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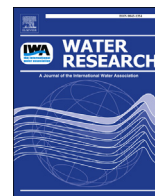
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## Photochlorination-induced transformation of graphene oxide: Mechanism and environmental fate



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### ABSTRACT

With the increasing production and wide utilization of graphene oxide (GO), the nanomaterials are expected to be released into the environment, and end up in surface waters, and/or wastewater treatment plants. This study explored the changes in the physicochemical properties of GO resulting from photochlorination—simulating the reactions that occur during water and wastewater treatment. Photochlorination resulted in significant changes in the surface oxygen-functionalities of the nanomaterials, and fragmenting of the graphenic carbon sheets was observed. We found that photochlorination can enhance the decomposition of GO through the formation of reduced GO. The changes in surface oxygen-functionalities of GO were attributed to the oxidation by chlorine of the nanomaterials' quinone groups, and further oxidation by  $\text{Cl}\cdot$  and/or  $\text{ClO}\cdot$  radicals. The surface charge of GO, measured by its zeta potential, increased in magnitude with chlorination but decreased in magnitude with photochlorination, leading to the decrease in the colloidal stability of the photochlorinated nanomaterials. The antibacterial effect of the nanomaterials increased with both chlorination and photochlorination. This study clearly shows how the physicochemical properties, and environmental fate and effect of GO are modified by photochlorination.

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### 1. Introduction

Graphene oxide (GO), a member of the graphene family nanomaterial, is a functionalized form of graphene, containing epoxy, hydroxyl, carbonyl and carboxyl functional groups (Bai et al., 2011; Lu et al., 2011). GO nanosheet is a unique carbonaceous nanomaterial that possesses excellent electrochemical, optical and mechanical properties. Hence, it has a wide range of applications, including energy storage, biomedicine, and pollution treatment, etc. (Chen et al., 2012; Adeleye et al., 2016; Petridis et al., 2016). The annual production of graphene family nanomaterials was estimated to be 120 metric tons in 2015, and expected to reach 1200 metric tons by 2019 (Zurutuza and Marinelli, 2014; Ciriminna et al., 2015). With the continued increase in production and application of graphene family nanomaterials, it is expected that the nanomaterials will be released into the environment, and may end up in

landfills, surface waters, and/or wastewater treatment plants. Accordingly, GO's potential adverse effects to the environment and human health, and its environmental transformations and implications need to be well-understood (Schmidt, 2009).

To date, only a few studies have focused on the transformation of GO in conditions similar and/or relevant to the natural environment (Chowdhury et al., 2015; Feng et al., 2015; Hou et al., 2015, 2016). Some of these studies have shown that sunlight is able to induce some physicochemical changes to the nanomaterials when present in aqueous media. For instance, solar irradiation caused GO to degrade and reduce (to reduced graphene oxide or rGO) (Hou et al., 2015, 2016). The production of  $\text{CO}_2$ , polycyclic aromatic hydrocarbons (PAHs), and reactive oxygen species (ROS) during solar irradiation of GO was also reported. Despite these existing studies, it is still impossible to predict the fate of these unique materials in most actual aquatic systems. As an example, if present in water treatment systems, GO will also be exposed to other chemicals used in water treatment (e.g. chlorination) in addition to sunlight irradiation.

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Chlorination is widely used in drinking water and wastewater (tertiary) treatment (Jiang et al., 2017; Li et al., 2017a, 2017b). If GO is not fully removed during earlier stages of (waste)water treatment, it will reach the chlorine disinfection step where it will be exposed to both disinfection and, perhaps, sunlight irradiation (Gottschalk et al., 2009; Wang et al., 2012b). Chlorination can potentially change the morphology and the oxygen-containing functional groups of carbonaceous materials, including GO (Li et al., 2010; Liu and Zhang, 2014; Stratakis et al., 2014; Savva et al., 2014; Zhai et al., 2014; Du et al., 2016; Li et al., 2016). Concurrent or subsequent sunlight irradiation of chlorinated GO may result in further transformation, affecting further the physico-chemical properties (and thus, environmental implications) of GO. This potential transformation of GO by photochlorination has not received much attention as most existing studies on the environmental fate of GO focused on its colloidal stability and sunlight-induced transformations (Chowdhury et al., 2013, 2015; Wu et al., 2013; Hou et al., 2015).

The objectives of the study were to (1) investigate the photochemical transformation of GO during traditional disinfection via photochlorination; and (2) evaluate the colloidal stability and toxicity of GO after photochemical transformations.

## 2. Material and methods

### 2.1. Materials and stock preparation

GO nanosheets used in this study were purchased from Planano Technology Co. (Tianjin, China), and used without further modifications. Based on the information provided by the supplier, the nanosheets were synthesized from graphite using a modified Hummers method (Kotov, 2006). To prepare GO stock suspensions, 400 mg of pristine GO nanosheets were added into 500 mL deionized (DI) water and then ultrasonicated for 4 h at 100 W using a Vibra-Cell VCX800 sonicator (Sonics & Material, Newtown, CT). The GO stock suspension was stored in the dark at 4 °C until use.

The stock of free chlorine solution (1000–3000 mg/L) used in the experiments was prepared from 5% sodium hypochlorite (NaClO; Sigma-Aldrich, St. Louise, MO). The chlorine stock was stored in a flask covered with aluminum foil and kept in the dark at 4 °C. The stock solution was periodically standardized by diethyl-*p*-phenylene diamine (DPD) ferrous titration (American Public Health Association, 1998).

### 2.2. GO chlorination and photochlorination

Two photochlorinated GO samples were prepared, and two chlorinated GO samples were prepared as a comparison. GO concentration was 10 mg/L for all experiments and the chlorine dosage was either 10 mg/L or 100 mg/L as chlorine (Cl<sub>2</sub>). The 10 mg/L Cl<sub>2</sub> dosage was used to simulate the typical chlorine concentration used in water or wastewater disinfection processes (Wang et al., 2006), while the 100 mg/L Cl<sub>2</sub> dosage was used to amplify the effects of chlorination on GO to clearly understand the mechanism. Chlorinated GO samples were kept in the dark throughout the experiment, while photochlorinated GO samples were kept under simulated sunlight for 8 h each day. The samples were buffered at pH 7 with 10 mM phosphate, and total reaction time was 5 d (similar to a previous study) (Li et al., 2016) for all samples. Simulated sunlight irradiation was achieved by placing the reaction quartz flask under a PL-XQ 500W adjustable xenon lamp (Lanpu Co., Shanghai, China) as a light source. The power of xenon lamp used in the study was 420 W and light intensity was 168 mW/cm<sup>2</sup>. The light spectrum from the xenon lamp was similar to the summer sunlight spectrum collected in Tianjin, China on

May 28, 2015 (Fig. S1).

To stop the chlorination reaction, ascorbic acid (at concentrations 1.2 times of the residual Cl<sub>2</sub>, measured with a HACH Cl<sub>2</sub> colorimeter) was added after which the GO samples were cleaned via ultrafiltration and then characterized. Ultrafiltration was done with stirred ultrafiltration cells (Millipore Corporation, Billerica, MA) to remove phosphate buffer, free Cl<sub>2</sub> residuals and other dissolved ions. Detailed information on the ultrafiltration procedure is provided in [Supplementary Material section S1.0](#). The GO samples treated with 10 mg/L and 100 mg/L Cl<sub>2</sub> are hereafter referred to as LCGO and HCGO, respectively. The simulated sunlight-exposed GO samples treated with 10 mg/L and 100 mg/L Cl<sub>2</sub> are hereafter referred to as PLCGO and PHCGO, respectively.

### 2.3. Material characterization

The functional groups of GOs were characterized via X-ray photoelectron spectroscopy (XPS; PHI 5000 VersaProbe, Japan) and Fourier transform infrared spectroscopy (FTIR; Bruker TENSOR 27, Bruker Optics Inc., Germany). Raman spectra (RM2000 Raman spectrometer, UK) of samples were used to evaluate the defects/changes of GO before and after (photo)chlorination. UV–Vis absorbance spectra of the suspensions were obtained with a UV-2401 spectrophotometer (Shimadzu Scientific Instruments, Columbia, MD). Atomic force microscopy (AFM; MMAFM/STM, D3100M, Digital Ltd., USA) and scanning electron microscopy (SEM; FEI Nova Nano 230, Hillsboro, OR) studies were carried out to analyze the morphology of samples. Total organic carbon (TOC) and inorganic carbon (IC) (multi N/C 3100, Analytik Jena AG, Germany) of GO suspensions were also determined before and after (photo)chlorination.

### 2.4. Zeta potentials and hydrodynamic diameter (*D<sub>h</sub>*) determination

The surface charge of GO nanomaterials before and after (photo)chlorination treatments were estimated by determining their zeta (ζ) potential using a ZetaSizer Nano ZS (Malvern Instruments, UK). Hydrodynamic size distributions of GO samples were determined before and after (photo)chlorination treatments via dynamic light scattering (DLS) using the ZetaSizer. The pH of GO suspensions was measured using a SevenCompact S230-Basic pH Meter (Mettler-Toledo International Inc., Zurich, Switzerland).

### 2.5. Antibacterial tests

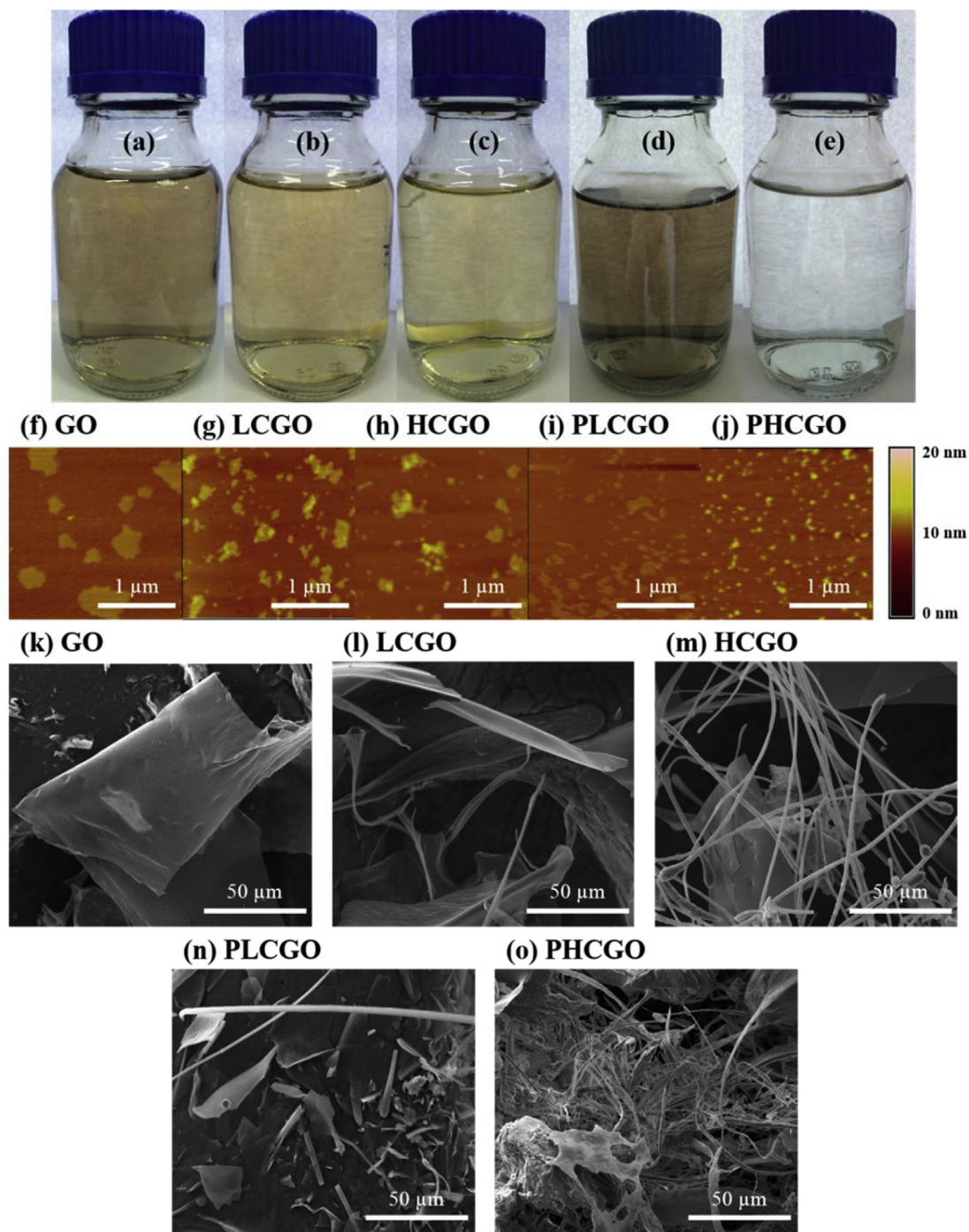
To investigate the toxicity of (photo)chlorinated GO, a Gram-negative bacterial strain, *Pseudomonas aeruginosa*, was exposed to the transformed nanomaterials in isotonic saline solution. A physiological saline solution containing the same concentration of bacteria and pristine GO was used as a control. The details of bacterial strains cultivation procedure were provided in [Supplementary Material section S2.0](#). After cultivation, the bacteria and GOs mixtures were incubated in a constant temperature (37 °C) incubator with gentle shaking. The total incubation time was 1 h, after which a 1 mL aliquot was drawn out for analyses. Additional experiments were carried out after incubation times of 2 and 3 h for comparison. Antimicrobial activity was analyzed via flow cytometry (FCM) using a CyFlow Space (Sysmex Partec GmbH, Germany) equipped with a 50 mW solid-state laser. Detailed information on FCM measurements was provided in [Supplementary Material section S3.0](#).

### 3. Results and discussion

#### 3.1. Effects of photochlorination on physical changes of GO

Changes in the physical appearance and morphology of GO after photochlorination were examined. Fig. 1a–e shows the appearance of GO suspensions before and after (photo)chlorination. The appearance of GO suspension clearly changed upon treatment with different concentrations of chlorine, with or without light. The color of GO faded with increased chlorination, as can be seen in the

images of LCGO (Fig. 1b) and HCGO (Fig. 1c). This is mainly due to oxidation of GO functional groups by chlorine, which may destruct some chromophore units (such as C=C and C=O groups on the graphitic carbon) (Szyrkowicz et al., 2001). However, the color trend changed upon the exposure of GO suspensions to light alongside chlorination. While PLCGO was slightly darker than pristine GO (Fig. 1d), but PHCGO was completely clear (Fig. 1e). According to a previous study, sunlight irradiation may reduce GO to reduced GO (rGO), which has a darker color than GO (Hou et al., 2015). Formation of rGO explains the color of PLCGO but not



**Fig. 1.** The photograph of GO suspensions before and after chlorination with or without simulated sunlight exposure: (a) pristine GO, (b) LCGO, (c) HCGO, (d) PLCGO, and (e) PHCGO). AFM micrographs of GOs upon chlorination with or without light irradiation: (f) pristine GO, (g) LCGO, (h) HCGO, (i) PLCGO, and (j) PHCGO. SEM images of GOs upon chlorination with or without light irradiation: (k) pristine GO (l) LCGO, (m) HCGO, (n) PLCGO and (o) PHCGO. (Reaction time = 5 d at 8 h/d).



PHCGO.

The AFM micrographs (Fig. 1f–j) show that GO nanoplatelets were damaged at the edges after mild or heavy (photo)chlorination, and the GO nanoplatelets were heavily fragmented by photochlorination (Fig. 1i–j), especially with high  $\text{Cl}_2$  concentration (PHCGO sample). SEM images (Fig. 1k–o) further prove the changes in the morphology of GO after treatment with chlorination and photochlorination. It should be noted that several of the extremely small fragments formed after the treatments may have been removed during the ultrafiltration step in the sample preparation. Scrolled GO sheets in the LCGO and HCGO samples, which were also found in our previous study (Li et al., 2010), may be due to asymmetric distributions of cations, creating a polarization that is relieved by curling (Schaak and Mallouk, 2000). The fragmentation of chlorinated GO nanosheets appeared to be intensified upon irradiation as can be seen in the much smaller sizes of PLCGO and PHCGO samples (Fig. 1n and o), which agrees well with AFM data. However, much more scrolled GO in the PHCGO sample also emphasize the importance of chlorination procedure in the photochlorination procedure.

### 3.2. Effects of photochlorination on chemical properties of GO

The influence of photochlorination on the surface chemistry of GO was evaluated via XPS, FTIR, Raman, and UV–Vis spectrometric analyses. Comparisons of the XPS survey and high resolution spectra of GO, LCGO, HCGO, PLCGO, and PHCGO are shown in

Fig. 2a–f. In the high resolution spectra, the C1s peaks located at binding energy (BE) of 284.5 eV, 286.7 eV, 287.8 eV, and 288.8 eV were assigned to the carbon atoms or oxygen-functional group in aromatic rings (C–C/C=C), hydroxyl or epoxy (C–OH/C–O–C), carbonyl (C=O), and carboxyl (O–C=O), respectively (Fig. 2a–e) (Chandra et al., 2010).

Quantitative analyses of these carbon species (Fig. 2g) showed that (in comparison to pristine GO) the fraction of C–OH/C–O–C and O–C=O groups increased in the LCGO samples, whereas the fraction of C=O group decreased. However, in HCGO samples, only the fraction of O–C=O groups increased, while C–OH/C–O–C and C=O groups decreased. The O/C ratio of LCGO and HCGO samples was similar to that of pristine GO, indicating that little or no further addition of new oxygen groups to the graphitic layers occurred upon 10 and 100 mg/L chlorine treatments. The decrease of C–OH groups and increase of O–C=O groups in HCGO sample suggests further oxidation of C–OH functional group to O–C=O functional group in the presence of higher concentrations of chlorine.

In contrast, the ratio of C–C/C=C groups in PLCGO and PHCGO increased significantly (Fig. 2g), which suggests decomposition (loss) of some of the oxygen-containing functional groups in GO (e.g., C–OH/C–O–C) due to sunlight irradiation. As shown in the XPS survey spectra of the different GOs in Fig. 2f, a peak for C–Cl (or Cl–C=O) bond with BE of ~200.3 eV emerged after (photo)chlorination (Kim et al., 2013; Zhang et al., 2013). The strongest C–Cl/Cl–C=O peak was found in the PHCGO sample, which indicates that simulated sunlight irradiation enhanced the chlorination of GO at

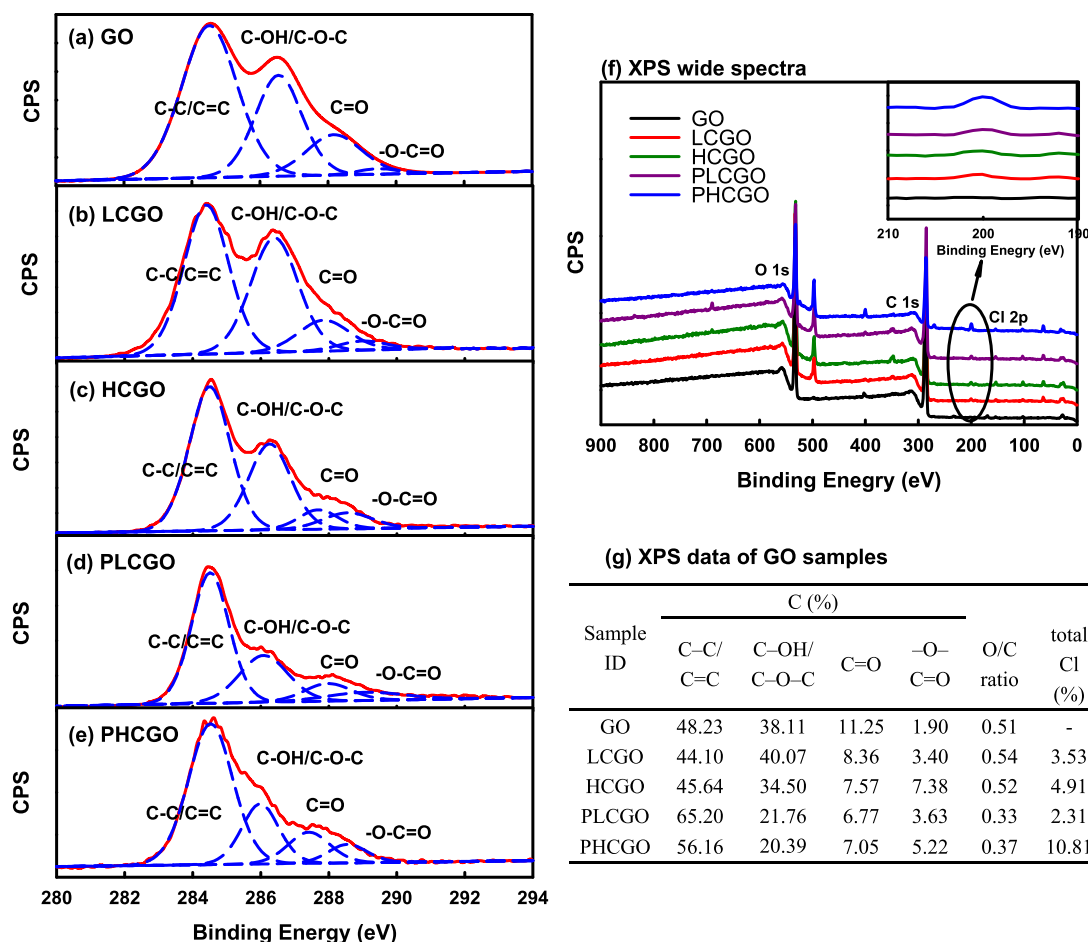


Fig. 2. X-ray photoelectron spectroscopy (XPS) analysis of GO before and after chlorine treatment with or without light irradiation. High resolution XPS spectra of the C 1s peaks of pristine GO (a), LCGO (b), HCGO (c), PLCGO (d) and PHCGO (e). (f) XPS survey spectra of all the GO nanosheets. (g) Quantitation of C1s peaks of the GO samples.

the high chlorine concentration treatment.

Changes in the surface chemistry of GO upon photochlorination were further characterized by FTIR (Fig. 3a and b). Pristine GO exhibited four characteristic peaks located at 612, 1383, 1628 and 1735  $\text{cm}^{-1}$ , corresponding to C–C vibration (Kratschmer et al., 1990), hydroxyl group (O–H bending band), carboxyl group stretching, and C=O stretching, respectively (Hontoria-Lucas et al., 1995; Acik et al., 2011; Li et al., 2012). After GO reacted with free  $\text{Cl}_2$  (10 mg/L or 100 mg/L) for 5 d, the peak for C=O stretching in GO decreased, while the peaks for C–OH and O–C=O functional groups increased, indicating a transformation of C=O groups to O–C=O and C–OH groups. In addition, two new peaks were observed (at 798 and 1100  $\text{cm}^{-1}$ , which are typical of C–Cl stretching and aryl C–Cl, respectively (Salavagione and Martinez, 2011; Wang et al., 2012a). This suggests that one of the most important outcomes of the reaction of GO with  $\text{Cl}_2$  is the incorporation of Cl atoms into unsaturated ethylic sites of GO aromatic rings. Changes in the surface chemistry of GO after photochlorination, observed via FTIR, were very similar to the changes observed with only chlorination. The most obvious difference observed between chlorination and photochlorination was a significantly larger peak at 1100  $\text{cm}^{-1}$  (assigned to aryl C–Cl group) on the PHCGO sample, which may be due to enhanced chlorination of GO in the presence of irradiation as also observed with XPS analysis.

Raman spectra of pristine, chlorinated and photochlorinated GOs are shown in Fig. 3c. The ratio of the D band (1360  $\text{cm}^{-1}$ ), which measures the presence of disorder in  $\text{sp}^2$ -hybridized carbon systems, to G band (1590  $\text{cm}^{-1}$ ) that evaluates stretching of C–C bonds in graphitic materials, ( $I_D/I_G$ ), were compared (Dresselhaus et al., 2010). The  $I_D/I_G$  value of all treated GO increased compared to that of pristine GO. The highest increases in  $I_D/I_G$  values were observed in PLCGO (increased by 0.15) and PHCGO (increased by

0.16) samples, which indicates that many more defects were formed on GO after photochlorination (compared to chlorination alone where the increase in  $I_D/I_G$  was <0.1).

The UV–Vis spectra of all the samples are shown in Fig. 3d. The characteristic absorbance peak of GO at 230 nm was detected in pristine GO, LCGO and HCGO samples. However, the peak (and thus, GO concentration) decreased sharply in samples treated with chlorine and simulated sunlight (i.e., PLCGO and PHCGO), which is quite different from the results observed when GO was only subjected to irradiation in a previous study (Hou et al., 2015). This result suggests that some decomposition of GO occurred in the presence of both chlorine and sunlight, but not (or much less so) in chlorine alone. In addition, a peak around 300 nm (which represents residual chlorine) was observed in the HCGO sample, mainly due to high  $\text{Cl}_2$  concentration used in that condition (Watts and Linden, 2007). However, a similar residual chlorine peak was not observed in PHCGO (which was also treated with a similar level of  $\text{Cl}_2$  as HCGO), suggesting a transformation of residual chlorine under sunlight irradiation.

### 3.3. Mechanisms controlling GO transformation by photochlorination

Based on the spectroscopic and electron microscopic analyses, we believed that chlorination and/or sunlight irradiation are effective in changing the surface chemistry and morphology of GO.

$\text{HClO}$ , a weak acid produced from chlorine in water, cannot substantially further increase the oxygen content of GO upon chlorination treatments although we observed the oxidation of quinone/carbonyl groups on the basal plane or edges of GO to epoxy, hydroxyl, and carboxyl groups (as shown by XPS and FTIR). Rather, chlorination of GO led to electrophilic additions of chlorine

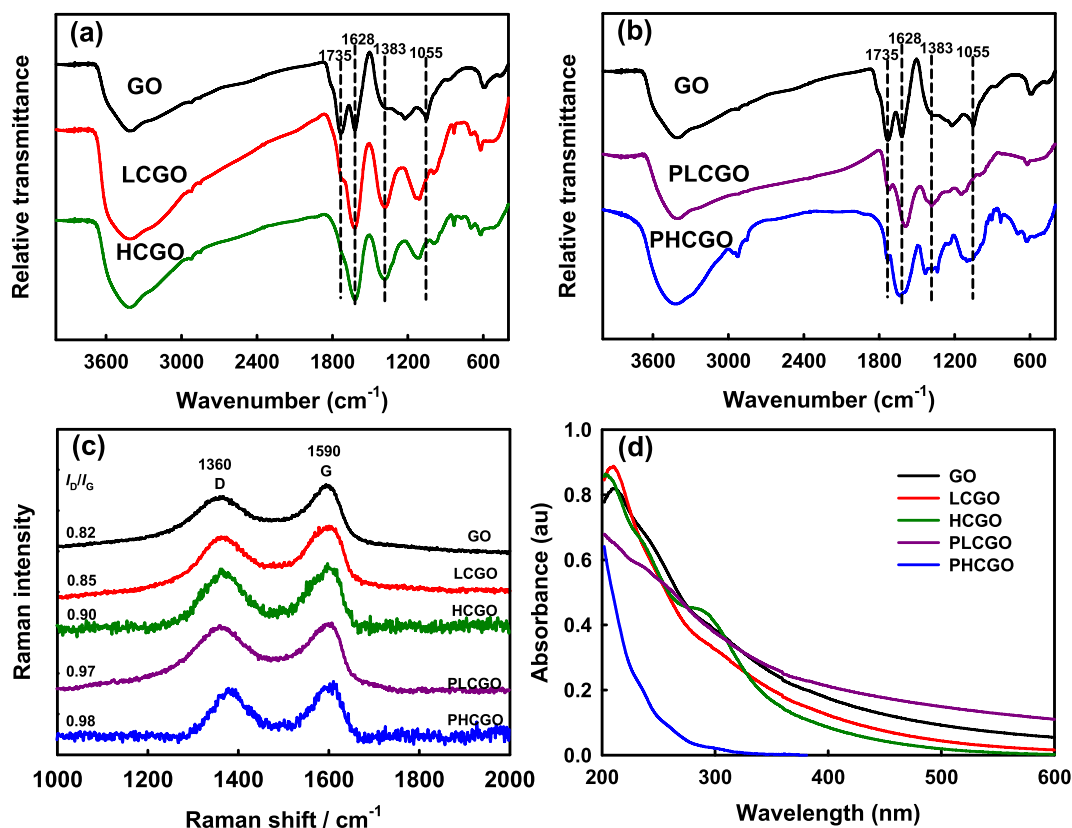


Fig. 3. (a–b) Fourier transform infrared (FT-IR) spectra, (c) Raman spectra, (d) UV–Vis spectra of GO, LCGO, HCGO, PLCGO and PHCGO.

atoms at the unsaturated sites of GO's benzene ring (Li et al., 2016). Higher HClO dosage had additional effects on GO's surface chemistry, such as further oxidation of hydroxyl groups to carboxyl groups (Sivey and Roberts, 2012). However, neither HClO nor aqueous Cl<sub>2</sub> was reactive enough to break GO's carbon platelets and add new oxygen functional groups onto them.

According to Hou and co-workers, irradiation led to the disproportionation of GO to CO<sub>2</sub>, photoproducts similar to reduced GO, and PAH-like materials (Hou et al., 2015). Similarly, we observed the production of CO<sub>2</sub> (inorganic carbon) and reduction of GO to rGO upon photochlorination of GO in this study (Fig. S2). CO<sub>2</sub> was formed from the oxidation of the unsaturated oxygen-containing functional groups on GO. The formation and release of CO<sub>2</sub> from GO platelets led to the fragmentation of the platelets, and the reduction of the oxygen-containing functional groups. However, the characterization results indicate that the effects of photochlorination on GO were much complicated, including other important reactions.

As a photo-reactive semiconductor, irradiation of GO results in the formation of oxidative (valence band holes, h<sub>vb</sub><sup>+</sup>) and reductive (conduction band electron, e<sub>aq</sub><sup>-</sup>) transients (Hou et al., 2015). These co-occurring transients react with dissolved oxygen in water to yield reactive oxygen species (ROS) such as <sup>1</sup>O<sub>2</sub>, •OH and O<sub>2</sub><sup>-•</sup>. The h<sub>vb</sub><sup>+</sup> may easily affect the oxidation of O-functional groups, while e<sub>aq</sub><sup>-</sup> promotes the formation of C–Cl. Meanwhile, photolysis of aqueous Cl<sub>2</sub> occurs when irradiated at wavelengths below 511 nm to form chlorine radicals (Cl•) and hydroxyl radicals (•OH) as shown in Equation (1) (Carrión et al., 1972; Nowell and Hoigné, 1992; Cataldo, 1994; Wang et al., 2012a; Zhao et al., 2011):



Cl• and •OH could further react with aqueous Cl<sub>2</sub> to form

oxychlorine radicals (ClO•) (Carrión et al., 1972; Nowell and Hoigné, 1992). Unlike HClO, ClO<sup>-</sup> and aqueous Cl<sub>2</sub> species that are formed from NaClO in the absence of light, Cl• and ClO• that are formed during irradiation are much more reactive with GO particles during photochlorination. Weak oxidants, HClO, ClO<sup>-</sup> (formed during chlorination), contribute to the formation of C–Cl and –COOH bonds at the edges of GO. Cl• and ClO•, which are formed when irradiation is coupled with chlorination, are stronger oxidants, and may easily attack GO at its reactive sites (including e<sub>aq</sub><sup>-</sup>) and even some nonreactive sites to form C–Cl and Cl–C=O bonds. The deconvolution of Cl 2p XPS peak showed the presence of C–Cl (BE = 200.3 eV) and Cl–C=O (BE = 202 eV) bonds in PLCGO and PHCGO samples; but Cl–C=O bonds were not present in samples treated only with chlorination (Fig. 4). As such, under sunlight irradiation, Cl<sub>2</sub> species/radicals species can attack both the edges and basal plane of GO sheets, while the basal plane is less vulnerable without irradiation.

Based on the results in this study and analysis above, we believe that the chlorination and irradiation effects are both important during the photochlorination procedure, but the dominant factor depends on chlorine concentrations. At low concentrations (such as 10 mg/L), chlorine was completely decomposed by irradiation to form Cl• and ClO• radicals. Although Cl• and ClO• are strong oxidants, their low concentrations limited their effects and the formation of C–Cl/C–Cl=O bonds. As a result, irradiation was the more dominant factor in the transformation of PLCGO samples. At much higher chlorine dosage (100 mg/L), however, much more Cl• and ClO• were formed during the irradiation of chlorine. High concentrations of HClO, Cl<sub>2</sub>, Cl• and ClO• resulted in the formation of C–Cl/C–Cl=O bonds on the reactive sites within GO's fragmented sheets. The possible scheme of photochlorination of GO is shown in Fig. 5. In the natural environment, other factors including pH, ions, and natural organic matter (NOM) will also play a role in the photochlorination of GO. In

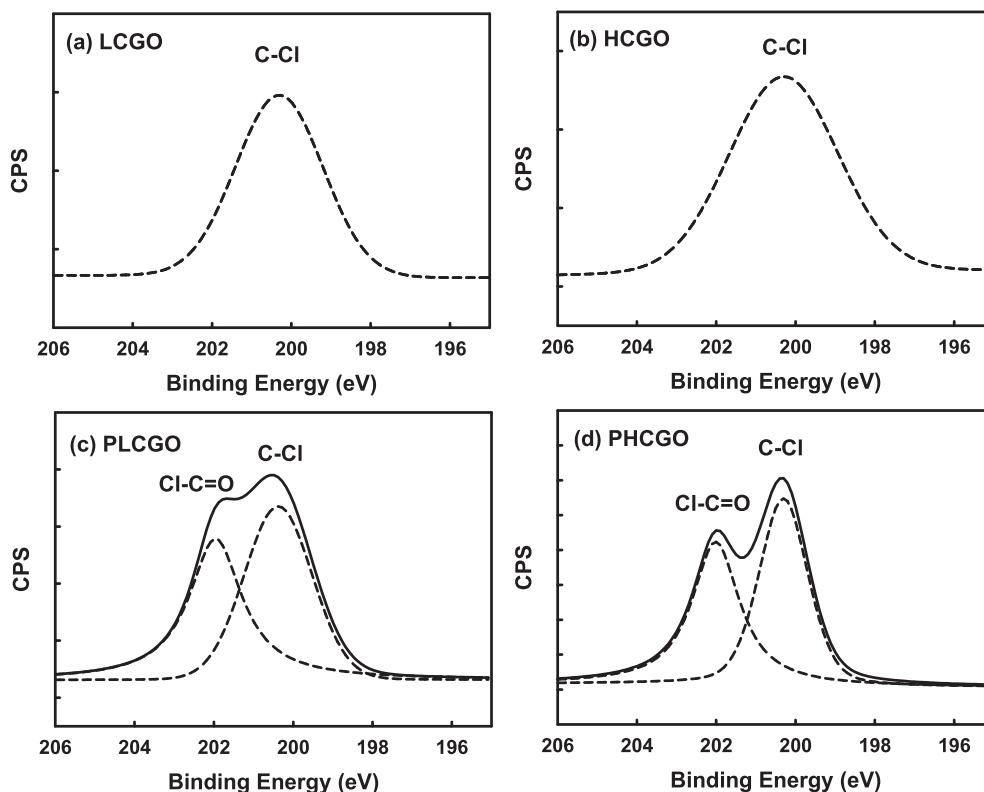


Fig. 4. High-resolution XPS spectra of the Cl 2p peak for (photo)chlorinated GO. Deconvolution revealed the presence of Cl–C (200.3 eV) and Cl–C=O (202 eV) groups.

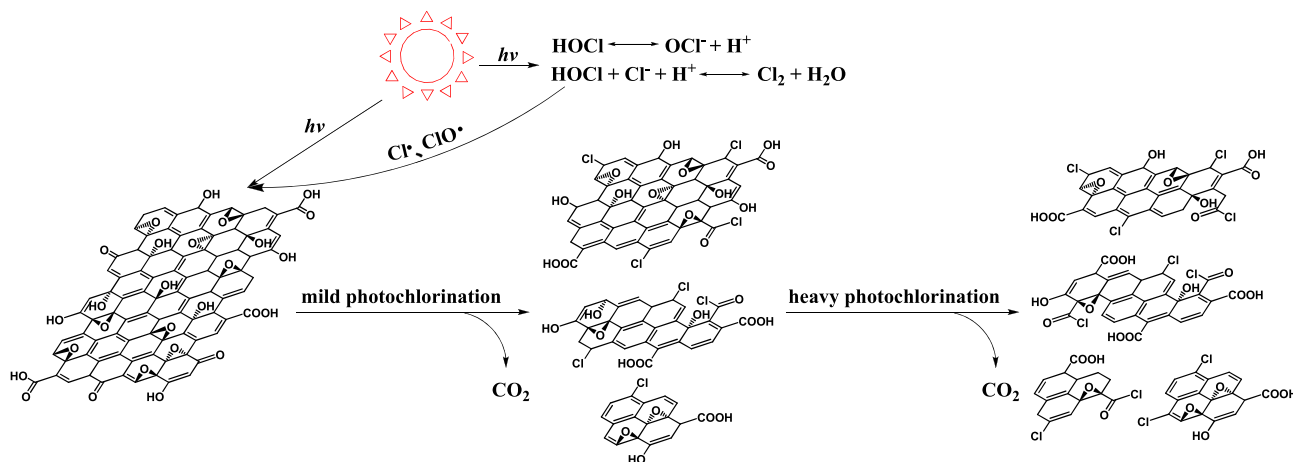


Fig. 5. Proposed mechanism of photochlorination of graphene oxide nanosheet.

particular, NOM may affect the transformation of GO in two possible ways: (1) NOM may coat GO nanosheets (Chowdhury et al., 2015) and thereby attenuate the effects of direct irradiation, and thus, decomposition of GO; and (2) NOM could also consume aqueous chlorine and other chlorine radicals (Li et al., 2017a), minimizing the formation of C-Cl/C-Cl=O bonds on the surface of GO.

#### 3.4. Effect of (photo)chlorination on colloidal stability of GO suspensions

The  $\zeta$  potential and size distribution of GO before and after the (photo)chlorination were shown in Fig. 6.  $\zeta$  potential of GO increased in magnitude from  $-45$  to  $-50$  mV after chlorination, and decreased to  $-40$  mV after photochlorination (Fig. 6a). Chlorination led to the oxidation of the  $-OH$  group on the surface of GO to  $-COOH$  group, which may lead to increase in the pH of GO suspension. Photochlorination of GO may also lead to increase in pH since  $-OH$  group was further oxidized. The pH of GO suspensions increased from 5 to a range of 8–10 (after chlorination) or a range of 5–7 (after photochlorination) as shown in Fig. S3. Both increase in  $-COOH$  group and elevated pH led to increases in the magnitude of  $\zeta$  potential observed in the (photo)chlorination treated samples. However, light-induced disproportionation of GO led to decrease in the  $\zeta$  potential of GO, making the nanosheets potentially less stable in aquatic systems. However, the  $\zeta$  potential of GO exceeded  $-30$  mV (the typical threshold of colloidal stability

(Everett, 1988)) in all the treatments indicating that the GO photo-transformation products were still reasonably stable.

DLS data showed that the hydrodynamic diameter of GO was not substantially changed upon (photo)chlorination. As shown in Fig. 6b, the size distributions of GO suspensions were similar before (GO) and after chlorination (LCGO and HCGO). Bimodal size distribution was observed in PLCGO and PHCGO with the new size distribution peak appearing at sizes smaller than that of pristine GO. This also agrees with microscopic data that showed degradation of GO sheets to smaller sizes after photochlorination.

#### 3.5. The toxic effects of (photo)chlorinated GO

The toxic effects of all the GO samples, measured by death rate, to *P. aeruginosa* was determined and shown in Fig. 7. In order to clearly see the antibacterial effect of GO samples, 100 mg/L GO was used in the experiment. Antimicrobial activity which was analyzed via flow cytometry (FCM) was shown in Fig. 7a. Bacterial death rate, resulting from 1 h of contact time with the nanomaterials, increased significantly from 15% when exposed to pristine GO to  $\sim 20\%$  ( $p < 0.05$ ) and  $\sim 30\%$  ( $p < 0.05$ ) when the cells were exposed to GO nanosheets treated with low dosage of  $Cl_2$  (LCGO) and high dosage of  $Cl_2$  (HCGO), respectively. Toxicity did not increase any further when the cells were exposed to GO samples that were simultaneously exposed to chlorination and sunlight irradiation (Fig. 7b). In fact, death rates were significantly less ( $p < 0.05$ ) in PHCGO than in HCGO. The highest toxicity was observed in HCGO

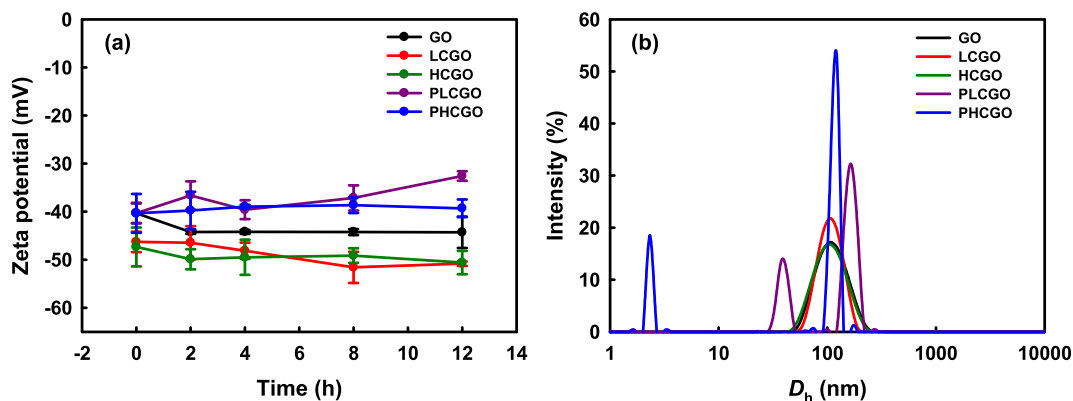
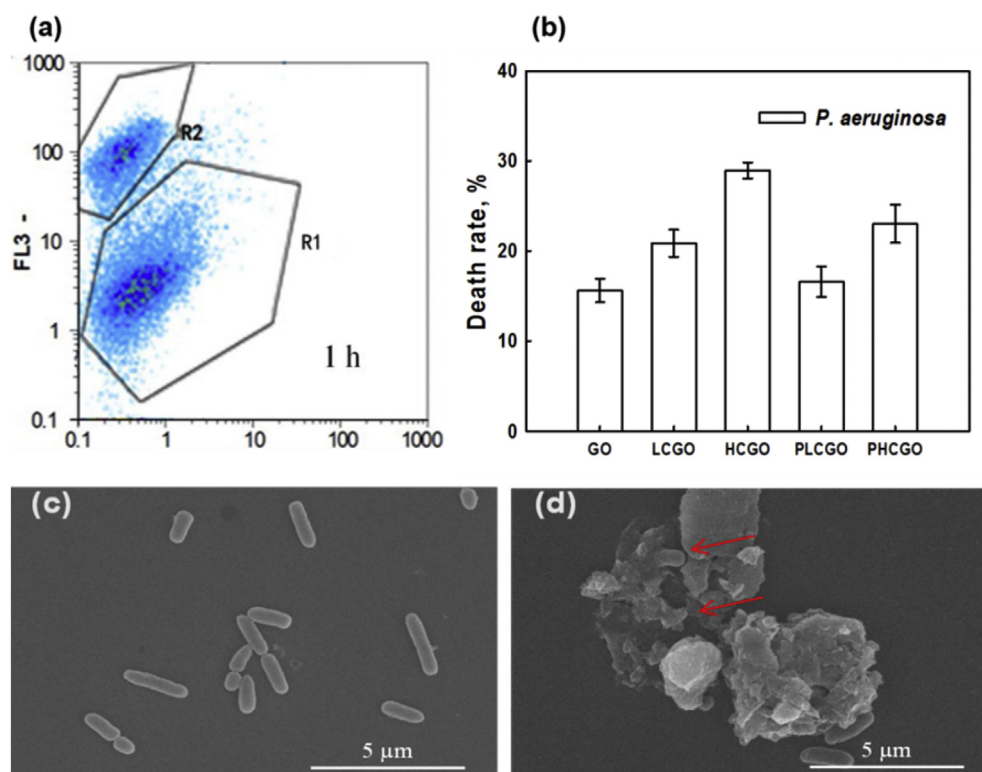


Fig. 6. (a) Zeta potential, and (b) Size distribution of GO before and after (photo)chlorination. GO concentration = 10 mg/L.





**Fig. 7.** (a) A typical flow cytometry (FCM) image of antibacterial activity of GO samples to *P. aeruginosa*. R1 represents live bacteria and R2 represents dead bacteria. (Contact time = 1 h), and (b) Effect of (photo)chlorination on the toxicity of GO to *P. aeruginosa* cells. (GO concentration = 100 mg/L; contact time = 1 h; pH = 7). SEM images of *P. aeruginosa* before (c) and after (d) contact with chlorinated GOs (arrows show some sliced bacterial cells).

samples. Further studies showed that the antibacterial effect of GO to the bacterium was both concentration- and exposure time-dependent (Figs. S4 and S5).

Generally, the antimicrobial effects of GO materials have been proposed to occur via three major mechanisms: (1) acting as *nanoknives* due to their sharp edges (Chen et al., 2013, 2014; Wang et al., 2013; He et al., 2015), (2) oxidative stress mediated with or without the production of ROS (Lewinski et al., 2008; Liu et al., 2013; Smith and Rodrigues, 2015), and (3) reduction of nanomaterials, which may decrease their stability, and thus the concentration on the bacteria due to heteroaggregation (Carpio et al., 2012; Perreault et al., 2015). As shown in Fig. 7c–d, SEM images provided evidence for slicing of bacterial cells by GO in this study (*nanoknives* effect). As explained earlier, chlorination of GO resulted in scrolling and increased defect at the edges; while photochlorination mainly led to defects on the edges and basal planes as well as fragmentation. In addition, photochlorinated GO was also reduced compared to the GO exposed to only chlorination. The scrolling of GO nanosheets may decrease their ability to act as a *nanoknives* but fragmentation and increased edge defects enhanced the antimicrobial effects of GO (Poland et al., 2008; Yang and Ma, 2010; Perreault et al., 2015). The results indicate that the defects from chlorination procedure may highly increase the antibacterial effects of GO, but the reduction (with photochlorination) may inhibit them. In this study, we only provided a preliminary investigation of the effect of (photo)chlorination-induced transformation on the antimicrobial properties of GO but further studies are needed to fully understand the mechanisms.

#### 4. Conclusions

Release of graphene oxide nanosheets into water and

wastewater treatment plants will facilitate some physicochemical transformations of the nanomaterials due to the exposure of the nanomaterials to chlorination and irradiation. This study showed that the oxygen-containing groups of graphene oxide were either transformed during chlorination, or simultaneously transformed and reduced when chlorination was combined with sunlight irradiation (as we would expect to occur in wastewater treatment plants). These chemical changes led to physical changes in the nanosheets—mainly scrolling and fragmentation to much smaller sizes. These changes influenced media pH, and nanomaterial size distribution, surface charge, and antibacterial effects. Although not tested in this study, the changes will also affect other processes such as nanomaterial aggregation, adsorption, transport, and interactions with other surfaces (geogenic and biogenic), etc.

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#### Appendix A. Supplementary data

Supplementary data related to this article can be found at <http://dx.doi.org/10.1016/j.watres.2017.07.054>.

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