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Luminescent group 113, 5-dimethyl pyrazolate complexes/titanium oxide composites for photocatalytic removal and degradation of 2, 4-dichlorophenoxyacetic acid

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Keywords: degradation, 2, 4-dichlorophenoxyacetic acid, group 11 metal complexes, luminescence, titanium oxide composite

Abstract

PAPER

We report the successful paration of new luminescent group 11 pyrazolate complex/titanium exide composites for the significant improvement of photocatalytic activity in the removal and degradation of 2, 4-dichlorophenoxyacetic acid (2, 4-D). Photocatalyst composites were successfully prepared using an impregnation method of all complexes (M(I or II)) in weight ratios of 0.1-1.0 wt% to T_{SR} . In contrast to anatase TiO₂ with 49% for removal and 10% for degradation, the composites $Au(\overline{I})/TiO_2$ and $Ag(I)/TiO_2$ with 0.4 wt% gave the highest removal and degradation in 85% and 36% in 1 h. Interestingly, the composite Cu(I)/TiO₂ with 0.4 wt% gave even better removal in 99% and almost the same percentage of degradation. By using the same ligand for the complexation, the photocatalyst composite Cu(II)/TiO₂ with the same composition was found to show enhancement in the degradation from 36% to 48% with similar percentage of the removal. These showed that group 11 pyrazolate complexes in TiO₂ composites could able to significantly improve the photocatalytic activity due to the reduction of electron-hole recombination of TiO2. Moreover, Cu(I) pyrazolate with longer metal-metal distance (emission peak at 590 nm) for electron transfer from the conduction band is responsible for such performance in the removal. Indeed, dinuclear Cu(II) pyrazolate complex with less rigid structure has more longer metal-metal distance (566 nm) to give such better performance in the degradation. This work demonstrated that modification of TiO₂ with group 11 pyrazolate complexes at the molecular level as photocatalyst composites is a new approach to enhance photocatalytic activity.

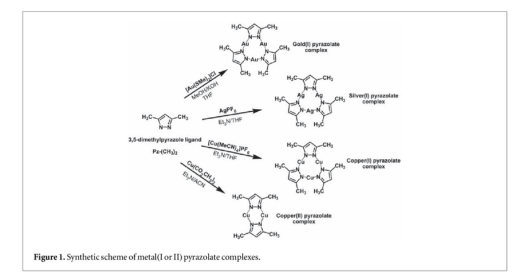
1. Introduction

Since last few decades, titanium oxide (TiO_2) has been widely used as a photocatalyst for the degradation of organic pollutants due to its high stability and non-toxicity as well as bandgap energy [1–3]. However, the activity cannot be significantly improved so far [3]. Thus, a lot of studies have been developed to modify TiO_2 as photocatalyst composites with high photocatalytic activity. These composites are generally prepared with the expectation that electron-hole recombination process can be reduced while energy band gap can be improved [4–7].

Recently, transition metal oxides have received special attention among the researchers in the development of photocatalyst composites with good catalytic activity [8–14]. These reports have not yet significantly improved the photocatalytic activity. We have recently reported a new approach to enhance the performance of

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TiO₂ by using columnar assembly of luminescent copper(1) pyrazolate complexes [15]. In this finding, Degussa P25 [27] composite with less rigid structure of 3, 5-dimethyl pyrazole ligand 26 pwed more active as a photocatalyst with improvement of photocatalytic activity (in 60%) for 44 degradation of 2, 4-dichlorophenoxyacetic acid (2, 4-D) as model pollutants compared to the bulk one of 4-(3, 5-dimethoxybenzyl)-3, 5-dimethyl pyrazole ligands (49%) [15]. Since other group 11 pyrazolate complexes also showed similar luminescent properties with copper complexes based on oxidation number of metal ions and distance of the weak metal-metal interactions [16–20], new luminescent composites with a cheaper precursor and stable structure will be highly required as a new photocatalyst. Therefore, we report the first example of photo 33 lyst composites of modified anatase TiO₂ from group 11 metal complexes for the photocatalytic removal and degradation of 2, 4-D. In this study, we parti 43 rly highlight the contribution of different metal complexes from group 11 with different molecular structures as well as more photocatalyst composites.

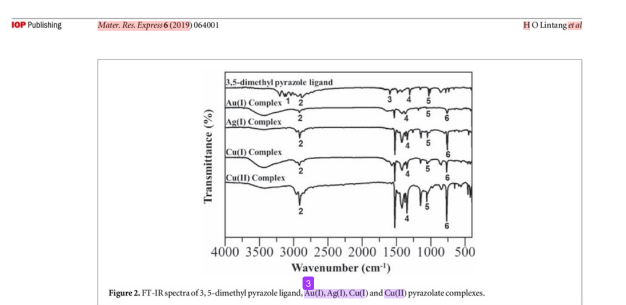
2. Materials and methods

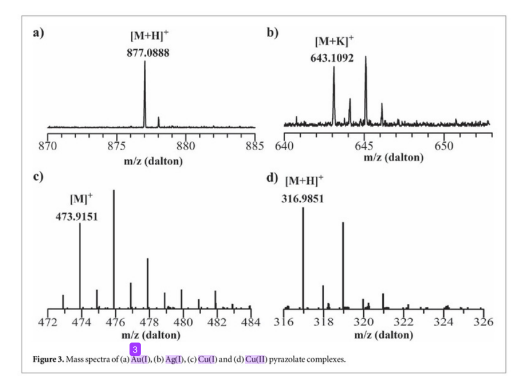
2.1. Materials

Commercial Hombikat UV100 TiO₂ (anatase TiO₂) from Sachtleben Chemie was used without further treatment. 3, 5-dimethylpyrazole ligand (Sigma-Aldrich, $C_5H_8N_2$, 99%) and metal salts of tetrakis(acetonitrile) copper(I) hexafluorophosphate (Sigma-Aldrich, $C_8H_{12}CuF_6N_4P$, 97%), copper(II) acetate (Sigma-Aldrich, $C_4H_6CuO_4$, 98%), chloro(dimethylsulfide) gold(I) (Sigma-Aldrich, C_2H_6SAuCl), and silver hexafluorophosphate (Sigma-Aldrich, AgPF₆, 98%) were used as precursors to synthesize group 11 metal (M(I or II); M = Au, Ag and Cu) pyrazolate complexes. Dry acetonitrile, methanol and tetrahydrofuran for the synthesis were directly used. Triethylamine was firstly distilled under vacuum and an inert condition where it was collected in the round bottom flask containing potassium hydroxide. 2, 4-D ($C_8H_6Cl_2O_3$, 98%) from Sigma-Aldrich was used as a model of organic pollutant.

2.2. Synthesis of luminescent group 11 metal pyrazolate complexes

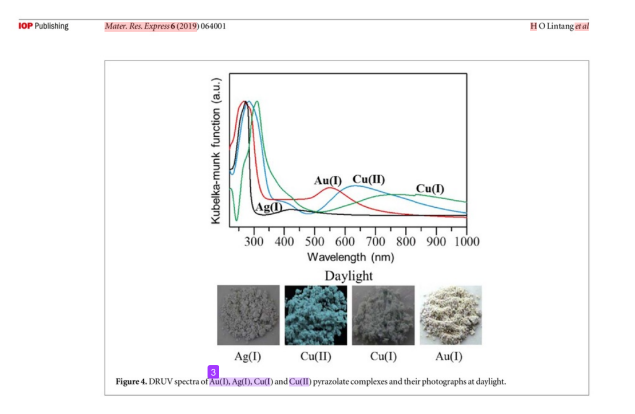
M(1) pyrazolate complexes were synthesized by in sing 3, 5-dimethylpyrazole ligand (Pz-(CH₃)₂) with metal salts in dry solvents using a Schlenk technique under an inert condition as shown in figure 1 [15–19]. Moreover, copper(II) pyrazolate complex was synthesized using copper (II) acetate as the source of metal ion with a certain modification in the synthesis procedure [20]. Typically, the mixture solution of Pz-(CH₃)₂ and metal salts was stirred for 5 mins and followed by addition of potassium hydroxide in methanol or fresh distilled triethylamine as a base. After overnight reaction, the solvent was completely removed from the residue using a vacuum pump in high pressure. The remaining residue was further purified using *in situ* recrystallization to obtain targeted complex by adding dichloromethane in small amount for dissolving the residue (desired product). The mixture solution was transferred into dry cold methanol (around 10 times of dichloromethane) to obtain a precipitate. The remaining solvent was removed by *in situ* filtration and further dried using vacuum pump pressure to give white powder of Au(I) pyrazolate complex in 82%, light grey powder of Ag(I) pyrazolate complex in 81%, light

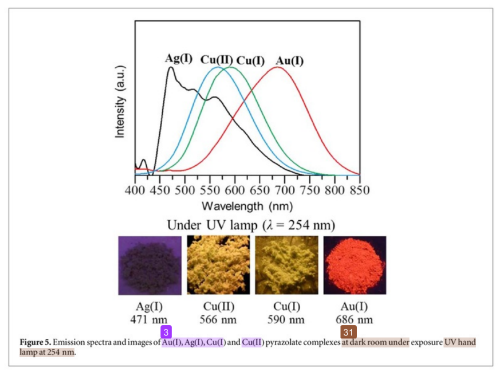




greenish powder of Cu(I) pyrazolate complex in 80%, and light blue-greenish powder of Cu(II) pyrazolate complex in 78%.

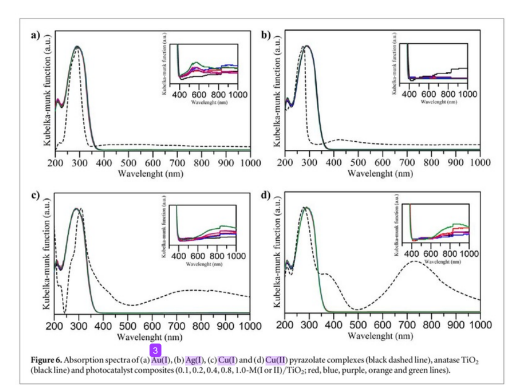
2.3. Synthesis of luminescent group 11 metal pyrazolate complexes/titanium oxide composites Group 11 metal pyrazolate complexes/titanium oxide composites (x-M(I or II)/TiO₂) were prepared as photocatalyst composites using an impregnation method where x represents the weight ratio of M(I or II) pyrazolate complexes to anatase TiO₂ from 0.1 to 1.0 wt%. In this step, TiO₂ (1.0 g) was dispersed to the dichloromethane solution (25 ml) of M(I or II) pyrazolate complexes. The m²⁵ re was then sonicated for 20 mins for better dispersed to the complex and TiO₂ [15, 17, 19]. Then, the mixture was stirred until dried at room temperature and calcined at 373 K for 4 hs with heating rate of 10 K min⁻¹.





2.4. Characterization of luminescent group 11 metal pyrazolate complexes and titanium oxide composites The M(I or II) or razolate complexes and x-M(I or II)/TiO₂ composites were characterized with various type of instruments. Fourier transform infrared spectroscopy (FT-IR) on a model of Nicolet iS50 Thermo Scientific was used to analyze M(I or II) pyrazolate complexes using potassium bromide pellet technique. Mass spectra of pyrazolate complexes were analyzed for exact mass analysis using an Agilent 6560 high performance liquid

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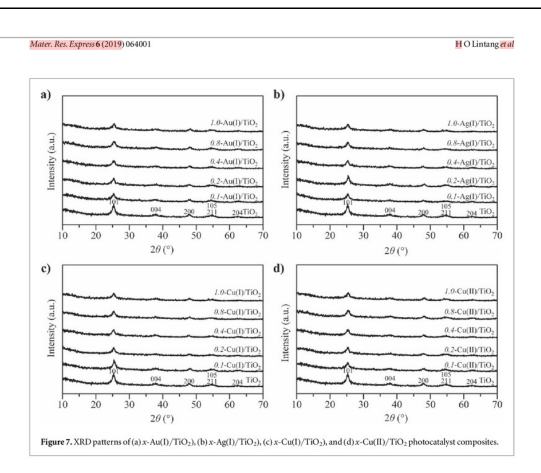


chromatography tandem mass spectrometry with quadrapole time-of-flight (HP LCMS-MS Q-TOF). The diffraction pattern and crystallite structure of the composites were chara 20 ized using Bruker D8 X-ray Diffractometer with a scan speed of 0.05 $^{\circ}s^{-1}$. The absorption properties were recorded on 3 R-UV spectrophotometer (Shimadzu UV-2600) where barium sulphate was used as the reference. JASCO FP-8500 spectroflourophotometer was used to analyze emission properties of the composites.

The morphology of the composites was studied using field emission scanning electron microscopy (FESEM, JEOL on a model of JSM-6701F) operated 135 kV on samples coated with Pt. FESEM was attached with a spectrometer for element measurement (energy 30 persive x-ray, EDX). Transmission electron microscopy (TEM, JEOL) was also performed on a model of JEM-2100 at an accelerating voltage of 200 I_{6}° . In the dark room, digital camera on a model of DMC–FZ38 (Panasonic) were used to take photographs under exposure to a hand-held UV lamp ($\lambda_{ex} = 254$ nm).

Photocurrent measurement was performed using an Interface 1000 potentiostat/galvanostat (Gamry Instruments Inc.) with chronoamperometry method. Screen-printed electrode (SPE) as a working electrode was prepared in saturated Na₂SO₄ (0.1 M, 6 ml) solution as an electrolyte. A Xe–Hg UV lamp (200 W, 8 mWcm⁻²) was used as the light source to irradiate the SPE substrate coated with sample. The SPE substrate containing of coated sample was prepared by depositing the mixture of homogeneous colloidal suspension (20 μ L) of Nafion (10 μ L) with photocatalyst composite (10 mg) in 1 ml of deionized water [12]. The droplet sample was naturally dried under ambient temperature until it was dried-off enough for that photocurrent measurement. The SPE substrate coated with the sample was then applied in the electrolyte system with light-on and off event in every 30 s.

2.5. Characterization of luminescent group 11 metal pyrazolate complexes and titanium oxide composites The prepared photocatal 12 uninescent composites of group 11 metal pyrazolate complexes/titanium oxide (x-M(I or II)/TiO₂) were tested for the remotal and degradation of 2, 4-D under a UV light irradiation [6, 13]. x-M(I or II)/TiO₂ in 50 mg was added to the 50 ml solution of 2, 4-D with concentra 14 of 0.5 mM. The mixture was stirred under dark condition for 1 h to reach the adsorption equilibrium. After that, the mixture was further stirred under UV irradiation (200 W Xe-Hg lamp equipped with IR cut off filter) with light intensity of 8 mWcm⁻² for 1 h. The mixture (3 ml) was collected and purified using nylon membrane filter (0.2 μ m) after adsorption proces 27 d lamp exposure. The filterate solution was further analyzed using Shimadzu prominence LC-20 (a high performance liquit 15 romatography; HPLC equipped with a UV detector and Hypersil GOLD PFP column). The concentration of 2, 4-D and it's intermediate (2, 4-dichlorophenol;



2, 4-DCP) were examined with an eluent mixture of acetonitrile/distilled water 60:40 (v/v) at 283 nm. The removal percentage and degradation percentage of 2, 4-D was derived from the following equations.

Removal of 2, 4-D =
$$\frac{[2,4-D]i - [2,4-D]f}{[2,4-D]i} \times 100$$

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Degradation of 2, 4-D = $\frac{[2,4-D]i - [2,4-D]f}{[2,4-D]i - [2,4-D]f - [2,4-DCP]} \times 100$

For the above equations, $[2, 4-D]_i$ is concentration of 2, 4-D before UV irradiation, $[2, 4-D]_f$ is concentration of 2, 4-D after UV irradiation, and [2, 4-DCP] is concentration of 2, 4-DCP.

o 3. Results and discussion

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3.1. Molecular structure and optical properties of synthesized group 11 metal pyrazolate complexes The molecular structure of synthesized all M(I or II) pyrazolate complexes were confirmed by using FT-IR spectroscopy. Figure 2 shows that vibration bands of pyrazole ligand for N–H group at 3163 (1), 3113 (1), 3058(1) and 1595 c 53¹ (3) were disappeared compared to that of all pyrazolate complexes. In addition, the new vibration band was observed at 767 cm⁻¹ (6), which is assigned to the formation of N–Cu–N [18]. These results indicated the complexation reaction of pyrazolate complexes with the base were completely occured.

QTOF LCMSMS is one of the HRMSs that can be used for the exact mass analysis of synthesized organic compounds. Figures 3(a) and (d) shows the observed molecular weight with molecular formula of $C_{15}H_{21}N_6Au_3$ and $C_{10}H_{14}N_4Cu_2$ at 877.0888 and 316.9851 Da for both Au(I) and Cu(II) complexes, which were similar to the isotopic patterns of the calculated molecular formula ($[M + H]^+$) at 877.0903 and 316.9889 Da for Au(I) and Cu(II) complexes. For Ag(I) pyrazolate complex with molecular formula of $C_{15}H_{21}N_6Ag_3$ ($[M + K]^+$) at 644.8613 Da, the particular peak of molecular weight was observed at 643.1092 Da (figure 3(b)). Moreover, the monoisotopic pattern for Cu(I) pyrazolate complex with molecular formula $C_{15}H_{21}N_6Ag_3$ showed a peak at 473.9151 Da (figure 3(c)) for observed molecular weight of $[M]^+$, which was closed to 473.9716 Da of the calculated one. Therefore, M(I or II) pyrazolate complexes were successfully synthesize givith high yields.

Figure 4 shows the absorption spectra with a peak tops of 267, 273, 309 and 285 nm for Au(I), Ag(I), Cu(I) and Cu(II) pyrazolate complexes, respectively. These peak tops can be assigned as an absorption edge of the

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Table 1. Crystallite size of x-Au(I)/TiO ₂ ,	
x-Ag(I)/TiO ₂ , x-Cu(I)/TiO ₂ , and	
x-Cu(II)/TiO2 photocatalyst composites.	

Composites	Crystallite size ^a (nm)
10	0.2
TiO ₂	9.3
0.1-Au(I)/TiO ₂	10.0
0.2-Au(I)/TiO ₂	10.2
0.4-Au(I)/TiO ₂	10.3
0.8-Au(I)/TiO ₂	10.2
1.0-Au(I)/TiO ₂	10.4
0.1-Ag(I)/TiO ₂	9.9
0.2-Ag(I)/TiO ₂	10.0
0.4-Ag(I)/TiO2	10.2
0.8-Ag(I)/TiO2	10.4
8-Ag(I)/TiO2	10.2
0.1-Cu(I)/TiO ₂	9.6
0.2-Cu(I)/TiO ₂	9.5
0.4-Cu(I)/TiO ₂	9.7
0.8-Cu(I)/TiO ₂	9.8
1.0-Cu(I)/TiO2	10.1
0.1-Cu(II)/TiO ₂	9.0
0.2-Cu(II)/52 ²	9.0
0.4-Cu(II)/TiO ₂	9.4
0.8-Cu(II)/TiO ₂	9.3
1.0-Cu(II)/TiO ₂	9.2

^a Calculated using Scherrer equation based on 101 lattice plane of anatase TiO₂.

metal complexes containing luminophores with π - π stacking. Moreover, the broad absorption bands at the visible region were usually appeared for the compounds with d-d transition. Such absorption spectra were also sup 18 ted by the photographs in daylight as shown in the boy 18 m part of figure 4.

By using a peak top from their absorption spectra for the excitation wavelength as shown in figure 4, the luminesent spectra were more for all complexes. Figure 5 shows the emission spectra with a peak centered at 471, 686, 590 and 566 nm for Au(1), Ag(1), Cu(1) and Cu(II) pyrazolate complexes, respectively, indicating phosphorescent compounds with the weak more properties [15–19]. To support these emission properties, the photographs were captured in the dark room at 254 nm using a UV 3 mp. As shown at the bottom part of figure 5, the images with dark purple, red, light orange greenish and orange for Au(1), Ag(1), Cu(1) and Cu(II) pyrazolate complexes.

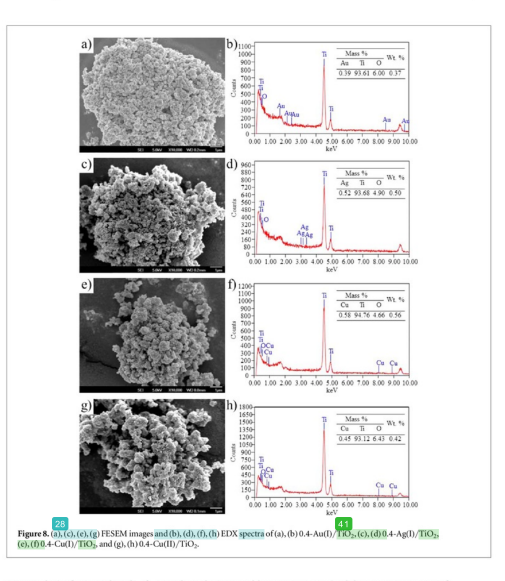
3.2. Luminescent properties, crystalline structure and morphology of photocatalyst composites

Figure 6 shows absorption spectra of all photocatalyst composites. Absorption band edge of the composites exhibited almost similar wit that of anatase TiO₂ (black line). These results demonstrated that pyrazolate complexes might be existed on the surface of the TiO₂. Moreover, absorption peaks of x-Au(1)/TiO₂, x-Cu(1)/TiO₂, and x-Cu(II)/TiO₂ at visible region from 400 to 1000 nm for d-d transition slightly increased with addition of pyrazolate complexes into TiO₂ (inset graph). In contrast, absorption peak at visible region of x-Cu(I)/TiO₂ decreased and remain unchanged upon addition of Ag(1) pyrazolate complexes into anatase TiO₂. These results demonstrated that all complexes are stable enough in the preparation of composites. Figure 7 displays XRD patterns of all photocatalyst composites. It was observed that all diffractograms showed diffraction peaks at 2θ of 25.35°, 38.10°, 48.05°, 54.55° and 62.60° (standard value, JCPDS file No.: 21-1271) which indicates the presence of an anatase phase for TiO₂. As shown in all samples, the intense diffraction peak at 2θ of 25.35° is generally assigned to 101 lattice planes. Estimated crystallite sizes were in the range of 9 to 10 nm where it was almost similar to that of the bulk TiO₂. Hence, these results suggested that addition of M(I or II) pyrazolate complexes have not affected the structural properties of anatase TiO₂ in their composites [12–15]

Figure 8 shows FESEM images of selected 0.4- $M(I \text{ or II})/TiO_2$ composites. The visualization showed that 0.4- $Au(I)/TiO_2$ composite displayed some particles with same shape and size (figure 8(a)), which is almost similar to the morphology of anatase TiO₂[12]. Moreover, all other 0.4- $M(I \text{ or II})/TiO_2$ composites with Ag(I), Cu(I) and Cu(II) showed some aggregation of particles with different shapes and sizes (figures 8(c), (e), (g)). Further investigation on the dispersion of M(I or II) pyrazolate complexes in the composites was carried out with

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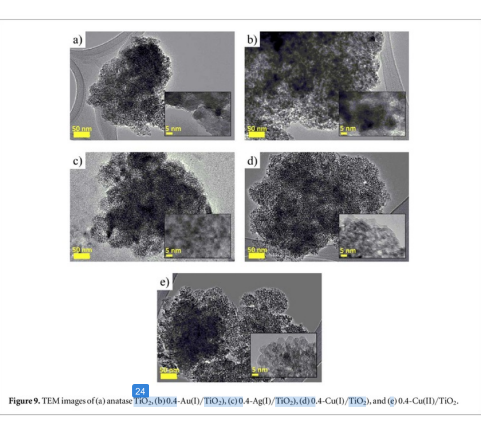


EDX analysis (figures 8(b), (d), (f), (g)) where the insert tables were summarized the mass percentage of respective elements. The insert tables showed that weight percentage of 0.4-M(I or II)/TiO₂ composites were almost closed to the loading of the complex to TiO₂. Hence, such impregnation method is good enough on the preparation of luminescent photocatalyst nanocomposite.

TEM measurements were performed to study the morphology of the composites. In figure 9, the morphology of selected 0.4-M(I or II)/TiO₂ composites (figures 9(b)–(e)) showed algost similar to that of anatase TiO₂ (figure 9(a)). Moreover, the particle size in all images of the composites was in good agreement with the crystallite sizes value as shown in table 1. These results indicate that small amount of complexes was dispersed well in anatase TiO₂ composites.

Figure 10 shows the emission properties of all photocatalyst composites. As discussed above, upon excited at the peak op of its excitation spectrum, it gave an emission peak centered (blaces ashed line) at 686, 471, 590 and 566 nm for Au(1), Ag(1), Cu(1) and Cu(11) pyrazolate complexes, respectively, du 59 the intermolecular metalmetal interaction [15–22]. On the other hand, photocatalyst composites showed a broad emission band with two peaks at 420 and 466 nm, which are corresponding to the emission of TiO₂. When loading amount of M(I or II) pyrazolate complexes were increased, the emission intensity of the composites at 420 and 466 nm were reduced for all composites. Such phenomena might be due to the surge period of TiO₂ will be transferred to the columnar assembly of metal complexes. Although all pyrazolate complexes were incorporated onto the surface of TiO₂, the

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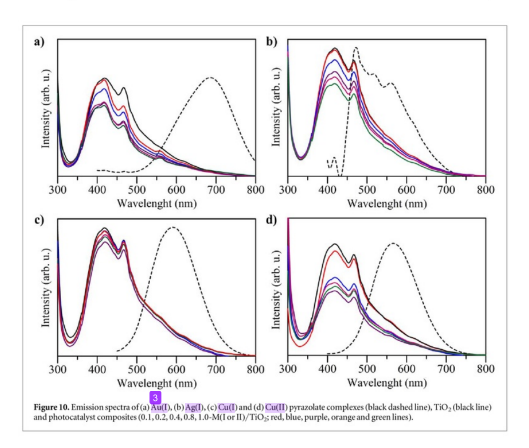


emission peak of M(I or II) pyrazolate complexes was not observed on all composites. These results suggested that low loading amount of pyrazolate complexes might not influenced $\rm TiO_2$ structure.

3.3. Photocatalytic activity of photocatalyst composites

The performance of luminescent group 11 117 al pyrazolate complexes/titanium oxide composites (x-M(I or II)/TiO₂) as a photocatalyst were evaluated using 2, 4-D as a model pollutant under a UV irradiation at ambient temperature in 1 h. Prior to this photocatalytic reaction, the equilibrium of adsorption-desorption process was firstly performed so that the adsorption efficiency was not involved in the calculation of photocatalytic activity. Based on stability 39 under UV light irradiation with M(I or II) pyrazolate complexes only and without photocatalyst, the degradation of 2, 4-D was not occurred due to lack of hydroxyl radical (•OH⁻) in the reaction system. By using standard calibration curve equation of 2, 4-D and 2, 4 CP with the previous equations above (y = 2,716,584x + 1263, y = 2,915,622x + 5829; respectively), the concentration of 2, 4-D and 2, 4-DCP in the testing can be determined.

Figures 11 (a) and (b) shows the photocatalytic testing of the photocatalyst composites where the removal and degradation percentages were evaluated. It showed that addition of M(I or II) pyrazolate complexes was found to give significant improvement in the photocatalytic activity of anatase TiO₂ even in 1 h. In particular, the composites 0.4-Au(I)/TiO₂ and 0.4-Ag(I)/TiO₂ gave the highest removal in 85% and degradation in 36% compared to anatase TiO₂ (49% and 10%, respectively). Interestingly, the composite Cu(I)/TiO₂ with 0.4 wt% gave even better removal in 99% and almost the same percentage of the degradation. In contrast, composites with complexation of Cu(II) ions by using the same ligand was found to show an enhancement in the degradation of 2, 4-D from 36% to 48%. The efficiency in removal of 2, 4-D (99%) for the composite Cu(II)/TiO₂ with 0.4 wt% as a photocatalyst was involved by its degradation 50 ficiency (48%) where the produced 2, 4-DCP was found to be around 51%. These results shot 231 that trinuclear Au(I), Ag(I) and Cu(I) pyrazolate complexes in anatase TiO₂ composites will signific 22 y improve the photocatalytic activity of TiO₂ due to the 22 uction of electron-hole recombination of TiO₂ (as shown in figure 10). Moreover, trinuclear Cu(I) pyrazolate complexes with longer metal-metal distance in composites as shown from the peak top of its emission spectrum (590 nm) for electron transfer from conduction band of TiO₂ is responsible for such performance in



the removal. Indeed, dinuclear Cu(II) pyrazolate complex with less rigid structure has more longer metal-metal distance (emission wavelength at 566 nm) to give such better performance in the degradation. In addition, further addition up to 1.0 wt% of M(I or II) pyrazolate complexes slightly decreaded the activity of TiO₂. Based on emission spectra in figure 10 and photocatalytic activity in figure 11, it can be proposed that the presence of small amount of M(I or II) pyrazolate complexes slightly a metal-metal distance in the columnar structure for light emission properties might play an important role to reduce the electron-hole recombination on TiO₂. The columnar assembly of the metal complexes with a *chair-like* coordination geometry as reported previously [15–22] is designed to trap the electron from the valance band of TiO₂ while the molecular structure promotes the performance of the slop to the degradation of the pollutant.

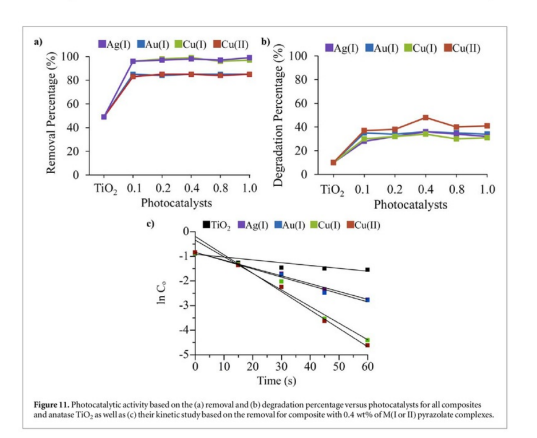
Figure 11(c) shows kinetic study of the com 37 ites for the removal of 2, 4-D at 15, 30, 45 and 60 mins. It was found that the removal of 2, 4-D could be fitted with the first order kinetic equation,

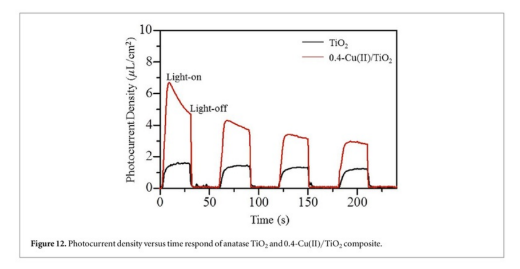
$$\frac{dC}{dt} = -kC \tag{1}$$

$$\ln C = -\frac{kt}{l} + \ln C_0 \tag{2}$$

which *C* and *C*₀ is final and initial concentrations of 2, 4-D, respectively and *k* is rate constant. This kinetic study showed that addition of M(I or II) pyrazolate complexes increased the *k* of TiO₂ (0.0105 min⁻¹) about 3 times for 0.4-Au(I)/TiO₂ (0.0327 min⁻¹) and 0.4-Ag(I)/TiO₂ (0.0317 min⁻¹) composites and 6 times for 0.4-Cu(I)/TiO₂ (0.0619 min⁻¹) and 0.4-Cu(II)/TiO₂ (0.0655 min⁻¹). Moreover, the above composite 0.4-Cu(II)/TiO₂ gave the highest *k* value compared to other composites where it was in good agreement with its highest photocatalytic activity (figure 11(a)). Indeed, such first order kinetic in the photocatalytic removal of 2, 4-D was also found with other photocatalysts in the previous reports [23–25].

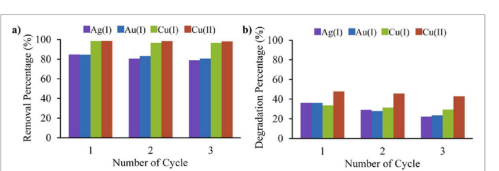
Photocurrent measurem 55 was carried out to study the higher activity of the 0.4-Cu(II)/TiO₂ composite compared to anatase TiO₂. A shown in figure 12, the photocurrent density of SPE substrate coated with 0.4-Cu(II)/TiO₂ composite showed a significant improvement around 2 times in the photocurrent density compared to anatase TiO₂. This photocurrent measurement for the composite was only dropped at first cycle and then was stable enough for each light-on/off event. Such higher density in the composite indicates that electron-hole pairs were generated in longer lifetime compared to in the anatase TiO₂ [12, 26]. These results also





47 indicate that the presence of Cu(II) pyrazolate complex in the composite effectively decreased electron-hole recombination process, thus gave much higher photocatalytic activity compared to anatase TiO₂.

In development of good photocatalysts, the stability is an important factor in the photocatalytic reactions as shown from unchanged photocatalytic activity while the chemical structure and morphology properties should remain the same. A series of experiment were done to study the stability of the photocatalyst composites with the evaluation of removal and degradation percentages for three cycles. As shown in figure 13, the photocatalytic activity of all composites almost provided the same results with only reduce in less than 5%. These results indicate the complexes in the TiO₂ composites are stable enough to work as photocatalysts.



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Figure 13. Photocatalytic stability experiments based on the (a) removal and (b) degradation percentage versus number of cycle for all composites with 0.4 wt% of M(I or II) pyrazolate complexes.

4. Conclusion

In conclusion, M(I or II) pyrazolate complexes in high yields were successfully synthesized by the complexation of 3, 5-dimethyl pyrazole ligand with different metal salt of group 11 where all complexes showed phosphorescent properties originating from the weak metal-metal interactions. A series of M(I or II) pyrazolate complexes/titanium oxide composit $_{36}^{36}$ -M(I or II)/TiO₂) with various loading amount of M(I or II) pyrazolate complexes (x = 0.1, 0.2, 0.4, 0.8, and 1.0 wt%) were prepared by using a simple impregnation method. Addition of M(I or II) pyrazolate complexes was not much affected the morphology and crystallite size of anatase TiO₂. In addition, presence of M(I or II) pyrazolate complexes $_{13}^{40}$ oreduced the emission intensity of TiO₂. 0.4 x-M(I or II)/TiO₂ composite showed the highest removal and degradation of 2, 4-D compared to TiO₂. This result demonstrated that optimum amount of M(I or II) pyrazolate complexes might act as an efficient modifier for 13 tase TiO₂, thus improved the photocatalytic activity. Interestingly, x-Cu(II)/TiO₂ showed the better degradation of 2, 4-D with 48% compared to other x-M(I)/TiO₂ and TiO₂ composites (36% and 10%; respectively). This work demonstrated that modification of TiO₂ with phosphorescent group 1 $_{45}^{45}$ razolate complexes at the molecular level (different metal ions and rigidity) as photocatalyst composites is a new approach to enhance photocatalytic activity of organic pollutants.

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