



# Degradation of Organic Methyl Orange (MO) Dye Using a Photocatalyzed Non-Ferrous Fenton Reaction

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Abstract: Removal of recalcitrant organic pollutants by degradation or mineralization from industrial waste streams is continuously being explored to find viable options to apply on the commercial scale. Herein, we propose a titanium nanotube array (based on a non-ferrous Fenton system) for the successful degradation of a model contaminant azo dye, methyl orange, under simulated solar illumination. Titanium nanotube arrays were synthesized by anodizing a titanium film in an electrolyte medium containing water and ethylene glycol. Characterization by SEM, XRD, and profilometry confirmed uniformly distributed tubular arrays with 100 nm width and 400 nm length. The non-ferrous Fenton performance of the titanium nanotube array in a minimal concentration of H<sub>2</sub>O<sub>2</sub> showed remarkable degradation kinetics, with a 99.7% reduction in methyl orange dye concentration after a 60 min reaction time when illuminated with simulated solar light (100 mW cm<sup>-2</sup>, AM 1.5G). The pseudo-first-order rate constant was 0.407  $\mu$ mol<sup>-1</sup> min<sup>-1</sup>, adhering to the Langmuir–Hinshelwood model. Reaction product analyses by TOC and LC/MS/MS confirmed that the methyl orange was partially fragmented, while the rest was mineralized. The facile withdrawal and regeneration observed in the film-based titanium nanotube array photocatalyst highlight its potential to treat real industrial wastewater streams with a <5% performance drop over 20 reaction cycles.

**Keywords:** methyl orange; titanium nanotube; non-ferrous Fenton reaction; advanced oxidation process; photocatalysis

## 1. Introduction

Advanced oxidation processes (AOPs) are an emerging method of treating organic contaminants in wastewater, characterized by the in situ generation of highly active oxidizing free radicals [1]. Free radical formation can be induced by many media such as  $H_2O_2$  [2],  $O_3$  [3],  $Cl_2$  [4], UV–Visible radiation [5], heat, persulfates and peroxymonosulfates [6], carbonates [7], and nitrate–nitrite mixtures [8]. Photocatalysis-based AOPs are driven by Fenton or Fenton-like chemistry [9], ozonation [10], and sulfate radical-based processes [11]. Herein, Fenton processes generate in situ hydroxyl radicals at a redox potential of 2.8 V, non-selectively attacking a wide array of chemical centers, leading to fragmentation and even mineralization of the parent molecules [12]. The classical Fenton process involves Fe(II) species that react with  $H_2O_2$  to produce  $\cdot$ OH radicals in situ. However, this process has limitations, primarily due to the precipitation and sludge formation of Fe(III), with the former process reducing the available Fe(II) content in the system over time and significantly lowering the reaction efficiency. The replacement of Fe(II) with other metals, alloys, and metal oxides for organic substance decomposition through the subsequent generation of  $\cdot$ OH radicals has recently gained much attention [13,14].

Unlike polymer composites for treating aqueous waste streams [15], metals with variable oxidation states act as promising non-ferrous Fenton catalysts in hydrogen peroxide



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**Copyright:** © 2023 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (https:// creativecommons.org/licenses/by/ 4.0/). solution due to their multiple oxidation states, which can be catalytically converted from inert to active or vice versa via a simple redox cycle [16]. In this regard, zero-valent aluminum in an acidic medium (pH < 4) has demonstrated Fenton-like activity that mineralizes phenol, 4-chlorophenol, and nitrobenzene [17]. This study found that the decomposition of bisphenol-A was achieved with an efficiency of 75% within 12 h in a highly acidic medium (pH~1.5) and that the rapid formation of reactive OH radicals caused by the addition of trace amounts of Fe(II) increased the decomposition rate exponentially [18]. Another study using a  $Ce^{3+}/H_2O_2$  system was found to act as an alternative to the  $Fe^{2+}/H_2O_2$  system, but the extent of ·OH radical generation was highly dependent on the surface defects and surface oxide states of  $CeO_2$  [19,20]. The degradation kinetics of the acid orange 7 dye compound indicated that the oxidation proceeds through an intermediate hydroxyl adduct followed by adsorption of the dye to the  $CeO_2$  surface [20]. However, the acute cytotoxicity of Ce in humans does not favor the use of Ce in large-scale applications [21]. Unlike most non-ferrous metals for the Fenton process, the Cr(VI) species has been found to generate highly active Cr(V) during the in situ formation of stable OH radicals over a wide pH range [22]. The rapid dissolution of Cr(VI) and extreme carcinogenic properties of Cr(VI) in aqueous media hinder its use in water treatment, as Cr(VI) is released after treatment, which opens up another threat that requires post-treatment again [23,24]. Furthermore, Co [25,26], Mn [27], Cu [28,29], and Ru [30,31] species have been studied in non-ferrous Fenton processes in mild acid, neutral, or weakly alkaline pH ranges.

TiO<sub>2</sub> has been used in UV-activated AOP systems with ozone, where hydroxyl radicals were produced by ozone decomposition in water [32,33]. The resulting system was highly active in decomposing recalcitrant 2,4-dichlorophenoxyacetic acid, bisphenol A, butylnaphthalenesulfonate, and benzyldodecyldimethyl ammonium bromide, leading to efficient mineralization as seen in total organic carbon (TOC) measurements [32]. Herein,  $TiO_2$  is non-toxic, but using ozone makes the process less environmentally benign [34]. The rapid reactivity of ozone induces undesirable side reactions that generally lead to toxic byproducts. Hydroxide radicals generated from  $H_2O_2$  are generally short-lived and formed under normal temperature and pressure conditions [35], so there are no major limitations on reactor design and apparatus. Thus, a simple and uncomplicated reactor design is sufficient to operate as a stand-alone single system or to work with another treatment method in a hybrid process. Hence, a  $TiO_2/H_2O_2$  system replacing ozone would be very attractive if it can produce surface active radicals to drive AOPs. Recently, Choi et al. showed that inorganic As(III) was oxidized to As(V) in a  $TiO_2/H_2O_2$  system along with  $WO_3/H_2O_2$ and  $ZrO_2/H_2O_2$  [36]. The authors suggested that the reaction proceeds through surface complexation between  $TiO_2$  and  $H_2O_2$  to form hydroperoxyl radicals on the surface of  $TiO_2$  by an inner sphere electron transfer. In support of this, the authors found that the rate of As(III) to As(V) conversion was least affected by hydroxyl radical scavengers, while there was a significant decrease in the reaction rate for hydroperoxyl or superoxide radical scavengers. Recently, a N-Ti $O_2/H_2O_2/v$  is ble light system was evaluated for the degradation of diclofenac in a submerged photocatalytic membrane reactor [37]. Even when the reaction proceeded with high mineralizing efficiency, the authors found that recovering the suspended catalyst powders was costly and laborious. This inherent downfall can be easily overcome by using a film-based catalyst that can easily enable the scalability of the reaction system.

As discussed above, AOPs are indispensable for treating organic or inorganic contaminants in ferrous or non-ferrous Fenton-type processes. Regardless of the type, most work has focused on catalysts in powder form. Catalysts grown on their own in a film undoubtedly have the highest level of stability that can be easily separated from the reaction system. However, the summary provided in Table 1 clearly indicates that a suitable catalytic system for treating common industrial pollutants in a process that can be easily operated utilizing abundant sunlight is still to be improved from a pilot-scale system to an industrial-scale system. It is known that  $TiO_2$  can be synthesized in the form of nanotubular arrays on Ti film by an anodization process, rendering fewer interfacial grain boundaries [38] to improve charge separation and enhance redox activity. In this study, the non-ferrous Fenton behavior of  $TiO_2$  was explored with a vision to be tested on a pilot scale in the future, with the addition of a small amount of  $H_2O_2$  under light irradiation for the degradation of methyl orange (MO), one of the major organic contaminants. Therefore, in this study,  $TiO_2$  nanotube arrays were fabricated and systematically utilized with a non-ferrous Fenton approach with pharmaceutical grade 6%  $H_2O_2$  to degrade the model contaminant, MO, under natural sunlight irradiation. The catalysts were characterized by surface analysis techniques such as XRD, XPS, SEM, and profilometry. The MO degradation rate was studied as a function of  $H_2O_2$  concentration, the intensity of solar light, and the type and composition of  $TiO_2$ material. A plausible reaction mechanism was derived from controlled experiments using hydroxyl and hydroperoxyl radical scavengers. In addition, the catalyst was reused for 20 cycles to compare the recyclability of the  $TiO_2$  film catalyst for each MO degradation.

#### 2. Experimental Methods

## 2.1. Materials

Phosphoric acid ( $H_3PO_4$ ), acetone, ethanol, sodium fluoride (NaF), ethylene glycol (EG), methyl orange dye (MO), and  $H_2SO_4$  (98%) were purchased from Sigma Aldrich (St. Louis, MO, USA) in the highest purity grade available and used as received. Titanium sheets (99.7% purity, Yunjie Metal Co., Baoji, China), stainless steel bars (Yunjie Metal Co., China), and polishing cloth (Struers LLC, Cleveland, OH, USA) were used to fabricate titania nanotube arrays. The  $H_2O_2$  used in the experiments was pharmaceutical grade with 6% solution in water (Meliorate Health, Mumbai, India). The water used in all experiments was deionized ultrapure water (DI), attained from a Milli-Q direct water purification system (Fischer Scientific, Waltham, MA, USA) with a resistance of 18.2 M $\Omega$  at 25 °C.

## 2.2. Catalyst Film Preparation

Titania nanotube array (TNA) films were fabricated using a modified anodization technique stated elsewhere [39,40]. In short, titanium sheets were cut into small pieces of  $2 \times 4$  cm<sup>2</sup> and then polished with sandpaper (400 grit) and a wet polishing cloth with an alumina micro polish slurry to create a flat, defect-free surface and remove the oxide layer on the surface. The polished Ti film was then ultrasonically cleaned in acetone and ethanol for 10 min each, then completely dried under a N<sub>2</sub> gas stream. Next, the cleaned Ti film was electrochemically anodized in an electrolyte solution containing 4:1 (v/v) of DI: EG, 0.26 M of NaF, and 0.94 M H<sub>3</sub>PO<sub>4</sub>. In the electrochemical cell, the Ti film was used as the anode and paired with a stainless-steel sheet as the cathode, and the distance between the two electrodes was less than 3 cm. After the electrochemical cell was set up, a direct current (DC) voltage of +30 V and 0.04 A was applied to the cell for 4 h and continuous stirring of the electrolyte was ensured during the anodization period to provide proper mass transport of charged ion particles in solution. The as-anodized Ti nanotube array film (anoTNA) was washed thoroughly with deionized water and ethanol, dried in ambient air, and annealed at 400 °C for 2 h to crystallize the anatase phase, and the resulting film was labeled as annTNA.

#### 2.3. Dye Degradation Tests

A clean, dry reactor with a quartz window was used for MO dye degradation tests. In a typical reaction, the total liquid volume was 45 mL, containing 20 ppm (61  $\mu$ M) MO concentration and 1 mL of 6% H<sub>2</sub>O<sub>2</sub>, with 2 mM H<sub>2</sub>SO<sub>4</sub> acid used to maintain a medium pH of 3.0. The protonation or deprotonation of the dye is important depending on the mechanistic pathway of degradation and affect the adsorption capacity of the dye to the catalyst [41]. A 2  $\times$  1 cm<sup>2</sup> area of the annTNA catalyst film was immersed in the solution facing the quartz window to maximize light absorption. In experiments studying the H<sub>2</sub>O<sub>2</sub> effect, the amount of H<sub>2</sub>O<sub>2</sub> was varied appropriately. Radical scavengers such as isopropyl

alcohol (IPA), formic acid (FA), tertiary butyl alcohol (TBA), and ascorbic acid (AA) were used in an amount equivalent to that of the  $H_2O_2$  in the medium. In a comparative study with other types of titania, 200 mg of powdered TiO<sub>2</sub> catalyst was used, which is equivalent to the Ti content of the immersed annTNA film catalyst. The photoelectrochemical reaction was carried out for up to 60 min at a light intensity of 100 mW cm<sup>-2</sup> irradiated with a solar simulator (ABET Technologies, Milford, CT, USA) equipped with a Xe Arc lamp. Test aliquots of 1 mL were collected at regular intervals, diluted with 2 mL of DI, and tested directly using a UV–Visible spectrometer (Biochrom, Cambridge, UK).

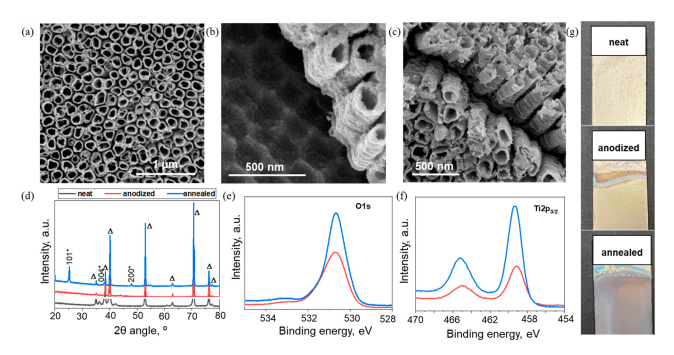
## 2.4. Characterization of Solid and Liquid Samples

The surface morphology and crystalline pattern of the synthesized TNA films were analyzed using scanning electron microscopy with energy dispersive X-ray spectroscopy (SEM/EDX, NOVANANOSEM 450, FEI company, OR, USA) and X-ray diffraction (XRD, PANalytical Empyrean, Malvern Panalytical, Malvern, UK), respectively. Additionally, the surface roughness and depth of numerous photocatalyst films were investigated using a profilometer (Leica DMC 8, Leica Microsystems, Wetzlar, Germany). X-ray photoelectron spectroscopy (XPS) (Kratos, Axis Ultra DLD, Kratos Analytical Ltd, Manchester, UK) was used to collect elemental information and oxidation states of electrode samples in Al K $\alpha$  monochromator mode. The photoexcitation behavior of annTNA was assessed by photoluminescence (PL) data recorded on an F-7000 FL (Hitachi, Tokyo, Japan) spectrometer with excitation at 375 nm, and emission was measured in the 400–800 nm range. After degradation testing, the degradation products of the MO dye were examined by LC/MS analysis using an Agilent 6460 (Agilent Scientific Instruments, Santa Clara, CA, USA) LC/MS/MS tandem mass unit with an electrospray ionization detector in negative mode. High-performance liquid chromatography (HPLC) analysis was performed on a Waters Acquity (Agilent Scientific Instruments, Santa Clara, CA, USA) instrument equipped with an RP C18 column and a photodiode array (PDA) detector for parallel recognition of MO dye degradation products.

## 3. Results and Discussion

## 3.1. Structural Characterization of the Prepared TNA Films

The annealed titania nanotube array films (annTNA) synthesized in this study showed very regular nanotube arrays with an inner tube diameter of ~100 nm, as shown in Figure 1a. To visualize the average tube length and the alignment of the tubes, the SEM sample specimen was scratched purposefully. As given in Figure 1b,c, annTNA tubes were regularly aligned with similar tube lengths of ~400 nm. The shape of the tube did not change at all from the anodizing process to the annealing step, so the tubular morphology was intact during the crystallization process. XRD analysis showed similar diffractograms for the pure Ti film and the anodized TNA film. Given that the photograph of the anodized TNA film (Figure 1g) shows full coverage of the TNA material on the Ti film, it is less likely that the diffraction pattern seen on the anodized TNA comes from the substrate Ti metal. Since the d-spacing of the  $TiO_2$  rutile phase coincides with that of the Ti metal [42], it is reasonable to assume that the TNAs formed on the anodized Ti film were a rutile phase, which was partially transformed into an anatase phase in the annealing step. This was confirmed by the diffractogram of annTNA, that exhibited new peaks at 25.30°, 38.57°, and 48.02° related to (101), (004), and (200) planes of the anatase phase, as shown in Figure 1d. XPS analysis of anoTNA film and its annealed TNA counterpart showed no differences in the binding energies of the  $Ti2p_{3/2}$  and O1s. The highest peaks of the  $Ti2p_{3/2}$  and O1s XPS spectra were centered at 459.3 eV and 530.7 eV, respectively (Figure 1e,f).

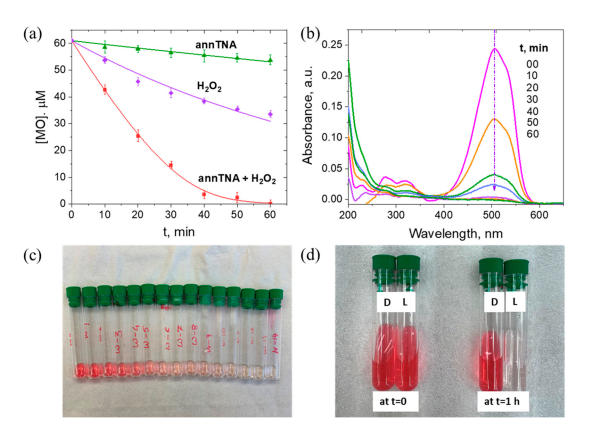


**Figure 1.** SEM images of (**a**) the top surface of annTNA (scale bar 1  $\mu$ m) and (**b**) the tube length and bottom surface after etching (scale bar 500 nm), and (**c**) the uniform height of nanotubes (scale bar 500 nm) visualized by a custom scratch on the TNA surface. (**d**) XRD analysis of anoTNA and annTNA; the indices '\*' and ' $\Delta$ ' represent the anatase phase and the rutile phase, respectively. XPS spectra of (**e**) O1s and (**f**) Ti 2p<sub>3/2</sub> of anoTNA and annTNA. (**g**) Photographs of bare Ti film (neat) and Ti film after the anodization and annealing steps.

## 3.2. Photocatalytic Degradation of MO Dye

The validity of the hypothesis that the titanium center of the annTNA arrays synergizes with  $H_2O_2$  to generate oxidative radicals capable of decomposing organic pollutants was examined by the photocatalytic efficiency of annTNA in degrading 20 ppm MO dye using a 6%  $H_2O_2$  solution under solar irradiation (AM 1.5 G, 100 mW cm<sup>-2</sup>). The degradation kinetics of the MO dye were also compared to those of the individual H<sub>2</sub>O<sub>2</sub> and annTNA counterparts over 1 h, with measurements taken at regular time intervals. Figure 2a-c shows that the annTNA– $H_2O_2$  coupled system degraded more than 99% of the dye molecules within 1 h of light irradiation, with 94% MO degradation after 40 min of reaction. The dependence of the MO dye concentration on the resulting degradation rate was thoroughly analyzed with zero-, half-, first- and second-order rate equations. Consistent with several reported works in the literature, the kinetics of MO dye degradation by the annTNA– $H_2O_2$ coupled system followed the Langmuir-Hinshelwood (L-H) reaction kinetics. Several reaction parameters govern the L–H kinetics, including the incident light intensity (I), the light-exposed surface area of the catalyst (s), the concentration of the MO dye (C), and the adsorption coefficient of the dye on the catalyst surface (K). Considering the effect of the parameters specified above, the reaction rate ( $\Gamma$ ) can be written as in Equation (1).

$$\Gamma = -\frac{\mathrm{dC}}{\mathrm{dt}} = \mathrm{f}(I, s, \mathrm{C}, K) \tag{1}$$



**Figure 2.** (a) Time-profiled MO degradation: (purple)  $H_2O_2$  only, (green) annTNA film alone, and (red) annTNA film +  $H_2O_2$ . Experimental conditions: initial [MO] = 20 ppm, [ $H_2O_2$ ] = 55 mM, and medium pH = 3.4, under solar irradiation (AM 1.5 G, 100 mW cm<sup>-2</sup>). Symbols represent experimental kinetic data and lines are nonlinear regressions with the Langmuir–Hinshelwood (L–H) model. (b) UV–Visible light absorbance measurements of the change in MO dye concentration at 5 min intervals for the annTNA +  $H_2O_2$  system. Photographs of MO dye solutions degraded by the annTNA +  $H_2O_2$  system: (c) taken at regular time intervals in light and (d) before and after 1 h in the dark (D) and light (L).

More specifically, the photocatalytic degradation of the MO dye, which proceeded through the L–H mechanism, follows a pseudo-first-order reaction rate where a single molecule of MO dye is attached to the active site of the TNA photocatalyst per occurrence. It can be expressed as Equation (2):

$$\Gamma = -\frac{\mathrm{dC}}{\mathrm{dt}} = k \frac{KC}{1+KC} \tag{2}$$

where t is the reaction time and *k* is the L–H rate constant. MO dye concentrations over time were fitted using the nonlinear regression function, 'nlinfit', in MATLAB. The resulting L–H rate constant value (*k*) indicates how fast the dye molecule decomposes, which was dependent on the band gap energy of the photocatalyst, the accessible active radical species, such as hydroxyl, hydroperoxyl, or superoxide radicals, and the lattice oxygen defects. In the photo-irradiation process, the first step in the degradation pathway involves the photo-interaction of titanium centers in TNA, which produces charge carrier pairs within the moiety when the photon energy of incident light is higher than the bandgap of the titania phase (R-3). In the next step, excited titania centers abstract electrons in H<sub>2</sub>O<sub>2</sub> to form hydroxyl radicals (·OH) and hydroxyl anions (R-4). During the degradation/mineralization process, a radical attack on the dye center is more subtle and facile than an expedition by positive or negative ions. Hydrogen peroxide, known as an oxidant by itself [43], degraded the MO dye by only 45% after 1 h of reaction under sunlight illumination. H<sub>2</sub>O<sub>2</sub> is known to produce ·OH radicals when exposed to photon energies below 380 nm (R-5). However,

natural sunlight (or simulated) emits about 7% UV radiation in its spectrum, which explains the underlying reason for the weak but possible degradation activity. It took 6 h to degrade 88% of the MO dye, and prolonging the reaction time did not increase the degradation rate further. The annTNA alone showed a degradation rate of 10% after a reaction time of 1 h. In a light-irradiated TiO<sub>2</sub>-only system, hydroxyl radicals can be produced from water molecules in aqueous media, as represented in R-6. However, the density of  $TiO_2(h_{VB}^+)$  in a TiO<sub>2</sub>-only system is low because the anatase-phase TiO<sub>2</sub> has a bandgap energy of 3.6 eV, lying in the UV region as previously stated, and there is no trigger for charge separation of (h\_{VB}^+ ) and (e\_{CB}^- ) as in the case of H\_2O\_2 (R-4). The degradation rate of the MO dye by the combined system annTNA +  $H_2O_2$  is more than twice that of its components alone, presumably due to a synergistic effect in the oxidative degradation reactions similar to the Fenton process (R-4 and R-6). This makes the  $TiO_2$  substrate open, available, and accessible to proceed with reactions that continuously form radical species in situ, which is noteworthy for further exploration in this study. The respective dark reaction performed under the same conditions did not show any form of degradation of the dye molecules (Figure 2d).

$$TiO_2 + h\nu \rightarrow TiO_2 (e_{CB}^- + h_{VB}^+)$$
(3)

$$\mathrm{TiO}_{2}(\mathbf{e}_{\mathrm{CB}}^{-} + \mathbf{h}_{\mathrm{VB}}^{+}) + \mathrm{H}_{2}\mathrm{O}_{2} + \mathbf{h}\nu \to \mathrm{TiO}_{2}(\mathbf{h}_{\mathrm{VB}}^{+}) + \mathrm{OH} + \mathrm{OH} - \tag{4}$$

$$H_2O_2 + (h\nu < 380 \text{ nm}) \rightarrow 2^{\circ}OH$$
(5)

$$\mathrm{TiO}_{2}(\mathbf{h}_{\mathrm{VB}}^{+}) + \mathrm{H}_{2}\mathrm{O} \to \mathrm{TiO}_{2} + \mathrm{H}^{+} + \mathrm{OH}$$

$$\tag{6}$$

$$\operatorname{TiO}_2(\mathbf{e}_{\operatorname{CB}}^-) + \operatorname{O}_2 \to \operatorname{TiO}_2 + \operatorname{O}_2^{\cdot\cdot -} \tag{7}$$

$$OH + MO \rightarrow MO' + H_2O \rightarrow degraded products$$
 (8)

$$O_2^{\cdots} + MO \rightarrow MO' + H_2O \rightarrow \text{degraded products}$$
 (9)

$$H_2O_2 + OH \rightarrow H_2O + OH_2$$
(10)

$$\mathrm{TiO}_{2}\left(\mathrm{e}_{\mathrm{CB}}^{-}+\mathrm{h}_{\mathrm{VB}}^{+}\right)\to\mathrm{TiO}_{2}+\mathrm{heat} \tag{11}$$

Table 1 compares the annTNA material studied here with other catalytic systems for MO dye degradation. The dye discoloration efficiency by the "annTNA +  $H_2O_2$ " system strongly displays a substantial enhancement in the MO degradation rate.

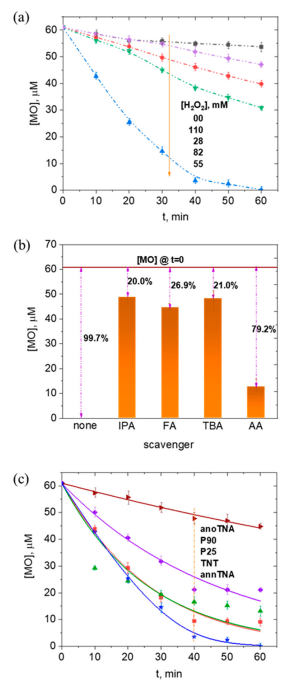
Entry	Catalyst		Experimental	Initial	Decoloration		
	Material	Form	Conditions [M		Efficiency/ %	Time/ min	Ref.
1	TiO <sub>2</sub>	film	external bias: 0.0 V vs. SCE Arc lamp, 165 mW/cm <sup>2</sup> pH~6.0	20	20	180	[38]
2	CdS/g-C <sub>3</sub> N <sub>4</sub>	powder	0.3 g/L catalyst W lamp, 100 W pH 3.4	2	40	60	[44]
3	ozone	gas	O <sub>3</sub> gas concentration 68.8 mg/L ultrasound irradiation 20 kHz heat	400	>99	20	[45]
4	Ag-P25	film	UV-A irradiation system, 350 nm, 71.7 μW/cm <sup>2</sup> pH 9.2	6	>99	120	[46]
5	Ag <sub>2</sub> CrO <sub>4</sub> /SnS <sub>2</sub>	powder	50 mg catalyst Xe lamp, 500 W pH not given	10	71	120	[47]

Entry	Catalyst		Experimental	Initial	Decoloration		
	Material	Form	Conditions	[MO]/ ppm	Efficiency/ %	Time/ min	Ref.
6	Fe from steel Industry waste	suspension	200 mg/L catalyst 34 mM H <sub>2</sub> O <sub>2</sub> pH 2	20	98	30	[48]
7	CuO on nanosized zeolite-X	powder	0.1 g/L catalyst Hg lamp, medium pressure pH 6	10	60	120	[49]
8	B-doped g-C <sub>3</sub> N <sub>4</sub>	powder	200 mg catalyst Xe lamp, 300 W pH 6.8	4	70	300	[50]
9	g-C <sub>3</sub> N <sub>4</sub>	powder	200 mg catalyst Xe lamp, 300 W pH 6.8	4	95	300	[50]
10	Au-TiO <sub>2</sub>	powder	20 mg catalyst W lamp, 500 W, 300 k lux pH not given	10	95	160	[51]
11	Ag–TiO <sub>2</sub> porous polymer	powder	50 mg catalyst Xe light, 100 mW/cm <sup>2</sup> pH not given	100	81.4	180	[52]
12	MoS <sub>2</sub> /YVO <sub>4</sub>	powder	100 mg catalyst Xe lamp, 300 W pH not given	10	98	60	[53]
13	cellulose-TiO <sub>2</sub>	sheet	2.5 cm × 2.5 cm film UV, 100 W lamp pH 3	5	95	150	[54]
14	80% BiOCl/BiOI	powder	20 mg catalyst Xe lamp, 300 W pH not given	50	75	375	[55]
15	Cu(OH) <sub>2</sub> -ZnO <sub>2</sub>	film	1.77 cm <sup>2</sup> film Hg lamp, 125 W pH not given	8	43	360	[56]
16	ZnO NPs	powder	50 mg catalyst UV lamp, 10 W bulb pH not given	15	85	180	[57]
17	$Nb_2O_5$	powder	5 mg catalyst UV lamp pH not given	20	99.9	20	[58]
18	TiO <sub>2</sub> –PLA	powder	50 mg catalyst UV lamp, 350 nm pH 8	320	81	300	[59]
19	TiO <sub>2</sub>	film	$55 \text{ mM H}_2\text{O}_2$ Xe lamp, 100 mW/cm <sup>2</sup> pH 3.4	20	>99	60	This work

Table 1. Cont.

# 3.3. Effect of $H_2O_2$ Concentration

In the annTNA–H<sub>2</sub>O<sub>2</sub> system, continuously and in situ generated hydroxyl radicals had a crucial role in MO dye degradation, similar to their role in a classical Fenton reaction, where Fe(II) is oxidized to Fe(III), generating ·OH. Hence, the MO degradation kinetics were evaluated over a range of H<sub>2</sub>O<sub>2</sub> concentrations. Keeping all other experimental parameters constant, increasing the concentration of the H<sub>2</sub>O<sub>2</sub> from 28 mM to 55 mM showed an exponential increase in the MO degradation rate. As shown in Figure 3a, the highest degradation of 99.7% was observed at 55 mM H<sub>2</sub>O<sub>2</sub>, further increasing the H<sub>2</sub>O<sub>2</sub> concentration to 82 mM and 110 mM had a detrimental effect on the overall dye degradation rate. It has previously been observed that exceeding a certain concentration of H<sub>2</sub>O<sub>2</sub> reduces the oxidative power of the whole system as it does not participate in the



**Figure 3.** (a) Effect of  $H_2O_2$  concentration on MO dye degradation. The legend indicates the concentration of  $H_2O_2$  in mM. (b) MO dye degradation rate plotted with equivalent quantities of different radical scavenges. IPA, FA, TBA, and AA refer to isopropanol, formic acid, tert-butanol, and ascorbic acid, respectively. The numbers represent the MO dye degradation rate in the solution after light irradiation for 1 h. (c) Effect of different types of titania catalyst materials on MO degradation. The symbols indicate kinetic measurements and the solid lines are nonlinear regressions with the L–H model.

# 3.4. Effect of Radical Scavengers

Most  $H_2O_2$ -catalyzed reactions are driven by  $\cdot OH$  radicals generated in situ. However, this is not the only possibility, and in certain multi-component systems, the reaction is

driven by superoxide radicals  $(O_2 \cdot \overline{})$ . Hence, to gain an insight into the MO degradation mechanism in this study, the photocatalytic reaction was performed under optimized conditions (20 ppm MO and 55 mM H<sub>2</sub>O<sub>2</sub>) in the presence of both hydroxyl and superoxide radical scavengers. Isopropyl alcohol (IPA), formic acid (FA), and tert-butanol (TBA) are known as ·OH radical scavengers [36], and ascorbic acid (AA) is known to quench superoxide radicals ( $O_2$ ·<sup>-</sup>). The MO dye degradation rates shown in Figure 3b were affected by all scavengers used regardless of their type. However, the reduction in the MO degradation rate was much greater with the hydroxyl radical scavengers, and it declined by more than 70% compared to the original MO degradation rate (without scavengers). The MO degradation rates were 20, 26.9, and 21% in the systems with IPA, FA, and TBA, respectively, whereas the rate reduction was relatively low at 79.2% in the system with AA. The experimental results are consistent with the known fact that  $H_2O_2$  acts as a precursor to the formation of  $\cdot$ OH and O<sub>2</sub> $\cdot$ <sup>-</sup>, clearly demonstrating that the present degradation reaction is mainly driven by  $\cdot OH$ , while  $O_2 \cdot \overline{\phantom{0}}$  affects the reaction insignificantly. Zhao et al. found that the surface complexation of  $TiO_2$  and  $H_2O_2$  leads to the formation of (Ti(IV)-OOH), resulting in •OH formation under visible light irradiation [63]. Kim et al. employed a TNA film catalyst combined with  $H_2O_2$  to oxidize As(III) to As(V) [36]. Their observations with radical scavengers contrasted with those observed in the current MO dye degradation system, where hydroxyl radical scavengers did not affect the overall reaction rate, while the oxidation of As(III) was significantly reduced in the presence of superoxide radical scavengers. Many studies have shown that most of the oxidation reactions of  $TiO_2$ -based photocatalysts are induced by superoxide radicals when anatase is the only phase present in the TiO<sub>2</sub> catalyst [64,65]. However, the presence of trace amounts of the rutile phase has been shown to deviate from the conversion of superoxide radicals from  $H_2O_2$  to hydroxyl radicals [66–68].

## 3.5. Effect of Titania Type

In this regard, other forms and polymorphs of titania were also evaluated for their ability to catalyze the oxidation process synergistically. P25, P90, titanate nanotubes (TNT, synthesized in the lab) powders, and as-anodized TNA (anoTNA) were employed in the MO degradation system under otherwise optimized conditions, as shown in Figure 3c. A comparison of all titania materials showed that the MO decomposition activity of anoTNA film with rutile only was the lowest, possibly due to the absence of a complete anatase phase. In addition, the degradation rate of MO by P25 was better than P90. P25 is a physical crystalline mixture with an anatase-to-rutile ratio of 75:25, while P90 has a ratio of 10:90, meaning that P90 has a higher rutile content than P25. After 10 min of reaction, P25 showed the highest MO degradation rate, which could be attributed to faster charge carrier separation and coupling with H<sub>2</sub>O<sub>2</sub> than other titania materials. However, as the reaction time continued, annTNA provided the superior MO degradation kinetics, indicating that the catalytic activity of titania increases with less rutile in the titania material. Romanos et al. modified commercial P25 with well-dispersed Cu nanoparticles, and the system was found to degrade 50% of 12 ppm initial dye after exposure to UV light in the 350–390 nm range for 3 h [69]. In another process, a bulk P25 composite using Posidonia oceanica fibers showed 100% degradation of phenol after 4 h of UV light irradiation [70]. It is noteworthy that the annTNT-H<sub>2</sub>O<sub>2</sub> system outperforms commercial P25 in this study. While anatase plays a crucial role in the light-driven process, the trace presence of rutile increases the abundance of hydroxyl radicals, which are the active attacker. TNT powder shows a very good performance, with an 85% MO degradation rate, second only to annTNA, revealing the importance of morphological effects, especially for regular tubular channels. Therefore, based on the observations of various radical scavengers combined with various titania materials, we make a reasonable conclusion that the annealed TNA films (the active catalyst in this study) contain a trace amount of rutile phase in the main anatase moiety. The L–H rate constant and adsorption coefficient values for the previously discussed systems are given in Table 2. The annTNA system has an L–H rate constant of 0.407  $\mu$ mol<sup>-1</sup> min<sup>-1</sup>, which is almost 10 times higher than that of anoTNA, at 0.045  $\mu$ mol<sup>-1</sup> min<sup>-1</sup>, while that of TNT is 73% higher than anoTNA, at 0.078  $\mu$ mol<sup>-1</sup> min<sup>-1</sup>.

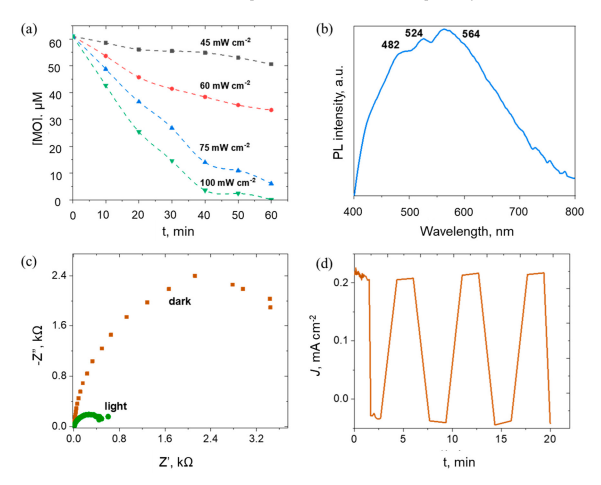
Table 2. The L-H rate constants of MO dye degradation by different titania materials.

TiO <sub>2</sub> Material	L−H Rate Constant, k (µmol <sup>−1</sup> min <sup>−1</sup> )	Adsorption Coefficient, K (mL μmol <sup>-1</sup> )	TiO <sub>2</sub> Material	L–H Rate Constant, <i>k</i> (µmol <sup>-1</sup> min <sup>-1</sup> )	Adsorption Coefficient, <i>K</i> (mL μmol <sup>-1</sup> )
annTNA	0.407	59.04	P25	0.00026	0.01
anoTNA	0.045	0.25	P90	0.005	0.11
TNT	0.078	3.36	ann TNA *	0.167	0.39

<sup>\*</sup> annTNA system in the absence of  $H_2O_2$ , all other systems consist of 55 mM  $H_2O_2$  in each, 20 ppm MO dye, and exposed to AM 1.5 G light with 100 mW cm<sup>-2</sup> intensity for 60 min.

## 3.6. Effect of Light Intensity

As already mentioned for light-induced reactions, light intensity makes a pivotal contribution, as it is directly linked to the number of photons reaching the titania catalytic centers of TNA, influencing charge carrier generation and separation (R-3). The effect of light intensity on the overall MO dye degradation rate in the optimized system was studied at four different light intensities: 45, 60, 75, and 100 mW cm<sup>-2</sup>. As shown in Figure 4a, the light intensity has a positive non-linear relationship with the MO degradation rate. As the light intensity increased to 60, 75, and 100 mW cm<sup>-2</sup>, the MO degradation rate of 17% at 45 mW cm<sup>-2</sup> was improved to 45, 90, and 99%, respectively.



**Figure 4.** (a) Effect of light intensity on the MO dye degradation rate. (b) PL intensity of annTNA film for excitation at 375 nm. (c) EIS performed under light/dark conditions from 50 mHz to 500 kHz in 0.1 M Na<sub>2</sub>SO<sub>4</sub> electrolyte. (d) Chopped chronoamperometry test performed at 0.0 V (vs. Ag/AgCl).

The photoluminescence (PL) spectra presented in Figure 4b provide meaningful insights into the excitation behavior of the fabricated annTNA film. The measured PL emission upon excitation at 375 nm shows three peaks at 482, 524, and 562 nm, all falling in the visible region. These blue and green level emissions can be partly attributed to defect sites in the annTNA film material arising from oxygen vacancies, which greatly contribute to improving the photoactivity of the annTNA film in the visible region.

## 3.7. Electrochemical Characterization of annTNA Film

The important role of light in the MO degradation reaction was further assessed and affirmed by a set of electrochemical tests employing annTNA as the working electrode in a three-electrode configuration. Charge carrier transport resistance in light and dark-assisted media was studied by electrochemical impedance spectroscopy (EIS) tests performed in 0.1 M Na<sub>2</sub>SO<sub>4</sub>. As shown in Figure 4c, the charge carrier transport resistance was low under the light illumination condition, indicated by a lower arc radius than in the dark (larger arc radius). The charge carrier generation with intermittent light illumination was analyzed by chopped chronoamperometry (Figure 4d). The strong and fast response of annTNA to the incident light was demonstrated by the troughs and crests of the chopped chronoamperogram.

#### 3.8. Tentative Reaction Mechanism

Based on the experimental results, it can be suggested that the main cause of MO dye degradation in the annTNA and  $H_2O_2$  combined system is the reactive hydroxyl radicals formed repeatedly under light illumination (Figure 5).

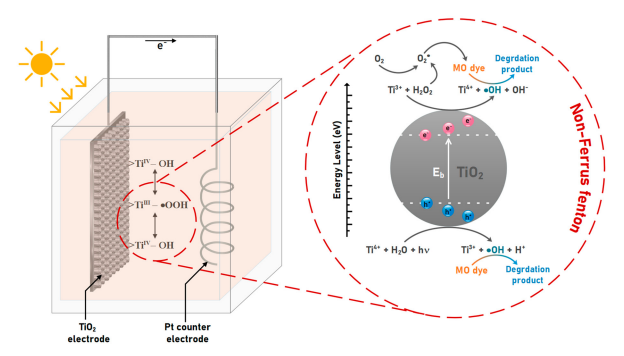
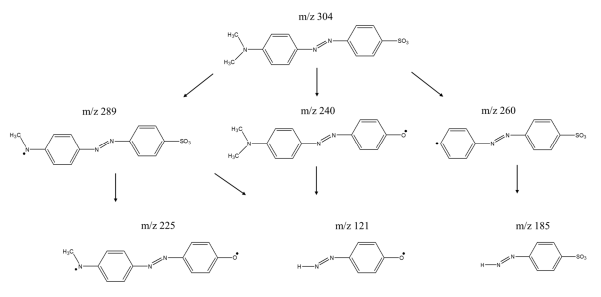


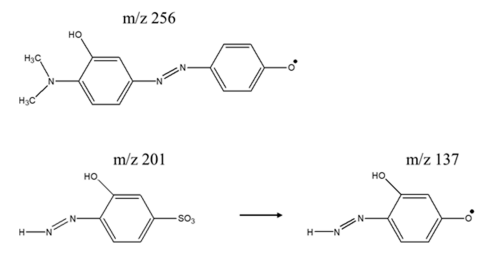
Figure 5. (Left) Experimental setup and (right) proposed reaction pathway.

Reaction products were also studied by total organic carbon (TOC) analysis on samples collected at regular intervals. As the reaction time increased, the TOC decreased, while the total carbon (TC) content remained consistent, indicating that the MO dye was partially mineralized with fragmentation. During mineralization, carbon components of the dye were transformed into  $CO_2$  or became bicarbonate (HCO<sub>3</sub><sup>-</sup>) in an acidic medium (pH 3), which was detected as inorganic carbon during the test. An LC/MS/MS tandem analysis revealed fragmented masses corresponding to m/z peaks at 304, 290, 260, 240, 225, and 201 in the first step, and 289, 275, 240, 256, 225, 185, 137, 121, and 112 in the next step (Figure S1).

According to the prior literature [71], fragmentation products related to the breakdown of methyl orange dye molecules were proposed, as shown in Scheme 1. Other fragmentation products were associated with impurities in the initial dye and are summarized in Scheme 2.



Scheme 1. Fragmentated products based on the LC/MS/MS analysis.

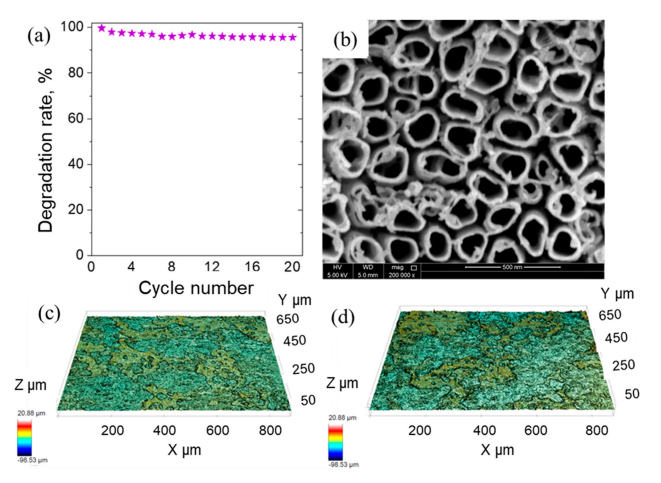


Scheme 2. Identified fragmentation products of impurities in the MO dye, assessed by LC/MS/MS.

## 3.9. Reusability of annTNA Photocatalyst

The recycling ability of the TNA electrode was studied for 20 repeated MO degradation cycles in an optimized environment. After each experiment, the TNA electrode was gently washed with DI and air-dried overnight to reuse in subsequent cycle experiments. As shown in Figure 6a, there was no significant difference in MO dye degradation rates during the 20 cycles. Furthermore, excellent structural integrity over repeated cycles was best evidenced by the SEM image of the TNA electrode evaluated after repeated degradation cycles, proven by undisturbed tubular arrays (Figure 6b). The stability of the annealed film under the consecutive dye degradation runs was evaluated by profilometry, as shown in Figure 6c,d, and a high stability of the film was observed.

In addition to its high catalytic activity, the use of TNA films has significant advantages in separating and recycling the catalyst from the medium. This is mainly due to the ease of managing the operational conditions. This system operated in an environment close to ambient temperature and pressure, indicating the cost-competitive nature of the system when scaled up and its ability to utilize abundant sunlight for wastewater treatment, particularly of textile waste. The system requires an in depth tolerance analysis [72]. Thus, the developed facile regenerative and self-supporting titania nanotube array– $H_2O_2$  coupling process will be evaluated for other contaminants in similar categories, such as phenol, bisphenol A, methyl blue, SRB dye, etc., to understand the capabilities of the TNA– $H_2O_2$  system. The vision is to upscale the system to an industrial level to reach Technology Readiness Level (TRL)-7 or higher to treat real complex industrial wastewater.



**Figure 6.** (a) MO degradation (%) over 20 cycles. (b) SEM image of the TNA film at 500 nm scale bar after 20 cycles of operation. Comparison of surface topography of the annTNA film (c) before and (d) after use in the MO degradation test: Color codes in (c) and (d) indicate depth relative to the ground plane.

## 4. Conclusions

Herein, we demonstrated the potential of a film-based annTNA and  $H_2O_2$  coupled system to degrade methyl orange, an azobenzene-type commercial textile dye, via a non-ferrous Fenton-type advanced oxidation process (AOP) under solar illumination with remarkable reaction kinetics. Methyl orange dye was decomposed by almost 99.7% after 60 min of solar irradiation (100 mW cm<sup>-2</sup>, AM 1.5G). TOC, LC/MS/MS, and UV–Vis spectroscopic analysis of the reaction products confirmed that the photocatalytic degradation proceeded by partial fractionation and mineralization. The experimental data were in good agreement with the Langmuir–Hinshelwood model, following pseudo-first-order reaction kinetics. The simple regeneration ability of the self-supporting catalyst through gentle washing of the film with deionized water and its consistent degradation performance (<5% change) over 20 cycles encourages the use of the system on a pilot scale to treat real wastewater.

**Supplementary Materials:** The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/nano13040639/s1, Figure S1: LC/MS/MS chromatograms.

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Conflicts of Interest: The authors declare no conflict of interest.

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