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Photo-Induced Charge Recombination Kinetics in MAPbI$_3$-$\chi$Cl$_\chi$ Perovskite-like Solar Cells Using Low Band-Gap Polymers as Hole Conductors.

Jose Manuel Marin-Beloqui,$^a$ Javier Pérez Hernández,$^a$ Emilio Palomares$^a$,$^b$

We measured the charge recombination kinetics using Transient Photovoltage (TPV) and Laser Transient Absorption Spectroscopy (L-TAS) in MAPbI$_3$-$\chi$Cl$_\chi$ perovskite solar cells using low band gap polymers as hole transport materials (HTM). Unequivocally, we assigned both transient decays to the recombination process between photo-injected electrons at the TiO$_2$ and the oxidised polymers.

Hybrid inorganic/organic double heterojunction solar cells have been the focus of intense research mainly to replace the liquid red/ox electrolyte in mesoscopic TiO$_2$ dye sensitized solar cells (DSSC)$^1$. In most cases, the same sensitizer is used for both architectures but the liquid electrolyte is replaced by a solution process hole transport material (HTM), generally either the spiro-OMeTAD$^2$ (chemical name: 2,2',7,7'-tetrakis(N,N-di-p-methoxyphenylamine) or a semiconductor polymer such as P3HT (chemical name: Poly(3-hexylthiophene))$^3$. However, the light-to-energy conversion efficiency ($\eta$) of these, so called, solid-state DSSCs have been generally below the efficiency values of their liquid counterparts mainly due to the lower photocurrent produced in the thinner mesoporous TiO$_2$ layers needed for solid-state DSSCs or the desorption of the sensitizer after the spin-coating of the HTM which frequently requires organic solvents such as chlorobenzene.

A breakthrough in the double heterojunction was reported in 2013 by several research groups$^4$ working independently by replacing the sensitizer using an organolead halide, either CH$_3$NH$_3$PbI$_3$ or CH$_3$NH$_3$PbI$_3$Cl$_3$ (MAPbI$_3$ or MAPbI$_3$Cl$_3$), and using spiro-OMeTAD as the hole conductor with outstanding efficiencies surpassing 15% under standard measurement conditions.

These results fuelled the interest of many research groups devoted to the study of DSSC and OSC (organic solar cells) and many different device configurations have been studied with published efficiencies ranging from 10% for thin-film approaches that avoid the use of HTM (see Scheme 1) to over 16% for those device structures that use HTM.

Most studies have focussed on replacing the spiro-OMeTAD HTM material with alternative HTM molecules with great success but studies on the photo-induced interfacial charge recombination reactions in these novel solar cells are more scarce$^5$. However, the study of the charge recombination processes under operation conditions are paramount to advance towards the theoretical maximum efficiency of MAPbI$_3$Cl$_3$ solar cells$^6$ by minimizing the interfacial losses, as for example the charge recombination reaction between the photo-injected electrons at the TiO$_2$ and the oxidised HTM (reaction 3 in Scheme 2).

In this communication we have fabricated efficient mp-TiO$_2$/MAPbI$_3$Cl$_3$ devices using as HTM material two semiconductor low-band gap polymers, PTB1 and PTB7, that show excellent hole mobility in OPV devices (4.5 $10^{-5}$ cm$^2$V$^{-1}$s$^{-1}$ and 5.8 $10^{-4}$ cm$^2$V$^{-1}$s$^{-1}$ for PTB1 and PTB7 respectively).

Scheme 1. Different device structures for MAPbI or MAPbICl based solar cells. (a) Thin-film approach without the use of a HTM. (b) Thin-film approach without the use of mp-TiO$_2$ and (c) the meso-structured approach using both mp-TiO$_2$ and HTM.
Figure 1 shows the IV curves for both type of solar cells and the standard solar cell using spiro-OMeTAD for comparison purposes.

To simplify our discussion the use of chemical additives such as 1,8 di-iodooctane (DIO), 4-tert-Butyl Pyridine (tBuPyr) or Lithium Bis(Trifluoromethanesulfonyl)Imide (LiTFSI) on the semiconductor polymers was avoided. Nonetheless, efficiencies as high as the ones reported for other semiconductor polymers in similar device structures have been achieved as listed in Table 1.

![Scheme 2](image)

**Scheme 2.** a) Energy diagram for the materials used in our devices (energy numbers in eV). The arrows correspond to the interfacial charge transfer reactions occurring during working conditions: (1) charge injection, (2) charge recombination between the photo-injected electrons in the TiO₂ and the holes in the perovskite material, (3) as in (2) but the recombination reaction occurs with the oxidised HTM. b) Molecular structure of the HTM materials used in our study. c) High resolution scanning electron microscopy image of the sections of our PTB1 based solar cell. H1= Gold contact (thickness: 89.6nm), H2=mp-TiO₂ (thickness: 261.0nm), H3=d-TiO₂ (thickness: 53.7nm), H4= PTB1 (thickness: 12.8nm), H5= perovskite (thickness: 189.4nm).

Table 1. IV curve parameters for our best devices.

<table>
<thead>
<tr>
<th>HTM</th>
<th>Jsc (mA/cm²)</th>
<th>Voc (mV)</th>
<th>FF (%)</th>
<th>PCE, η (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spiro-OMeTAD</td>
<td>21.6</td>
<td>839</td>
<td>60.5</td>
<td>10.9</td>
</tr>
<tr>
<td>PTB1</td>
<td>19.3</td>
<td>543</td>
<td>57.2</td>
<td>6.0</td>
</tr>
<tr>
<td>PTB7</td>
<td>18.0</td>
<td>699</td>
<td>59.8</td>
<td>7.5</td>
</tr>
<tr>
<td>PCPDTBT</td>
<td>10.3</td>
<td>770</td>
<td>66.7</td>
<td>5.3*</td>
</tr>
<tr>
<td>PCDTBT</td>
<td>10.5</td>
<td>920</td>
<td>43.7</td>
<td>4.2*</td>
</tr>
</tbody>
</table>


It is important to note that the devices were stable during the time necessary to characterize the recombination kinetics using TPV and L-TAS as shown in the ESI.

The devices were fabricated using a 50 nm thick layer of dense TiO₂ (d-TiO₂) and a 300 nm thick mesoporous TiO₂ (mp-TiO₂) layer all deposited onto FTO (fluorine doped Tin Oxide) glass with a resistance of 8Ω/cm² following previous literature. The CH₃NH₃PbI₃Clₓ was afterwards deposited on the mp-TiO₂ spin-coating a PbCl₂:CH₃NH₂I solution in DMF with a molar ratio of 1:3. The deposition of the metal oxide layers were carried out under ambient conditions but the subsequent deposition of the MAPbI₃Clₓ was carried out in a glove box ([H₂O] < 0.1 ppm and [O₂] < 100 ppm content). Each semiconductor polymer was spin-coated onto the FTO/d-TiO₂/mp-TiO₂/MAPbI₃Clₓ at 2000rpm for 60 s from a solution of chlorobenzene (10 mg/mL). The devices made using spiro-OMeTAD as HTM were fabricated using an identical protocol. **Scheme 2c** shows the cross-sectional scanning electron microscopy image for a PTB1 device. **Figure 2** shows the different L-TAS transient decays for the films upon excitation using our nanosecond Nd:YAG laser system (see ESI).
moreover, the decay lifetime for the spiro-OMeTAD is also slower, which implies that the back-electron transfer from the photo-injected electrons in the TiO₂ to the oxidised spiro-OMeTAD (reaction 3 in Scheme 2) is much slower than for the polymers. This can help explain why spiro-OMeTAD based devices are more efficient. We could estimate qualitatively the efficiency on the regeneration of the perovskite ground state from these signal amplitudes. However, the unknown molar absorption coefficient of the HTM polaron for both spiro-OMeTAD and the semiconductor polymers makes it difficult to ensure that the larger signal amplitude for spiro-OMeTAD corresponds to a better regeneration of the perovskite ground state. Further work is in progress regarding this. Nonetheless, additional data using Time Correlated Single Photon Counting (TCSPC) (see ESI) shows differences in luminescence quenching of the perovskite for the different HTM materials in perfect agreement with the L-TAS signal amplitude.

In order to validate our hypothesis that the L-TAS signal corresponds to the recombination between the photo-injected electrons and the oxidised HTM, we carried out TPV measurements in complete devices and compare the TPV decay with the L-TAS decay (Figure 3). This assumption cannot be done without the combination of both time-resolved techniques.

Figure 2. L-TAS decays, under ambient conditions, for the FTO/d-TiO₂/mp-TiO₂/MPB₁ₓCl₁₋ₓHTM films exciting at λₑₓ=500nm for the spiro-OMeTAD films and λₑₓ=700nm for both semiconductor polymer films. The probe wavelengths were λₚᵣₓ=1400nm for the spiro-OMeTAD, λₚᵣₓ=1100nm for the PTB7 and λₚᵣₓ=980nm for the PTB1 films.

As can be seen, the signal amplitude for the spiro-OMeTAD and the PTB1 films are greater than that for the PTB7 film and,

![Graph showing L-TAS decays for different HTMs](image)

Figure 3. TPV and L-TAS decays measured for the different devices. Notice the different Voc (right Y axis) for the different devices. (a) Spiro-OMeTAD, (b) PTB7, (c) PTB1 and (d) different TPV decays at different light bias (solar cell photovoltage at different light intensities) for the PTB1 device.

TPV is a technique that allows the measurement of the photo-induced recombination kinetics in complete devices under device working conditions. Upon light illumination the solar cells generate a photovoltage (Voc) that can be perturbed using a fast-pulsed laser. Upon excitation it generates a small increment of charges injected from the organolead halide to the

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TiO₂ raising the photovoltage to Voc₂. This increase in Voc, ΔV, is proportional to the raise in the Fermi level of the TiO₂. After the laser pulse, the original Voc₁ is restored with a lifetime (τ) that is the time that it takes the excess photoinjected charges (resulting from the laser pulse) to recombine and return the solar cell to the initial photovoltage Voc₁. It is important to note that the laser pulse must be small in energy (ΔV<15mV) to obtain a mono-exponential decay. Figure 3 shows the TPV decays for the devices studied in this work.

As can be seen, the TPV and L-TAS decays fit in good agreement implying that, indeed, in both cases the charge transfer reaction measured corresponds unequivocally to the recombination reaction between the photo-injected electrons, from the MAPb₁ₓCl₁₋ₓ, to the oxidised HTM. Moreover, as shown in Figure 3d, the increase of the light intensity (higher Voc) leads to faster TPV decays (faster charge recombination), which implies a direct relationship between increasing the charge density, increasing the solar cell Voc and faster recombination kinetics between the photo-injected electrons at the TiO₂ and the oxidised materials used as HTM.

Conclusions

We have fabricated efficient solar cells using MAPb₁ₓCl₁₋ₓ as the light harvesting material and low band gap semiconductor polymers such as PTB1 and PTB7 as HTM. The interfacial charge transfer dynamics were probed using micro-second to millisecond L-TAS. The L-TAS signals suggest better regeneration kinetics for the spiro-OMeTAD and the PTB1 polymer in complete MAPb₁ₓCl₁₋ₓ solar cells. Moreover, the L-TAS kinetics appear slower for the spiro-OMeTAD films. To confirm our hypothesis we measured TPV in complete devices, at different light bias, and we found that for low light intensities, close to the ones used in our L-TAS experiments, the TPV and L-TAS decays fit in perfect agreement. The increase of the light intensity leads to higher device Voc with faster TPV decays in all cases, suggesting that there is a direct correlation between photo-induced charge density, solar cell open circuit voltage (Voc) and faster recombination kinetics. Thus, in order to compare recombination kinetics in organolead halide based solar cells, it will be necessary to know beforehand the charge density in the devices under illumination as different device charge lifetime may well be due to the different charge density at the solar cell.

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Notes and references


Electronic Supplementary Information (ESI) available: Experimental section for device preparation. The double heterojunction scheme, the stability IV curve measurement as well as the description of the systems used to acquire the data using L-TAS, TPV and TCSPC.

See DOI: 10.1039/c000000xc

REFERENCES