL lysis buffer (CLS-TC) and transferred to a 2 mL centrifuge tube. For the final filtrate (4) samples, 500 mL was filtered through a 0.2 μm filter. The filter was placed in a 2 mL centrifuge tube and 1.0 mL lysis buffer was added. The resuspended pellet and filter with lysis buffer were stored at -20 °C until DNA was extracted.

Molecular methods

DNA extraction. Samples for ARG quantification, filter or pellet, containing lysis buffer (CLS-TC) underwent three freeze–thaw cycles utilizing liquid nitrogen for improved lysis.³¹ DNA purification from lysed samples was performed according to manufacturer instructions (FastDNA SPIN kit, MP Biomedicals, Santa Ana, CA). Extracted DNA was quantified using a spectrophotometer (NanoDrop Lite, ThermoFisher Scientific, Waltham, MA).

qPCR. Quantitative polymerase chain reaction (qPCR) using previously published primers was employed to enumerate total bacterial 16S rRNA gene,²⁴ antibiotic resistance genes (*erm(B)*,³² *tet(O)*,³² and *sul1*³³) and horizontal gene transfer related gene (*int1*²⁴). All qPCR reactions were run with 20 μL reaction volumes and consisted of 1× PowerUp SYBR Green Master Mix (Applied Biosystem; Foster City, CA) and primers at a final concentration of 1.0 μM. DNA template was added as 5 μL of DNA diluted by 1:100 or greater per reaction. Each sample was run in triplicate and results averaged. If only two of the three replicates were quantifiable, then the average was taken of the two quantifiable results. Single quantification of three replicates was reported at the quantification limit. Measurements of a no-template control and standards containing target gene DNA between 10⁰ and 10⁸ copies were also performed in triplicate. Cycling conditions were as follows: 2 min at 50 °C to activate the uracil-DNA glycosylase (UDG), 10 min at 95 °C to inactivate UDG and activate the DNA polymerase, 40 cycles of 95 °C for 30 s, 60 °C for 30 s, followed by 72 °C for 30 s. Following amplification, melting curves were acquired. Reactions were carried out using a LightCycler® 96 (Roche Applied Science, Mannheim, Germany). Results were reported as copy number (CN) per L or standardized to total 16S rRNA CN. Specificity of amplification of target genes in samples was confirmed by melt curves consistent with that of the standard. Amplification efficiency was determined by the resulting standard curve and was considered acceptable be-

tween 0.9 and 1.1. Efficiency was consistent with previously published ranges for this designed assay.³⁴ Coefficient of determination in the linear regression of the standard curves were greater than 0.99, consistent with the designed assay.³⁴ The qPCR limit of quantification for ARGs and related genes was 5 copy numbers (CNs), which was equivalent to 5 × 10⁴ CN L⁻¹ for final filtrate and 5 × 10⁵ CN L⁻¹ for samples taken at primary influent feed, FBR effluent, and membrane feed (Fig. 1) based upon the volume of sample, resuspension volume following DNA extraction, and volume of DNA used in PCR. The 16S rRNA gene limit of quantification was 500 CN in a reaction.

Microbial sequencing and bioinformatic analysis. Amplicon sequencing of the V4 hypervariable region (primers: GTGCCAGCMGCCGCGGTAA and GGAC TACHVGGGTWTCTAAT, excluding barcodes) of the 16S rRNA gene from bacteria was conducted at the University of Minnesota Genomic Center. After V4 amplification, indexing, and quantification of PCR products, 250 cycles on an Illumina MiSeq (v2 chemistry) produced paired-end reads. Raw paired-end reads were processed using Mothur (v1.35.1)³⁵ as described by Kozich *et al.*³⁶ on the Pere cluster of the MUGrid at Marquette University. Taxonomic assignment of processed sequences was performed using the SILVA v123 reference database.³⁷ Results were converted to a BIOM formatted file and used for statistical analysis.

Analysis and statistics. Statistical analyses were performed using the open statistical program 'R'.³⁸ Analysis of variance (ANOVA) by the 'aov' function was used to determine significant differences among samples in the qPCR results for each gene quantified. The 'transformTukey' function of the 'rcompanion' package was used to determine if transformation of data was needed prior to ANOVA. *Post hoc* multiple pairwise comparisons were conducted using Tukey's honestly significant difference (HSD) test by the 'TukeyHSD' function. For analysis of sequences, the BIOM file generated *via* Mothur was imported into R using the 'phyloseq' package.^{39,40} 'Phyloseq' was used for general visualization of sequence data. Alpha-diversity metrics and plots were generated *via* the 'plot_richness' function. ANOVA was used to determine significance among the alpha-diversity metrics. Abundance data were transformed for equal sampling depth prior to generation of the distance matrix used for the non-metric multidimensional scaling (NMDS) plot and multivariate analysis. The Horn–Moristia index^{41,42} was used as the distance matrix, which is a measure of the similarity in taxa compositions and relative abundance values among samples. The 'adonis' function, permutational multivariate analysis, from the 'vegan' package was used to determine significance between the membrane feed and filtrate taxa. The NMDS plot was generated from the distance matrix using 'phyloseq'. Parametric fitting of dispersions to the mean intensity and Wald significance test of coefficients in a negative binomial in the 'DESeq2' package were used to determine significant differential abundance in individual taxonomy between the membrane feed and filtrate bacterial



community.⁴³ Rarefaction curves were generated using the 'ggrare' function from the 'phyloseq-extended' package of scripts.⁴⁴

Results and discussion

Enumeration of antibiotic resistance and related genes

All antibiotic resistance genes (*erm(B)*, *tet(O)*, and *sul1*) and *intI1* abundance values in the final filtrate were between 3 and 4 log units lower than those in primary influent, FBR effluent and the membrane feed ($p < 0.01$) (Fig. 2). In contrast, ARG and *intI1* abundance values were not considerably different among primary influent, FBR effluent or membrane feed water samples; however, the *tet(O)* and *sul1* gene abundance values were greater in the primary influent and membrane feed water compared to the FBR effluent (Fig. 2, $p < 0.05$).

There was a significant 3.94 log reduction of abundance in the final filtrate compared to the other sampled locations in the *tet(O)* gene ($p < 0.01$) conferring resistance to tetracycline through protection of the ribosome.⁴⁵ A significant 3.7 log reduction in *erm(B)* abundance was determined compared to the other sampled locations ($p < 0.01$) indicating a loss in resistance to macrolides, lincosamides, and streptogramin B antibiotics through modification of 23S rRNA.⁴⁶ There was a modest but significant 0.66 log₁₀ greater abundance of the *erm(B)* gene within the primary influent compared to FBR effluent (Fig. 2, $p < 0.05$), suggesting a loss of

resistance within the fluidized bed reactor and before membrane filtration. The final filtrate has a significant 3.6 log reduction in *sul1* abundance compared to all other sampled locations (Fig. 2, $p < 0.01$) indicating a loss of sulfonamide resistance by expression of a dihydropteroate synthase with less affinity for the drug.⁴⁷ The *sul1* gene is normally found linked to other resistance genes in class 1 integrons.⁴⁷ The class 1 integrons have been utilized to quantify the potential of horizontal gene transfer in the bacteria community, specifically in the spread of antibiotic resistance.^{47,48} The class 1 integron (*intI1*) had a similar log reduction as the *sul1* gene (Fig. 2) indicating a reduction in the potential for horizontal gene transfer in the bacterial community of the final filtrate.

Membrane filtration alone led to a 3.3–3.6 log reduction in ARGs and *intI1* in the final filtrate compared to the membrane feed which is consistent with the removal of antibiotic resistant bacteria (ARB) and cell-free ARG-bearing plasmids (extracellular DNA) observed by Cheng and Hong for AnMBRs.⁴⁹ However, the type of wastewater treated and the temperature employed by Cheng and Hong differed from those employed herein; Cheng and Hong operated an AnMBR at 35 °C and fed synthetic wastewater that was spiked with bacteria containing ARGs.⁴⁹ Additionally, the methods of sampling in the current study, low-speed centrifugation with disposal of supernatant and filtration reduces the probability of acquiring free-extracellular DNA. However, the methods are not efficient enough to distinguish between bound-extracellular DNA and intracellular DNA.⁵⁰ Extracellular DNA containing ARGs has the potential for horizontal gene transfer through transformation of bacteria within the environment, including wastewater treatment plants.^{51–56} The detection of bound-extracellular DNA and intracellular DNA would have been similarly dependent on the ability of amplification, therefore representing a risk of dissemination of antibiotic resistance through vertical or horizontal gene transfer. The results herein demonstrate that AnMBR treating actual wastewater at ambient temperature reduce ARGs.

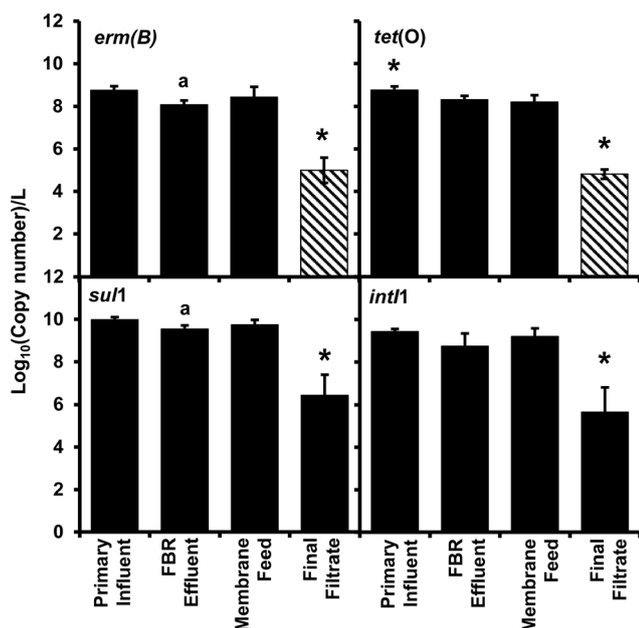


Fig. 2 Abundance of antibiotic resistance genes *erm(B)*, *tet(O)*, and *sul1* and the class 1 associated integrase gene (*intI1*) per liter from the AnMBR. Bars represent the average value and error bars represent the standard deviation from four sampling events. Striped bars represent reported results equivalent to the detection limit because only one or two samples out of four were quantifiable. The asterisk (*) denotes significant difference from all other sample locations, the (a) denotes significant difference compared to primary sample.

Enumeration of 16S rRNA genes

There were no significant differences among 16S rRNA gene concentration measurements for primary influent, FBR effluent, and membrane feed samples ($p \geq 0.34$; Fig. 3). There was a significant 3-log decrease in biomass indicated by 16S rRNA gene copy number in the final filtrate compared to samples from all other locations within the AnMBR system ($p < 0.01$). The 16S rRNA gene copy log removal was consistent with bacterial removal *via* anaerobic digestion as reviewed by Avery *et al.*⁵⁷ and an AnMBR operated at 35 °C that also demonstrated a 3-log reduction in total bacteria copy numbers.⁵⁸

Changes in abundance of the ARGs and *intI1* normalized to 16S rRNA gene copies (Fig. 4) showed no significant difference (ANOVAs, $p > 0.85$) suggesting no selective removal of any specific gene over another gene.



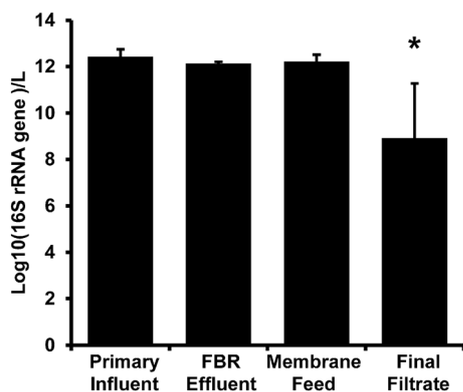


Fig. 3 16S rRNA gene abundance per liter from the AnMBR (Fig. 1). Bars represent the average value and error bars represent the standard deviation from four sampling events. The asterisk (*) denotes significant difference from all other sample locations.

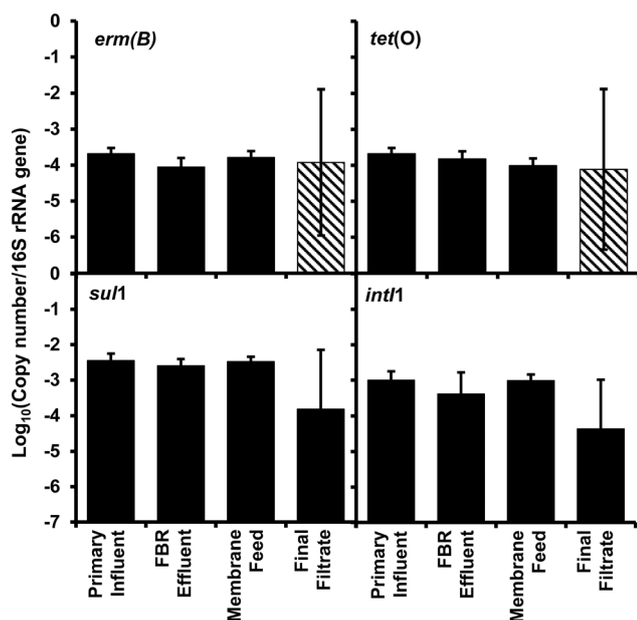


Fig. 4 Relative abundance of antibiotic resistance genes *erm(B)*, *tet(O)*, and *sul1* and the class 1 associated integrase gene (*int1*) per 16S rRNA gene collected from the AnMBR (Fig. 1). Bars represent the average value and error bars represent the standard deviation from four sampling events. Striped bars represent reported results equivalent to the detection limit because only one or two samples out of four were quantifiable.

Membrane feed and membrane filtrate microbial communities

A total of 1 730 988 16S rRNA gene reads were generated, with reads per sample ranging from 970 to 359 868. Taxonomic assignments of 12 713 unique OTUs designated 47 Phyla, 227 Families, and 588 Genera. Alpha-diversity metrics measuring diversity at each sampling location (Fig. 5), showed significantly greater diversity in sequences in the membrane feed compared to the final filtrate (ANOVA, $p < 0.05$) in the Chao1, ACE, Shannon, Simpson, Inverse Simpson, and Fisher

indices. The change of diversity in sequences suggests that the AnMBR is selective for specific bacteria in the membrane feed through the membrane.

In addition, membrane feed and the membrane final filtrate communities were significantly different as determined using the Horn–Moristia index,^{41,42} visualized by NMDS (Fig. 6) and calculated by permutational multivariate analysis of variance ($p = 0.029$). These results are consistent with previous observations by Harb and Hong who reported distinct communities within the influent and effluent from an AnMBR operated at 35 °C treating real wastewater.⁵⁸

Significant differences ($p < 0.01$) in dominant genera between membrane feed and membrane final filtrate were determined by DESeq (Fig. 7). Genera that include known opportunistic pathogens and showed increased abundance included: *Alcaligenes*, *Staphylococcus*, *Sphingomonas*, *Stenotrophomonas*, *Arcobacter* and *Pseudomonas*.^{53,57,59–61,63–67} While the membrane final filtrate showed significant decreases in biomass, the increased relative abundance of these opportunistic pathogens suggests positive selection for these genera. In the US Center for Disease Control and Prevention (CDC) Antibiotic Resistance Threat Report in 2013, methicillin-resistant *Staphylococcus aureus* (MRSA) and vancomycin-resistant *Staphylococcus aureus* (VRSA) were each listed as a serious and concerning level of hazard.²² Interestingly, the *Stenotrophomonas* genus contains a multidrug resistant opportunistic pathogen.⁶³ Further study by quantification of specific virulence genes associated with these and other opportunistic pathogens would yield information on the potential to select for pathogens.

Of those genera that showed significant increases in relative abundance between the membrane feed and membrane filtrate (Fig. 7), *Alcaligenes* and *Arcobacter* are known to perform anaerobic respiration,^{67,68} *Staphylococcus*, *Sulfuricurvum*, and *Pseudomonas* are facultative anaerobes,^{66,68} while *Syntrophomonas* are identified as anaerobic.^{69,70} Similarly, the aerobic genera of *Sphingomonas*, *Stenotrophomonas*, *Leucobacter*, *Thiovirga*, and *Comamonas* also showed significant increases in relative abundance in the membrane filtrate compared to the membrane feed.^{62,63,71–75} Only two genera showed significant decreases in relative abundance between the membrane feed and membrane filtrate, consisting of the genus *Sulfurospirillum*, able to ferment fumarate under anaerobic conditions⁶⁸ and the strictly anaerobic *Desulfuromonas*.⁷⁶ The decrease in *Desulfuromonas* relative abundance might have been due to its lack of tolerance to oxygen. However, the increases in both anaerobic and aerobic bacteria suggest that selection based on oxygen tolerance was absent or weak.

The genera *Alcaligenes*, *Sphingomonas*, *Syntrophomonas*, *Thiovirga*, *Stenotrophomonas*, *Leucobacter*, *Pseudomonas*, *Comamonas*, and *Desulfuromonas* have been associated with wastewater treatment and resource recovery processes including anaerobic digestors and AnMBR treatment. *Alcaligenes* taxa have been identified in anaerobic digestors.⁷⁷ *Sphingomonas*, while being aerobic, have been identified in anaerobic digestors, though their role remains unclear.⁷¹ The



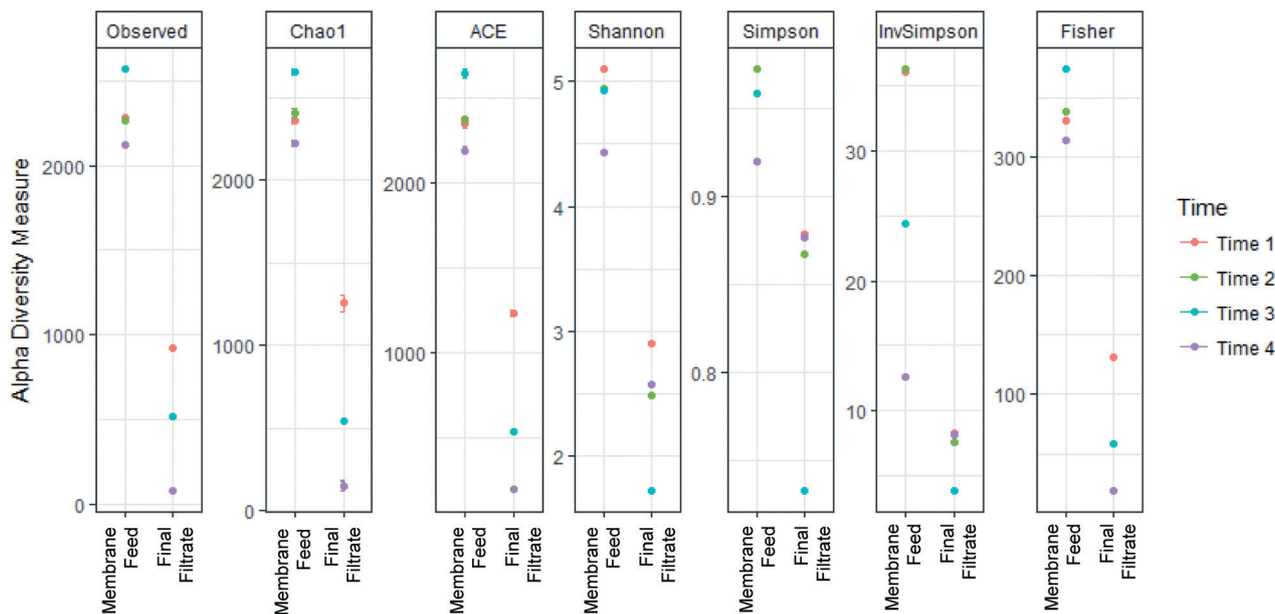


Fig. 5 Membrane substantially decreased richness and diversity between the membrane feed and membrane final filtrate.

Syntrophomonas genus include anaerobic fatty-acid-oxidizing bacteria that are associated with methanogens and are common to anaerobic bioreactors when long-chain fatty-acids are available^{69,70} indicating they are present in the studied FBR. *Thiovirga* are chemolithoautotrophic, sulfur-oxidizing bacteria which were first isolated from a microaerobic wastewater biofilm while investigating its sulfur cycle.⁷³ *Stenotrophomonas*, while aerobic, has been previously

detected in the effluent of a lab-scale AnMBR treating municipal wastewater.⁵⁸ *Leucobacter* genus have been isolated from activated sludge. *Pseudomonas* genera have been detected and the species *Pseudomonas aeruginosa* has been quantified previously from the filtrate of an AnMBR.⁵⁸ *Comamonas* are aerobic bacteria which have been identified in AnMBR, activated sludge, and anaerobic digesters.^{74,75} *Desulfuromonas* were identified in an AnMBR feed with synthetic textile industry wastewater.⁷⁸

Differences in primary effluent and membrane filtrate microbial communities showed that the AnMBR system selected for specific bacteria. This selection was presumably due to size exclusion, however, the sizes of the bacteria represented in the membrane feed and membrane filtrate are comparable based on published direct observation which are usually taken under ideal growth conditions.⁷⁹ The bacterial cell wall structure is an important factor in bacteria breakthrough of small-pore-size membranes.^{80–82} The bacterial cell wall can be elastic and deform under pressure or become less rigid dependent on the environment, such as the presence of antibiotics.^{83–85} The presence of foulant is known to improve the removal of extracellular DNA containing ARGs and ARB,⁴⁹ which most likely led to the variability of the microbial community and differential removal of ARGs found in the final filtrate. Optimization of the control of the level of foulant could improve ARB and ARG removal.⁴⁹

Collectively, the survey of the relative abundance of 16S rRNA genes in membrane final filtrate and membrane feed samples demonstrated a significant increase in the relative abundance of genera containing opportunistic pathogens as well as decreased diversity in the membrane filtrate compared to the membrane feed samples. The changes indicated selective removal at the membrane and/or selective growth or regrowth of bacteria in the membrane filtrate tank.

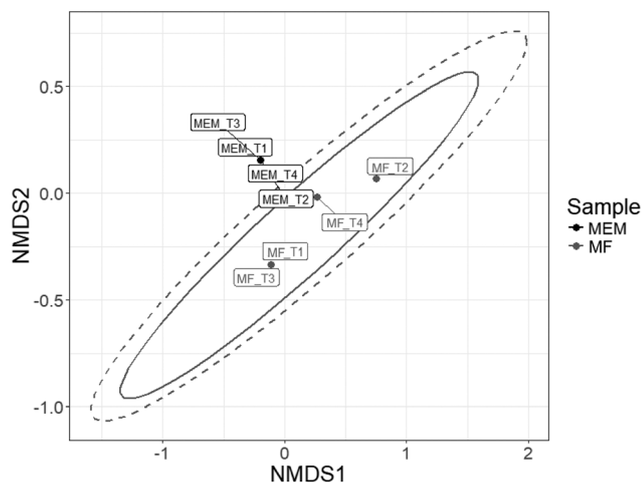


Fig. 6 Nonmetric multidimensional scaling (NMDS) plots of 16S rRNA genus data from membrane feed (MEM) and membrane final filtrate (MF). MF samples do not overlap with the MEM samples indicating these microbial communities are different. The microbial communities that thrive in filtrate represent microbes that made it through membrane at a disproportionately higher rate than the rest of the community or represent microbes that infiltrated the effluent tube from the collection basin. Ellipses represent 95% confidence interval assuming a multivariate normal distribution (dotted) or multivariate t-distribution (solid).



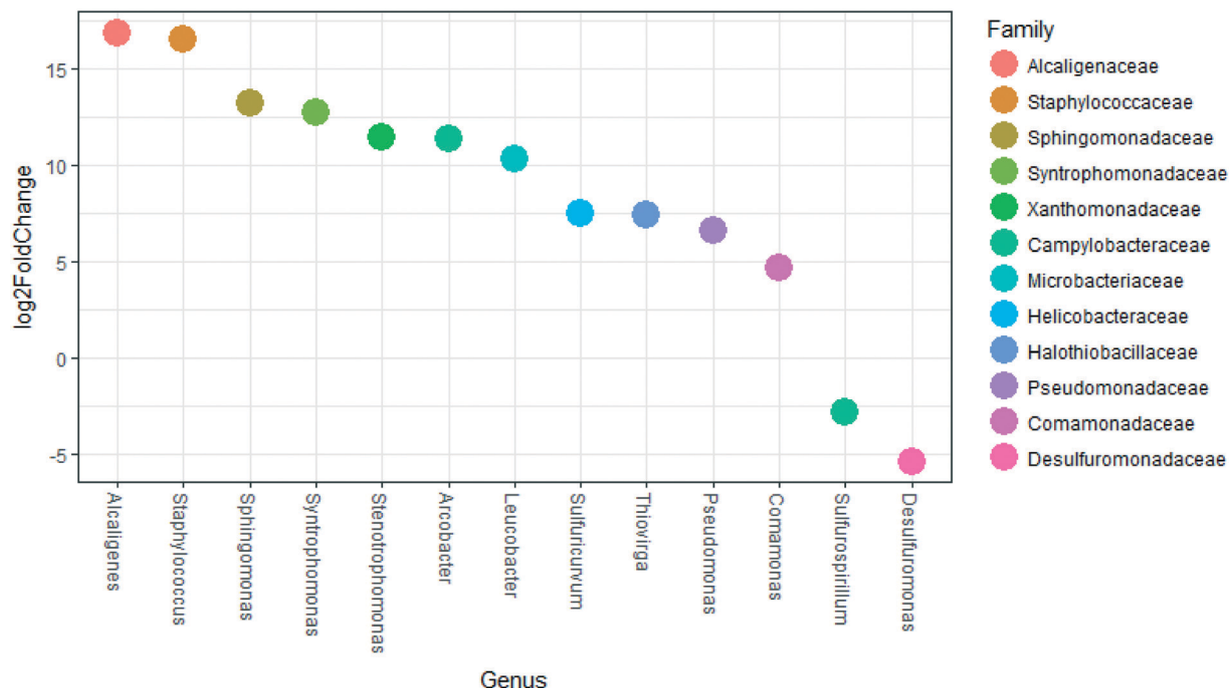


Fig. 7 Genera demonstrating a significant ($p < 0.01$) increase or decrease in relative abundance within the final membrane filtrate (MF) compared to the membrane feed (MEM). DESeq2 differential abundance analysis expressed as log₂ fold-change comparison of membrane feed and final filtrate.

Conclusions

This research revealed that AnMBR significantly reduce ARGs in primary clarifier effluent. ARGs are an important emerging contaminant as antibiotic resistance is a major public health concern.⁸⁶ We no longer have the luxury to view drinking water, wastewater, and stormwater as isolated entities, but rather as one water. It is important to develop and implement sustainable wastewater treatment and resource recovery technologies that produce high quality water, recovery energy, and generate value-added products as the population continues to increase and WRRFs receive more stress. AnMBRs produce methane that can be recovered on-site and offer potential for down-stream nutrient recovery. This research revealed that they substantially reduce concentrations of ARGs, likely to a much greater and consistent extent than conventional activated sludge reactors.

Conflicts of interest

There are no conflicts of interest to declare.

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