Solar-to-hydrogen peroxide conversion of photocatalytic carbon dots with anthraquinone: Unveiling the dual role of surface functionalities

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Abstract
Solar-driven photocatalytic production of hydrogen peroxide (H\textsubscript{2}O\textsubscript{2}) requires only sunlight, oxygen, and water, making it a green and sustainable alternative to conventional H\textsubscript{2}O\textsubscript{2} production processes. We present photocatalytic carbon dots (CDs) as a new candidate for high-performance H\textsubscript{2}O\textsubscript{2} production. Owing to the generation of an excellent charge carrier and the presence of various oxygen-containing functional groups, CDs showed an outstanding H\textsubscript{2}O\textsubscript{2} production capability of 609.4 \textmu mol g\textsuperscript{-1} h\textsuperscript{-1} even in the absence of an electron donor, demonstrating promising self-electron-donating capabilities. Hydroxyl groups on their surface, in particular, serve a dual role as photocatalytic active sites and as electron and proton donors toward the oxygen reduction reaction (ORR). The photocatalytic activity of CDs was significantly improved to 1187.8 \textmu mol g\textsuperscript{-1} by functionalizing their surfaces with anthraquinone (AQ) as a co-catalyst; it promoted the charge carrier separation and electrochemically favors the two-electron pathway of ORR. These carbon-based metal-free nanohybrids that are a unique combination of CDs and AQ could offer insights into designing efficient photocatalysts for future solar-to-H\textsubscript{2}O\textsubscript{2} conversion systems.

Introduction
Hydrogen peroxide (H\textsubscript{2}O\textsubscript{2}) is widely used as a clean oxidant in various fields such as organic synthesis, disinfection, and bleaching as it leaves no toxic byproducts and generates only water and oxygen.\textsuperscript{1} As demand for sustainable energy sources grows, H\textsubscript{2}O\textsubscript{2} has recently received renewed attention as a promising alternative liquid fuel for fuel cell system owing to its higher energy density compared to hydrogen.\textsuperscript{2} The mass production of H\textsubscript{2}O\textsubscript{2} is currently based on the anthraquinone (AQ) process, which involves the hydrogenation and auto-oxidation reactions between AQ and anthrahydroquinone (AHQ).\textsuperscript{3} However, this method involves costly noble metal catalysts, toxic organic solvents, and hydrogen gas under high pressure and temperature conditions.

In this regard, the photocatalytic production of H\textsubscript{2}O\textsubscript{2} has emerged as a promising and environmentally friendly alternative to the conventional AQ process because H\textsubscript{2}O\textsubscript{2} can be produced from both the oxygen reduction reaction (ORR) and water oxidation by harnessing solar energy.\textsuperscript{4, 5} Considering the high overpotential required for water oxidation to produce H\textsubscript{2}O\textsubscript{2} (1.78 V vs. RHE) than that for oxygen evolution reaction (OER) (1.23 V vs. RHE), the majority of H\textsubscript{2}O\textsubscript{2} formation by photocatalysts is derived from the reductive pathway through the ORR.\textsuperscript{6} In general, the ORR is categorized by the number of electrons that participate in the reaction, which depends on the reaction kinetics and energy levels of the active materials.\textsuperscript{7} For the efficient photocatalytic production of H\textsubscript{2}O\textsubscript{2}, high selectivity for the two-electron pathway is thus critical to prevent side reactions that occur through the single-electron pathway and decompose the H\textsubscript{2}O\textsubscript{2} produced.\textsuperscript{8}

Recently, carbon nitride (C\textsubscript{3}N\textsubscript{4})-based photocatalysts have attracted attention for H\textsubscript{2}O\textsubscript{2} production because of their high selectivity for the two-electron pathway in the ORR.\textsuperscript{9} However, C\textsubscript{3}N\textsubscript{4}-based
photocatalysts suffer from limited visible-light absorption due to their large bandgap energy, high rate of charge carrier recombination, and inefficient water oxidation capability. To overcome these intrinsic limitations of C₃N₄, C₃N₄-based nanocomposites have been developed with various complementary supporters or co-catalysts such as pyromellitic diimide,¹⁰ polyoxometalates,¹¹ black phosphorus,¹² cobalt phosphide,¹³ carbon nanotube,¹⁴ reduced graphene oxide,¹⁵ and AQ.¹⁶

As an alternative to semiconducting quantum dots, carbon dots (CDs) possess interesting physical, optical, and chemical properties for various energy- and bio-applications¹⁷-²⁰ due to their aqueous solubility, abundant functional groups, excellent light absorption, and carrier generation. In this study, we report the design and a synthesis of new class of CDs as high-performance H₂O₂ photocatalysts (Fig. 1). The unique photoelectron donating capability of CDs with oxygen-bearing functionalities (e.g., hydroxyl, hydroquinone, and carboxyl moieties) allows them to present the outstanding photocatalytic performance toward H₂O₂ evolution, outperforming most of C₃N₄- and semiconducting nanoparticle-based photocatalysts in the absence of electron donors (i.e., alcohols). More importantly, by taking advantage of the unique surface chemistry of CDs, we functionalize their surfaces with AQ molecules (CD-AQ) to further enhance their photocatalytic performance. As a co-catalytic supporter, AQ serves to promote the two-electron reaction for H₂O₂ production. We envision that this unique combination of CD coupled with AQ will provide new insights into the design of metal-free carbon-based photocatalytic nanohybrids.

**Results**

**Synthesis of CD and CD-AQ.** CDs were initially prepared via a hydrothermal reaction using citric acid (CA) as a carbon source in the presence of ethylene diamine (EDA). EDA acts as both a nitrogen source and a surface-passivating agent in this reaction (Fig. 1a). The as-synthesized CDs showed high aqueous stability with a ζ-potential of +5.2 mV at pH 7 owing to the presence of functional amine groups on the CD surface derived from EDA. In order to introduce AQ molecules as a co-catalyst, we employed N-ethyl-N’-(3-dimethylaminopropyl)carbodiimide methiodide (EDC) to mediate the surface functionalization of CDs through the reaction between amines and carboxylic acid-containing AQ molecules. Considering the solubility of all reactants, ethanol was used as a suitable solvent for the functionalization.

High-resolution transmission electron microscopy (HR-TEM) analysis revealed the nearly spherical morphology of CD and CD-AQ with an average diameter of 4.5 ± 0.8 nm and 9.9 ± 2.3 nm (Fig. 1b and Fig. S1 in the Supplementary Information). The bandgap energy (E₉) and energy levels of the conduction band (CB) and valence band (VB) of CD and CD-AQ were respectively determined by using Tauc’s plot from UV/vis spectra and ultraviolet photoelectron spectroscopy (UPS) (Fig. S2 and S3). Interestingly, there was no significant difference in the bandgap energies (3.2–3.3 eV) or in the energy level of CB and VB regardless of the types of CDs used.

Successful functionalization by AQ was also monitored by UV/vis and photoluminescence (PL) spectroscopy (Fig. 1c and 1d). The CDs exhibited characteristic absorption peaks at 237 nm (π–π*
transition of sp²-carbon network) and 341 nm (n−π* transition of carbonyl groups), confirming the presence of various surface functional groups. The AQ molecules used for the functionalization displayed distinct absorbance at 258 nm and 335 nm owing to the π−π* transition of benzoid and quinoid. The spectra of the CD-AQ displayed the presence of absorption peaks of both CD and AQ molecules, indicating successful surface functionalization. The characteristic PL spectra of both CD and CD-AQ were found to depend on the excitation wavelength (Fig. S4). The maximum emission of CD was observed at 429 nm upon excitation at 335 nm, while that of CD-AQ was located at 434 nm upon excitation at 345 nm. In addition to shorter PL lifetime of CD-AQ than CD obtained by time-correlated single photon counting (TCSPC) measurements (Fig. S5), the PL intensity of CD-AQ was decreased by 87% upon conjugation with AQ on the surface of CD (Fig. 1d), indicating the efficient transfer of the photoexcited charge carriers from CD to AQ.

The successful functionalization of CD-AQ was further investigated by Fourier transform infrared (FT-IR) spectroscopy and X-ray photoelectron spectroscopy (XPS) (Fig. 2). In the FT-IR spectra of both CD and AQ molecules, a peak at 1700 cm⁻¹ was observed corresponding to the stretching vibration of C=O (Fig. 2a). However, the C=O stretching vibration peak of CD-AQ shifted slightly to 1656 cm⁻¹, indicating the successful amide coupling between CD and AQ. In addition, a new peak attributed to the amide bond was also observed in 3300 to 3480 cm⁻¹ in the XPS spectrum of the CD-AQ. In the C 1s XPS spectra, the peak intensity of the oxygen-containing groups at a higher binding energy such as C-OH, C=O, and COOH decreased, while the peaks at 284.2 and 285.6 eV assigned to the C=C and C-N, respectively, increased after functionalization of AQ on CD owing to the formation of an amide bond. However, a simple mixture of CD and AQ (in the absence of the EDC agent) did not indicate the formation of an amide group, suggesting a critical role of EDC in the coupling reaction of CD-AQ (Fig. 2b). In addition, a new characteristic N 1s peak at 402 eV was observed for the CD-AQ, which was assigned to the amide bond without noticeable changes in the other N-configurations of graphitic-, pyridinic-, and pyrrolic-N (Fig. 2c). The peak shift of N1s of CD-AQ to a lower energy compared to pristine CD and CD+AQ (mix) is also attributed to the electron delocalization by amide coupling with AQ molecules, which can also offer a strong evidence for successful functionalization of AQ on CDs. Taken together, the successful chemical functionalization of AQ on CD to afford a CD-AQ was confirmed. Meanwhile, the relative mass ratio of AQ to CD was determined to be 1.7 in the CD-AQ nanocomposite based on UV/vis spectroscopy (Fig. S6).

**Photocatalytic production of H₂O₂ with CDs.** The molar ratio of the CD precursors CA to EDA, for instance, significantly affects the light absorption and photocatalytic properties of CDs. Thus, before investigating the photocatalytic performance of as-synthesized CD-AQ, we first evaluated the performance of CDs while varying the molar ratio of their precursors. All CDs were successfully synthesized, showing a characteristic n−π* transition peak of carbonyl groups at 341 nm (Fig. S7). Under solar-simulated light illumination, H₂O₂ was produced with an initial production rate of 148 (CA:EDA ratio of 5:2), 423 (5:5), and 89 μmol g⁻¹ h⁻¹ (5:7) for CDs prepared with a different molar ratio of CA to EDA, respectively (Fig. 3a). Interestingly, the highest photocatalytic performance was observed for CD with an equimolar ratio of CA to EDA (Fig. S8). This can be accounted for by considering the high degree of
carbonization with a higher fraction of C=C bond in reference to C-N in C 1s XPS spectra than those of CDs prepared from other precursor ratios (Fig. S9), resulting in enhanced light absorption under a wide range of wavelengths. Hereafter, CD prepared at a ratio of 5:5 was selected for further experiments owing to its high photocatalytic H$_2$O$_2$ production.

In the absence of light irradiation and O$_2$ source, CD showed a negligible photocatalytic performance for H$_2$O$_2$ generation, indicating that the reaction required to produce H$_2$O$_2$ is a photoresponsive catalytic reaction through the ORR and consumes O$_2$ (Fig. 3b and Fig. S10). We also conducted the photocatalytic H$_2$O$_2$ production on CD in the presence of 1 mM of CA and EDA to evaluate the effect of the residual precursors in as-prepared CDs because these precursors are also known to serve as electron donors in conventional H$_2$O$_2$ production processes. However, the amount of H$_2$O$_2$ generated over 2 h decreased notably in the presence of each and both precursors (i.e., CA and EDA), with corresponding performances of 699.8 μmol g$^{-1}$ (CA only), 711.2 μmol g$^{-1}$ (EDA only), and 177.8 μmol g$^{-1}$ (both CA and EDA), respectively, compared to 845 μmol g$^{-1}$ for the bare CDs. The results indicate that the presence of residual CA and EDA interferes with the photocatalytic performance of CD instead of accelerating H$_2$O$_2$ production by acting as electron donors. It is inferred that CA and EDA are adsorbed on CDs by interacting with functional groups may interrupt surface reactions (e.g., substrate adsorption, electron transfer from donors and to acceptors) by covering the active reaction sites on the surfaces of the CDs.

Most semiconductor photocatalysts cannot achieve the H$_2$O$_2$ production and water oxidation simultaneously in the absence of sacrificial electron donors due to insufficient oxidation power and inefficient charge carrier separation. Thus, much effort has been made towards developing photocatalysts that enable the efficient H$_2$O$_2$ production only from water and molecular oxygen by tuning their structural, chemical, and electrical properties. As one of the most efficient photocatalysts for H$_2$O$_2$ production owing to its unique chemical process through the formation of a superoxo radical and 1,4-endoperoxide species, C$_3$N$_4$ generated H$_2$O$_2$ at a rate of 606.7 μmol g$^{-1}$ h$^{-1}$. However, the CD prepared in this study demonstrated an even higher H$_2$O$_2$ product rate of 631.8 μmol g$^{-1}$ h$^{-1}$ in the presence of ethanol as a sacrificial electron donor. Most interestingly, there was negligible production of H$_2$O$_2$ in C$_3$N$_4$ (69.6 μmol g$^{-1}$ h$^{-1}$), in the absence of ethanol electron donor, while CD could efficiently produce H$_2$O$_2$ (609.4 μmol g$^{-1}$ h$^{-1}$) in the same conditions (Fig. 3c). The significant decrease in H$_2$O$_2$ production on CD in the presence of both CA and EDA supports the finding that these potential impurities merely act as retardants for H$_2$O$_2$ production, rather than as electron donors (Fig. 3d).

**Photocatalytic production of H$_2$O$_2$ with CD-AQ.** We further functionalized CDs with AQ molecules, which improved the photocatalytic effect of hybrid CD-AQ. In addition, we optimized the photocatalytic performance of CD-AQ via varying degrees of surface functionalization of AQ molecules on the surfaces of the former (Fig. S11). Surprisingly, the kinetics of H$_2$O$_2$ production improved significantly for highly functionalized CD-AQ (1187.8 μmol g$^{-1}$) compared to bare CD (845.4 μmol g$^{-1}$) under solar-simulated light
illumination for 2 h (Fig. 4a). Although AQ itself can produce H₂O₂ under UV illumination via an irreversible photochemical reaction, the photochemical production of H₂O₂ using bare AQ was merely 237.2 μmol g⁻¹ even at a much higher concentration of 0.5 g L⁻¹. This enhanced performance by functionalization with AQ agrees with our previous finding that the AQ accelerated the photocatalytic H₂O₂ production of C₃N₄ by acting as a co-catalyst. The combination of CD and AQ in the CD-AQ hybrid is unique and beneficial for efficient dual-photocatalytic process for H₂O₂ production, not only because it introduces other catalytic active sites, but also because it promotes efficient charge carrier separation, as demonstrated by the decreased PL intensity and lifetime of CD-AQ (Fig. 1d and Fig. S5) as well as the enhanced apparent quantum yield (AQY) of CDs upon functionalization with AQ; the AQY profile of CD-AQ analogously resembles its absorption spectrum and is more pronounced than that of CD regardless of the excitation wavelengths (Fig. S12).

The pH-dependent photocatalytic effect of CD-AQ for H₂O₂ production was investigated because the photocatalytic ORR is governed by the proton-coupled electron transfer (PCET) reaction as in the following reaction (1):

\[ \text{O}_2 + 2\text{H}^+ + 2\text{e}^- \rightarrow \text{H}_2\text{O}_2 \]  (1)

where the photoexcited electrons from the photocatalyst react with molecular oxygen and protons in the solution. As decreasing pH, therefore, H₂O₂ production rate of CD-AQ increased considerably, to 2512 μmol g⁻¹ h⁻¹ at pH 1, which was an approximately 4-fold increase compared to pH 6 (Fig. 4b and Fig. S13), outperforming other carbon-based photocatalysts for H₂O₂ production reported to date (Tables S1). This tendency clearly supports the fact that the formation of H₂O₂ on CD-AQ is based mainly on the photocatalytic ORR under solar-simulated light illuminations. Similar trends have been observed with other photocatalytic systems used for photocatalytic H₂O₂ production.

Long-term stability is another critical requirement in the development of efficient photocatalysts. We therefore performed a long-term performance and stability tests of CD-AQ. There was a continuous increase in H₂O₂ concentration for a 12 h reaction by CD-AQ up to 4,700 μmol g⁻¹ at pH 6 and 28,600 μmol g⁻¹ at pH 3, respectively (Fig. S14). The slight decrease in the H₂O₂ production rate was probably due to the increase of pH (e.g., from pH 6 to pH 7.26) due to the protons consumed during the photocatalytic formation of H₂O₂ through the PCET reaction, and in situ photodecomposition of produced H₂O₂ by the absorption of UV light. The decomposition of H₂O₂ by CD and CD-AQ was further examined because in situ produced H₂O₂ can be decomposed by both electrons from CB and holes from VB, which are generated during photocatalytic process. Unlike the gradual decrease of initial H₂O₂ concentration obtained with C₃N₄ for 2 h of reaction, a negligible decomposition (less than 1%) of H₂O₂ was observed with CD and CD-AQ (Fig. S15). Furthermore, there was no change in the specific absorbance of CD after the 2 h photocatalytic process. Collectively, these results indicate that the CD-
based photocatalysts are an ideal candidate for the photocatalytic generation of H$_2$O$_2$ owing to their outstanding performance for H$_2$O$_2$ evolution in electron donor-free systems as well as their superior photochemical stability with a limited sluggish decomposition of H$_2$O$_2$.

**Electrochemical properties of CD and CD-AQ.** Electrochemical analysis was conducted to verify the role of AQ molecules in enhancing the catalytic performance of CD-AQ nanocomposites. AQ exhibits distinct oxidation and reduction peaks and an extremely fast two-electron redox reaction with a redox potential that shifts depending on the functional groups on the AQ.$^{34}$ Therefore, we investigated the cyclic voltammetry (CV) of CD and CD-AQ in the potential range from -1.2 to 0.0 V vs. Ag/AgCl (Fig. S16). The CV curve of CD-AQ showed an anodic peak at -0.52 V and a cathodic peak at -0.60 V, corresponding to the redox peaks of AQ, whereas that of CD showed a rectangle shape associated with a non-Faradaic current (without distinct peaks). These results imply that AQ was successfully functionalized on the CDs, thus CD-AQ showed the higher current density than that of CD for ORR in O$_2$-saturated condition. As a result, the electrochemical kinetics of electrons transferred within CD-AQ nanocomposites can be affected by the functionalized AQ.

To evaluate the electrochemical kinetics of CD and CD-AQ toward the ORR, we used a rotating ring disk electrode (RRDE) to examine linear sweep voltammetry (LSV) (Fig. 5). The CD-AQ exhibited a ring and disk current density that was more than twice the current density of the CD (Fig. 5a and 5b). The average electron transfer number of CD was calculated as 2.4 and the average efficiency of H$_2$O$_2$ production was 78.4% in the potential ranges from -0.80 to -0.40 V (vs. Ag/AgCl), suggesting that CD is responsible for the two-electron ORR pathway (Fig. 5c). The two reduction plateaus observed in LSV curve can be a key evidence for electrochemical H$_2$O$_2$ production. The electrochemical ORR performance of CD is known to be governed through consecutive two-electron steps upon generation of H$_2$O$_2$ as an intermediate by slow kinetics on carbon surfaces containing chemical functional groups.$^{35}$ In addition, the imbalance in charge density also increases the oxygen adsorption on the surface of CD, thereby increasing oxygen solubility. It also participates in the release of H$_2$O$_2$ as an intermediate. Interestingly, CD-AQ showed not only highly enhanced electrochemical kinetics with a lower charge transfer resistance (R$_{ct}$) measured by electrochemical impedance spectroscopy (EIS) (Fig. S17 and Table S2), but also increased selectivity compared to CD alone, with an average electron transfer number of 2.3 and an average H$_2$O$_2$ selectivity of 80.6% as the rotating speed in the same potential (Fig. 5d). These results indicate that the chemical functionalization of AQ onto CD creates a more efficient catalyst for generating H$_2$O$_2$ by enhancing the facile charge transfer and facilitating the selective two-electron pathway of the ORR.

**Mechanism of the CD-AQ.** The structural advantages of versatile CD-based catalysts (containing a unique sp$^2$/sp$^3$ hybrid carbon structure with various oxygen- and nitrogen-functional groups) facilitate and promote various chemical reactions, which suggests their widespread utility in a number of applications.$^{19,36-39}$ In this regard, we monitored the time profile of specific chemical compositions of CD during the photocatalytic process to clarify the underlying mechanisms behind the outstanding performance of CD-AQ in the evolution of H$_2$O$_2$ (Fig. 6b and Fig. S18).
The deconvoluted high-resolution C 1s spectra collected from XPS clearly revealed that the oxidative conversion of surface functional groups of CD took place during photocatalytic \( \text{H}_2\text{O}_2 \) production in which the C-OH group was diminished considerably whereas the C=O group was increased at the same time. This result agrees with our previous finding that the specific transformation of surface functional groups on CD, from hydroquinone to benzoquinone, was involved in and promoted the formation of metal nanoparticles onto the surface of CD by reducing metal ions under UV illumination.\(^{20,40}\) Moreover, hydroxyl and hydroquinone groups are known to be very effective for the formation of \( \text{H}_2\text{O}_2 \) through ORR in the presence of molecular oxygen.\(^{16}\)

In order to further elucidate the proposed mechanism and active sites on the surface of CDs, we conducted density-functional theory (DFT) calculation to investigate the adsorption energy and free energy of carbon surface like graphene nanoribbon as a model of CD and AQ-anchored carbon surfaces. In general, the two-electron ORR pathway follows two-step reactions for adsorption and desorption of oxygen with the formation of intermediate *OOH as in step 1 and 2:

\[
\begin{align*}
\text{O}_2 + \text{H}_2\text{O} + e^- + * & \rightarrow *\text{OOH} + \text{OH}^- \quad \text{(step 1)} \\
*\text{OOH} + e^- & \rightarrow \text{HO}_2^- + * \quad \text{(step 2)}
\end{align*}
\]

where * is an active site and *OOH is a key intermediate that plays a pivotal role in overall two-electron ORR performance. As a result, the adsorption energy of *OOH (\( \Delta G_{\text{OOH}} \)) can be a descriptor with corresponding the thermodynamic limiting potential (\( U^\text{L} \)) for determining the active sites. An ideal catalyst for \( \text{H}_2\text{O}_2 \) production has a \( \Delta G_{\text{OOH}} \) of 4.22 eV, which provides the highest activity with the thermodynamic equilibrium potential (\( U^0 = 0.70 \text{ V} \)) in the activity volcano plot. Thus, we calculated the energy barrier in terms of free energy (Fig. 6c and S19) and adsorption energy (Fig. 6d) of CD and CD-AQ. The corresponding atomic structures of the examined CD and CD-AQ are shown in Fig. 6e. The ORR on AQ showed a thermodynamically uphill reaction by high free energy of 0.728 eV, resulting in low adsorption of OOH on AQ anchored carbon matrix. Although AQ method is commonly used in industrial \( \text{H}_2\text{O}_2 \) production, it is because AQ mechanism is not a competitive pathway for carbon-based catalysts and AQ follows different catalytic mechanism reported in other studies using various quinone-enriched molecules with similar DFT results of AQ molecules.\(^{41}\) On the other hand, the oxidized CD surface (herein a form of CD-ketone for DFT calculation) exhibited the highest activity compared to bare CD and other carbon matrix, which suggests a plenty of oxygen functional groups on the surface of CD can serve as thermodynamically active sites in accord with experimental XPS results.

From these results, we propose the possible mechanisms that retain a number of elementary reaction steps occurring on the surface of both CD and CD-AQ systems to form \( \text{H}_2\text{O}_2 \) (Fig. 6a). Specifically, CD initially generates the photoexcited electrons and holes owing to its appropriate bandgap energy. The photoexcited electrons are then transferred to the active sites (hydroxyl groups) on the surface of CDs, resulting in the reduction of \( \text{O}_2 \) that produces \( \text{H}_2\text{O}_2 \) with the surrounding protons. This step is followed by
the conversion of the hydroxyl groups to carbonyl groups on the surface of CDs. These protons can not only be supplied by the dissociation of nearby carboxylic acid groups; they can also be generated by hydroxyl groups from the CDs themselves in the absence of electron- and proton donors such as ethanol (Fig. S20). This interpretation is also based on the poor performance of C$_3$N$_4$ in the absence of ethanol due to the lack of surface functional groups (Fig. 3c).

Considering the results from the reduced PL and electrochemical analysis, the role of AQ in CD-AQ for H$_2$O$_2$ production was proven to be that of a co-catalyst that promotes charge carrier separation and selectivity for H$_2$O$_2$ formation through the sequential hydrogenation-dehydrogenation of AQ. In CD-AQ, photoexcited electrons are transferred efficiently from CD to AQ molecules. O$_2$ is then reduced by a reversible reaction between AQ and AHQ with electrons and protons donated from the CDs or ethanol.

**Discussion**

We developed highly efficient CD-based metal-free photocatalysts towards the environmentally friendly production of H$_2$O$_2$. The CDs and CD-AQ nanocomposites exhibited a superior photocatalytic performance, and physical and chemical stability compared to C$_3$N$_4$-based photocatalysts as, with H$_2$O$_2$ production gradually increasing during the long-term light irradiation over 12 h. The AQ molecules were successfully functionalized on the surface of CDs, where they act as co-catalysts that promote efficient charge carrier separation from the photoexcited CDs and accelerate the two-electron ORR pathway. This results in significantly enhanced electron transfer kinetics for H$_2$O$_2$ production. The photocatalytic mechanism was elucidated by examining the change in the chemical compositions of the surface functional groups of CDs; a reversible reaction occurred between the hydroxyl and carbonyl groups during photocatalytic production of H$_2$O$_2$ under light illumination. Notably, CDs showed a considerable H$_2$O$_2$ production rate without the addition of any extra electron donors owing to their unique self-electron donating capabilities. We anticipate that this new photosystem, which boosts photocatalytic effects by combining two or more components, can be extended further to design other multi-functional hybrid photocatalysts that facilitate selective electron transfer and wide visible light absorption as a step towards highly efficient solar-to-fuel conversion systems.

**Methods**

**Materials**

Citric acid (CA), ethylenediamine (99%, EDA), anthraquinone-2-carboxylic acid (AQ), N,N-Diethyl-1.4-phenylenediamine sulfate (DPD), peroxidase, from horseradish (POD, type VI-A), and N-ethyl-N’-(3-dimethylaminopropyl)carbodiimide methiodide (EDC) were purchased from Sigma-Aldrich and used without further purification.

**Synthesis of carbon dots (CDs)**
The synthesis of the CDs involves a hydrothermal reaction of a mixture of CA and EDA in deionized (DI) water. Specifically, 960 mg of CA (5.0 mmol) and 347 μL of EDA (5.0 mmol) were dissolved in 10 mL of DI water. For comparison with 5:2 and 5:7 CDs, 2.0 mmol and 7.0 mmol of EDA were used, respectively. The solution was placed into a muffle furnace (MF-22G, JEIO TECH) and heated at 180 °C for 6 h. Thereafter, the obtained brownish suspension was dialyzed (SpectraPore MWCO 500–1,000) in DI water over 2 weeks to remove salts, unreacted chemicals, and much smaller CDs. Note that a thorough dialysis process is critical to overall performance for H₂O₂ production of CDs as shown in Fig 3. After dialysis process, the aqueous CD solution was evaporated by a rotary evaporator and the resulting CD powder was re-dispersed by DI water at a desired concentration (10 mg mL⁻¹).

**Synthesis of functionalized CDs with anthraquinone (CD-AQ)**

A suspension of 4 mL of the as-prepared CDs (conc. 10 mg mL⁻¹) was reacted with the excess amount of 100 mL of AQ solution dissolved in ethanol (2 mM) for 12 h in the presence of 500 mg of EDC. The resulting suspension was dialyzed (SpectraPore MWCO 500 – 1,000) in ethanol over 2 weeks to remove unreacted EDC and AQ.

**Characterization**

A UV/vis spectrophotometer (UV-2550, Shimadzu) was used to measure the absorbance. The photoluminescence (PL) emission was determined with a Varian Cary 5000 spectrophotometer. Ultraviolet Photoelectron Spectroscopy (UPS) (AXIS-NOVA, Kratos Inc.) was used to measure the energy level. The chemical structure was analyzed by X-ray photoelectron spectroscopy (XPS) (K-alpha, Thermo Fisher) and Fourier-transform infrared spectroscopy (FT-IR) (Varian, Cray 660). The morphology and size of the CDs and CD-AQ were measured using Transmission electron microscopy (TEM) (JEOL, JEM-F200).

**Photocatalytic H₂O₂ production**

Photocatalytic H₂O₂ production experiments were carried out by irradiating simulated solar light (AM 1.5, 100 mW cm⁻²) from a 150-W Xenon Arc lamp to the aqueous suspension of various CDs in the presence or absence of 10 vol% ethanol as an electron donor.
suspension was continuously purged with O$_2$ for 15 min before and during illumination. The incident light intensity was determined using a Newport calibrated Si solar cell. Briefly, a sample aliquot was collected from the reaction suspension using a syringe. A DPD colorimetric method was also used to measure the quantitative H$_2$O$_2$ production (at 551 nm, $\varepsilon = 21000$ M$^{-1}$ cm$^{-1}$, the detection limit in the range of 0.20 - 0.30 µg/L).\textsuperscript{42} For all photocatalytic tests, multiple experiments were conducted independently.

**Electrochemical analysis**

Electrochemical experiments were carried out in a standard three-electrode cell system connected to a computer controlled potentiostat (Autolab, PGSTAT302N). The reactor contained a glassy carbon electrode (GCE), a platinum wire, and a Ag/AgCl/KCl (sat) electrode as a working, counter, and reference electrode, respectively, with an aqueous solution of 0.1 M KOH (pH = 13) as an electrolyte. Catalyst inks were prepared by dispersing a catalyst in 107 µL of DI water and 93 µL of ethanol followed by sonication process for 15 min. The 75 µL of as-prepared catalyst ink (421.5 mg L$^{-1}$) was drop-casted on a rotating ring–disk electrode (RRDE, glassy carbon disk and Pt ring, AG), and dried at room temperature overnight. For good adhesion of catalyst on the electrode surface, 5 µL of 5 wt.% Nafion solution (Sigma-Aldrich) and 2.5 µL of a solution containing 1.2 mL of 5 wt.% Nafion solution and 10 mL of acetonitrile (Sigma-Aldrich) were covered successively and dried at 70 °C for 6 h. The loading density of catalyst on the electrode was about 161 µg·cm$^{-2}$. Cyclic voltammetry (CV) was performed at 10 mV s$^{-1}$ with a potential range between -1.2 V and 0.0 V (vs. Ag/AgCl), with a GCE having a geometric area of 7.07 mm$^2$. The rotating disk electrode (RDE) and rotating ring disk electrode (RRDE) analyses were carried out by sweeping the disk potential from -1.2 V to 0.0 V vs. Ag/AgCl at 10 mV s$^{-1}$ with different rotation rates ranging from 400 to 2500 rpm, while holding the Pt ring at 0.5 V vs. Ag/AgCl to oxidize the *in situ* produced H$_2$O$_2$ formed on the disc electrode (for only RRDE), in an O$_2$-saturated electrolyte. The number of transferred electrons and H$_2$O$_2$ yields and were calculated using the following equation:
\[ n = 4 \times \frac{I_d}{I_d + I_r} \]  
\[ \text{H}_2\text{O}_2(\%) = \frac{200 \times I_r}{I_d + I_r} \]

where \( I_d \) is the disk current, \( I_r \) is the ring current, and \( N \) is the collection efficiency (24.9%).

**Computational calculation**

Spin-polarized density functional theory (DFT) calculations were performed by using Vienna Ab-initio Simulation Package (VASP) code.\(^{43}\) The plane wave cutoff energy of 400 eV was adopted within the PBE parameterization of GGA exchange-correlation functional.\(^{44}\) All structures were fully optimized in the 1x1x1 k-mesh with the van der Waals correction by Grunich's D3 method\(^ {45}\) until all the atomic forces are less than 0.01 eV/Å. For the nitrogenated and oxygenated edge carbons, the graphene nanoribbon model was adopted with 20 Å and 15 Å vacuum regions along the transverse and vertical directions, respectively. For Gibbs free energy calculations, we used \( \Delta G = \Delta E^{\text{DFT}} + \Delta Z\text{PE} - T\Delta S + \Delta G_U + \Delta G_{pH} \) equation.\(^ {46,47}\) \( \Delta E^{\text{DFT}} \) refers to the adsorption energy difference between intermediates by DFT. \( \Delta Z\text{PE} \) and \( T\Delta S \), the zero-point energy and vibrational entropy, were calculated by using vibrational frequencies of intermediate (\(^ *\)OOH), where we assumed \( T = 298.15 \) K. \( \Delta G_U = -eU \) denotes the correction by applied electrode potential, and \( \Delta G_{pH} = -k_B T \ln[H^+] \) the correction by solution’s pH.

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**Declarations**

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**Author contributions**

M.G and D.-Y.L contributed equally to experimental work and conducted data analysis. D.K. assisted with the synthesis of CDs. J.M. and G. L. conducted DFT calculation. H.-i.C assisted the electrochemical analysis. B.-S.K and H.-i.K coordinated and supervised the overall project. M.G., D.-Y.L, B.-S.K., and H.-i.K cowrote the manuscript. All authors discussed the results and participated in manuscript preparation.
Competing interests

The authors declare no competing interests.

Additional information

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