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Balance of photon management and charge collection from carbon-quantum-dot layers as self-powered broadband photodetectors

Highly sensitive, rapid-response and self-driven photodetectors based on single-layer carbon QD layers (CQDL) are visualized, and they demonstrate a remarkable detectivity above 5.9×10^{12} under broadband light illumination.

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Balance of photon management and charge collection from carbon-quantum-dot layers as self-powered broadband photodetectors†

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Semiconductor colloidal quantum dots (QDs) have been regarded as promising fluorescent materials for chemical sensing, bio-detection and optical communications; yet it still remains challenging to bring out self-powered photodetectors based solely on QDs because the excited charges within QDs are extremely immobile due to their reduced dimensionalities and they hardly form effective photocurrents. Hence, we have attempted to decouple the light-absorption and charge-transport criteria in order to feature highly-sensitive, rapid-response and self-driven photodetectors based on single-layer carbon QD layers (CQDLs) *via facile in situ* self-assembling deposition with fine control over thickness. We show explicit dark-current suppression by visualizing charge blocking phenomena and screen effects due to layered CQDL structures, which alleviate the movement of leakage carriers crossing over the CQD interlayers. By examining the distribution of electric fields within CQDLs under light excitation, the spatial dependence of the light-trapping effect within CQDLs was confirmed. These features are strongly associated with the thickness tuning of CQDLs, while 65 nm of CQDL thickness could manifest remarkable photoresponsivity above 9.4 mA W⁻¹ and detectivity above 5.9 × 10¹² under broadband light illumination. These results demonstrate the insights gained from an understanding of broadband optoelectronics, which might potentially pave the way for further employment in functional photodetection.

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Introduction

Solution-processed fabrication has been regarded as a potential strategy that paves the way toward the advanced development of functional devices and applications due to the integration of compatibility with commercial microelectronics, low cost and achievable large-area synthesis.^{1–5} With respect to photo-detection, the involvement of a monolayer of graphene has emerged as a compelling solution that does not merely allow efficient sensing performances covering a wide range of illumination wavelengths, but provides further methods for the realization of a flexible photodetection platform due to its ultrathin nature and high mobility (up to 10⁵ cm² V⁻¹ s⁻¹).⁶ Nevertheless, its inherent limitations set against its practical applications correlated with diminishing effective bandgaps and large dark currents, resulted in a low detection responsivity of 10⁻³ to 10⁻¹ A W⁻¹.^{7,8} To overcome these demanding

problems, a hybrid strategy combining light sensitizers with desirable band gap energy and graphene has been extensively investigated. Several promising photo-absorbing media, such as ZnO,⁹ TiO₂,¹⁰ Si,¹¹ and MoS₂, have emerged in graphene-based photodetectors and been found to greatly improve sensing performances.¹² However, the involvement of complicated heterointerfaces between graphene and sensitizer materials created sophisticated charge-transfer characteristics across graphene/sensitizer interfaces,¹³ and thus more in-depth investigations still need to be undertaken.

In addition, the effective thickness of light-absorptive nanostructures should be extremely thin to pursue the efficient extraction and collection of photogenerated carriers due to limited carrier lifetime and diffusion length.¹⁴ These features raised the issue of fabrication complexity and further restricted their applicability for the large-scale synthesis and preparation of sensing devices. Moreover, it remains highly challenging to uniformly and tightly adhere sensitizer nanostructures to graphene in a reliable way. These synthesis problems will inevitably cause friction with their mechanical robustness and long-term operational stability for photosensing operations. There is, therefore, a high demand to explore the facile utilization of solely carbon-based nanomaterials while retaining the capabilities of both efficient photon utilization and charge

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transport. Therefore, in this study, thickness-controllable carbon-quantum dot layers (CQDLs) were rationally formed, featuring an effective light-absorption and electron-collection layer that could substantially improve the photodetection capabilities of Si-based photodetectors. Facile *in situ* self-assembling deposition formed a CQDL with a tailored layer thickness which behaved as a self-powered photodetector, manifesting a remarkable photoresponsivity above 9.4 mA W^{-1} , and detectivity above 5.9×10^{12} under broadband-light illumination, which were among the best values of reported single-QD based photodetectors. Explorations of the spatial dependence of light-trapping effects on layered QD structures were performed to elucidate the underlying broadband aspects of the photodetection performance.

Experimental details

Preparation of the thickness-controllable CQDLs

Glucose powder (purity = 99.5%) without further purification was calcined in an autoclave at $400 \text{ }^\circ\text{C}$ for 2 h. After that, the as-prepared samples were finely ground into fine powders, and then dispersed in deionized water under gentle sonication treatment for 1 h. Subsequently, the CQD solutions were filtered with a standard membrane with a pore size of $0.22 \text{ }\mu\text{m}$, and then dried at $70 \text{ }^\circ\text{C}$ in air. Monodispersed CQDLs on Si substrates were then prepared by drop coating while the thickness of the deposited CQDL could be adjusted with cycles of drop coating; then the samples were annealed in air at $100 \text{ }^\circ\text{C}$ for 30 min. To fabricate the photodetectors, 100 nm Ag and 150 nm Al as top and bottom electrodes, respectively, were directly deposited on CQDL surfaces using electron gun evaporation.

Characterization

The microstructures and surface morphologies of the CQDs were characterized with transmission electron microscopy (TEM, JEM-2100F) and scanning electron microscopy (SEM, Hitachi S-4800), respectively. The functional features of CQDs were analyzed with a Fourier transform infrared (FTIR, PerkinElmer Frontier) spectrometer. Crystallographic examinations were conducted with an X-ray diffractometer (Bruker AXS GmbH) using Cu-K radiation as the X-ray source. Photoluminescent characterizations were conducted with a photoluminescent spectral system equipped with a light-emitting diode lamp (output power 780 mW and wavelength of 365 nm). X-ray photoelectron spectroscopy (XPS, PHI 5000 Versa Probe) was utilized to understand the surface compositions and chemical states through a monochromatic X-ray source ($400 \text{ }\mu\text{m}$ spot size, 36 W). Light-reflection spectra were measured with a UV-vis-NIR spectrometer (Hitachi U-3900H). Spatial field distributions of the devices under various light illuminations were modeled with Finite-Difference Time-Domain (FDTD) simulation.

Photocurrent characterizations were performed in a source meter (Keithley 2400), where LED light sources with various wavelengths, including 352 , 580 and 850 nm , were utilized. Examinations of transient photoresponse employed the

chronoamperometry method under pulsed monochromatic light through variation of light intensity.

Results and discussion

Characterizations and synthetic mechanism of CQDL

Fig. 1(a) displays a cross-sectional TEM view of the formed CQDL, where closely packed CQDs without the creation of voids or structural defects *via in situ* self-assembled layer deposition were confirmed, visualizing the successful formation of regularly layered structures rather than distributed quantum dots. This arrangement of features played an important role in charge collection and transport that responded with sound photo-detection characteristics. In addition, the microstructures constituting CQDs were analyzed with HRTEM examination, as shown in Fig. 1(b). It could clearly be observed that the cores of CQDs showed good crystallinity, showing the (002) lattice of graphite with a crystallographic fringe of 0.348 nm . It should be noted that the resulting fringe spacing was slightly larger than the typical value for a graphite lattice (0.345 nm),^{15–17} which will be discussed later. Apart from that, the distinctive features observed in CQD shells might indicate the existence of abundant functional groups on CQD surfaces with a noncrystalline nature. To further identify them, FTIR analysis was conducted, as demonstrated in Fig. 1(c). The results indicated surface features containing OH- functional groups located at wavenumber 3400 cm^{-1} , which explained the good water solubility of the CQDs. In addition, the spectral dips at 1640 , 1360 , 1027 and 672 cm^{-1} were assigned to the stretching vibrations of C=O, C=C, C-O and out-of-plane bending vibration of CH_2 , which provided the affinity of carbon-to-carbon adhesion and contributed to the dense formation of CQDLs.^{18–20} Fig. 1(d) demonstrates the XRD patterns of various carbon-based

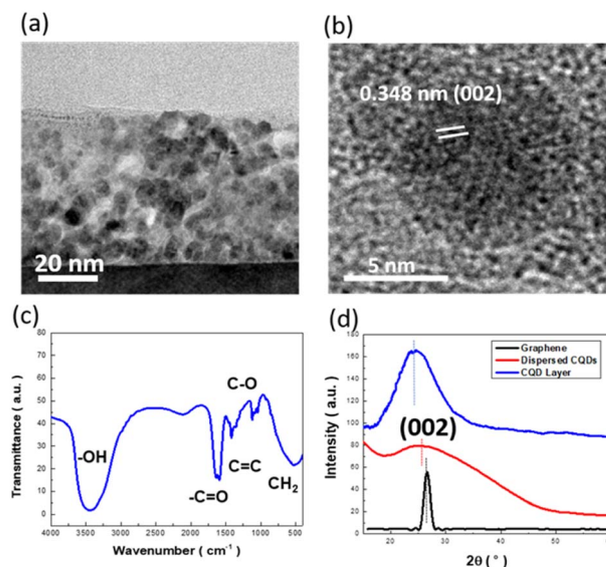


Fig. 1 (a) Cross-sectional TEM image of CQDL directly formed on Si substrate. (b) HRTEM image of a representative CQD. (c) FTIR results of CQDL with a thickness of 65 nm . (d) XRD patterns of graphene, dispersed CQDs and CQDL.



nanostructures: multiple-layer graphene, a dispersed CQD suspension and as-formed CQDLs. The sole diffraction peak of graphene located at a Bragg angle of 26.6° corresponding to the (002) graphite configuration could be observed. Although the similar diffraction facets of the (002) graphite lattice with comparably broad bandwidth existed in dispersed CQDs, one could see that the diffraction pattern was slightly shifted to smaller Bragg angle (25.6°). These results indicated the fact that CQD surfaces containing COO⁻ features as electron-withdrawing groups can capture electrons from CQD cores, thus slightly expanding the core graphite lattice and giving rise to a reduction in the diffraction angle of the (002) graphite configuration by increasing the corresponding lattice spacing. Interestingly, in closely packed CQDLs, a comparably substantial shift in diffraction angle reaching 24.6° was encountered. These findings could be interpreted by the fact that the surficial COO⁻ features with electron-withdrawing affinity could further capture electrons from neighboring CQDs due to their densely attached configuration. Thus, the lattice dimensions of CQD cores were forced to expand to 0.348 nm [Fig. 1(b)] that reflected the obvious reduction in correlated Bragg angle. The correlated crystallographic examinations from both HRTEM and XRD analysis confirmed the feasibility of varying CQD-based structures through transition from distributed features toward arranged layered structures.

Explorations of thickness control over CQDL preparation are displayed in Fig. 2(a). Our findings disclose the reliability of tuning the CQDL thickness by adjusting the cycles of drop coating, where the linear regression of CQDL thickness with respect to the cycling depositions with R^2 of 0.998 could be observed. These characteristics displayed sound control over CQDL thickness for the optimization of photodetection capabilities. Another striking property of the CQDL synthesis correlated with consistent film crystallinity by varying the CQDL thickness, as shown in Fig. 2(b). Raman peaks located at 1425 and 1581 cm^{-1} could be observed, corresponding to the D band and G band of the carbon structures,^{21–24} respectively, whereas approximately similar intensity ratios, in terms of I_D/I_G , with a value of 0.66 were in evidence, regardless of a CQDL thickness ranging from 10 to 100 nm. These results indicated sound reliability for further engineering CQDLs in their practical employment in functional optoelectronic applications.

On the basis of the microstructural, chemical and crystallographic information attained from detailed material

characterizations, we attempted to provide insights into the possible mechanism for CQDL formation *via in situ* self-assembled deposition, where the correlated schematic illustrations are presented in the ESI.† When the drop of CQDs was in contact with Si substrates, a convex surface of the CQD suspension facing Si was formed, which brought about the faster evaporation of the water solvent at the edge of the liquid drop than in the center. This induced a radial outward convective flow from the center along with CQDs to compensate for the partial loss of water solvent due to instant evaporation at the periphery. Moreover, two types of capillary force were created by the deformation of the liquid/air interfaces facing in opposite directions, hydrostatic pressure and surface tension, that were exerted on CQDs at the boundary of the Si substrate and tended to push the self-assembly layered formation of CQDs. While residual water continued to evaporate, the balance of forces was broken, leading to a black surrounding ring formed from an aggregation of CQD clusters at the edge of the substrate surface. This phenomenon has also been observed and interpreted from another two-dimensional ordering of polystyrene sphere nanoparticles by N. Denkov *et al.*²⁵ Afterwards, a concave meniscus surface of the water solvent forced the CQDs to spread out toward the center of the Si surface. It should be noted that another capillary force, termed sucking capillary pressure, was taking place due to the inherently high hydrophilicity, which mediated the radial inward capillary flux that drove the movement of CQDs toward the unoccupied regions, whereas the involvement of capillary attraction captured the dispersed CQDs towards the ordered region of CQDs.^{26,27} With such layered formation of CQDs and liquid level drawdown approximately toward the diameter of the CQDs, the hydrostatic pressure and surface tension force were exerted again and applied to an inward convective flow, leading to the multilayered film formation constituted by CQDs.

The spectral light-reflection characteristics of CQDLs with various thicknesses are displayed in Fig. 3(a). To clarify the underlying light/matter interaction, FDTD examinations on light reflectivity were also performed, as shown in Fig. 3(b). It was explicitly found that the experimental results corresponded well with the calculated spectra for bare Si, graphene/Si and five different CQDL/Si. Specifically, the measured average light reflectivity (R_{av}) of CQDL/Si was found to be fairly low for CQDL thicknesses of 13 nm ($R_{av} = 28.0\%$) and 26 nm ($R_{av} = 23.8\%$), where the substantial suppression of light reflectivity was validated compared with either bare Si ($R_{av} = 40.1\%$) or graphene/Si ($R_{av} = 36.0\%$) structures, indicating the extraordinary light trapping and utilization capabilities of CQDLs. In addition, the correlated R_{av} could be further reduced to 17.7%, 16.2% and 13.2% in the cases of 39 nm, 65 nm and 91 nm in CQDL thickness, respectively. All these results essentially reveal the broadband light capture/utilization of CQDLs employed in broadband photosensing applications. To elucidate these findings, the electric-field distributions of CQDL (thickness 65 nm)/Si were calculated under three different light illuminations, of 352 nm, 580 nm and 850 nm, as demonstrated in Fig. 3(c). It could be speculated that the spatial distributions of light trapping effects induced by CQDLs depended on the

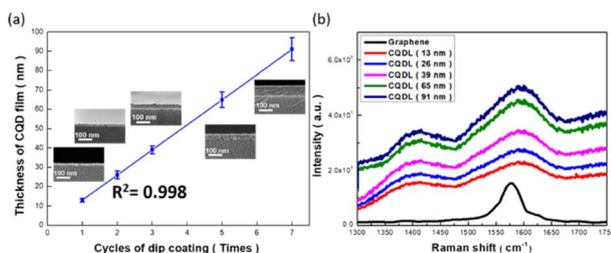


Fig. 2 (a) Correlation of CQDL thickness with respect to the cycles of drop coating. (b) Raman results of CQDL with variations in layer thickness.



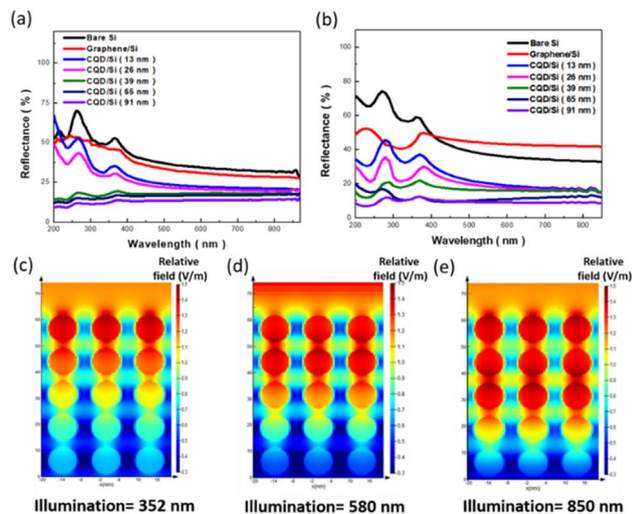


Fig. 3 (a) Measured and (b) simulated light-reflection spectra of various samples. FDTD examinations of electric-field distributions of CQDLs (65 nm in thickness) under three various light illuminations: (c) 352 nm, (d) 580 nm and (e) 850 nm.

illumination wavelengths. In general, under short-wavelength (352 nm) illumination, the excited electric fields were substantially confined in the top portions of the CQDLs within the upper two layers of CQDLs owing to the low penetration depth of the incident light. In addition, the regions for field localization of CQDLs extended spatially from the top to the middle range (approximately 1–3 stacked layers and mainly in the 2nd and 3rd stacked layers) and from the top to the bottom range (approximately 1–5 stacked layers and mainly in the 4th and 5th stacked layers) of CQDLs under 580 nm and 850 nm light illumination, respectively. These results explained that the broadband light capturing effects indeed arose through the spectral superposition of light-absorption bands due to the establishment of layered CQD structures rather than the light-induced direct interband transition of individual CQDs, which provided sound evidence for the reflection measurements shown in Fig. 3(a).

Photodetection performance

By simply integration with electrode pairs as Ag/CQD/Si/Al sandwiched structures, the CQDL-based photodetectors were created without any sophisticated surface treatment or introducing any additional photonic/collection components. The correlated dark current (I)-voltage (V) relationships of various CQDL-based devices were examined, as shown in Fig. 4(a), and the evaluations of dark currents, and ratio of reverse saturation current and forward/reverse current without the employment of an external bias are summarized in the ESI.† In addition, the dark I - V results of graphene/Si-based devices were also compared. We found that the graphene/Si samples possessed the largest existing dark current (6.02×10^{-7} A) and reverse saturation current (6.67×10^{-3} A), which were an order of magnitude larger in both dark and reversed saturation currents than the correlated results of all CQDL/Si devices. These

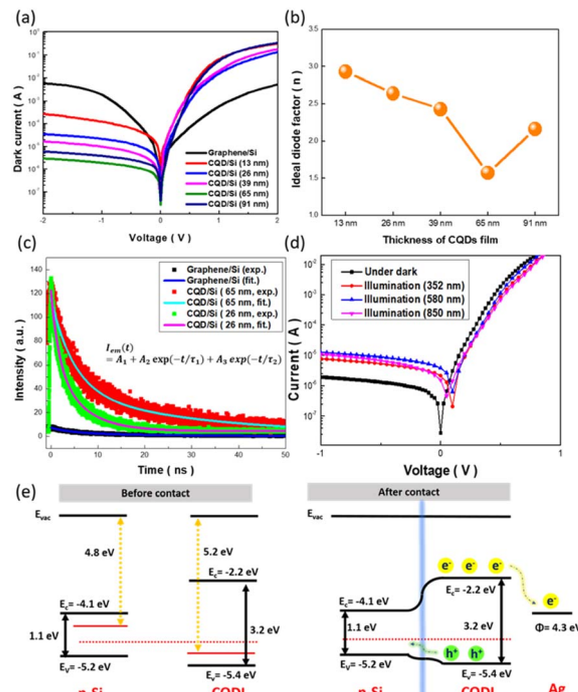


Fig. 4 (a) Dark I - V curves of various CQDL/Si and graphene/Si-based devices. (b) Extracted ideal diode factor with respect to the thickness of the CQDL. (c) TRPL examinations of graphene/Si, CQDL (26 nm)/Si and CQDL (65 nm)/Si. The correlated exponential fittings are also shown. (d) Light-driven I - V results of CQDL (65 nm)/Si under various light illuminations. (e) Schematic presentation of band diagrams before and after CQDLs coming into contact with Si substrates.

features could be attributed to the high carrier concentration and large charge mobility inherently existing in the highly conductive graphene layer, which thus inevitably contribute to the service leakage currents that could significantly degrade the photodetection performance.²⁸ By further comparing the device performances of CQDL/Si under dark conditions, the forward/reverse current ratios, functioning as an evaluation of the rectifying behaviors of heterojunction contacts, and depending distinctly on CQDL thickness, are displayed in detail in the ESI.† Specifically, the results indicated that with a CQDL thickness of 13 nm, the corresponding ratio for forward/reverse current reached 527, where the findings closely corresponded to those in the reported literature.²⁹ Nevertheless, the optimal rectifying condition from CQDL/Si photodetectors was achieved when the CQDL thickness increased to 65 nm, reaching a superior ratio of 2.96×10^4 , and then gradually decreased to 1.58×10^4 when the CQDL thickness was 91 nm. These features visualized the possible charge blocking phenomena arising from the vertical stacking of CQDs that could facilitate the inhibition of leakage formation reaching the electrode sides. Another possible mechanism that could assist leakage inhibition came from the screening effect from layered CQDLs, because similar charge concentrations appeared in different layered CQDLs and thus could not induce the movement of carriers crossing over the interlayers due to the alleviated driving forces under dark conditions.³⁰



Another striking piece of evidence regarding the diode behavior of the CQDL/Si heterojunction could be understood through examination of the ideality diode factor, presented as follows,^{31–33}

$$n = qdVKT \ln I \quad (1)$$

where q is the charge of a dark electron, K is the Boltzmann constant, V is the voltage employed in the diode device and I is the current formed. The comparative results are demonstrated in Fig. 4(b), showing that the ideality diode factor of CQDL/Si devices depending on CQDL thickness was around 2.9 when the CQDL thickness was designed as 13 nm, and then gradually reduced to 1.3 when the CQDL thickness was engineered to 65 nm. It has been reported that a resulting ideality diode factor approaching 1 indicates an ideal diode performance that facilitates the rectification of charges, where the results demonstrated here agree well with the indication from the forward/reverse current ratio presented in Fig. 4(a). We also note that when the CQDL thickness was too thick (91 nm), the clear offset of the ideality diode factor from 1 implied that the rectifying capability was weakened because the mobile charges within CQDL layers close to electrode pairs tended to readily diffuse into electrode sides rather than across thick CQDL interlayers, thus causing the unwanted device noise.

In addition, the measured results of time-resolved PL traces at room temperature for the emission peak are displayed in Fig. 4(c). Based on the spectral deconvolution with respect to the PL response, a double exponential equation, $I = A \exp(-\tau_1) + B \exp(-\tau_2) + C$, was utilized for the fitting.^{34,35} Accordingly, τ_1 (τ_2) were extracted as 4.95 ns (8.8 ns), 4.77 ns (28.8 ns) and 4.86 ns (16.7 ns) from graphene/Si, CQDL (65 nm)/Si and CQDL (26 nm)/Si, respectively, which were regarded as nonradiative and

radiative lifetimes, respectively. A remarkable improvement in radiative carrier lifetime in the case of CQDL (65 nm)/Si could be observed, validating the sound heterojunction involved in the efficient separation of photogenerated carriers that prevented direct charge recombination. In addition, the correlated band diagram is shown in Fig. 4(e), while experimental examinations of the band diagram are demonstrated in the ESI.† When the CQDL was in contact with an n-type Si substrate, interfacial band bending was caused to balance the Fermi-level difference in both sides. Specifically, the edge of the valence band in the CQDL bent upward, which energetically favored the driving of photogenerated holes across interfaces to reach the Si side, which benefited charge separation and in turn avoided direct recombination. On the other hand, a large deviation (2.1 eV) of the conduction band minimum in the CQDL and the work function of the Ag electrode was encountered compared with the energy difference between the conduction band edges of Si and CQDL, termed $\Delta E_c = 1.9$ eV. Such circumstances involve a larger rate constant for electron transfer from the CQDL toward the Ag electrode, thus preferentially driving the movement of photogenerated electrons for the generation of photocurrents. Notice that the large potential difference involved could even attain photocurrent formation even when no external bias was employed, and therefore the CQDL/Si could potentially be operated *via* a self-powered configuration.^{36,37} These features were evidenced in the measurements of I - V curves under various light illuminations, as shown in Fig. 4(d). Compared with I - V phenomena under dark conditions, remarkably enhanced photoexcited currents covering all the sweeping voltages from -1 to 1 V appeared under the detection of a broad band of light, including 352, 580 and 850 nm.

To evaluate the response time under broadband illumination, repetitive on/off cycles were also recorded, as

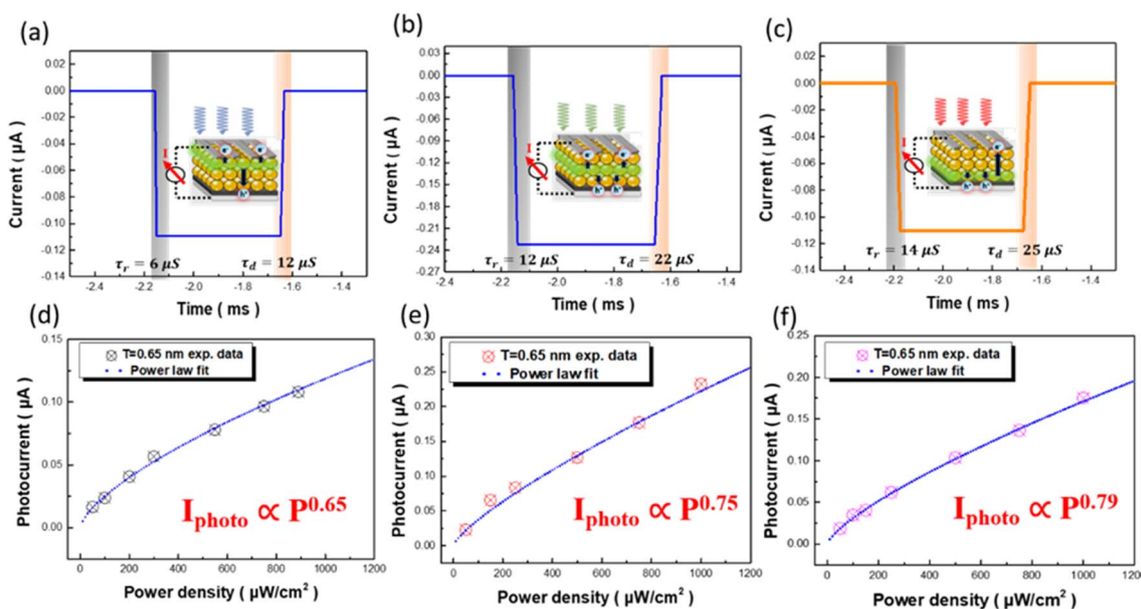


Fig. 5 On/off switching of photoresponse for CQDL (65 nm in thickness)/Si photodetectors under light illumination with wavelengths of (a) 365 nm, (b) 580 nm and (c) 850 nm. Relationship between power density of illuminated light with respect to the excitations of photocurrents under illumination wavelengths of (d) 365 nm, (e) 580 nm and (f) 850 nm.



demonstrated in Fig. 5(a)–(c). Specifically, both the rise time and decay time, defined as the time required for the response to rise from 10% to 90% of the on/off switch peak, and the time required for the response to decay from 90% to 10% of the switch peak, respectively. Accordingly, the evaluated rise time/decay time of devices under 580 nm illumination [Fig. 5(b)] were 12 μ s/22 μ s, whereas both time parameters turned out to be shorter [6 μ s/12 μ s, Fig. 5(a)] and longer [14 μ s/25 μ s, Fig. 5(c)] under 365 nm and 850 nm illumination, respectively. It has been reported that a shorter rise time in terms of switching response correlates with effective photoexcitation and shortened transport length of carriers, and a longer decay time is strongly associated with a reduced probability of carrier recombination, which could be facilitated by charge separation arising from the existence of a heterojunction.

Therefore, we speculated that the distinct dynamic photoexcitation processes of CQDLs depending on illumination wavelengths could be interpreted with the schematic illustrations shown in the insets to Fig. 5(a)–(c), respectively. Under short wavelengths [Fig. 5(a)], the active region for light-absorption/photoexcitation lay in the upper region of the CQDL (highlighted in green) on the basis of FDTD examinations [Fig. 4(c)], and therefore the involvement of rising photocurrent was rapid and responded to the fairly short rise time (6 μ s); in contrast, the diminished charge separation, owing to the fact that the existing active CQDL region was far away from the CQDL/Si heterojunction, increased the charge recombination within the CQDL which was in turn reflected in the comparably short decay time. These features could be further supported by the TRPL explorations on thinner CQDL (26 nm)/Si samples, where the reduced radiative carrier lifetime compared with the result of a 65 nm thick CQDL coupled with Si was in evidence, as shown in Fig. 4(c). On the other hand, under long wavelength [Fig. 5(c)], the relatively long transport length for photo-generated electrons to reach the upper electrode and effective hole separation toward the Si side were due to the fact that the active CQDL region for long-wavelength light trapping lay in the proximity of CQDL/Si interfaces. These caused the results of a relatively long rise time and decay time, respectively, in such a spectral situation. Aside from these two spectrally operating edges, one could observe the balance of photoelectron collection and hole separation coming from light illumination with a wavelength of 580 nm, which thus resulted in intermediate rise/decay dynamics compared with short- and long-wavelength excitation. In addition, we also found that the device responsivity (R) and detectivity (D) for sensing 580 nm light reached the highest values among three tested spectral ranges, as detailed in the ESI.† Yet, all the device performances in terms of R and D remained remarkably sound for employment in broadband photodetection applications.

To explore the detection capabilities according to the power intensity of the incident light, the measured photocurrents at zero bias with respect to light power ranging from 0 to 1400 mW cm^{-2} were examined under three different light illuminations, as shown in Fig. 5(d)–(f), respectively. Mathematical fitting, in terms of power law, was employed to elucidate the underlying

power-dependent photosensing characteristics, as presented below,^{38,39}

$$I_{\text{photo}} \propto P^r \quad (2)$$

where I_{photo} is the measured photocurrents, P is the power density of the light, and r indicates the trapping and recombination processes of the carrier in light-driven devices.⁴⁰ Accordingly, the extracted exponents, r , were found to be 0.65, 0.75 and 0.79, where all the results were within the range $0.5 < r < 1$ indicating reliable photodetection performances as r is close to 1 representing the ideal photodiode design. It should be pointed out that the value of r was lowest under 365 nm illumination among the three tested conditions, which could be associated with relatively obvious charge recombination occurring in the CQDL in such situations, where these findings were fairly well explained by the far distance of the active CQDL region from the CQDL/Si heterojunction, as evidenced in Fig. 5(a).

Device reliability

Finally, the environmental reliability of the photodetectors was examined, as shown in Fig. 6. Long-term operational tests were performed by conducting 2000 on/off cycles, as shown in Fig. 6(a). The consistent on/off switching without the creation of photocurrent hysteresis within each switch again confirmed the instant signal creation/recovery responding to the spectral variations of the detected light. Aside from that, two practical device parameters, the normalized photocurrent to dark current ratio (NPDR) and noise equivalent power (NEP), were estimated, which clearly validated the maintenance of device reliability for broadband detection, as shown in the ESI.† The influence of environmental temperature on CQDL/Si-based devices was explored by monitoring the device responsivity under a wide temperature range from 0 $^{\circ}\text{C}$ to 400 $^{\circ}\text{C}$, as shown in Fig. 6(b). It seemed quite promising that the photodetectors could sustain abrupt temperature variations while maintaining a highly stable photosensing response with a slight deviation of less than 1%. These findings indicated the involvement of a quenching effect for the thermal activation of QD architectures, which caused phonon scattering at interparticle edges and thus tended to initiate the non-radiative decay of thermally excited charges that readily diminished the external thermal influences.⁴¹ Fig. 6(c) presents examinations of the

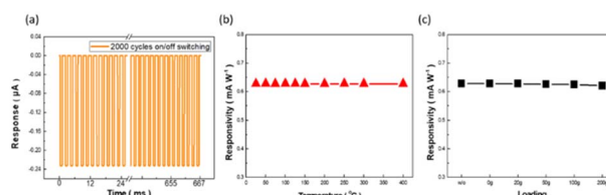


Fig. 6 (a) Long-term on/off switching performances of CQDL (65 nm in thickness)/Si photodetectors. Device responsivity of CQDL (65 nm in thickness)/Si photodetectors by examining the influence of (b) environmental temperature and (c) normal loadings on CQDL sides.



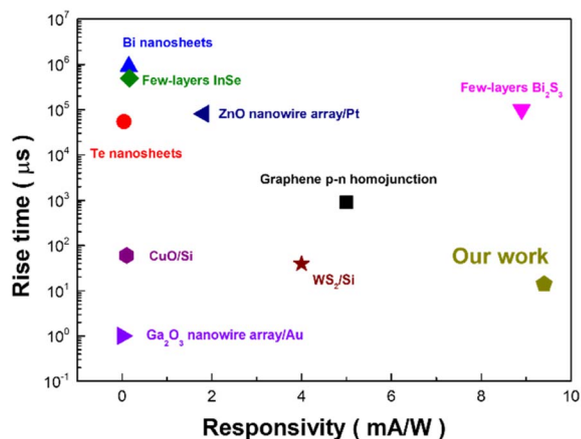


Fig. 7 Comparison of device performance of broadband photodetectors in terms of rise time and specific responsivity.

photodetectors under abrasion, where the CQDL/Si devices were subjected to various weight loadings from 0 to 200 g and the CQDL sides were directly polished on a piece of PET film for a length of 50 cm, respectively. The device responsivity recorded with respect to the employment of loading weight explicitly suggested reliable photodetection performances with a trivial reduction in responsivity values of less than 2%, which could be attributed to structural robustness arising from densely layered QD films and approximately defect-free CQDL/Si interfaces. Overall, the results clearly validated the environmental reliability and wearing capability of the devices for potential use in harsh conditions. To shed light on the improvement in device performance, two critical photodetection parameters, the rise time and R of solely nanostructure-based self-powered photodetectors were compared, as shown in Fig. 7, and the detailed results are summarized in the ESI.† The remarkable improvement in device R (9.4 mA W^{-1}) among all the comparative self-powered devices is evident. In addition, the present rise time of CQDL photodetectors in the order of 10^{-5} s also fell at the top level compared with those in the reported literature. Thus, we would point out that the CQDL/Si self-powered photodetectors could provide excellent sensing responsivity and promising rapid photoresponse dynamics, which might have quite a potential impact on the advanced development of ultrahigh-speed optical communications and further broaden the use of photosensing applications.

Conclusions

Novel self-powered photodetectors based on thickness-tunable CQDLs were readily manufactured *via* facile solution deposition. By performing detailed examinations of carrier kinetics, band structures, power-dependent photoresponse characteristics and operational reliability, we found that the CQDL/Si could manifest remarkable photoresponsivity above 9.4 mA W^{-1} and detectivity above 5.9×10^{12} , and exhibited extremely rapid response with rise time/decay time of $<14 \mu\text{s}/<25 \mu\text{s}$ under broadband light illumination. The superior photodetection

performances could be attributed to the synergistic contributions from spatial trapping effects within CQDLs and the established heterojunction that facilitated charge separation, which involved a larger rate constant for electron transfer from a CQDL toward an Ag electrode, thus driving photocurrent generation without the employment of an external bias. Thus, this work could have an impact on the development of single-layer-based photodetectors, and provide further inspiration for other functional photodetection applications.

Conflicts of interest

There are no conflicts of interest to declare.

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