Sub-Millisecond Response Time in a Photorefractive Composite Operating under CW Conditions

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Extensive study of photorefractive polymeric composites photosensitized with semiconductor nanocrystals has yielded data indicating that the inclusion of such nanocrystals enhances the charge-carrier mobility, and subsequently leads to a reduction in the photorefractive response time. Unfortunately, the included nanocrystals may also act as a source of deep traps, resulting in diminished diffraction efficiencies as well as reduced two beam coupling gain coefficients. Nonetheless, previous studies indicate that this problem is mitigated through the inclusion of semiconductor nanocrystals possessing a relatively narrow band-gap. Here, we fully exploit this property by doping PbS nanocrystals into a newly formulated photorefractive composite based on molecular triphenyldiamine photosensitized with C60. Through this approach, response times of 399 μs are observed, opening the door for video and other high-speed applications. It is further demonstrated that this improvement in response time occurs with little sacrifice in photorefractive efficiency, with internal diffraction efficiencies of 72% and two-beam-coupling gain coefficients of 500 cm−1 being measured. A thorough analysis of the experimental data is presented, supporting the hypothesized mechanism of enhanced charge mobility without the accompaniment of superfluous traps. It is anticipated that this approach can play a significant role in the eventual commercialization of this class of materials.

Owing to their large optical nonlinearities, low permittivity and low cost, organic photorefractive (PR) composites are potentially useful in a variety of real-time optical applications1–5. Polymer-based PR composites are attractive due to the ease with which their constituents may be independently modified, allowing the composite to be tailored for a specific application. Despite the flexibility associated with organic PR composites, the lack of organic photosensitizers exhibiting sufficiently large photo-charge generation at near infrared wavelengths, λ, proved to be a significant hurdle. It was primarily this challenge which inspired the photosensitization of otherwise all-organic PR composites through the inclusion of semiconductor nanocrystals, also known as quantum dots or Q-dots6–23. Although motivated by the ability to extend the spectral range accessible to this class of materials, most studies have been conducted at visible wavelengths, primarily λ = 633 nm, using Q-dots such as nano-sized CdS (QCdS) as well as QCdSe and QCdTe6,7,9–14,17,20–22. Although far fewer examples exist, narrower band-gap semiconductor materials such as QPbS or QPbSe have been generally used for λ such as 1.31 μm! and 1.51 μm, however, reported efficiencies and response times have been substandard relative to those reported for PR composites photosensitized with more traditional organics, most notably C608,15,16,23.

In addition to acting as a photosensitizer, a significant amount of experimental data, particularly from time-of-flight characterizations, indicate the included Q-dots also increase the charge-carrier mobility, μ, within the PR composite16. This enhancement in μ is primarily attributed to the ability of the free charge-carriers to enter into, and subsequently be transported through the included Q-dots, where they experience a faster μ relative to that attributed to the organic charge-transporting species. Not surprisingly, it has also been demonstrated that this improvement in μ may be accompanied by a reduction in the PR response time, τ17,18,23. Notwithstanding the enormous insights gained into the fundamental processes of charge-generation and charge-transport occurring

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In inorganic–organic hybrid composites, there has existed an inability to capitalize on these understandings with, until recently, $\tau > 100$ ms representing the best PR response time reported for an organic composite photosensitized with Q-dots\(^{17,23}\). This $\tau$ is significantly larger than those reported for PR composites photosensitized with organic photosensitizers such as C\(_{60}\)\(^{24-35}\). Furthermore, for studies involving visible $\lambda$, the data suggest that while the inclusion of Q-dots introduces a secondary charge-transport species imparting enhanced $\mu$, they may also act as a trapping species\(^{17,18,23}\). While a certain concentration and depth of traps is necessary for the PR effect, excessive trapping of positive free charge-carriers can have a detrimental influence over the $\tau$, operational voltage, $E$, as well as the overall diffraction efficiency\(^{1-3,36}\). In an effort to circumvent this issue, a recent study investigated the photosensitization of a PR composite at $\lambda = 633$ nm through the inclusion of QPbS where the QPbS exhibited significant, but off-resonance, absorption\(^{21}\). In a separate study it was established that by exchanging the relatively wide band-gap QCdSe with QCdTTe, which possesses a comparatively narrower band-gap, an improvement in the $\tau$ was realized\(^{17}\). These studies provided strong evidence that in addition to the photosensitization of PR composites, Q-dots may alternatively be used specifically for enhancing the $\mu$ and thereby substantially reducing the $\tau$ of a PR composite.

In this communication we present a novel PR composite specifically designed to exploit the insights gained with regard to the influence of Q-dot inclusion over charge-carrier $\mu$ and subsequent reduction of the $\tau$. In many cases, organic PR composites described in literature use poly(N-vinylcarbazole) (PVK) due to its efficient hole-transporting capability. More recently PR composites based on polymers which replace the charge-transporting carbazole group with that of triphenylidine (TPD), or similar structures, have been described\(^{26-33}\). Of these polymers perhaps the most notable is poly(acrylic tetraphenylidamino-phenyl) (PATPD) which is composed of a TPD molecule attached to acrylic backbone through a pendant linkage\(^{28,30-33}\). In this study, a molecular form of TPD acts as the primary organic charge-transport species. Through this approach, the inert backbone, as well as the pendant linkage associated with polymerized forms of TPD can be eliminated. Furthermore, there is no need for the addition of an inert plastiziser, greatly increasing the loading content of the inert backbone, as well as the pendant linkage associated with polymerized forms of TPD can be eliminated.

This has been shown, however, that the elimination of shallow traps can result in an enhancement in the PR $\tau$\(^{38}\). In addition to the charge-transporting molecular TPD, the composite described herein also contains the non-linear optical (NLO) dye 4-homopiperidinobenzylidenemalononitrile (7-DCST) providing the electro-optic activity required for the PR effect\(^{42}\). 7-DCST has been used in conjunction with PVK as well as PATPD and is a common choice of NLO dye for PR composites in which the optimization of $\tau$ is of interest\(^{24,25,30-36}\). Solid phase crystallization of the molecular TPD as well as that of the 7-DCST, also well known for its susceptibility to phase-separation, was successfully suppressed through the inclusion of a small amount of polymer. Rather than employ a functionally passive polymer, it was found that PVK in concentrations as low as 10 wt% was especially effective at preventing detectable aggregation of the molecular constituents. Since the primary goal of this study was to study the performance of a PR composite in which the charge-carrier $\mu$ is enhanced through the inclusion of narrow band-gap Q-dots, there was no need for the Q-dots to perform as the photosensitizer as in most of the previous studies involving the inclusion of Q-dots in an otherwise all-organic PR composite. This, in conjunction with the experimental $\lambda = 633$ nm, permitted the use of C\(_{60}\) as the photosensitizer in the current study. To study the enhancement of the $\tau$ as a result of the inclusion of narrow band-gap Q-dots, QPbS was introduced into the previously described composite at various concentrations. It will be demonstrated that the inclusion of the QPbS in the PR composite resulted in a decrease in $\tau$ by a factor of 6, to $\tau = 399 \mu$s, the fastest $\tau$ reported for PR composites under CW conditions. This significant decrease in $\tau$ is especially exciting due to the ability to avoid the detrimental effect over PR efficiency, with $\eta_{\text{ext}}$ decreasing by only 4%, to 59%, as a result of introducing the QPbS into the composite.

To characterize the performance of the composites, time-resolved degerate-four-wave-mixing (DFWM), two-beam-coupling (TBC), visible absorption spectroscopy, and conductivity experiments were employed, the results and implications of which will be presented. In addition to the insights gained into the fundamental mechanisms relevant to this class of materials, a significant advancement in PR performance is realized. It is anticipated that the advances illustrated herein represent significant progress towards the eventual practical application of PR composite materials.

**Results**

The goal of this study is to examine the feasibility of using narrow band-gap Q-dots to affect charge-carrier $\mu$ in a PR composite so as to improve the $\tau$ without detrimentally affecting the PR efficiency of the composite. The first objective to these ends involved the choice of semiconductor material\(^{17,18,23}\). In this regard, it is noted that in all previous studies concerning the inclusion of Q-dots in a PR composite, the Q-dots were included to serve as the photosensitizer. However, data accumulated in these studies demonstrate that in addition to creating free charge-carriers within the photoconductive (PC) matrix, the Q-dots also enhance the $\tau$ of the PR composites. It was speculated that this enhancement in $\tau$ was attributable to the faster charge-carrier $\mu$ associated with inorganic semiconductors as compared to the organic matrix, typically PVK. Definitive evidence of the charge-carrier $\mu$ being enhanced in a PC polymer through the inclusion of Q-dots came in a study in which QCdS was doped into a thin film of PVK\(^{31}\). Using time-of-flight experimental techniques, the QCdS/PVK composite exhibited a faster charge carrier $\mu$ than undoped PVK. Moreover, it was demonstrated that the $\mu$ scaled with Q-dot concentration.

Despite these promising results, there is also evidence that Q-dots may play a third role in addition to photoncharge-generator and charge-transport species, that of a charge trapping species. Although a certain concentration and depth of charge traps are required for the PR effect, impurities and imperfections associated with
the polymer have traditionally fulfilled this requirement, and a separate component is not usually included. For the case of Q-dots, data indicate that their propensity to act as charge traps can have a detrimental influence over the efficiency of a PR composite into which they are included. In an effort to overcome this hindrance, recent work has focused on the use of Q-dots with a narrower band-gap. These studies have demonstrated the ability to photosensitize a PR polymeric composite with QPbS where, unlike in the majority previous studies, the QPbS had a band-gap energy (1.02 eV) which was considerably less than that associated with the optical radiation employed in the PR characterizations (λ = 633 nm, corresponding to 1.96 eV). Here, using the relatively narrow band-gap QPbS resulted in τ = 34 ms, which is among the fastest reported for a polymeric PR composite photosensitized with Q-dots, as well as for polymeric PR composites in general. In addition, this τ was achieved without sacrificing efficiency, as indicated by the internal diffraction efficiency, η_int = 83% reported for this PR composite. Based on these results, it was determined that QPbS would serve as the Q-dot used in the current study. It is noted that the QPbS used in the current study originate from the same batch as those used in this cited work and were passivated with oleic acid (OA) and 1-octadecene (ODE).

TEM images of the QPbS were obtained and can be found elsewhere. The particles are irregularly shaped, but approximately spherical with a mean diameter of 3.62 ± 0.58 nm. The UV-Vis absorption spectrum of the QPbS dispersed in toluene was also collected, showing that the first exciton occurs at ~1220 nm. At a normalized concentration of 1 mg/mL, the QPbS exhibits an absorption coefficient of α_{1220} = 0.0296 cm⁻¹ at 1220 nm and α_{633} = 0.234 cm⁻¹ at 633 nm. The presence of the well defined peak at ~1220 nm also indicates a high degree of monodispersity. To quantify the ratio of organic capping material to inorganic PbS present in the QPbS, TGA was conducted. In summary, the inorganic PbS accounts for 33.6 wt% of the QPbS, with the remainder of the mass being primarily attributed to OA and ODE.

The next objective in this study was the formulation of the organic components constituting the PR composite. Because this study is largely concerned with the PR τ, it was this parameter which motivated the choices made in this regard. Perhaps the most influential component over the PR τ is that of the charge-transporting species. This functionality is often provided through the inclusion of a PC polymer, with PVK being one of the most prevalent. Because this study is largely concerned with the PR τ, it was this parameter which motivated the choices made in this regard. Perhaps the most influential component over the PR τ is that of the charge-transporting species. This functionality is often provided through the inclusion of a PC polymer, with PVK being one of the most prevalent. More recent research indicates, however, that PR composites which contain TPD as the charge-transporting moiety exhibit superior PR performance compared to their PVK counterparts, attributed in part to a faster charge-carrier μ associated with TPD. The superior performance associated with TPD-based composites can also be traced to the relationship among the various highest-occupied-molecular-orbitals (HOMO) associated with the components, especially when used in conjunction with certain NLO dyes. Specifically, TPD is more easily ionized than is PVK since the HOMO of TPD lies higher in energy at approximately –5.4 eV, than that of PVK which occurs at –5.92 eV. Furthermore, when PVK is used in conjunction with certain NLO dyes, such as 7-DCST which has a HOMO occurring at –5.9 eV, it is energetically favorable for the positively charged holes to become trapped at a 7-DCST molecule, whereas in the case of TPD this issue is avoided. It is also noted that because the HOMO of TPD is higher in energy than that of the HOMO in PVK, it is energetically favorable for photogenerated holes to remain associated with the TPD and therefore it is not anticipated that the PVK plays any significant role in the charge-transport process.

The TPD-based PR composites typically contain polymers in which the TPD is either part of the polymer backbone or alternatively as a pendant unit grafted to a polymer backbone. When employed in the pendant geometry as in PATPD, TPD-based PR composites have exhibited τ = 10 ms with η_int = 80% under CW conditions. These figures-of-merit are remarkable, especially considering that PATPD is 78.9 wt % TPD, with the remainder of the mass being attributable to the polymer backbone and pendant structure, which is essentially inert with respect to the PR process. Furthermore, PATPD has a Tg well above RT, dictating the need to include an inert plasticizer allowing for the orientational mobility of the NLO dye. There are also questions as to whether the TPD pendant are able to achieve the optimal stacking conformation which is conducive to the hopping mechanism of charge-transport ubiquitous to these types of PC materials. Faster τ have been reported for other PR composites utilizing polymerized forms of TPD, however these fast τ often come at the expense of efficiency. Several studies exist in which τ as fast as 1–3 ms are reported, though diffraction efficiencies were either not reported or did not exceed 0.5%. To date, very few examples of a sub-millisecond τ have been reported for PR composites characterized under CW conditions. A sub-millisecond of τ = 300 μs has been described for a PR composite characterized under pulse conditions.

Based on these considerations, a PR composite rootined in a molecular form of TPD is desirable. Nevertheless, polymeric forms of TPD are typically employed due to the very large propensity for TPD, as well as its various molecular derivatives, to crystallize within the solid solution of the PR composite. After screening a substantial number of TPD derivatives in conjunction with several NLO dyes as well as selected photosensitizers, it was established that a PR composite, stable against aggregation of any components for at least 12 months up until now, could be achieved using the N,N-Bis(3-methylphenyl)-N,N-bis(phenyl)benzidine (DMTPD) derivative of TPD as the PC species. Although fraught with a pair of methyl groups to aid in phase stability, this is a relatively small cost in terms of additional inert mass. As will be discussed in greater detail, this allowed for ~45 wt% loading of DMTPD, a relatively high concentration that is favorable for charge-carrier μ. In addition to the DMTPD, a relatively small amount of PVK totaling ~10 wt% was introduced to assist in phase stabilization. Although difficult to confirm, it is likely that charge-transport occurs in the PVK as well as the DMTPD, and as such the PVK may not be inert with regard to the PR process. However, based on the HOMO considerations, a positively charged hole is lower in energy when associated with DMTPD than when with PVK, and as such, once a hole becomes associated with DMTPD, it is unlikely to re-associate with the PVK.

To impart NLO activity to the PR composite, 7-DCST was employed as it has been in many recent studies which have highlighted the PR τ. The PR composites described herein contained ~45 wt% 7-DCST which is similar to concentrations employed in previously described PR composites. The photosensitizer used in the current study was C60 and is well known to be one of the most efficient PR photosensitizers for λ = 633 nm. With
a HOMO occurring at $-6.40\text{ eV}$, C$_{60}$ can also behave as a trapping species within the PR medium. In previous studies C$_{60}$ has been used in a range of concentrations, but typically on the order of 0.2–2.0 wt%. In this study C$_{60}$ was included at 0.2 wt% which is a relatively low concentration. Although attempts were made to increase the C$_{60}$ concentration in an effort to improve the PR performance, it was observed that unlike DMTPD and 7-DCST, C$_{60}$ exhibited a very high susceptibility to aggregation within the PR composite at higher concentrations. The resulting PR device is referred to as DC60PBS and its composition is detailed in Table 1.

As described in the Experimental section several control devices were also fabricated and their compositions as well as the $\alpha_{633}$ are also detailed in Table 1. The absorption spectra of the devices employed in this study are provided in Fig. 1. The device void of C$_{60}$ and QPbS, D0, exhibits $\alpha_{633} = 2.46\text{ cm}^{-1}$ whereas the addition of 0.223 wt % of QPbS, as in the DPBS device, increased the $\alpha_{633}$ by 0.29 cm$^{-1}$. Similarly, the addition of 0.200 wt % of C$_{60}$, as in DC60, resulted in an increase in $\alpha_{633}$ of 5.35 cm$^{-1}$. Finally, in the DC60PBS device, the presence of both C$_{60}$ and QPbS at concentrations commensurate with those of the control devices produced an increase in $\alpha_{633}$ by 5.58 cm$^{-1}$.

Figure 1. Visible absorption spectra of the PR devices used in this investigation; DC60PBS (solid line), DC60 (dashed line), and DPBS (dotted line).

For the DC60PBS device, C$_{60}$ is intended as the primary sensitizer, however, QPbS, included for purposes of enhanced charge-transport, has also been shown to photosensitize similar composites with $\lambda = 633\text{ nm}$

$\sigma$ are measured for each device and are shown as a function of $E$ in Fig. 2. Looking first at the $\sigma_p$, it is observed that this parameter is nearly constant for all devices across the entire range of investigated $E$. While somewhat surprising, it indicates that the presence of C$_{60}$ and/or QPbS does not have a significant effect on the $\sigma_p$. Conductivities in general, $\sigma$, depend explicitly on mobility as

\[
\sigma = pe\mu,
\]

where $p$ is the density of mobile charge carriers (holes) and $e$ is the fundamental electric charge, it can be assumed that the unilluminated QPbS does not enhance charge carrier mobility under these conditions. This situation is favorable for the PR effect because the establishment of a space-charge field depends on large ratio of $\sigma_p/\sigma_d$. Turning to the $\sigma_p$, it is apparent from the figure that the DC60PBS device exhibits the best performance in this regard. This is not surprising since this composite contains both C$_{60}$ and QPbS. The DC60 device shows a slightly diminished $\sigma_p$ relative to that of the DC60PBS device. This is again expected since DC60, though void of QPbS, still contains the primary photosensitizer, C$_{60}$. The DPBS device, although decreased by nearly an order of magnitude relative to the DC60PBS device, still exhibits a significant increase $\sigma_p$ relative to its $\sigma_d$, again demonstrating the ability for QPbS for photosensitize this composite, though significantly less effectively than that of the C$_{60}$ at the relevant concentrations. It is noted that for this study the concentration of QPbS was chosen so as to optimize

### Table 1. Compositions and $\alpha_{633}$ of the photorefractive devices used in this study.

| Device     | C$_{60}$ (wt%) | QPbS (wt%) | DMTPD (wt%) | 7-DCST (wt%) | PVK (wt%) | $\alpha_{633}$ (cm$^{-1}$) |
|------------|----------------|------------|-------------|--------------|-----------|--------------------------|
| DC60PBS    | 0.199          | 0.223      | 44.8        | 44.8         | 9.96      | 8.04                     |
| DC60       | 0.200          | NA         | 44.9        | 44.9         | 9.98      | 7.81                     |
| DPBS       | NA             | 0.223      | 44.9        | 44.9         | 9.98      | 2.75                     |
| D0         | NA             | NA         | 45.0        | 45.0         | 10.0      | 2.46                     |
the \( \tau \) of the DC60PBS device and not the \( \sigma_\text{p} \) of the DPBS device, and therefore it is not possible to draw a direct comparison of the photosensitization ability between the QPbS and C\textsubscript{60} from these data. The DC0 device (not shown in the figure) did not exhibit a measurable \( \sigma_\text{p} \).

From the \( \sigma_\text{p} \) it was possible to determine \( \Phi \) for the relevant devices and are presented as a function of \( E \) in Fig. 3. As seen, the \( \Phi \) are of the same order of magnitude at each \( E \) for all compositions. The \( \Phi \) of the DC60 device does consistently exceed that of the DPBS device, suggesting that C\textsubscript{60} provides superior photosensitization relative to that of QPbS in this composite. With \( E = 100 \text{ V/\mu m} \), the DC60 shows \( \Phi = 0.30\% \) whereas the DPBS device exhibits \( \Phi = 0.19\% \). These data indicate, however, that with further optimization, the \( \Phi \) of QPbS, or some similar material, may exceed that of C\textsubscript{60}. It is also observed that the \( \Phi \) of the DC60PBS (\( \Phi = 0.44\% \) with \( E = 100 \text{ V/\mu m} \)) device exceeds that of the DC60 device at all \( E \). Because the DC60PBS device contains both C\textsubscript{60} and PbS, and because at the relevant concentrations there is no reason to anticipate any interactions between the C\textsubscript{60} and the QPbS, to a first approximation, its \( \Phi \) should be intermediate between that of DC60 and that of DPBS. Therefore, the reason for this observation is not immediately apparent but may indicate an as of yet unknown synergistic effect between the QPbS and the C\textsubscript{60}. The available data do not warrant further speculation.
index grating generated in the PR medium and the interference pattern associated with the writing beams. The conventional TBC experiments. This exchange of energy is attributed to a spatial shift between the refractive indices photosensitized with Q-dots. For the DC60PBS device, the performance of the DPBS was far inferior with DC60, also shows a similar trend, indicating that this behavior is not unique to compositional variation of such deep traps does not vary significantly among the various compositions. This reinforces the hypothesis that narrow band-gap Q-dots have a reduced propensity to act as deep traps in these PR composites relative to their wider-band-gap counterparts. The magnitude of \( \Gamma \) is also noteworthy with DC60 exhibiting \( \Gamma = 510 \text{ cm}^{-1} \), and DC60PBS showing a slightly diminished \( \Gamma = 500 \text{ cm}^{-1} \), both at \( E = 100 \text{ V/\mu m} \). Although the DPBS device experienced dielectric breakdown prior to \( 100 \text{ V/\mu m} \), it still showed \( \Gamma = 420 \text{ cm}^{-1} \) at \( E = 90 \text{ V/\mu m} \). For practical applications the condition \( \Gamma > \alpha \) should be met for a given device. Due to its relatively small \( \alpha_{\text{DC60}} \), DPBS meets this condition at \( E \) as low as \( 10 \text{ V/\mu m} \). For DC60 and DC60PBS, their \( \Gamma \) exceeds their respective \( \alpha_{\text{DC60}} \) prior to \( 20 \text{ V/\mu m} \). Remarkably, these figures-of-merit are among the best reported for any PR material. The inset in Fig. 4 depicts the optical amplification factor, \( \gamma \), as a function of \( t \) for the DC60PBS device at \( E = 80 \text{ V/\mu m} \) and provides qualitative insight into the temporal characteristics associated with the TBC in these composites.

Having established the PR nature of the observed gratings, the internal diffraction efficiencies, \( \eta_{\text{int}} \), were evaluated. These data are presented as a function of \( E \) in Fig. 5. In this figure the solid lines represent the best fit of the data to a simple sine-squared function as predicted by theory. Although the fits are wanting, an over-modulation \( E \) where \( \eta_{\text{int}} \) achieves a maximum is clearly observed for all three devices. For DC60PBS and DC60 this occurs at \( E \approx 50 \text{ V/\mu m} \) and for DPBS at \( E \approx 40 \text{ V/\mu m} \), which are typical values for organic PR composites. Moreover, there exists a subsequent minimum in \( \eta_{\text{int}} \) for DC60PBS and DC60 at \( E \approx 90 \text{ V/\mu m} \) and for DPBS at \( E \approx 80 \text{ V/\mu m} \). With \( \eta_{\text{int}} = 75\% \), it was the DC60 device which exhibited the highest internal efficiency at it over-modulation \( E \). However, the addition of the QPbS in the DC60PBS device lowered the efficiency only slightly to 72\% at the same \( E \). It is central to this study that the diffraction efficiency is maintained in the presence of the QPbS. The performance of the DPBS was far inferior with \( \eta_{\text{int}} = 9.5\% \) at its maximum. However, it is again noted that the composition of this sample was not optimized for performance and is only for comparison purposes, and other studies employing
the same QPbS species at higher concentrations in similar composites have shown significantly higher diffraction efficiencies.

While the external diffraction efficiencies, $\eta_{\text{ext}}$, are fundamentally important, it is the $\eta_{\text{ext}}$ which are of functional significance and are depicted as a function of $E$ in Fig. 6. As in Fig. 5, the solid lines represent the best fit of the data to a simple sine-squared function. Due to their relatively high $\eta_{\text{ext}}$, in conjunction with their reasonably low $\alpha_{633}$, the DC60 and DC60PBS devices also exhibit a favorable $\eta_{\text{ext}}$ at their over-modulation $E$, with $\eta_{\text{ext}} = 63\%$ and $\eta_{\text{ext}} = 59\%$, respectively.

Time resolved DFWM techniques were used in the quantification of the $\tau$. An example of such DFWM data is presented in Fig. 7, where $\eta$ is presented as a function of $t$ for the DC60PBS with $E = 100\,\text{V/\mu m}$. The solid line in the figure represents the best fit to Equation 6 and a reasonably good fit to the data is obtained. In this case, the fast time constant, $\tau_f = 0.399\,\text{ms}$ was derived from the fitting process. This figure-of-merit represents one of the best reported for any PR composite and is nearly an order of magnitude faster than that obtained for the DC60 composite, with $\tau_f = 2.42\,\text{ms}$. It is noted that even this $\tau_f$ is among the best for any PR composite under these conditions, which demonstrates the practicality of the DMTPD-based composite. Figure 8 depicts the $\tau_f$ as a function of $E$ for the relevant devices. Obvious is that DC60PBS has a significantly faster $\tau_f$ than DC60, and both exhibit superior $\tau_f$ relative to DPBS, for the entire range of $E$ investigated. It is speculated that DPBS has a relatively...
slow $\tau_f$ due to the exceedingly low concentration of photosensitizer present in this composite. As such, it takes longer to produce sufficient number of free charge-carriers to achieve a steady-state space-charge field. These data strongly suggest that the presence of the QPbS has a significant effect on $\tau_f$. These $\tau_f$ indicate the potential for the use of these composites in video-rate applications.

It has been speculated that it is the $\tau_f$ which is associated formation of the space-charge field and it is the slow time constant, $\tau_s$ which is attributed to the reorientation of the NLO chromophore, 7-DCST in this case. The $\tau_s$ are plotted as a function of $E$ in Fig. 9. The $\tau_s$ exhibit a similar functional form as the $\tau_f$ however there appears to appear to be more noise in these data. Also evident is that there are no $\tau_s$ reported for DC60PBS for $E > 80$ V/$\mu$m or for DC60 when $E > 70$ V/$\mu$m. The reason for this is evident from the inset of Fig. 9. As mentioned in the Experimental section, the weighting factor, $m$, was also allowed to float in the fitting process. Apparent from the figure is that the $m$ are highly scattered, but display the unmistakable trend of more heavily weighting $\tau_f$ at the expense of $\tau_s$ as $E$ is increased for all devices. At relatively low voltages $m \approx 0.5$ and the $\tau_s$ and $\tau_f$ contribute almost equally to the grating formation. As higher voltages are attained, the data suggest that it is the formation of the space-charge field which becomes the limiting process. For relatively high voltages $m$ tends toward unity and the rise in DFWM signal can be modeled by a single exponential. For DPBS, $m$ does increase with $E$, but $m \neq 1$ even at higher $E$. This is again attributed to the relatively slow build up in the density of free charge carriers, and the

**Figure 7.** Temporal evolution of the diffracted probe beam, $I_s$, in the DFWM experiment for the DC60PBS device at $E = 100$ V/$\mu$m. The solid line is a fit to a weighted biexponential function (see text). The inset depicts an expanded view of the same data (the fit is eliminated for clarity).

**Figure 8.** Electric field dependence of the fast time constant, $\tau_f$, for DC60PBS (squares), DC60 (circles), and DPBS (triangles) at $\lambda = 633$ nm. The lines are guides for the eye.
QPbS, as exemplified by the exceedingly small increase in the charge-carrier density. Furthermore, if QPbS were significantly contributing to the photo-charge generation, the inability of the NLO chromophore to align quickly in the absence of a reasonably large space-charge field. Though given the scatter in the data, it is best to not draw a strict interpretation concerning this fact.

As outlined in the introduction, QPbS was included in the DC60PBS device with the expectation of enhancing the charge-carrier μ, thus improving the τ of this PR composite relative to a composite void of QPbS, i.e. DC60. The data, as summarized in Table 2, leave little doubt as to the success of the approach, as τ showed an enhancement of nearly one order of magnitude and remained relatively constant. This distinction corresponds to current PR theory in that the presence of the QPbS was included to enhance charge-carrier μ, which would directly improve the speed at which the space-charge field is established, and it is this process which is commonly associated with the τ. Conversely, there is no reason to expect that the presence of QPbS would influence the rate at which the NLO chromophores, 7-DCST in this case, would reorient within the established space-charge field. Because it this process which is associated with τ, it is expected that τ should not be effected by the presence of QPbS and thus remain relatively constant between the DC60PBS and DC60 devices. It is noteworthy that the observed enhancements in τ, as well as the various other PR parameters occur at a relatively low concentration of QPbS. While the mechanism responsible for the anticipated improvement in μ has been hypothesised in the literature, exactly how such a significant effect can be attributed to such a relatively low concentration of QPbS may not be entirely explained through this proposed mechanism and is the subject of current research. It is further noted that enhancements of similar magnitude have been reported for similarly low Q-dot concentrations.

Although the inclusion of QPbS dramatically improved the τ, this observation does not in itself confirm that this enhancement was accomplished through the envisioned mechanism, an enhancement in charge-carrier μ, and this conclusion requires some further analysis. Ideally, the μ should be measured directly through time-of-flight experiments. An attempt was made in this regard, however, as is typically the case the presence of the highly dipolar 7-DCST resulted in highly diffuse charge-packets precluding such measurements. In an effort to circumvent this issue, samples were fabricated void of 7-DCST however the DMTPD crystalized before such experiments could be completed. Although not possible with the current compositions, previous time-of-flight characterizations conducted with PVK doped with QCdS clearly demonstrated a correlation between the concentration of QCdS and the charge-carrier μ. While it is very probable that the observed improvement in τ can be primarily attributed to an enhancement in μ, other mechanisms may be considered. Perhaps the most obvious mechanism is one involving the augmentation of charge-carriers with QPbS acting primarily as a photosensitizer. Although QPbS does certainly act as a photosensitizer it is unlikely that such a significant improvement in τ could be attributed to this effect. Although the DPBS device exhibits respectable Φ, the loading content of QPbS, as exemplified by the exceedingly small increase in α_{633} in the DPBS device relative to that of the D0 device (2.75 cm^{-1} - 2.46 cm^{-1} = 0.29 cm^{-1}), indicates that the QPbS is not likely responsible for any significant increase in the charge-carrier density. Furthermore, if QPbS were significantly contributing to the photo-charge generation.

![Figure 9](image_url)  
Figure 9. Electric field dependence of the slow time constant, τ_s, for DC60PBS (squares), DC60 (circles), and DPBS (triangles) at λ = 633 nm. The lines are guides for the eye. The inset depicts the electric field dependence of the weighting factor, m, for the same PR devices. The lines are guides for the eye.

### Table 2. Summary of the PR figures-of-merit attributed to the various composites relevant to this study.

| Device     | τ_int (ms) | τ_s (ms) | η_int (%) | η_s (%) | Γ (cm⁻¹) | Γ_{633} (cm⁻¹) |
|------------|------------|----------|-----------|---------|----------|----------------|
| DC60PBS    | 0.40 ± 0.01 | 17.5 ± 0.7 | 72°       | 59°     | 500°     | 492°           |
| DC60       | 2.4 ± 0.2  | 156 ± 3   | 75°       | 63°     | 510°     | 502°           |
| DPBS       | 6.3 ± 0.1  | 310 ± 50  | 9.5°      | 8.9°    | 420°     | 417°           |

| Electric field, E [V/μm] | τ_int (ms) | τ_s (ms) | η_int (%) | η_s (%) | Γ (cm⁻¹) | Γ_{633} (cm⁻¹) |
|--------------------------|------------|----------|-----------|---------|----------|----------------|
| 100                      | 0.40 ± 0.01 | 17.5 ± 0.7 | 72°       | 59°     | 500°     | 492°           |
| 90                       | 2.4 ± 0.2  | 156 ± 3   | 75°       | 63°     | 510°     | 502°           |
| 80                       | 6.3 ± 0.1  | 310 ± 50  | 9.5°      | 8.9°    | 420°     | 417°           |
| 70                       | 0.40 ± 0.01 | 17.5 ± 0.7 | 72°       | 59°     | 500°     | 492°           |
| 0.1                      | 2.4 ± 0.2  | 156 ± 3   | 75°       | 63°     | 510°     | 502°           |
| 0.2                      | 6.3 ± 0.1  | 310 ± 50  | 9.5°      | 8.9°    | 420°     | 417°           |

\[ \mu = \frac{\alpha_{633}}{\epsilon_0} \]

where \( \alpha_{633} \) is the molar absorption coefficient at 633 nm. The lines are guides for the eye. The inset depicts the electric field dependence of the weighting factor, m, for the same PR devices. The lines are guides for the eye.
process, an accompanying increase in $\eta$ would be expected, but this was not observed. It was also speculated that QPbS may enhance PR performance in ways not related to its ability to act as a photosensitizer. Based on the propensity for Q-dots to act as traps in PR composites, it may be possible that the included QPbS enhances the trapping of holes in the dark fringes. However, because the presence or absence of QPbS in the respective composites had little if any effect on $\sigma_d$ and only a favorable effect on $\eta$, it seems unlikely that QPbS is acting as a significant trapping species. Interestingly, the fact that the addition of QPbS results in an improvement in $\sigma_d$ without affecting $\eta$, indicates that enhancements in charge-carrier $\mu$ may manifest exclusively in illuminated regions and not in dark regions. This performance would be favorable for PR performance because charge-carriers are ideally immobilized within the dark fringes. Lastly, because TBC is highly sensitive to trap density, the similarity in the TBC performances across the various composites further refutes the speculation that QPbS is acting as a significant source of traps.

Discussion

The goal of this study was to examine the feasibility of using Q-dots to affect charge-carrier $\mu$ in a PR composite so as to improve the $\tau$ without detrimentally affecting PR efficiency of the composite. It has been demonstrated that through the inclusion of a narrow bang-gap Q-dot, specifically QPbS, sub-millisecond response times have been achieved for the first time in a PR composite under CW conditions. With a $\tau_{f} = 400 \mu s$, this approach makes the use of PR composites viable materials for video and other high-speed applications. It is noteworthy that the enhancement in $\tau_{f}$ is not accompanied by a loss in PR efficiency, with $\eta_{\mu} = 72\%$ and $\Gamma = 500 \text{ cm}^{-1}$ observed in the same composite, indicating that the included QPbS does not act as a significant source of traps. An analysis of the experimental data strongly suggests that the enhancement in $\tau_{f}$ can be attributed to an improvement in enhanced charge $\mu$, which may even be exclusively manifest in the illuminated regions of the PR sample. It is anticipated that this approach can play a significant role in the eventual commercialization of this class of materials.

Experimental Section

All chemicals were obtained from Aldrich and used as received unless otherwise noted. QPbS: QPbS was synthesized based on a procedure found in the literature and is described in detail elsewhere. The QPbS were characterized using visible absorption spectroscopy, thermal gravimetric analysis (TGA), and TEM. The data obtained from these characterizations can also be found elsewhere.

PR Composite Devices.

DMTPD was obtained from Magical Scientific USA and purified by recrystallization in hot CH$_2$Cl$_2$ prior to use. 7-DCST was synthesized in our lab according to a procedure in the literature. For the composite samples, PVK (secondary standard), DMTPD, 7-DCST and the appropriate quantities of capped QPbS and C$_{60}$ were dissolved in toluene and, after thorough mixing, filtered to remove any undissolved solids. This solution was stored in a vacuum oven at 50 °C for 24 h to remove the solvent. The solid residue was subsequently recovered, placed between two pieces of glass coated with indium tin oxide and heated above its melting temperature on a hot-plate. The sample was then mechanically pressed forming the typical “sandwich” geometry using glass spacers to control the thickness of the device, $d$, at 100 μm.

Although many compositions were considered, the results for four devices are presented here. For these PR devices, the TPD:7-DCST:PVK ratio was held constant at 45:45:10 wt%, respectively. The first PR device contained only these components, constituting a control device and is herein referred to as D0. A second device contained these components in this ratio as well as being photosensitized through inclusion of 0.200 wt % of C$_{60}$. This device is referenced as DC60. A third PR device was fabricated nearly identical to the second with the exception of the addition of 0.223 wt% of QPbS. It is noted that this wt% includes both inorganic PbS as well as the organic capping groups. This device is referred to as DC60PBS. Finally a fourth PR device was fabricated which was identical to the third, with the exception that the C$_{60}$ was not included. This device was fabricated so that the effect of the QPbS over the charge-generation process could be isolated and is referred to as DPBS. To determine the weight percentage of QPbS in each composite, several QPbS:solvent solutions of known absorbance and volume were evaporated to dryness and the mass of the solid residue was measured. It is noted that this residue includes the inorganic PbS as well as the organic passivating layer. The weight percent of QPbS in each device was thus calculated indirectly from the known absorbance and volume of the aliquot used in the fabrication of the respective composite. The compositions and the spectroscopically measured $\alpha_{\text{abs}}$ of all devices are presented in Table 1. The PR devices have not shown any change in their optical properties or degradation in PR performance over the course of 12 months. All UV-Vis absorption spectra were recorded on a Beckman DU 640B spectrophotometer.

Photorefractive Characterizations.

The PR properties of the composite devices were studied via TBC and temporally resolved DFWM techniques using a standard tilted geometry. Holographic gratings were written through the intersection of two coherent beams generated by a helium-neon (HeNe) laser operating at 633 nm with incident angles of $\theta_i = 45^\circ$ and $\theta_s = 75^\circ$ (in air) relative to the sample normal. In the TBC experiments, both writing beams were $p$-polarized with intensities of $I_p = 0.05 \text{ mW}$ and $I_s \approx 8 \text{ mW}$. The external bias was applied such that $I_0$ would experience gain at the expense of $I_s$. Asymmetric energy transfer was observed by monitoring the intensities of the writing beams after the PR device with a photodiode. In the DFWM experiment the writing beams were $s$-polarized with intensities of $I_p \approx 3 \text{ mW}$ and $I_s \approx 9 \text{ mW}$. In addition, a $p$-polarized probe beam propagated in a direction opposite to $I_p$ with an intensity of $I_p \approx 2 \times 10^{-3} \text{ mW}$. Through the use of a polarizing beam splitter placed in the path of $I_s$ in conjunction with a photodiode, the diffracted portion of $I_p$, also referred to as the signal beam, $I_s$, could be quantified. In all PR experiments, $I_p$ and $I_s$ had beam diameters of ~280 μm while $I_p$ possessed a beam diameter of ~120 μm. Beam diameters were measured by using a fractional irradiance of 1/e$^2$. The $\Gamma$, was determined in terms of the experimentally measured quantities $\gamma$ and $\beta$, as
\[ \Gamma = \frac{\ln(\gamma \beta) - \ln(\beta + 1 - \gamma)}{L}, \]  

where \( L \) is the path length of the beam experiencing gain inside the sample, \( \beta \) is the ratio of the writing beam intensities before the sample, and \( \gamma \) is the ratio of the intensity of the beam experiencing gain with and without the pump beam. The \( \eta_{int} \) were quantified according to the equation

\[ \eta_{int} = \frac{I}{I'p}, \]  

where \( I_p \) is the intensity of the probe beam after the device with no bias applied. Similarly, the \( \eta_{ext} \) accounting for reflections, were determined according to the equation

\[ \eta_{ext} = \frac{I}{I'p}. \]  

Time resolved DFWM techniques were used in the quantification of the \( \tau \). For this experiment, \( E \) was applied to the device while blocking one of the writing beams. The device was permitted to settle for approximately 30 s and the blocked writing beam was unblocked. The diffracted portion of the probe beam was then recorded as a function of time, \( t \). The determination of the \( \tau \) was accomplished by fitting the data obtained in DFWM experiments to an equation of the form

\[ \eta(t) = \sin^2 [A [1 - m \exp - t/\tau_f - (1 - m) \exp - t/\tau_s]], \]  

where \( A \) is a fitting constant, \( m \) is a weighting ratio, \( \tau_f \) is the fast time constant and \( \tau_s \) is the slow time constant. For the fitting, all four variables were allowed to float.

Conductivity characterizations were made using a dc-photocurrent technique with a Keithley electrometer used to measure the current passing through the sample as a function of \( E \). The beam intensity for all \( \sigma_p \) characterizations was \( \sim 10 \text{ mW} \) with a beam diameter of 0.98 mm. To measure the \( \sigma_p \), the sample was initially illuminated for several minutes followed by the application of the \( E \) and the current was allowed to achieve a steady state (typically \( \sim 5 \text{ min} \)). To measure the dark conductivity, \( \sigma_d \), an \( E \) was applied and the current was allowed to reach a steady state. In calculating the \( \sigma_p \), the dark current density was subtracted from the photocurrent density. For calculations involving the photocurrent density, the diameter of the laser was employed, and for those related to the dark current density, the area of electrode overlap was used. These data were calculated using the equation

\[ \sigma = \frac{I}{E}, \]  

where \( I \) is the experimentally determined current density.

From the \( \sigma_p \) data it is possible to determine the internal photocurrent efficiency, \( \Phi \), of the photosensitizer using the equation

\[ \Phi = \frac{N_c}{N_{ph}} = \frac{\sigma_p \hbar c E}{k e \lambda \alpha_d d}, \]  

where \( N_c \) is the number of charge-carriers generated per unit volume, \( N_{ph} \) is the number of photons absorbed per unit volume, \( h \) is Planck’s constant, \( c \) is the speed of light, and \( e \) is the fundamental unit charge.

References
1. Gunter, P. & Huignard, J.-P. Photorefractive Materials and Their Applications, I & II, Topics in Applied Physics, Vols. 61 and 62, Eds.; Springer-Verlag: Berlin (1988).
2. Ostroverkhova, O. & Moerner, W. E. Organic Photorefractives: Mechanisms, Materials, and Applications. Chem. Rev. 104, 3267–3314 (2004).
3. West, D. & Binks, D. J. In Physics of Photorefractor in Polymers; Garito, A. F., Kajzar, F., Boardman, A., Eds.; Advances in Nonlinear Optics & CRC Press: Boca Raton, Florida (2005).
4. Lynn, B., Blanche, P.-A. & Peyghambarian, N. Photorefractive Polymers for Holography. J. Polym. Sci., Part B: Polym. Phys. 52, 193–231 (2014).
5. Tsutsumi, N. Recent Advances in Photorefractive and Photoactive Polymers for Holographic Applications. Polym. Int. doi: 10.1002/pi.5096 (2016).
6. Winiarz, J. G., Zhang, L., Lal, M., Friend, C. S. & Prasad, P. N. Photogeneration, charge transport, and photoconductivity of a novel PVK/CdS-nanocrystal polymer composite. Chem. Phys. Lett. 245, 417–428 (1999).
7. Winiarz, J. G., Zhang, L., Lal, M., Friend, C. S. & Prasad, P. N. Observation of the photorefractive effect in a hybrid organic-inorganic nanocomposite. J. Am. Chem. Soc. 121, 5287–5295 (1999).
8. Winiarz, J. G., Zhang, L., Park, J. & Prasad, P. N. Inorganic: Organic hybrid nanocomposites for photorefractivity at communication wavelengths. J. Phys. Chem. B 106, 967–970 (2002).
9. Winiarz, J. G. & Prasad, P. N. Photorefractive inorganic polymer-dispersed liquid-crystal nanocomposite photosensitized with cadmium selenide quantum dots. Opt. Lett. 27, 1330–1332 (2002).
10. Binks, D. J., Bant, S. P., West, D. P., O’Brien, P. & Malik, M. A. CdSe/CdS core/shell quantum dots as sensitizer of a photorefractive polymer composite. J. Mod. Opt. 50, 299–310 (2003).
11. Choudhury, K. R., Winiarz, J. G., Samoc, M. & Prasad, P. N. Charge carrier mobility in an organic-inorganic hybrid nanocomposite. Appl. Phys. Lett 82, 406–408 (2003).
12. Puentes-Hernandez, C., Suh, D. J., Kippelen, B. & Marder, S. R. High-performance photorefractive polymers sensitized by cadmium selenide nanoparticles. Appl. Phys. Lett. 85, 534–536 (2004).
13. Aslam, E. et al. Photorefractive performance of a CdSe/ZnS core/shell nanoparticle-sensitized polymer. J. Chem. Phys. 122, 184713 (2005).
14. Aslam, E., Binks, D. J., Daniels, S., Pickett, N. & O’Brien, P. Spectroscopic studies of nanoparticle-sensitised photorefractive polymers. Chem. Phys. 316, 171–177 (2005).
15. Choudhury, K. R., Sahoo, Y., Ohulchansky, T. Y. & Prasad, P. N. Efficient photoconductive devices at infrared wavelengths using quantum dot-polymer nanocomposites. Appl. Phys. Lett. 87, 073110 (2005).
16. Cho, N. et al. Efficient Photodetection at IR Wavelengths by Incorporation of PbSe–Carbon Nanotube Conjugates in a Polymeric Nanocomposite. *Adv. Mater.* **19**, 232–236 (2007).

17. Winiarz, J. G. Enhancement of the photorefractive response time in a polymeric composite photosensitized with CdTe nanoparticles. *J. Phys. Chem. C* **111**, 1904–1911 (2007).

18. Fears, T., Anderson, C. & Winiarz, J. G. Photorefractivity in a polymeric composite photosensitized with NiS nanocrystals. *J. Chem. Phys.* **129**, 15470 (2008).

19. Sliwińska, E., Mansurova, S., Hartwig, U., Buse, K. & Meerholz, K. Effect of Co-Sensitization in New Hybrid Photo-Refractive Materials Based on PVK Polymer Matrix and Inorganic LiNbO₃-Nano-Crystals. *Appl Phys B* **95**, 519–524 (2009).

20. Anczykowski, A., Bartkiewicz, S., Nyl, M. & Mysliwiec, J. Study of Semiconductor Quantum Dots Influence on Photorefractivity of Liquid Crystals. *Appl. Phys. Lett.* **101**, 101107 (2012).

21. Li, C., Li, X., Cao, L., Jin, G. & Gu, M. Exciton-Plasmon Coupling Mediated Photorefractivity in Gold-Nanoparticle- and Quantum-Dot Dispersed Polymers. *Appl. Phys. Lett.* **102**, 251115 (2013).

22. Dradzhal, K., Bartkiewicz, S. & Miniewicz, A. Electrooptical Properties of Hybrid Liquid Crystalline Systems Containing CdSe Quantum Dots. *Appl. Phys. Lett.* **105**, 231903 (2014).

23. Moon, J.-S. et al. Off-Resonance Photosensitization of a Photorefractive Polymer Composite using PbS Nanocrystals. *J. Phys. Chem. C* **119**, 13827–13835 (2015).

24. Wright, D., Díaz-García, M. A., Casperson, J. D., DeClue, M. & Moerner, W. E. High-speed photorefractive polymer composites. *Appl. Phys. Lett.* **73**, 1490–1492 (1998).

25. Díaz-García, M. A. et al. Photorefractive properties of poly(N-vinyl carbazole)-based composites for high-speed applications. *Chem. Mater.* **11**, 1784–1791 (1999).

26. Ostrauskaitė, J., Karickal, H. R., Leopold, A., Haerer, D. & Thelakkat, M. Poly [bis (triphenylamine) ether] s with low glass transition temperatures as photoconductors in fast photorefractive systems. *J. Mater. Chem.* **12**, 58–64 (2002).

27. Wright, D., Gubler, U., Moerner, W. E., DeClue, M. S. & Siegel, J. S. Photorefractive properties of poly (siloxane)-trialkylamine-based composites for high-speed applications. *J. Phys. Chem. B* **107**, 4732–4737 (2003).

28. Thomas, J. et al. Bistriarylamine Polymer Photorefractive Polymers for Photorefractive Applications. *Adv. Mater.* **16**, 2032–2036 (2004).

29. Jung, G. B. et al. High-speed TPD-based photorefractive polymer composites. *Sen’i Gakkaishi* **60**, 193–197 (2004).

30. Fuentes-Hernandez, C. et al. Video-rate compatible photorefractive polymers with stable dynamic properties under continuous operation. *Appl. Phys. Lett.* **85**, 1877–1879 (2004).

31. Tay, S. et al. High-performance photorefractive polymer operating at 1550 nm with near-video-rate response time. *Appl. Phys. Lett.* **87**, 171105 (2005).

32. Eralp, M. et al. Photorefractive polymer device with video-rate response time operating at low voltages. *Opt. Lett.* **31**, 1408–1410 (2006).

33. Eralp, M. et al. Submillisecond response of a photorefractive polymer under single nanosecond pulse exposure. *Appl. Phys. Lett.* **89**, 114105 (2006).

34. Tsutsumi, N., Kinashi, K., Masumura, K. & Kono, K. Photorefractive Dynamics in Poly(triarylamine)-Based Polymer Composites. *Opt. Express* **23**, 25158–25170 (2015).

35. Tsutsumi, N., Kinashi, K., Masumura, K. & Kono, K. Photorefractive Performance of Poly(triarylamine)-Based Polymer Composites: An Approach from the Photocative Properties. *J. Polym. Sci., Part B: Polym. Phys.* **53**, 502–508 (2015).

36. Grunnet-Jepsen, A. et al. Spectroscopic determination of trap density in C₆₀-sensitized photorefractive polymers. *Chem. Phys. Lett.* **291**, 533–536 (1998).

37. Blanch, P.-A. Photorefractive Polymers for 3D Display Application. In *Optical Properties of Functional Polymers and Nano Engineering Applications*; Jain, V.; Kokol, A., Eds.; CRC Press: Boca Raton, Florida, pp 27–60 (2015).

38. Tsujimura, S., Fujihara, T., Sassa, T., Kinashi, K., Sakai, W., Ishibashi, K. & Tsutsumi, N. Enhanced Photoconductivity and Trapping Rate Through Control of Bulk State in Organic Triphenylamine-Based Photorefractive Materials. *Org. Electron.* **15**, 3471–3475 (2014).

39. Smith, A. M. & Nie, S. Semiconductor nanocrystal structure, properties, and band gap engineering. *Acc. Chem. Res.* **43**, 190–200 (2010).

40. Barkhouse, D. A. R., Pattantyus-Abraham, A. G., Levina, L. & Sargent, E. H. TiO₂ passivates recombination centers in colloidal quantum dots leading to enhanced photocatalytic device efficiency. *ACS Nano* **2**, 2356–2362 (2008).

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**Author Contributions**

J.-S.M and J.G.W designed the project, performed device fabrication, measurements, data analysis and wrote the main manuscript text and figures. T.E.F, T.C.M and D.L.H synthesized the PbS and assisted with data analysis. J.-W.O and S.-H.J assisted with data analysis. All authors reviewed the manuscript.

**Additional Information**

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