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Characterization of biofoulants illustrates different membrane fouling mechanisms for aerobic and anaerobic membrane bioreactors

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Abstract

This study compares the membrane fouling mechanisms of aerobic (AeMBR) and anaerobic membrane bioreactors (AnMBR) of the same reactor configuration at similar operating conditions. Although both the AeMBR and AnMBR achieved more than 90% COD removal efficiency, the fouling mechanisms were different. Molecular weight (MW) fingerprint profiles showed that a majority of fragments in anaerobic soluble microbial products (SMP) were retained by the membrane and some fragments were present in both SMP and in soluble extracellular polymeric substances (EPS), suggesting that the physical retention of SMP components contributed to the AnMBR membrane fouling. One of the dominant fragments was comprised of glycolipoprotein (size 630-640 kD) and correlated in abundance in AnMBR-EPS with the extent of anaerobic membrane fouling. In contrast, all detected AeMBR-SMP fragments permeated through the membrane. Aerobic SMP and soluble EPS also showed very different fingerprinting profiles. A large amount of adenosine triphosphate was present in the AeMBR-EPS, suggesting that microbial activity arising from certain bacterial populations, such as unclassified Comamonadaceae and unclassified Chitinophagaceae, may play a role in aerobic membrane fouling. This study underlines the differences in fouling mechanisms between AeMBR and AnMBR systems and can be applied to facilitate the development of appropriate fouling control strategies.

Keywords: membrane fouling; molecular weight of biopolymers; soluble microbial product; extracellular polymeric substances; microbial community
1. Introduction

The membrane bioreactor (MBR) is a treatment process that couples membrane separation to the biological process for solid-liquid separation of the mixed liquor [1]. The integration of a membrane filtration unit achieves better effluent quality and de-couples the sludge retention time (SRT) from the hydraulic retention time (HRT) of the reactor, enabling higher biological oxygen demand (BOD) and chemical oxygen demand (COD) removal efficiency as compared to conventional activated sludge systems [2]. The MBR system can be applied to both aerobic (AeMBR) and anaerobic (AnMBR) treatment processes. Unlike the AeMBR, which has already found widespread application in low to medium strength municipal wastewater treatment [3], AnMBRs have been considered impractical for similar applications due to the perception that comparable transmembrane flux rates are not achievable. Nonetheless, the AnMBR has continued to gain consideration as an alternative treatment technology for municipal wastewater due to its potential advantages in reduced energy input, energy generation by methane production, and low sludge production [4-7].

Despite each MBR type’s potential advantages, membrane fouling remains the major obstacle hindering their extensive application. It has been reported that the primary contributor to membrane pore blockage in MBRs is the deposition of the dissolved fraction of activated sludge [8, 9]. This form of less-reversible membrane fouling can lead to more rigorous forms of membrane maintenance (e.g. chemical cleaning, backwashing and high cross-flow) being necessary, diminishing the economic viability of operating MBRs for municipal wastewater treatment.
The MBR, being a biological treatment process, contains microbial cells as part of the activated sludge that produces soluble microbial products (SMP) and extracellular polymeric substances (EPS). SMP was defined by Namkung and Rittmann as being comprised of utilization-associated SMP (i.e., UAP) and biomass-associated SMP (i.e., BAP) [10]. On the other hand, EPS is comprised of a matrix of polysaccharides, proteins and other macromolecules, which collectively provide adhesion, aggregation and stabilization functions for microorganisms on a membrane surface [11]. A unified theory put forward by Laspidou and Rittmann further stated that the soluble fraction of EPS is actually SMP [12]. Based on these technical definitions of SMP and EPS, much of the existing research has focused on quantification of SMP and EPS components (e.g. proteins, polysaccharides, nucleic acids and so on) and abundance ratios between those individual components in their soluble phase [13-17]. Along with the organic matter from influent wastewater, these non-settleable organic components are one of the primary culprits inhibiting MBR performance based on their role as biofoulants [15, 18, 19].

Most of the studies that focus on elucidating the role of the soluble components of EPS and SMP on membrane biofouling were carried out in AeMBRs [15, 18-20]. Little is known about the differences between AeMBR and AnMBR fouling mechanisms as there have not been any studies that have specifically examined soluble foulants in both systems under the same operating conditions. Instead, it has conventionally being presumed that the fouling mechanisms between aerobic and anaerobic systems are similar and that most of the knowledge related to the fouling mechanisms in AeMBR systems can be extrapolated to the AnMBR [21].
Furthermore, no significant attempts have been made in previous studies to investigate the specific bacterial populations present on MBR membrane foulant layers and their correlation with the occurrence of specific soluble biofoulants contributing to membrane fouling.

In this study, it is hypothesized that AeMBRs and AnMBRs may be subject to different fouling mechanisms arising from the differences in the soluble foulant components generated by the microbial communities of both systems. To address this hypothesis, an AeMBR and AnMBR were operated using a similar reactor configuration that combined an upflow attached-growth (UA) reactor with three-successive PVDF membrane filtration units that were operated for varying time periods of up to 9 weeks to reflect different extents of membrane fouling. Quantification of the protein and carbohydrate concentrations was coupled with high performance size exclusion chromatography to determine the molecular weight distributions of proteins and polysaccharides in SMP in the permeate and retentate streams of both the AeMBR and AnMBR. The same quantification procedure was also performed on the soluble EPS from biomass attached on both aerobic and anaerobic membranes. Specific microbial populations that correlated with the extent of membrane fouling and the biofoulants were further examined using high-throughput sequencing, and their bioactivity levels evaluated by measuring adenosine triphosphate (ATP) and quorum-sensing signal molecule concentrations. Ultimately, this study aims to provide a direct multifaceted comparison of the differences in biofouling mechanisms between AeMBR and AnMBR systems.
2. Materials and methods

2.1. Reactor configuration and operating conditions

To facilitate the comparison of aerobic and anaerobic systems, the same upflow-attached (UA) reactor configuration was applied to both AeMBR and AnMBR systems (Figure 1). This reactor configuration was evaluated in this study as it was previously found to have a positive role in controlling membrane fouling in MBRs [22]. Both UA reactors were filled with ceramic ring media with an average 1.5 cm diameter and length. The seed sludge in both systems originated from the same source, and comprised of camel manure and anaerobic sludge from a wastewater treatment plant in Riyadh, Saudi Arabia [23]. No oxygen or air supply was provided in the AnMBR. Aeration was supplied to the AeMBR by two air diffusion stones at the bottom of the reactor to achieve more than 2 mg/L O₂ throughout the AeMBR system. Both reactors were fed with a synthetic wastewater of 750 mg/L carbon oxygen demand (COD) [24], and operated at 35 °C and pH of 7. Hydraulic retention times (HRTs) of both reactors were 18.5 h.

Prior to connection with membrane separation units, both UA reactors were at steady-state operation and stable performance conditions. The UA reactors were individually connected to three PVDF microfiltration (MF) membrane modules in external cross-flow mode. Membrane modules were connected in series along the recirculation line with a recirculation to effluent flow ratio of 500:1. The MF membranes were JX model MF PVDF (GE Osmonics, Minnetonka, MN, USA) and had a nominal pore size of 0.3 μm. Constant flux was maintained at 6 to 8 L/m²/h (LMH) while changes in transmembrane pressure (TMP) were recorded by a pressure
gauge connected to each membrane unit. There was no sludge wasted in the anaerobic system, while 150 mL of sludge suspension was taken from the aerobic reactor per day, resulting in a sludge retention time (SRT) of 13 days. This was done to maintain a MLSS level comparable to that in the anaerobic MBR.

2.2. Soluble microbial products (SMP) and soluble extracellular polymeric substances (EPS) sampling procedure

Permeate from each membrane cassette as well as the retentate in both systems were sampled weekly throughout the course of the experiment, and were collectively referred to as soluble microbial products (SMP) in this study. All SMP samples were centrifuged at 9,400 g for 10-30 min. The supernatant was filtered with a 0.2 µm cellulose acetate membrane and stored at -20 °C prior to analyses. Analyses for SMP included determination of COD, protein and carbohydrate concentrations and the corresponding fingerprinting profiles of proteins and carbohydrates based on molecular weight (MW) fragments. Membranes were harvested from the AeMBR at time intervals of 3, 4 and 5 weeks. Membranes were harvested from the AnMBR at time intervals of 3, 6 and 9 weeks. The sampling intervals were decided based on the measured TMP and chosen to represent different extents of fouling on the membranes (Figure 2). To harvest the membrane at each time point, one of membrane cassettes was removed from the successive filtration unit, and replaced with a new membrane module to ensure constant operating conditions throughout the system. Each harvested membrane with an area of 20 cm by 2.5 cm was sectioned into three equal parts, namely inlet, mid and outlet, based on the flow direction of the wastewater stream. Soluble EPS was extracted from each section as follows: the membrane with
an area of 4 cm by 2.5 cm was cut into small strips and dispersed into two 2mL microcentrifuge tubes. 2 mL 1X PBS was added into each tube. The tubes were ultrasonicated by a QSonica Q500 Sonicator (QSonica LLC, Newton, CT, USA) for 5 min at 25% amplitude and with 2 s pulsating intervals. The membrane strips were then removed and the remaining suspension centrifuged at 9,400 g for 30 min. The supernatant after centrifugation was recovered, filtered with 0.2 μm cellulose acetate membranes and stored at -20 °C prior to analyses. The cell pellet obtained after centrifugation was also stored for DNA extraction and molecular analyses.

2.3. Protein, carbohydrate and COD quantification

Total carbohydrate concentration in the SMP and soluble EPS was determined by the phenol-sulfuric acid method with glucose as standard [25], measured in duplicate. Since the method cannot differentiate between the varying types of carbohydrates (e.g. polysaccharides, oligosaccharides and so on), measurements obtained from this method are subsequently referred to as carbohydrate concentrations in this study. Total protein concentration in the SMP and soluble EPS were detected using a protein kit from Sigma-Aldrich (TP0300, Sigma-Aldrich) with Bovine Serum Albumin (BSA) as standard and measured in triplicate. COD from all MBR reactor feeds, retentates, and permeates was determined by a HACH DR2800 Spectrophotometer (Hach, Loveland, Colorado, USA). Prior to determination of COD, the COD sample was digested via either HACH LCK 314 (15-150 mg/L) or LCK 514 COD (100-2000 mg/L) cuvette test vials (Hach-Lange, Manchester, UK) depending on the concentration to be measured.
2.4. Molecular weight (MW)-based fragment analysis of proteins and carbohydrates in SMP and soluble EPS

High Performance Liquid-Size Exclusion Chromatography (HPL-SEC) was employed to perform a MW-based fragment analysis of proteins and carbohydrates within the SMP and soluble EPS of both AeMBR and AnMBR systems. A Waters Breeze TM 2 HPLC System (Waters Chromatography, Milford, MA, USA) composed of Binary HPLC pump (Waters 1525), auto sampler (Waters 2707) and UV/Visible Detector (Waters 2489) was applied to separate the proteins in SMP and EPS. 20 µL of each sample was injected for fragment separation on a Shodex KW-802.5 column (Showa Denka America, NY) at a 20 min retention time. The mobile phase was composed of 50 mM phosphate buffer with 300 mM NaCl (pH = 7.0). All samples were detected with a mobile phase flow of 1 mL/min at room temperature. Bigger MW fragments have a shorter retention times than those with smaller MW, hence achieving fragment separation on the HPLC chromatogram. A UV detector at 280 nm was employed to detect the protein components [26]. The size calibration standard curve of $\log (\text{MW}) = - (\text{Retention Time}) * 0.4483 + 8.294$ ($R^2 = 0.995$) was generated by a series of protein standards of various MW (i.e., 1.3 kDa, 6.5 kDa, 12.4 kDa, 29 kDa, 66 kDa, 150 kDa, 200 kDa, 443 kDa, 669 kDa) and their respective retention times. A 1200 Series GPC-SEC Agilent HPLC separation system with a Refractive Index Detector (RID) was applied for monitoring the MW fingerprints of carbohydrates. The RID was heated to 35 °C prior to connection with the HPLC-SEC system. All other operational parameters were same as those used for protein separation with the exception of the mobile phase. Deionized water was used as
mobile phase as the RID detection exhibited high background noise signal when phosphate saline solution was used. The size calibration standard curve of log (MW) = -0.4786 * (Retention Time) + 8.003, with $R^2 = 0.993$, was generated by a series of Pullulan standards of various MW (i.e., 324, 1320, 6000, 21700, 48800, 113000, 210000, 366000, 708000 Dalton) and their respective retention times. Prior to performing MW size distribution of carbohydrates, the extracted sample in PBS was dialyzed with regenerated cellulose dialysis tubing of 3500 kDa molecular weight cut-off (Thermo Fisher Scientific, Waltham, MA, USA). The dialyzed soluble EPS was then analyzed based on above-mentioned procedure to examine the HPLC fingerprinting profiles of carbohydrates in soluble EPS.

2.5. Peptide sequencing of a protein fragment in AnMBR-EPS

The 633 kDa protein fragment within the soluble EPS of anaerobic membranes was further separated from other fragments for protein sequencing. The protein fragment was first separated by an Agilent 1260 Infinity Preparative Scale Purification System (Agilent Technologies, Santa Clara, CA, USA) equipped with a UV detector. The fragment was detected at an excitation wavelength of 280 nm, and was fractionated at 5-6 min retention time for every 50 μL of injected sample. Sufficient amount of the protein fragment was obtained after repeating the entire HPLC-based purification procedure more than 10 times. The eluted fragment was then dialyzed with regenerated cellulose dialysis tubing of 3500 kDa molecular weight cut-off (Thermo Fisher Scientific, Waltham, MA, USA). After four repeats of dialysis with deionized water, the sample was freeze-dried, and then submitted to KAUST Proteomics Core lab for protein digestion and peptide sequencing.
2.6. Adenosine triphosphate (ATP) and autoinducer-2 (AI-2) quantification

Bioactivity on the membranes was measured based on ATP and AI-2 concentrations. Adenosine triphosphate (ATP) is considered to have a core role in respiration and metabolism, and is the most important energy supplier in many enzymatic reactions [27]. Similarly, autoinducer-2 (AI-2) is a furanosyl borate diester signal molecule secreted and responded to by many Gram-negative and Gram-positive bacteria [28] during biofilm development [29, 30].

ATP was extracted from a 0.5 cm by 2.5 cm membrane segment. The membrane was placed into a microcentrifuge tube containing 2 mL of deionized water, and ultrasonicated for 1 min at 25% amplitude. After a brief vortex, the membrane was removed from the tube, and the suspension was further ultrasonicated for 1 to 2 min. ATP content in the suspension was quantified with the Celsis amplified-ATP reagent kit on an Advance luminometer (Celsis, Westminster, London, UK) with deionized water as a negative control. All samples were measured in triplicate.

The concentration of AI-2 present in the biocake from the harvested membranes was estimated by a protocol previously described [31]. Briefly, the AI-2 indicator *Vibrio harveyi* ATCC® strain BB170 grew overnight with autoinducer bioassay (AB) medium. After overnight growth, the AI-2 culture was diluted 1:5000 with fresh AB medium. 20 μL of extracted soluble EPS to be tested was placed into 96-well solid white microplate prior to addition of 180 μL of diluted AI-2 reporter. The 96-well plate containing the samples was incubated in dark on a 150 rpm shaker incubator platform at 30 ºC. The bioluminescent intensity was detected with the
Infinite M200 PRO microplate reader over time (Tecan, Männedorf, Switzerland).

Sterile 1X PBS that was used to extract for the soluble EPS from membranes was also measured as a negative control. Varying concentrations of (S)-4,5-dihydroxy-2,3-pentanedione (DPD) (Omm Scientific, Dallas, TX, USA) diluted in deionized water were used as standard. The bioluminescent intensities from the samples, standards and negative control increased with incubation. However, the increment rates for the bioluminescent intensities in samples and standards were higher than that for the negative control. The intensities measured for each respective standard were determined after 5 h of incubation, and normalized against the intensities measured from the negative control. This ratio was used to generate a calibration curve that plotted the intensity ratio versus the concentration of (S)-4,5-dihydroxy-2,3-pentanedione (DPD). The calibration curve was represented by the following equation:

\[ y = 19.844 \times x + 311.62 \text{, with } R^2 = 0.994 \]

where \( y \) denotes intensity ratio, and \( x \) denotes the concentration of AI-2 in DPD equivalence. Similarly, the intensities measured for each respective sample were determined after 5 h of incubation and normalized against that from the negative control. The ratio was then substituted into the equation to determine the AI-2 concentration present in that sample. AI-2 was measured in quadruplicates per sample.

2.7. DNA extraction and barcoded amplification of 16S rRNA genes

The cell pellets obtained based on procedures described in Section 2.2 were extracted for their genomic DNA using the UltraClean® Soil DNA Isolation Kit
(MoBio Laboratories, Carlsbad, USA) with slight modifications [32]. PCR amplification for the 16S rRNA genes was performed with barcoded forward primer 515F (5’-GTGYCAGCMGCCGCGGTA-3’) and reverse primer 909R (5’-CCCGYCAATTTCMTTTRAGT-3’) based on thermal cycling conditions previously described [33]. All amplicons were of the correct size of ~450 bp and all negative controls had no detectable amplification. Gel-purification of PCR amplicons was performed with Wizard SV Gel and PCR Clean-up system (Promega, Madison, USA). The amount of DNA in the purified amplicons was measured by Qubit® broad range dsDNA assay (Invitrogen, Carlsbad, CA, USA). Equal amounts of the samples were mixed together and submitted to Macrogen Korea for Ion Torrent sequencing (Life Technologies, Carlsbad, USA) on 316 chips.

2.8. Ion Torrent sequencing data analysis

The sequencing data from Ion Torrent platforms were first sorted by the KAUST Bioinformatics Team based on a Phred score of >20. All primers, barcodes and adapters as well as the sequences with <350 nt were removed. Sequences that passed the initial quality check were then evaluated for presence of chimeric sequences using UCHIME [34]. The remaining chimera-free sequences were then analyzed for their taxonomic affiliations at a 95% confidence level using the RDP Classifier [35]. Chimera-free sequences were also collated with an in-house written Perl script, and then sorted for unique operational taxonomic units (OTUs) at 97% 16S rRNA gene similarity using CD-Hit [36]. OTUs were blasted against the NCBI 16S rRNA gene nucleotide database using blastn to check for their closest matching identities.
2.9. Statistical analysis

The extent of similarities between the microbial communities attached to the aerobic and anaerobic MBR membranes were represented on a non-metric multidimensional scaling (nMDS) plot. To generate the nMDS plot, the relative abundances of the bacterial and archaeal genera were calculated, collated and then square-root transformed. The square-root transformed dataset was then computed for Bray-Curtis similarities and plotted on the nMDS. Vectors illustrated on the nMDS exhibited a Pearson correlation of > 0.9 to the sample distribution. All measured ATP, AI-2, protein and carbohydrate concentration data were also collated, log-transformed and normalized prior to principal component analysis (PCA). The four measured parameters (i.e., ATP, AI-2, protein and carbohydrate) were displayed as vectors on the PCA to illustrate the contribution of each parameter in the distribution of samples on the plot. All statistical analysis described in this section was performed on Primer-E v7 software [37].

2.10. Nucleotide sequence accession numbers

All high-throughput sequencing files were deposited in the Short Read Archive (SRA) of the European Nucleotide Archive (ENA) under study accession number PRJEB9458.
3. Results

3.1. AeMBR and AnMBR operational performances

The AeMBR and AnMBR systems achieved an average COD removal efficiency of 94% and 90%, respectively (Figure S1). The TMP in the AeMBR system increased to 52 kPa at the end of 3 weeks of continuous operation (Figure 2), while the TMP measured for the membranes connected to the AeMBR at the end of 4 weeks and 5 weeks was 88 kPa for both. For the AnMBR system, the TMP gradually increased from 0 kPa to 23 kPa and 62 kPa at the end of 3, 6 and 9 weeks, respectively (Figure 2). Membranes were harvested from the AeMBR at time intervals of 3 weeks, 4 weeks and 5 weeks, and at time intervals of 3 weeks, 6 weeks and 9 weeks for the AnMBR to represent comparable levels of fouling for both systems.

3.2. Protein and carbohydrate concentration in retentate and permeate streams of AeMBR and AnMBR

The amount of total protein and carbohydrate in retentate from the AeMBR was on average 7.2 ± 1.3 mg/L and 21.8 ± 2.7 mg/L, respectively, throughout operation (Figure 3A). The amount of protein and carbohydrate in AeMBR retentate was lower than that detected in the AnMBR retentate. The average amount of protein in AnMBR retentate was 94.1 ± 24.7 mg/L while the amount of carbohydrate in the AnMBR retentate exhibited a gradual increase from 22.6 ± 2.9 mg/L during days 0 to 40 of operation to 51.2 ± 7.5 mg/L from day 40 onwards (Figure 3B). The amount of total carbohydrate in AeMBR permeate was 76.5% of that detected in the retentate, and was significantly lower in the permeate (t-test, P < 0.05). However, the amount of
protein in permeate and retentate of the AeMBR was not significantly different (t-test, P = 0.65). In contrast, up to 89.2% of protein and 91.3% of carbohydrate were removed from the AnMBR retentate, and a significant reduction for both protein and carbohydrate was observed in the AnMBR permeate compared to retentate (t-test, P < 0.05).

3.3. Protein and carbohydrate concentration in soluble EPS of attached biomass

The amount of carbohydrate measured in soluble EPS extracted from different portions of the 3-week aerobic membranes ranged from 70-102 mg/L. There was a comparatively higher concentration of carbohydrate than protein in the soluble EPS of aerobic membranes. Furthermore, the amount of carbohydrate increased to 102-126 mg/L and 173-197 mg/L in the 4-week and 5-week aerobic membranes, respectively (Figure 3C). These carbohydrate concentrations detected in both 4-week and 5-week membranes were significantly higher than that detected in 3-week membranes (t-test, P < 0.05). The average protein content in 3-week and 4-week aerobic membranes was 40.5 ± 5.2 and 36.1 ± 6.0 mg/L, respectively, and significantly increased (t-test, P < 0.05) to 67-74 mg/L in the 5-week aerobic membranes (Figure 3C). In contrast, more protein than carbohydrate was present in the soluble EPS of anaerobic membranes (Figure 3D). There was a significant increase in both carbohydrate and protein concentrations detected in the soluble EPS of 6-week and 9-week anaerobic membranes compared to the 3-week membrane (t-test, P <0.05). A slight increase of carbohydrate was also observed between the 6-week and 9-week anaerobic membranes (t-test, P = 0.09) but no significant difference was observed for protein (t-test, P = 0.46).
3.4. Molecular Weight (MW)-based fingerprinting profiles of SMP and soluble EPS in AeMBR

A single-peak fragment of 0.9 kDa was detected in the retentate and permeate throughout the course of AeMBR operation (Figure 4A). The peak area for this fragment was similar for both AeMBR retentate and permeate, indicating that this fragment had completely permeated through the PVDF MF membrane. Unlike the MW-based fingerprinting profile of AeMBR SMP, the protein components present in the soluble EPS of aerobic membranes had a bimodal distribution of the MW fragments (Figure 4C). A single-peak fragment of 487.8 ± 43.9 kDa was observed in the 3 and 4 weeks membranes, in contrast to two peaks on the 5 weeks aerobic membrane that were comprised of the original fragment and an additional 837.7 kDa fragment. In addition, fragments of small MW ranging from 0.01-4 kDa were observed on all aerobic MBRs but were not detected in the SMP of AeMBR (Figure 4C). The carbohydrates detected in both SMP and soluble EPS of AeMBR shared different fingerprinting profiles (Figure 4B and 4D). A single peak associated with an average MW of 180.1 ± 5.5 kDa was also observed in the SMP (Figure 4B) but not in the soluble EPS fraction (Figure 4D). Instead, two main fragments of MW 3396.6 ± 64.4 kDa kDa and 94.9 ± 9.7 kDa were the main polysaccharide compounds detected on the soluble EPS of aerobic membranes.

3.5. Molecular weight (MW)-based fingerprinting profiles of SMP and soluble EPS in AnMBR

SMP and soluble EPS of the AnMBR exhibited different fingerprinting profiles compared to those of the AeMBR (Figure 5). The pattern of MW fingerprints
of protein in SMP in the AnMBR was similar to those in EPS of anaerobic membranes (Figure 5A and 5C). Both had a single peak and a bigger MW size ranging from 630-640 kD. This fragment in SMP was completely rejected by the MF membrane. Given that this specific fragment was ubiquitously detected in the AnMBR system, the fragment was further extracted for peptide sequencing. Out of the 80 total peptide sequences obtained, 57.5% of the sequences were identified to be vitellogenin, which is a glycolipoprotein (Supplementary Material).

Among the three detected polysaccharide fractions in the SMP of AnMBR, only the first eluted fragment with the biggest MW (average 2526.3 ± 483.4 kDa) was completely rejected by the MF membrane (Figure 5B). However, even though the 2526.3 kDa polysaccharide fragment was rejected by the anaerobic membranes, this fragment was not detected in the soluble EPS of the membranes (Figure 5D). The majority of the fragment with a MW size of 0.17 kDa was retained in the reactor. The fragment with a size of 158.5 ± 6.5 kDa was low in abundance in the SMP but increased in abundance within the soluble EPS of the anaerobic membranes with time, and constituted one of the two peaks detected (Figure 5D).

3.6. ATP and AI-2 in AeMBR and AnMBR

The amount of ATP measured in the biocake layer on aerobic membranes was at least 3-fold higher than that measured on anaerobic membranes (Figure 6A and 6B). The ATP content for all three sessions in AnMBR membranes was less than 100 pmol/cm², while 300 to 1500 pmol/cm² ATP was measured in the biocake on the 3-week to 5-week AeMBR membranes. The amount of ATP on the aerobic membranes increased with time but there was no significant difference between the amount of
ATP on the 5-week aerobic membrane compared to 3-week aerobic membrane (t-test, P = 0.07). There was also no significant change of AI-2 content among the different aerobic membranes (t-test, P > 0.25). The average AI-2 in the biocake of all aerobic membranes was 0.2 nmol/cm², while a temporal increase in AI-2 content was observed for the anaerobic membranes. The AI-2 concentration in the 3-week anaerobic membrane was less than 0.2 nmol/cm² while the average amount of AI-2 in the 6-week anaerobic membrane was 0.4 nmol/cm². The average amount of AI-2 further increased in the 9-week anaerobic membrane to 0.83 nmol/cm², and was significantly higher than that detected from the 3-week anaerobic membrane (t-test, P = 0.03).

3.7. Microbial community on AeMBR and AnMBR membranes

The aerobic and anaerobic membranes were evaluated for their extent of dissimilarities based on the measured concentrations of ATP, AI-2, carbohydrate and protein in the biocake (Figure 7A), as well as relative abundance and occurrence of bacterial populations attached to the membranes (Figure 7B). Based on both evaluation approaches, it was observed that the biocake properties and the microbial communities of the aerobic membranes were distinctly different from those of the anaerobic membranes. To illustrate, aerobic and anaerobic samples were spatially clustered apart on the PCA plot along the PC1 axis, which accounted for 54.9% of the total variance. The main vectors that accounted for the spatial distribution of the aerobic samples on the PCA were the concentrations of ATP and carbohydrate. In particular, the amount of ATP and carbohydrate in the biocake of aerobic membranes was higher than that detected in the anaerobic membranes. A similar distribution of
the microbial communities attached to aerobic and anaerobic membranes was observed on the MDS plot. Unclassified Syntrophaceae was one of the two bacterial populations that was highly correlated (ρ = 0.9) with the positioning of 9-week anaerobic samples on the nMDS. Further evaluation showed that an OTU with a best-matched identity to *Smithella propionica* (i.e., a syntroph) increased in its relative abundance from 0.27% on a 3-week anaerobic membrane to 1.2% on a 9-week anaerobic membrane. In contrast, the abundance of unclassified Comamonadaceae (e.g. *Comamonas testosteroni*) and unclassified Chitinophagaceae (e.g. *Ferruginibacter lapsinanis, Terrimonas rubra*) highly correlated with the spatial distribution of the aerobic membranes. Collectively, these three OTUs accounted for up to 13.6% of the total microbial community attached to the most fouled 5-week aerobic membrane (Table 1). The concentration of ATP in the biocake was further identified as the single variable that best linked the spatial distribution of samples on the PCA plot with those on the MDS plot (correlation value = 0.698, p = 0.001).

4. Discussion

In recent years, there has been increasing interest in AnMBRs due to the various advantages associated with the coupling of membrane separation to anaerobic fermentation processes. As is the case with all membrane bioreactors, membrane fouling remains a significant obstacle limiting their widespread use. It has conventionally being presumed that the fouling mechanisms between aerobic and anaerobic systems are similar, at least at the macro scale [21]. Unlike past studies, which examined fouling rates in CSTR-type AeMBRs, this study evaluated the differences in the fouling rates of AeMBR and AnMBR operated based on an
attached-growth reactor configuration. This is built upon past observations which reported a lower MLSS resulting from attached-growth reactors playing a positive role in controlling membrane fouling in AeMBR systems [22]. However, our results showed that when an AeMBR was evaluated against an AnMBR system with the same reactor configuration, the fouling rates and mechanisms experienced by their respective membranes were very different, despite both systems achieving similar COD removal efficiencies.

Compared to the AeMBR, more gradual membrane fouling was observed in the AnMBR. TMP in the AeMBR increased to over 90 kPa after 4 weeks of operation, while TMP in the AnMBR did not reach similar levels until after 9 weeks of operation (Figure 2). Correlating with the TMP increment, aerobic membranes were more severely fouled after 4 and 5 weeks of operation as compared to the 3-week membrane, with significantly higher protein and carbohydrate contents in the 4-week and 5-week aerobic membranes than the 3-week membrane. Similarly, the 6-week and 9-week anaerobic membranes exhibited significantly higher protein and carbohydrate contents than that of the membrane removed after 3 weeks of operation (Figure 3).

The slower fouling rates in the AnMBR system may be due to a lower solids deposition rate on the anaerobic membrane surface than the aerobic membrane [5]. This was in spite of the overall SMP levels in the AnMBR being almost 5 times higher than in the AeMBR reactor (Figure 3), and that a high abundance of SMP had been shown previously to correlate to membrane fouling [38]. A previous study comparing AeMBR and AnMBR systems reported SMP levels that were up to 6
times higher in the AnMBR, and that the AnMBR experienced more rapid membrane fouling [39]. However, our findings did not show any correlation between the abundance of SMP and extent of membrane fouling. This implies that, although SMP is known to contribute significantly to membrane fouling rates, absolute quantification of SMP may not be as important as its specific components.

Conversely, soluble EPS (i.e., SMP associated with the attached biomass) examined in this study had levels that correlated positively with the membrane fouling rates, both in terms of raw quantification (Figure 3) and specific protein and polysaccharide fractionation profiles (Figure 4 and 5).

Based on fractionation profiles, it was observed that the AnMBR produced bigger molecular weight fractions in its SMP than the aerobic process. This, in turn, resulted in the majority of the proteins and carbohydrates being retained in the anaerobic retentate. Protein subsequently accumulated on the fouled anaerobic membranes, and was comprised of a single fraction with a MW size of 633 kDa (Figure 5C). This protein fragment was further determined from peptide sequencing to be made up by glycolipoprotein (Supplementary Material). It is to be noted that although the MW 633 kDa was less than the physical pore size of 0.3 μm of PVDF membrane, HPLC analysis indicated that this fragment was totally rejected by the AnMBR membrane. A similar occurrence was also observed in a study by Jang et al., where it was found that a MF membrane with pore size of 0.4 μm could retain biopolymers of MW as low as 40 kDa [40]. Electrostatic repulsion may be a factor in causing the rejection of biopolymers of sizes smaller than the physical pore size of membranes [41]. Another contributing factor could be related to the deposited
biofoulants on the membrane surface, as seen in studies where the biofilm layer acted as a secondary self-forming dynamic membrane that modified the membrane’s rejection properties [42-45].

This phenomenon would also likely explain the partial rejection of a polysaccharide compound of 0.17 kDa MW that was also found in the AnMBR-SMP, implying that in certain instances properties such as particle charge, hydrophobicity, and aggregation completely supersede physical rejection by the porous membrane. Although the 0.17 kDa compound was rejected by the anaerobic membrane, an associated fragment of the same size was not observed within the soluble EPS of the anaerobic membrane (Figure 5). Instead, the 158 kDa polysaccharide fragment, which was observed in low abundances in the SMP, built up within the soluble EPS associated with the attached biomass on anaerobic membranes. In addition, a polysaccharide component of 4308 kDa MW increased in abundance as the anaerobic membrane became increasingly fouled with time. This compound was not observed in the SMP and could not be easily traced. It is possible that this large size MW fragment may be comprised of a complex of inorganic and organic foulants that combined to form a fragment of this size. Past studies have shown that anaerobic MBRs are more prone to inorganic fouling due to the high ammonium and phosphate content within the reactor [46]. Inorganic foulants such as struvite, otherwise also known as magnesium ammonium phosphate hexahydrate (MgNH₄PO₄·6H₂O), were thought to contribute to polysaccharide accumulation due to the free positive ions binding to the negative charges of OH⁻ and COO⁻ that may be present among extracellular polysaccharide molecules [47].
Unlike the anaerobic membrane, the aerobic membrane displayed SMP MW profiles that were very different from those of the soluble EPS of the fouled membranes in terms of both proteins and carbohydrates. To illustrate, there were no apparent soluble EPS fragments that could be traced to SMP counterparts in the AeMBR reactor (Figure 4). The sole protein and polysaccharide fragments detected in the SMP were completely permeated through the membrane in the AeMBR, suggesting that the aerobic sludge produced SMP fragments with physical properties that reduced interactions with the MF membranes used.

Results of ATP quantification of the biofilms further suggest that the mechanism for biofouling in the AeMBR system could be completely different from that of the AnMBR system. Instead of SMP compound deposition on the membrane like what had been observed for the anaerobic membrane, it appeared that activity of the localized microbial communities and their EPS production could be responsible for the majority of the soluble EPS on the AeMBR membrane surface. To illustrate, ATP levels per surface area in the AeMBR were significantly higher (> 30 fold) than those in the AnMBR (Figure 6A). This relatively higher ATP content in the AeMBR membrane biofilms demonstrates that microbial activity is also likely much higher [48, 49]. A large fraction of the EPS on the AeMBR membranes could be attributed to this microbial activity as higher biomass activity is generally associated with higher specific soluble EPS production rates [50]. Considering these observations, in combination with the previously discussed protein and polysaccharide fragment analyses that showed little similarity between the soluble EPS and SMP profiles for
the AeMBR, it seems likely that the main contributors to membrane fouling in the AeMBR arise from microbial activity on the membranes.

An evaluation of the microbial communities showed that the abundance of unclassified Comamonadaceae (e.g. *Comamonas testosteroni*) and unclassified Chitinophagaceae (e.g. *Ferruginibacter lapsinanis, Terrimonas rubra*) were the main bacterial vectors that resulted in the separation of the aerobic membrane samples from those of the anaerobic membranes in MDS analysis (Figure 7B). Collectively, the three OTUs most likely representing these bacterial groups accounted for up to 13.6% of the total microbial community attached to the most seriously fouled 5-week aerobic membrane (Table 1), suggesting that they may play a role in the aerobic membrane fouling process. Past studies have reported the presence of *Comamonas* spp. in the biofilms of aerobic membranes [51]. Furthermore, a significantly higher abundance of *Comamonas* spp. and *Ferruginibacter* spp. have been observed in aerobic sludge samples taken from an MBR fed with influent wastewater of a higher COD/N ratio than ones taken from an MBR fed with a lower COD/N ratio [20]. These observations suggest that these two bacterial populations could preferentially attach to aerobic membrane surfaces, where dense organic matter is likely to accumulate. In a related study, clone library analysis similarly revealed that Betaproteobacteria (*Comamonas*) and Bacteroidetes (*Ferruginibacter*) were the dominant groups in aerobic MBR membrane biofilms [52]. Given the positive correlation between ATP concentrations and the abundance of *Comamonas* spp. and Chitinophagaceae (*p* = 0.698, *p* = 0.001), it is likely that these two bacterial
populations were contributing to the bioactivity on the aerobic membrane surfaces, resulting in an increase in ATP levels.

This study did not look into the insoluble portions of membrane biofilm EPS, which generally account for a relatively larger portion of the EPS matrix than the soluble fractions. Neither did the study comprehensively characterize the bound EPS from the biofilm cake layers and suspended mixed liquor in the fouling of both MBR types. Furthermore, it is likely that the extraction method used in this study did not recover all of the protein and polysaccharide components present in the SMP and soluble EPS. In a study by Aquino and Stuckey that used liquid-liquid extraction protocols with various organic solvents, up to 20 MW fragments ranging from 10 kDa to over 300 kDa in the soluble microbial products of AnMBR effluent were identified [53]. Further studies on the insoluble EPS and tightly-bound EPS extracted from different protocols may develop new and improved insight on the different membrane fouling mechanisms. Despite these limitations, this study revealed significant differences between aerobic and anaerobic MBRs in the fractions of SMP and soluble EPS measured that shed light on the mechanisms potentially responsible for fouling.

Given the observed differences in the fouling mechanisms of both systems, different membrane fouling control approaches can be recommended accordingly. Since the mechanisms most responsible for membrane fouling in AnMBRs are SMP deposition, as shown in this study, and cake formation [54], conventional chemical or physical cleaning methods could be effective AnMBR fouling control techniques. For example, addition of FeCl₃ reduced the accumulation of protein and carbohydrate on AnMBR membrane surfaces, leading to lower fouling rates. It was proposed that
FeCl₃ acts as a coagulant for colloidal and soluble substances, and prevented the development of a strongly-attached cake layer on the anaerobic membranes [55]. Recent studies have also utilized granular activated carbon (GAC) to reduce AnMBR membrane fouling by providing mechanical scouring to the membrane surfaces [56-58] and have demonstrated that the AnMBR can be operated continuously for up to 6 months at 6-11 LMH with no significant increases in TMP [57]. Similarly, mechanical air scouring methods have been used extensively for mitigation of membrane fouling in aerobic MBRs [59, 60]. To compliment these methods, biological-based approaches can be used to further prolong membrane life in AeMBRs. Examples of such approaches include quenching of acyl-homoserine lactone (AHL) and AI-2 signal molecules to inhibit cell-to-cell communication within the biofilm matrix and addition of enzymes to break down proteins and polysaccharides of the biofilm [61-65]. One study showed that by encapsulating Rhodococcus sp. or recombinant E. coli in microbial vessels as a live source of quorum quenching inside the MBR, the emitted quorum quenchers were able to delay the occurrence of maximum TMP by up to a day as compared to the control AeMBR [61, 64].

5. Conclusion

Although high COD removal efficiency was achieved for both aerobic and anaerobic MBR systems, faster membrane fouling was observed in the AeMBR. The MW fingerprints of SMP and EPS in the AnMBR showed that a majority of EPS fragments could be derived from the retained SMP fractions, implying that the major contributor to AnMBR membrane fouling was physical retention of SMP components
by the microfiltration membrane. In the case of the AeMBR, all SMP fragments were completely permeated through the membrane and EPS did not correlate with SMP profiles. Furthermore, significantly higher levels of ATP were present in AeMBR membrane biofilms than in those of the AnMBR, suggesting that a larger fraction of the EPS on the AeMBR membrane was a product of microbial activity arising from certain bacterial populations on the membrane surfaces, such as *Comamonas* and unclassified Chitinophagaceae. This study shows that potentially very different fouling mechanisms controlled AeMBR and AnMBR membrane biofouling and reiterates the importance of a multifaceted approach in studying membrane fouling and developing control strategies for different MBR systems.

6. **Acknowledgements**

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References


Figure captions

Figure 1. Operational setup for both aerobic MBR and anaerobic MBR. An attached-growth reactor configuration was used and the MBR systems were operated at 18.5 h hydraulic retention time (HRT) and with flux of 6-8 L/m²/h through the microfiltration (MF) membrane.

Figure 2. Transmembrane pressure (TMP) measured throughout the operation of AeMBR and AnMBR. Aerobic membranes were harvested after 3, 4 and 5 weeks of operation at different final TMP. Anaerobic membranes were harvested after 3, 6 and 9 weeks of operation at different final TMP.

Figure 3. The concentration of proteins, PN, and carbohydrates, CH, in (A) retentate and permeate of aerobic MBR, AeMBR (B) retentate and permeate of anaerobic MBR, AnMBR (C) soluble EPS from different sections of aerobic membranes (D) soluble EPS from different sections of anaerobic membranes.

Figure 4. Molecular weight fingerprinting profiles of (A) proteins in SMP of aerobic MBR retentate and permeate, (B) carbohydrates in SMP of aerobic MBR retentate and permeate (C) proteins in soluble EPS of aerobic membranes, and (D) carbohydrates in soluble EPS of aerobic membranes.

Figure 5. Molecular weight fingerprinting profiles of (A) proteins in SMP of anaerobic MBR retentate and permeate, (B) carbohydrates in SMP of anaerobic MBR retentate and permeate (C) proteins in soluble EPS of anaerobic membranes, and (D) carbohydrates in soluble EPS of anaerobic membranes.

Figure 6. Bioactivity measured in the biocake on aerobic and anaerobic membranes. Evaluation was made based on (A) ATP content of aerobic membranes, (B) ATP content of anaerobic membranes, (C) AI-2 content of aerobic membranes, and (D) AI-2 content of anaerobic membranes.

Figure 7. Ordination analysis of aerobic and anaerobic samples. (A) Principal component analysis of the ATP, AI-2, protein PN and carbohydrate CH contents, (B) non-metric multidimensional scaling plot of microbial communities.
Table 1. Operational taxonomic units (OTUs) that were increased in relative abundance with time on the (A) aerobic membranes, and (B) anaerobic membranes. OTUs were blast for their best-matched identities based on partial 16S rRNA gene sequences.

### A) Aerobic membrane

<table>
<thead>
<tr>
<th>Best-matched species</th>
<th>3-week Avg. (%)</th>
<th>4-week Avg. (%)</th>
<th>5-week Avg. (%)</th>
<th>Identity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Prolixibacter bellariivorans</em></td>
<td>0.52</td>
<td>2.68</td>
<td>2.14</td>
<td>86</td>
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<tr>
<td><em>Ferruginibacter lapsinanis</em></td>
<td>0.44</td>
<td>3.93</td>
<td>7.46</td>
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<tr>
<td><em>Cloacibacterium rupense</em></td>
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<td>0.57</td>
<td>0.59</td>
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<tr>
<td><em>Meiothermus granaticius</em></td>
<td>1.13</td>
<td>1.88</td>
<td>3.90</td>
<td>93</td>
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<tr>
<td><em>Terrimonas rubra</em></td>
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<td>0.88</td>
<td>0.86</td>
<td>98</td>
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<td><em>Ferruginibacter lapsinanis</em></td>
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</tr>
<tr>
<td><em>Melioribacter roseus</em></td>
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<td>1.42</td>
<td>1.12</td>
<td>82</td>
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<tr>
<td><em>Comamonas testosteroni</em></td>
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<td>2.05</td>
<td>1.53</td>
<td>98</td>
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<td><em>Zoogloea resiniphila</em></td>
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<td>0.49</td>
<td>0.48</td>
<td>97</td>
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<tr>
<td><em>Candidatus Nitrospira defluvi</em></td>
<td>0.074</td>
<td>0.96</td>
<td>1.09</td>
<td>94</td>
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</table>

### B) Anaerobic membrane

<table>
<thead>
<tr>
<th>Best-matched species</th>
<th>3-week Avg. (%)</th>
<th>6-week Avg. (%)</th>
<th>9-week Avg. (%)</th>
<th>Identity (%)</th>
</tr>
</thead>
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<tr>
<td><em>Smithella propionica</em></td>
<td>0.27</td>
<td>0.57</td>
<td>1.22</td>
<td>96</td>
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<tr>
<td><em>Melioribacter roseus</em></td>
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<td><em>Ignavibacterium album</em></td>
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<td>0.20</td>
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<tr>
<td><em>Cloacibacterium haliotis</em></td>
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<td>0.42</td>
<td>0.50</td>
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<tr>
<td><em>Melioribacter roseus</em></td>
<td>3.04</td>
<td>4.17</td>
<td>10.3</td>
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<td><em>Methanothermobacter tenebrarum</em></td>
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<td>0.54</td>
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<tr>
<td><em>Oceanibaculum indicum</em></td>
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<td>0.65</td>
<td>87</td>
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<tr>
<td><em>Halothiobacillus Neapolitanus</em></td>
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<td><em>Melioribacter roseus</em></td>
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<tr>
<td><em>Cloacibacterium haliotis</em></td>
<td>0.64</td>
<td>0.98</td>
<td>1.29</td>
<td>96</td>
</tr>
</tbody>
</table>
Fig. 1
Fig. 2

![Graph showing TMP (kPa) vs Duration (d) for different MBR treatments.](image-url)
Fig. 3
Fig. 4
Fig. 5

(A) Day 0: 644.5±12.0
   Day 14:
   Day 27:
   Day 42:
   Day 56:
   Day 63:

(B) Day 0: 2526.3±483.4
   Day 14:
   Day 27:
   Day 42:
   Day 56:
   Day 63:

(C) 3-week: 633.3±49.2
     6-week:
     9-week:

(D) 3-week: 157.0±44.1
     6-week:
     9-week:
Fig. 6
Fig. 7

(A)

(B)

Legend
- Aerobic MBR – 2-week
- Aerobic MBR – 4-week
- Aerobic MBR – 5-week
- Anaerobic MBR – 3-week
- Anaerobic MBR – 6-week
- Anaerobic MBR – 9-week

2D Stress: 0.05

Unclassified Comamonadaceae
Unclassified Bacteroidetes
Unclassified Chitinophagaceae
Unclassified Bacteria
Unclassified Syntrophaceae
Highlights
- Aerobic and anaerobic MBR with same configuration achieved > 90% COD removal
- Different membrane fouling mechanisms in aerobic and anaerobic MBR
- SMP and EPS protein profiles similar in anaerobic but not in aerobic system
- Physical retention of SMP contributed to anaerobic membrane fouling
- Microbial activity predominant in aerobic membrane fouling