



Carbogenically coated silica nanoparticles and their forensic applications†

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Carbogenically coated silica nanoparticles (C-SiO₂) exhibit color-tunability and carry great promise for two important forensic applications. First, the C-SiO₂ nanopowders are ideal for fingerprint development, yielding strong contrast against multicoloured and patterned backgrounds. Second, spontaneous nanoparticle aggregation leads to non-duplicable, inexpensive nanotags that can support sustainable technologies to combat counterfeiting.

The remarkable photophysical properties of carbogenic nanoparticles (otherwise known as C-dots) are systematically explored in a variety of applications including printing inks, photocatalysis, biologically labelling, bioimaging and chemical sensing.^{1–5} Their non-toxic and biocompatible nature^{6,7} endows distinct advantages over conventional polyaromatic dyes and heavy metal-based quantum dots.

Large-scale and cost-effective synthesis of C-dots is realised *via* pyrolytic decomposition of virtually any type of carbon-rich precursors including plants tissue,⁸ natural saccharides,^{9,10} animal products¹¹ and synthetic polymers.^{12,13} Alternatively, C-dots with varying graphitization degree are derived through disintegration,¹⁴ hydrothermal treatment¹⁵ and electrooxidation¹⁶ of carbon nanotubes, carbon nanofibers and graphite. In principle, quantum yield can be improved *via* surface passivation,¹⁷ heteroatom doping¹⁸ and the *in situ* formation of organic chromophores.^{19,20}

In this report we focus on the exploitation of the advanced photophysical properties of carbogenically coated silica nanoparticles (C-SiO₂) in two distinct forensic applications. First, we demonstrate that the colour-tuneable C-SiO₂ can be used as high performance nanopowders for latent fingerprint enhancement.

Second, spontaneous aggregation of the C-SiO₂ during solvent evaporation generates non-duplicable photoluminescent motives that are ideal nanotags to mark and authenticate products. As reviewed recently a number of studies report the use of SiO₂ based nanomaterials for fingerprint enhancement.²¹ Notable contributions focus on the use of surface modified SiO₂ nanoparticles to gain insights into the mechanisms responsible for fingermark detection²² and the use of fluorescent dye-doped SiO₂ nanoparticles to improve the image homogeneity and reduce donor inter-variability.²³

To synthesize C-SiO₂, colloidal silica was treated with dimethyloctadecyl[3-(trimethoxysilyl)propyl]ammonium chloride to give a product with 34 wt% organic content as determined by TGA (ESI,† Fig. S1). Subsequently, the product was pyrolysed at 250 °C before being subjected to surface oxidation *via* HNO₃, amine functionalization and excessive dialysis against water. Elemental analysis of C-SiO₂ thus received suggests the presence of 26% C, along with minor amounts of H and N (4 and 5%, respectively). The FTIR spectrum (ESI,† Fig. S2) shows peaks attributed^{24,25} to the vibrational stretching of C–H (1380 cm⁻¹), the anti-symmetric and symmetric stretching vibrations of N–C=O (1550 cm⁻¹ and 1658 cm⁻¹, respectively) and symmetric (2850 cm⁻¹) and anti-symmetric (2920 cm⁻¹) stretching vibration of sp² C–H.

The TEM image of C-SiO₂ (Fig. 1) indicates the presence of spherical nanoparticles with average diameter 22 ± 2 nm. The nanoparticles are easily dispersed in water forming stable colloidal dispersions with pH = 10.2, consistent with the presence of surface functionalities generated through the acid/amine treatment. Fig. 2 suggests that the aqueous dispersions exhibit excitation wavelength dependent photoluminescent emission, in a manner similar to that observed for other carbon-based nano-emitters such as C-dots and graphene dots. The emission mechanism is not thoroughly understood, however contributions stemming from surface defects and the conjugated π-domains have been identified.^{26,27}

The freeze-dried fine powder shows excellent flowability and can be easily applied to a variety of non-porous surfaces (ESI,† Fig. S3). The nanopowder adheres strongly to the fingerprints

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† Electronic supplementary information (ESI) available: Detailed experimental section, TGA thermograph, FTIR spectra, AFIS analysis of fingerprints, hydrodynamic diameter and fluorescence images of fingerprints and nanotags. See DOI: 10.1039/c6cc02556k



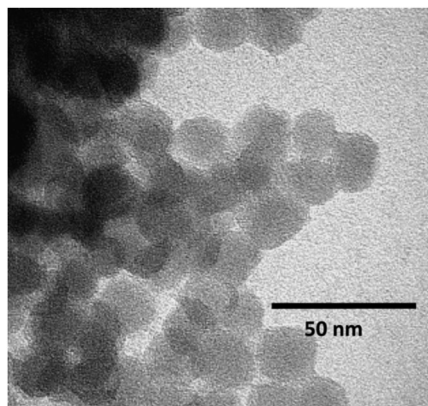


Fig. 1 TEM image of the C-SiO₂ nanoparticles.

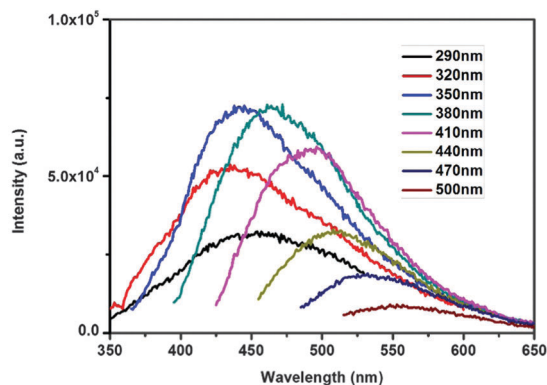


Fig. 2 Photoluminescence spectra of C-SiO₂ aqueous solutions under different excitation wavelengths.

revealing well-resolved patterns that meet the standard for individual identification (ESI,† Fig. S3 and S4). Automated Fingerprint Identification System (AFIS) analysis of a fingerprint developed by C-SiO₂ reveals 73 minutiae (details of a fingerprint such as ridge ending, ridge bifurcation, etc) compared to 65 minutiae for an identical fingerprint developed by a standard white fingerprint powder (WFP) (ESI,† Fig. S5).

Significantly, as shown in Fig. 3a the nanopowders exhibit colour-tunability and emit within the blue, green and red region when they are irradiated by violet, blue and green light, respectively. The supreme photophysical properties of C-SiO₂ can open new horizons in forensic investigation by bypassing problems related to strong background interference from overlapping letters or fluorescent, multi-coloured and textured substrates. To demonstrate this behaviour, a set of fingerprints deposited on a highly fluorescent cardboard were developed under identical conditions using the C-SiO₂ nanopowder and a commercial WFP, respectively. High quality images are collected for the C-SiO₂ enhanced fingerprints (left image in Fig. 3b) under a 445 nm wavelength laser on a crime-lite imager. In contrast, the fingerprints developed using the standard WFP suffer from strong background interference leading to poor image quality (right image in Fig. 3b). A full set of images under the different wavelengths available on the crime lite imager can be seen in ESI,† Fig. S6. Moreover,

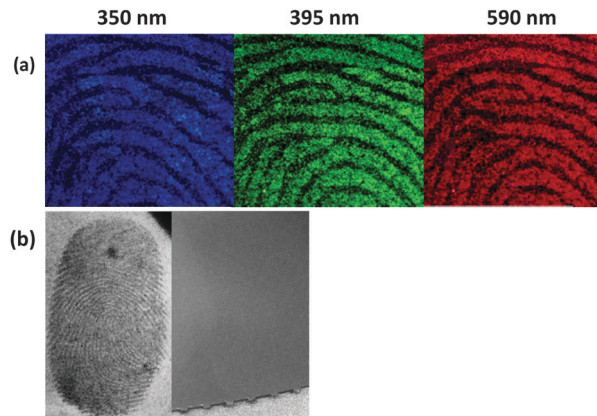


Fig. 3 (a) Fluorescent microscopy images of a fingerprint developed with the C-SiO₂ nanopowder. (b) Comparison between the C-SiO₂ and a commercial fingerprint powder on a fluorescent background investigated on a crime lite imager (445 nm laser with a 455 nm filter).

a fingerprint developed with C-SiO₂ compares favourably with that developed by a commercial fluorescent powder, in the sense that the latter shows strong contrast only under certain illumination wavelengths (ESI,† Fig. S7).

Fingerprint matching remains by far the most reliable biometric method used by law enforcement for individual identification, but its accuracy critically depends on the quality of the recovered impressions. To that end, a large number of fingerprint powders have been developed to maximize contrast against a variety of radically different substrates. In that sense, the use of a single, yet colour-tunable C-SiO₂ nanopowder offers significant advantages in terms of dusting speed and image quality.

We have recently introduced a close related class of nanopowders comprising minor amounts of C-dots that also exhibit colour-tunability with respect to the incident radiation and, thus, are suitable for fingerprint enhancement.²⁸ Another study describes the use of aqueous dispersions of C-dot/poly(dimethylacrylamide) nanocomposites for fingerprint detection.²⁹ Notably, nanopowders with high C-dot content are not photoactive, presumably due to extensive self-quenching effects.³⁰ A recent study suggests that blue, green and tan silicon-carbon dots/silica nanocomposite powders show adjustable optical properties as a function of their composition and structure.³¹ Here we demonstrate colour-tunable nanopowders containing a single type of nanoparticles and are, thus, distinctly different from the nanocomposite and hybrid materials reported previously. The main advantage of C-SiO₂ nanoparticles compared to C-dots based nanocomposites refers to their much larger carbogenic surface that holds great promise for selective and sensitive sensing of forensically important compounds found in fingerprints. We note that this report is a proof-of-concept investigation corresponding to phase 1 pilot studies as described by the International Fingerprint Research Group.³²

When the pH of the aqueous suspension of C-SiO₂ drops below 8.5, the colloidal stability is compromised as evident by the evolution of large aggregates (ESI,† Fig. S8). During water evaporation from an acidified dispersion, the C-SiO₂ were seen to undergo spontaneous and random self-assembly, giving rise



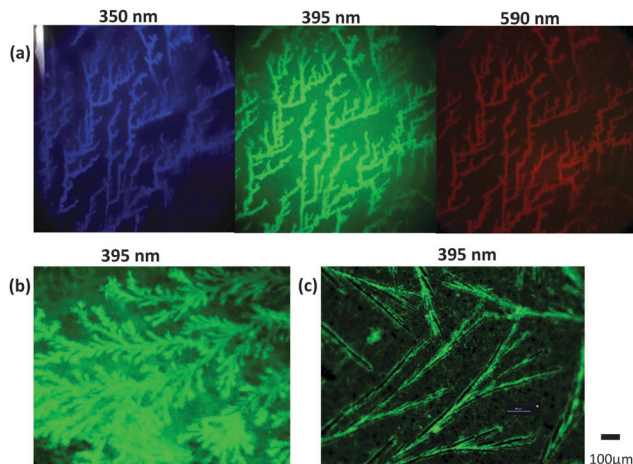


Fig. 4 Fluorescent microscopy images of the nanotags; (a) deposited in glass and illuminated under different excitation wavelengths, (b) deposited in glass from a more concentrated suspension, (c) deposited on a polymeric surface.

to motives that are highly fluorescent under laser radiation as shown in Fig. 4a. For comparison, we display two more images; one derived from a more concentrated dispersion (Fig. 4b) and one formed on a polymeric surface (Fig. 4c). (In contrast, no patterns are formed from dispersions with pH = 10, as shown in ESI,† Fig. S9).

The SEM images displayed in Fig. 5 reveal a high level of structural complexity confirming that the motives are nearly impossible to duplicate and can serve as nanotags for object authentication. The nanotags adhere strongly to a variety of substrates (polymers, metals, glass) and remain unaltered even after prolonged exposure to high temperature (ESI,† Fig. S10). We attribute this remarkable structural stability and durability to their organic-free and polymeric-free composition.

Suffice to say that counterfeit accounts for a 5–7% world trade, causing a severe financial drain to the global economy. This crime deprives taxable income from *bona fide* business, discourages investment for innovation and inhibits social progress. Counterfeit drugs not only fail to cure, but they can cause severe health issues. Recent trends to address this problem rely on the use of upconverting nanocrystals³³ and dye-doped nanoparticles³⁴ in integrated safety features and security graphics; however, those sophisticated compounds are often toxic and prohibitively expensive. In contrast, C-SiO₂ based nanotags are cost-effective and non-toxic for humans and the environment. Moreover, by virtue of their colour-tuneable nature, C-SiO₂ based nanotags impart enhanced security features, compared to commonly used dyes that display only one fixed colour.

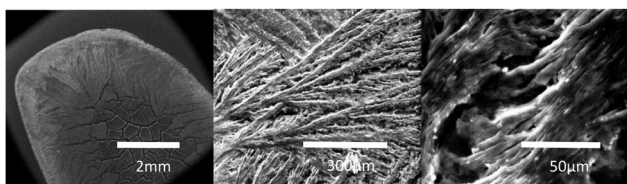


Fig. 5 SEM images of the formed patterns under different magnifications.

In conclusion, we introduce a new class of colour-tuneable nanomaterials (C-SiO₂) and we demonstrate their great potential in two distinct forensic applications. As fingerprint powders they circumvent problems stemming from poor image quality against multicoloured substrates. As nanoparticles that undergo spontaneous and random aggregation, they generate non-repeatable motives with a high level of structural complexity that can be used to mark and authenticate products. Those inexpensive and non-toxic nanotags are viable alternatives to current solutions with prohibitively high cost and they can support sustainable technologies to combat counterfeiting.

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