Carbon-dot–hydrogel for enzyme-mediated bacterial detection†

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A hybrid carbon-dot (C-dot)–hydrogel matrix was constructed and employed for detection of bacteria. The transduction mechanism is novel, based upon cleavage of ester bonds within the hydrogel scaffold by bacterially-secreted esterases; the ensuing fluidization of the hydrogel resulted in aggregation of the embedded C-dots and consequent quenching of their fluorescence. We show that the C-dot–hydrogel exhibits high sensitivity and can distinguish among bacterial species through modulation of the emitted fluorescence, depending upon their esterase secretions.

Introduction

Hydrogels, three dimensional flexible networks of water-soluble polymers, are useful host materials enabling encapsulation of high concentrations of varied guest molecules and nanoparticles.1–7 Hydrogels are generally classified according to two categories: (i) thermally-irreversible hydrogels in which the gelator molecules are crosslinked by covalent bonds (defined as chemical gels),7 and (ii) thermally-reversible physical gels in which the gelator molecules exhibit weak physical interactions.3 Hydrogels have been used in varied applications including tissue engineering, drug delivery and controlled-release, and medical implants.5

Hydrogel composites have been used as useful biosensing platforms. Redox-based sensing schemes utilizing hydrogel-encapsulated enzymes were reported.6,9 A peptide hydrogel consisting of a Fmoc-diphenylalanine-encapsulating enzyme receptor (e.g., glucose oxidase or horseradish peroxidase) and semiconductor quantum dots (QDs) was reported for detection of glucose and toxic phenolic compounds through quenching of the QDs’ fluorescence.8 A hydrogel cross-linked with aptamers was reported to detect very low amount of cocaine through visual detection.9 Porous silica polyacrylamide hydrogel system served as an optical bacterial sensor.12

The hydrogel-based sensors discussed above largely rely upon molecular recognition and/or redox reactions involving embedded enzymes as the components in the transduction mechanism. In such systems, the hydrogel serves only as a host matrix for immobilization of guest biomolecules. Exploiting changes in the physical state of the hydrogel as a sensing strategy could be, in fact, a promising sensing avenue. Varied externally-added molecules were shown to affect the structural features and mechanical properties of hydrogel matrices.12–14 Hydrogel framework modulation has been employed as a sensing platform.15,16

Carbon dots (C-dots) are newly discovered fluorescent carbonaceous nanoparticles (<10 nm), and have attracted broad interest in light of their intriguing luminescence properties, particularly a broad range of excitation-dependent emission range.17–24 C-dots are biocompatible and less cytotoxic than semiconductor quantum dots, thus constituting promising vehicles for biological studies.19,25,26 Modulation of C-dots’ fluorescence has been used as a powerful transduction mechanism in varied sensing applications, for example targeting metal ions,27,28 organic molecules,29,30 and radical species.31–33 C-dots have been recently used as a platform for bacterial sensing and imaging.34 Hybrid carbon-dot hydrogel systems were also used for detection of heavy metal ions.34,35

Herein, we report construction of a new C-dot–hydrogel hybrid material which was used for bacterial detection through modulation of the fluorescence emission of the hydrogel-embedded C-dots. In particular, we found that cleavage of hydrogel scaffolds’ ester bonds by bacterially-secreted esterases and lipases resulted in fluidization of the hydrogel, consequent aggregation and concomitant fluorescence quenching of the embedded C-dots. The C-dot–hydrogel constitutes a sensitive bacterial detection platform since the transduction mechanism is based upon secreted enzymes rather than the bacterial cells themselves. Furthermore, the new assay can be employed for distinguishing bacterial species based upon the extent of their esterase/lipase secretion.

Experimental

Materials

D-Glucose, sodium sulfate, pyridine, lauric acid and sodium chloride were purchased from Sigma Aldrich, USA. L-(+)-Tartaric acid was purchased from Acros Organics, USA. D-Glucose, sodium bicarbonate, sodium hydrogen carbonate, sodium glutamate, sodium citrate and sodium benzoate were purchased from Acros Organics, USA. Sodium chloride was purchased from Gladen, Israel. Ethanol and toluene were purchased from Riedel de Haen, Germany. A 2,2′-bipyridine solution in ethanol and tetrahydrofuran (THF) was purchased from Sigma Aldrich, USA. Hydrogen peroxide (35%) was purchased from Fisher Scientific, USA. Dithiothreitol (70%) was purchased from Merck, Germany. All other reagents were of analytical grade and were used without further purification.
acid was purchased from Alfa-Aesar, England. Lauroyl chloride was purchased from TCI, Japan. Chloroform and n-hexane were bought from Daejung chemicals, Korea. Dimethylformamide (DMF) and acetone were purchased from Frutarom (Haifa, Israel). Ethyl acetate and concentrated hydrochloric acid were purchased from Bio-Lab Ltd (Jerusalem, Israel). Luria–Bertani (LB) agar was purchased from Pronadisa (Spain). All chemicals were used without further purification.

C-dot–hydrogel synthesis

The C-dots and hydrogel were prepared from 6-O-(O’-dilauroyl tartaric acid)α-D-glucose (compound 1, Fig. 1A) according to a published procedure. Briefly, we prepared 6-O-acylated fatty acid ester of α-glucose by reacting it with O’-dilauroyl tartaric acid anhydride. In a Teflon film tightened, septum-capped test tube 100 mg of 1 and 330 μL DI water were placed and then heated in an oven to 125 °C for 2.5 h. Upon completion of the carbonization, the reaction mixture was cooled to room temperature yielding a brown precipitate indicating the formation of carbon dots and purified for further work. For preparation of the C-dot–hydrogel, 5 mg of C-dots were mixed with 50 μg of compound 1 and dissolved in 0.5 mL DI water in a sealed vial by sonicating for 30 min at 50 °C and further heated for 2–3 min until the gelator compound was dissolved completely. Upon cooling to room temperature for about 24 hours, the aqueous solution completely gelified, tested by the “stable to inversion” method.

Bacterial growth

The bacteria used in the studies were Bacillus cereus, Bacillus subtilis (stain Bacillus 3610), Pseudomonas aeruginosa PA01 wild type, and Staphylococcus aureus wild type. The bacteria were grown aerobically at 37 °C in sterilized solid LB medium composed of 13.5% yeast extract, 27% peptone, 27% NaCl, and 32.5% agar at pH 7.4. After overnight growth, a colony from each bacterial strain was taken and added to 10 mL sterilized LB medium and incubated at 37 °C. Bacterial growth was monitored in the specific time points through measuring the concentration of the bacteria by visible spectroscopy (10⁶ cells per mL when optical density at 600 nm was 1.0). Bacterial cells were separated from the growth medium through centrifugation followed by washing with sterilized water several times and resuspending in distilled water.

Confocal fluorescence microscopy

Confocal microscopy images of the C-dot–hydrogel were acquired on an UltraVIEW system (PerkinElmer Life Sciences, Waltham, MA) equipped with an Axiovert-200M microscope (Zeiss, Oberkochen, Germany) and a Plan-Neofluar 63×/1.4 oil objective. Excitation wavelengths of 405, 488 and 514 nm were produced by an argon/krypton laser. For the experiment the C-dot–hydrogel was added on a glass slide.

Scanning electron microscopy (SEM)

Scanning electron microscopy (SEM) images were recorded on a Quanta 200 SEM. For the SEM study, the dilute solution of gel materials was dried on a glass cover slip and gold coating was carried out.

Fluorescence spectroscopy and detection of bacteria

A Varioskan 96-well plate reader was used for the fluorescence analysis. 50 μL of C-dot–hydrogel was placed in each well and gelation was carried out. After 24 hours, fluorescence measurement and titration with different bacterial stains at 10⁶ cells per mL bacterial concentrations were carried out using λex = 375 nm at 37 °C by addition of 15 μL aqueous bacterial suspension. Quenching efficiencies (Q. E.) of the C-dot–hydrogel hybrid were calculated by the Stern–Volmer equation:

\[
Q. \ E. = \frac{I_0 - I}{I}
\]

in which I₀ and I refer to the fluorescence intensities of the system (λem = 430 nm) in the absence and presence of bacterial suspensions, respectively.

Thermal and chemical treatments on B. cereus

The aqueous suspensions of 10⁶ cells per mL bacterial cells were heated to 100 °C and then cooled to room temperature. From
this suspension, 15 μL bacteria were added to the C-dot–
hydrogel to observe the change in fluorescence intensity
(430 nm emission wavelength at 37 °C, where λex: 375 nm).
To determine the contribution of divalent metal ions to
the enzymatic activity of B. cereus, 15 μL of 5 mM EDTA solution was
added together with 15 μL of aqueous bacterial suspension to
the C-dot–hydrogel.

Rheology measurements
Rheology experiments were carried out using an Advanced
Rheometer AR 2000 (TA Instruments) by cone and plate geome-
try in a Peltier plate. The cone diameter was 40 mm, cone angle
4°, and truncation 104 μm. The rheology analysis was per-
formed by changing the shear rate (at 37 °C).

NMR studies were carried out on a Bruker DPX 400 MHz spec-
trometer at 300 K. Compound concentrations were 10 mm in
methanol-d4. Chemical changes were monitored through
incubation of suspensions of Bacillus cereus with hydrogel
precursor 1 for 30 minutes. The reaction product was lyophi-
lized for 2 h, and 1H and 13C NMR experiments were carried out
without further purification.

FT-IR analyses
All the FT-IR measurements were performed on a Thermo
Scientific Nicolet 6700 spectrometer. The organic compound
before and after addition of bacteria were identified by using
this technique. The extraction of the organic compound was
similar like the NMR experiments.

Results and discussion
Fig. 1 depicts the preparation scheme of the C-dot–hydrogel
hybrid and the enzyme-mediated bacterial detection concept.
The hybrid hydrogel was assembled through mixing the
precursor molecule 6-O-(O-C’-dilauroyltartaryl)-D-glucose and
amphiphilic Cdots in water (Fig. 1A). Electron microscopy
analysis confirmed the crystalline structure of the graphitic
cores of the Cdots (Fig. S1, ESI†). The Cdots/precursor solution
was gently heated until a homogeneous mixture was formed,
followed by gelation which occurred at room temperature.
Importantly, in the initial gel state prior to addition of bacterial
suspensions (Fig. 1B, left), the embedded amphiphilic Cdots
were dispersed within the hydrogel host matrix due to affinity
to amphiphilic and hydrophobic domains within the internal
porous surface. The hydrogel-embedded Cdots exhibited
enhanced fluorescence as they were not aggregated, and thus
fluorescence quenching was blocked.14,15

However, the hydrogel network was disrupted upon addition
of bacterial suspensions through ester bond cleavage by
bacterially-secreted esterases (enzymatic cleavage sites are
indicated by the arrows in Fig. 1B). The collapse of the hydrogel
network gave rise to aggregation of the embedded amphiphilic
Cdots and concomitant quenching of their fluorescence emis-
sion (Fig. 1B, right). The visual and fluorescence photographs in

Fig. 1C depict the bacterially-induced transformation of the
hybrid C-dot–hydrogel. As shown in Fig. 1C, left, the as-
prepared dense gelatinous substance was fluorescent (λex: 375
nm) due to the embedded dispersed Cdots. Following incuba-
tion with a bacterial suspension, the gel material became more
fluid and non-fluorescent (Fig. 1C, right).

Fig. 2 presents microscopic characterization of the C-dot–
hydrogel composite. The representative scanning electron
microscopy (SEM) image in Fig. 2A reveals a highly porous sheet
morphology of the hydrogel displaying considerable internal
surface area. The fluorescence microscopy data in Fig. 2B
confirm encapsulation of abundant Cdots within the hydrogel
matrix. Specifically, the confocal fluorescence microscopy
images in Fig. 2B reveal a highly fluorescent hydrogel scaffold
due to the Cdots bound to the hydrogel surface. In particular,
the different colors of the hydrogel in Fig. 2B reflect the
excitation-dependent fluorescence emissions of the embedded
Cdots.21,22 The excitation-dependent emission spectra of the
hydrogel-embedded Cdots was distinctive from soluble Cdots
(see ESI, Fig. S2†), confirming their encapsulation within the
hydrogel scaffold.

We examined the effect of bacterial suspensions upon the
fluorescence properties of the C-dot–hydrogel. Fig. 3 shows
the fluorescence modulation (λex = 375 nm) induced by addition of
Bacillus cereus (cell concentration 105 cells per mL) to the C-dot–
hydrogel. Fig. 3A reveals significant reduction of the fluores-
cence emission recorded 15 minutes after incubating the
composite gel with the bacteria (concentration of 105 cells per
mL) at 37 °C. The direct relationship between the degree of
fluorescence quenching and bacterial concentration is depicted
in Fig. 3B. Indeed, the dose–response curve in Fig. 3B points to
a detection threshold of around 105 cells per mL reflecting
excellent sensitivity of the C-dot–hydrogel platform.

While the data presented in Fig. 3A and B were recorded
upon incubation of bacterial cell suspensions with the C-dot–
hydrogels, we examined whether the fluorescence quenching
effect was related to molecules secreted by the bacteria. Indeed,
Fig. 3C reveals that the fluorescence quenching effect could be
effectively induced by the supernatant (i.e. after discarding the

Fig. 2 Microscopic images of the C-dot–hydrogel: (A) scanning electron microscopy (SEM) image of the C-dot–hydrogel. Scale bar corresponds to 10 μm. (B) Confocal fluorescence microscopy images of the C-dot–hydrogel. Bright field (i); and confocal fluorescence microscopy images recorded using λex = 405 nm, emission filter EM 445/60 (ii); λex = 488 nm, emission filter EM 525/50 (iii); λex = 514 nm, emission filter EM 525/50 (iv). Scale bar corresponds to 10 μm.
cell precipitate), indicating that molecules secreted by the bacterial cells were responsible for fluorescence modulation. No quenching was recorded upon mixing of bacteria and soluble C-dots (data not shown), confirming that the quenching phenomenon was due to hydrogel disruption rather than direct interactions between the C-dots and bacterial cells.

To account for the bacterially-induced modulation of hydrogel-embedded C-dots’ fluorescence we carried out experiments aimed at characterization of the mechanical gel properties before and after addition of the B. cereus suspension (Fig. 4). Specifically, Fig. 4 depicts rheology analysis of the hydrogel before and after addition of B. cereus (10^8 cells per mL) at 37 °C. The data in Fig. 4 present frequency sweep experiments in which the storage modulus (G’), and loss modulus (G’’) were measured, reflecting the gel-phase of the material in which G’ is greater than G’’. The curves in Fig. 4ii and iv demonstrate that both G’ and G’’ significantly decreased after addition of the bacterial suspensions, reflecting reduction of the hydrogel’s visco-elasticity (e.g. greater fluidity). The difference between G’ and G’’ was retained, however, pointing to retention of the intrinsic gel organization of the C-dot–hydrogel in the frequency range measured.

Nuclear magnetic resonance (NMR) and Fourier transform infrared (FTIR) spectroscopy experiments in Fig. 5 were designed to elucidate the molecular factors responsible for the bacterially-induced fluidization of the C-dot–hydrogel. The ^13C NMR and ^1H NMR data in Fig. 5A and B reveal that the molecules secreted by B. cereus induced distinct chemical transformations within the hydrogel scaffold. Specifically, the ^13C NMR spectra in Fig. 5A demonstrate the occurrence of ester bond cleavage following addition of the B. cereus suspensions, resulting in release of glucose molecules. This is reflected in the appearance of a new downfield ^13C peak at 63 ppm in the B. cereus-treated sample (Fig. 6Aii) corresponding to the C6 atom of free glucose. Similarly, ^1H NMR experiments presented in Fig. 5B provide evidence for bacterially-induced hydrolysis of the ester bonds linking the lauryl side chains (Fig. 1B). Indeed, inspection of Fig. 5B reveals generation of free lauric acid, reflected in the appearance of the Cx-attached hydrogen at 2.25 ppm following addition of the bacteria.

The fluorescence data, mechanical, and spectroscopic analyses in Fig. 3–5 suggest that ester-hydrolyzing enzymes, specifically esterases and lipases secreted by B. cereus are responsible for the structural and luminescent transformations observed in the C-dot–hydrogel system. Esterase constitute a diverse enzyme family secreted by both mammalian and bacterial cells, which function through hydrolysis of ester bonds in target substrates. Notably, the hydrogel scaffold contains a multitude of ester bonds, both positioned in polar environments (e.g. ester bonds linking the glucose residues to the modified tartaric acid-based anhydride moieties, Fig. 1A) which serve as cleavage sites for bacterially-secreted esterases, as well as ester bonds in non-polar regions of the hydrogel (e.g. between the acyl chains and the central anhydride, Fig. 1A), constituting substrates for lipases, considered as esterase subfamily which functions in non-polar lipidic environments.

Fig. 6 presents experiments designed to provide evidence for the assignment of bacterially-secreted esterases and lipases as inducers for hydrogel reorganization and fluorescence quenching of the embedded C-dots. Fig. 6A highlights the effects of heat and ethylenediaminetetraacetic-acid (EDTA) upon fluorescence quenching. Specifically, the fluorescence emission data in Fig. 6A demonstrate that heating the B. cereus
supernatant to 100 °C essentially cancelled the fluorescence quenching effect (long-dash spectrum in Fig. 6A). This result is ascribed to heat-induced denaturation of bacterially-secreted proteins, thereby eliminating enzymatic digestion of the hydrogel scaffold.

Fig. 6A also shows that addition of EDTA to the bacterial suspension reduced fluorescence quenching (short-dash spectrum in Fig. 6A); EDTA binds divalent ions such as Ca$^{2+}$, which are cofactors that are necessary to lipase functioning, thus inhibiting enzymatic cleavage of the hydrogel ester bonds. It should be noted that esterases generally do not require metal cofactors for activity thus the residual quenching still recorded after EDTA addition (Fig. 6A) might provide an indication as to the relative contribution of esterases vs. lipases to the observed C-dot–hydrogel fluorescence quenching.

Fig. 6B depicts quenching of the C-dot–hydrogel fluorescence following addition of commercially-obtained esterase extracted from B. subtilis (B. subtilis belongs to the same subgroup of Bacillus species as B. cereus). The fluorescence quenching apparent in Fig. 6B both confirms the role of esterases in hydrogel disruption, as well as the fact that the bacterially-induced quenching phenomena reported here involve contributions from both esterases and lipases.

The C-dot–hydrogel fluorescence quenching assay was applied for analysis of several bacterial species (Fig. 7). Specifically, the graph depicted in Fig. 7 illustrates the degree of fluorescence quenching induced by selected bacteria (10$^8$ cells per mL) added to the C-dot–hydrogel. The significant differences in quenching magnitudes highlighted in Fig. 7 likely reflect the variability of esterase/lipase secretions by the bacteria tested. Importantly, while some esterase and lipase enzymes secreted by bacteria have been identified, the complexity and diversity of these enzyme families have been recognized and manifested in many studies; in this context, the C-dot–hydrogel fluorescence assay provides a simple platform for evaluating the overall secretion of these enzymes by bacteria. Moreover, the distinct quenching degrees could be employed for distinguishing and identification of different bacteria.
Conclusions

We have developed a new hybrid carbon-dot (C-dot)–hydrogel system capable of detecting bacteria through quenching of the C-dots fluorescence. Bacterial sensing was demonstrated in case of *Bacillus* and *Staphylococcus* strains, although the detection concept is generic and might be implemented for other bacterial species. The sensing mechanism is based upon cleavage of ester bonds within the hydrogel scaffold, induced by esterases and lipases secreted by the bacteria. Cleavage of the ester bonds disrupted the hydrogel framework, resulting in fluidization and aggregation and quenched fluorescence of the embedded C-dots. A series of physical and chemical experiments provide evidence for the gel transformation and reaction mechanism. The C-dot–hydrogel matrix enabled both high-sensitivity detection of bacteria (*B. cereus*) as well as distinguishing among different bacteria through the extents of fluorescence quenching.

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