Tailoring surface charge and wetting property for robust oil-fouling mitigation in membrane distillation

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ABSTRACT

The efforts of developing anti-oil-fouling membranes have so far been focusing on tailoring membrane surface wetting properties, whereas the impacts of surface charge are often eclipsed. In this study, we investigated the impacts of surface charge and wetting property on oil fouling kinetics in membrane distillation (MD). Two composite membranes with in-air hydrophilic and underwater oleophobic surfaces, one of positive charge and the other of negative charge, were fabricated by modifying a hydrophobic polyvinylidene fluoride (PVDF) membrane with hydrophilic polyelectrolytes with different charges. The modified composite membranes were compared with the reference PVDF membrane for their contact angles, adhesion force curves, and fouling kinetics in MD processes. It was found that the negatively charged composite membrane performed the best in mitigating fouling by the negatively charged oil emulsion, followed by the positively charged composite membrane, with the pristine PVDF membrane being the most susceptible to oil fouling in MD experiments. The results from underwater oil CA measurements and oil probe force spectroscopy corroborated the results in the fouling experiments. The impact of surface charges on oil-membrane interaction and associated mechanism were also discussed.

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1. Introduction

Membrane distillation (MD) is an emerging membrane-based thermal desalination process capable of utilizing low-grade heat. In an MD process, a microporous hydrophobic membrane is employed as the barrier for liquid water transport and the medium for water vapor transport. The temperature difference between hot salty water (the feed solution) and cold deionized water (the distillate) results in a partial vapor pressure difference which drives the vapor to transport from the feed stream to the distillate stream, thereby desalinating the feed water [1–3]. Research interests in MD have recently grown significantly due to the increasing demand for desalination and the potential of MD to become a competitive technology in certain challenging desalination applications. For example, a distinct advantage of MD is the ability to harness low-grade thermal energy such as waste heat, geothermal energy, and solar thermal energy for desalination [4–6]. Such an advantage, together with the small system footprint and low capital cost, renders MD an appealing process for sustainable and off-grid desalination [7–11]. On the other hand, MD has also been identified as one of the only few promising technological candidates for desalinating shale gas produced water which, in certain cases, is too saline for reverse osmosis to treat [12–14].

Most existing MD membranes are only applicable to desalinate relatively ‘clean’ water (e.g. seawater) with a very low concentration of hydrophobic contaminants. This is because the commercially available hydrophobic membranes are inherently susceptible to fouling by hydrophobic contaminants due to the strong and long-range hydrophobic-hydrophobic interaction [15–17]. The direct consequence of MD membrane fouling is flux decline attributed to the blocking of membrane pores by the accumulated foulants. This problem of membrane fouling by hydrophobic contaminants is particularly conspicuous when MD is employed to desalinate saline industrial wastewaters enriched in hydrocarbons. One prominent example is shale gas produced water, which is not only hyper-saline but in many cases abundant with hydrocarbons. Previous MD studies using a hydrophobic polyvinylidene fluoride (PVDF) membrane to treat a feed solution with oil reported rapid flux decline that led to process failure in a relatively short time [18,19]. Another example is the application of MD to treat...
anaerobic digestate which, though not as potent as oil emulsion as a foulant, still led to considerable flux decline in a long term operation [20].

An effective approach to mitigate oil fouling in MD is to develop membranes with an underwater oleophobic surface that resists oil adhesion when submerged in water. The underwater oleophobicity is to be distinguished from the in-air oleophobicity which stands for resistance to oil adhesion in air. Inspired by biological surfaces such as sharkskin and clamshells that are resistant to biological or organic adhesion underwater, the materials science community has developed a plethora of interfacial materials with underwater superoleophobicity for different applications such as marine biofouling control [21,22] and oil-water separations [23–33]. In general, there are two requirements for constructing an underwater superoleophobic surface: rough (textured) surface morphology and hydrophilic surface chemistry. Following this principle, composite MD membranes with a hydrophobic substrate and an underwater oleophobic top surface have been developed using hydrogel–nanoparticle composites for oil fouling mitigation [18,19]. The presence of highly hydrophilic surface coating renders the feed-facing surface of the MD membrane in-air superhydrophilic and underwater superoleophobic, enabling MD to desalinate oily feed water with robust anti-fouling performance.

It is well known in surface science that a hydrogen-bond forming, superhydrophilic surface can resist the adhesion of hydrophobic substances underwater due to the formation of a surfactant hydration layer that serves as an energetic barrier for hydrophobic foulants to attach onto substrate surface [34–38]. This mechanism, namely the hydration force, has been a pivotal guiding principle for developing fouling resistant membranes and anti-oil-adhesion surfaces [39,40]. Although the effect of hydration force is indeed prominent, there is another important factor that plays a critical role in oil-membrane interaction but is nonetheless often neglected in developing underwater oleophobic surfaces [28,41–43]. This factor is surface charge, which plays a significant role in controlling both electrical double layer (EDL) interaction and interfacial tension [44,45].

In a recent study where a PVDF MD membrane was coated with titanium dioxide (TiO₂) nanoparticles and polyethylene glycol (PEG) to acquire resistance to mineral oil fouling, it has been observed that varying the pH around the isoelectric point of the modified PVDF slightly affected the fouling kinetics [19]. However, such an effect was not appreciable, which may be explained by the fact that both the modified PVDF membrane and the mineral oil emulsion had similar isoelectric points and their interaction was thus always repulsive regardless of the solution pH. On the other hand, many studies conducted to develop underwater superoleophobic meshes or membranes for oil-water separation only focused on engineering the surface wetting properties but largely overlooked the impact of the surface charge. Because both the electrostatic interaction and the hydration force may be important as suggested by the extended Derjaguin Landau Verwey Overbeek (x-DLVO) theory [46], it is our objective in this study to systematically assess the impacts of these two interactions on the oil-membrane interaction and consequently on the propensity of an MD membrane to be fouled by an oil-in-water emulsion.

In this study, we compare the performance of three membranes with different surface wetting and charge properties in direct contact membrane distillation (DCMD) experiments with feed water containing crude oil. A hydrophobic and negatively charged PVDF membrane was used as the reference. The other two membranes are composite membranes prepared by modifying the hydrophobic PVDF membranes with nanocomposite coating comprising silica nanoparticles and hydrophilic polymers with different surface charges. We characterize these membranes to acquire information regarding their morphological and wetting properties and conduct DCMD experiments with these membranes to compare their fouling kinetics. We also conduct force spectroscopy using a force tensiometer to probe the underwater interactions between a crude oil droplet and different membranes, and apply these force curves to elucidate the difference of these membranes in their fouling propensity.

2. Materials and methods

2.1. Materials and chemicals

The hydrophobic microporous membranes, used as the reference in fouling experiments and as the substrates for surface modification, were made of PVDF with a nominal pore size of 0.45 μm (GE Healthcare Life Sciences, Pittsburg, PA). Dopamine hydrochloride, Trizma hydrochloride buffer, poly(diallyldimethylammonium chloride) solution (PDDA, 20% in H₂O), aceton, sodium chloride (NaCl), potassium chloride (KCl), sodium hydroxide (NaOH), and hypochloric acid (HCl) were all procured from Sigma Aldrich (St. Louis, MO). Silica nanoparticles (SiNPs) with diameters ranging between 20 and 60 nm were purchased from SkySpring Nanomaterials (Houston, TX). The crude oil as the fouling agent was acquired from Texas Raw Crude Oil (Midland, TX).

2.2. Fabrication of the underwater oleophobic membranes

We prepared two different composite membranes by modifying the PVDF substrate membrane. Both membranes were underwater oleophobic due to the presence of the highly hydrophilic coating. However, due to the different hydrophilic surface modifiers used, the two membranes were of opposite charges at environmentally relevant pH. This subsection describes the experimental details of fabricating the two composite membranes.

2.2.1. Composite membrane coated with polydopamine (PDA) and SiNPs

The first composite membrane was prepared by coating a PVDF membrane with a layer of negatively charged nanocomposite of PDA and SiNPs (PDA/SiNPs). We prepared such a composite membrane by first spray-coating the substrate membrane with SiNPs and then dip-coating the SiNPs coated membrane with PDA [47,48]. The SiNPs, which are inexpensive and chemically inert, were used to increase the surface roughness and create hierarchical morphology that are essential for in-air superhydrophilicity and underwater superoleophobicity. For spray-coating, 0.5 g SiNPs were dispersed in 100 mL acetone under sonication for 5 min to obtain a 5 g/L SiNPs suspension. 20 mL of SiNPs suspension were spray-coated onto a PVDF membrane substrate with an area of 20 cm × 10 cm using a spray gun with 0.2 MPa air.

After briefly dried in air, the SiNPs coated membrane was submerged in a dopamine solution prepared by dissolving 0.1 g dopamine powder in 100 mL of a 15 mM Trizma hydrochloride buffer solution with the pH adjusted to 8.5. A layer of PDA formed as a result of self-polymerization of dopamine, covering the SiNPs and the top surface of the PVDF membrane. During the dip-coating process, the PVDF membrane was taped around on a glass substrate to ensure that only the SiNPs coated surface of the membrane was exposed to the dopamine suspension. Because the hydrophilic PDA solution could not wet the hydrophobic pores of the PVDF membrane, the hydrophobicity of the inner pore structure of the PVDF substrate was not compromised, i.e., only the wetting property of the membrane surface, not that of the entire membrane, was altered by this membrane modification approach. After
24-hour self-polymerization of dopamine, the PDA/SiNPs coated PVDF (PDA/SiNPs-PVDF) membrane was washed with DI water and dried at room temperature.

2.2.2. Composite membrane coated with PDDA and SiNPs
   The second composite membrane with PDDA and SiNPs nanocomposite (PDDA/SiNPs) was fabricated via layer-by-layer spray-coating [49]. All the spray-coating steps were conducted using a spray gun with 0.2 MPa pressurized air. First, a PVDF membrane substrate with an area of 20 cm × 10 cm was sprayed-coated a layer of PDDA with 5 mL aqueous PDDA solution (52 g/L) and then dried at room temperature. Second, 10 mL SiNPs suspension (5 g/L in acetone) was sprayed onto the PDDA coated membrane to form a SiNPs layer. The function of first PDDA layer was to impart positive charge to the PVDF substrate so that the SiNPs can attach onto the PDDA coated membrane via electrostatic interaction. Lastly, the air-dried SiNPs decorated membrane was spray-coated with another layer of PDDA solution (1 mL, 52 g/L). The resulting membrane was heated at 100 °C for 1 h. The coating applied using this approach has been tested to be mechanically robust [49].

2.3. Characterizations of morphology and wetting property
   The surface morphology of the pristine PVDF membrane and the two fabricated composite membranes was observed using scanning electron microscopy, or SEM (Merlin, Zeiss, Thornwood, NY) and atomic force spectroscopy, or AFM (Dimension Icon, Bruker, Billerica, MA). The pore size distribution and the liquid entry pressure, LEP, of each membrane were determined with a Bruker, Billerica, MA. The water vapor flux and salt rejection were calculated from the hydraulic pressure in the feed stream than in the distillate stream [13,18]. This operation conditions resulted in a slightly higher hydraulic pressure in the feed stream than in the distillate stream in this specific test cell, so that the liquid flow through any potentially wetted pores would always be driven from the feed stream to the distillate stream and thereby increase the salinity of the distillate stream. The mass and conductivity of the solution in the distillate reservoir were constantly measured and recorded. The water vapor flux and salt rejection were calculated from the temporal change of mass and conductivity of the distillate.

2.6. Membrane fouling tests
   DCMD experiments were conducted with the saline crude oil emulsion to compare the fouling behaviors of the pristine PVDF membrane and the fabricated composite membranes. In these experiments, membrane coupons with an area of 2.5 cm × 8 cm were used in a cross-flow test cell using a co-current flow configuration. In the experiments with composite membranes, the modified, underwater oleophobic surfaces of the membranes were in contact with the feed solution, whereas the unmodified sides faced the distillate. The influent temperatures of the oily feed solution and deionized distillate were 60 °C and 20 °C, respectively.

   To facilitate the detection of membrane wetting, the flowrates of feed and distillate streams were controlled to be 0.45 L/min and 0.2 L/min (9.6 cm/s and 4.3 cm/s in our test cell), respectively [13,18]. This operation conditions resulted in a slightly higher hydraulic pressure in the feed stream than in the distillate stream in this specific test cell, so that the liquid flow through any potentially wetted pores would always be driven from the feed stream to the distillate stream and thereby increase the salinity of the distillate stream. The mass and conductivity of the solution in the distillate reservoir were constantly measured and recorded. The water vapor flux and salt rejection were calculated from the temporal change of mass and conductivity of the distillate.

2.7. Underwater oil-adhesion force measurements
   To mechanistically understand the fouling behaviors of the pristine PVDF membrane and the fabricated composite membranes, force spectroscopy was conducted with a force tensiometer (Sigma701, Attention, Finland) to probe the underwater interaction between a crude oil droplet and the membrane surfaces [25,56–58]. In these force spectroscopic experiments, the measurements of underwater CA with crude oil were carefully hung on to a submerged force probe (T114, Attention, Finland) using a syringe. For collecting the force curves, the force probe with the attached oil droplet approached the membrane sample surface at a constant speed of 0.3 mm/min until the oil droplet contacted with the membrane surface. Once physical contact between the oil droplet and the membrane sample occurred, the force probe retracted from the membrane surface at the same speed as approaching. Compressing the oil droplet against the sample surface was deliberately avoided, because the force probe was an open ring that can be readily forced into the crude oil phase and generate data that is not directly relevant to the oil-membrane interaction. The results from the force spectroscopy were force curves that quantify the force, measured by the micro-electro-mechanical sensor, as a function of the force probe position.

3. Experimental results
3.1. Membrane surface properties
3.1.1. Membrane surface morphology
   The SEM and AFM images of the pristine PVDF membrane (Fig. 1(A), (D) and (G)), the PDA/SiNPs-PVDF membrane (Fig. 1(B), composed of the two parallel surfaces covered by the membrane samples. This flow of electrolyte solution generated streaming potentials from which the zeta potential of the membrane surface was calculated using the Helmholtz–Smoluchowski equation [54,55]. The electrolyte solution was 1 mM KCl and the pH was automatically adjusted by the addition of KOH and HCl.
(E) and (H)), and the PDDA/SiNPs–PVDF (Fig. 1(C), (F) and (I)), show that surface modifications drastically affected the morphology of the PVDF membranes. Specifically, both the top view (Fig. 1(B) and (C)) and the cross-section view (Fig. 1(E) and (F)) of the SEM images suggest that the nanocomposite coatings thoroughly cover the surface of the PVDF substrate and block most of the membrane pores on the surface. However, the modifications did not significantly affect the pore size distribution (Fig. A1) of the membrane because the thickness of coating layer is very small compared to the thickness of the PVDF membrane which was 176.4 ± 7.3 μm.

The surface modifications did not lead to any appreciable change in LEP either. Compared to the PVDF membrane with an LEP of 2.93 bar, the LEP of the PDDA/SiNPs and PDA/SiNPs modified PVDF membranes were 2.98 and 3.01 bar, respectively. Although the surface modification significantly reduced the size of the membrane surface pores, the strong hydrophilicity of the coating layer facilitated the permeation of water with little extra resistance. Consequently, the LEP should still be primarily determined by the wetting property and pore size of the hydrophobic substrate which in all three cases were the same PVDF membrane. For the similar reason, the coated membranes still maintained significant water vapor permeability, which will be further discussed in Section 3.3. The hydrophilic nanocomposite PDA/SiNPs and PDDA/SiNPs coatings exhibited hierarchical roughness which, according to Wenzel theory, enhances the hydrophilicity of the surface [59,60]. The results from PM-IRRAS spectra confirmed the chemical compositions of the surface coatings on the modified composite membranes (Fig. A2).

Fig. 1. SEM images featuring the local surface morphology of (A) the commercial PVDF membrane, (B) the PDA/SiNPs–PVDF composite membrane, and (C) the PDDA/SiNPs–PVDF composite membrane. The inset images show the membranes surface at a larger scale. The PDA/SiNPs and PDDA/SiNPs coatings covered most of the membrane pores. Cross-section SEM images of (D) the commercial PVDF membrane, (E) the PDA/SiNPs–PVDF composite membrane, and (F) the PDDA/SiNPs–PVDF composite membrane. Surface AFM images of (G) the commercial PVDF membrane, (H) the PDA/SiNPs–PVDF composite membrane, and (I) the PDDA/SiNPs–PVDF composite membrane.
3.1.2. Membrane surface potential

Surface potential is an important membrane property affecting the electrostatic interaction between membranes and oil droplets. In this study, we measured the zeta potentials of each membrane at different pH (Fig. 2). The zeta potential varied with pH due to protonation or deprotonation of surface functional groups. Throughout the entire pH range tested, the pristine PVDF membrane was negatively charged (green triangles in Fig. 2) and the PDDA/SiNPs-PVDF composite membrane was positively charged (red circles in Fig. 2). The point of zero charge (PZC) for the PDA/SiNPs-PVDF membrane was slightly below 4, significantly lower than the pH of the saline crude oil emulsion, which was between 5–6.

3.1.3. Membrane surface wetting property

The wetting properties of the composite membranes were drastically different from that of the pristine PVDF membrane. The in-air water CA (blue bars in Fig. 3) of the pristine PVDF, the PDA/SiNPs-PVDF composite membrane, and the PDDA/SiNPs-PVDF composite membrane were 106.7 ± 3.7°, 35.1 ± 5° and 17.9 ± 2°, respectively. Both nanocomposite coatings imparted strong hydrophobicity to the PVDF membrane that was originally hydrophobic, which can be explained by the strong attractive interaction between water and the hydrophilic polymer on the modified membrane surfaces.

To obtain the membrane wettability information directly relevant to oil fouling in MD, we also measured the underwater CA with crude oil for the three membranes (orange bars in Fig. 2). The unmodified PVDF membrane was instantly wetted by the crude oil with a negligibly small oil CA, indicating that the membrane was underwater superoleophilic. In comparison, the PDA/SiNPs-PVDF composite membrane exhibited underwater superoleophobicity with a very high oil CA (162.8 ± 2.4°). Such underwater superoleophobicity is attributable to the strong interaction between the water and the hydrophilic surface coating (i.e., the hydration force). As a highly hydrophilic polymer, PDA has a variety of amide and alcohol functional groups that can form hydrogen bonds with water and create an interfacial hydration layer on the membrane surface. Wetting the hydrogel coated membrane by the crude oil requires surface dehydration, which is energetically highly unfavorable. With a similar principle, underwater superoleophobicity was also expected for the PDDA/SiNPs-PVDF composite membrane, because PDDA is also a highly hydrophilic polymer that forms abundant hydrogen bonds with water. Indeed, the initial underwater oil CA for PDDA/SiNPs-PVDF membrane was also very high (157.7 ± 1.5°), indicating the composite membrane was superoleophobic upon immediate contact with water. However, the oil CA decreased over time to 57.1 ± 2° in about 30 min, which implies that the PDDA/SiNPs-PVDF composite membrane became oleophilic if sufficient time was given for the oil-membrane interaction to reach equilibrium. In comparison, the underwater oil CA for the PDA/SiNPs-PVDF membrane was very stable with no observable change in 1 h.

Such a difference between the two composite membranes in their dynamic wetting behavior is most likely attributable to the difference in their surface charges and hence the electrostatic interaction with the crude oil. In fact, the x-DLVO theory accounts for the electrical double layer, van der Waals, and the Lewis acid-based (AB) interactions, two of which depend on surface charge. The x-DLVO theory can in part explain the observed difference in the wetting behaviors between the two composite membranes, as the electrostatic interaction between the PDA/SiNPs-PVDF membrane and the oil droplet was repulsive and that for between the PDDA/SiNPs-PVDF membrane and the oil droplet was attractive. More detailed discussion will be given in Sections 4.1 and 4.2 regarding the applicability of the x-DLVO theory in explaining the oil-membrane interaction.

3.2. Properties of the saline crude oil emulsion

The crude oil emulsion (1000 ppm wt%), prepared as oil feed solution, was highly turbid (Fig. 4(A) inset). The emulsion was stable without observable phase separation for at least 24 h. Such kinetic stability of the emulsion can be attributed to the presence of naturally occurring emulsifier such as asphaltenes, naphthenic acid, carboxylic acid, and other oil-soluble acids and bases in crude oil.

Observation using optical microscopy suggested that the diameters of the oil droplets in the emulsion were in the range of few microns (Fig. 4(A)), which was confirmed by DLS measurements that yielded a distribution of hydrodynamic diameter of 4.9 ± 0.8 µm (Fig. 4(B)). Since the nominal pore size of the PVDF membrane substrate was only 0.45 µm, the oil droplets are expected to effectively block the membrane pores once they attach onto the membrane surfaces. The zeta potential of the oil droplets...
in the emulsion was $-16.4 \pm 0.8$ mV derived from the measured particle electrophoretic mobility. The pH of the prepared feed solution was measured to be between 5 and 6, probably due to the various amphiphilic acids present at the water-oil interface.

3.3. Membrane performance in MD experiments

DCMD experiments with the saline crude oil emulsion using the pristine PVDF membrane and the two composite membranes showed drastically different fouling behaviors (Fig. 5). In each experiment, the conductivity of the distillate was observed to be decreasing with more feed water recovered, yielding a salt rejection rate of over 99.9% for every membrane tested. The fouling of the pristine PVDF membrane was very fast as indicated by drastic decline of water vapor flux in a short period of time (green triangles in Fig. 5). In fact, the water flux dropped to almost zero in only about two hours. Since the PVDF membrane was in-air hydrophobic and underwater oleophilic as suggested by the CA measurements, the very strong long-range hydrophobic-hydrophobic interaction between the crude oil droplets and the membrane surface promoted the retention of oil droplets on the membrane surface and resulted in blockage of membrane pores that served as the pathways for water vapor flux.

In comparison, the DCMD performance of the PDA/SiNPs-PVDF composite membrane was appreciably more stable. With the same crude oil emulsion that severely fouled the pristine PVDF membrane, the PDA/SiNPs-PVDF composite membrane was able to maintain a nearly constant water vapor flux along the 12 h MD operation (blue squares in Fig. 5). The more robust MD performance of the PDA/SiNPs-PVDF composite membrane can be explained by the underwater superoleophobicity of the composite membrane as suggested by its underwater oil CA. The strong hydration force due to the presence of the highly hydrophilic PDA polymer, and the electrostatic repulsion between the nanocomposite and the oil droplets, both contributed to mitigating the adhesion of the oil droplets to the membrane surface.

The PDDA/SiNPs nanocomposite coating slowed down the membrane fouling as reflected by the appreciably slower flux decline of the water vapor flux (red circles in Fig. 5) as compared to that with the pristine PVDF membrane. However, the fouling control was significantly less effective with the PDDA/SiNPs nanocomposite than with the PDA/SiNPs nanocomposite. The difference in anti-fouling performance between these two nanocomposite membranes is in good accordance with the results observed in underwater oil CA measurements. The initial underwater superoleophobicity of the PDDA/SiNPs-PVDF membrane, which was a result of the repulsive hydration force, contributed to the slower flux decline as compared to the pristine PVDF membrane. However, the PDDA/SiNPs-PVDF membrane gradually lost its oleophobicity possibly due to electrostatic attraction, which eventually led to significant membrane fouling within a relatively short time. In comparison, the PDA/SiNPs-PVDF was able to sustain a much more robust MD performance due to the presence of both hydration force and electrostatic repulsion.

The initial fluxes of the PDDA/SiNPs-PVDF composite membrane (23.6 LMH) and PDA/SiNPs-PVDF composite membrane (24.3 LMH) were lower than that of the pristine PVDF membrane (31.0 LMH). The flux reduction was obviously consequent of the surface modification that blocked a significant portion of the membrane. On the other hand, the percentages of flux reduction...
were clearly not in proportion to the fraction of pore blockage shown in the SEM images. Because the nanocomposite layers were porous hydrophilic networks, the liquid feed water should be able to permeate through layer with limited resistance. We therefore speculate that the reduction of vapor flux was mostly attributable to the enhanced temperature polarization in the coating layer. The hydrated nanocomposite coating is a hydraulically stagnant layer insusceptible to the cross-flow induced mixing in the feed channel [2,67,68]. Similar to the internal concentration polarization in forward osmosis which has a detrimental impact on mass transfer [69], the temperature polarization across the hydrated nanocomposite layer was also “internal” in the sense that it occurred within the membrane structure. This additional temperature polarization served to reduce the driving force for vapor transfer across the MD membranes.

3.4. Force spectroscopy with oil droplet probe

To acquire more fundamental understanding about the interaction between the oil droplet and the membrane surface and to explain the different fouling behaviors with different membranes, we conducted force spectroscopy with an oil droplet as the force probe and water as the medium. The position dependent interaction forces between an oil droplet and different membrane surfaces are presented in Fig. 6 for comparison. A complete interaction curve includes two stages, the advancing and receding stages, and five distinct events, including “approach”, “contact”, “retract”, “split” and “detach”. The details for general interpretation of the force curves were given in our previous publication [18].

Before the oil droplet force probe contacted the surface, the “adhesion force” was set to zero by default, as there was no physical interaction between the oil droplet and the membrane. Upon contact between the oil droplet force probe and the pristine PVDF membrane, a strong and sudden attraction was detected. This strong attraction can be attributed to the hydrophobic interaction between the oil droplet and the PVDF membrane [15,16,70]. As the force probe retracted, it experienced an attractive force until the split of the oil droplet. The oil droplet split was reflected on the force curve as a precipitous drop in the “adhesion force”.

A finite “adhesion force” was measured after the split of the oil droplet, which was an artifact resulting from a smaller oil droplet remaining on the force probe. The new “baseline” was non-zero because the difference between buoyancy and gravity for the remaining oil droplet was smaller than that of the original oil droplet [18]. However, there was no physical oil-membrane interaction or “adhesion” corresponding to the finite “adhesion force”, because the oil droplet had already detached from the surface. Although such an “adhesion force” measured by the tensiometer was an artifact in essence, its magnitude positively correlates with the size of the oil droplet remaining on the force probe, which still yields a useful piece of information regarding the affinity between oil droplet and the membrane surface.

For the PDA/SiNPs-PVDF composite membrane, no strong and sudden attraction between the oil droplet and the membrane was observed. On the contrary, a slight repulsive interaction was detected as suggested by a negative “adhesion force” measured upon the contact between the oil droplet and the composite membrane. We did not attempt to quantify such a repulsive force by further pressing the oil probe against the membrane, as the ring-shaped force probe could be forced into the compressed oil droplet and generates irrelevant interaction information. Therefore, the force probe was retracted almost instantly after the oil droplet contacted the membrane.

Nonetheless, the repulsion between the oil droplet and the PDA/SiNPs-PVDF composite membrane upon their first contact was conspicuous from the recorded force curve. The retraction of the force probe resulted in attractive interaction between the oil droplet and the membrane as evidenced by the increasing adhesion force during the receding stage, before the oil droplet split. However, such an adhesion force was very small for the PDA/SiNPs-PVDF composite membrane, as revealed by the force curve (blue curve in Fig. 6). Furthermore, the remaining part of the force curve after oil droplet split only shifted slightly from the original baseline, indicating that only a small proportion of the oil droplet remained on the membrane surface.

The shape of the force curve for the interaction of an oil droplet with the PDDA/SiNPs-PVDF composite membrane resembled that for the interaction with the PDA/SiNPs-PVDF membrane. For instance, repulsion was also observed upon the initial contact between the oil droplet and the composite membrane. However, the attractive interaction between the oil droplet and the PDDA/SiNPs-PVDF was significantly stronger than that between the oil droplet and the PDA/SiNPs-PVDF membrane during the force probe retraction. Since the primary difference between the two composite membranes was the surface zeta potential, it is likely that electrostatic interaction played an important role that differentiated two composite membranes in their interactions with the oil droplet probe.

4. Discussion

4.1. Implications of electrostatic interaction

The results from the force spectroscopy can be employed to explain the different fouling behaviors observed with different MD membranes. The pristine PVDF membrane was most susceptible to
fouling as evidenced by the fastest flux decline among three membranes. This can be explained by the instant and very strong attraction between the oil droplets and the PVDF membrane via the strong hydrophobic-hydrophobic interaction.

On the other hand, the other two underwater oleophobic membranes both mitigated membrane fouling due to the presence of the repulsive hydration force. However, the extents to which membrane fouling was alleviated by the two composite membranes were very different due to the different surface charges and consequently different electrostatic interactions. The attractive electrostatic interaction between the positively charged PDDA/SiNPs-PVDF membrane and the negatively charged crude oil particles offset the repulsive hydration force, leading to a membrane fouling rate between that with the hydrophobic and underwater oleophilic PVDF membrane and that with the underwater oleophobic and negatively charged PDA/SiNPs-PVDF membrane.

The response for the wetting-based interaction seemed to be much faster than that for the electrostatic interaction. This phenomenon was observed in both the underwater oil CA measurements and the oil droplet force probe spectroscopy. In the underwater oil CA measurement using the PDDA/SiNPs-PVDF membrane, the initial underwater oil CA was very high due to the in-air hydrophilic surface coating of the PDDA/SiNPs. The gradual transition of the membrane surface from being underwater superoleophobic to underwater oleophilic can be attributed to electrostatic interaction which did not take effect immediately after the initial oil-membrane contact.

Different responses based on the interaction mechanisms were also observed in force spectroscopy. For example, the precipitous attraction upon the first contact between the membrane and the oil droplet was manifest for PVDF membrane but absent for the PDDA/SiNPs-PVDF membrane, which can be explained by their different surface wetting properties. The relatively significant electrostatic interaction between the positively charged PDDA/SiNPs-PVDF membrane and the negatively charged oil droplet did not become effective until the oil droplet force probe was being retracted from the membrane surface.

The experimental evidences from CA measurements, force spectroscopy, and fouling experiments consistently lead to a cogent conclusion that surface potentials (charges) played a critical role, in addition to wettability, in determining the oil-membrane interaction in water. However, electrostatic interaction is a rather broad concept with many manifestations. For example, ionic bonding, electrical double layer interaction, or even the Keesom type of vdW interaction (e.g. hydrogen bond) can all be considered as electrostatic interaction [70]. The exact mechanism of electrostatic interaction relevant to our experimental observations in this study still needs to be explicated. In the following subsections, we will discuss the impact of the surface potentials on the oil-membrane interaction from both the DLVO and the interfacial tension perspectives.

4.2. Mechanism of electrostatic interaction

When particle-membrane interaction (colloidal deposition) is involved, the DLVO theory has been often cited for theoretical explanation [71–74]. The DLVO theory states that the affinity between a particle and a surface is primarily determined by the electrical double layer (EDL) interaction and the van der Waals (vdW) interaction. The attachment efficiency, defined as the probability for successful attachment per collision, has been employed to quantify such an affinity [74].

However, the applicability of the DLVO theory, for explaining the experimental results of membrane fouling in our study, is questionable. One major problem is that the EDL interaction, which comprises both osmotic and Coulombic contributions [70,75], should theoretically have negligible impact on the overall interaction due to the very short Debye length (calculated to be \(0.4\,\text{nm}\) for the feed solution used). Within such a short Debye length, the vdW force should dominate the overall interaction and eclipse any difference in the EDL interaction. Therefore, the DLVO interaction was unlikely the manifestation of the differential electrostatic interactions between the PDA/SiNPs-PVDF and PDDA/SiNPs-PVDF membranes that led to drastically different fouling behaviors.

To account for the impact of surface wetting properties, the extended DLVO (x-DLVO) theory has often been cited to quantify the affinity between a particle and a membrane surface [76]. The x-DLVO theory incorporates the Lewis acid-base (AB) interaction that accounts for the wetting property based short-range interactions such as hydration force. The short range vdW and AB interactions contribute to the interfacial tension and the free energy of adhesion that not only determine how easy a particle can attach onto the membrane surface (i.e. the attachment efficiency) but also how readily a particle can detach from the membrane surface upon local perturbation.

According to van Oss, the AB component of the interfacial tension involves interaction between electron acceptors and electron donors and only takes effect when the two interacting surfaces are both polar [76–78]. The surface charges of the membranes and the oil droplets have significant impact on the electron donor and electron acceptor parameters, which can explain the drastically different wetting, adhesion, and fouling behaviors between the positively charged PDDA/SiNPs-PVDF membrane and the negatively charged PDA/SiNPs-PVDF membrane. In other words, the electrostatic interaction most likely exerted its impact via the polar interaction component in surface adhesion, rather than the EDL interaction.

However, as free energy of adhesion is a thermodynamic property, it still cannot explain the dynamic behaviors of the PDA/SiNPs-PVDF membrane in the CA measurement and force spectroscopy. We speculate that such dynamic wetting behavior involves the redistribution of natural anionic surfactants at the interface between crude oil and water. These anionic surfactants have lateral mobility along the interface and thus can migrate under the influence of electric field. When the oil droplet was in contact with a positively charged surface, these anionic surfactants gradually concentrated toward the contact area and thereby increased the local electronegativity, which resulted in a higher electron donor parameter of the oil surface and thus a stronger attractive AB interaction between the oil droplet and the membrane.

5. Conclusions

In this study, we tested a hydrophobic PVDF membrane, a PDA/SiNPs-PVDF composite membrane with a hydrophilic and negatively charged surface, and a PDDA/SiNPs-PVDF composite membrane with a hydrophilic and positively charged surface in membrane distillation experiments with a saline crude oil emulsion. While the fouling resistance of the PVDF membrane was unexpectedly poor, both composite membranes were effective in fouling control, but to different extents. The negatively charged PDA/SiNPs-PVDF membrane was highly effective and robust in mitigating the fouling of the negatively charged crude oil emulsion; whereas the positively charged PDDA/SiNPs-PVDF membrane only offered limited improvement over the hydrophobic PVDF membrane. The difference in the fouling behaviors between the three membranes was, to a high degree, corroborated by results obtained from underwater oil CA measurement and from oil droplet probe force spectroscopy.
Dynamic behaviors of the wetting and adhesion were observed when the negatively charged crude oil droplet interacted with the hydrophilic but positively charge PDIA/SiNPs-PVD membrane. The fundamental mechanism underlying this dynamic behavior remains uncertain. We also the relative irrelevance of the colloid deposition approach and the strong relevance of the adsorption energy approach for elucidating the impact of electrostatic interaction on oil-membrane interaction. One possible explanation for the dynamic wetting and adhesion behaviors was the redistribution of charge bearing surfactants at the oil water interface under the electrostatic influence of the oppositely charged membrane, which led to increasing attractive interaction between the crude oil droplet and the PDIA/SiNPs-PVD membrane.

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Appendix A. Supplementary material

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References
