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Photocatalytic generation of hypochlorous acid on plasmonic Au/AgCl catalysts prepared by microwave-assisted reduction[†]

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Plasmonic Au particles loaded on AgCl powders (Au/AgCl) prepared by microwave-assisted reduction, when photoirradiated under visible light in aerated chloride solutions, efficiently generate hypochlorous acid owing to the large Au-AgCl periphery sites by the formation of small Au particles.

Hypochlorous acid (HClO) is a strong, water-soluble oxidant that is commonly used for disinfection, bleaching, and sterilization.¹ HClO is industrially produced from Cl_2 gas manufactured by the electrolysis of Cl^- solutions,² which involves the oxidation of Cl^- at the anode (eqn (1)) and the reduction of O_2 (eqn (2)) at the cathode.³ The total electrolysis can be expressed as shown in eqn (3).⁴ The dissolution of Cl_2 gas in water generates HClO by disproportionation (eqn (4)). The electrolysis (eqn (3)) is an up-hill reaction with a large Gibbs free energy gain and requires a huge amount of electrical energy.⁵ Therefore, a clean and sustainable method is necessary for HClO production.

$$2\text{Cl}^- \rightarrow \text{Cl}_2 + 2e^- (+1.36 \text{ V} \nu s. \text{ NHE})$$
 (1)

$$O_2 + 4H^+ + 4e^- \rightleftharpoons 2H_2O \quad (+1.23 \text{ V} \nu s. \text{ NHE}) \qquad (2)$$

$$2\text{Cl}^{-} + 1/2\text{O}_2 + 2\text{H}^+ \to \text{Cl}_2 + \text{H}_2\text{O} \quad (\Delta G^{\circ} = +105 \text{ kJ mol}^{-1} \text{ at pH 7.0})$$
(3)

$$Cl_2 + H_2O \rightleftharpoons HClO + H^+ + Cl^- \quad (K = 3.94 \times 10^{-4} \text{ M}^2 \text{ at } 298 \text{ K})$$

(4)

^b Innovative Catalysts Science Division, Institute for Open and Transdisciplinary Research Initiatives (ICS-OTRI), Osaka University, Suita 565-0871, Japan Semiconductor photocatalysis can generate HClO in Cl⁻ solutions with O₂ at ambient temperature (eqn (3)). The photogenerated valence band holes (h_{VB}^+) oxidize Cl⁻ to form Cl₂ (eqn (1)), which generates HClO *via* disproportionation (eqn (4)), while the conduction band electrons (e_{CB}^-) reduce O₂ (eqn (2)). However, early reported photocatalysts produced low-HClO-concentration solutions (<3 ppm).⁶⁻⁹ This is because (i) the oxidation of bulk Cl⁻ by h_{VB}^+ is difficult to promote due to its deep oxidation potential (eqn (1)); and (ii) these catalysts are mainly photoexcited by UV light (λ < 400 nm), which readily promotes photolysis of HClO into O₂ and Cl⁻ (eqn (5)).⁴ The design of photocatalysts that efficiently oxidize Cl⁻ under visible light is therefore necessary.

$$2\text{HClO} \rightarrow \text{O}_2 + 2\text{H}^+ + 2\text{Cl}^- \tag{5}$$

Recently, we found that visible light irradiation of Au particles loaded on semiconductor AgCl powders, synthesized by a deposition-precipitation (DP) method [Au(DP)/AgCl], efficiently generated HClO in Cl⁻ solutions under air flow.⁴ As shown in Scheme 1A(a), visible light activation of the localized surface plasmon resonance (LSPR) of Au particles produces the hot electrons (e_{hot}⁻) and hot holes (h_{hot}^{+}) . As shown in (b), the e_{hot}^{-} is injected into the AgCl CB and reduces O_2 (eqn (2)). The h_{hot}^+ oxidizes the lattice Cl^- of AgCl (Cl_{L}) in the periphery of Au particles and generates Cl_{2} , which is disproportionated to afford HClO (eqn (4)). As shown in Scheme 1B, the Cl_L⁻ oxidation occurs at ca. +0.5 V [vs. reversible hydrogen electrode (RHE)], which is more negative than that for bulk Cl⁻ oxidation (1.36 V), owing to the strong Ag-Cl orbital hybridization. This results in efficient HClO generation under visible light. As shown in Scheme 1A(c), the eliminated Cl_L⁻ is compensated from the solution. The Cl_L⁻ oxidation/compensation cycle on the Au(DP)/ AgCl catalysts generated 38 ppm of HClO, which was 10-fold higher than that generated by earlier reported systems.6-9

The next challenge is the enhancement of catalytic activity. The rate-determining step is the Cl_L^- oxidation on the peripheral sites at the Au-AgCl interface (Scheme 1A(c)). We hypothesized that creating a large number of small Au particles on AgCl would

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Scheme 1 Proposed mechanism for (A) HClO generation on plasmonic Au/AgCl catalysts in Cl⁻ solutions with O₂, which includes (a) LSPR activation, (b) oxidation/reduction, and (c) compensation of Cl_L^- . (B) Band structures at the Au–AgCl interface.

increase the number of peripheral sites, which may increase the number of Cl_L^- that can be oxidized by e_{hot}^- photogenerated on the Au particles. The DP method used for deposition of Au particles in our previous work⁴ involves the adsorption of Au precursors (HAuCl₄·4H₂O) onto AgCl and subsequent calcination at a high temperature (673 K).¹⁰ In this case, Au(DP)/AgCl with 2.7 wt% Au showed the highest activity. However, higher Au loadings created larger Au particles owing to their migration and coalescence.¹¹ This decreases the number of peripheral Cl_L^- and showed lower activity.⁴ Therefore, a method to create a large number of small Au particles is necessary.

Microwave (MW)-assisted reduction is a simple method for the deposition of metal particles. This facilitates the deposition of several kinds of metal particles (Au,¹² Pt,¹³ Pd¹⁴) onto several inorganic/organic supports by a relatively short time (~30 min) MW irradiation of a mixture containing metal precursors, supports, and reducing reagents such as ethylene glycol at low temperatures (373–453 K). In the present work, we deposited Au particles on AgCl by the MW method using ethylene glycol as a reducing reagent. The obtained Au(MW)/AgCl powders contained a large number of small Au particles. This increases the number of peripheral Cl_L^- and exhibits enhanced activity.

AgCl powder was prepared by a precipitation method using AgNO₃ and KCl in water.¹⁵ The AgCl powder was stirred in ethylene glycol with different amounts of HAuCl₄·4H₂O. MWs were irradiated onto the mixture for 17 min at 433 K under magnetic stirring. The resultant product was recovered by centrifugation, washed with water, and dried *in vacuo*, yielding Au(MW)_x/AgCl powders, where *x* denotes the amount of Au loaded, which is determined by X-ray fluorescence (XRF) analysis [*x* (wt%) = Au/AgCl × 100]. The powder X-ray diffraction (XRD) patterns of Au(MW)_x/AgCl (Fig. S1, ESI†) showed a peak at 38° assigned to the {111} diffraction peak of metallic Au (JCPDS 04-0784) as is the case for Au(DP)_x/AgCl,⁴ indicating the formation of Au⁰. They also showed peaks assigned to cubic



Fig. 1 Amount of HClO generated on the respective catalysts during 24 h of photoirradiation in (red) water and (blue) 550 mM NaCl solution. Reaction conditions: solution (50 mL), catalyst (0.1 g), air flow (1.0 L min⁻¹), $\lambda > 420$ nm (Xe lamp), and temperature (303 K).

AgCl (JCPDS 31-1238) as is the case for pristine AgCl and $Au(DP)_x/AgCl$, indicating that the AgCl structure was maintained even after the MW irradiation.

Photoreactions were performed in 550 mM NaCl solutions whose Cl⁻ concentration was similar to that of seawater.⁴ A NaCl solution (50 mL) containing catalyst (0.1 g) was photoirradiated under visible light ($\lambda > 420$ nm) using a Xe lamp under air flow (1.0 L min⁻¹) with magnetic stirring at 303 K, where the emission spectrum of the lamp is shown in Fig. S2 (ESI†). Fig. 1 (blue bars) shows the amount of HClO generated on the respective catalysts during 24 h of photo-irradiation. In the case of Au(DP)_x/AgCl, Au(DP)_{2.7}/AgCl showed the highest activity, and further Au loading decreased the activity.⁴ In contrast, in the case of Au(MW)_x/AgCl, Au(MW)_{4.9}/AgCl and Au(MW)_{6.1}/AgCl showed the highest activity with the amount of HClO generated being about twice those generated on Au(DP)_{2.7}/AgCl. In addition, Au(MW)_{2.7}/AgCl also showed higher activity than Au(DP)_{2.7}/AgCl. This indicates that the MW method produces more active Au/AgCl catalysts than the DP method.

The action spectrum for HClO generation on Au(MW)4.9/AgCl (Fig. S3, ESI⁺) agrees with the Au LSPR band, as is the case for $Au(DP)_{2.7}/AgCl_{4}^{4}$ confirming that the Au LSPR activation by visible light triggers the reaction (Scheme 1A(a)). The apparent quantum yields (Φ_{AOY}) of Au(MW)_{4.9}/AgCl are higher than those of Au(DP)_{2.7}/ AgCl at the entire wavelengths, suggesting that the h_{hot}^+ and $e_{hot}^$ generated on Au(MW)4 9/AgCl are consumed more efficiently. Cyclic voltammetry (CV) was performed using the catalyst-loaded fluorinated tin oxide (FTO) electrodes to confirm the Cl_L⁻ oxidation/ compensation cycle (Scheme 1A(b) and (c)). The CV of $Au(MW)_{4.9}$ AgCl measured in NaCl solution under O2 (Fig. S4, ESI⁺) showed the anodic current for Cl_{L}^{-} oxidation at >0.6 V (vs. RHE) and the cathodic current for O2 reduction at <0.2 V. These currents appeared reversibly, as was the case for Au(DP)2.7/AgCl.4 In contrast, the absence of Cl⁻ in solution showed a weak current for Cl₁⁻ oxidation, which disappeared after the second cycles. The results suggest that the catalyst photoexcited in Cl- solution generates HClO via the Au LSPR activation by visible light and Cl_L⁻ oxidation/compensation cycle (Scheme 1A). The electrochemical impedance spectroscopy (EIS) Nyquist plots of Au(MW)4.9/AgCl⁻ and Au(DP)2.7/AgCl⁻ loaded FTO

measured in NaCl solution in the dark (Fig. S5, ESI[†]) showed similar charge transfer resistance ($R_{\rm CT}$). Visible light irradiation decreased the $R_{\rm CT}$ of both catalysts with Au(MW)_{4.9}/AgCl showing lower $R_{\rm CT}$. This indicates that the $h_{\rm hot}^+$ photogenerated on Au(MW)_{4.9}/AgCl is consumed more efficiently by oxidation of Cl_L⁻: the efficient consumption of $h_{\rm hot}^+$ by Cl_L⁻ results in higher photocatalytic activity of Au(MW)/AgCl (Fig. 1). It is noted that Au(MW)_{4.9}/AgCl, when reused for photoreactions, showed activity similar to that of the fresh catalyst (Fig. S6, ESI[†]), suggesting that the catalyst is reusable without significant loss of activity.

The higher photocatalytic activity of Au(MW)/AgCl originates from the formation of smaller Au particles. Fig. 2a shows the results of scanning transmission electron microscopy (STEM) observation of Au(MW)4.9/AgCl. The energy-dispersive X-ray spectroscopy (EDS) elemental maps indicate that Au particles are dispersed on the AgCl surface, as is the case for $Au(MW)_{1,3}$ AgCl (Fig. S7, ESI[†]), Au(DP)_{2.7}/AgCl (Fig. S8, ESI[†]), and Au(DP)_{4.5}/ AgCl (Fig. S9, ESI⁺). The size distributions of the Au particles (Fig. 2b) show that Au(DP)_{4.5}/AgCl contains Au particles with an average diameter of 50 nm, which is larger than that of $Au(DP)_{2.7}$ AgCl (41 nm). This indicates that, in the DP method, higher Au loadings create larger Au particles due to the migration and coalescence of the particles by high temperature calcination (673 K).^{4,11} In contrast, Au(MW)_{4.9}/AgCl (Fig. 2c) contains smaller Au particles (36 nm), which are similar to those of the lower-Auloading Au(MW)1 3/AgCl catalyst (38 nm). This indicates that the MW method creates smaller Au particles even with higher Au loadings. This may be because rapid heating by MW irradiation creates numerous nuclei¹⁶ and low temperature heating (433 K) suppresses their migration and coalescence.



Fig. 2 (a) STEM-EDS results of Au(MW)_{4.9}/AgCl. Size distributions of Au particles on (b) Au(DP)_x/AgCl and (c) Au(MW)_x/AgCl catalysts.

The smaller Au particles created by the MW method increase the number of peripheral sites (Scheme 1A(c)), which efficiently promotes the oxidation of Cl_L⁻ by h_{hot}⁺ photogenerated on Au particles, enhancing photocatalytic activity. The number of peripheral Cl_L⁻ can roughly be determined by the photoreaction in pure water without Cl⁻. In this case, Cl_{L}^{-} of AgCl oxidized by h_{hot}^{+} is not compensated from the solution; therefore, the amount of HClO generated reflects the number of peripheral Cl₁⁻ that can be oxidized by h_{hot}⁺ on Au particles. As shown in Fig. 1 (red bars), the amount of HClO generated on Au(MW)_x/AgCl is much larger than that generated on Au(DP)2.7/AgCl and increases with Au loading. These results agree with the photocatalytic activity (blue bars) and the EIS Nyquist plots (Fig. S5, ESI⁺). This suggests that the MW method produces small Au particles on the AgCl surface and creates a larger number of peripheral Cl_L⁻ that are active for the consumption of the h_{hot}⁺ photogenerated, thereby enhancing photocatalytic activity.

The amount of HClO generated on Au(MW)_x/AgCl (blue bars in Fig. 1) is saturated with \geq 4.9 wt% Au, although the number of peripheral Cl_L^- increases with ≥ 4.9 wt% Au (red bars). This is because of the decrease in the LSPR band of the Au particles. The diffuse-reflectance (DR) UV-vis spectra of Au(MW)x/AgCl (Fig. 3) indicate that the intensity of the LSPR band decreases with Au loading, together with a red shift of the band. This is ascribed to the decrease in the electron density of Au particles.¹¹ X-ray photoelectron spectroscopy (XPS) analysis (Fig. S10, ESI⁺) was used to clarify the electronic states of Au, Ag, and Cl. The Ag 3d (Fig. S11, ESI†) and Cl 2p (Fig. S12, ESI[†]) spectra of Au(DP)_{2.7}/AgCl and Au(MW)_x/AgCl are similar, indicating that the Ag and Cl components of these catalysts have similar electronic states. The Au 4f spectra of the catalysts (Fig. S13, ESI[†]) show Au⁰ components.¹⁷ The increase in the Au loading shifts the peaks to higher binding energy, confirming the decrease in the electron density of the Au particles. The Au loading leads to a transfer of AgCl e_{CB}^{-} to the Au particles to balance their Fermi levels, thereby increasing the electron density of the Au particles.18 However, loading of excess amounts of Au may lead to a charge dispersion on the Au particles.¹⁹ This decreases the electron density of the Au particles and weakens the LSPR band (Fig. 3). The weak LSPR band suppresses the activation by visible light and decreases the photocatalytic activity, although the number of peripheral Cl₁⁻ increases with Au loadings (red bars in Fig. 1). The results indicate that loading an appropriate amount of Au particles is necessary for high photocatalytic activity.



Fig. 3 Diffuse-reflectance UV-vis spectra of Au(MW)_x/AgCl.



Fig. 4 Change in the amount of HCIO generated and the SCC efficiency with time under simulated sunlight irradiation of the respective catalysts ($\lambda > 420$ nm). Conditions: 550 mM NaCl (50 mL), catalyst (0.1 g), air flow (1.0 L min⁻¹), and temperature (303 K).

Active Au(MW)/AgCl catalysts can be synthesized at relatively low temperatures (\geq 413 K). The catalysts synthesized by MW irradiation at \geq 413 K showed similar Au loadings and activities (Fig. S14, ESI†). In contrast, lower temperature synthesis created lower-Au-loading catalysts with lower activity: they showed weaker LSPR band (Fig. S15, ESI†) and Au{111} diffraction (Fig. S16, ESI†) because the reduction of Au³⁺ to Au⁰ was suppressed at lower temperatures.²⁰ Also note that a relatively short time MW irradiation (\geq 7 min) produces active catalysts. As shown in Fig. S17 (ESI†), the catalysts synthesized by different MW irradiation times (7–25 min) showed similar Au loadings and activities, where they showed similar LSPR absorption (Fig. S18, ESI†) and Au (111) diffraction (Fig. S19, ESI†). These results indicate that active Au(MW)/AgCl catalysts can be synthesized by short time MW irradiation (\geq 7 min) at relatively low temperatures (\geq 413 K).

The solar-to-chemical conversion (SCC) performance of Au(MW)₄ ₉/AgCl was evaluated under irradiation with AM1.5G simulated sunlight²¹ over the wavelength range of 420–2500 nm (Fig. S2, ESI⁺). Fig. 4 shows the change in the amount of HClO generated and the SCC efficiency with time. Au(MW)_{4.9}/AgCl (square) generates more HClO than than of Au(DP)2.7/AgCl (circle). The SCC efficiency for HClO generation on Au(MW)_{4.9}/AgCl ($\sim 0.05\%$) is higher than that on $Au(DP)_{2.7}/AgCl (\sim 0.03\%)^4$ and remains almost constant during the photoirradiation, indicating stable and efficient HClO generation on Au(MW)_{4.9}/AgCl. Note that the irradiation of UV region light (300-400 nm) is ineffective: as shown in Fig. S20 (ESI⁺), irradiation with entire wavelength light ($\lambda > 300$ nm) decreases the HClO amount and SCC efficiency because UV light promotes photolysis of HClO generated (eqn (5)). The results indicate that visible light irradiation is effective for HClO generation. The visible light irradiation of Au(MW)_{4.9}/AgCl for 24 h generated 63 ppm of HClO, which was higher than that obtained on Au(DP)_{2.7}/AgCl (38 ppm).⁴ This HClO concentration (63 ppm) exceeds the concentration that is recommended by the WHO for the disinfection of drinking water $(>3 \text{ ppm})^{22}$ and the concentration that was reported effective for the inactivation of several types of bacteria and viruses such as SARS-CoV-2 (60 ppm).²³ Furthermore, as shown in Fig. S21 (ESI⁺), the high oxidation capability of the HClO solution (63 ppm) obtained by Au(MW)_{4.9}/AgCl is confirmed by the bleaching tests using a black mold and a piece of denim pants.

In summary, we demonstrated that the MW method rapidly ($\geq 7 \text{ min}$) created a large number of small Au particles ($\sim 40 \text{ nm}$) on AgCl powders under mild temperature ($\geq 413 \text{ K}$) conditions. Visible light irradiation of the obtained Au(MW)/AgCl catalysts in aerated Cl⁻ solutions generated high-HClO-concentration solutions *via* the Cl_L⁻ oxidation/compensation cycles owing to the enlarged Au–AgCl periphery site that efficiently promotes Cl_L⁻ oxidation. The results based on the creation of small Au particles by MW irradiation may contribute to the clean and sustainable production of HClO using solar energy.

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Data availability

The data supporting this article have been included as part of the ESI.†

Conflicts of interest

There are no conflicts to declare.

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