Chemically-specific time-resolved surface photovoltage spectroscopy: carrier dynamics at the interface of quantum dots attached to a metal oxide

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Abstract

We describe a new experimental pump-probe methodology where a 2D delay-line detector enables fast (ns) monitoring of a narrow XPS spectrum in combination with a continuous pump laser. This has been developed at the TEMPO beamline at Synchrotron SOLEIL to enable the study of systems with intrinsically slow electron dynamics, and to complement faster measurements that use a fs laser as the pump. We demonstrate its use in a time-resolved study of the surface photovoltage of the m-plane ZnO (10 10) surface which shows persistent photoconductivity, requiring monitoring periods on ms timescales and longer. We make measurements from this surface in the presence and absence of chemically-linked quantum dots (QDs), using type I PbS and type II CdSe/ZnSe (core/shell) QDs as examples. We monitor signals from both the ZnO substrate and the bound QDs during photoexcitation, yielding evidence for charge injection from the QDs into the ZnO. The chemical specificity of the technique allows us to observe differences in the extent to which the QD systems are influenced by the field of the surface depletion layer at the ZnO surface, which we attribute to differences in the band structure at the interface.

1. Introduction

The study of ‘real-time’ charge carrier dynamics at material interfaces is becoming ever more important as next generation technologies increasingly access nanoscale dimensions. This is particularly the case for solar harvesting technologies, where a clear understanding of the dynamics of photoexcited carriers is of key fundamental importance. One such example is in devices utilising cost-reducing photovoltaic materials such as light-harvesting colloidal quantum dots (QDs) [1, 2] where a transparent conducting oxide (TCO) such as ZnO is used as photoanode [3-5]. These QDs have been shown to exhibit multiple exciton generation (MEG, creation of more than one pair of charge carriers from a single photon) for photon energies greater than around twice the effective band gap. The band gap is typically 1.0 eV, meaning MEG is present at wavelengths within the solar spectrum, giving potential for photovoltaic devices with enhanced efficiencies [1]. Considerable effort is currently being devoted to extending the photoexcited electron-hole pair lifetime in order to maximise the benefit of MEG. Quantum dot cores (known as type I QDs) that have been surrounded by a shell are known as type II QDs, and these are engineered such that the band structure leads to the photoexcited electrons and holes being separately trapped in the core and shell or vice versa, reducing the wavefunction overlap and extending radiative recombination lifetimes. Hence there is great interest in these type II structures for photovoltaic applications [6-8]. Here we study the carrier dynamics in heterojunctions formed both from type II QDs, and from type I QDs that have been shown to exhibit MEG, attached to a semiconductor photoanode, ZnO.

Carrier dynamics at the surface of semiconductors may be probed using time-resolved surface photovoltage (SPV) spectroscopy [9-11]. The space charge layer at the surface of semiconductors where the Fermi level is pinned by surface states within the band gap leads to band bending in this region. For n-type semiconductors the bands bend upwards due to the presence of a depletion layer, and downwards in p-type semiconductors due to an accumulation layer. A laser is used to promote carriers across the band gap, and, in the
case of an n-type semiconductor, electrons and holes migrate away from and towards the surface respectively (and vice versa in a p-type semiconductor). This additional electric field reduces the band bending and changes the energy of the valence band maximum in the space charge region. A synchronised X-ray source is then used to excite core level photoelectrons (using X-ray photoelectron spectroscopy, XPS), as the binding energy (BE) of the core levels in the space charge region at the surface shift as the valence band maximum shifts [9-11]. In n-type semiconductors these core levels shift to higher BE and in p-type semiconductors to lower BE [10]. By varying the time delay between the laser pump and XPS probe, the recombination dynamics is monitored over time, because the band bending and core level binding energies return to equilibrium as carriers recombine. Thus the recombination dynamics at the surface are elucidated [10-18].

For the case of colloidal QDs chemically linked to a semiconductor, this technique then provides a way to monitor the carrier dynamics in the semiconductor substrate upon photoexcitation of the QDs, i.e., as additional carriers are injected into the semiconductor conduction band upon laser illumination of the QDs [19]. Moreover, photoemission allows chemical specificity because the substrate and QDs can be monitored separately or in some cases simultaneously, simply by monitoring different core level photoemission lines. The technique then gives a chemically-specific diagnostic of charge transfer between the QD and the substrate.

Laser-pump X-ray-probe SPV experiments using pulsed lasers, where the pump laser is synchronized to individual X-ray pulses from a synchrotron, are limited by the time window available, determined by the synchrotron pulse length and repetition rate [11-13]. In the case of work by us at the SRS, UK, with the synchrotron operating in single-bunch mode, the maximum time window available was 320 ns, i.e., laser pump and X-ray probe pulses were incident every 320 ns with a set delay time between the two [10]. When measuring the transient SPV at the Si (111) 7x7 surface the dark carrier lifetime was in excess of the time window, and so a multi-pulse model was required to fit the carrier dynamics since the observed SPV transient was influenced by the residual SPV induced by preceding laser pump pulses [10]. Recently the use of angle-resolved time-of-flight (ARToF) analysers (such as at the BL07LSU beamline at Spring-8, Japan [14]), has allowed synchronisation of the pump laser pulse with hybrid mode pulses in multiple synchrotron periods, enabling SPV measurements extending to μs timescales, in studies of Si (111) [14-16], ZnO (0001) [17], and TiO_2 (110) [18].

A different approach is necessary to enable SPV measurements over much longer timescales (up to ms), which may be required where the electronic dynamics are intrinsically slow, for example due to persistent photoconductivity (PPC) as in ZnO. This has been enabled at the TEMPO beamline at Synchrotron SOLEIL, France [20], where pump-probe measurements are now possible over timescales ranging from 50 ps to ms or longer. Fast (ps - ns) measurements use a pulsed fs laser in combination with hybrid mode synchrotron radiation [21]. Dynamics with slower characteristic times are conveniently monitored using a CW laser. This is modulated using the output of a signal generator, where the period and duty cycle are easily controlled, and a fast XPS detection system is used to monitor XPS spectra at small time intervals over this modulation period [21], e.g. a core-level photoemission peak is monitored at 50 ns intervals as the CW laser is switched on and then off over ms timescales (typically on for 0.5 ms, and off for 0.5 ms). The key advantage of this system is the access this affords to long monitoring periods, from μs to ms, s and beyond. This is important for the SPV at the ZnO (10 00) surface where ms dynamics are exhibited, due to PPC controlled by trapping at band gap states associated with ionized oxygen vacancies [10, 19, 22, 23].

2. Experiment

In order to access relatively wide time domains in pump-probe XPS spectroscopy experiments, a two-dimensional (2D) delay-line detector was installed on a SCIENTA SES 2002 analyser at the TEMPO beamline at SOLEIL, [21, 24]. This allows XPS spectra over a narrow BE range (~2 eV) to be captured at least every 50 ns. The delay-line detector was developed in combination with custom-built software in order to monitor XPS spectra over time windows easily controlled by a signal generator. The triggering of this fast XPS detector is also used to control and modulate a CW laser (Coherent CUBE, 10 mW, 372 nm). The complete experimental setup is shown in Fig. 1. The laser is modulated as fast XPS spectra are captured every 50 ns over the period, T, of the laser modulation, e.g., for a period of 1 ms with a 50% duty cycle, 20,000 XPS spectra are captured as the laser is on for 0.5 ms and off for 0.5 ms. When the laser is switched on, the SPV shift is monitored using a shifting core level photoemission peak as charge carriers are injected into the conduction band, and a new equilibrium BE is established. The recombination time of these charge carriers is measured when the laser is turned off. Whilst the time resolution of the delay-line detector is 5 ns [21], the time resolution of the detection system is limited by the time spread of photoelectrons within the analyser and the speed of the electronics, which has been determined to be 150 ns. The signal-to-noise ratio (SNR) of a single XPS spectrum is low, and so typically in excess of 10,000 accumulations are taken (over 2-3 hours of data collection time) to obtain an adequate SNR. Very small SPV shifts on the order of meV changes in binding energy are easily detected under these conditions as shown in Fig. 2.
X-ray photon energies of 200 eV (for the Zn 3d core level) and 280 eV (for the Pb 4f core level) were chosen to maximize surface sensitivity (for examination of the space charge layer) without undue loss of flux. High-quality XPS spectra of the monitored photoemission peaks were taken in static mode in order to extract peak fitting parameters (e.g., spin-orbit splitting, full width at half maximum (FWHM)). These parameters were then applied to each fast XPS spectrum across the laser modulation time window, and the BE positions of the photoemission peaks obtained from the fitting were used to measure the SPV shift upon, during and after photoexcitation. Static XPS with and without laser illumination was also used to ensure that no long-term sample charging was occurring, i.e., that the photoemission peaks returned to the original BE positions after the laser was turned off.

Spatial overlap of the laser pump and X-ray probe beam at the sample position was achieved using a charge-coupled device (CCD) camera, and a photodiode was scanned across the sample position to measure the sizes of the laser pump and X-ray probe beams. The X-ray beam measured approximately 150 μm vertically by 100 μm horizontally, and the laser beam radius was approximately 3 mm. Ensuring the photoexcitation beam is substantially larger than the X-ray probe ensures an even photoexcitation fluence across the area being probed.

![Schematic diagram of the laser-pump X-ray-probe experiment at the TEMPO beamline at Synchrotron SOLEIL.](image_url)

**Fig. 1.** Schematic diagram of the laser-pump X-ray-probe experiment at the TEMPO beamline at Synchrotron SOLEIL. [20, 21]. A signal generator is used to modulate a continuous wave (CW) laser over a period \(T\) (Coherent CUBE, 10 mW, 372 nm), and fast XPS spectra are captured every 50 ns over the modulation period using a 2D delay-line detector installed on a SCIENTEA SES 2002 electron energy analyser [21]. The signal generator is used both to trigger the detection period and to control the time interval between spectra. Typically in excess of 10,000 accumulations over the modulation period are collected to obtain an adequate signal-to-noise ratio.

3. Materials

The non-polar \(m\)-plane ZnO (10\(\overline{1}0\)) surface was prepared according to an established recipe as has been detailed elsewhere [10, 19]. The final part of the cleaning procedure includes a period of annealing in 10\(^7\) mbar O\(_2\) at 700 K, which is required to heal an excess of oxygen vacancies created by previous argon ion sputtering and electron bombardment annealing cycles. This oxygen annealing period controls the concentration of oxygen vacancies, which is critical in controlling the final conductivity of the surface and the timescales for PPC. A change in the length of the oxygen anneal (for example from ten to twenty minutes) changes the dark carrier lifetime substantially [10, 19]. The SPV at the ZnO surface was monitored using the Zn 3d core level.

Two colloidal QD samples, type I PbS [2] and type II CdS/ZnSe core/shell QDs [19], were chemically-linked to a clean ZnO substrate using 3-MPA (3-mercaptopropionic acid) ligands, where the thiol links to the QD (Pb atoms in PbS QDs, Zn atoms in the ZnSe shell in CdS/ZnSe QDs) and the acid group to Zn in the substrate. In the PbS QD experiment, the substrate was initially prepared with a twenty minute oxygen anneal cycle, and for the CdS/ZnSe QD experiment the substrate was first prepared with a ten minute oxygen anneal. The preparation and characterisation of the QDs have been detailed elsewhere [19]. The type I (core-only) PbS QD sample used has been shown to exhibit MEG for photon energies greater than ~2.5 times the effective band gap [1]. The PbS QD sample was monitored using XPS of the Pb 4f core level, whilst the CdS/ZnSe QD was monitored simultaneously with the ZnO substrate due to the presence of zinc in both substrate and QD; a Zn 3d species associated with the QD (chemically shifted to a higher BE) was fitted along with the dominant Zn 3d peak associated with the substrate.

The laser photon energy (3.33 eV) was chosen for photoexcitation of the ZnO substrate and for photoexcitation of the PbS QDs and the ZnSe shell of the CdS/ZnSe QDs. For the PbS sample, this photon energy means that MEG is present since the photon energy is greater than 3 times the effective band gap.

4. Results
The SPV at the ZnO (10 10) surface upon laser photoexcitation was monitored with and without chemically-linked PbS and CdS/ZnSe QDs. The energy level line up diagrams for these two QD samples at the surface of ZnO are shown in Figs. 2 (a) and (e). The transient SPV, $\Delta V_{sp}(t)$, is fitted with a decelerating exponential decay as detailed elsewhere [10, 12, 13, 19]:

$$\Delta V_{sp}(t) = -\alpha kT \ln \left[ 1 - \exp \left( -\frac{t}{\tau_e} \right) \left( 1 - \exp \left( -\frac{\Delta V_{sp}^{tot}}{kT} \right) \right) \right].$$

Here, $\alpha$ is a material parameter likened to the ideality factor in a Schottky diode [9], $\tau_e$ is the dark carrier lifetime (the lifetime of carriers in the absence of a SPV), and $\Delta V_{sp}^{tot}$ is the total SPV shift.

![Diagram of ZnO and PbS QDs](image)

**Fig. 2.** Transient surface photovoltage (SPV) of quantum dots (QDs) chemically linked to ZnO (10 10) with 3-MPA (3-mercaptopropionic acid) ligands. The parameters extracted from fitting are shown. An X-ray photon energy of 200 eV was used for the Zn 3d core level (b, c, f-h), and a photon energy of 280 eV was used for the Pb 4f core level (d). Arrows denote laser modulation; the laser is turned on at $t = 0$ ms, and switched off at $T/2$, where the period $T$ is 1 ms for the PbS QD case, and 4 ms for the CdS/ZnSe QD case. In all cases laser photoexcitation at 372 nm ($h\nu = 3.33$ eV) was used, meaning that both QDs and the ZnO substrate were photoexcited (and for the CdS/ZnSe QDs, the shell of the QDs was photoexcited directly, enabling charge injection from the shell into the substrate as illustrated in (e)). Energy level diagrams (a) and (e) are adapted from [25, 26]. Dashed lines indicate a reduction in the band bending upon photoexcitation as carriers are promoted (or injected) into the CB.
Type I (core only) PbS QDs: (a) energy level line up diagram of the highest occupied molecular orbital (HOMO) and lowest unoccupied molecular orbital (LUMO) of the QDs with the ZnO conduction band (CB), valence band (VB) and Fermi level (EF); (b) transient SPV for the clean ZnO substrate, prepared with a twenty minute oxygen anneal cycle; (c) transient SPV of the Zn 3d core level after QD attachment; (d) transient SPV of the Pb 4f core level of the PbS QDs.

Type II (core/shell) CdS/ZnSe QDs: (e) energy level line up diagram, showing that for photon energies less than 2.7 eV, electrons and holes are trapped in the core and shell of the QD respectively; (f) transient SPV for the clean substrate. A shorter oxygen anneal cycle of ten minutes was used, giving an extended dark carrier lifetime compared with (b); (g) transient SPV of the Zn 3d core level component associated with the substrate (at a binding energy of ~10.18 eV) after QDs have been attached to the surface; (h) transient SPV of the Zn 3d core level component associated with the ligand/QD (chemically shifted to a higher binding energy of ~10.65 eV), where no SPV shift is observed.

Figs. 2 (b) and (f) show that the dark carrier lifetime, $\tau_\infty$, extracted from fitting equation (1) increases from 150 μs to 510 μs, and the magnitude of the SPV shift, $\Delta V_{\text{spV}}^\text{tot}$, decreases from 48 to 35 meV, as the oxygen anneal cycle length is halved, i.e., the dynamics are slowed as the oxygen vacancy concentration in the substrate is increased (as oxygen annealing acts to reduce the number of oxygen vacancies in the naturally nonstoichiometric ZnO$_{1-x}$) [10]. The timescale of the onset of the SPV upon laser illumination, modelled in an analogous way to the SPV decay (Eq. (1)), also increases from 45 to 90 μs.

Figs. 2 (c) and (g) show the changes in the carrier dynamics at the ZnO surface when the QD samples have been chemically linked to the surface. In both cases, the dynamics at the substrate surface speed up, even though the magnitude of the total SPV shift is reduced. For the PbS case, the total SPV shift is reduced to 15 meV, the dark carrier lifetime is reduced to 65 μs, and the SPV onset time is reduced to 5 μs. For the CdS/ZnSe case, the total SPV shift is reduced to 9 meV, the dark carrier lifetime is reduced to 140 μs, and the onset time appears instantaneous, i.e., the measured onset time is now limited by the time resolution of the experiment (0.15 μs).

Figs. 2 (d) and (h) show the transient SPV shifts now recorded using a core level photoemission line associated with the QDs rather than the substrate: in (d) the Pb 4f core level for the PbS QDs, and in (h) a Zn 3d core level species chemically-shifted to higher binding energy, associated with Zn atoms in the QDs attached to the ligand [19]. The Zn 3d core level associated with the CdS/ZnSe QDs does not shift upon photoexcitation, whereas a small but clear SPV shift of 3 meV is observed in the PbS QDs, with a dark carrier lifetime of 19 μs. The onset time of this SPV shift is also within the time resolution of the experiment, appearing as an instantaneous shift at $t = 0$ ms.

5. Discussion

The change in the magnitude and carrier recombination lifetimes for the clean ZnO substrate with two different oxygen anneal cycle lengths (20 minutes in Fig. 2 (b) and 10 minutes in Fig. 2 (f)) is consistent with the PPC being controlled by the oxygen vacancy concentration; a longer oxygen anneal cycle reduces the oxygen vacancy concentration. Dark carrier lifetimes recorded under similar conditions, but in different experiments are found to be reproducible to around a factor of two [10]. Band gap states associated with ionized oxygen vacancies have been implicated in this PPC, with recent experimental evidence supporting the theory proposed by Lany and Zunger [10, 19, 22, 23]; in this scenario, the PPC persists for longer as the number of oxygen vacancies (which, in their unionized state, act as efficient hole traps) increases [10].

Deposition of QDs chemically linked to the surface with 3-MPA ligands reduces the total SPV shifts observed in the ZnO substrate, but also reduces the dark carrier lifetimes and the onset times of the SPV shifts (Figs. 2 (c) and (g)). The presence of QDs linked to the surface is expected to change the equilibrium band bending, $V_g$, if the QDs donate or withdraw charge from the surface, and the total SPV shift, $\Delta V_{\text{spV}}^\text{tot}$, depends on the equilibrium band bending according to [27]:

$$\frac{\Delta V_{\text{spV}}^\text{tot}}{kT} \exp\left(\frac{\Delta V_{\text{spV}}^\text{tot}}{kT}\right) = n_r \exp\left(\frac{V_g}{kT}\right),$$

(2)

where $n_r$ and $n_o$ are the photoexcited carrier density and the doping carrier concentration respectively. In all cases the laser photoexcitation fluence was the same, indicating a change in equilibrium band bending. The reduction in the total SPV shift observed on QD deposition in both cases here suggests that the QDs donate electrons into the ZnO surface. In the absence of any new mechanism for carrier injection or recombination, this reduction in the total SPV shift is expected to give rise to an increase in carrier lifetime, rather than the decrease observed in Figs. 2 (c) and (g), since (for carriers recombining by thermionic emission across the surface depletion layer) the photoexcited carrier lifetime, $\tau$, varies with SPV shift, $\Delta V_{\text{spV}}$, according to [12]:
\[ \tau = \tau_0 \exp \left( \frac{-\Delta V_{sp}}{\alpha kT} \right) \]

In this model, the dynamic change in the photoexcited carrier lifetime with SPV shift requires the decelerating exponential model given in Eq. (1), where the lifetime increases as SPV shift is reduced. The fact that the onset times for the SPV shifts are faster, rather than slower, when the equilibrium band bending is reduced by attachment of QDs suggests direct charge injection from the QDs to the ZnO conduction band (CB) as anticipated according to the energy level line up diagrams shown in Figs. 2 (a) and (e). In the case of CdS/ZnSe QDs the onset appears instantaneous suggesting that the onset time is now within the experimental time resolution, and indeed there is evidence that charge transfer to an oxide substrate in similar QD systems may occur on ps and fs timescales [28-30]. The reduced dark carrier lifetime may indicate that the QD attachment is providing additional pathways for electron-hole pair recombination that avoids the long-lived PPC state (e.g., via new interface states). These changes in the recombination dynamics are important when considering light harvesting QDs with ZnO as photoanode as model systems for next-generation photovoltaics.

Transient SPV shifts upon laser illumination are caused by a change in the band bending at the interface, which is caused by the presence a depletion layer in the case of n-type ZnO. In the case of QDs, which are nm-scale semiconductor crystals, intrinsic band bending is not necessarily expected as the size of the QDs is much less than typical depletion layer widths. Therefore, no intrinsic SPV shift is expected when monitoring core level photoemission lines associated with the QDs. However, the electric field associated with the surface depletion layer in the ZnO substrate permeates the medium both above and below the surface, to an extent dependent on factors such as the dielectric constant and carrier concentration of the medium. We might expect to observe a SPV shift in the core levels of atoms associated with the QD if those atoms lie within this field; the size would depend on the position of the atom in the field, but is clearly expected to be smaller than the shift we observe at the ZnO surface. It is interesting that a small SPV shift of 3 meV (in the same direction as the ZnO substrate, i.e., to higher BE) is observed when monitoring the Pb 4f photoemission line in the PbS QD sample (Fig. 2 (d)). This indicates that the Pb atoms in the QD lie in the tail of the field associated with the surface depletion layer in the ZnO substrate. Analysis of this transient gives an onset time within the time resolution of the experiment, and the decay of the SPV shift yields a fitted dark carrier lifetime of 19 μs, faster than that observed in the substrate (65 μs). Both are consistent with fast intrinsic charge dynamics in PbS, not subject to the PPC observed in ZnO. Interestingly, a recent study by Neppel et al. [31] observed SPV shifts in N3 dye molecules attached to nanoporous ZnO. It is then interesting that we observe no SPV shift in the shell of the CdS/ZnSe type II QDs, monitored using a chemically-shifted component of the Zn 3d core level (Fig. 2 (h)); this may be associated with the potential barrier provided by the ZnSe shell, which may act to truncate the field associated with the ZnO substrate. We note that in this case photoexcited electrons are more stable in either the ZnO substrate or the CdS core of the QD than in the ZnSe shell (Fig. 2(e)). The difference in behaviour between the two QD systems points to the need for more experiments in this field, particularly using photon frequencies resonant with the lowest absorption energies of the QDs. However, the measurements presented here show the power of soft X-ray pump-probe experiments in allowing us to probe electron dynamics at these complex interfaces with atomic specificity.

6. Conclusions

A fast XPS technique, where a 2D delay-line detector enables fast (ns) monitoring of a narrow XPS spectrum at the TEMPO beamline at Synchrotron SOLEIL [20, 21], in combination with a CW laser, has been developed to enable the study of systems with intrinsically slow electron dynamics. This has allowed the charge carrier dynamics at the interface of colloidal QDs chemically-linked to ZnO (10 ̅10 ) to be monitored on ms timescales, required due to the PPC at the ZnO (10 ̅10 ) surface [10, 19, 22, 23]. The shift in core level photoemission lines upon photoexcitation, the SPV shift, has been measured over time windows of several ms. The chemical specificity of XPS allows for monitoring of not only the substrate but also of the QDs bound to the surface. In the case of PbS QDs, the observations of a small SPV shift of the Pb 4f core level indicates that the QDs lie within the field associated with the depletion layer at the ZnO surface. However, no transient SPV is observed when monitoring a Zn 3d core level component associated with the shell of type II CdS/ZnSe QDs, which may be due to the effect of the potential barrier at the QD surface provided by this shell. While further study is required to better understand the charge transfer and recombination dynamics at these interfaces, our work highlights the power of laser-pump X-ray probe techniques in measuring photoexcited carrier dynamics at these interfaces with chemical specificity.

Acknowledgments
References


