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Cite this: Environ. Sci.: Water Res. Technol., 2019, 5, 482 Emerging investigator series: photocatalysis for MBR effluent post-treatment: assessing the effects of effluent organic matter characteristics[†]

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Dissolved organic matter (DOM) poses a serious challenge to applied photocatalysis. Membranes may offer a promising synergistic opportunity to enable efficient photocatalysis in the presence of DOM. Membrane bioreactor (MBR) effluent from a municipal treatment plant was studied to elucidate the effects of filtration and organic matter composition on photocatalysis. Effluent samples were collected from MBR units during routine operation and before/after chemical cleaning. Additional DOM samples from the bulk supernatant were separated into colloidal, hydrophobic and transphilic fractions, providing a novel examination of the inhibition potential of DOM. These DOM fractions and the effluent organic matter (EfOM) samples were then characterized utilizing three-dimensional excitation-emission matrix (3DEEM) fluorescence spectroscopy and assayed for their potential to inhibit TiO2-mediated photocatalytic degradation of a probe compound, para-chlorobenzoic acid (pCBA). The colloidal fraction of DOM was found to exert the strongest inhibition, followed by the transphilic, then the hydrophobic fractions; at 5 mgC L^{-1} , these fractions reduced the photodegradation rates by approximately 75%, 27%, and 17%, respectively. Of the effluent samples, EfOM from the recently-cleaned membrane caused the greatest inhibition of photocatalysis (~100% reduction at 0.5 to 2.0 mgC L^{-1} , whereas the effluent from the fouled membrane provided the least inhibition (~33% reduction at 2.0 mgC L⁻¹). The 3DEEM analysis predicted inhibitory action of both DOM and EfOM, based on total fluorescence volumes. Results here demonstrate the prospective utility of combining membrane technologies with photocatalytic processes

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Water impact

This work addresses critical challenges to the application of photooxidation processes to waters with significant organic matter content. Fractionation and characterization tools were used to better understand the effects of organic matter on these oxidation pathways. The results guide wastewater reuse efforts by identifying critical impediments to photooxidation processes in wastewater and demonstrating the potential use of membranes to overcome the challenges.

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1. Introduction

As the human population continues to grow, careful utilization of natural resources becomes increasingly more important. Water usage, and particularly reuse, is a critical topic for many communities.^{1,2} The development of membrane bioreactor (MBR) technology has been an important step towards wastewater reuse, given substantial advantages over conventional activated sludge systems in terms of improved efficiency and effluent quality.³ Despite these benefits, which stem from the use of a physical barrier (membranes), several types of contaminants can pass through the membranes and pose significant health risks upon release of the effluent into the environment. Indeed, viruses have been found in the effluents of state-of-the-art MBR treatment plants.^{4,5} In

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addition, pharmaceuticals^{6,7} and, more recently, antibiotic resistant genes have drawn attention as significant concerns posing environmental and health risks.⁸ Thus, MBR systems require a disinfection step post-filtration to provide a safeguard. Although UV disinfection can be used in lieu of chlorination, and thereby avoid the necessity for an added dechlorination step, there are disadvantages: UV treatment does not significantly degrade antibiotic resistant genes⁸ and many pollutants are recalcitrant to UV at commonly applied UV doses.⁹

Given these post-filtration challenges, an integration of innovative technologies could provide the key functionality to eliminate the remaining hazards from MBR effluent. In particular, photocatalytic materials applied in conjunction with existing UV dosing systems could produce reactive oxygen to destroy these contaminants via advanced oxidation processes.9-12 Germicidal UV radiation, a subset of the spectrum of short-wavelength UV light, often called UVC,^{13,14} can be enhanced by the addition of photocatalytic processes to promote the production of reactive oxygen species (ROS). These ROS are known to be particularly effective at inactivating viruses compared to bacteria which have protective cell membranes.¹⁵⁻¹⁸ For example, the inactivation kinetics of Escherichia coli by hydroxyl radicals ('OH) or by singlet oxygen $({}^{1}O_{2})$ have been shown to have a lag-phase where the cell membrane protects the bacteria's intracellular components against ROS attack,^{15,16,18} whereas the genetic and essential components of viruses, such as MS2 bacteriophage, have very little protection and therefore no delay in their inactivation kinetics.^{15,16} A combined UVC-photocatalytic system is a plausible conception that could serve as an advanced oxidation process to oxidize pharmaceutical compounds in addition to providing disinfection activity.¹⁰

Application of photocatalytic processes to natural or waste waters faces a significant challenge in the form of non-target organic matter interferences. In the case of MBRs, effluent organic matter (EfOM) contains a variety of molecules that are known to quench hydroxyl radicals ('OH),¹⁹ which are generally the most important ROS in any advanced oxidation process-including TiO2 mediated photocatalysis. MBR EfOM is a complex mixture of organic molecules such as proteins, polysaccharides, humic substances and nucleic acids.²⁰⁻²³ These molecules originate primarily from microbial activity (soluble microbial products, SMPs), produced during secondary biological treatment (via suspended or attached growth processes), and are typically found at concentrations ranging from 3 to 25 mgC L^{-1} $^{21,22,24-27}$ EfOM can interfere with photocatalytic treatment through different inhibitory mechanisms. First, EfOM absorbs significant amounts of UV light, limiting the amount of photons available for catalyst excitation.²⁸ Second, EfOM quenches ROS, preventing reactions with the target compounds or microorganisms.^{29,30} This competition for ROS between the non-target EfOM and the target constituents can occur in two ways: scavenging of surfacebound ROS by EfOM and quenching of bulk phase ROS.^{15,29,31,32} Within the complex mixture of EfOM, less than

2% of the dissolved and colloidal organic materials are considered target contaminants, such as viruses or pharmaceuticals that originate from the influent wastewater;³³ thus, most photocatalytically generated 'OH radicals will be quenched by reactions with non-target EfOM. Indeed, EfOM has been reported to scavenge between 65 and 95% of 'OH in conventional effluents and is considered the most important 'OHscavenger in such systems.^{34,35} EfOM constituents, such as fulvic acid and humic acid (HA), have a net negative charge above pH 3 due to the presence of phenolic and carboxylic groups.^{36,37} These molecules can therefore interact favorably with and adsorb onto the polar surface of TiO₂, reacting directly with ROS production sites.

It is important to understand the factors that control surface and bulk quenching mechanisms; ROS-EfOM reactivity and EfOM-photocatalyst adsorption affinities drive bulk and surface quenching routes, respectively. Different ROS have differential reactivities; for example, singlet oxygen ($^{1}O_{2}$) is less reactive and more selective than 'OH.³⁸ Likewise, EfOM constituents may also vary in propensity to react with ROS, with some compounds being recalcitrant to strong oxidants, while others readily react with weaker ROS, such as $^{1}O_{2}$.^{30,39} With regard to adsorption interactions, the nature of the photocatalyst surface will determine the type of EfOM molecules that will adsorb onto the photocatalyst. These types of interactions have been studied in depth for the case of membrane fouling by organic matter,^{40–42} and offer potential insights into DOM-photocatalyst interactions.

Membrane technology can be used to selectively remove fractions of organic matter. In the case of an MBR treating municipal wastewater, the membrane's material, pore size, and fouling state affect its selectivity and, therefore, the composition of the EfOM.⁴³ It is known that, in general, hydrophilic macromolecular and colloidal portions of organic matter cause more reversible membrane fouling than other fractions in MBR systems by forming a cake layer.^{41,44,45} Fouling changes the effective pore size and surface characteristics of membranes; consequently, permeate quality changes over the operational timeline, since the last chemical cleaning event.46-48 Membrane operation may control DOM retention and thereby the composition of DOM that passes through (EfOM); therefore, the time since last cleaning event could be an important parameter when considering the use of effluent disinfection strategies. The extent to which membrane operation time can be used as a control EfOM quality is not well known. A better understanding of the variability of EfOM constituents as a function of membrane operational parameters is critical for applying post-filtration disinfection technologies. Elucidating the effects of membrane operation on EfOM content provides an excellent opportunity to scrutinize the effects of EfOM constituents on photocatalytic processes, a significant area of need for the field of photocatalytic water treatment.

While TiO₂ systems have been studied extensively, the mechanisms driving ROS inhibition by DOM are poorly understood. A recent literature review quantified the number of

research articles investigating "photocatalysis" and "natural organic matter" and found that of the 17 500 papers found when searching for photocatalysis, only 0.8% (137) also referenced DOM.⁴⁹ The segregation of DOM into fractions to discern phenomenological effects of constituents on photocatalytic processes is therefore a critical step towards practical application of photocatalysts. A study completed in 2014 on the effects of size-fractionation of DOM on the photocatalytic degradation of DOM by TiO₂ is perhaps the first report to scrutinize the inhibitory mechanism by analyzing fractionated DOM samples.⁵⁰ The approach in the present study utilizes EfOM from differentially fouled bioreactor membranes and functionally fractionated bioreactor DOM to provide a novel assessment of inhibitory mechanisms of DOM in TiO₂ photocatalysis.

Bulk supernatant DOM and EfOM samples collected in 2015 and 2016 from an operational MBR in a municipal wastewater treatment plant (WWTP) are studied here. Fractionation of samples in terms of DOM size and hydrophobicity, a method commonly used to isolate organics, was applied to MBR bulk supernatant. Here, the effects of different fractions of bulk supernatant DOM and EfOM samples on photocatalytic processes are assessed to identify the most important fractions to reject during filtration. Threedimensional fluorescence excitation-emission matrix (3DEEM) analysis is employed to characterize the resultant DOM from fractionation procedures and the MBR effluent samples, to better forecast and understand their effect on photocatalysis processes. 3DEEM is increasingly employed to understand DOM evolution in wastewater systems.^{51,52} A recent study also highlighted that 3DEEM can be used to distinguish proteins from biopolymers and humic substances and to quantify building blocks, with potential use as an on-line indicator to describe DOM fate and behavior.53 Further, this technique has distinguished the effects of different types of DOM on water treatment technologies (i.e., membrane fouling, UV attenuation, and disinfection byproduct formation).43,54-56 Inhibitory profiles of the DOM fractions and EfOM samples are established by measuring the photodegradation of a molecular probe as a function of total organic carbon (TOC) concentration. Inhibition mechanisms are discussed in the context of an experimentally validated model that accounts for surface and bulk phase quenching processes simultaneously.29 Finally, comments are made on the prospective utility of photocatalytic membrane reactors (PMRs)⁵⁷⁻⁵⁹ as a combined treatment process.

2. Materials and methods

2.1. Chemicals

Humic acid and 4-chlorobenzoic acid were obtained from Alfa Aesar (Haverhill, MA). Titanium dioxide (99.9% Anatase) was purchased from Alfa Aesar with a nominal particle size of 32 nm and surface area of 45 m² g⁻¹. Ultrapure water (>18.2 M Ω cm) was produced using a Nanopure Infinity system (Thermo Fisher Scientific Inc., Waltham, MA). HPLC solvents were HPLC-grade and obtained from Alfa Aesar.

2.2. EfOM sampling

EfOM samples were collected from a full-scale MBR wastewater treatment plant (La Grande Motte, France), which treats municipal wastewater and serves a population of approximately 60 000. The plant performs biological removal of nitrogen (nitrification and denitrification) and phosphorus. The plant comprises four MBR tanks, each equipped with KUBOTA Submerged Membrane Units® (SMUs, KUBOTA, Japan), which are flat sheet microporous membranes made of chlorinated polyethylene with an average pore size of 0.2 µm and a nominal pore size of 0.4 µm. Only two MBR tanks were studied. Here, we define MBR1 as the unit which underwent chemical cleaning and MBR2 as a reference unit that did not undergo chemical cleaning during the sampling period. MBR2 was two months into a three- to four-month cycle and therefore was chosen to represent a membrane during normal operation. To assess the cleaning effect, activated sludge (AS) and permeate samples were taken from MBR1 and MBR2 one day before and one day after the cleaning procedure took place for MBR1 (June 2016). After sampling, AS samples were filtered with a 1.2 µm glass microfiber filters (Whatman GF/C) to collect the dissolved portion of the AS, labeled as the bulk supernatant (BSN). Hence, four samples from the MBR1 cleaning campaign and two samples from MBR2 were collected and analyzed for this study, and each sample was given a reference name as shown in Table 1.

In addition to the samples taken to assess the effects of membrane cleaning, 500 L of AS were also collected from MBR1 in June 2015 to perform DOM fractionation using dialysis and XAD-resins. Prior to fractionation the AS was filtered successively through 50 µm and 2 µm polypropylene filters to collect BSN. Next, softening was performed using a sodium cation-exchange resin (Purolite, France) to remove calcium and magnesium ions, to avoid ion complexation with DOM and scaling during the following step: reverse osmosis (RO).⁶⁰ DOM in the BSN sample was concentrated via RO in order to minimize the time required for the fractionation step. A Filmtec TW 30 membrane was used for the RO process, since it is known to be more resistant to DOM adsorption.⁶¹ The RO process effectively concentrated the BSN by 100-fold which was subsequently used to perform DOM fractionation.

2.3. DOM fractionation

The first fractionation step consisted of isolating the colloidal portion of DOM by size exclusion, using dialysis (3.5 kDa, Spectra/Por 6 Dialysis Membrane) against HCl (0.01 mol L^{-1} , pH 2). Next, organic colloids were separated from colloidal silica and precipitated salts by dialysis (3.5 kDa) against 0.2 mol L^{-1} HF.⁶² The dialysate, approximately 30 L of HCl solution containing DOM compounds with a molecular weight smaller than 3.5 kDa, was then passed through XAD8 and

Table 1 Nomenclature of samples and fractions based on their respective MBR units, sampling period, or fractionation procedure					
Label	Collection	Description			
BSNf-MBR1	MBR1, bulk supernatant	Fouled membrane (pre-wash)			
BSNw-MBR1	MBR1, bulk supernatant	Washed membrane (post-wash)			
BSN-MBR2	MBR2, bulk supernatant	Midpoint between chemical cleaning events			
Pf-MBR1	MBR1, permeate	Fouled membrane (pre-wash)			
Pw-MBR1	MBR1, permeate	Washed membrane (post-wash)			
P-MBR2	MBR2, permeate	Midpoint between chemical cleaning events			
С	MBR1, bulk supernatant	Colloidal fraction			
HPO	MBR1, bulk supernatant	Hydrophobic fraction			
TPI	MBR1, bulk supernatant	Transphilic fraction			

XAD4 resins (Amberlite, Sigma Aldrich) arranged in tandem. This step allowed for the collection of hydrophobic (HPO) and transphilic (TPI) fractions.^{42,63} The hydrophilic (HPI) fraction, composed of low molecular weight hydrophilic DOM and salts, was collected in the outlet of the resins tandem. This fraction, however, was not used in the study because the solution contained highly concentrated salts, which co-precipitate with the organic matter. Removing these salts, while possible, would have required a complex purification step called azeotrophic distillation.⁶⁴ To collect HPO and TPI fractions adsorbed onto XAD resins, elution with an acetonitrile/MQ water solution (75/25% v/v) was performed, followed by evaporation and freeze-drying of the respective organic matter samples.

2.4. DOM characterization

2.4.1 TOC and UV_{254} absorbance measurements. TOC analysis was performed using a TOC-VCSH Shimadzu analyzer (Shimadzu Japan). The UV_{254} absorbance was measured in a 1 cm quartz cuvette using a UV-vis spectrophotometer (UV-2401PC, Shimadzu, Japan). The specific UV absorbance (SUVA₂₅₄) was then calculated as the ratio of UV_{254} absorbance and TOC value.⁶⁵ These analyses are reported in Table S1.[†]

2.4.2 3DEEM analysis. Fluorescence spectra were obtained using a Perkin-Elmer LS-55 spectrometer (USA) and a procedure described elsewhere.⁵³ Spectra were divided into five regions as defined by Chen *et al.*,⁵⁴ corresponding to different groups of fluorophores. The regions were categorized by excitation–emission ranges, as noted in Table 2. Region I is associated with aromatic protein-like fluorophores type I (tyrosine type); region II is associated to aromatic protein-like fluorophores type II (tyrosine type); region III corresponds to fulvic acid-like fluorophores; and region IV and V are associated with SMP-like fluorophores (tryptophane type) and humic acid-like fluorophores, respectively.

Table	2	Excitation	and	emission	wavelength	classifications	of
fluorop	hor	res					

	Region I	Region II	Region III	Region IV	Region V
Excitation, nm	200-250	200-250	200-250	250-350	250-500
Emission, nm	280-330	300-350	380-600	280-380	380-600

For qualitative analysis, spectra are represented in A.U. (arbitrary unit) and rejected fraction spectra (R) were calculated by subtracting permeate spectra from the BSN spectra, in order to better visualize the constituents that are rejected by the membrane. For semi-quantitative analysis, the volume of fluorescence $\Phi(i)$ (Raman unit nm²) normalized by the Raman spectra,⁶⁶ consisting of the integration of the spectral regions, was calculated in the different spectral regions using the following equation taken from the fluorescence regional integration (FRI)⁵⁴ method:

$$\Phi(i) = \mathrm{MF}(i) \sum_{\mathrm{ex}} \sum_{\mathrm{em}} I(\lambda_{\mathrm{ex}} \lambda_{\mathrm{em}}) \Delta \lambda_{\mathrm{ex}} \Delta \lambda_{\mathrm{em}}$$
(1)

where MF(i) is a multiplication factor, $\Delta \lambda_{ex}$ is the excitation wavelength interval (2 nm), $\Delta \lambda_{em}$ is the emission wavelength interval (0.5 nm) and $I(\lambda_{ex}\lambda_{em})$ is the fluorescence intensity at each excitation–emission pair (Raman units). $\Phi(i)$ normalization was necessary to compare values from different regions of the 3DEEM response. To do so, MF(i) was calculated using eqn (2).

$$MF(i) = \frac{\text{Total spectra area}}{\text{Specific region area}(i)}$$
(2)

For percentage analysis, the ratio between the volume of fluorescence of each region and the total volume was used.

2.5. Photochemical experiments

Photochemical experiments were conducted in an enclosed UV cabinet with a magnetically stirred photoreactor at room temperature. A 15 W low pressure mercury lamp (Sankyo Denki Co., Japan) was used as a UVC light source. The distance between the light source and reaction vessel was 20 cm. The irradiance at 254 nm at the location of the vessel was measured to be 295 μ W cm⁻² with a BLUE-Wave UVNb-25 Spectrometer (StellarNet Inc., Tampa, FL). The UV/Vis emission spectrum for the lamp, shown in Fig. S1,† was also recorded. The DOM fractions and EfOM samples described above, along with HA, were used to show the inhibitory effect of organic matter on photocatalytic degradation of target pollutants. Experiments utilized 15 ml of solution, containing 5 μ g L⁻¹ TiO₂ with 10 μ M *para*-chlorobenzoic acid (*p*CBA) as a probe compound that has a known reaction rate constant

with 'OH.³⁰ HA, DOM fractions, or EfOM samples in various concentrations were added to the reaction solutions to assess the quenching potential of each fraction. Sample aliquots of 0.5 mL were taken at fixed time points and analyzed for pCBA concentration via HPLC, according to methods reported elsewhere.⁶⁷ Briefly, this analysis was conducted with an Agilent HPLC (Agilent technology, 1260 infinity) using a C18 (125 mm) column using acetonitrile and 10 mM phosphoric acid as mobile phase solvents (40:60). The flow rate was 0.5 mL min⁻¹ and the detection wavelength was 234 nm. For all photochemical reactions, pCBA degradation rates were obtained by linear regression of plots of pCBA concentration versus radiant fluence (µJ cm⁻²). Fluence values were calculated according to Bolton and Linden (2003),⁶⁸ as described previously.57 Importantly, these calculations account for reductions in UV₂₅₄ transmission by using sample-specific UV₂₅₄ absorbance values and the transmission distance inside the reactor. The resulting observed photodegradation rates (k_{obs}) were, to a good approximation, first order with respect to radiant exposure (H, μ J cm⁻²) such that the units of $k_{\rm obs}$ are reported as (cm² μ J⁻¹), according to eqn (3)-(6):

$$\frac{\mathrm{d}C}{\mathrm{d}t} = k'_{\mathrm{obs}}C,\tag{3}$$

$$\frac{1}{H\left(\mu J \operatorname{cm}^{-2}\right)} \frac{\mathrm{d}C}{\mathrm{d}t} = \frac{1}{H\left(\mu J \operatorname{cm}^{-2}\right)} k_{\mathrm{obs}}' C, \qquad (4)$$

$$\frac{\mathrm{d}C}{\mathrm{d}H} = k_{\mathrm{obs}}C,\tag{5}$$

and

$$k_{\rm obs}\left(\rm cm^{2}\,\mu J^{-1}\right) = \frac{k_{\rm obs}^{\prime}\left(\frac{1}{\rm s}\right)}{E\left(\mu \rm W\,\rm cm^{-2}\right)}.$$
(6)

Here, $k'_{obs}(s^{-1})$ is the first-order degradation rate constant of *p*CBA, *C* is the molar concentration of *p*CBA, and *E* is the irradiance (μ W cm⁻²) at 254 nm. The differences in the k_{obs} in the presence or absence of organic compounds were used to quantify the inhibitory effect of these compounds. Control experiments were also conducted in the absence of organic matter, TiO₂, or light.

3. Results

3.1. Isolated DOM fractions

Prior to performing photocatalytic experiments, DOM fractions were characterized using 3DEEM to identify molecular characteristics of DOM within each fraction. The 3DEEM spectra compiled for the fractions are available elsewhere²³ and were used here to quantify the volume of fluorescence and the percentage of fluorescence of each region in Fig. 1.

3DEEM analysis showed that each of the three DOM fractions contained both classes of fluorescent compounds: proteins (regions I, II and IV) and humic substances (regions III and V) identified with red and blue patterns in Fig. 2, respectively. However, as seen in Fig. 1a, DOM fractions exhibited different fluorescent properties, reflecting differences in their compositions. The percentages of fluorescence of HPO and TPI fractions were similar for all regions and had a dominant proportion of aromatic protein-like type II and fulvic-like fluorophores (Fig. 1a). That HPO and TPI compositions did not vary significantly in terms of fluorophore content was expected; a study on EfOM of wastewater treatment plants also found that these fractions were similar in terms of fluorophore composition.⁶⁹ For the colloidal fraction, 80% of the fluorescent compounds were aromatic protein-like type I and II fluorophores. Recent studies showed that both protein-like and humic-like fluorophores impact photocatalytic performance. Protein-like constituents were found to react with 'OH radicals in bulk solution,⁵⁵ with reported reaction rate constants of amino acids, proteins, and peptides with 'OH ranging from 1.7×10^7 to 1.05 $\times 10^{10}$ M⁻¹ s⁻¹ (the rate constant between *p*CBA and 'OH is similar at 5.2 \times 10⁹ M⁻¹ s⁻¹),⁷⁰⁻⁷² while humic-like compounds, having a large number of carboxylic groups, adsorbed onto TiO₂ surfaces, particularly at low pHs.⁷³ Thus, the high proportion of fluorescing compounds in regions I and II in colloids, suggests that the colloids may be more reactive with 'OH than HPO and TPI. On the contrary, HPO and TPI are expected to exhibit more surface-phase quenching by adsorbing more strongly onto TiO₂ and than the colloids.

The volume of fluorescence is an indicator proportional to the concentration of fluorophores contained in each region. The higher electron density of fluorophores compared with other moieties could yield higher reactivity with ROS, and it would follow, then, that the higher the volume of fluorescence, the higher the quenching of photocatalysis. Thus, from Fig. 1b, and hypothesizing that surface-phase quenching is the most problematic for photocatalysis, the DOM quenching potency could be expected in the following order: HPO > TPI > C. A similar analysis can be conducted by measuring the SUVA₂₅₄ values as a representation of average aromatic moiety content, which is known to loosely indicate DOM hydrophilicity.⁷⁴ In general, DOM compounds with higher SUVA₂₅₄ values are considered to be more hydrophobic than those with lower values.⁷⁵ In addition, higher SUVA₂₅₄ values correspond to more aromaticity, which could indicate higher reactivity with ROS, given the electron rich moieties. The SUVA₂₅₄ values for the colloidal, HPO, and TPI fractions were measured to be 1.8, 2.2, and 1.6 L mg⁻¹ m⁻¹, respectively. Based on this method of analysis, and assuming that electron-dense functional groups are the primary factor in determining ROS quenching, the inhibition capacity of the fractions could be expected in this order: HPO > C > TPI. Neither of these methods are expected to conclusively predict the true inhibition potential, given the many additional factors involved with quenching mechanisms.

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Fig. 1 (a) Percentage of fluorescence and (b) volume of fluorescence of the colloidal (C), HPO and TPI fractions prepared at $1 \text{ mgC } L^{-1}$. Region I, region II, region II, region IV and region V correspond to aromatic proteins-like type I, aromatic proteins-like type II, fulvic-like, SMP-like and humic-like fluorophores, respectively.



Emission (nm)

Fig. 2 Fluorescence spectra of DOM from BSN and EfOM from permeate samples, with I, II, III, IV, V corresponding to region I (aromatic proteinslike type I), region II (aromatic proteins-like type II), region III (fulvic-like), region IV (SMP-like) and region V (humic-like). R spectra correspond to the mathematical subtraction of the permeate spectra from the bulk supernatant spectra allowing the identification of compounds retained by the membrane. Note the different color scale for the R spectra.

3.2. EfOM composition and effect of membrane fouling

To estimate the effect of membrane fouling and cleaning on the retention of fluorophores, 3DEEM spectra of MBR bulk supernatant and permeate were compared (Fig. 2). The membranes rejected most compounds from regions I and II in the three MBR cases studied. This selectivity was apparent in the 3DEEM spectra obtained by subtracting the permeate spectrum from the BSN spectrum. It is likely that most of these aromatic-like fluorophores were associated with organic colloids since they represented 80% of the overall colloidal content. This observation is consistent with a previous study that demonstrated that colloids were major membrane foulants.²³

Fig. 3a and c show the fluorophore compositions in EfOM samples as percentages of the different regions; these data showed a preferential rejection of the fluorophores from region I and region II. Indeed, for the three samples, the membrane reduced the fluorescence by $11 \pm 2\%$ in both regions I and II. This reduction corresponded to an increase of fluorescence percentage of the regions III and V in the permeate. The relative increase of the humic substance-related fluorophores confirmed that the membrane preferentially retains colloids, since they are typically high molecular weight molecules associated with protein-like fluorophores (Fig. 1a).53 Membrane fouling clearly affected the type of fluorophores retained in the MBR (Fig. 3b and d). The three fouling stages present similar bulk supernatant volumes of fluorescence (Fig. 3b) and permeate percentage of fluorescence profiles (Fig. 3c), but different volumes of fluorescence in the permeate (Fig. 3d). The TOC normalized volume of fluorescence for Pf-MBR1 was reduced by 60% (Fig. 3b and d), while the volume of fluorescence was only reduced by 14% and 10% for P-MBR2 and Pw-MBR1, respectively (Fig. 3b and d). Membrane fouling therefore has a clear effect on fluorophore quantity, via restricting EfOM permeation (Fig. 3b). Indeed, more fluorescent compounds are retained, on a per carbon basis, by a fouled membrane. This result shows that the fouling layer on the membrane surface selectively removes compounds rich in functional groups with high electron density, which are more reactive with ROS than other moieties. Pw-MBR1 and P-MBR2 are therefore expected to quench photocatalysis to a greater extent. This assumption is supported by the SUVA₂₅₄ data: values for the Pf-MBR1, Pw-MBR1, and P-MBR2 samples were measured to be 0.8, 2.0, and 2.0 L mg⁻¹ m⁻¹, respectively. Pf-MBR1, having a SUVA₂₅₄ value of 0.8 L mg⁻¹ m⁻¹, is characterized by non-aromatic organic compounds and therefore fewer potential functional groups reactive with ROS. On the contrary, Pw-MBR1 and P-MBR2, with SUVA₂₅₄ values of 2.0 L mg⁻¹ m⁻¹, contain more aromatic compounds, which may preferentially compete with ROS.

Control of membrane fouling may provide an opportunity to increase photocatalysis process efficiency by regulating the chemical makeup and concentration of EfOM. Less frequent cleaning events could be ideal, since the fouled membranes provided the highest DOM retention. From the fluorescence volumes, it is expected that TiO₂ photocatalysis would be quenched to a greater extent by Pw-MBR1 and P-MBR2, than by Pf-MBR1.



Fig. 3 Percentage (a and c) and volume (b and d) fluorescence values for bulk supernatant (a and b) and permeate (c and d) samples.

3.3. Inhibition of 'OH by DOM fractions

Segregation of MBR DOM into functional categories allowed for a unique examination of the inhibition potential of these functional classes of compounds. Colloidal, HPO, and TPI fractions were each examined for concentration-dependent inhibitory activity. Control tests confirmed the photocatalytic action of TiO₂ and differentiated the role of ROS from the direct photolysis by UV_{254} light (Fig. S3[†]). The action by UV_{254} alone represented the lower bound of $k_{\text{obs},pCBA}$, where 'OH radicals were completely quenched by DOM. Likewise, the case of TiO₂ and pCBA in pure water served as the upper bound of photocatalytic efficiency, with no interfering quenching agents. The $k_{obs,pCBA}$ values plotted in Fig. 4 showed that of the three DOM fractions, colloids exerted the strongest inhibition by far. The corresponding $k'_{obs}(s^{-1})$ data is shown in Fig. S4.† The TPI and HPO portions were similar in their effect on $k_{obs,pCBA}$, and exerted mild inhibition at low TOC concentrations. Interestingly, for both TPI and HPO, the $k_{\text{obs.}p\text{CBA}}$ increased from 7.5 to 10 mgC L⁻¹. This increase in photodegradation efficacy was surprising but not unprecedented; it was recently reported that natural organic matter (NOM) actually enhanced the TiO₂-driven photodegradation of carbamazepine, pharmaceutical compound, at specific TiO₂: NOM ratios, by up to 8%.⁷⁶ Favorable NOM-carbamazepine interactions explained the increased effectiveness; these interactions draw the compound closer to the active surface sites of TiO₂, where 'OH are present at higher concentrations. The colloidal fraction did not increase the photoactivity at any concentration. Examination of the inhibition profiles of



Fig. 4 *p*CBA degradation rate constants in the presence of 5 mg L⁻¹ TiO₂ and various concentrations of colloids, TPI, HPO, and HA are depicted here. The rate constant for *p*CBA degradation by UVC without TiO₂ is also shown. Ambient temperature was measured at 24 °C.

the three DOM fractions in the context of 3DEEM analysis (Fig. 1) suggested that the quenching action of the DOM fractions is correlated to higher concentration of colloids, which are characterized by a higher proportion of fluorescence in region I and region II (Fig. 1a). This observation suggests that despite higher volumes of fluorescence, HPO and TPI are less potent inhibitors of photocatalysis than the colloids. The surface interactions, and therefore inhibition mechanism, of the colloids with the TiO₂ surface could be fundamentally different from that of the HPO and TPI fractions, because the colloidal fraction was not segregated based on surface character, but rather by size only. Control of membrane surface properties and fouling could reduce the colloidal content-much of which consists of high molecular weight molecules that can be preferentially retained-in EfOM and thereby mitigate the quenching of photocatalytic processes by DOM.23

3.4. Inhibition of 'OH by EfOM

The EfOM of the three MBR permeate samples, as described above, was tested for inhibition potential of 'OH-mediated *p*CBA degradation. The samples were examined on a TOC basis to discern changes in inhibition potential caused by qualitative differences in EfOM composition. A low concentration of TiO₂, relative to that used in similar studies on photocatalyst–DOM interactions,^{29,38,77,78} was selected to avoid the effects of EfOM transformation by oxidation. Experiments shown in Fig. S2† showed that hour long UVC irradiation experiments with 10 mg L⁻¹ HA and various concentrations of TiO₂ showed that *p*CBA photodegradation kinetics were linear for the TiO₂ concentration of 5 mg L⁻¹. Tests with TiO₂ concentrations of 100 mg L⁻¹ or higher showed accelerating kinetics and suggested that HA was itself being degraded by 'OH radicals so that its inhibition potential changed with time.

The inhibition capacities of MBR EfOM samples were evaluated by measuring $k_{\text{obs},pCBA}$ as a function of individual EfOM sample concentrations. These rates were calculated across concentrations ranging from 0 to 2.3 mgC L^{-1} (Fig. 5). The corresponding $k'_{obs}(s^{-1})$ data is shown in Fig. S5.[†] Comparing $k_{\text{obs},p\text{CBA}}$ values for the same TOC content reveals that the state of membrane fouling drove clear distinctions in inhibitory activity of the EfOM. While it was expected that a fouled membrane would reject more DOM than a clean membrane, the inhibition capacity on a per carbon basis was not known. Here, it was observed that EfOM from a fouled membrane system inhibited the photocatalytic process much less than EfOM from a cleaned membrane. At just 0.5 mgC L^{-1} , Pw-MBR1 quenched the photocatalytic process completely, while no quenching was observed by Pf-MBR1 EfOM at the same concentration. This result provides evidence that the changes in EfOM composition caused by membrane fouling; the reduction of colloid concentration and total fluorophores is especially beneficial for photocatalytic operation. 3DEEM confirmed that molecules containing fluorescent groups in regions I and II impact photocatalytic performance more

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Fig. 5 pCBA degradation rate constants in the presence of various concentrations of HA, effluents from Pf-MBR1, from Pw-MBR1, and from P-MBR2 with 5 mg L⁻¹ TiO₂ are depicted here. The rate constant for pCBA degradation by UVC without TiO₂ is also shown.

than other compounds. Qualitative changes in DOM retention by the membrane, therefore, impacted the photocatalytic quenching process. Considering these results in the context of the DOM fractions analysis, retention of organic colloids by the fouled membrane was likely enhanced by the formation of a fouling layer.^{3,22} Inhibition by P-MBR2, sourced from a membrane at the midpoint between chemical cleanings, was between the two extremes of Pw- and Pf-MBR1, with a ~75% reduction in $k_{\text{obs},p\text{CBA}}$ at 0.5 mgC L⁻¹. Alternatively, it may be possible to choose or modify membrane materials to selectively reject the organic colloidal materials regardless of the fouling state. HA served as a reference material, which represents NOM found in drinking water sources more closely than EfOM, and exhibited stronger quenching than the P-MBR2 case but less inhibition than Pw-MBR1. It is noteworthy that HA inhibits TiO₂ driven photocatalysis to a greater extent than EfOM from a fouled MBR on a carbon basis. This finding contradicts a 'common sense' assumption that could be made based solely on TOC values: that photocatalysis would be more applicable for drinking water applications than for WWTP effluent.

The 3DEEM analyses (Fig. 3) of the MBR EfOM samples predicted that the fouled membrane would reduce the quantity of fluorescent compounds in the EfOM and therefore lead to less inhibition of photocatalysis. However, for cases of similar fluorescence volumes, as for Pw-MBR1 and P-MBR2 in particular, the use of 3DEEM did not explain differences in inhibitory action. In these cases, other factors, such as the hydrophobic/hydrophilic character of the EfOM, may have been altered by the membrane fouling but not detected by 3DEEM or TOC analysis. It is well known that membrane fouling affects rejection of DOM components^{41,43,79} and that the mechanism of action is not simply size exclusion alone: changes in the surface characteristics (*i.e.* charge and hydrophobicity), due to fouling layer formation, are also important.^{3,22}

3.5. Inhibitory mechanisms for DOM samples

Identification of the mechanism of inhibition by DOM on TiO₂ photocatalysis is the key to designing processes to overcome the problem of ROS quenching. Numerous studies have evaluated the adsorption interactions of NOM onto TiO₂, fitting experimental findings to Freundlich⁸⁰ or Langmuir-Hinshelwood^{32,77,81} isotherms. Only recently, however, was a model developed that accounted for both bulk- and surfacephase quenching interactions.²⁹ In their work, Brame et al. experimentally validated a model that combined a multisolute Langmuir model⁸² with bulk phase competitive reaction rates by assuming steady-state ROS concentrations.²⁹ Based on this dual-phase model, the mode of inhibition (bulk or surface reactions) was successfully predicted by analysis of the dependency of k_{obs} on TOC. A linear dependence of k_{obs} , on TOC implied that inhibition primarily occurred in the bulk phase and surface interactions were unimportant; alternatively, an exponential decay of k_{obs} with increasing TOC indicated that surface sorption and reactions played a significant role in the inhibitory process.²⁹ Note that the aforementioned report used Suwannee River humic acid as an NOM source, which consists of a wide range of molecules;²⁹ applying Brame's model in experiments with fractionated DOM samples is an important extension of the earlier work allowing for a discriminating analysis of inhibition mechanisms across the DOM spectrum. Here, all experiments were performed with the same probe compound, photocatalyst concentration, and UV254 lamp, so normalization of $k_{\text{obs},p\text{CBA}}$ was not necessary. The inhibition profile for HA was non-linear and therefore depended on surface interactions, in line with previous reports for TiO₂ inhibition by NOM.^{29,38,83,84} Upon examination of the inhibitory profiles of the MBR effluents, trends for Pf-MBR1 and P-MBR2 were noted to be nearly linear, whereas Pw-MBR1 showed an exponential relationship. These observations suggest that the membrane fouling layer played a critical role by rejecting DOM that adsorbs favorably onto the surface of TiO_2 , thereby exerting a strong quenching effect on photocatalytic processes. These observations correlate well with the observed inhibition profiles of the fractionated DOM.

As discussed, the colloidal fraction of BSN DOM exerted the strongest inhibitory action of any of the fractions (Fig. 4). The $k_{obs,pCBA}$ inhibition profiles of the DOM fractions reveal that the colloids quenched the photocatalytic process *via* sorption onto the TiO₂ surface and reacting with surfacebound 'OH. The HPO and TPI fractions, however, displayed a linear dependence—if the spurious enhancement of $k_{obs,pCBA}$ at the 10 mgC L⁻¹ mark is neglected—on TOC. The HPO and TPI samples, therefore, primarily reduced $k_{obs,pCBA}$ through bulk phase reactions limited by diffusion and relative



Fig. 6 $k_{obs,pCBA}$ inhibition profiles of HA, Pf-MBR1, and the colloidal fraction. Data from Fig. 3 and 4 are used here.

reaction rates. Note that these remarks on quenching mechanisms are generalizations: even the fractionated DOM samples contain a wide variety of molecules, each with specific adsorption affinities and reaction rates. Still, results of both fractionation and membrane fouling conditions showed significant changes to inhibitory action of DOM. The inhibitory action of the colloidal fraction was particularly interesting, given the lack of inhibitory action by effluent from the fouled membrane. These observations taken together in Fig. 6 (data replotted from Fig. 4 and 5) suggest that fouled membranes reject key organic colloids that would otherwise adsorb strongly to TiO₂ surfaces and greatly reduce photodegradation rates. The corresponding $k'_{obs}(s^{-1})$ data is shown in Fig. S6.† The prospective utility of a membrane for pretreatment is clearly demonstrated by these results: if a membrane can be selected or optimized to reject problematic colloids, photocatalysis may indeed be effective for disinfection of MBR effluent.

4. Conclusions

The challenge of unwanted ROS-DOM reactions has long plagued photocatalysis, particularly for applications dealing with high TOC concentrations such as in a typical MBR effluent. 3DEEM can be used to predict the inhibitory effects of DOM composition, and the experiments shed new light on the quenching of photocatalysts by DOM. First, the total fluorescence volume correlated well with the extent of photocatalytic inhibition on a carbon basis, further the DOM fractionation demonstrated that the colloidal fraction of DOM exerted stronger quenching action than HPO and TPI. Assessment of the effects of membrane fouling showed that permeate from a fouled membrane caused less inhibition than that of clean and moderately fouled membranes. In fact, DOM from fouled membrane appeared to quench 'OH primarily via bulk-phase scavenging, whereas DOM from a clean membrane showed an inhibition profile consistent with surfacephase reactions,²⁸ suggesting that the membrane fouling layer rejected materials that would otherwise adsorb strongly to the TiO₂ surface. To enhance photocatalysis efficiency, it might be possible to select a membrane with a "built-in" selectivity similar to that of the fouled membrane in order to remove the problematic colloidal fraction. Analysis of the inhibition profiles of the EfOM described here suggests that for the operation of a PMR a trade-off can be made between the operational pressure and the photocatalytic efficiency; by reducing the (chemical) cleaning frequency and thereby maintaining a minimal level of fouling, inhibition of photocatalysis by organic colloidal inhibitors would be mitigated at a cost of higher trans-membrane pressures. Further, the surface coverage of TiO2 on PMRs can be tuned to optimize photocatalyst surface area⁸⁵ and may not be limited to the DOM: TiO₂ ratios explored here.

Further research on the fundamental surface interactions between these organic colloidal materials and photocatalyst or membrane surfaces should be pursued in order to develop mitigation strategies for DOM-related ROS inhibition. Specifically, the assessment of the potential effects of the hydrophilic fraction and dissolved ions (*i.e.*, multivalent cations and halides), which were not retained by the fractionation processes, should be examined. The results of the present study may be applicable to the use of photocatalytic materials in systems containing other DOM sources, therefore additional investigations on systems such as potable water supplies or industrial waste streams would be timely and important.

Conflicts of interest

There are no conflicts of interest to declare.

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