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Triple optically modulated and enzymatically responsive organic afterglow materials for dynamic anti-counterfeiting†

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Luminescent anti-counterfeiting is one of the extensively used strategies to combat the growing serious global forgery issue in modern society. Although many luminescent security labels have been developed, the majority of them are static and can be readily cloned due to their deterministic decoding mechanism. Therefore, an advanced dynamic anti-counterfeiting strategy is urgently needed. Herein, a multiple optical and biological response based label is developed for the first-time using o-phenylenediamine doped cyanuric acid as the information carrier that shows artificial enzyme activity and triple optical responses, *i.e.*, fluorescence, thermally activated delayed fluorescence (TADF) and phosphorescence. The optical properties of the security label can be reversibly modulated with excitation-wavelength, temperature and time, respectively, and irreversibly modulated by enzymatic discoloration, making it difficult to duplicate. The fabricated organic information carrier shows multiple optical properties compared to the traditional TADF or phosphorescent materials with limited information encryption, making it more favorable for anti-counterfeiting. The visually verifiable afterglow label bearing multiple dynamic encrypted messages offers bright prospects for high-security-level anti-counterfeiting applications.

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Introduction

In recent years, anti-counterfeiting techniques have attracted tremendous attention from governments, business organizations, and customers due to the increasing number of fake and shoddy products. The counterfeiting business is widespread in the manufacturing of products ranging from counterfeit currency to daily necessities, to pharmaceuticals, luxuries, and high-tech devices. The related products from illegal channels have caused great economic losses. For example, as claimed by the Global Brand Counterfeiting Report 2018, the total global economic losses from counterfeiting reached \$1.2 trillion in 2017 and rocketed to \$1.82 trillion in 2020. In response to the severe situation, many security labels, such as

To address this issue, afterglow materials have been proposed and introduced in optical encryption as a promising alternative. These afterglow material-based labels possess merits of dynamic patterns and tunable color under irradiation, making them difficult to duplicate by counterfeiters. ^{32,33} Although a large number of afterglow materials have been fabricated, metal-free afterglow materials are particularly interesting in the field of anti-counterfeiting owing to the

topographical markers,6 luminescent labels,7-10 luminescent magnetic labels, 11-13 and plasmonic security taggants, 4,14,15 have been developed to protect valuable documents and brands. Among these, luminescence anti-counterfeiting is one of the commercialized techniques with benefits of relatively high photostability, multicolor, facile design, and easy to visualize by eye or read using an authentication tool such as a digital camera or a spectrometer. 16 A luminescent security label is usually produced by printing specific optical materials, such as carbon quantum dots, 17-19 metal-organic frameworks, 20-22 coordination polymers, 23-25 organic dyes, 26,27 and up-conversion nanoparticles, 28-30 as information carriers. Unfortunately, fluorescent security labels usually fall into the class of static and single-encrypted messages, which are at high-risk of being replicated, thus offering compromised protection for the products.³¹

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advantages of abundance and low price of raw materials, lowtoxicity and easy-modulation of luminescence properties. For instance, carbonyl functional group room temperature phosphorescence (RTP) materials have made remarkable achievements in advanced dynamic information encryption. 34,35 However, phosphorescent carbonyl compounds still suffer from obvious limitations, e.g., short lifetime, photobleaching, and poor water stability.36 Compared to the phosphorescent carbonyl compounds, pure organic afterglow materials with longer lifetime, higher stability and resource of eco-friendly material makes it possible to encode optical information on space or time scales.37-39 More encouragingly, a color-tunable dualmode afterglow label proposed by the Zhao group and the Xu group further improves the anti-counterfeiting performance of traditional labels. 40,41 Ren and Ou designed a time-dependent phosphorescent color material to create a 3D code for dynamic information encryption. 42 Huang et al. have reported a colorful phosphorescent material to create a 4D code for information encryption. 43 These examples show that the more information a material can carry, the higher will be its encoding capacity and level of security. Although significant progress in optical encryption based on dual-/multi-mode responses of the composite afterglow materials under various external stimuli (such as excitation light, 44-47 heat, 48 mechanical force, 49 chemical reagents, 50 etc.) has been made, achieving all these properties in a single material remains a major challenge. Inspired by the recent study using dopant-matrix collaboration, molecular doping may lead to the enhancement of phosphorescence of an organic material and prolongation of its phosphorescence lifetime.⁵¹ Cyanuric acid (CA) may be a great candidate as a host material for molecular doping to achieve water-resist afterglow materials with long lifetime because it forms crystals that have weak RTP and are chemically stable in water via intermolecular hydrogen bonds.^{52,53} Based on the above considerations, we speculated that if we introduce a suitable molecule into the skeleton of CA through hydrogen bonding, its phosphorescence emission and lifetime could be greatly enhanced in both air and aqueous environments. o-Phenylenediamine (oPD) is considered as a good doping molecule that carries two amino groups and a benzene ring, which can easily form hydrogen bonds and provide impurity energy levels, respectively. As a verification experiment, a novel oPD doped CA organic afterglow material with triple optical response and enzymatically responsive was synthesized.

Experimental

Materials and chemicals

All chemicals and solvents were commercially purchased and used as received without further purification. Potassium chloride (KCl), o-phenylenediamine (C₆H₈N₂), cyanuric acid (C₃H₃N₃O₃), lithium chloride (LiCl) and methyl sulfoxide-d₆ (C₂D₆OS) were purchased from Energy Chemical (China). Melamine (C₃H₆N₆), formic acid (CH₂O₂) and ployvinyl acetate (PVCA) were purchased from Sinopharm Chemical Reagent

Co., Ltd (China). 3,3',5,5'-Tetratmethylbenzidine (C₁₆H₂₂Cl₂N₂) was purchased from Mackmin (China).

Preparation of oPD Doped g-C₃N₄ supramolecule

Melamine (2.523 g, 20 mmol) and o-phenylenediamine (0.06 g, 0.5 mmol) were fully ground with potassium chloride (2.7 g, 63.7 mmol) and lithium chloride (3.3 g, 63.7 mmol) in a mortar. The mixed precursors were transferred into a porcelain boat and heated to 350 °C using a muffle furnace with a heating rate of 5 °C min⁻¹ and kept at this temperature for 4 hours. The whole process was protected by a N2 atmosphere (flow rate 50 mL min⁻¹). Finally, yellow powders were obtained, named PCN.

Preparation of oPD-CA

0.5 g of PCN was dissolved in 10 mL formic acid with the aid of ultrasonication for 10 minutes. Then, the homogeneous solution was transferred to a 25 mL of stainless-steel teflonlined autoclave and heated to 180 °C for 24 hours. The product was cooled down to room temperature and washed with ultrapure water three times. Finally, the product was dried in an oven at 60 °C for 12 hours and pale-yellow crystals were obtained, named oPD-CA.

Preparation of the solution of TMB-H₂O₂

Take 40 µL of H₂O₂ droplets into 40 mL of water and shake them to obtain a mixed solution. Then add 20 mg of TMB to the mixed solution, stir until it is completely dissolved, and get 0.5 g L^{-1} of TMB solution.

Preparation of the dynamic anticounterfeiting labels

PVCA was used as the matrix for the oPD-CA material. First, 0.5 g oPD-CA powders were dispersed in 0.5 g PVCA and mechanical stirring was performed for 10 min. Then, the mixture was printed on a A4 paper by film printing. Finally, after curing at room temperature for 10 min, PVCA-based multiple-dimension luminescence images for dynamic anticounterfeiting labels were obtained.

Characterizations

Powder X-ray diffraction (XRD) patterns of the as-prepared samples were recorded with a Bruker D8 advance X-ray diffractometer using Cu Ka radiation. FTIR spectra were recorded in the range 400-4000 cm⁻¹ on a Nicolet iS50 FT-IR spectrophotometer. Thermogravimetric-differential scanning calorimetric analysis (TG-DSC) was done using a TGA/DSC3+ system (METTLER TOLEDO) in the nitrogen atmosphere to investigate the thermal stability and difference of the samples. X-ray photoelectron spectroscopy (XPS) measurements were performed on a VG ESCALAB 250 spectrometer with a monochromatic Al Ka X-ray source. Nuclear magnetic resonance spectroscopy was performed on a 500 MHz spectrometer (JNM-ECZ500R/S1). The morphology and microstructure were studied using a high-resolution transmission electron microscope (FEI Talos F200S G2). The absorption of the samples was measured by a UV-visible diffuse reflectance spectrometer (Cary 500, Varian Co.) using BaSO₄ as the reference. Steady-state

fluorescence/phosphorescence spectra, excitation spectra lifetimes and time-resolved emission spectra were obtained using a fluorophotometer (Edinburgh FL/FS980) equipped with a xenon arc lamp (Xe 900) and a microsecond flash-lamp (μF900), respectively. The photoluminescence quantum efficiency was recorded by a Hamamatsu Absolute PL Quantum Yield Spectrometer C11347. The phosphorescence quantum yield was obtained by eqn (1),

$$\varphi_{\text{Phos.}} = \frac{B}{A} \times \varphi_{\text{PL}} \tag{1}$$

In eqn (1), $\varphi_{\text{phos.}}$ and φ_{PL} represent the phosphorescence quantum yield and absolute photoluminescence quantum yield, respectively. A and B are the integral area of total photoluminescence and phosphorescence spectra, respectively. The photos of fluorescence and phosphorescence emission were recorded by a digital camera SONY DSC-HX400.

Results and discussion

Synthesis and characterization of small molecules doped CA

It has been previously reported that graphite-phase carbon nitride (g-C₃N₄) can be transformed to colorless CA crystals through hydrothermal synthesis. 54 Therefore, it is believed that

small organic molecule doped CA crystals could also be synthesized in the same way. To probe this notion, o-phenylenediamine (oPD) doped graphitic carbon nitride (named PCN) was fabricated by molten salt treatment (350 °C, 4 h) of melamine and oPD, the resultant mixture was then subjected to a hydrothermal treatment (180 °C, 24 h) (see Scheme S1 for the synthesis process), yielding faint yellow columnar crystals (named oPD-CA) (see Fig. S1a and b, ESI†). Sub-micron particles with bulk morphology were observed from the TEM image (see Fig. S1c and d, ESI†). It should be noted that these particles can be easily damaged upon electron beam irradiation during the TEM measurement (see Fig. S1d, ESI†), which indicates that they are organic materials. 55,56 The oPD-CA crystals were ground into powder to examine the powder X-ray diffraction (XRD) pattern, ¹³C-NMR and Fourier transform infrared (FTIR) spectra in comparison to those of CA and oPD samples (Fig. 1ac). (1) Strong diffraction peaks were observed at about 19.8°, 22.44°, 26.4°, and 29.8°, which correspond to (111), (002), (020) and (202) crystal facets of the CA matrix, respectively. They all shift to smaller diffraction angles ($\sim 0.3^{\circ}$) compared to those of the pristine CA crystals. (2) In the ¹³C-NMR spectra, oPD-CA shows a major chemical shift for CA located at 135 ppm and three weak chemical shifts for oPD located at 135, 117 and 115 ppm, which confirms the doping of oPD into CA.

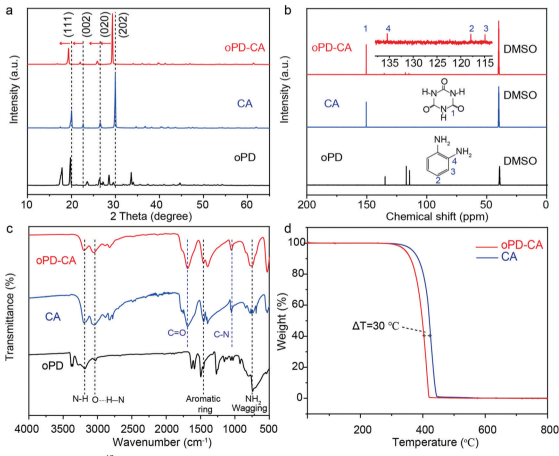


Fig. 1 (a) Powder XRD patterns, (b) ¹³C-NMR spectra, (c) FTIR spectra of as-synthesized oPD-CA, pristine CA, and oPD, and (d) TG curves of oPD-CA, and the pristine CA in the nitrogen environment.

According to the external standard method (see Fig. S2, ESI† for calculation details), the experimental molecular doping concentration is calculated to be 96:1 (molar ratio of CA: oPD), which is lower than the theoretical value of 40:1. (3) All the diffraction peaks and FTIR absorption peaks of oPD-CA are well matched with those of the pristine CA crystals, but a distinct amino (-NH₂) wagging band (741 cm⁻¹) and an aromatic ring (1262 cm⁻¹ and 1500 cm⁻¹) were observed. Moreover, the peak intensity of the hydrogen bond at 3046 cm⁻¹ is significantly increased (see Note S1 and Fig. S3, ESI† for calculation details). Thermogravimetry (TG) measurements were carried out to determine the weight percentage of the oPD dopant. Fig. 1d compares the TG curves of oPD-CA and CA. Obviously, both of oPD-CA and CA are thermally stable below 300 °C in air. The main difference between the two samples is that the complete-decomposition temperature of oPD-CA decreases by ~ 30 °C compared to CA. This indicated that oPD was successfully doped into the CA matrix and it reduced the thermal stability of oPD-CA. Other molecules (e.g., catechol) can also be doped in CA, forming an organic afterglow material with different optical properties (see Fig. S4, ESI†).

X-Ray photoelectron spectroscopy (XPS) analysis was carried out to gain insight into the chemical structure of oPD-CA and

CA samples (Fig. 2). In the survey spectra, C, N, and O elements are observed for both of oPD-CA and CA (Fig. 2a). The peak fitting of the high-resolution C_{1s} peaks reveals that there are three carbon species with binding energies at 284.8 eV, 286.3 eV, and 289.9 eV, which are assigned to C=C/C-C carbon bonds, C-N species and sp²-bond carbon (C=O), respectively (Fig. 2b). 54 Fig. 2c shows the high-resolution N_{1s} XPS spectra of oPD-CA and CA, in which two types of nitrogen species with binding energy at 400.7 eV and 402.0 eV are observed for the two samples. However, a new peak of Ph-NH2 with binding energy at 399.2 eV is only detected for oPD-CA. In addition, the O_{1s} can be deconvoluted into two peaks, responding to C=O species at 532.2 eV and C-O at 533.3 eV, respectively (Fig. 2d). The higher C_{1s} intensity of $C = C (sp^2)$ carbon bonds and the presence of Ph-NH2 species for oPD-CA reveal the successful doping of oPD into the CA matrix, which is in good agreement with the XRD, 13C-NMR spectra, FTIR and TG data.

Optical properties of oPD-CA

The synthesized oPD-CA crystals exhibit three different optical responses, i.e., fluorescence, TADF and phosphorescence (Fig. 3a-f). The wavelength-dependent luminescence spectra of oPD-CA at room temperature were recorded wherein the

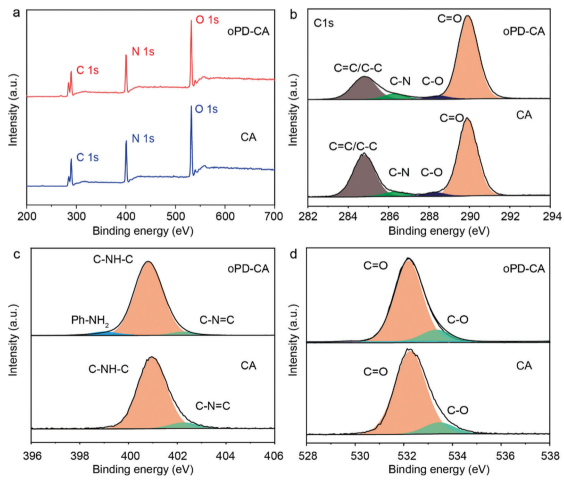


Fig. 2 XPS spectra of oPD-CA and CA for (a) survey scan, (b) C1s, (c) N1s, and (d) O1s.

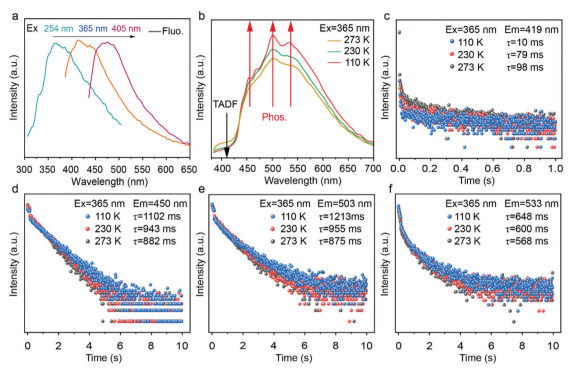


Fig. 3 (a) Excitation wavelength dependent room-temperature fluorescence spectra, (b) temperature-dependent afterglow spectra, (c) TADF decay kinetics, (d) phosphorescence decay kinetics for 503 nm, and (f) phosphorescence decay kinetics for 533 nm.

red-shift of an emission peak happens upon an increase in excitation wavelength (Fig. 3a). Interestingly, a strong green and cyan afterglow of the oPD-CA powder in air or in water was observed by naked eyes after the removal of an ultraviolet (UV) lamp for 0.1 s and 1 s, respectively (see Fig. S5, ESI†). In contrast, only a very weak blue afterglow of the pristine CA powder was observed only in air but not in water (see Fig. S6, ESI†). Obviously, the doping of oPD has significantly enhanced the afterglow of the CA matrix. Under 365 nm light excitation, the fluorescence quantum yield and phosphorescence quantum yield of oPD-CA were recorded as 10.5% and 14.2%, respectively (see Table S1, ESI†). To understand the temperature-dependent afterglow behavior of oPD-CA, its afterglow spectra were obtained under UV (365 nm) excitation at different temperatures (Fig. 3b). Four emission peaks at 419 nm, 450 nm, 503 nm and 533 nm were observed. The position of the shoulder peak (Fig. 3b, black curve) is almost identical to that of its fluorescence emission peak under the same excitation conditions. Surprisingly, the emission intensities of the three peaks shown in Fig. 3b evolve in an opposite trend with the decrease of temperature. More specifically, the emission intensity of the shoulder peak drops, while that of the main peak increases, indicating that the origin of the first peak is different from the other three peaks. It is believed that the shoulder peak is assigned to TADF, while the other peak is attributed to the phosphorescence.^{57,58} Fig. 3c-f show the emission decay profiles of TADF and phosphorescence components of oPD-CA at different temperatures. All the afterglow-lifetime decay curves can be well fitted by triexponential decay (see Tables S2 and S3, ESI†). The average lifetimes were calculated by the following eqn (2):

$$\tau = \sum a_i \tau_i^2 / \sum a_i \tau_i \tag{2}$$

where τ_i represents a luminescent component with different lifetimes (*i.e.*, τ_1 , τ_2 and τ_3). a_i is the initial strength of each component (amplitudes) of the lifetime (τ_i).⁵⁹

The calculated lifetimes are 10 ms, 79 ms, and 98 ms for TADF, 1102 ms, 943 ms and 882 ms for the phosphorescence of 450 nm, 1213 ms, 955 ms and 875 ms for the phosphorescence of 503 nm and 648 ms, 600 ms and 568 ms for the phosphorescence of 533 nm at 273 K, 230 K, and 110 K, respectively, under excitation at 365 nm and the slit size for the excitation and emission wavelength 7 nm and 7 nm, respectively. Due to the shorter lifetime at emission wavelengths 419 nm and 533 nm, the yellow-green composite afterglow can be observed first. Conversely, the cyan complex afterglow can be observed later due to longer lifetimes at emission wavelengths of 450 nm and 503 nm. It explains well the reason why oPD-CA has a yellow-green and a cyan afterglow, after 365 nm light irradiation. These unique triple-mode emission phenomena (i.e., fluorescence, TADF and phosphorescence) of the novel organic oPD-CA material pave a good way for advanced anticounterfeiting applications due to the encoding of multidimensional dynamical optical information.

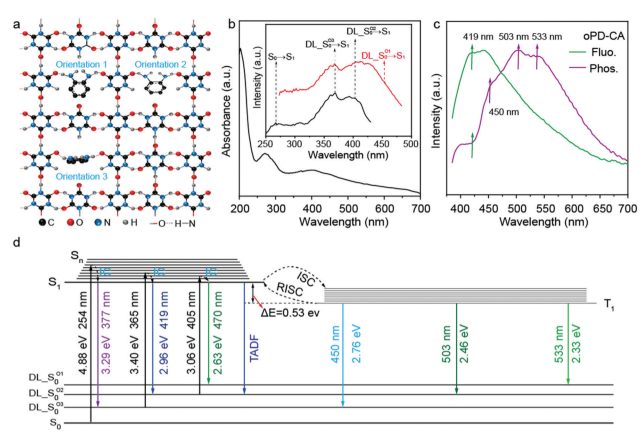


Fig. 4 The role of the oPD dopant for the triple-mode emissions: (a) The proposed chemical structure, (b) diffuse reflectance spectrum of oPD-CA, and the inset represents the excitation spectrum of oPD-CA at 450 nm and 503 nm emission wavelength, (c) afterglow spectra of oPD-CA at room temperature, and (d) luminescence mechanism of oPD-CA.

Luminescence mechanism of oPD-CA

Fig. 4a illustrates the proposed molecular structure of oPD-CA crystals. As shown in Fig. 4a, the oPD molecules are randomly doped in the CA matrix with different orientations by replacing some CA molecules in the lattice. To understand the role of the dopant oPD for the observed triple-mode emissions, we studied the diffuse reflectance spectroscopy (DRS), excitation, fluorescence and phosphorescence properties of oPD-CA at the room temperature (Fig. 4b and c). As shown in Fig. 4b, there are two broad absorption peaks at around 270 nm and 400 nm. Four excitation peaks correspond to the transitions from the new ground states (i.e., So and the three doping levels) to So of CA. (see the inset of Fig. 4b). Under the excitation of 365 nm light, the maximum fluorescence emission peak of oPD-CA is located at 419 nm, and the phosphorescence peaks are located at 450 nm, 503 nm and 533 nm, respectively. Moreover, a weak delayed fluorescence can be observed around 419 nm (Fig. 4c).

Since CA itself shows a weak blue afterglow in air (see Fig. S6, ESI \dagger), both the singlet state (S₁) and the triplet state (T₁) must co-exist in the system. The doping of oPD into the system inevitably introduces new doping levels, which may originate from the orientation of the dopant molecule. 60 According to Kasha's rule, 61 photon emission should occur only from the lowest excited states of the luminescent material (S1 and/or triplet state T_1) to its ground states (S_0) . The observance of the

red-shift of the fluorescent (S1-S0 transitions) and phosphorescent (T_1 - S_0 transitions) emissions for the doped CA indicates that the doped energy levels should locate close to the original So level of CA, forming new ground states. Based on the luminescence properties of oPD-CA shown in Fig. 3a and b as well as the absorption and excitation spectra shown in Fig. 4b, we proposed a possible luminescence mechanism of oPD-CA to explain the observed triple optical responses. As shown in Fig. 4d, under UV light irradiation, the excited electrons are relaxed to the S₁ state through the direct internal conversion (IC) process, and partial electrons return to the new ground states (i.e., $S_1 \to DL_S_0^{O1}$, $S_1 \to DL_S_0^{O2}$ and $S_1 \to DL_S_0^{O3}$) via emitting fluorescence (Fig. 3a). At the same time, some cross to the triplet excited state (T₁) through the intersystem crossover (ISC) and return to the new ground states (i.e., $T_1 \rightarrow DL_S_0^{O1}$, $T_1 \rightarrow DL_S_0^{O2}$ and $T_1 \rightarrow DL_S_0^{O3}$), via emitting phosphorescence (Fig. 3b and 4c). Notably, it has been reported that when the energy band difference of S₁ and T₁ is sufficiently small, the competition rate of the reverse intersystem crossover (RISC) at room temperature will be large enough. 62,63 The energy difference between S1 and T1 for the oPD-CA system is calculated to be as small as 0.53 eV. Such a small energy difference allows some electrons at T₁ to go back to S₁ through the RISC process by absorbing the surrounding thermal energy and eventually de-excite to the new $DL_{S_0}^{O2}$ ground state, thus emitting TADF.

Obviously, high temperature is favorable for TADF to occur, which explains the decrease of TADF intensity as the temperature drops (Fig. 3b). The unique triple-mode optical responses are dynamically tunable, making the oPD-CA material an excellent information carrier for anti-counterfeiting applications.

More encouragingly, the obtained oPD-CA also displays artificial enzyme activity besides its optical excellence. The

OPD-CA powder can make 3,3',5,5'-tetramethylbenzidine (TMB) gradually turn blue in the presence of hydrogen peroxide (H₂O₂). Two absorption peaks at about 370 nm and 650 nm were observed and intensified with increasing reaction time, from the UV-Vis absorption spectrum (Fig. S7, ESI†). The silica powder does not change the color of the solution. The peroxidase-like catalysis was revealed by triggering the colorimetric reaction using the TMB substrate in the presence of

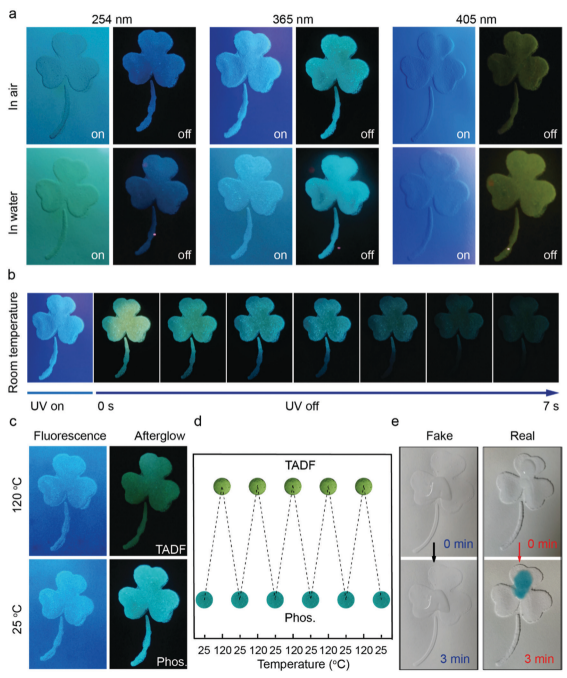


Fig. 5 Photograph of the fabricated dynamic security labels modulated with (a) excitation wavelength in air (top column) and in water (bottom column), (b) time, (c and d) temperature, and (e) TMB colorimetric detection of the fake label manufactured using SiO₂ (left) and real label manufactured using oPD-CA (right).

H₂O₂ (Fig. S8, ESI†). This is mainly attributed to the confinement of oPD inside the CA matrix-based host, which plays a vital role in the generation of catalytic activity.64-66 In the present oPD-CA system, the -C=O groups could serve as the catalytic activity site and substrate-binding site, respectively.⁶⁷

Triple optically modulated and enzymatically responsive dynamic anti-counterfeiting

Intrigued by the fascinating optical properties and artificial enzyme activity of oPD-CA, we developed a quad-modulated dynamic anti-counterfeiting strategy using oPD-CA as an information carrier. The dynamic security labels with various patterns were fabricated by stencil printing of a slurry composed of polyvinyl acetate and oPD-CA on A4 paper (Fig. 5). First, the emission fluorescence and afterglow colors of the fabricated security label can be dynamically tuned with different excitation wavelengths. Fig. 5a shows the optical images of a clover pattern that is placed in air and water with UV light on and off, respectively. The pattern exhibits bright blue fluorescence at room temperature under excitation with 254 nm UV light. After turning off the UV light, the pattern area remains blue due to the phosphorescent emission, while the area surrounding the pattern turns black. When the excitation wavelength shifted to 365 nm, the color of the pattern turns cyan under both UV on and off conditions. With a further increase of excitation wavelength of 419 nm, the blue fluorescence can still be observed, while the yellow-green phosphorescence is detected by a camera. These results are consistent with the optical spectra shown in Fig. 3 wherein the emission peak red-shifts with the increase of the excitation wavelength. Such excitation wavelength modulated, and water-resistant optical properties can be used as the first layer of security (see Fig. 5a and Videos S1 and S2, ESI†). Furthermore, the excellent time-resolved encryption performance of the security label can be used as the second layer of security. As shown in Fig. 5b, the color of the clover pattern immediately changed from blue fluorescence to yellow-green phosphorescence, followed by an instant change to cyan phosphorescence within 1 s, and the phosphorescence can last for 7 s, after turning off the 365 nm UV light (see Video S3, ESI†). The intensity of the afterglow color of the security labels is able to be dynamically tuned with time, which can serve as the second layer of security. Finally, the afterglow colors of the security labels are also able to be dynamically modulated by changing the external temperatures (Fig. 5c). Although there is no obvious change in fluorescence colors under 365 nm excitation when switching the label's temperature from 25 °C to 120 °C, the afterglow color changes from cyan to blue (Fig. 5c and Video S4, ESI†). An advanced anti-counterfeiting label requires that the optical material owns not only a great color tunability but also an excellent reversibility. We, therefore, monitored the afterglow color change of the oPD-CA material by repeatedly switching the temperature between 25 °C and 120 °C using a heating plate. It is found that the afterglow color change is highly reversible (Fig. 5d) between cyan and blue. This temperature modulated afterglow colors are used as the third layer of security. The system of oPD-CA-TMB-H2O2 we

present can be applied to the fourth layer of security. After adding TMB-H2O2 mixed solution to the label, the true label could turn blue within three minutes (Fig. 5e, right), while the false label could not (Fig. 5e, left). Moreover, the developed anti-counterfeiting labels offer four layers of security and can be massively produced using stencil printing techniques at low cost and visual authentication. These merits make the security label difficult to counterfeit and meet the criteria and demand for advanced anti-counterfeiting.

Conclusions

In summary, a type of unprecedented organic afterglow material, small molecule doped CA, was successfully fabricated. Our results indicate that small molecules (e.g., oPD and catechol) can be doped in the lattice of CA with different orientations. Not only the phosphorescence intensity and lifetime are greatly enhanced, but also oPD-CA is endowed with the optical properties of TADF and polychromatic phosphorescence due to the formation of new ground states stemming from molecular doping. By simply changing the doping molecules, the optical properties of CA can be tailored. As proof-of-concept, we demonstrated a universal synthetic method for the fabrication of small molecule doped CA afterglow materials and a costeffective, easily authenticated, quad-modulated (i.e., artificial enzyme responsive, excitation-wavelength/temperature switchable and time-resolved) dynamic anti-counterfeiting label using oPD-CA as an information carrier. Such quadruple multiresponsive anti-counterfeiting labels are distinct from the traditional static security labels with mono-optical information encryption, making them harder to counterfeit.

Author contributions

Q. H. J., W. S. W., C. J., and Y. H. Z. conceived the idea; Q. H. J. designed the experiments and synthesized materials; H. R. C. and Y. F. L. performed the transmission and fluorescence spectra test; M. L., X. F., and H. X. C. assisted in data analysis; Q. H. J., C. J, and Y. Z. wrote the manuscript and all authors reviewed it.

Conflicts of interest

There are no conflicts to declare.

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Research Article

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