

# **Regulating the absorption spectrum of polydopamine**

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Synthesis and characterization of TEMPO-doped PDA. (A) Schematic illustration of the TEMPO-doped PDA with narrower bandgap and improved light absorption ability compared to conventional PDA. (B) Polymerization of dopamine and TEMPO, together with their molecular structures and powder photographs. (C) SEM image of PDA-3. (D) EELS mapping analysis of PDA-3 (Scale bars, 100 nm). (E) XPS survey spectra of PDA-i (i = 0 to 3). a.u.,



arbitrary units. (F) C 1s peaks, (G) N 1s peaks, and (H) O 1s peaks in XPS spectra of PDA-3. Credit: Science Advances, doi: 10.1126/sciadv.abb4696

Polydopamine (PDA) is an advanced functional material and its emergent light absorption properties make it crucial for applications in materials science. However, it is challenging to rationally design and regulate PDA absorption properties due to its complex architecture. In a new report, Yuan Zou and a team of researchers in polymer science, optoelectronic materials and physical chemistry in China proposed a simple method to regulate the light absorption behaviors of PDA. To accomplish this, they constructed donor-acceptor pairs in the microstructures via connections between specific chemical moieties. They then used detailed structural and spectral analysis as well as density functional theory (DFT) simulations to confirm the existence of such donor-acceptor molecular pairs. The molecular pairs could decrease the energy bandgap (or energy gap where no electrons exist) and increase electron delocalization to enhance light absorption across a broad spectrum. The rational design of PDA nanoparticles with tunable absorption properties allowed an improved photothermal effect, which the team demonstrated with excellent performances during solar desalination. The work is now published on *Science Advances*.

### Polydopamine

Inspired by the bio-macromolecular pigments of melanin, polydopamine (PDA) has received increasing attention for applications in surface engineering, photothermal therapy and bioimaging. The strong adhesive and light absorption properties of PDA can also benefit interfacial engineering during water remediation. Scientists have proposed many synthetic methods to prepare PDA nanomaterials, although with limited attention to regulate its absorption spectrum. The dopamine



polymerization process is composed of several complicated pathways and therefore not fully understood. As a result, Zou et al. assumed that the construction of highly conjugated structures relative to donoracceptor pairs in the PDA nanostructures could regulate the absorption spectrum of the sample. To accomplish that in this work, they developed a <u>one-pot synthesis strategy</u> to synthesize PDA nanoparticles (NPs) with tunable light absorption properties.

### Synthesis and characterization

During the synthetic process, they conducted direct copolymerization of 2.2,6,6-Tetramethylpiperidine-1-oxyl (TEMPO) – a typical nitroxyl radical, onto dopamine in an aqueous solution. They doped the TEMPO moiety to the polydopamine microstructure by covalently connecting the molecule with 5,6-dihydroxyindole (DHI) and Indole-5,6-quinone (IQ) oligomers to narrow the energy band gaps of the material and improve light absorption behaviors of conventional polydopamine nanoparticles (PDA NPs). The scientists confirmed the outcomes by using electrochemical analysis, density functional theory simulations and spectral measurements. The work demonstrated an outstanding photothermal efficiency for the product that can be used in interfacial solar steam generation and <u>seawater desalination</u>.





Proposed reaction pathways and intermediates formation during the polymerization of dopamine and TEMPO. (a) Proposed reaction pathways and mechanism during the polymerization of dopamine and TEMPO. (b) ESI-MS spectrum of the crude product solution after 5 min reaction; (c) Proposed intermediate molecular structures assigned to main peaks in (b). Credit: Science Advances, doi: 10.1126/sciadv.abb4696

The scientists developed three types of PDA NPs (classified between 1 to 3) with different doping contents and similar particle sizes by tuning the initial concentration of TEMPO. They synthesized conventional PDA NPs through self-polymerization of dopamine in the presence of ammonium using a well-established method. They observed the resulting PDA sample characteristics using scanning electron microscopy, dynamic light scattering and Fourier transform infrared spectra (FTIR).



Using X-ray photoelectron spectroscopy (XPS), they confirmed the existence of carbon, nitrogen, and oxygen elements in all PDA samples highlighting the successful preparation of TEMPO-doped PDA NPs. Based on the results, Zou et al. hypothesized two possible pathways to form the cross-linked macromolecular structure.

## **Enhanced light absorption and photothermal behavior of TEMPO-doped PDAs.**

The team examined the light absorption capacity and total photothermal effect of these TEMPO-doped PDA NPs, where the product strongly absorbed light by catching and converting solar energy to heat energy efficiently with wide-ranging applications. During further tests, they dispersed PDA-3 in water at several concentrations for irradiation under lasers. When compared to many other outstanding <u>photothermal</u> <u>materials</u>, the TEMPO-doped PDA NPs showed better photothermal behaviors. For instance, Zou et al. noted how gold nanoparticles could suffer substantial loss of light absorbance after long-term irradiation due to structural destruction via the accompanying heat of the experimental conditions. The team contrastingly demonstrated how TEMPO-doped PDA NPs maintained enhanced light absorption capabilities with improved photothermal behaviors compared to conventional photothermal nanomaterials. The resulting material can serve as new generation photothermal agents to complete a variety of applications.





Enhanced light absorption and photothermal behavior of TEMPO-doped PDA. (A) Photographs of PDA aqueous solutions with concentration of 50 and 100  $\mu$ g ml–1. Photo credit: Yuan Zou, Sichuan University. (B) L\* values of different PDA aqueous solutions. (C) UV-vis-NIR spectra of PDA-i (i = 0 to 3) ranging from 300 to 1500 nm. (D) Temperature elevations of PDA-3 at different concentrations under 808-nm laser irradiation. (E) The photothermal response of PDA-i (i = 0 to 3) aqueous solutions (100  $\mu$ g ml–1) for 600 s with 808-nm laser irradiation, and then the laser was shut off. (F) Temperature curves of PDA-3 (100  $\mu$ g ml–1) under four on/off cycles and under 808-nm laser irradiation. The light intensity of 808-nm laser was 2.0 W cm–2. (G) Molar extinction coefficient,  $\Delta$ T, and total photothermal efficiency of PDA-i (i = 0 to 3). Photo credit: Yuan Zou, Sichuan University. Credit: Science Advances, doi: 10.1126/sciadv.abb4696



### Analyzing the structure and absorption property regulation of TEMPO-doped PDA

The scientists noted the spontaneous formation of donor-acceptor microstructures in the TEMPO-based PDA system due to chemical conjugation between TEMPO and DHI, IQ and their oligomers during the polymerization process. This reaction contributed to the lower energy bandgap and enhanced light absorption of the product. To verify this, they calculated the optical bandgap value of different PDA samples in their aqueous solution forms alongside their electrochemical cyclic voltammetry (CV) to investigate energy bandgaps of all samples. They established the highest occupied molecular orbital (HOMO) and lowest unoccupied molecular orbital (LUMO) using the CV measurements and established the TEMPO unit as a donor fragment. As the doping concentration of TEMPO increased, the IQ moiety proportion also gradually increased, resulting in better delocalization of electrons for improved light absorption. The team hypothesized an increase in free radicals during PDA synthesis via TEMPO doping, which they tested and verified using <u>electron paramagnetic resonance</u> (EPR) measurements. Since the exciton-induced absorption (EIA) spectra did not rely on the amount of TEMPO doped to form the compound, the team credited it broadly to the presence of excitons in its additional constituents (such as DHI, IQ).





The light energy conversion process within PDA. (A) EPR spectra of PDA-i (i = 0 to 3) with the same mass in the solid states. (B) Transient absorption kinetics traces for PDA-i (i = 0 to 3). (C) Transient absorption spectra of PDA-3 at indicated delay times. (D) The EIA kinetic trace of PDA-3. mOD, mean optical density. Credit: Science Advances, doi: 10.1126/sciadv.abb4696

#### **Applications of water desalination**

The excellent photothermal and light absorption properties of TEMPObased PDA made the material well-suited for applications in water steam generation and seawater desalination. Of the variety of samples, Zu et al.



selected PDA-3 as the most promising candidate to develop the evaporation device. To accomplish this, they deposited the PDA-3 aqueous solution onto a cellulose membrane as a hydrophilic light absorber and prevented direct contact with water by using a thermal insulation layer such as polystyrene. When Zou et al. exposed the experimental setup to solar irradiation, they purified water by collecting condensed water from the solar steam. The PDA-3-coated cellulose membrane showed improved light absorption compared to control samples. The construct absorbed a majority of solar energy in the UV and visible regions. To understand solar vapor generation and photothermal evaporation performance, they measured the weight loss of water during evaporation and considered the energy conversion efficiency as an important index. The results indicated the feasibility of the device for desalination alongside efficient and durable activity.





Water desalination experiment. (A) A schematic diagram of the solar steam evaporation device based on PDA-3. (B) Photograph of the CM and PDA-3–coated CM. (C) SEM cross-section image of the double-layered film structure. (D) UV-vis-NIR diffuse reflection spectra of the CM and PDA-3–coated CM in the wavelength range of 250 to 2500 nm. (E) IR image of PDA-3–based device under one sun for 15 min. (F) Time course of water evaporation performances of saline water, CM, and PDA-3–coated CM under one sun irradiation. (G) Solar steam efficiency and evaporation rate of saline water, CM, and PDA-3–coated CM. (H) Photograph of the water steam generated under solar illumination of four suns with the PDA-3–coated CM. (I) The ion concentration of the saline water and seawater obtained from Bohai Bay before and after desalination. The dashed lines refer to the standard for drinkable water from the World Health Organization (WHO) and U.S.



Environmental Protection Agency (EPA), respectively. (J) Evaporation cycle performance of solar desalination devices over 30 cycles, with each cycle sustained more than 1 hour. The inset showed the photograph of absorber after 30 cycles. (K) Evaporation rate among different PDA-based evaporators under one-sun illumination. PVDF, polyvinylidene difluoride. Photos credit: Yuan Zou, Sichuan University. Credit: Science Advances, doi: 10.1126/sciadv.abb4696

In this way, Yuan Zou and colleagues proposed a simple method to regulate the absorption spectrum of polydopamine (PDA) in a one-pot polymerization process in the presence of dopamine and TEMPO. The resulting nanoparticles had improved light absorption ability and photothermal effects when compared to conventional PDA nanomaterials due to the donor-acceptor structures in the PDA system. When they coated the resulting TEMPO-based PDA on the cellulose film, the construct acted as a sunlight absorber suited for water evaporation with a high efficiency of solar conversion and excellent rate of evaporation. The work will offer new opportunities for structural and functional PDA nanomaterials suited for light-harvesting applications.

**More information:** Yuan Zou et al. Regulating the absorption spectrum of polydopamine, *Science Advances* (2020). <u>DOI:</u> <u>10.1126/sciadv.abb4696</u>

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