Critical Issues and Guidelines to Improve the Performance of Photocatalytic Polymeric Membranes

Marta Romay, Nazely Diban, Maria J. Rivero, Ane Urtiaga and Inmaculada Ortiz *

Department of Chemical and Biomolecular Engineering, ETSIIyT, University of Cantabria Avda, Los Castros s/n, 39005 Santander, Spain; romaym@unican.es (M.R.); dibann@unican.es (N.D.); mariajose.rivero@unican.es (M.J.R.); urtiaga@unican.es (A.U.)
* Correspondence: ortizi@unican.es; Tel.: +34-942-201-585

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Abstract: Photocatalytic membrane reactors (PMR), with immobilized photocatalysts, play an important role in process intensification strategies; this approach offers a simple solution to the typical catalyst recovery problem of photocatalytic processes and, by simultaneous filtration and photocatalysis of the aqueous streams, facilitates clean water production in a single unit. The synthesis of polymer photocatalytic membranes has been widely explored, while studies focused on ceramic photocatalytic membranes represent a minority. However, previous reports have identified that the successful synthesis of polymeric photocatalytic membranes still faces certain challenges that demand further research, e.g., (i) reduced photocatalytic activity, (ii) photocatalyst stability, and (iii) membrane aging, to achieve technological competitiveness with respect to suspended photocatalytic systems. The novelty of this review is to go a step further to preceding literature by first, critically analyzing the factors behind these major limitations and second, establishing useful guidelines. This information will help researchers in the field in the selection of the membrane materials and synthesis methodology for a better performance of polymeric photocatalytic membranes with targeted functionality; special attention is focused on factors affecting membrane aging and photocatalyst stability.

Keywords: membrane functionality; persistent organic pollutants; photocatalytic membranes; photocatalytic membrane reactor (PMR); composite polymeric membranes; wastewater treatment

1. Introduction

The wide use of chemicals in our society contributes to the accumulation of a huge amount of pollutants in the environment. Many of these recalcitrant pollutants listed in the Stockholm Convention on Persistent Organic Pollutants (POPs) [1] are persistent in the environment due to their strong and stable structure; therefore, conventional technologies of wastewater treatment are not able to degrade or remove them [2].

Advanced oxidation processes are based on the degradation of organic or inorganic contaminants in water and wastewater through oxidation reactions. Among all, heterogeneous photocatalysis is a promising technology in the field of environmental applications because it is driven by UV or visible light [3,4]. Photocatalysis needs a semiconductor material that is excited mainly with UV light. Oxidant species produced during the photocatalytic process can attack the pollutant and break the molecule in smaller compounds. The main advantages of the photocatalytic technology are (i) low operational and installation costs and null post-treatment cost as there is no sludge production except in suspended systems, (ii) it is a technology capable of degrading non-biodegradable pollutants, and (iii) advances in the development of new composite photocatalysts, by doping the primary semiconductor with a co-catalyst, could open up the possibility to use low-cost
visible light. However, among certain drawbacks it is worth mentioning the short service life and high power consumption of the light source, and the requirement of a facility for the photocatalyst recovery in suspended systems [3].

Hybrid processes, such as photocatalytic membrane reactors, PMRs, integrate the activity of the photocatalyst and membrane separation in the same device, this configuration shows great potential to effectively solve the separation problem of harsh systems and the recovery of the catalyst particles in wastewater treatment. Thus, PMRs not only maintain the advantages of photocatalyst technology for degrading high concentration of refractory organic wastes, but they also have the benefit of non-selectivity, fast reaction speed, and complete degradation [5].

Furthermore, the potential harmful effects of nanoparticles are still under debate. If they are not correctly handled and they are released to the environment, they could interact with living organisms. They have similar dimensions to biological molecules such as proteins, so they can enter the human body. Small concentrations of about 5–50 μg/L may cause physiological changes, chromosomal alterations and oxidative stress [6]. Photocatalyst immobilization on membranes can significantly contribute to avoid nanoparticles hazards.

Different classifications have been followed to better characterize PMRs [7–9]. The first one distinguishes between the catalyst being suspended in solution (SPMR) and the catalyst being immobilized on the membrane (IPMR). SPMR primarily aimed at catalyst separation and recovery. The membrane can be submerged in the photocatalytic reactor, Figure 1A [10]. This configuration fails from protecting the membrane for UV irradiation after long operation times (Figure 1A1), although this effect could be reduced with the use of a light-tight baffle in between the light source and the membrane (Figure 1AII). Alternatively, the membrane can be placed out of the photo-reactor, Figure 1B [10], but this configuration is prone to higher pressure drops in the catalyst flow and instability of the catalyst concentration inside the reactor. Otherwise, the catalysts can be fixed to the membrane, IPMR. In this configuration, the membrane can act as support with the only function of immobilizing the catalyst (Figure 1C), or the membrane can integrate the catalyst and simultaneously have a separation function, Figure 1D,E [10]. In this latter system, the membrane acts both as selective barrier for the contaminants to be degraded, thus maintaining them into the reaction environment, and as the support for the photocatalyst. Here, two different flow configurations are possible, dead-end, Figure 1D; or cross-flow, Figure 1E [7]. External illumination is usually applied for these configurations [5,10].

IPMR configuration is raised as an advanced solution. Although suspension photocatalysis results in great pollutant degradation yields due to the large surface area of the nano-size particles, this is at the same time the main drawback that makes difficult the catalyst separation and recovery. Moreover, nanoparticles tend to agglomerate which can lead to a reduction of the reaction yield. This could be somehow avoided if they are properly fixed to some supporting material as it is promoted in IPMRs. In addition, membrane fouling in SPMRs due to photocatalyst deposition is an additional problem. In line with a process intensification philosophy, photocatalyst immobilization can avoid the necessity of the photocatalyst post-recovery stage. Furthermore, it could simultaneously overcome fouling phenomena associated with the photocatalyst and pollutants [4,5,8].
In advanced IPMRs, the role of the membrane is to perform as a barrier to retain the pollutants as well as photocatalyst support. The membrane support can be made of ceramic, metallic, or polymeric materials [8]. Although ceramic and metallic membranes have great chemical stability and high mechanical strength, their applications are limited because of the manufacturing costs [4,8]. Taking into account the economic factor, polymeric membranes are cheaper and easier to apply on an industrial scale in spite of their limitations. Moreover, they are easy to process, and their properties and microstructure can be tuned. The selection of the polymeric membrane material is not trivial. Due to the nature of polymers as long-chain carbon molecules, they are susceptible to UV light and oxidative species degradation that can damage the membrane during the photocatalytic process. The challenge is to develop low-cost polymeric membranes with adequate mechanical strength and flexibility, which withstand UV and oxidative conditions [4] while supporting the photocatalyst particles. Furthermore, the conformed polymeric membrane should prevent the pollutant from crossing the membrane before photocatalytic degradation while still performing with sufficiently high total flux.

Previous reviews [11–14] have addressed the most recent advances on membrane synthesis and applications and have detected major challenges that need to be overcome to develop technically competitive IPMRs for future scaling-up and expansion of the technology. One of these challenges results from the intrinsic nature of an immobilized nanoparticle system. The photocatalyst can be immobilized either in the membrane matrix or on the membrane surface as a coating layer. The method of photocatalyst immobilization importantly affects the photocatalyst distribution in the membrane, its stability and accessibility to light-source. Therefore, a rigorous and methodological analysis of how the immobilization method of the photocatalyst in the membrane affects all those parameters needs to be done to ascertain their impact on the overall performance of the functional membrane, in terms of flux properties and photocatalytic activity [4].

This is the first attempt reported in the literature to critically review the information on photocatalytic polymeric membranes so far applied to the treatment of waters and wastewaters polluted with POPs or model organic pollutants; the analysis has been focused to identify the influence of (i) the polymer and photocatalyst selection and, (ii) the methodology of membrane...
synthesis, on the functional properties of the resulting membranes. As a major novelty, this review is conceived to not only point out the major challenges collected after reviewing the literature, but it also extracts the reasons of current failures and proposes guidelines and solutions to overcome the reported limitations for the successful development of photocatalytic polymeric membranes. Thus, this review aims at becoming a practical guide for scientists in the field of photocatalytic membranes to decide the best methodological approach to deal with the synthesis of new functional photocatalytic membranes and to detect possible experimental gaps that future research would need to fill.

2. Materials in Polymeric Photocatalytic Membranes

Tables 1–3 show a literature survey of the porous photocatalytic polymer membranes synthesized so far. Tables 1 and 2 collect the photocatalytic membranes classified as mixed matrix membranes (MMM), considering that the photocatalyst is dispersed in a polymeric matrix. Table 1 presents the membranes synthesized by phase inversion methods and Table 2 presents the membranes synthesized by electrospinning. On the other hand, Table 3 collects the works that propose different synthesis techniques of thin-film composite membranes (TFCM). Moreover, the materials employed (i.e., polymer, non-solvent, additive, and photocatalyst), components composition and the application evaluated in each study are presented. These data will be analyzed to acquire knowledge to define the guidelines that will help to decide which is the adequate material and processing technique for the desired application.
## Table 1. Mixed-matrix membrane systems synthesized by phase inversion: materials, compositions, and applications.

| Polymer (wt. %) | Solvent | Non-Solvent | Additive (wt. %) | Photocatalyst | wt.% (Polymeric Solution) | wt. %/cm² (Membrane) | Application (Removed Pollutant) | Author |
|-----------------|---------|-------------|-----------------|---------------|---------------------------|----------------------|--------------------------------|--------|
| PVDF (20)       | DMAC    | deionized water | PEG (5)         | TiO₂          | 0–7, 5 *                 | 0–0.612, 0.478 *     | BSA                            | Méricq, J.P [15] |
| PVDF (20)       | DMAC    | deionized water | PEG (5)         | TiO₂          | 4                        | 0.30                 | (-)                            | Tran, D. [16]    |
| PVDF (19)       | DMAC    | distilled water | PEG (7)         | Ag-TiO₂       | 0.01-0.06, 0.06 *        | (-)                  | BSA, E.Coli                   | Chen, Q. [17]   |
| PVDF (18)       | DMAC    | deionized water | PEG (2)         | TiO₂          | 0–1, 1 *                 | 0–0.33, 0.33 *       | Estrone, 17β-estradiol          | Wang, M. [18]   |
| PVDF (16)       | DMAC    | water: isopropanol (70:30) | (-) | TiO₂ | 0–3, 2* | 0–0.04, 0.027 | RTB | Sakarkar, S. [19] |
| PVDF (15)       | DMAC    | Water         | PVP (1)         | GO-TiO₂       | 1                        | 0.32                 | BSA                            | Xu, Z [20]       |
| PVDF (15)       | DMAC    | distilled water | PVP (1)         | GO-OMWCNTs    | 1% (carb/pol)            | 0.106                | BSA                            | Zhang, J. [21]  |
| PVDF (14)       | DMAC    | tap water     | (-)             | AC-TiO₂       | (0–0.5)–(0–0.1)          | 0.042–0.297,0.083 *  | NOM, HA                       | Liu, Q. [22]    |
| PVDF (12)       | DMAC    | tap water     | PEG (1–5), 2 *  | TiO₂          | 0.25–2, 0.5 *            | 0–0.23, 0.083 *      | NOM, HA                       | Song, H. [23]   |
| PVDF (12)       | DMAC    | Tap water     | LiCl (0–4), 0.5 | TiO₂          | 0–1.5, 0.5 *             | 0–0.33, 0.33 *       | NOM, HA                       | Song, H. [24]   |
| PVDF-PDMA (12)  | TEP     | Water         | PEG (25) + PEG (5) | TiO₂ | 0–0.5, 0.5 * | 0–0.5, 0.5 * | MB | Benhabiles, O. [25] |
| PVDF (10)       | NMP     | tap water     | (-)             | TiO₂          | 0–4, 2–4 *               | 0–0.189, 0.110–0.189 | E. Coli, RB5, BSA | Damodar, R.A. [26] |
| PMMA-g-PVDF/PAN (-) | DMAC | Ethanol: deionized water (1:3) | (-) | N-TiO₂ | 1,3,5 | (+) | Bentazon | Mungondori, H. [27] |
| PSF (20)        | NMP     | distilled water | (-)             | TiO₂          | 0–2.43, 1.96 *           | 0–0.553, 0.455 *     | Cr(VI)                         | Jyothi, M.S. [28] |
| PSF (18)        | DMAC:NMP | Ethanol:water (20:80) | (-) | TiO₂ | 0–5, 2 * | (+) | BSA | Yang, Y. [29] |
| PSF (18)        | DMAC:NMP | deionized water | PEG (8)         | Fe-TiO₂       | 0–4.5, 3.6 *             | 0–0.077              | BPA                            | Wang, Q. [30]   |
| PSF (18)        | DMAC    | Water         | (-)             | TiO₂-ZnO, TiO₂-SiO₂ | 0.16 | (+) | MO, phenol | El-Aassar, A. [31] |
| PSF (18)        | NMP     | deionized water | (-)             | N, Pd-TiO₂    | 0–1.26, (0–7)            | (+)                  | EY                            | Kuvarega, A.T. [32] |
| PSF (18)        | NMP     | Water         | PVP (2)         | N-rGO-TiO₂    | 0.5                      | 0.095                | DR 80, DB 15                  | Xu, H. [33]     |
| Catalysts | Volume | Year | Pages | Title | Authors | Reference |
|-----------|--------|------|-------|-------|---------|-----------|
| PSF (17)  | NMP    | tap water | PVP (0.5) | mpg-C3N4-TiO2 | 0–1, 1 * | 0–0.653, 0.653 * | SMX | Yu, S. [34] |
| PSF (12)  | DMF    | distilled water | (-) | CNTs-TiO2 | 1 | (-) | Amoxicillin | Muhulea, A. [35] |
| PES (27 g) | DMF, EtoH (1–4) | distilled water | (-) | TiO2 | 0.1–0.4, 0.1 * | 0.0062–0.024, 0.0062 * | HA | Sotto, A. [36] |
| PES (26)  | NMP    | Water | (-) | Co-TiO2 | 0.5–1, 1 * | 0.065–0.129, 0.129 * | 2-DCP | Hoseini, S.N. [37] |
| PES (21)  | DMAC   | distilled water | PVP (1) | rGO-TiO2 | 0.05–0.2, 0.1 * | 0.237–0.943, 0.497 * | DY 12, RG 19, RB 21, BSA | Safarpour, M. [38] |
| PES (20)  | DMAC   | distilled water | PVP (1) | B-TiO2-SiO2/CoFe2O4 | 0–1, 0.5 * | 0–0.497, 0.239 * | DR 16, POME | Zangeneh, H. [39] |
| PES (18)  | NMP    | tap water | SMM (1) | O-g-C3N4 | 1 | (-) | Phenol | Salim, N. [40] |
| PES (18)  | NMP    | tap water | SMM (1–5, 4 *) | O-g-C3N4 | 1 | (-) | Phenol | Salim, N. [41] |
| PES (15)  | DMAC   | Water | PVP (5) | mNi-TiO2 | 0–1, 1 * | 0–0.083, 0.083 * | BSA, YEF, SA, HA, MB | Sun, T. [42] |
| PES-F-COOH (20) | DMF | deionized water | PVP (10) | TiO2 | 1–5, 5 * | 0.123–0.519, 0.519 * | PAM | Geng, Z. [43] |
| CA-PS (-) | Acetone | distilled water | (-) | ZnO | 0.1 g | (-) | CR, RY 105 | Rajeswari, A. [44] |
| CA-PU (-) | Acetone-chloroform | distilled water | (-) | ZnO | 0.3 | (-) | RR 11, RO 84 | Rajeswari, A. [45] |
| P(VDF-TrFE) (10) | DMF | (-) | (-) | TiO2 | 8 | 0.32 | MB, CIP, IBP | Martins, P.M. [46] |
| P(VDF-TrFE) (10) | DMF | (-) | (-) | TiO2 | 8 | 0.017 | Tartrazine | Aoudjit, L. [47] |
| P(VDF-TrFE) (10) | DMF | (-) | (-) | TiO2, ZnO | 0–15,15 * | 0–3.75, 3.75 | MB | Teixeira, S. [48] |
| P(VDF-TrFE) (10) | DMF | (-) | (-) | TiO2 (NaY) | 0–8, (0–8), (0–8) * | 0–3.70, 3.70 * | MB | Martins, P.M. [49] |
| PVDF-HFP (15) | DMF | (-) | (-) | Ag-TiO2 | 0–10, 10 * | 0–0.56, 0.56 * | NOR | Salazar, H. [50] |
| PSF (16)  | NMP    | distilled water | PVP (2) | TiO2, MIP TiO2, NIP TiO2 | 2 | (-) | MB, MO | Melvin, H.K. [51] |
| PVDF Dul layer HF (18 in, 15 out) | DMAC | tap water | (-) | TiO2 | 0 (in) 3 (out) | (-) | 8 pharmaceutical mixture | Paredes, L. [52] |
| PVDF Dual layer HF (18 in, 15 out) | DMAc | tap water | (-) | TiO₂ | 0 (in) 3–15 (out), 3 * | 0 (in) 0.067–0.201 (out), 0.067 * | NOM | Dzinun, H. [53] |
|----------------------------------|------|-----------|-----|------|----------------------|---------------------------------|-----|----------------|
| PVDF Dual layer HF (18 in, 15 out) | DMAc | tap water | (-) | TiO₂ | 0 (in) 3 (out)       | 0 (in) 0.067 (out)               | NP  | Dzinun, H. [54] |
| PVDF Dual layer HF (18 in, 15 out) | DMAc | tap water | (-) | TiO₂ | 0 (in) 0–15 (out), 15* | 0 (in) 0–0.201 (out), 0.201 *   | NP  | Dzinun, H. [55] |
| PVDF Dual layer HF (18 in, 15 out) | DMAc | tap water | (-) | TiO₂ | 0 (in) 3 (out)       | 0 (in) 0.067 (out)               | NP  | Dzinun, H. [56] |
| PVDF HF (18–19) | NMP | tap water | PEG (5 in, 0 out) | TiO₂ | 0.5 | 0.071 | MB | Galiano, F. [58] |
| PVDF HF (18) | DMAc | Water | PVP (5) | TiO₂ | 0–4, 2 * | (-) | Oil wastewater | Ong, C.S. [59] |
| PVDF (18) | DMAc, NMP, DMF | Water | (-) | TiO₂ | 0.001, 0.01, 0.1 g/L | (-) | HA | Teow, Y.H [60] |
Table 2. Mixed-matrix membrane systems synthesized by electrospinning: materials, compositions and application.

| Polymer (wt. %) | Solvent          | Additive | Photocatalyst          | wt. % (Polymeric Solution) | Application (Removed Pollutant) | Author          |
|----------------|------------------|----------|------------------------|-----------------------------|---------------------------------|-----------------|
| P(VDF-TrFE)    | DMF/MEK, 85/15   | (-)      | GO-TiO₂                 | 0–20, 5*                    | MB                              | Almeida, N.A. [61]|
| PVDF (-)       | (-)              | PEO      | TiO₂-MWCNTs (20:1)      | 0–40                        | HA                              | Chen, J. [62]   |
| PTFE: PVA (6:1)| water            | PVA      | ZnO                    | 0–30, 20*                   | RhB                             | Huang, Y. [63]  |
| PTFE (15)      | (-)              | PVA (1), BA (0.0025) | TiO₂ (-) | 0.9 | MG | Kang, W. [64] |
| PA6 (12)       | AA: FA (2:1)     | (-)      | TiO₂                  | 25                           | E. coli, RBB | Blanco, M. [65] |
| PAN (8)        | DMF              | (-)      | TiO₂-ZnO              | 2                            | MO                              | Yar, A. [66]    |
| PAN (6)        | DMF              | (-)      | ZnO                    | 0.9                          | MO                              | Tissera, N. D. [67] |
| PAN (7)        | DMF              | (-)      | TiO₂                  | 3.57                         | Nitrate                         | Suriyaraj, S.P. [68] |

* optimal concentration. Abbreviations: Polymers: cellulose acetate (CA), polyamide 6 (PA6), polyacrylonitrile (PAN), polyethersulfone (PES), polystyrene (PS), polysulfone (PSF), polytetrafluoroethylene (PTFE), polytrifluoroethylene (TrFE), polyurethane (PU), polyvinylidene fluoride (PVDF), PVDF-co-hexafluoropropylene (PVDF-HFP); Solvents: acetic acid (AA), dimethylacetamide (DMAc), dimethylformamide (DMF), ethanol (EtOH), formic acid (FA), methyl ethyl ketone (MEK), N-methylpyrrolidone (NMP), triethyl phosphate (TEP); Photocatalysts: activated carbon (AC), silver (Ag), boron (B), bismuth (Bi), carbon nanotubes (CNTs), cobalt (Co), cobalt ferrite (CoFe₂O₄), carbon quantum dots (CQDs), iron (Fe), graphene oxide (GO), molecular imprinting polymer (MIP), mesoporous graphitic carbon nitride (mpg-C₃N₄), multi-walled carbon nanotubes (MWCNTs), nitrogen (N), sodium Y zeolite (NaY), magnetic nickel (mNi), non-imprinting polymer (NIP), oxygen doped graphitic carbon nitride (O₃g-C₃N₄), oxygenated MWCNTs (OMWCNTs), palladium (Pd), reduce graphene oxide (rGO), silicon oxide (SiO₂), titanium dioxide (TiO₂), zinc oxide (ZnO); Additives: boric acid (BA), lithium chloride (LiCl), polyethylene glycol (PEG), polyethylene oxide (PEO), polyvinyl alcohol (PVA), polyvinyl pyrrolidone (PVP), surface molecule modifier (SMM); Pollutants: 2,4 dichlorophenol (2-DCP), bovine serum albumin (BSA), bisphenol A (BPA), ciprofloxacin (CIP), congo red (CR), direct blue (DB), direct red (DR), direct yellow (DY), Escherichia coli (E. Coli), eosin yellow (EY), humic acid (HA), ibuprofen (IBP), indigo carmine (IC), methylene blue (MB), malachite green (MG), methylene orange (MO), Natural Organic Matter (NOM), norfloxacin (NOR), nonylphenol (NP), polyacrylamide (PAM), palm oil mill effluent (POME), reactive black 5 (RB5), reactive blue 21 (RB21), Remazol black B (RBB), reactive green (RG), rhodamine B (RhB), reactive orange (RO), reactive red (RR), remazol turquoise blue (RTB), reactive yellow (RY), ammonium alginate (SA), sulfamethoxazole (SMX), yeast extract fermentation (YEF).
### Table 3. Literature review on thin film composite membranes: synthesis methods, materials, compositions, and pollutant application.

| Synthesis Method                  | Support Material                  | $A_{\text{eff}}$ (cm²) | Photocatalyst       | Photocatalyst Mass (mg) | Photocatalyst per Membrane Area (mg/cm²) | Application (Removed Pollutant) | Author                  |
|-----------------------------------|-----------------------------------|-------------------------|----------------------|-------------------------|------------------------------------------|--------------------------------|--------------------------|
| Vacuum deposition                 | PES (com)                         | 4.3                     | nAg-GO-TiO₂          | 3                       | 0.697                                    | E. coli, B. subtilis           | Jiang, Y. [69]           |
|                                   | Pretreatment: PAAM                |                         |                      |                         |                                          |                                |                          |
| Vacuum deposition                 | CA (com)                          | 12.56                   | Ag-rGO-TiO₂          | 2.5–20, 10 *            | 0.199–1.59, 0.796 *                    | MB, RhB, oil water             | Chen, Q. [70]            |
|                                   | Pretreatment: PEG + GA            |                         |                      |                         |                                          |                                |                          |
| Vacuum deposition                 | CA (com)                          | 12.56                   | rGO-g-C₃N₄           | 10–100, 25 *            | 0.796–7.96, 1.99 *                     | RhB                            | Zhao, H. [71]            |
| Vacuum deposition                 | MCE (com)                         | 1.54                    | GO-TiO₂              | 10                      | 6.49                                    | DP, MO                         | Pastrana-Martínez, L [72] |
| Vacuum deposition                 | PC (com)                          | (-)                     | GO-TiO₂              | (-)                     | (-)                                      | DR 80, DB 15                   | Xu, C. [73]              |
| Vacuum deposition                 | PC (com)                          | (-)                     | GO-TiO₂              | (-)                     | (-)                                      | MO, RhB                        | Xu, C. [74]              |
| Vacuum deposition                 | PAN (8wt. %, DMF)                 | 12.56                   | rGO-α-Fe₂O₃          | (-)                     | (-)                                      | MB, MO, RhB, R6G, MG, GV       | Sun, K. [75]             |
|                                    | (Support: Electrospinning)       |                         |                      |                         |                                          |                                |                          |
| Filtration                        | CA (com)                          | 14.6                    | GO-TiO₂              | 50–400, 100 *           | 3.42–27.4, 6.85 *                      | CR                             | Nair, A.K. [76]          |
| Filtration                        | CA (com)                          | 11.94                   | GO-TiO₂              | 50–300, 200 *           | 4.18–25.12, 16.75 *                    | RhB, AO7                       | Gao, P. [77]             |
| Immersion                         | PSF (com)                         | 17.34                   | AC-N-rGO-TiO₂        | 10–160, 120 *           | 0.57–9.22, 6.92 *                      | MO                             | Wu, T. [78]              |
|                                    | Pretreatment: PVA (2.5mg)         |                         |                      |                         |                                          |                                |                          |
| Immersion                         | CA-GO (15–(0–1.5) wt. %, DMF)    | (-)                     | NH₂-TiO₂             | 0.005 g/ml              | (-)                                      | MB, IC                         | Aboamera, N. M. [79]     |
|                                    | Pretreatment: GA                  |                         |                      |                         |                                          |                                |                          |
| Immersion                         | PAN-CNT (10 wt. %, DMF)           | (-)                     | NH₂-TiO₂             | 40.6 wt. % (CNT-TiO₂)   | (-)                                      | Cr (VI)                        | Mohamed, A. [80]         |
| Method                                       | Material Details                                                                 | Performance | Medium | Source |
|----------------------------------------------|----------------------------------------------------------------------------------|-------------|--------|--------|
| Immersion - In situ growth (Support: Electrospinning) | PAN (9 wt. %, DMF) Pretreatment: PDA                                             | 20          | Ag-TiO$_2$ | (-)    | MB, Phenol | Shi, Y. [81] |
| Immersion (Layer by Layer (LbL))            | PSF (com)                                                                        | (-)         | GO-TiO$_2$ | (-)    | 0.062  | MB | Gao, Y. [82] |
| Immersion (Support: Phase inversion)         | PEI/P25 (24/1.23 wt. %, NMP)                                                    | 12.56       | TiO$_2$ nw | 250    | 19.9   | RhB | Jiang, R. [83] |
| Immersion (Plasmagrafted)                   | PVDF-g-PAA (com)                                                                 | 4.5         | TiO$_2$   | 0.5 1.53 % m/v | (-) | RB5, BSA | You, S.-J. [84] |
| Immersion (UV-grafted)                       | PA-g-PAA (com)                                                                   | 13.4        | Ag-ZnO-FeO$_2$-MWCNTs | 8.7  | 0.649  | Amoxicilin | Irani, E. [85] |
| Electrospraying (Support: Electrospinning)  | PVDF (18 wt. %, DMF:acetone 60:40)                                              | 45          | TiO$_2$   | 4.5-27, 27* | 0.1-0.6, 0.6* | BPA, 4-CP, CMT | Ramasundaram, S. [86] |
| Coaxial electrospinning                     | PAN (10, 15 core) DMac                                                          | (-)         | CQDs-Bi$_{201}$-TiO$_{25}$ | 5, 10, 15 wt/v% | (-) | Isoproturon | Xie, R. [87] |
| Hot pressing (Support: Phase inversion)     | PVDF (-wt. %, DMac)                                                             | (-)         | TiO$_2$   | (-) | (-) | BPA | Nor, N.A.M. [88] |
| Sputtering + Anodization                    | PES (com)                                                                        | 17.35       | TiO$_2$   | (-) | (-) | diclofenac | Fischer, K. [89] |

*Optimal concentration. Abbreviations: Polymers: cellulose acetate (CA), commercial (com), mixed cellulose esters (MCE), polyamide (PA), polyacrylonitrile (PAN), polycarbonate (PC), (polyetheramide (PEI), polyethersulfone (PES), polysulfone (PSF), polyvinylidine fluoride (PVDF); Solvents: dimethylacetamide (DMAc), dimethylformamide (DMF), N-methyl pyrrolidone (NMP). Additives: glutaraldehyde (GA), lithium chloride (LiCl), polyacrylic acid (PAA), polyallylamine (PAAM), polyethylene glycol (PEG), polydopamine (PDA), polyvinyl alcohol (PVA); Photocatalysts: activated carbon (AC), silver (Ag), cobalt (Co), graphitic carbon nitride (g-C$_3$N$_4$), graphene oxide (GO), nitrogen (N), nano-silver (nAg), amine group (NH$_2$), nanowire (nw), polyoxometalate (POM), reduce graphene oxide (rGO), titanium dioxide (TiO$_2$), hematite (α-Fe$_2$O$_3$); Pollutant: 4-chlorophenol (4-CP), acid orange 7 (AO7), Bacillus subtilis (B. subtilis), bisphenol A (BPA), bovine serum albumin (BSA), cimentine (CMT), congo red (CR), chromium VI (Cr VI), crystal violet (CV), direct blue (DB), Diphenhydramine (DP), direct red (DR), Escherichia coli (E. coli), gentian violet (GV), humic acid (HA), indigo carmine (IC), methylene blue (MB), malachite green (MG), Methyl Orange (MO), reactive black 5 (RBS), rhodamine B (RhB), rhodamine 6G (R6G).
2.1. Nano-Photocatalyst

As it can be observed in Tables 1–3, the photocatalytic membranes prepared so far were mostly synthesized using semiconductor materials such as TiO$_2$ [15,16,18,23–26,28,29,36,47,52–60,64,65,68,83,84,86,88,89], ZnO [44,45,48,63,67], g-C$_3$N$_4$ [34,40,71], and Fe$_2$O$_3$ [75]. Most of them are metal oxides such as TiO$_2$, ZnO, and Fe$_2$O$_3$. Titanium dioxide (TiO$_2$) is the most commonly used semiconductor for photocatalytic membranes, due to its chemical and thermal stability, low cost, high reusability, and excellent yield in the degradation of organic pollutants. The main disadvantages of TiO$_2$ are the high electron-hole recombination ratio, and large bandgap (3.2 eV) that implies a low adsorption capacity for visible light [3,90]. To avoid these limitations some strategies such as doping with a co-catalyst are being tested. The introduction of doping agent into the bulk photocatalyst provides a large dipole moment to change the electron transfer kinetic, and more electrons can be transferred from valence band (VB) to conduction band (CB) of the photocatalyst, hence narrowing the E$_g$ value. The low E$_g$ value indicates a better absorption ability in the visible light or natural sunlight. The incorporation of two semiconductors forms a hybrid photocatalyst, where the photo-generated electrons can flow from more negative to less negative fermi energy (E$_F$) in CB, while holes flow from more positive to less positive E$_F$ in VB at the interface to prevent charge recombination. These new materials have been mostly used in slurry-type reactors and when it comes to photocatalytic membranes inorganic supports [91] or self-standing catalysts [92]. Silver [17,50,69,70,81,85] and iron [30,39,75] are the metals most commonly used, but also palladium [32], cobalt [37,39], and magnetic nickel [42] have been tested. Other elements used are nitrogen [27,32,33,78–80] and boron [39].

Similarly as in suspension systems, in immobilized photocatalysis, the combination of the photocatalyst with other materials is a widely used strategy to enhance their photocatalytic properties and to redshift its bandgap. Composite photocatalysts are synthesized combining the semiconductor photocatalyst with other materials. Many authors have reviewed the use of composites based on TiO$_2$ in batch suspension systems [3,93], including composites synthesized with carbonaceous materials (GO-TiO$_2$, rGO-TiO$_2$, MWCNTs-TiO$_2$, AC-TiO$_2$) [94,95] and g-C$_3$N$_4$ based photocatalyst (rGO-g-C$_3$N$_4$, g-C$_3$N$_4$-TiO$_2$) [96]. Readers are referred to the just mentioned thorough reviews to get deeper insight on the photocatalytic activity of those materials, as it is not the objective pursued with the present review. As Tables 1–3 show, the use of composite materials has been extended to the preparation of photocatalytic membranes. Carbon materials that include graphene oxide (GO) [69,73,74,76,77,79,82], reduced graphene oxide (rGO) [20,33,38,61,70–72,75], activated carbon (AC) [22], carbon nanotubes (CNTs) [35,80], multi-walled carbon nanotubes (MWCNTs) [21,62,85], graphitic carbon nitride (gC$_3$N$_4$) [34,40,41,71], and carbon quantum dots (CQDs) [87] have been immobilized in polymeric membranes. Certain carbon materials develop high surface area and in general provide good mechanical properties, together with antimicrobial properties [69].

Additionally to the photocatalytic activity, the introduction of semiconductor nanoparticles in the membrane matrix can also lead to: (i) the enhancement of the membrane hydrophilicity due to the photocatalyst polarity, and (ii) the modification of the membrane morphology during the synthesis. The influence on membrane morphology will be further discussed in the section covering the membrane synthesis methods of the present review. Some authors have observed a so-called superhydrophilicity of polyvinylidene fluoride (PVDF) [15,18,59], polysulfone (PSF) [28,29], polyethersulfone (PES) [97], PES-F-COOH [43], and polyetherimide (PEI) [83] membranes that incorporate TiO$_2$. Superhydrophilicity is defined as an intrinsic property of TiO$_2$ surface that generates a significant increase in water flux under UV or sunlight conditions [15,16,59,98]. The mechanism of TiO$_2$ nanoparticles superhydrophilicity is depicted in Figure 2. As the TiO$_2$ is UV-irradiated, the photo-generated electrons reduce Ti$^{4+}$ to Ti$^{3+}$, and the O$^{2-}$ anions are oxidized to O$_2$ in the photocatalyst holes. Oxygen vacancies are produced on the surface, so the empty sites can be occupied by the water molecules, and OH$^-$ groups are adsorbed on the surface, which increases the surface hydrophilicity [97]. In consequence, the water permeate flux is increased [18,28,29,43]. Composites with ZnO particles also provided improved permeation flux in references [44,45].
To find out the optimum photocatalyst concentration to be immobilized in a membrane is not trivial. Multiple factors can influence this variable as stated in the revised literature, such as the selection of the photocatalyst and the polymer materials, the synthesis method and the membrane macrostructure, among others. As Tables 1–3 show, a wide variety of photocatalyst concentration ranges has been studied, from 0 wt.% to 30 wt.% for similar systems. Particularly, the synthesis method and the processing variables exert an important influence. Therefore, the critical analysis of the optimal nanoparticle concentration observed by different authors will be addressed in the specific membrane synthesis section.

2.2. Polymer

The selection of the polymer matrix material is a key point in the performance of photocatalytic membranes. As it was already mentioned in the introduction, the support material must be resistant to the degradation caused by UV irradiation and by the generated oxidants. Therefore, this section overviews the different polymers reported in the literature to develop photocatalytic membranes, focusing on the attention on their chemical, and particularly, photochemical resistance.

Tables 1–3 show that several polymers have been used to produce photocatalytic membranes. These polymers can be grouped in:

1. Fluorine-based: polyvinylidene fluoride (PVDF), poly(vinylidenefluoride–trifluoroethylene) P(VDF-TrFE), polytetrafluoroethylene (PTFE), and a copolymer of PVDF and hexafluoropropylene (PVDF-HFP)
2. Sulfur-based: polysulfone (PSF), and polyethersulfone (PES)
3. Nitrogen-based: polyacrylonitrile (PAN), polyethylenimine (PEI), polyamide (PA), and polyamide 6 (PA6)
4. Cellulose derivatives: cellulose acetate (CA), and mixed cellulose esters (MCE)
5. Other polymer: polycarbonate (PC)

Among all polymers, PVDF is the most widely employed material for photocatalytic applications in literature, followed by PAN, PES, PSF, and CA. Other polymers such as PTFE, PC, PDA, and MCE are less frequently used. PVDF, P(VDF-TrFE), PTFE, PSF, and PES are hydrophobic polymers while PAN, PEI, CA, MCE, and PC are considered hydrophilic. PVDF, PTFE, PES, PSF, and PAN are polymers usually employed to synthesize mixed matrix membranes because they are easy to process. On the other hand, porous CA, MCE, PC, and PEI membranes are usually employed as supports for thin-film composite photocatalytic membranes.

For mixed-matrix membranes fabrication (Tables 1 and 2), PVDF is the most commonly used polymer [15–18,20–23,25,26,52–60,62,84,86,88] due to the strength and stability of the C-F bond. Other fluorine-based polymers are P(VDF-TrFE) [46] and PVDF-HFP copolymer [50], which presents similar properties than PVDF, and to a lesser extent, PTFE [63,64]. PVDF is commonly used in microfiltration or ultrafiltration membranes as it presents good thermal and chemical resistance [15]. One of the major drawbacks of these fluoropolymers is their high hydrophobicity and therefore low water filtration fluxes. A strategy to improve the wettability of P(VDF-TrFE) membranes was developed by incorporating hydrophilic fillers, such as NaY zeolite type [49].

Sulfonated polymers, i.e., PSF [28–35,51,82] and PES [36–43,69,89] have been widely used to manufacture photocatalytic membranes [28,38]. Both are commercially available polymers with
excellent thermal, chemical, and biological stability, good mechanical properties, and high rigidity. However, due to their hydrophobic nature, membranes made with PSF and PES provide low water flux and fouling problems arise, because of the strong interaction between hydrophobic compounds and the membrane surface. Moreover, sulfur-based polymers have poor resistance to UV irradiance [28–30,32,34,38,39,43], as the C-S bond is weak and unstable under UV light and oxidative conditions [99,100].

PAN is a non-toxic polymer and presents UV and chemical resistance [66,80,81]. It is commercially available and environmentally stable. Moreover, it provides good mechanical properties, particularly its flexibility. PEI, whose chemical formula is (C₆H₁₂O₆N₆)ₙ [83], PA [85], and PA6 also known as nylon 6 [65] are other nitrogen-containing polymers that have been attempted to a lesser extent for preparation of photocatalytic membranes.

CA and MCE are polymers widely used in the preparation of membranes for water purification and biomedical applications due to their good mechanical strength, water affinity and especially their easy availability and low cost. Among their drawbacks, these materials have low chemical and thermal resistance [44,45]; to overcome these disadvantages, CA is combined with other polymers such as polystyrene (PS) [44], polyurethane (PU) [45], or with nanomaterials such as graphene oxide [79].

Finally, PC [73,74] commercial membranes are mostly used as supports, for the case of thin-film photocatalytic membranes.

The resistance to UV light of several polymeric PAN, CA, PVDF, PTFE, PSF, PES, and PC commercial membranes in the presence of TiO₂ under photocatalytic conditions was studied by Chin et al. [99]. Membranes made of PSF, PES, PC, and CA broke before 30 days of exposure to light irradiance. The rest of the materials were tested again by adding hydrogen peroxide. The authors concluded that PVDF and PTFE showed the highest resistance to oxidative conditions. Meanwhile, PAN membranes reported reduced mechanical properties, leading to the breakup of the PAN membrane. However, there is still an important lack of information about how photocatalytic membranes produced by different processing techniques and polymer materials perform under long-term exposures to UV-light. In the following sections, a specific analysis will be directed to address this issue.

3. Membrane Synthesis Method

The membrane morphology and nanoparticles distribution will depend on the synthesis method employed to incorporate the photocatalyst in the membrane. Below, a thorough analysis of the influence of processing variables on membrane morphology will be performed. The objective pursued is to establish processing guidelines to attain the desired membrane morphology.

3.1. Mixed Matrix Composite Membranes Synthesis Methods

As it is shown in Tables 1 and 2, two different synthesis methods have been reported to immobilize photocatalyst nanoparticles embedded in the polymer matrix: (i) phase inversion and (ii) electrospinning, respectively. For both techniques, the photocatalyst nanoparticles are dispersed in the polymer solution before the membrane synthesis; thus, during precipitation of the polymer photocatalyst nanoparticles remain embedded into the polymer matrix.

3.1.1. Phase Inversion

Phase inversion is the mechanism that takes place in the membrane formation following different polymer coagulation routes such as i.e., Non-solvent Induced Phase Separation (NIPS) or Evaporation Induced Phase Separation (EIPS) [101]. These techniques applied to classical systems of three components can be extrapolated to quaternary systems. These techniques produce two configurations: flat membranes and hollow fibers, that have been reported in the literature as indicated in Table 1. Flat photocatalytic membranes are simple to produce and very useful for laboratory-scale evaluation [101], and mainly incorporate bare TiO₂ or TiO₂ composites combined
with other substances. However, the large surface area provided by hollow fibers might be required to accomplish scalable technologies. Hollow fibers are made by extrusion techniques [52,58,59]. In particular, co-extrusion allows to create two membrane layers with different functional characteristics, the inner part acting as support and the outer part that contains TiO₂ nanoparticles and gives to the membrane photocatalytic and/or antifouling function [52–57].

When the membrane is synthesized by NIPS, a solution formed by a polymer dissolved into an organic solvent and blended with the photocatalytic nanoparticles is forced to demix via its introduction into a bath of a non-solvent, also known as coagulant. At this moment, there is an exchange between the solvent and non-solvent, which must be miscible. The polymer solution separates in two phases at equilibrium, one of them is rich in polymer (solid containing the photocatalyst) and the second one is poor in polymer (liquid). The thermodynamics of this process is explained through the Flory–Huggins theory based on Gibbs free energy equations and interaction parameters of the components. Figure 3 shows an exemplary diagram with the main thermodynamic elements and a typical precipitation pathway of a polymer in NIPS membrane fabrication process. Binodal and spinodal curves are represented in ternary-phase diagrams. As a result of the addition of photocatalytic nanoparticles, such as TiO₂, the diagram of the binodal curve of the ternary system is displaced as depicted in the red line plotted in Figure 3 [102,103]. This will ultimately affect the porous morphology of the polymer membrane and it should be analyzed in detail for each quaternary system.

![Figure 3. Ternary phase diagram with a precipitation pathway. Adapted from [102].](image-url)

Additionally, the kinetics of the exchange between the solvent and non-solvent would affect the membrane morphology. When the exchange is fast, the pathway followed by the change in the mixing composition would cross the binodal line earlier; thus, big finger-like pores and asymmetric membrane morphology are expected. Otherwise, when the exchange kinetics is slow, small sponge-like pores are usually obtained.

PVDF and sulphur-based polymers i.e., PSF and PES are the polymers mostly used to develop photocatalytic membranes by NIPS. TiO₂-based photocatalyst has been introduced into these membranes to remove some model compounds such as dyes (MB, MO), BSA, or humic acid and organic pollutants as bisphenol A or tartrazine.

Regarding the solvent/non-solvent pairs, as can be seen in Table 1, water is the non-solvent most commonly used. Although a large number of solvent and non-solvent combinations are possible, DMAc/water [15–18,20–23,31,38,39,42,52–57,59] and NMP/water [26,28,32–34,37,40,41,51,58] are the most popular pairs in NIPS (Table 1). DMF/water [35,36,43] and TEP/water [25] are also used but to a lower extent. Membranes from ternary systems polymer/solvent/ non-solvent, where the polymer is PVDF, PSF, or PES, the solvent is DMAc, NMP, or DMF and the non-solvent is water, present the typical porous asymmetric structure formed by a thin dense top layer, supported by a finger-like structure [15,17,18,20–23,28–30,33,34,36–39,43,51], which is characteristic of instantaneous demixing.
when water is used as coagulant. Moreover, due to the binodal curves of the systems PVDF/DMAc/water, PVDF/NMP/water, and PVDF/DMF/water being very close, similar membrane structures with the three solvents were obtained [104]. The use of a water: isopropanol (70:30) mix as coagulation bath for TiO2/PES membranes produce a sponge-like structure [19].

Regarding quaternary systems, the most popular photocatalysts dispersed on the polymer solution are TiO2 and ZnO as shown in Table 1. The photocatalyst concentration affects importantly the rheology of the polymeric solution [15,19,29]. Figure 4 gathers the effects of the photocatalyst concentration on the polymeric solution rheology and the consequent membrane performance. On the one hand, at low concentrations of the photocatalyst, the viscosity of the polymer solution slightly and progressively increases. The hydrophilic groups of the catalyst attract the water molecules that diffuse faster towards the polymer phase, so as previously mentioned, the binodal curve shifts to the left of the ternary phase diagram (see Figure 3). This will ultimately accelerate the demixing rate, producing bigger and distorted finger-like pores and an increase of the membrane porosity. On the other hand, at high concentration of photocatalyst, due to the strong interaction between the photocatalyst nanoparticles and the polymer molecules, the viscosity increases abruptly, the polymer solution changes its nature from Newtonian to non-Newtonian fluid and the membrane formation is governed by the kinetics, that is radically slowed down reducing the pore radius and membrane porosity. This phenomenon is directly related to the membrane performance. At the photocatalyst concentration of the rheological change (Figure 4), the flux, the porosity, the pore size, the hydrophilicity, and the mechanical properties, such as breaking strength, reach a maximum (‘optimum’) value [15,19,23,26,28–30,36,38,39,43,53,59,61,63] From this point on, the membrane properties, i.e., flux, pore size, porosity, etc., decrease and additionally the photocatalyst nanoparticles tend to aggregate. The nanoparticles aggregation could be avoided if a strong chemical bonding is achieved between the nanoparticles and the polymer chains, for instance, using silane (γ-aminopropyltriethoxysilane) as a coupling agent to form a covalent link between the photocatalyst and the polymer [43].

**Figure 4.** Rheological change of the polymeric solution when the photocatalyst concentration reaches certain value.

Additionally to the photocatalyst, some works incorporate other additives or fillers to the polymer solution [15,17,18,20,21,23,24,30,33,34,38,39,43,51,57–59,62]. Some of these additives are i.e., leachable agents such as polyvinyl pyrrolidone (PVP) [17,18,20,21,33,34,38,39,43,51,58,59] and polyethylene glycol (PEG) [15,23,30,57] and inorganic salt as lithium chloride (LiCl) [24], which increase the membrane pore size and the number of pores. Surface modifier macromolecules (SMM) are tailormade polymers with tuneable hydrophobicity. As hydrophilic SMMs have lower surface
energy than polymers, they migrate to the air/membrane interface to minimize the total free energy of the system. Its addition to the polymer solution forms a denser skin layer and forces the photocatalyst migration to the active layer of the membrane [40,41]. Another interesting strategy to arrange the position of the photocatalyst is the use of magnetic nanoparticles. Sun et al. [42] applied a magnetic field over a casted solution of PES and magnetic Ni-TiO$_2$ catalyst before the immersion to the bath coagulation. The particles moved to the surface of the membrane instead of being homogeneously dispersed in the membrane matrix, confirmed by Ni and Ti mapping test of SEM images.

Finger-like microstructure usually produces membranes with low mechanical stability. On the other hand, the sponge-like structure would allow the membranes to avoid membrane compaction during filtration. This structure can be obtained with slow polymer precipitation techniques. For instance, in EIPS [101] the polymer precipitation is induced by solvent evaporation, which retards the polymer solidification. A slow precipitation mechanism usually produces membranes with a dense homogeneous structure. However, it has been reported the EIPS synthesis, using DMF as a solvent, of homogeneous sponge-like photocatalytic membranes with controlled porosity and pore size of P(VDF-TrFE) [46–49] and PVDF-HFP [50] copolymers when adding TiO$_2$ [46–49], Ag-TiO$_2$ [50], and/or ZnO [48] as photocatalysts.

The sponge-like structure can be also obtained by controlling the solvent evaporation time before the introduction of the polymer solution into the coagulation bath in a NIPS process [25,40,41]. When the solvent (NMP) evaporation time was below 5 min, g-C$_3$N$_4$ doped polymer membranes with an asymmetric structure formed by a dense top-layer and finger-like structure sub-layer were obtained. Above this evaporation time, the structure changed from finger-like to sponge-like [40,41]. In other works, using TEP as the solvent with an evaporation time of 2.5 min [25] or using isopropanol: water as the coagulation bath [19], TiO$_2$ containing polymer membranes presented sponge-like structure.

The variables employed during the phase inversion, such as i.e., components, composition, and solvent evaporation time, importantly affect the ultimate membrane morphology in terms of pore size and pore structure. Figure 5 depicts schematically the different types of variables that mainly influence the membrane structure when phase inversion techniques are used.

![Figure 5](image-url)

**Figure 5.** Schematic overview of the influence of phase inversion variables on membrane structure.

### 3.1.2. Electrospinning

Electrospinning is a simple, versatile, and low-cost way to synthesize polymeric nanofibers [66,80]. A high potential gradient is applied between the grounded collector and the polymer solution droplet. When the electrostatic potential overrides the polymer solution droplet surface tension,
charged threads of the polymer solution are formed with fiber diameters in the order of some hundreds of nanometers [105]. During this processing technique, as it has been revised in Tables 2 and 3, micro/nano-fibers have been usually collected in flat mats. In most cases, photocatalysts such as bare TiO₂ [64,65,68], and ZnO [63,67] or composites as GO-TiO₂ [61], and TiO₂-ZnO [66] were incorporated to the fiber (Table 2). In addition, electrospun mats could be used as supports for TFCM as indicated in Table 3 [75,81,87,88].

The morphological structure of the fibers can be tailored by changing different processing variables, such as the polymer molecular weight, polymer concentration, and flow rate of the polymer solution, tip to collector distance, and applied electric voltage [61,66,67,79,88]. In general, solid (non-porous) polymer fibers are produced. The bulk porosity of the mat is formed by the interstitial separation among the deposited micro/nano-fibers. Electrospinning allows high aspect ratio (length/diameter) of the fibers and uniform diameter, which means large specific surface area [61,80].

As can be seen in Tables 2 and 3, DMF is the principal solvent used [66–68,79,80] and to a lesser extent water [63], DMAc [87] and mixtures of acetic acid: formic acid [65] have been also used.

Regarding the polymer materials usually employed to produce photocatalytic mats by electrospinning, PAN is the most widely used polymer [66–68,75,80,81,87] due to its flexible nature and because it is easy to process. PAN fibers have high mechanical strength and are chemically resistant [66,67,81,87]. Other polymers used, although to a lower extent, are PTFE [63,64], P(VDF-TrFE) [61], PA6 [65], CA [79] and PVDF [86].

The photocatalytic nanoparticles of TiO₂, ZnO, among others (see Table 2) are mostly incorporated and homogeneously dispersed in the polymer solution [61,63–68], and subsequently electrospun.

The rheological change of the polymer solution caused by the photocatalyst incorporation at high concentrations also causes the agglomeration of the nanoparticles and consequently the formation of beads which could affect the polymer structure [106]. In most cases the fiber diameter and porosity increase with the photocatalyst concentration due to the increase of the solution viscosity [63,65–67], as can be extracted from scanning electron microscopy (SEM) images. When using ZnO concentration higher than 20 wt. %, there is adhesion between the nanofibers. As a result, the mat porosity decreases blocking the light access inside the mat [63]. However, Almeida et al. [61] found that the fiber diameter and porosity decreased with the increase of the TiO₂/GO concentration due to mechanical stretching during the material processing caused by an increase in the solution electrical conductivity when doping GO to form composite TiO₂/GO photocatalyst nanoparticles.

The selection of the polymer material limits the dosage of nano-photocatalysts in the polymer solution. When fibers are synthesized with fluorine polymers it is possible to work with photocatalyst concentration up to 20 wt. % of TiO₂/GO and ZnO [61,63]. Membranes that use PAN usually work with photocatalyst concentration below 2 wt. % including TiO₂-ZnO [66], and ZnO [67].

3.2. Thin Film Composite Synthesis Methods

Table 3 shows different coating techniques to prepare thin-film photocatalytic membranes, including vacuum deposition also called vacuum filtration [69–75], filtration [76,77], immersion [78–85], electrospaying [86], co-electrospinning [87], hot-press deposition [88], and sputtering followed by anodization [89]. Being immersion and vacuum deposition, the techniques mostly used. In this case the main photocatalysts incorporated to the membrane were TiO₂ [83,84,86,88,89] and composites of TiO₂ with carbon based compounds [69,70,72–74,76–80,82], among others (see Table 3).

Commercial microfiltration membranes with a pore size between 0.2–0.45 μm are frequently used as supports to be coated with the photocatalyst [70,71,73,74,76,77,84,85,89]. The use of ultrafiltration membranes is less common [69,78,82]. Commercial supports can be made of different materials as can be seen in Table 3, but the most used polymer is CA [70,71,76,77]. However, tailor-made supports have also been produced by phase inversion [83,88] or electrospinning [68,75,79–81,86,87].

The photocatalyst loading capacity depends on the synthesis technique. For instance, Table 3 shows that it is possible to successfully coat membranes with a photocatalyst (nAg-GO-TiO₂) [69], Ag-
membranes. PVDF grafting trigger Ti
For [87]. amino concentration the coating.

The work is popular approach to improve the long-term stability of the nanoparticles in the membranes. Related to this, only one work analyzed in this review applied in situ deposition, one of them using polydopamine (PDA) as an additive that facilitates the in situ immobilization of Ag-TiO2 nanoparticles [81]. High temperatures are required for the in situ growth of the photocatalyst in the membrane, so this could be the reason why only one work uses this technique in polymeric membranes.

The surface modification of the polymeric substrate is employed to create binding sites for the photocatalyst or facilitate its adhesion. For instance, plasma-induced graft polymerization is a technique that eliminates the need for a chemical initiator and is followed by immersion. Commercial PVDF membranes were modified by plasma-grafting which generates activated species that can trigger polymerization reactions [84]. The liquid grafting was made by the introduction of the plasma pre-treated membrane in a polycrylic acid (PAA) polymer solution. Binding sites were created in the membrane surface that facilitated the self-assembly of TiO2 nanoparticles as a thin and uniform coating. The number of binding sites was controlled and maximized by the modification of plasma-grafting conditions (plasma treatment: 100 W for 120 s; liquid graft: 70% acrylic acid solution at 60°C for 2 h). Similarly, the grafting can be initiated by UV irradiation [85], in this case the Ag-ZnO-Fe3O4-MWCNT/PAA was grafted to a PA membrane support.

More elaborated coating techniques also ensure the good adhesion of the photocatalyst to the support as electro-deposition techniques that includes electrospaying [86] and co-electrospinning [87]. Hot-press is a process in which pressure and temperature are applied to adhere the photocatalyst fibers, by a partial fusion, to a polymeric membrane surface; the field emission scanning electron microscopy (FESEM) cross-section images indicate a great adhesion between TiO2 as photocatalyst and PVDF as the membrane matrix [88]. Sputtering is a physic process where there is vaporization of Ti atoms from a solid material “blank” by bombarding it with energetic ions then, by anodization, Ti atoms crystalize forming TiO2 nanotubes which are firmly bonded to the PES commercial support [89].

However, in the cited literature, most of the works [71–77,82,83] do not use any additive or additional technique to ensure the photocatalyst attachment. To further analyze the photocatalyst stability, chemical evidences of the particle leaching from the membrane should be followed during the experimental evaluation of the membrane performance. Works complying this study are analyzed in the sections below.
4. Membrane Functionality

The incorporation of photocatalytic nanomaterials can lead to the following functional improvements in polymeric membranes: (1) membranes with antifouling properties and the consequently improved filtration capacity and reusability, (2) membranes with photocatalytic activity, or 3) a combination of the above two functional features. Below, a detailed analysis of the filtration and photocatalytic properties of the reported membranes will be addressed in terms of the influence of the membrane synthesis method. Additionally, general guidelines about the processing variables advisable to achieve improved membrane performance and membrane long-term stability will be emphasized.

4.1. Filtration Performance

Membranes intended for filtration applications were made by NIPS and EIPS in the case of MMM, while both commercial polymeric supports and NIPS-homemade supports were employed for TFCM. The functionality of MMM and TFCM membranes was analyzed in terms of hydraulic permeability and antifouling properties and compared to neat polymeric membranes. The hydrophilic enhancement provided by TiO$_2$ automatically benefited the permeability and antifouling properties of TiO$_2$-functionalized membranes [21,22,38], without the necessity of applying further UV irradiation (and therefore additional photocatalytic action). In other cases, the organic deposits on the functional membranes were cleaned by photocatalytic degradation after membrane filtration [17,23,43] using cycles of membrane filtration-UV irradiation cleaning. Below a detailed analysis of the membrane features and their influence on the hydraulic permeability and antifouling effect of the modified membranes will be addressed.

4.1.1. Hydraulic Permeability

The effect of adding TiO$_2$, AC, GO, and TiO$_2$ carbon-based nanomaterials on the structure and filtration properties of PVDF and PSF membranes, prepared by NIPS are shown in Table 4.
Table 4. Permeability, pore size, and porosity comparison between neat and composite membrane.

| Polymer (%) | Permeability (L/hm²bar) | Mean Pore Size (nm) | Porosity (%) | Nanoparticles Added (%) | Permeability (L/hm²bar) | Mean Pore Size (nm) | Porosity (%) | Literature |
|-------------|--------------------------|---------------------|--------------|-------------------------|--------------------------|---------------------|--------------|------------|
| PVDF (15)   | 150                      | 48.1                | 69.6         | TiO₂ (1)                | 300                      | 52.6                | 75.1         | [20]       |
|             |                          |                     |              | GO (1)                  | 400                      | 55.7                | 78.3         |            |
|             |                          |                     |              | GO-TiO₂ (1)             | 490                      | 65.2                | 83.1         |            |
| PVDF (14)   | 90                       | 18.6                | 47.2         | AC (0.5)                | 170                      | 18                  | 56           | [22]       |
|             |                          |                     |              | TiO₂ (0.1)              | 280                      | 28.2                | 54.3         |            |
|             |                          |                     |              | AC-TiO₂ (0.5:0.1)       | 255                      | 30.6                | 55.4         |            |
| PSF (18)    | 115                      | 56.2                | 62.5         | GO (0.5)                | 150                      | 61.4                | 69.4         | [33]       |
|             |                          |                     |              | TiO₂ (0.5)              | 155                      | 62.8                | 71.6         |            |
|             |                          |                     |              | rGO-TiO₂ (0.5)          | 180                      | 67.9                | 77.2         |            |
|             |                          |                     |              | N-rGO-TiO₂ (0.5)        | 230                      | 70.5                | 81.8         |            |
In Table 4, the values of permeability are directly related with the pore size of the membrane, which is defined by the initial system polymer/solvent/additive. The higher values of pore size reported in Table 4 [20,33] are due to the use of pore formers during membrane synthesis. It can be seen that, as previously indicated, the incorporation of TiO₂ increased the membrane permeability. In the MMMs the increase of membrane permeability could be mainly attributed to the increase of the pore size and the porosity produced by the inclusion of the nanomaterials on the ternary system. The use of TiO₂ modified with carbon-based nanomaterials even enlarged membrane pore size and porosity and therefore higher permeability was attained. As it was previously explained in Section 3.1.1., for MMM synthesized by NIPS, the optimal selection of photocatalyst concentration and its effect on the rheological properties of the polymer solution resulted in obtaining membranes with adequate porosity and morphology. Above the nanoparticles optimal concentration, they tend to aggregate, and it has detrimental effects in the membrane permeability and hydrophilicity. For example, the permeability for a bare PES membrane was found to be 1.1 L/m²hbar [38]. The addition of nanoparticles increased the pore size, the porosity, and the hydrophilicity (measured by contact angle), thus the permeability improved to 2.25 L/m²hbar, corresponding to a 1 wt. % of nanoparticles. Above this nanoparticle concentration, the aggregates clogged the pores of the membrane, which increased the water transport resistance of the membrane surface [59] hence, the permeability decreased.

The works that analyze the optimum concentration of TiO₂ [15,19,26,28,29,59] and Fe-TiO₂ [30] conclude that the optimum concentration of TiO₂ in the polymeric solution will be between 2–5 wt. % according to the concentration before the rheological change (see Figure 4 and Section 3.1.1), which gives the highest permeability value. This range usually corresponded to a 0.2–0.5 wt. %/cm². However, the optimum value should be experimentally found for each system as it will depend on the particular rheological properties of the polymer solution that incorporate the photocatalyst.

Additionally, Tran et al. [16] showed an efficient way to maximize pure water flux, in absence of pollutants, with fewer energy consumption using irradiation cycles of non-UV and UV periods for TiO₂/PVDF flat membranes.

In TFCM, the incorporation of an additional thin layer with low-size pore to a microfiltration or ultrafiltration support typically increases the resistance to the mass transport and even reduces further the support membrane pore size, therefore the flux is reduced [69–74,76,77,82,89]. This effect has been frequently observed when vacuum [69–74], filtration [76,77], immersion through layer by layer [82], and sputtering [89] deposition were used, using mostly GO-TiO₂ as the catalyst, but also rGO-g-C₃N₄ and TiO₂, see Table 3. On the contrary, three works reported an increase in the membrane flux after the photocatalyst deposition. Nor et al. [88] used the hot-press technique, under controlled temperature conditions (160 °C), to partially melt the TiO₂ photocatalyst to the PVDF membrane. In that case, the porosity of the support remained unaltered and the use of hydrophilic TiO₂ as photocatalyst increased the flux in contrast to the plain support. On the other hand, the flux enhancement is also achieved modifying the membrane support hydrophilicity through two ways, using the hydrophilic PVA as a crosslinker between the AC-N-rGO-TiO₂ coating and the PSF membrane support prepared by immersion [78] or using plasma grafting, which modifies the hydrophilicity of the PVDF support combined with the immersion in a TiO₂ solution [84].

4.1.2. Antifouling Properties

The antifouling behavior has been studied for MMM systems synthesized by phase inversion [15,20–23,26,33,36,38,43,53,57] and for TFCM systems synthesized by vacuum deposition [70,71] and immersion [83–85]. Model organic foulant solutions, i.e., bovine serum albumin (BSA) protein [15,17,20–22,26,84], humic acid (HA) [23,36], and polyacrylamide (PAM) [43] are usually employed for studying antifouling phenomena.

The benefits of immobilizing photocatalysts for the membrane antifouling property can be associated with: (i) an increase of the hydrophilicity and (ii) a decrease of the membrane roughness. As it has been previously explained, the use of bare TiO₂ produced an enhancement of the hydrophilicity [15,23,24,36,43,53,57,83,84], which can be enhanced even more when the material is
irradiated with UV light. In addition, this property might be intensified when carbon based composite photocatalysts are used [20–22, 33, 38, 70, 71] considering the synergy between carbon compounds and TiO₂ [20]. This synergy is produced because the presence of oxygen-containing functional groups (hydroxyl, epoxy, carboxyl, and carbonyl) in the surface of the photocatalyst facilitates the interaction of carbonaceous materials with a wide variety of organic and inorganic materials that helps avoiding the aggregation of the nanoparticles [72, 75]. Furthermore, in MMM the use of additives, i.e., PEG [57], also improved the anti-fouling property because this additive generated smoother membrane surface.

As it was previously mentioned for MMM, at the photocatalyst concentration of rheological change, the permeability, the porosity, the pore size, and the hydrophilicity of the membrane reach a maximum [15, 26, 36]. In most of the cases, this concentration matches with the one that gives the best antifouling performance [15, 29]. The best antifouling performance occurs at the concentration where the flux recovery ratio is the highest and the irreversible fouling is the lowest.

4.2. Photocatalytic Activity

4.2.1. Comparison between Suspended and Membrane Immobilized Systems

In general, it has been reported that the SPMRs permit to achieve higher degradation yields of POPs when compared to IPMRs, due to the larger active surface area, which guarantees a good contact between the photocatalysts and the pollutants. As a consequence, there are more studies of SPMRs. However, fouling, which is caused by deposition of the photocatalyst nanoparticles on the membrane surface with a consequent flux decline, and light scattering, still limit the performance of this type of PMRs configuration [4, 108, 109]. Meanwhile, in the IPMRs, the photocatalyst/pollutants contact is hindered by the mass transfer limitation over the immobilized photocatalyst. However, in this configuration, catalyst recovery and reuse can be more easily achieved than in SPMR and it is potentially less hazardous.

In general, few works have been found in the literature comparing the photocatalytic activity of (i) systems with immobilized photocatalyst, and (ii) the corresponding amount of suspended photocatalyst. In terms of membrane functionality, this comparison is a key point. It will help to determine the synthesis method that preserves the photocatalyst activity once it is immobilized or improve the dispersibility of the immobilized photocatalyst; the latter will favor enhanced performance when compared to suspended systems. When MMMs are used, a slight reduction of the photocatalytic performance compared with suspended systems has been so far observed. This reduction, in most of these works, can be considered inside the experimental error (<5%–10%). For instance, flat membranes made by EIPS showed a negligible reduction in the photocatalytic degradation yield of MB of 3% using TiO₂-NaY/P(VDF-TrFE) [49]; in the case of NIPS membranes, the photocatalyst performance decayed down to 6% (Co-TiO₂/PES) [37] and 15% (O-g-C₃N₄/PES) [41] for the degradation of 2,4 dichlorophenol and phenol, respectively. For electrospun nanofibers, using TiO₂-MWCNTs/PVDF nanofibers to degrade MB, the performance loss was 12% compared to the suspended photocatalyst [62]. Finally, degrading MO with ZnO/PAN nanofiber mat, the efficiency loss was about 4%, which means that the nanofiber photo-activity is 1.6 times less than the photocatalyst (ZnO) powder [67]. As it can be seen, in general low performance reductions on immobilized photocatalysts on MMMs were always reported (<15%) compared to suspended systems.

However, Ramasundaram et al. [86] showed an interesting comparison of the photocatalytic activity between TiO₂ (i) in suspension, (ii) deposited by electro-spraying on PVDF nanofibers mat (TFCM), and (iii) immobilized in PVDF MMM by NIPS; these authors analyzed different synthesis methods. Total degradation of MB and IC was achieved with TiO₂: suspension in 40 min, while for the electrospayed nanofibers mat, the time to achieve the degradation increased up to 60 and 100 min for each pollutant respectively. Furthermore, after 140 min of irradiation, the pollutants studied were not completely degraded with the MMMs. From these results, a lower photoactivity of MMM
is observed, attributed to a lower photocatalyst active surface area in contact with the pollutant, in contrast to the suspended system.

Overall, attending to the aforementioned results, it could be that the dispersibility on the polymer matrix of MMMs of bare TiO₂ might be more difficult than for other photocatalysts, such as composite TiO₂ photocatalyst or other type of semiconductors. However, more detailed studies should be done to confirm this preliminary observation.

4.2.2. Influence of the Synthesis Method on the Membrane Photocatalytic Performance

As previously remarked, it has been observed in the literature that the different photocatalyst immobilization methods resulted in different photocatalytic performances. This effectiveness might be attributed to different photocatalyst accessibility to light sources. Therefore, the in-depth analysis of the effect of the method of membrane synthesis on the allocation of the photocatalyst in the membrane matrix is; this analysis is focused to evaluate the potential relationship between the photocatalyst entrapment or shielding effect within the membrane and its photocatalytic activity, as well as its long-term stability.

In the case of phase inversion MMMs, the nanoparticles could be homogeneously dispersed in the whole membrane matrix, so they could be embedded in the membrane and therefore the number of nanoparticles accessible to light could be importantly reduced. To overcome this problem, dual-layer hollow fibers provide an external photocatalytic layer, while the inner layer is not photocatalytic. Within this strategy, similarly than in TFCM strategies, the filtrating performance of the membrane might be hampered with the additional photocatalytic layer. Therefore, it is often necessary to reach a compromise on the photocatalyst concentration and be careful with the pore structure of the coating layer. Different works have observed that the best photocatalytic performance of double layered TiO₂/PVDF hollow fibers [56] and flat membranes of TiO₂/P(VDF-TrFE) prepared by EIPS by Teixeira et al. [48] was achieved with a 15 wt. % of photocatalyst concentration. Further concentration increase would limit the photocatalytic activity due to the nanoparticles agglomeration with the consequent decrease the surface area. However, regarding the filtration performance of the double layered TiO₂/PVDF hollow fibers, Dzinun et al. [53] observed that at a concentration of 3 wt. % of TiO₂: the highest membrane flux was achieved as further concentration of photocatalyst blocked the membrane pores. Overall, Dzinun et al. [49] observed that a concentration of 3 wt. % of TiO₂ could be considered adequate to perform simultaneous filtration and photocatalytic degradation of NP.

Usually, the electrospun MMMs have been only used to immobilize the photocatalyst and never with filtration purposes as they usually have large macro-pores and therefore, low pollutant rejection. The mat is submerged in the polluted solution and, after the degradation process, it is easily recovered [61,63–68,87]. Similarly as in blended MMMs by NIPS, in electrospun membranes, the photocatalytic nanoparticles, such as ZnO and TiO₂, were homogeneously located in the membrane matrix [63,65]. On the other hand, the nanoparticles of the CQDs-Bi₂O₃-TiO₂ photocatalyst could be also anchored on the surface of the electrospun nanofiber using co-electrospinning [87]. Salazar et al. [50] compared the photocatalytic performance of Ag-TiO₂/PVDF-HFP membranes prepared by solvent casting and electrospinning. Solvent casting membranes present higher degradation rates of norfloxacin (64%) than the nanofibers (51%).

Only 2 out of the 21 works dealing with photocatalytic TFCMs study the influence of the photocatalyst concentration in the degradation of some dyes. On the other hand, most of the works of TFCMs select the optimum photocatalyst concentration based on the point of highest flux, but not based on results of the membrane photocatalytic activity. For instance, for GO-TiO₂/CA membranes prepared by filtration the catalyst concentration that provided the highest permeation flux was 6.85 mg/cm² but the best photocatalytic performance to degrade CR was achieved with 27.4 mg/cm² [76]. Similarly, for AC-N-rGO-TiO₂/PSF membranes prepared by immersion the optimum photocatalyst concentration was fixed in 6.92 mg/cm², although the higher degradation of MO was achieved with 9.22 mg/cm² [78]. Most of works in the literature of TFCM synthesis applied the criteria of the highest filtration performance to set the optimum photocatalyst concentration highlighting the importance of this property on the overall photocatalytic system.
On the other hand, the method of deposition of the thin photocatalytic film on the membranes is of utmost importance on the efficient anchorage of the photocatalyst on the membrane surface. While using vacuum deposition on CA membranes, low photocatalyst concentration could be deposited, i.e., 0.796 mg/cm² of Ag-rGO-TiO₂ [70] and 1.99 mg/cm² of rGO-g-C₃N₄ [71], with filtration and immersion as synthesis techniques, higher concentration of photocatalyst was attained, 16.75 mg/cm² (GO-TiO₂/CA) and 13.95 mg/cm² (Ag-PtO/PAN). Consequently, lower photocatalytic degradations were achieved with the vacuum deposited TFCMs, however, attention should be paid to the different type of photocatalysts that were used in the works reported above.

The comparison between the photocatalytic performance of TFCM and MMM membranes has been scarcely reported. Wu et al. [78] prepared TFCM with a PSF commercial support coated with a PVA solution to immobilize the nanoparticles (AC-N-rGO-TiO₂) by a surface deposition method. The MMM was synthesized by NIPS, a solution of PSF, PVP, and AC-N-rGO-TiO₂ was prepared in NMP using water as coagulation bath. TFCM presented higher photocatalytic performance (95.2%) than MMM (31.1%). In addition, Ramasundaram et al. [86] compared TFCM with MMM. They deposited TiO₂ by electro-spraying on PVDF nanofibers mat and immobilized TiO₂ in PVDF MMM by NIPS. Surface deposition by electro-spraying was more favorable than NIPS to maintain the photocatalyst accessibility to light. In the latter works, the lower photocatalytic performance of the MMM was explained because the blending method encapsulated the nanoparticles into the membrane matrix, which protects the nanoparticles but hinders light irradiation. However, as previously analyzed in Section 4.2.1, many works also reported similar photoactivity of photocatalytic MMMs than suspended systems. Therefore, more attention to the methodology of MMM membranes preparation should be paid; further works comparing the structural features and functional performance of photocatalytic TFCMs and MMMS should be done discerning the advantages and disadvantages offered by the two membrane types.

4.2.3. Membrane Aging

Polymer Stability

Certain polymers are materials susceptible to photolytic and photocatalytic degradation. As photocatalytic membranes are going to be exposed to UV irradiation, membrane aging, and long-term resistance are crucial to their application. It must be remarked that only few works have explored this long-term membrane behavior.

As in MMMS, the photocatalyst is embedded inside the matrix, and there is no additional protective layer over the membrane, the polymer is highly exposed to light effects. In terms of membrane aging, PVDF membranes have demonstrated stability against photocatalytic reaction and UV irradiation in long-term operation [53–55]. After 30 days of UV irradiation, the membranes upheld their integrity despite many cracks were formed on the surface, which increased the membrane roughness, analyzed with atomic force microscopy (AFM). FTIR tests showed a change of the PVDF crystalline phase, besides the formation of –CF=CH- double bond by dehydrofluorination. The dehydrofluorination reaction of PVDF is presented in Figure 7, following the pathway of carbocation reaction using H₂O as a polar solvent and UV as the heat source, described by Dzinun et al. [54]. There was a slight decrease in tensile strength that increased with the increase of the UV exposure time, causing negative impact on the overall stability. SEM images of an mp-g-C₃N₄-TiO₂/PSF membrane [34] showed that the membrane structure was maintained without changes after its exposure to simultaneous photocatalytic degradation and filtration of the antibiotic sulfamethoxazole (SMX) during 30 h. Some loss in tensile strength was observed, although the membrane integrity and flexibility were maintained. In the case of MMM synthesized by electrospinning, ZnO/PTFE nanofibers, they only lost 3.8% of mechanical strength after 5 cycles of operation (25 h) [63].
Figure 7. Dehydrofluorination reaction of PVDF through a carbocation reaction where H₂O acts as a polar solvent and UV light as the heat. Adapted from [54].

On the other hand, the TFCM configuration might incorporate a photolytic protective layer to low UV-resistant polymeric supports such as PP [99]. This solution was also pointed out by Tsehaye et al. [100] in their study of the PES membrane stability under oxidative conditions.

Photocatalyst Detachment or Leaching

During operation, photocatalytic membranes might suffer the loss or leaching of photocatalytic nanomaterials. The methodology employed to synthesize the membranes might induce different nanomaterials anchorage on the polymer matrix. However, only two authors evaluate the amount of photocatalyst that leached out from the membrane after each cycle using inductively coupled plasma (ICP). Ramasundaram [86] did not detect any titanium in solution after 10 cycles (17.7 h) of use of a TFCM of electrospayed TiO₂ on a PVDF nanofibers mat and they observed that the photo-activity was preserved during all cycles. Tissera et al. [67] analyzed the Zn leached from a MMM made of electrospun ZnO/PAN nanofiber after 3 cycles (30 h). Concentrations of Zn of 40, 20, and 17 ppb (which accounted respectively by 0.08, 0.04 and 0.034 μg of ZnO leached per mg of membrane) were measured in the feed tank solution during the first, second and third degradation cycles, respectively. This meant than less than 0.0052% of the photocatalyst embedded in the membrane matrix was leached out of the membrane after 30 h of use.4.2.3.3. Long term membrane performance

Table 5 compiles the works that reused the membrane in several cycles and the change in the degradation rate of organic pollutants. Only 2 out of 5 works of TFCM found a loss of 10% in photocatalytic activity during the cycles of photocatalyst reuse while the others reported stable membrane photocatalytic activity. Meanwhile 6 out of 12 works of MMMs presented a decline in the photocatalytic activity with membrane reuse between 8 and 45% independently of the synthesis method. This means that, similarly, approximately 40% of both TFCM and MMM works report a decline in photocatalytic activity during the reuse of the photocatalytic membranes, although the loss of activity could be more pronounced in the case of MMMs due to the protective effect that the photocatalytic layer could exert on the polymer substrate. Bare TiO₂ or ZnO were used in most membranes collected in Table 5.

For instance, in TiO₂/P(VDF-TrFE) MMMs synthesized by NIPS, the degradation rate of MB and the kinetic constants of three consecutive cycles of 5 h each were reported [48]. Particularly, when the membrane incorporated a TiO₂ concentration in the polymer solution of 5, 10, and 15 wt. %, the reduction in the degradation rate between cycle 1 and cycle 3 was 6%, 16%, and 13%, respectively. In this work, nanoparticles were located mainly on the membrane surface and weakly bonded to the polymer, so the authors attributed this effect to the nanoparticles leaching out of the membrane.

It is worth remarking that the important 45% decay in photocatalytic activity reported in Table 5 for some MMMs could be attributed to an important membrane degradation during operation caused by the low chemical and photochemical stability of CA polymer [41] and the high power of the light source employed [60].
Table 5. Influence of the synthesis method on the nanoparticles stability based on the degradation change of the membrane after its reuse.

| Photocatalyst Polymer | Synthesis Method | Targeted Pollutant | Number of Cycles | Total Irradiation Time (h) | Loss in Degradation Rate (Cycle 1 – Last Cycle) | Power of the Lamp | Ref. |
|-----------------------|------------------|--------------------|------------------|---------------------------|-----------------------------------------------|------------------|------|
| MMM                   |                  |                    |                  |                           |                                               |                  |      |
| TiO₂/ PSF             | NIPS             | Cr (VI)            | 4                | -                         | 0%                                            | Sunlight         | [28] |
| N-TiO₂/ PMAA-g-PVDF/PAN | NIPS             | Bentazon           | 3                | 10                        | 0%                                            | UV (5063 lux)    | [27] |
| ZnO/ CA-PS            | NIPS             | CR, RY             | 5                | 5                         | 45%                                           | Sunlight         | [44] |
| TiO₂/ PSF             | NIPS             | MB, MO             | 5                | 7.5                       | 0%                                            | UV-C 10W         | [51] |
| TiO₂/ P(VDF-TrFE)     | EIPS             | MB, CIP, IBP       | 4                | 20                        | 0%                                            | UV-A 48W         | [46] |
| TiO₂ or ZnO/ P(VDF-TrFE) | EIPS             | MB, Model organic  | 3                | 15                        | 13%                                           | UV-A 48W         | [48] |
| Ag-TiO₂/ PVDF-HFP     | EIPS             | NOR                | 3                | 15                        | 15.6%                                         | UV-A 8W          | [50] |
| Ag-TiO₂/ PVDF-HFP     | Electrospinning  | NOR                | 3                | 15                        | 8.8%                                          | UV-A 8W          | [50] |
| ZnO/ PAN              | Electrospinning  | MO                 | 3                | 30                        | 0%                                            | UV-A 40 W        | [67] |
| ZnO/ PTFE:PVA         | Electrospinning  | RhB                | 5                | 25                        | 20%                                           | UV 500 W         | [63] |
| TiO₂/ PTFE            | Electrospinning  | MB                 | 5                | 7.5                       | 45%                                           | UV 300 W         | [64] |
| TiO₂/ PA6             | Electrospinning  | RBB                | 3                | 12                        | 0%                                            | UV 6 W           | [65] |
| TFCM                  |                  |                    |                  |                           |                                               |                  |      |
| TiO₂/ PVDF            | Electrospinning  | BPA, 4-CP, CMT     | 10               | 16.7                      | 0%                                            | 4 W              | [86] |
| GO-TiO₂/ CA           | Filtration       | CR                 | 3                | -                         | 10%                                           | No data          | [76] |
| TiO₂/ PEI-P25         | Immersion        | RhB                | 6                | 12                        | 10%                                           | UV 18W           | [83] |
| NH₂-TiO₂/ PAN-CNT     | Immersion        | Cr (VI)            | 5                | 5                         | 0%                                            | 125 W (420 nm)   | [80] |
| rGO-α-Fe₂O₃/ PAN      | Vacuum deposition | MB, MO, RhB, R6G, MG, GV | 5              | 4.2                       | <10%                                          | UV-vis 275 W     | [75] |
Overall, the characterization of the membranes before and after a long-term exposure to UV would help to better analyze membrane aging. The morphological characterization should be done with SEM or FESEM, to obtain images of the membrane surface [55] and AFM, to analyze changes in the membrane surface roughness [54,55]. Additionally, other chemical analytical techniques can be used, X-ray diffractive dispersion (XRD) to observe changes in the crystallinity of the materials employed and energy dispersive X-ray (EDX) to observe changes in the chemical structure of the membrane. Finally, analysis of the functionality of the photocatalytic membrane with filtration tests and photocatalytic reuse tests through ICP analysis of the liquid medium would complete the photocatalytic membrane stability characterization.

5. Guidelines and Recommendations for Researchers

This review aims to evaluate the materials, synthesis methods, and their relationship with the functional performance of photocatalytic polymeric membranes. PVDF is the recommended material to fabricate flat or hollow fiber MMMs by phase inversion techniques due to its photocatalytic resistance; this is supported by the large information gathered in the reported works that use this polymer. However, the use of PES and PSF is also popular due to their easy manufacture. In the case of nanofibers made by electrospinning, PAN is the polymer mostly used because it is easy to process by this technique, albeit it is not the most stable polymer under UV-light exposure. On the other hand, in TFCM it seems that the photocatalytic coating generates a protective layer, so supporting materials with low photocatalytic resistance, such as PES or PSF, could be used.

Many works use TiO₂ as photocatalyst for photocatalytic membranes, however, it has been demonstrated that carbon based composite photocatalysts not only improve the photocatalytic performance in both MMMs and TFCMs but also contribute to enhancing between 2 and 3.26 times the filtration performance of the membrane. Moreover, from preliminary observations in this review, it could be stated that TiO₂-based composite photocatalysts could be better anchored and dispersed by linkers on TFCMs than bare TiO₂ nanoparticles.

After an exhaustive review of the synthesis methods for photocatalytic polymeric membranes, not enough evidence was found to select the best methodology and the choice should ultimately be based on the aimed application of the material. The few works that compare the photocatalytic performance of MMM and TFCM have observed that TFCM provided higher photocatalyst surface area of contact with the pollutant, so the degradation is favored with respect to MMM. Nevertheless, in MMM some researchers investigate the synthesis of dual-layer hollow fibers. In this case, the outer layer would have photocatalytic properties and the inner one would act as support to give mechanical strength to the membrane.

Photocatalyst leaching out of the membrane is an important issue. Higher stability of the photocatalyst entrapped in the MMM than in TFCM is expected. However, the use of photochemically non-stable polymers and/or very high powerful light sources could cause the polymer membrane degradation in MMMs and the consequent release of the photocatalyst. Furthermore, in TFCM the use of additives and the application of modification techniques such as surface oximation and plasma-grafting improves the photocatalyst adherence. Furthermore, no reliable evidence has been found to demonstrate any direct photocatalyst loss attributed exclusively to the membrane synthesis technique.

Finally, although ideally, a good method of photocatalyst immobilization on a membrane could reduce the aggregation of suspended systems, it has been well established in a variety of studies that the photocatalytic degradation activity is normally reduced when the photocatalyst is immobilized in a membrane due to the reduction of the available contact area. This reduction has been studied and quantified in several works giving a range of 3%–30%. Therefore, the challenge still persists on the search for a membrane synthesis method that helps reducing nanoparticles aggregation and maximize the area of the photocatalyst exposed to the light source. Herein some guidelines are proposed to help authors in decision-making on the synthesis methodology to produce a membrane with the best performance depending on the required function.
If a membrane with high filtration and antifouling performance is sought, MMM synthesized by phase inversion is recommended, and particular attention should be paid to the solution rheology to obtain maximum porosity and permeability. The photocatalyst concentration when the rheological change of the polymer solution occurs should be found in the range of 0.2–0.5 wt. % for MMM, albeit each system should be studied in detail. Furthermore, TiO₂-based composites of metal or carbon based materials are recommended.

If high pollutant rejection is desired, either MMM can be produced with techniques to slow down phase inversion or TFCM techniques, as these would reduce the mean pore size and surface porosity. Among TFCM deposition techniques, vacuum filtration is the least recommended method as it produces the lowest performance in the deposition of the photocatalyst and the least photocatalytic activities.

Based on this literature review, we consider that there are important methodological aspects that should be considered and/or studied more in depth to facilitate future comparison among photocatalytic membranes. Some recommendations can be suggested:

- The influence of the lamp irradiance is crucial on photocatalytic membrane systems, in particular on the effects on membrane aging. In the case of photocatalytic membranes, UV light sources of low power (less than 50 W) or light-emitting diodes (LEDs) are highly recommended to reduce polymer aging and thus, ensuring long-term stability of the membrane.

- Long-term stability is a key issue for process scalability. Therefore, high attention should be paid on this aspect during the membrane viability study. Performing (i) long-term experiments to verify that the membrane maintains its integrity under the experimental conditions, and (ii) reuse experiments to ensure good stability of the photocatalyst is therefore encouraged. It should be considered the effect of elevated heat, UV irradiation and moisture which can lead to polymer degradation.

- The use of analytical techniques such as EDX, XRD, AFM, and SEM or FESEM, before and after the UV exposure, are highly recommended to analyze the aging of the membrane. The quantification of the leaching of nanoparticles from the membrane to the medium should be analyzed with ICP.

- High efforts are being adopted on synthesizing novel and more active photocatalysts. Their physico-chemical characteristics can be notably different from conventional semiconductors such as commercial TiO₂. Nanoparticle dispersions in polymeric matrix present different rheology so the membrane processing can suffer significant changes. These changes could play a key role on the improvement of nanoparticles dispersibility on the polymer matrix so far encountered on MMMs.

- When the membrane has a simultaneous filtration and photocatalytic function, experimental reactors should integrate both filtration and photocatalytic degradation. Membrane functionality characterization should consider: (i) pure water flux test to obtain permeability values (L/hm².bar); (ii) photocatalytic degradation comparison with the suspended system to evaluate the change in the photocatalytic activity when the nanoparticles are immobilized, and (iii) pollutant rejection under dark and UV irradiation conditions.

- In the particular case of TFCM synthesis, the influence of the photocatalyst concentration using different coating techniques has not been sufficiently evaluated. There is expected to be an optimal concentration to maintain the balance between the membranes photocatalytic and filtration functions. While, as aforementioned, in MMMs the recommended concentration range is 0.2–0.5 wt. %, in TFMC this evaluation has not yet been assessed and should also be addressed.

- It is noteworthy that most of the works studied in this review analyze the photocatalytic activity using dyes as model organic pollutants. These types of molecules are photosensitive and can be adsorbed on the catalysts and/or on the membranes leading to results that are not representative of other pollutants. Therefore, it is advisable to select
other types of organic model compounds, such as, acetic acid, or certain non-biologically degradable organic compounds as those contained in hospital effluents, such as antibiotics, (cautiously) organohalogens, etc., to generalize the obtained conclusions. In this regard, considering the important presence of persistent compounds, antibiotics and disinfectants likely causing bacterial inhibition in the on-site hospital wastewater treatment, the use of photocatalytic membrane reactors for the on-site treatment of hospital wastewaters is envisaged as a promising alternative.

Finally, here we would like to remark that for instance regarding the membrane nature, most PMRs make use of polymeric membranes either in SPMR or IPMR configuration. Although ceramic membranes are less common, their positive characteristics such as high permeability, superior chemical, mechanical, and thermal resistance than polymeric membranes, resistance to UV irradiation, long life, and excellent antifouling properties counterbalance their higher price and there is an expanding trend of applications to wastewater remediation [110].

Overall, the development of fouling resistant and more photo-catalytically active membranes that could maximize the energy adsorbed from the visible light wavelength, together with a deeper knowledge and control of mass transfer limitations in advanced configurations of PMRs and the continuous research of motivating applications will facilitate process design and scale-up, thus, paving the way to establish PMR as one of the best available technologies (BAT) for remediation of wastewaters containing persistent organic pollutants.

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