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Exploration of CdTe quantum dots as mesoscale pressure sensors via time-resolved shock-compression photoluminescent emission spectroscopy

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The nanometer size of CdTe quantum dots (QDs) and their unique optical properties, including size-tunable narrow photoluminescent emission, broad absorption, fast photoluminescence decay, and negligible light scattering, are ideal features for spectrally tagging the shock response of localized regions in highly heterogeneous materials such as particulate media. In this work, the time-resolved laser-excited photoluminescence response of QDs to shock-compression was investigated to explore their utilization as mesoscale sensors for pressure measurements and in situ diagnostics during shock loading experiments. Laser-driven shock-compression experiments with steady-state shock pressures ranging from 2.0 to 13 GPa were performed on nanocomposite films of CdTe QDs dispersed in a soft polyvinyl alcohol polymer matrix and in a hard inorganic sodium silicate glass matrix. Time-resolved photoluminescent emission spectroscopy was used to correlate photoluminescence changes with the history of shock pressure and the dynamics of the matrix material surrounding the QDs. The results revealed pressure-induced blueshifts in emitted wavelength, decreases in photoluminescent emission intensity, reductions in peak width, and matrix-dependent response times. Data obtained for these QD response characteristics serve as indicators for their use as possible time-resolved diagnostics of the dynamic shock-compression response of matrix materials in which such QDs are embedded as in situ sensors. Published by AIP Publishing.

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I. INTRODUCTION

The shock-compression behavior of highly heterogeneous materials, such as particulate media, is dominated by mesoscale effects arising from collapse of voids, as well as shock wave interactions with heterogeneities (e.g., particle boundaries and interfaces between constituents with dissimilar material properties). For applications involving materials subjected to shock-compression or dynamic high strain rate loading, understanding the influence of mesoscale heterogeneities on the bulk behavior/response of such materials is of paramount importance. There is, therefore, a significant need for diagnostic methods and sensor materials that can provide direct in situ measurements of pressure (and other parameters) with appropriate time- and spatial resolutions relevant to the understanding of the dynamics of shock compression processes in inert and reactive particulate materials. The limitations of shock-compression experiments in providing the mesoscopic details of deformation and mass flow in particulate materials have long been recognized.1 Conventional diagnostics, such as in situ piezoresistive/piezoelectric stress gauges,2,3 provide time resolutions down to the nanosecond scale but have spatial resolutions spanning the area of the sensor element of several millimeters. The current state-of-the-art in velocity interferometry, namely, “line VISAR” or “ORVIS” (Optically recording velocity interferometer system), allows for the collection of velocity profiles along a line,4 providing information inherent to the heterogeneous effects along the linear dimension (typically up to couple of millimeters); however, it remains a surface-sensitive diagnostic method that averages the effects of the volume of the material being diagnosed.

Mesoscale sensing in shock-compression experiments requires the sensor material to be small in size and possess pressure-dependent physical properties which can be captured and recorded at the nanosecond scale to achieve high temporal resolution.5,6 Quantum dots (QDs) are semiconducting nanoparticles of a few to tens of nanometers. These robust inorganic photoemitters exhibit size-dependent emission spectra in the whole visible and near-infrared range, due to quantum confinement effects.7 Heterogeneous materials consisting of different constituents can have each component tagged with different-sized QDs. The QDs can be excited with a single wavelength, while the photoluminescent emission can consist of multiple discrete narrowband components corresponding to each tagged constituent, thereby providing

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a great advantage for multiplexed sensing at different locations or different interfaces within the heterogeneous sample. The photoluminescence decay time of QDs is in the picosecond to nanosecond range; thus, a fast temporal response can be achieved. Nanoparticles that are small compared to optical wavelengths exhibit negligible light scattering when incorporated into transparent matrices. The optical properties of QDs have been reported to be pressure- and strain-dependent as a result of changes in their bandgap with particle size.\textsuperscript{8,9} Therefore, QDs can potentially be used as sensor materials for investigating the time-resolved mesoscale effects of shock-compression of highly heterogeneous materials, including compacts of inert or reactive powders, in order to obtain unprecedented information about local particle or ensemble behavior, as well as the dynamics of the surrounding materials during high-strain-rate or shock-compression loading.

Hydrostatic compression of QDs has been observed to produce photoluminescent emission blueshifts that increase monotonically with pressure.\textsuperscript{10–12} In our recent theoretical study\textsuperscript{13} on QDs subjected to shock-compression and associated non-hydrostatic loads, pressure-induced blueshifts were observed to initially increase to a maximum and then decrease. The effect was also confirmed through experimental studies\textsuperscript{14} of the photoluminescent emission spectra of shocked ~3 nm CdTe QDs in a hydrophilic polymer matrix (poly-vinyl alcohol; PVA). However, the time response of the measured blueshift from the QDs in the PVA matrix was not very fast (of the order of ~60 ns), indicating that the highly sluggish mechanics of the surrounding (polymer matrix) medium plays an important role in influencing the optomechanical response of the QDs. Therefore, the effect of embedding the QDs in different matrix media, on the time resolution of the QD response, needs to be studied. In the present work, we have investigated and compared the shock-compression response of CdTe QDs dispersed in both a soft PVA polymer matrix and a hard sodium silicate glass matrix. Studies of these two types of matrix media, hereafter referred to as “polymer” and “glass,” respectively, are also relevant for the understanding of the response of two widely-used embedded shock sensors, namely, the polyvinylidene fluoride polymer (PVDF)\textsuperscript{15} and quartz stress gauges.\textsuperscript{15}

In addition to matrix effects, many other issues need to be addressed and understood prior to the possible use of QDs as pressure sensors for \textit{in situ} probing of the shock-compression response of heterogeneous materials. The QD structure and surface chemistry need to be tuned to maintain compatibility with the media of interest. Previous studies have indicated that positively-charged QDs are needed with the PVA matrix, since negatively-charged QDs heavily aggregate in PVA, leading to a complex and abnormal spectroscopic response. As shown in our previous theoretical study,\textsuperscript{13} the overall shock-induced blueshift mechanism can be understood based on strain tuning of the QD bandgap using a simplified model of a QD without surface capping groups in a matrix. However, factors such as shock-induced photoluminescent emission intensity change, peak width change, and the possibility of dynamic phase transformations, require more detailed experimental studies. In the present work, the response of CdTe QDs of different sizes and capping groups dispersed in a polymer or glass (matrix) media was investigated to understand the effects of dynamic high-pressure (shock-compression) loading on their photoluminescence spectral response.

II. SYNTHESIS of CdTe QDs AND NANOCOMPOSITE FILMS

Colloidal CdTe QDs were synthesized by adding freshly prepared NaHTe solution to N\textsubscript{2}-saturated Cd(ClO\textsubscript{4})\textsubscript{2} solutions in the presence of 2-mercaptoethaneamine (MEA) or 3-mercaptopropionic acid (MPA) as the stabilizing agents and water as the solvent. The solutions were then heated to the boiling temperature and refluxed for different times to grow and obtain QDs of controlled particle sizes. The detailed synthesis procedure is reported elsewhere.\textsuperscript{16,17} The surface chemistries of the QDs were altered to prepare non-aggregated nanocomposite films. For incorporation in the PVA matrix, we used MEA-capped QDs, which had net positive charge due to –NH\textsubscript{4}\textsuperscript{+} (ammonium) capping groups. For incorporation in the glass matrix, a liquid sodium silicate solution (obtained from Sigma-Aldrich) with MPA-capped QDs, which had net negative charge due to the –COO\textsuperscript{−} (carboxylate) end groups, was used.

The MEA-capped QDs in the nanocomposite polymer films were ~3 nm in size and emit at a characteristic wavelength of 586 nm. The MPA-capped QDs in the nanocomposite glass films were ~4 nm in size and emit at a characteristic wavelength of 674 nm. A dual-emitting composite film sample with a PVA matrix was also prepared using both 582 nm and 632 nm emitting MEA-capped QDs. After the QDs were dispersed at room temperature in the respective polymer, or glass (water-based sodium silicate solution), matrix to form solutions of appropriate concentration and viscosity, nanocomposite thin film samples were prepared by drop-coating onto 25.4 × 25.4 × 0.5 mm\textsuperscript{2} glass substrates (Delta Technologies) followed by air drying to form homogeneous non-aggregated ~30 μm thick films with ~0.15 wt. % QD concentration.

III. SETUP OF LASER-DRIVEN SHOCK-COMPRESSION EXPERIMENTS

The shock-compression experiments involved a laser-launched Al-flyer plate impacting on QD-containing nanocomposite films, using the setup developed by the Dlott group.\textsuperscript{18} A schematic illustration of the experimental set-up is shown in Figure 1, with details described previously.\textsuperscript{19} Photon Doppler velocimetry (PDV)\textsuperscript{19} was used to monitor the flyer plate velocity history via a laser beam that transmitted through the transparent sample and reflected back into a detector from the flyer surface, while simultaneously, time-resolved spectroscopic photoluminescence spectra were obtained from the laser-excited QDs. The flyer plates were launched from an assembly consisting of a glass substrate 6.35 mm in thickness and an aluminum foil (Alufoil) 5 × 5 cm\textsuperscript{2} in area and 50 μm in thickness which were bonded together using low-viscosity Eccobond 24 water-clear epoxy (Henkel Loctite). A spatially homogenized Q-switched Nd:YAG laser (Quanta-Ray Pro-350) was
used to launch the flyer, producing shock pulses of \(\sim 20\) ns in duration following the impact on the sample. The flyer plate velocities measured with PDV were controlled by varying the launch pulse energy up to a maximum of 2.5 J/pulse. The laser launch beam has a central 0.5 mm diameter region whose intensity is uniform to within \(\pm 2\%\). An all-reflective beam stretcher was used to stretch the nominal 10 ns laser pulse to 20 ns full-width half-maximum.\(^{20}\) The PDV and spectroscopic measurements probed smaller regions of 0.07 mm and 0.2 mm in diameter, respectively, at the center of the uniform launch pulse.\(^{18}\) The QD photoluminescent emission was excited by 250 ns FWHM pulses from a Q-switched 527 nm Nd:YLF laser (Quantronix 527 DP) and recorded with 0.5 ns effective time per CCD pixel using a streak camera. Displayed data show each row of the streak camera output spaced at 0.5 ns intervals. The overall time resolution of the system was 2 ns.

The PDV measurements probed the velocity history of the impact surface of the 50 \(\mu\)m thick Al flyer plates. After the initial acceleration transient, the flyer plate travels freely (in vacuum) at a velocity denoted as \(U_f\). When the plates impact the targets, the velocity drops to a value denoted as \(U_p\) which is the velocity of the flyer/target interface, or alternatively the material velocity of the shocked target at the impact face. The measured values of \(U_f\) averaged over the time interval of the steadily-driven shocks and averaged over ten shock events, typically varied by \(\pm 2\%\) (one standard deviation). The calculated pressure uncertainty then depends on the Hugoniot for each medium, and a reasonable estimate is \(\pm 5\%\).

IV. RESULTS AND DISCUSSIONS

Two examples of time-stream photoluminescent emission spectra of samples impacted by 50 \(\mu\)m thick flyers, recorded by a spectrograph and streak camera, are shown in Figure 2, with one time streak for 586 nm emitting CdTe QDs in polymer (with \(U_f = 1.12\) km/s) and the other for 674 nm emitting CdTe QDs in glass with \(U_f = 0.89\) km/s. Figures 2(a) and 2(b) present the data as contour plots of the streak camera output, and Figures 2(c) and 2(d) display the corresponding photoluminescent emission spectra. The QDs in both samples exhibit shock-induced blueshift and intensity loss. The CdTe QDs are direct band gap semiconductors with strong quantum confinement effects, and shock-compression leads to increases in band gap energies and corresponding blueshifts in photoluminescent emission wavelength.\(^{10,12}\) Note that the shock response of the QDs, measured in terms of the time required for attaining the peak wavelength shifts and intensity changes, is much faster in glass than in the polymer.

The photoluminescent emission spectra were processed to extract time-dependent intensity changes, wavelength blueshifts, and spectral (or peak) widths, using the method of moments proposed by Brown et al.,\(^{18}\) which is convenient...
for automated analysis of time-streams of spectra. The zeroth moment is the wavelength-integrated photoluminescent emission intensity, the first moment is the average wavelength blueshift, and the square-root of the second moment is the spectral width. The effects of the polymer or glass matrix are presented and discussed in Sections IV A–IV E.

### A. Effect of matrix material on response time

Figure 3 shows typical plots of flyer velocity profiles, fractional photoluminescent emission intensity decreases, and blueshifts for QDs in the respective polymer and glass matrices. The initial part of the flyer velocity profiles (dashed red curves) shows the flyer approaching the sample at $U_f = 0.6 \text{ km/s}$ in the polymer case and at $U_f = 0.79 \text{ km/s}$ in the glass case. Upon impact at $t = 0$, the velocities drop to the level denoted $U_p$ over a time interval denoted $\tau_{drop}$. $U_p$ denotes the velocity of the flyer/sample interface. The value of $\tau_{drop}$, typically a few nanoseconds, arises from the imperfect planar contact process due to the curvature and roughness on the order of microns of the flyer and target surface pair. Despite the different values of $U_f$ shown in Figures 3(a) and 3(b), the values of $U_p$ for the polymer and glass samples are nearly identical ($0.49 \text{ km/s}$ and $0.47 \text{ km/s}$), due to the differing shock impedances of the samples. The value of $U_p$ remain steady for a time interval denoted $\tau_{pl}$ before decaying gradually to zero over several tens of nanoseconds. In the experiments reported here, $\tau_{pl} = 13.9 \text{ ns}$, and it represents the duration of steady-state shock drive in the samples.

The value of the steady-state pressure in the material can be determined from the measured value of $U_p$ and tabulated properties\(^{18,21}\) of the polymer and glass via the shock and particle velocity empirical relationship $U_s = a + b U_p$ and the Rankine-Hugoniot jump condition

$$P - P_0 = \rho_0 U_s U_p,$$

where $U_s$, $\rho_0$, and $P$ are the shock velocity, initial density, and pressure, respectively. Note that the Hugoniot pressure, $P$, is the steady-state stress, although it will be referred synonymously as pressure throughout the paper. For the PVA polymer, $a = 2.46 \text{ km/s}$ and $b = 1.565.\(^{21}\)$ Corresponding parameters for the liquid-deposited sodium silicate glass were not available, so we estimated the glass pressures based on parameters for a related soda-lime glass.\(^{22}\)

The calculated pressure in the polymer sample in Figure 3(a) is $2.0 (\pm 0.1) \text{ GPa}$, and in the glass sample in Figure 3(b) it is $6.5 (\pm 0.2) \text{ GPa}$. Since the QD-doped glass was on a closely impedance-matched glass substrate, shock reflections were minimal so the sample was subjected to a single-stage shock. The PVA has acoustic impedance lower than glass. When shock waves propagate from PVA to glass, there is a shock reflection back into PVA. Therefore, in PVA we have both direct and reflected waves and the pressure rises in the PVA as a two-stage process.

The end of the steady shock drive in Figure 3, at $19 \text{ ns}$ for the polymer and $15.5 \text{ ns}$ for the glass, is indicated by vertical green lines. With the glass sample (Figure 3(b)), the blueshift and intensity loss are close to their maximum values at the end of the shock drive, but with the polymer sample, the blueshift and intensity loss are just beginning, and these parameters continue to increase significantly until about $60 \text{ ns}$. The slower time response of the polymer is possibly due to its sluggish viscoelastic shock-compression behavior.\(^{23}\) The glass response is much faster because glasses are elastic below the Hugoniot elastic limit (HEL, reported to be $\sim 3.5 \text{ GPa}$ by Alexander and co-workers.), and approximately fluid above this limit.\(^{24}\)

### B. Shock-induced spectral blueshift

Time-dependent QD spectral blueshifts at a series of pressures are shown in Figures 4(a) and 4(b) for polymer- and glass-matrix samples. The values of the blueshifts at the end of the steady-driven shocks (defined as the blueshift at $t = 20 \text{ ns}$ for simplicity) and the maximum blueshift values are shown in Figures 4(c) and 4(d). Due to the slower response in polymer, the blueshift at $20 \text{ ns}$ is noticeably smaller than the maximum value. In the faster-responding glass, the blueshift at $20 \text{ ns}$ is already close to its maximum value.
Similar to experimental results we reported previously\textsuperscript{13} in both glass and polymer, the maximum blueshift does not increase monotonically with increasing shock pressure. Instead there is a turnover pressure where the blueshift reaches a peak and then starts decreasing at $4$ GPa in the polymer and $9$ GPa in the glass. The blueshift pressure turnover under shock is in stark contrast to primarily the monotonic blueshift increase (with no subsequent decrease) observed under static compression reported in Ref.\textsuperscript{12}.

There are two mechanisms that can lead to the turnover pressure dependence. The first one is due to the coupled temperature and pressure effect. Specifically, the energy gap of the QD is affected by elevated temperature during shock compression, especially after the “ring-up” process in the sample. According to the empirical Varshni equation, the temperature dependence of the energy gap ($E_g$) of semiconductor crystals can be described as

$$E_g = E_{00} - \alpha \frac{T^2}{T + \theta}, \quad (2)$$

where $E_{00}$ is the energy gap at $0$ K, $\theta$ is the Debye temperature (for CdTe $\theta = 160$ K), and $\alpha$ is the temperature sensitivity (for CdTe QD with $d \sim 3$ nm, $\alpha = 0.55$ meV/K).\textsuperscript{25} Eq. (2) suggests that the elevated temperature will decrease the energy gap (redshift). On the other hand, if the QD is under hydrostatic compression, the energy gap is related to the pressure as

$$E_g = E_{g0} + a_1 P - a_2 P^2, \quad (3)$$

where $E_{g0}$ is the energy gap at ambient pressure. For CdTe QD with $d \sim 3.4$ nm at room temperature, $E_{g0} = 2.01$ eV, the linear factor $a_1 = 46.0$ meV/GPa and nonlinear factor $a_2 = 2.1$ meV/GPa$^2$ are obtained from experiment.\textsuperscript{11} Evaluation of Eq. (3) indicates that $E_g$ increases monotonically with pressure (blueshift) before phase transformation occurs.\textsuperscript{11} Since temperature in the sample increases along with pressure during shock compression, there will be a competition between the temperature and pressure effects on the energy gap. Though the relation between shock temperature

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**FIG. 4.** Spectral blueshifts of QD photoluminescent emission in (a) polymer matrix with 586 nm emitting QDs and (b) glass matrix with 674 nm emitting QDs. The rise and fall of the blueshift are slower in the polymer. The transient blueshift dips in the rising edges for the polymer (in the 4.3–6.9 GPa range) possibly resulting from temporary appearance of small redshifted sideband (c.f. Fig. 2(a)). (c) and (d) Pressure-dependence of blueshifts near the end of steady-state shock stage (at 20 ns), depicted in blue, compared to the maximum blueshift, depicted in red. The maximum blueshift was reached much later, at $\sim 60$ ns, in the polymer. The blueshift versus pressure relation has a turnover at $\sim 4$ GPa in the polymer and $\sim 9$ GPa in the glass, possibly due to differences in their volumetric strains.
and pressure for PVA and glass is not available, a qualitative estimation of the effect can be obtained by using available experimental data of polymethyl methacrylate (PMMA) under shock compression. Calculated results based on the data of PMMA show a turnover point at 6.0 GPa with a blueshift of 19 nm. The real relation between shock temperature and pressure of PVA and glass should be different from that of PMMA and would lead to a turnover point at smaller or larger pressure level.

The second mechanism is related to non-hydrostatic deformation of the QD. As mentioned in the previous work and Ref. 12, non-hydrostatic strain states of the QD can result in lower blueshift and non-monotonic dependence on pressure. If the QD is embedded in elastic matrix material softer than the QD and under static uniaxial strain field, according to the solution of the Eshelby's inhomogeneity problem, the QD will undergo non-hydrostatic deformation with decreased dimension in loading direction and increased transverse dimension. However, when the matrix is under shock compression with pressure beyond its HEL, the QD will experience more hydrostatic loading conditions, especially after the pressure “ring-up” within the QD. So this mechanism will take effect with shock pressure below HEL. In the present study, the HEL for both the PVA and glass matrix is smaller than the examined pressure, so the first mechanism may be expected to play a more dominant role. The coupled temperature and pressure effects also explain the different turnover pressures observed with the QDs dispersed in the polymer (~4 GPa) and in the glass (~9 GPa) matrix.

The time dependence of the blueshift shown in Figures 4(a) and 4(b) reveals interesting features. At a given pressure, the blueshift is larger in the polymer than in the glass matrix. This difference can be attributed in part to different compressibilities and to different QD sizes, since the smaller-size QDs in the polymer have greater quantum confinement, which leads to larger bandgap changes at the same volumetric strain. In the polymer, the blueshift rise times and fall times were much greater than in the glass. Furthermore, in the polymer, the blueshift does not return to its original preshock value within the 450 ns time window of the experiment, whereas within this time window, the blueshift recovery is nearly complete in the glass. Clearly, there are two recovery processes in the glass samples, one faster and one slower, suggesting two different relaxation stages of volume deformation of QDs embedded in the shocked glass matrix.

An interesting effect is noted in the blueshift for the polymer matrix samples at intermediate (4.3–6.9 GPa) pressures in Figure 4(a). The blueshift first rises, dips, and increases again, within all within a few tens of nanoseconds. This effect is clearly real and cannot be attributed to data analysis alone. Note that a smaller, slightly redshifted peak or shoulder appears in Figures 2(a) and 2(c) at around 30 ns. Since the blueshift was computed by the method of moments that finds the average spectral wavelength, the appearance of a secondary peak slightly pulls the average value to longer wavelengths and leads to the transient blueshift dip. As discussed below, the appearance of a new peak or a peak shoulder would also increase the effective peak width determined via the second moment of the spectrum. This blueshift dip prominent in the polymer samples is not observed in the glass samples with larger QDs. The origin of this smaller, secondary peak is unclear based on photoluminescent emission spectra alone. It may, however, be related to fluorescence resonance energy transfer (FRET) between adjacent QDs of slightly different sizes or, alternatively, a shock-induced QD phase transformation. It has been reported that under hydrostatic conditions there is a zinc blende (ZB) to cinnabar structure phase transformation in bulk CdTe at ~4 GPa. For nano-sized CdTe QDs, the phase transformation pressure is expected to be size-dependent and higher. A shift of band gap energy to lower levels (longer wavelength) was also reported for ~3 nm CdTe QDs at 5.8 GPa and attributed to cinnabar to rock salt phase transformation. The phase transformation pressure of QDs under shock compression conditions is lower than that under hydrostatic compression, presumably due to the effects of shock induced shear stresses influencing the phase transformation kinetics.

C. Relative photoluminescent emission intensity decrease

Figures 5(a) and 5(b) show relative (fractional) intensity decrease as a function of time for the same experiments as in Figure 4. The corresponding end of shock (20 ns) and maximum intensity decreases are shown in Figures 5(c) and 5(d). Unlike the blueshift exhibiting a pressure turnover, the intensity loss increases monotonically with increasing shock pressure.

The intensity loss has an initial steeper drop followed by a slower recovery. The intensity loss does not show anything comparable to the blueshift dip seen in Figure 4(a) that arises due to the transient appearance of a redshifted photoluminescent emission band in the polymer samples. The initial intensity drop is slower in the polymer matrix than in the glass matrix, but the much slower intensity recovery is about the same in polymer and in glass. Additionally, the initial intensity loss is slower than the initial blueshift rise, which can be seen by comparing the earliest-time data in Figures 4 and 5 for the polymer and glass matrix samples. These features can be qualitatively understood by considering the different nature of the blueshift and intensity loss processes. The blueshift is an instantaneous electronic response to shock compression, whereas the intensity loss involves shock-induced changes in QD temperature and surface defect states as described in the following paragraph. Because of this reason, the intensity loss may be slower than the spectral shift.

To understand the intensity-loss process, we note that undoped QDs undergo significant intensity loss at temperatures above 100 °C, conditions that the QD composite samples can easily reach during the shock pressures reached in the present work. For example, the estimated shock temperature for PMMA increases from ~50 °C to ~220 °C when pressure increases from 2 GPa to 7 GPa pressure. This loss of photoluminescent emission intensity is most likely closely related to the same process that causes the well-known QD blinking effect. The prevailing explanation for blinking is...
that sometimes excited QDs undergo a transfer of either electrons or holes to surface state traps which are dark defect states for which back-transfer is slow. QD blinking is known to become more prominent at higher temperatures. In our shock experiments, some dark defect trap states may be present initially and more may be created by extreme conditions of shock loading such as shearing between the QD and matrix due to uniaxial compression; hence, shock-induced intensity loss may be much more dramatic than the infrequent blinking process.

In the dark state trapping picture, the initial intensity drop is slower in polymer (Figure 5(a)) than in glass (Figure 5(b)) due to the slower response of the polymer to shock-compression. The initial intensity drop in both the polymer and in the glass is slower than the blueshift because it would take several cycles of laser excitation and QD relaxation to achieve the maximum dark state population, so intensity loss would lag behind blueshift. The recovery of the QD photoluminescent emission intensity, which is slow on the experimental timescale and similar in the polymer and glass, is insensitive to the faster response of the glass to shock-compression because it is controlled by the dark state lifetime.

D. Shock-induced spectral-width reduction

Figures 6(a) and 6(b) show the spectral widths of the QD photoluminescent emission peaks obtained from the square-root of the second moment of the spectrum. Since the QD peaks are nearly Gaussian, this quantity is equivalent to the standard deviation. It is striking that shock-compression causes the peak widths to decrease. Despite the high pressure, the anisotropic stress, and the high temperatures, the QD photoluminescent emission peaks actually sharpen up due to shock-compression. Figures 6(c) and 6(d) show the maximum value of the shock-induced peak narrowing (reduction) plotted as a function of shock pressure. At a given pressure, the shock-induced peak width reduction is smaller in glass than in polymer. The onset of the peak width reduction is faster in glass but the return to the initial width is about the same in both, the polymer and glass. The peak width stops decreasing at ~4 GPa in the polymer but it keeps increasing up to 12 GPa in the glass matrix.

As shown in Figures 6(a) and 6(b), at the higher shock pressures, the time-dependence of the shock-induced peak width has a non-monotonic behavior. The width first undergoes a temporary increase and then a longer-lasting decrease. The temporary increase is likely due to the appearance of the redshifted sideband responsible for the dips in the shock-induced blueshift seen in Fig. 4(a).

At present, we do not have enough data for a definitive explanation of the shock-induced photoluminescent emission narrowing, but it may be a result of the size distribution of the QDs in the samples, since the emission peak width in QD emitters is ordinarily associated with a size distribution that produces a distribution of emission frequencies. The shock-induced peak narrowing effect occurs in association with a large shock-induced intensity loss, so one possible explanation for the peak narrowing might involve a size-dependence on the surface trapping process.

E. Multiplex probing with dual QDs

The use of multiple-wavelength emitters to study the behavior of the different particulates that comprise
heterogeneous samples would be possible only if there is minimal energy transfer between the different-color QD probes. To test the feasibility of multiplex probing with dual QDs, we performed an exploratory experiment in which two different kinds of MEA-capped CdTe QDs, emitting either at 582 nm or 632 nm, were dispersed in a PVA polymer matrix sample. Figure 7(a) shows the time-stream photoluminescent emission spectra of a dual-QD sample shock-compressed at a steady-state pressure of 4.2 GPa. It can be seen that both emission peaks are present and well-distinguishable throughout the experiment. The method of moments is not appropriate for a multiple-peak measurement, so the intensity losses, blueshifts, and peak width changes were determined by the more laborious procedure of fitting the spectra to a pair of Gaussian lines, with the results shown in Figures 7(b)–7(d).

It can be seen in Figures 7(b)–7(c) that the two types of QDs have similar time-dependent blueshift and intensity loss behaviors, except that the effects are always greater for the smaller (i.e., the blueshifted) QDs. The spectral peak width dynamics are, however, quite different for the two types of QDs. The larger QDs show a prominent and temporary peak width increase around 45 ns, followed by a small decrease in width, whereas the smaller QDs show a much smaller and temporary peak width increase before settling into a small peak width decrease. The dissimilar peak width responses are most likely related to the different size-dependent phase transition dynamics. These results confirm that the two types of QDs do not exchange significant quantities of electronic energy and behave independently, thereby providing a strong indication for the feasibility of multi-wavelength sensing of shocked materials with embedded QDs.

V. SUMMARY AND CONCLUDING REMARKS

We have studied the time-resolved photoluminescence of QDs dispersed in PVA polymer matrix and silicate glass matrix under shock-compression at pressures from 2 to 13 GPa. Due to the different compositions, charges, and polarities of these different matrices, a positively-charged QD was needed for the polymer, and a negatively-charged QD for the glass matrix. Otherwise, incompatibility with the matrix would lead to QD aggregation and abnormal photoluminescence behavior. The 586-nm emitting QDs in the polymer were also slightly smaller than the 674-nm emitting QDs in the glass. We also carried out an experiment with QDs of two different sizes (582 and 632 nm emitting QDs) dispersed in the same PVA polymer matrix.

The arrival of a shock front in the samples causes the QD photoluminescent emission to blueshift, lose intensity and emit in a narrower spectral line. The blueshift arises due to the effects of shock-compression on quantum confinement. Compressing the QDs causes the bandgap to widen and the wavelength to decrease. With shock-compression, the decrease in wavelength (or blueshift) increases with pressure reaching a peak. Above the turnover (peak) pressure, the blueshift decreases, in contrast to the static high pressure blueshift, which increases monotonically with pressure. The blueshift turnover results from the coupled temperature and pressure effect on the band gap of the QDs.

The blueshift is expected to be an instantaneous electronic response to strain. In the limit of instantaneous mechanical response of the embedding matrix medium, the response time of the QDs in terms of their blueshift should be limited by the shock transit time across the sample. That or something close to it appears to occur with the sample.
containing QDs dispersed in the glass matrix. The 30 μm sample thickness corresponds to a transit time of ~6 ns, and the blueshift rise time is observed to be of a similar time scale. However, the blueshift response of QDs in the PVA polymer matrix is much slower, with rise times of ~70 ns. It appears that when the mechanical response of the matrix medium containing QDs is sluggish, as in the case of PVA, then the QD photoluminescent emission dynamics are controlled by the response of the surrounding medium.

The intensity loss process was observed to be always slower than the blueshift process. This is because the intensity loss requires both the extreme conditions associated with shock-compression, plus a finite-duration photophysical process. In the case of QDs, we believe that the intensity loss arises from effects similar to those that cause QD blinking, which is usually attributed to electron, or hole, transfer from the QD to dark surface state traps. Once the QDs are subjected to extreme conditions, it takes multiple cycles of excitation and decay to populate a surface trapped state. Even though the intensity loss effect is slower than the blueshift process, intensity losses of up to 90% are much easier to measure than the observed blueshifts, which in the pressure range studied in the present work, are a fraction of the ambient pressure photoluminescent emission linewidth.

The spectral line narrowing (or peak width reduction) is more difficult to measure accurately than the blueshift and intensity loss. However, it is a useful observation and it is remarkable that the photoluminescent emission narrows under shock-compression. Physical properties such as x-ray diffraction peak widths and optical emission spectra are much more likely to broaden under shock-compression due to the increased temperature and pressure gradients causing changes in lattice spacing. Thus, the peak narrowing (reduction) effect seen with the QDs must result from an interesting dynamical process. It seems possible that the peak reduction effect results from a shock-induced narrowing of the size distribution of the emitting QDs, likely due to the QDs losing intensity by populating dark surface states.

Finally, we have demonstrated the feasibility of multiplex detection of different species in a shock-compressed material, based on an experiment in which two different sized QDs were embedded in the same polymer matrix. Our results show that the necessary conditions for multiplex detection, namely, independent behavior of the different color QDs, can be met.

The dynamic response of QD photoluminescence has been shown in this work to depend on QD composition, size, concentration, and the mechanics of the embedding medium under shock-compression. The diversity of behaviors and the shock-compression responses of the matrix materials complicate the interpretation of photoluminescent emission spectra under shock-compression loading; but, it also offers the high potential for creating and employing diverse types of shock sensors for single and multiplex detection in complex heterogeneous materials, such as particulate media, as demonstrated by the present work.

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