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Heterogeneous photocatalysis using UV/VIS light or natural solar radiation as the energy source is one of the most efficient advanced oxidation processes (AOP) for the removal of chemical and bacterial pollutants from water. One of the least investigated oxides in this context, i.e. MnO₂ and its combination MnO₂/TiO₂ are examined as potential photocatalysts for the removal of Indigo carmine (IC) dye pollutant from the water. The catalysts are characterized by XRD, FTIR, SEM, TEM, adsorption and surface area measurements. While MnO₂ is very efficient for the decolorization of the dye, it is not effective enough for mineralization. MnO₂/TiO₂ as the photocatalyst at the optimized ratio of 9:1 combines the advantages of both the oxides, i.e., rapid decolorization and efficient mineralization. Persulphate $(S_2O_8^{2-})$ enhances the degradation while H_2O_2 inhibits the same. The degradation is dependent on pH with higher degradation under extremely acidic conditions. The influence of dissolved salts/anions in the water on the degradation varies from 'moderate inhibition' to 'no effect' or even 'enhancement' depending on the chemistry of the anion and reaction conditions. Effect of various parameters such as reaction time, substrate concentration, catalyst dosage, the presence of O2 recycling of the catalyst, etc. on the efficiency of degradation is investigated. The results are critically analyzed, and a tentative mechanism is proposed.

Introduction

Heterogeneous photocatalysis mediated by semiconductor (SC) oxides is an efficient advanced oxidation process (AOP) for the decontamination of water from chemical and bacterial pollutants.¹⁻⁵ When the SC oxide is irradiated with a light energy source (UV or visible) such that hv > Eg(band gap of the SC), electron-hole pairs are created. The electrons get excited from the valence band to the conduction band leaving positively charged holes in the valence band. These holes and electrons can participate in oxidative and reductive reactions respectively. In aqueous suspensions, highly reactive OH radicals are generated, and they can promote the oxidation and eventual mineralization of the pollutant to CO_2 , water, and inorganic salts.

The most widely investigated SC oxides in this respect are ZnO and TiO₂ in view of their favorable physicochemical properties, low cost, easy availability, high stability and low toxicity. The band gaps of ZnO (~3.2 eV) and TiO₂ (anatase $E_{\rm bg} = \sim 3.2$ eV, rutile $E_{\rm bg} = \sim 3.0$ eV) are fairly similar. Hence they can absorb light only below 400 nm, which is in the UV range that constitutes only <5 % of the sunlight. ZnO is relatively more active in the visible region and is hence preferred in the investigations aimed at the harnessing of solar energy.⁶⁻⁹ However, it has the disadvantage of corrosion under extreme acidic and alkaline conditions as well as photocorrosion making the scope of its commercial application limited.

A number of studies have been reported on the modification of semiconductor oxides in order to extend the absorption of light to the visible range. These include dye sensitization, semiconductor coupling, impurity doping, use of coordination metal complexes and metal deposition.^{4,10-14} Composites such as TiO₂/carbon have also been reported.¹⁵ Physical modification of TiO2 with small amounts of transition metal cations such as vanadium and chromium extends the absorption up to 550 nm and makes it active under UV as well as sunlight.¹⁶ Deposition of noble metals such as Pt, Pd, Au, Ag etc. on TiO₂ enhances the catalytic oxidation of organic pollutants due to the increased light absorption and retardation of the recombination of photogenerated electrons and holes.17-19

MnO₂ is one of the least investigated SC oxides in photocatalysis in spite of its excellent semiconductivity, porosity, and mixed-valent properties. The combination of MnO₂ with TiO₂ may offer advantages of the high adsorption and reactivity of the former and photochemical property of the latter. Considering these, the application of MnO₂ and MnO₂/TiO₂ as photocatalysts for the removal of recalcitrant toxic materials is examined here. One of the widely used special application dyes Indigo Carmine (IC) is selected as the test candidate.

Indigo carmine (3,3'-dioxo-1,3,1',3'-tetrahydro-[2,2']-biindolylidene-5,5'-disulphonic acid disodium salt) is an important dve used for dving of clothes and also for medical diagnostic purposes. It can also help to target biopsies even better since the homogeneously stained or unstained areas can be correlated with intraepithelial neoplasia.²⁰ IC is highly toxic and potentially carcinogenic.²¹ Extensive application of the dye leads to environmental contamination, in particular, water pollution.

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The chemical structure of IC is given in Figure 1.



Figure 1. Chemical structure of Indigo carmine

Only a few investigations aimed at the removal IC from water have been reported.²²⁻²⁶ To the best of our knowledge, the current study is the first instance of the application of MnO_2 mediated photocatalysis for the degradation/mineralization of IC from the water. The efficacy of the technique is evaluated under various reaction conditions, and the optimum parameters are identified.

Experimental

Materials

MnO₂ used in the study was supplied by Merck India Limited. The particles were approximately spherical and nonporous with > 99 % purity. The BET surface area was ~31m²/g. The pore volume was ~ 0.05 cm³ g⁻¹, and the average pore width was 74.2 Å. Degussa P-25 TiO₂ (> 99 % pure, average particle size ~ 15-20 µm, BET surface area ~15 m² g⁻¹) consisting of ~80 % anatase and ~20 % rutile was used as such without further purification.

Persulphate (PS) (purity ~99 %) and H_2O_2 (30.0 % w/v), both from Qualigen (India) and Indigo Carmine (Extra Pure Grade > 99.5 % purity) from Sisco Research Laboratories Pvt. Ltd (India) were used as such without further purification. Doubly distilled water was used in all the experiments. All other chemicals were of AnalaR Grade or equivalent.

Characterization of the catalyst

X-ray diffraction measurements were made by using Rigaku X-ray diffractometer with CuK_{α} radiation. Scanning Electron Microscopy (SEM) measurements were performed with JEOL Model JSM-6390LV. Diffuse Reflectance Spectra (DRS) were recorded with Varian Cary 5000 using BaSO₄ as the reference. Transmission electron microscopy (TEM) was done using JOEL/JEM/200; source LaB₆. FTIR spectra were recorded using Nicolet Avatar 370 instrument equipped with triglycine sulfate detector with a KBr window and a KBr beam splitter. The equipment was purged with dry air to prevent interference from atmospheric moisture. The transmission spectrum was obtained with a resolution of 4 cm⁻¹ by using 32 scans in the range 400 – 4000 cm⁻¹. The powdered samples were diluted with finely ground KBr to about 5 % by weight.

Photocatalytic experiments

Photocatalytic experiments in the presence of UV were performed in a jacketed Pyrex reactor. The dye solution together with weighed amount of the catalyst and other additives as required was taken in the inner compartment and cooling water from a thermostat $(27\pm1^{\circ}C)$ was circulated through the outer jacket. UV irradiation was done using a 400 W medium pressure mercury vapor lamp. The suspension was frequently stirred to ensure uniform mixing. Samples were drawn periodically, centrifuged and analyzed for the dye concentration by Spectrophotometry (608 nm). Suspension kept under identical conditions in the dark was used as the reference in each case to eliminate the contribution from adsorption towards the reduction in the dye concentration. H₂O₂ was determined by standard iodometry. Chemical oxygen demand (COD) measurements were made by the open reflux method.²⁷ Adsorption studies were made by the standard technique.²⁸

Detection of hydroxyl radicals

The formation of OH radicals during the UV irradiation of the reaction system was tested by the photoluminescence (PL) technique using terephthalic acid (TPA) as the probe molecule.^{29,30}

The hydroxyl radicals formed in situ in the system reacts with TPA to form the 2-hydroxyterephthalic acid (HTPA), which is a fluorescent molecule. The intensity of its PL is proportional to the concentration of in situ formed 'OH radicals in the system. MnO₂ at the standard experimental concentrations is suspended in an aqueous solution of TPA (2 x 10⁻⁴ M) and NaOH (2x10⁻³ M) and irradiated by UV light. The PL spectrum of the product HTPA is recorded in the range of 400 - 450 nm after every 5 minutes of irradiation. The excitation wavelength was 315 nm. The PL intensity at 425 nm corresponds to the concentration of HTPA and hence of the OH radicals formed in the system. model **RF-5301PC** Shimadzu fluorescence spectrophotometer is used for recording the spectrum.

Results and Discussions

Characterization of MnO₂

The catalysts MnO_2 and TiO_2 were characterized by XRD, SEM, TEM, Fourier transform infrared (FTIR) spectroscopy and DRS. Typical results with respect to MnO_2 are given in Figures 2 A, B, C, D and E. The characteristics of TiO_2 are same as those reported earlier.³¹

The XRD results show that the MnO_2 is amorphous in nature with weak diffraction pattern (Figure 2A). The morphology and particle size analysis were done using SEM (Figure. 2B). Particles were approximately rod-shaped with average particle size in the range of 220 nm. The characteristics of the particles are appropriate for efficient photocatalytic properties such as light absorption, diffusion as well as regeneration of the catalyst by suitable treatment.



Figure 2A. XRD image of MnO₂.



Figure 2B. SEM image of MnO₂.

The morphology and particle size are further confirmed by TEM analysis (Figure 2C) which shows platelet and rod-like structure. The FTIR pattern (Figure 2D) shows very week absorption intensity of OH (3100-3600 cm⁻¹) or H₂O (1600 and 3600 cm⁻¹). The general spectral pattern shows that MnO_2 is pure with no contamination from any of the precursors.



Figure 2C. TEM image of MnO₂.

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Figure 2D. FTIR spectrum of MnO₂.

Diffuse reflectance spectrum (Figure 2E) shows a sharp reflectance band between 200 and 300nm indicating that the material absorbs in the UV range. The optical absorption intensity of MnO_2 steadily increases from 300 to 800 nm with no clear band edge. This indicates that MnO_2 is not a typical semiconductor like TiO₂ or ZnO. The brown color of MnO_2 also indicates favorable absorption of visible light.³²



Figure 2E. DRS of MnO₂.

Photocatalytic degradation of IC on MnO₂

Photocatalytic degradation of IC in the presence of MnO_2 and UV irradiation is tested, and relevant parameters are optimized. MnO_2 is a good adsorbent of IC, and hence there is a significant decrease in the concentration of the dye initially even without irradiation. However, once the optimum adsorption under the experimental conditions is reached, no further decrease is noticed. When the suspension is exposed to UV irradiation, the concentration of IC decreases rapidly leading to eventual decolorization. In the absence of the catalyst also, moderate decolorization/degradation of the dye takes place on exposure to UV light. Preliminary results are plotted in Figure 3.



Figure 3. Photocatalytic degradation of IC in the presence of MnO2.

Effect of catalyst dosage

The effect of MnO_2 loading on the photocatalytic degradation of IC is tested by varying the dosage of MnO2, keeping other parameters constant. The results are plotted in Figure 4 which shows that the degradation of the dye increases with increase in catalyst dosage and stabilizes at 0.14 g in 100 mL under the reaction conditions.



Figure 4. Effect of MnO2 dosage on the photocatalytic degradation of IC.

The enhanced degradation at higher catalyst loadings can be attributed to the increased number of adsorption and interaction sites for the IC and/or the intermediates and more effective harvesting of light. This leads to increased formation of reactive hydroxyl and other radicals which are known to be responsible for the photocatalytic degradation of organic pollutants. Any increase in catalyst concentration beyond the optimum will only result in scattering and reduced passage of light through the suspension medium. Another reason may be the aggregation of catalyst particles causing a decrease in the number of available active surface sites. Beyond a particular loading, the catalyst particles cannot be fully and effectively suspended in a particular reactor which also leads to suboptimal penetration of light and reduced adsorption of the substrate on the surface. It is also possible that at higher loading, part of the originally activated MnO₂ is deactivated through collision with ground state catalyst according to the following equation.³³:

$$MO^* + MO \longrightarrow MO^{\#} + MO$$
 (1)

where MO represents MnO_2 , MO^* and $MO^{\#}$ are its activated and deactivated forms, respectively).

In the case of MnO_2 which is a good adsorber/absorber and reservoir for O_2 , higher dosage provides more photogenerated reactive oxygen species (ROS) such as 'OH radicals. The formation of 'OH radicals in $MnO_2/IC/UV$ is confirmed by PL spectral studies. The increase in fluorescence with irradiation time (Figure 5) corresponds to the progressive formation of 'OH radicals (See experimental section).



Figure 5. PL spectral changes during UV irradiation of MnO₂/terephthalic acid/NaOH.

Cao and Suib³² suggested from binding energy calculations that MnO₂ is a mixed-valent system consisting of Mn(4+) and Mn(3+). The multiple oxidation states enable it to function as good electron donor/acceptor and hence oxidation-reduction catalyst.³⁴ This property favors increased photocatalytic activity with an increase in catalyst dosage. This also helps the facile regeneration of the used catalyst by replenishing the depleted oxygen. Surface and lattice oxygen species in MnO₂ are known to take part in the catalytic oxidation process. Photolysis of MnO₂ is reported to result in a five-fold increase in the surface oxygen, either by migration of oxygen to the surface or by migration of Mn to the bulk or both.³² During the photocatalytic reaction, surface oxygen is consumed, and bulk oxygen moves to the surface. Hence, the more the catalyst dosage, the more the availability of reactive oxygen and consequently better the degradation. This can continue until other reaction parameters, as well as the size, shape, and geometry of the reactor assembly, are appropriate to facilitate the degradation process. Hence, for each reactor configuration, the optimization of catalyst dosage has to be made separately. In the present case, the optimum loading of MnO_2 is 0.14 g in 100 mL. Hence all further studies were carried out with this loading.

Effect of concentration of IC

The effect of initial concentration of IC in the range of 10-60 mg/L in the water on its photocatalytic degradation is investigated. The percentage degradation decreases as the concentration of IC increases. In contrast, the rate of removal increases steadily and steeply with an increase in the initial concentration of IC up to 40 mg L⁻¹ and stabilizes after that (Figure 6). The optimum kinetic rate of 1.9 mg L⁻¹ min⁻¹ is reached at 40 mg L⁻¹ of IC. This is followed by a slight decrease and eventual stabilization of the rate implying gradually decreasing and eventual zero order kinetics at higher concentration of >40 mg L⁻¹.



Figure 6. Effect of concentration of IC on the rate of photocatalytic degradation.

The photocatalytic degradation of many organic pollutants in water is reported to follow pseudo first order kinetics.³⁵⁻³⁷ The increase in degradation with an increase in concentration up to 40 mg L⁻¹may is due to increase in adsorption of IC on MnO2 which will continue until the surface is fully covered and all the active sites are occupied. Further, at higher concentration, there will be more IC molecules available in bulk as well as in the proximity of the surface which can also interact with the surface-generated active free radicals and ROS such as OH and H₂O₂. At higher concentration beyond the optimum, at least a part of the irradiating UV light may be blocked and/or absorbed by IC and the intermediates formed from it thereby reducing the photons available for catalyst activation. Another reason may be retardation in the penetration of light at a longer distance from the light source and consequent decrease in its path length. This results in decreased availability of photons per IC molecule in the system and hence decreased degradation.³⁸ As the initial concentration of IC increases, the requirement of catalyst surface needed for maintaining the corresponding rate of degradation also increases. Since illumination intensity and catalyst concentration are maintained constant, the relative number of ROS available per IC molecule is also presumably decreasing with increase in the concentration of IC.

This also leads to stabilized/ decreased rate of degradation. Once the concentration of the substrate is enough to interact with all the optimum available ROS and/or other reactive free radicals, any further increase cannot result in increased reaction, and the IC removal becomes independent of concentration. It is also possible that at higher substrate concentration, some of the reaction intermediates may get adsorbed onto the surface or remain in bulk for the relatively longer period leading to the less frequent interaction between fresh IC molecules and the ROS. At any point in time during the irradiation, there will be an optimum for the number of substrate molecules that can interact with the reactive free radicals generated in the system. This optimum will depend on some reaction parameters such as initial concentration of the substrate, pH, intensity of illumination, the wavelength of light, mass and type of photocatalyst, type and geometry of photoreactor, the presence of other contaminants, etc. Consequently, the optimization will apply only to the specific reaction conditions and cannot be generalized.

Since both catalyst and light are essential for the degradation, the negative effect of increasing concentration of IC on the rate of degradation implies that at higher concentration, the dye is inhibiting the action of the catalyst and/or light. At higher concentration, there will be better adsorption of the dye on the surface of the catalyst. This will prevent direct absorption of light by the catalyst thereby inhibiting the generation of reactive free radicals and ROS. In the absence of steady and significant degradation, the adsorbed dye will not leave the surface sites. This will prevent the adsorption of fresh dye molecules and continued degradation.

The simplest way to represent the rate of photocatalytic degradation is the Langmuir-Hinshelwood model, modified to accommodate reactions taking place at the solid-liquid interface. Accordingly, the rate of degradation is:

$$r_0 = \frac{-dC}{dt} = \frac{kKC_0}{1+KC_0} \tag{2}$$

where

 r_0 is the rate of disappearance of IC (mg L⁻¹ min⁻¹), C_0 (mg L⁻¹) is the initial concentration of the dye, *C* is the concentration at time t, *K* is the equilibrium adsorption coefficient (mgL⁻¹) and *k* is the reaction rate constant (min⁻¹) at maximum surface coverage.

Equation 2 can be integrated and presented as

$$t = \frac{1}{kK} \ln \frac{C_0}{C} + \frac{1}{k} \left(C_0 - C \right)$$
(3)

when C_0 is very small, (C_0-C) is negligible and equation (3) becomes

Removal of indigo carmine from water

$$\ln \frac{C_0}{C} = kKt = k't \tag{4}$$

In the present study, the logarithmic plot of $\ln(C_0/C)$ vs. time t did not give straight lines that pass through the origin. This shows that the degradation of IC in the presence of MnO₂ does not follow the expected pseudo first order kinetics as in the case of most photocatalytic processes. The Langmuir-Hinshelwood model and the pseudo-first order kinetics underline the key role of adsorption on the rate of reaction assuming that the adsorption-desorption kinetics is faster than the photochemical reaction.^{39,40} The rate of unimolecular surface reaction is proportional to the surface coverage, assuming that the reactant is more strongly adsorbed than the products on the catalyst particles.⁴¹

Adsorption studies of IC on MnO₂ under the reaction conditions showed that~32 % of the dye is adsorbed at the equilibrated stage. When K is large, the first term of equation (3) is negligible. i.e.

$$C_0 - C = kt \tag{5}$$

The concentration of the dye decreases with time until the optimum surface coverage is attained.

The pseudo first order kinetics, as in the case of typical semiconductor oxide photocatalysts such as ZnO (adsorption: ~ 11 %) and TiO₂ (adsorption: ~ 8 %), is not followed in the case of MnO_2 with high adsorption (~32 %). Hence it may be inferred that the degree of adsorption of the substrate on the catalyst does influence the degradation kinetics. This further indicates that the general assumptions implicit in the application of LH mechanism and related kinetic analysis are not fully applicable in the case of MnO₂ mediated photocatalysis. These assumptions are:42

- i) The reaction of adsorbed species is the ratedetermining step
- ii) Fractional surface coverage (θ_A) of the reactant species is the more appropriate concentration parameter to be applied in rate laws,
- iii) $\theta_A \rightarrow 1$ for saturation coverage corresponding to a monolayer of adsorbate on the surface and
- iv) Langmuir adsorption isotherm type equation represents the relationship between θ_A and the activity. In the case of highly adsorbing particles such as MnO₂, there will be competition between the solvent (water) and the solute (IC) molecules for adsorption sites which can complicate the kinetics of adsorption as well as degradation.

Effect of pH

The pH of the reaction medium is known to influence the surface characteristics and adsorptive properties and consequently the photocatalytic behavior of semiconductor oxides. The effect of pH on the photodegradation of IC in is observed at pH 2. However, under this condition, the decolorization is very facile even without UV irradiation.

the presence of MnO₂ is investigated, and the results are given in Figure 7. Maximum degradation (~97 % in 10 min)



Figure 7. Effect of pH on the photocatalytic degradation of IC over MnO₂

The pH has only negligible effect on the decolorization of the dye as such (in the absence of UV and MnO₂). However,

under UV, even the dye alone undergoes reasonable

decolorization at various pH, the maximum being at pH 2.

The degradation decreases steadily with the increase in pH

up to pH 10 and increases slightly after that. The fast decolorization in the presence of MnO₂ under acidic pH, even in the absence of UV irradiation, shows that simple adsorption of IC on MnO₂ also is facilitated at acidic pH. Similar pH effect on the adsorption of IC on TiO₂ catalyst has been reported by Barka et al."43 pH is known to influence the surface state of the catalyst and ionization state of the dye. The point of zero charges (PZC) of MnO_2 is ~ 4.7.^{26,44,45} Below this pH the surface is positively charged, and hence its oxidizing ability will be relatively higher. IC is a dianionic dye in aqueous solution, and this configuration is maintained in the pH range 3-11.^{24,45} Below the PZC of MnO₂, the electrostatic interaction between the positive catalyst surface and the dianions can lead to strong interaction between them and subsequent degradation of the dye. At higher pH when the surface is negatively charged, the adsorption is less due to the electrostatic repulsion between the negatively charged surface and the dye. It is also possible that the heterogeneous degradation in the presence of MnO₂ involves the formation of precursor complex between the dye and surface bound Mn.⁴⁶ Electrons from the activated dye move to the Mn(IV) on the surface of Consequently, the dye undergoes oxidative MnO₂ degradation. The Mn(II) thus formed reductively from MnO₂ dissolves in the bulk solution. Thus MnO₂ functions both as a catalyst as well as an oxidant. The Mn(II) is oxidized to Mn(IV) oxide again by the dissolved O₂. In this respect, MnO₂ plays the role of a catalyst and dissolved O₂

acts as the oxidant. The re-adsorption of free Mn(II) ions in

solution back onto MnO₂ surface is facilitated at pH beyond

the PZC. At the natural pH, ~5.5 of the reaction system, the

decolorization of IC in the presence of MnO_2/UV is 50% more efficient than simple adsorption. High degradation of IC at pH ~2 shows that it is induced by extremely acidic conditions. UV irradiation and MnO_2 accelerate the degradation. The slight increase in degradation at pH~ 11 may be due to increased number of reactive 'OH radicals formed from the higher concentration of OH⁻ ions.

The effect of pH on the photocatalytic efficiency of a highly adsorbing and reactive oxide such as MnO₂ may not be simple or straightforward because it depends on the complex interplay of multiple factors, i.e. chemistry of the substrate and the surface, extent, and mode of adsorption (of IC as well as the numerous intermediates), concentration and nature of interactions of reactive free radicals, etc. The adsorption of the substrates is more in the acidic range while the concentration of the 'OH radical is more in the alkaline region as explained earlier. Thus the effect of pH on these two parameters, which are important for the degradation of IC, is different and opposing. As a consequence, the combined net effect need not necessarily be the same quantitatively even at the same pH at different times. Hence precise correlation of the effect of pH on the adsorption/degradation of IC with the PZC of MnO₂ will not be fully consistent. Lack of direct correlation between the PZC and the adsorption/degradation rate can also be because the PZC itself depends on some factors including the size and nature of dispersion of the particles, chemistry of substrates and the intermediates and the type of catalyst itself.

Effect of H₂O₂

 H_2O_2 , which is concurrently formed in many photocatalytic processes is a powerful oxidant and has been widely investigated for its potential to enhance the mineralization of organic water pollutants. The enhancement is caused by the photocatalytic generation of extra 'OH radicals as follows:

$$H_2O_2 \rightarrow 2^{\circ}OH$$
 (6)

In this context, the effect of externally added H_2O_2 on the photocatalytic decolorization of IC is tested under different conditions. In the absence of MnO₂, under UV irradiation, the degradation of IC increases with increase in the concentration of H_2O_2 and reaches a plateau at 50 mg L⁻¹ (Figure 8A).

The initial decrease in concentration of IC in the presence of the only MnO_2 can be attributed to simple adsorption. In the presence of MnO_2 and H_2O_2 (at 10 mg L⁻¹), initially, there is inhibition in the % decolorization of IC compared to that in the presence of MnO_2 only (Figure 8A). With the increase in the concentration of H_2O_2 , the net concentration of IC in the system is stabilized. This indicates that H_2O_2 is getting competitively adsorbed on the surface of MnO_2 and replacing some of the adsorbed IC molecules. Once a steady adsorption-desorption equilibrium for both IC and H_2O_2 is reached further increase in H_2O_2 does not result in enhanced adsorption of H_2O_2 or desorption of IC.



Figure 8A. Effect of added H_2O_2 on the degradation of IC (Inset: Effect of time)

In the combined presence of MnO₂ and UV irradiation, H_2O_2 inhibits the degradation initially. At the H_2O_2 concentration of ~20 mg L-1, the degradation starts increasing moderately and stabilizes at ~30 mg L⁻¹. The stabilized level of degradation is almost the same as in the absence MnO_2 . This shows that in the presence of H_2O_2 , MnO₂ is not an effective photocatalyst for the degradation of IC. Hence MnO₂ may be simultaneously generating OH radicals as well as accelerating their deactivation by facilitating the interaction with H₂O₂ at the surface. The stabilization of degradation at higher concentration of H₂O₂ shows that the deactivation as well as the formation of OH and another ROS is more or less balanced. The stabilization is further evident from Figure 8A (inset) which shows that at later stages of the reaction when the concentration of IC becomes relatively less, the degradation remains more or less the same irrespective of the concentration of H_2O_2 .

At higher concentrations of added H_2O_2 some of the reactive OH radicals get quenched as follows:

$$H_2O_2 + OH \rightarrow HO_2 + H_2O$$
 (7)

Consequently, the number of \cdot OH radicals interacting with the substrate is less resulting in decreased degradation of IC. The quenching of the \cdot OH radicals (oxidation potential: 2.80 eV) results in the formation of less efficient oxidant HO₂· (1.70 eV). The decomposition of H₂O₂ into H₂O and O₂ and the recombination of HO₂· (reactions 8 and 9) are other possible reasons for the inhibition.

$$\begin{array}{ccc} UV \\ 2H_2O_2 & \rightarrow & 2H_2O+O_2 \end{array} \tag{8}$$

$$HO_2 + HO_2 \to H_2O_2 + O_2$$
(9)

Reaction 8 also results in a decrease in the UV radiation available for the catalyst which leads to decrease in the degradation of IC. Eventually, the rate of simultaneous formation and quenching of \cdot OH get balanced, at least in the concentration range studied here, resulting in more or less steady degradation, i.e. no significant effect by added H₂O₂.

The initial inhibiting effect of H_2O_2 is further confirmed by addition of H_2O_2 in-between to the system in which the degradation of IC is in progress. Results are shown in Figure 8B. Every extra addition of H_2O_2 (20 min and 40 min) decreases the % degradation of IC thereby reconfirming that in the presence of MnO_2 as a catalyst, H_2O_2 functions more as an inhibitor.



Figure 8B. Effect of in-between addition of H_2O_2 on the degradation of IC.

The effect of H_2O_2 on the adsorption of IC over MnO₂ as a possible cause of inhibition/stabilization of the degradation is tested by measuring the adsorption of the dye at various concentrations in the presence of H_2O_2 . The results presented in Table 1A show that irrespective of the concentration of IC, H_2O_2 inhibits the adsorption. Similarly, experiments on the effect of increasing H_2O_2 concentration on the adsorption of IC (Table 1B), show that the adsorption decreases initially and stabilizes after that.

Table 1A. Effect of H_2O_2 on the adsorption of IC at various concentrations on MnO₂. [MnO₂]:1400 mg L⁻¹, [H₂O₂]:40 mg L⁻¹, time:30 min

[IC] mg L ⁻¹	Adsorption of IC, %	
	with H ₂ O ₂	without H ₂ O ₂
20	23.5	40.7
30	24.4	33.5
40	22.0	33.0
50	19.5	28.9
60	17.0	27.5

Table 1B. Effect varying concentration of H_2O_2 on the adsorption of IC on MnO₂. [MnO₂]:1400 mg L⁻¹, [IC]: 40 mg L⁻¹, time: 30 min

[H ₂ O ₂] mg L ⁻¹	Adsorption of IC, %
0	33.0
10	25.5
20	24.0
30	23.5
40	22.0
50	24.2
60	24.5

Hence it is possible that in the presence of added and/or in situ formed H_2O_2 , at least some of the surface sites on MnO_2 are utilized for their adsorption and/or decomposition, and this may be one of the reasons for the initial inhibition by H_2O_2 . However, there is a limit to the number of sites that can be occupied by H_2O_2 at any point in time for a given amount of MnO_2 . This may be a reason for the stabilization of the adsorption as well as degradation of IC beyond a particular concentration of H_2O_2 .

The competition between H_2O_2 and IC for adsorption on the surface is further confirmed by using IC pre-adsorbed MnO₂ for H_2O_2 adsorption/ decomposition (IC: 60 mg L⁻¹, H_2O_2 : 50 mg L⁻¹, MnO₂: 1800 mg L⁻¹). In this case, the decrease in the concentration of H_2O_2 is only ~60 % against ~ 90 % on fresh MnO₂.

Effect of persulphate (PS)

PS (S₂O₈²⁻, $E^0 = 2.1$ V) is a powerful oxidant capable of mineralizing some recalcitrant pollutants under AOP conditions and is investigated as a potential enhancer of the photocatalytic degradation of IC. It has specific advantages such as high solubility and stability under ambient conditions. Further, the SO_4^{2-} ions, which are the major products of PS reduction are relatively harmless and considered to be environment-friendly. Preliminary experiments showed that PS as such does not cause any degradation of IC. However, irradiation by UV causes moderate degradation which increases with increase in the concentration of PS (Figure 9). The decolorization is ~35 % in the presence of MnO₂ in the absence of UV irradiation, which is due to simple adsorption and is practically unaffected by the presence of PS. This also shows that unlike in the case of H₂O₂, the adsorption of PS on MnO₂ is weak.



Figure 9. Effect of added $K_2S_2O_8$ on the degradation of IC (Inset: Effect of time).

PS enhances the degradation of IC moderately under MnO_2/UV in the beginning and stabilizes with an increase in the concentration of PS. The enhancement is almost ~20% with respect to the system with no added PS. The addition of PS in between to a reaction ($MnO_2/IC/UV$) in progress also enhances the rate of degradation of IC. Since there is no

negative effect in the presence of an excess of PS, any unused oxidant remaining in the system can be used for the treatment of fresh input of IC pollutant which is important from the angle of commercial application of the process.

The MnO_2/UV degradation of IC in the presence of the combination of H_2O_2 and PS showed that the degradation is only the average of the negative effect of H_2O_2 and the positive effect of PS. Hence this option is not pursued in detail.

PS is known to generate highly reactive SO_4 - radical anions in situ under photocatalytic conditions. These can interact with the substrate and initiate/enhance the degradation of organics under AOPs. More SO_4 - radicals will be formed with an increase in the concentration of PS, under UV irradiation and a series of chain reactions can follow as in equations 10-20.⁴⁷

$$S_2O_8^2 \rightarrow 2SO_4$$
 (10)

$$S_2O_8^{2-} + RH \rightarrow SO_4^{-} + HSO_4^{-} + R$$
 (11)

RH: organic pollutant

$$SO_4 + H_2O \rightarrow SO_4 + H^+ + OH$$
 (12)

$$SO_4 + RH \rightarrow R + HSO_4$$
 (13)

$$OH + RH \quad \rightarrow \quad R^{\cdot} + H_2O \tag{14}$$

$$R + OH \rightarrow$$
 Intermediates \rightarrow products (15)

$$SO_4 \cdot + R \cdot \rightarrow Chain termination$$
 (16)

$$SO_4 \rightarrow OH \rightarrow Chain termination$$
 (17)

 $2SO_4 \rightarrow S_2O_8^{2-}$ Chain termination (18)

$$2 \cdot OH \rightarrow H_2O_2$$
 Chain termination (19)

$$2R^{-} \rightarrow RR$$
 Chain termination/ (20)

Various ROS such as H_2O_2 , HO_2 , OH etc. and SO_4 . formed during the irradiation interact with IC on the surface of the catalyst as well as in bulk, leading to its degradation into various intermediates and eventual mineralization. However, the degradation is not increasing with increase in the concentration of PS, after the initial enhancement. The excess of SO_4 . radicals formed in situ at higher concentration of PS may be getting deactivated as in reactions (16) and (18). The reactive SO_4 . radicals can also interact with PS and form less reactive S_2O_8 . radicals as follows:

$$SO_4^{-} + S_2O_8^{2-} \rightarrow SO_4^{2-} + S_2O_8^{-}$$
 (21)

Thus the SO_4 - radicals which are responsible for the enhancement of the degradation of IC in the presence of PS can get deactivated by unproductive interactions. This leads to decrease/ stabilization in the rate of degradation at higher concentration of PS and with reaction time.

Effect of dissolved salts/anions

Many of the anionic contaminants which are naturally present in water inhibit the efficiency of degradation of organic pollutants under AOPs.^{36,48,49} Instances of anions functioning as enhancers of the degradation of pollutants also have been reported.⁵⁰ In this context, the effect of some of the commonly occurring anions in water, i.e. SO₄²⁻, Cl⁻, PO₄³⁻, CO₃²⁻, HCO₃⁻ and NO₃⁻ on the efficiency of UV/MnO₂ degradation of IC is tested at various concentrations and reaction times (Figure 10A).



Figure 10A. Effect of concentration of anions on the photocatalytic degradation of IC.

The results showed that the anions SO_4^{2-} , Cl^- and PO_4^{3-} at lower concentrations have only negligible effect on the degradation of IC while CH_3COO^- , CO_3^{2-} and HCO_3^- inhibit the degradation at all concentrations. SO_4^{2-} and Cl^- do not influence the degradation significantly even at higher concentrations while PO_4^{3-} becomes a good inhibitor at higher concentrations. NO_3^- enhances the degradation at all concentrations.

At lower concentrations of 5 mg L^{-1} , the anion effect is less pronounced and the inhibition is in the order:

 CO_3^{2-} (inhibition) > HCO_3^{-} (inhibition) > CH_3COO^{-} (inhibition) > PO_4^{3-} (mild inhibition to no effect) > SO_4^{2-} (no effect) ~ CI^{-} (no effect) > NO_3^{-} (enhancement)

(22)

At higher concentration of the anion $(15 \text{ mg } \text{L}^{-1})$ where the effect is significant, the inhibition effect is in the sequence;

 HCO_3^- (inhibition) $\ge PO_4^{3-}$ (inhibition) $> CO_3^{2-}$ (inhibition) $> CH_3COO^-$ (inhibition) $> SO_4^{2-}$ (no effect) $\sim Cl^-$ (no effect) $> NO_3^-$ (enhancement)

The inhibiting anions become better inhibitors, and the enhancing anion becomes better enhancer with an increase in concentration. However, the comparative effect of the anions remains more or less the same irrespective of the concentration. Another parameter that can influence the effect of anions is the duration of the reaction possibly due to the increasing presence of various in-situ formed intermediates and the decreasing concentration of the substrate. The relative concentration of the ion with respect to the substrate increases progressively with time leading to more inhibition. The effect is tested at 3 different time intervals, and the results are plotted in Figure 10B.



Figure 10B. Effect of reaction time on the photocatalytic degradation of IC in the presence of various anions.

The relative effect of the anions at different durations of reaction is summarized in (24) - (26).

10 min.:

 HCO_3^- (inhibition) $\ge PO_4^{3-}$ (inhibition) $> CO_3^{2-}$ (inhibition) $> CH_3COO^-$ (inhibition) $> SO_4^{2-}$ (no effect) $\sim Cl^-$ (no effect) $> NO_3^-$ (enhancement)

20 min.:

$$\begin{split} &HCO_3^{-1} \text{ (inhibition)} \geq PO_4^{3-1} \text{ (inhibition)} > CO_3^{2-1} \text{ (inhibition)} \\ &> CH_3COO^{-1} \text{ (inhibition)} > Cl^{-1} \text{ (slight inhibition to no} \\ &\text{effect)} \sim SO_4^{2-1} \text{ (no effect)} > NO_3^{-1} \text{ (enhancement)} \end{split}$$

(25)

30 min.:

$$\begin{array}{l} PO_4{}^{3^{-}}(\text{inhibition}) > CO_3{}^{2^{-}}(\text{inhibition}) \geq HCO_3{}^{-}\\ (\text{inhibition}) > CH_3COO^{-} (\text{inhibition}) > SO_4{}^{2^{-}}(\text{no effect}) \\ Cl^{-}(\text{no effect}) > NO_3{}^{-}(\text{enhancement}) \end{array}$$

The trend remains more or less the same irrespective of the duration of the reaction. The effect ranging from 'inhibition' to 'no effect' and 'enhancement' and the variation depending on the concentration indicate that many reaction parameters such as the characteristics of the anions, the chemistry of the substrate and the catalyst, relative concentration of different components, the nature of their interactions etc. influence the anion effect. Hence any general conclusions on the effect of anions may not be consistent.

The inhibition of the photocatalytic degradation of organics by anions/salts is often explained based on blocking of the active surface sites on the catalyst by competitive adsorption, surface layer formation and/or scavenging of reactive 'OH radicals. In this context, the adsorption of IC on MnO_2 in the presence of the anions is measured experimentally. The results are presented in Table 2.

The adsorption is the least in the presence of $PO_4^{3^-}$ followed by $CO_3^{2^-}$ and HCO_3^- while CI^- , CH_3COO^- (Ac⁻), $SO_4^{2^-}$ and NO_3^- do not influence the adsorption. The cation was kept the same in all cases, i.e., Na^+ . Had competitive adsorption of the anions been the main cause, the inhibition should have been in the order:

$$PO_4^{3-} > CO_3^{2-} \ge HCO_3^{-} > Ac^- \sim SO_4^{2-} \ge Cl^- \sim NO_3^{-}$$
(27)

Table 2. Adsorption of IC on MnO_2 in the presence of anions. [IC] : 40 mg L⁻¹, MnO_2 : 0.14 g in 100 mL, [Anion] : 10 mg L⁻¹

Anion	Adsorption of IC, %
None	32.2
PO4 ³⁻	14.3
HCO3 ⁻	21.3
CO ₃ ²⁻	19.4
Cl-	33.5
SO4 ²⁻	30.5
Ac ⁻	29.5
NO ₃ -	34.7

This sequence is consistent with the experimentally observed inhibition thereby suggesting that preferential adsorption by the anion is a major cause of inhibition. The anions are also known to scavenge the ROS, especially OH, which also could cause the inhibition of the degradation of IC. Scavenging of the OH by anions would yield respective radical anion species as follows:

$OH + HCO_3 \rightarrow$	$H_2O + CO_3$	(28)

$$\cdot OH + CO_3^{2-} \rightarrow OH^- + CO_3^{--}$$
(29)

$$\cdot OH + SO_4^{2-} \rightarrow OH^- + SO_4^{--}$$
 (30)

$$\cdot OH + NO_3^- \rightarrow OH^- + NO_3^-$$
(31)

$$OH + Cl^{-} \rightarrow OH^{-} + Cl^{-}$$
 (32)

 $Cl + Cl \rightarrow Cl_2 \rightarrow Cl_2 \rightarrow (33)$

The radical anions are also reactive towards organic compounds, though less efficiently than 'OH.⁴⁹ Moderate degradation in the presence of the anions can also be due to the interaction of the radical anions with the substrate on the surface as well as in bulk.

The presence of anions also reduces the diffusion coefficient of the organic pollutant in the reaction medium. Consequently, the pollutant molecules will be less accessible to the reactive \cdot OH and this also can contribute to the inhibition. Another possible reason for the inhibition by anions is the formation of an inorganic layer on the catalyst surface. The efficiency of layer formation depends on the solubility of the salts.⁵¹ The higher the solubility of the salt the lower will be the layer formation. The solubility of the salts (in mg/g of water at 20^oC) tested here is in the order;

 $\begin{array}{l} NaNO_3 \left(\ 94.9 \right) > CH_3 COONa \ (54.6) > Na_2 SO_4 \ (40.8) \geq \\ Na_2 CO_3 \ (39.7) > NaCl \ (36.1) > Na_3 PO_4 \ (16.3) > NaHCO_3 \\ (11.1) \end{array}$

(34)

Correspondingly the layer formation, if any, will be in the order:

$$HCO_3^{-} > PO_4^{3-} > Cl^{-} \ge CO_3^{2-} \ge SO_4^{2-} > Ac^{-} > NO_3^{--}$$

(35)

If layer formation is the major cause, the inhibition must be in the same order as in (35). The experimentally observed anion effect in the current instance is not fully consistent with equation (35). However, the trend is similar especially in the case of the maximum ($\text{HCO}_3^- > \text{PO}_4^{3^-}$) and the minimum (NO_3^-) inhibition. Hence, the layer formation also may be contributing partially to the anion effect. However, the relative contribution of various factors towards the anion effect may vary.

The inhibitive or 'nil' effect of various anions on the photocatalytic degradation of IC in the presence of MnO_2 can be explained to a great extent by any or all of the above interpretations. However, a universally consistent explanation to account for the effect of anions in different kinds of photocatalytic systems with different catalysts, substrates and reaction conditions is not possible, and it will be more appropriate to consider individual anions to explain specific effects. In this context, two specific examples, i.e. SO_4^{2-} for 'no effect' and NO_3^- for 'enhancement' are discussed here.

The primary step in the photocatalytic process is:

Catalyst + $h\nu \rightarrow$ Catalyst (e⁻, h⁺) (36)

(Formation of electrons and holes)

Catalyst (e^{-}, h^{+}) \rightarrow Catalyst (37)

(Recombination of electrons and holes)

In the case of SO_4^{2-} ions; they get adsorbed on the surface of the catalyst and interact with the photoproduced holes to generate sulfate radical anions as follows⁵²:

$$h^+ + SO_4^{2-} \rightarrow SO_4^{-}$$
 (38)

S is a strong oxidizing agent and hence the SO_4 can accelerate the degradation process according to reactions (39) and (40).

$$SO_4 \cdot + Pollutant \rightarrow SO_4^2 + Intermediate$$
 (39)

$$SO_4$$
 + Intermediate $\rightarrow SO_4^2$ + CO_2 + salts (40)

As seen in reaction (12), the presence of SO_4 can lead to the formation of more reactive OH radicals which can accelerate the degradation of the dye. Hence, unlike in the case of other anions which scavenge the OH, the SO_4^{2-} anions concurrently generate them. Thus, factors leading to inhibition and enhancement of the degradation coexist in the system. Under the reaction conditions used here, they may balance, and hence the degradation is practically unaffected. It is also possible that depending on the domination of either of the effects, the net effect may be 'inhibition,' 'no effect' or 'enhancement' at any point in time during the reaction.

The enhancement in the degradation of IC in the presence of NO_3^- shows that the factors responsible for the inhibition by other anions are not significant here. Hence both the surface and bulk processes are not affected. Further, under irradiation by UV light, NO_3^- leads to the formation of reactive OH radicals as follows⁵³:

 $NO_3^- + hv \rightarrow NO_2^- + O \tag{41}$

$$NO_3^- + H_2O + h\nu \rightarrow NO_2^- + OH^- + OH$$
(42)

$$O + H_2 O \to 2 \cdot O H \tag{43}$$

The enhancement by NO₃⁻ can be thus attributed to the direct and indirect formation of OH radicals. Zepp et al.⁵³ suggested that nitrate-induced photooxidation is a possible transformation mechanism for trace organic pollutants in shallow clear water bodies with inputs from ground waters or other sources with higher ratios of nitrate/dissolved organic carbon concentration. Zhu et al. also reported similar findings.⁵⁴

Chemical oxygen demand

Estimation of chemical oxygen demand (COD) of the optimized reaction system UV/MnO₂ /IC at various intervals of irradiation shows that the degradation and decolorization of IC do not lead to instant mineralization.

However, once the solution is decolorized, and the UV/MnO_2 irradiation is continued, mineralization becomes faster, and the COD reaches a stable level (Figure 11). After that, the COD decreases very slowly even on continuous UV radiation indicating that the intermediates are stable and cannot be mineralized under these conditions.



Figure 11. COD of the reaction system at different times of irradiation after decolorization of IC.

The photodegradation of IC is known to yield stable organics such as acetic acid, tartaric acid, malic acid, oxalic acid, anthranilic acid etc.^{22,25} Independent experiments in our laboratory with these compounds showed that they are not mineralized under MnO_2 mediated photocatalysis (not reported here). In this context, the possibility of combining a more efficient photocatalyst such as TiO_2 with MnO_2 in order to enhance the mineralization capability is examined.

Investigations using different weight ratios of MnO_2 and TiO_2 have shown that MnO_2/TiO_2 in the ratio 9:1 is optimum for the decolorization of IC under the standardized reaction conditions. The comparative decolorization and mineralization efficiency of MnO_2 , TiO_2 , and MnO_2/TiO_2 for IC are evaluated. The rate of decolorization and mineralization is more in the presence of MnO_2/TiO_2 compared to individual MnO_2 or TiO_2 . This is evident from the COD values in the presence of the three catalysts (Figure 11).

The complete and faster mineralization in the presence of MnO_2/TiO_2 shows clearly that the advantages of higher adsorption and oxidative properties of MnO_2 and the photocatalytic efficiency of TiO_2 can be beneficially combined for the mineralization of traces of IC in water. However, the precise mechanism of the process, as well as the nature of intermediates, may be different in the case of different catalysts, and these aspects need to be investigated in detail.

General mechanism

The primary step in semiconductor mediated heterogeneous photocatalytic processes is the formation of an electron-hole pair as in reaction (36) which may undergo concurrent recombination as in reaction (37). When the particles are small, the holes escape recombination in the bulk of the catalyst and can reach the surface. The average transit time τ_D of the charge carriers to reach the surface is calculated by Gratzel and Frank⁵⁵ as:

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$$\tau_{\rm D} = \frac{r^2}{\pi^2 D_{\rm c}} \tag{44}$$

where

r is the radius of the particle and

 $D_{\rm c}$ is the diffusion coefficient of the charge carrier.

If the photogenerated hole does not recombine with the electron, it can react with the organic pollutant (RH) or even with the solvent water as follows:

$$RH + h^+ \rightarrow R + H^+ aq$$
 (45)

$$H_2O + h^+ \rightarrow OH + H^+ aq$$
 (46)

The hole can also react with the OH⁻ ions either present in the system under alkaline conditions or formed on the surface by the dissociative adsorption of water to H^+_{ads} . and OH⁻ ads.⁵⁶. The OH⁻ads can act as traps for the holes at the surface;

$$h^+ + OH_{ad} \rightarrow OH_{ad}$$
 (47)

$$\cdot OH + \cdot OH \quad \rightarrow \quad H_2O_2 \tag{48}$$

 H_2O_2 formed in the photocatalytic system has been proven to undergo concurrent decomposition leading to the phenomenon of oscillation in its concentration.⁵⁷ The reductive decomposition of H_2O_2 by the photogenerated electron as in (49) results in the formation of reactive \cdot OH radicals. This also protects the holes from recombination with the electrons.

$$H_2O_2 + e^- \rightarrow OH^- + OH^- \tag{49}$$

According to Gerischer and Heller⁵⁸ reaction (47) is more likely than reaction (46) since the availability of holes on the catalyst surface is more under photocatalysis as explained above. Further, the \cdot OH_{ad} can exist as the stable product at specific surface sites. Consequently, the oxidation of organic pollutants can occur as:

$$RH + .OH_{ad} \rightarrow R' + H_2O$$
 (50)

The direct reaction between the organic pollutant and the hole as in reaction (45) can compete with reaction (50) if the concentration of RH is high or the RH molecules are strongly adsorbed on the surface.

If reaction (50) is the main path, the oxidation may be weaker compared to direct oxidation by holes as in reaction (45) since only some of the holes are available for the formation of $\cdot OH_{ads}$.⁵⁷ This will lead to decrease in the oxidation rate and the quantum efficiency in the case of stable organic molecules.

When there are more H_2O_2 in the system, it can also trap the h⁺ as in reaction (51) and quench $\cdot OH$ as in reaction (7) which also lead to decreased rate of degradation.

$$H_2O_2 + 2 h^+ \rightarrow O_2 + 2H^+$$
(51)

Oxygen plays an important role in photocatalytic degradation reactions in aqueous solutions by scavenging the electrons generated on the photocatalyst thereby inhibiting the recombination of electrons and holes. The formation of highly reactive superoxide radical anion which is responsible for the photocatalytic degradation is the outcome of this reaction.

$$O_2(g) + e^- \to (O_2^-)$$
(52)

 O_2 . participates in a series of reactions directly or indirectly as in (53) – (60) resulting in the formation of a number of reactive free radicals which interact with and degrade the pollutant.

 $H_2O \rightarrow OH^- + H^+ \tag{53}$

 $O_2^{-} + H^+ \rightarrow HO_2^{-} \tag{54}$

 $2HO_2 \rightarrow H_2O_2 + O_2 \tag{55}$

 $\mathrm{HO}_{2^{\bullet}} + \mathrm{e}^{\bullet} \to \mathrm{HO}_{2}^{\bullet}$ (56)

$$\mathrm{HO}_{2}^{-} + \mathrm{H}^{+} \to \mathrm{H}_{2}\mathrm{O}_{2} \tag{57}$$

 $H_2O_2 + O_2^{-} \rightarrow OH^{-} + OH^{-} + O_2$ (58)

 $H_2O_2 + e \rightarrow OH + OH \tag{59}$

$$OH^{-} + h^{+} \rightarrow OH^{-} \tag{60}$$

In order to confirm the role of dissolved O_2 in the photocatalytic degradation IC, the reaction system is deaerated with N_2 , and the experiments were carried out under otherwise identical conditions. The effect of deaeration is not significant thereby suggesting that the main source of O_2 for the reaction is not the dissolved form. The required O_2 is provided primarily by the surface (adsorbed) and the lattice of MnO_2 which cannot be easily removed by flushing with N_2 .

Amorphous MnO_2 is known to release bulk oxygen more easily to the surface which makes it a better catalyst regarding facile activation and regeneration.³² MnO_2 with multiple oxidation states together with its electron donoracceptor properties is an excellent oxidation – reduction catalyst. Photolysis of MnO_2 increases the oxygen species on the surface either by oxygen migration to the surface or by Mn migration to the bulk or both. Surface oxygen is consumed faster upon irradiation and oxygen from the bulk moves to the surface. In the case of MnO_2 , the loss of oxygen takes place at temperatures as low as 50 °C. The photo-initiated oxygen release from MnO_2 may be due to movement of O^{2-} (bulk) to the surface and subsequent weakening of MnO_2 bonds.⁵⁹ In the re-oxidation of oxides, atmospheric oxygen and/or dissolved O_2 is taken up by the surface of the partially reduced MnO_2 as follows³²:

$$O_2(g) + e^- \to (O_2^-) \tag{61}$$

$$(O_2^{-}) + e^{-} \rightarrow 2(O^{-}) \rightarrow O^{2-}(s)$$
(62)

Most of the oxygen radicals are present in the bulk of the catalyst while some of them may remain on the surface for a short period. These highly active oxygen radicals can regenerate reduced manganese species. Under UV irradiation, the bonds in MnO_2 are weakened and O^{2-} is released to the surface. Even though the lifetime of excited state oxygen species is short, they possess adequately high energy and electronegativity to facilitate the reduction or hydrogen abstraction from the substrate. Possible reaction pathways are:

$$O^{2-}(s) \xrightarrow{hv} (O^{-}) \xrightarrow{hv} O_{2}(surface)$$
(63)

 O_2 (surface) + RH (i.e., IC) \rightarrow HO₂·(surface) + R· (64)

 O_2 (surface) + R \rightarrow Intermediates \rightarrow

 \rightarrow Mineralization (CO₂ + H₂O) (65)

The abstracted H interacts with the hole and gets oxidized to form acid sites on the surface of MnO_2 .

$$\mathbf{H}^{\cdot} + \mathbf{h}^{+} \rightarrow \mathbf{H}^{+} \tag{66}$$

The regeneration of MnO₂ may be represented as:

$$-Mn^{(3+ \text{ or } 2+)} \xrightarrow{h\nu} -Mn^{(4+ \text{ or } 3+)}$$

$$-e^{-}$$
(67)

 $n(-Mn^{(4+ \text{ or } 3+)}-)+O^{2-}(\text{from reaction } 62) \rightarrow MnO_2 \text{ (or } Mn_3O_4)$

(68)

Unless the oxygen is replenished periodically, the activity of the catalyst will be lost faster. This is verified by recycling the catalyst immediately after use by simple filtration followed by quick drying at 120 °C for 1 h. The degradation of IC (in 30 minutes) decreased steeply from ~75 % in the presence of the fresh catalyst to ~30% in first recycling, ~11 % in second recycling and ~8% in third recycling. This confirms the drastic change in the surface characteristics, loss of adsorption sites and loss of oxygen from the lattice, bulk and/or the surface of MnO₂. This loss is not fully compensated by contact with atmospheric oxygen. The migration of bulk oxygen to the less energetic surface sites and its consumption by participation in catalytic reactions has been proven experimentally and from ESR studies.⁶⁰

The formation of H_2O_2 during the photocatalytic degradation of IC is verified by analyzing the reaction solution immediately after decolorization and periodically after that. The concentration of in situ formed H_2O_2 increases initially, reaches an optimum and decreases after that. This is consistent with the earlier reports⁵⁷ on the oscillation in the concentration of in situ formed H_2O_2 in semiconductor-mediated photocatalytic systems which are attributed to concurrent formation and decomposition [reactions (53)-(60)].

Dyes are known to absorb light efficiently in the visible region. Hence IC can absorb more visible light and act as a sensitizer thereby transferring electrons from the excited dye molecule to the conduction band of MnO_2 .

$$Dye_{(ads)} \rightarrow Dye_{(ads)}^{*}$$
 (69)

 $\text{Dye}_{(ads)}^{*} + \text{MnO}_2 \rightarrow \text{Dye}_{(ads)}^{+} + \text{MnO}_2(e_{CB})$ (70)

 $\text{Dye}_{(ads)}^+ \rightarrow \text{Intermediates} \rightarrow \text{Oxidation}/\text{Degradation}$

(71)

The similar pathway was suggested for the degradation of dyes on Ag/TiO₂ catalyst also.⁶¹ However, dye sensitization as a mode of catalyst activation is highly concentration-dependent. At lower concentration of the dye, there will be only monolayer coverage and the energy acquired by light absorption may not be adequate to sensitize the semiconductor. At higher concentration, there will be multilayer adsorption of the dye. Consequently, light absorption and photocatalytic efficiency may not increase significantly because the inner layers will tend to act as insulators with respect to outer layers.

Conclusion

Trace amounts of the toxic dye pollutant Indigo Carmine in water are degraded efficiently by MnO₂ mediated photocatalysis. However, the degradation does not lead to complete mineralization which requires more efficient photocatalyst such as TiO₂ or the combination MnO₂/TiO₂. The degradation does not follow the modified L-H mechanism or pseudo-first order kinetics as in the case of many AOPs probably because of the high adsorption and oxidation capability of MnO₂ and consequent faster degradation. The degradation is faster under acidic conditions. H₂O₂ which is a powerful oxidant for many photocatalytic reactions inhibits the degradation which is attributed to the quenching of the photoproduced holes and reactive 'OH radicals. However, another oxidant persulphate enhances the degradation through the in situ generation of reactive SO4- radical anion and OH. Salts/anions likely to be present in water influence the degradation differently varying from 'moderate inhibition' to 'no effect' or even 'enhancement' depending on the chemistry of the anion, its concentration, interactions with the substrate, catalyst, intermediates, etc. and other reaction conditions. The surface, bulk as well as lattice oxygen of MnO_2 participates in the photocatalytic degradation of IC, and this results in decreased activity with every recycling.

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