Ultrafast polariton dynamics in strongly coupled zinc porphyrin microcavities at room temperature

P. G. Savvidis,1,2 L. G. Connolly,3 M. S. Skolnick,3 D. G. Lidzey,3 and J. J. Baumberg1
1Department of Physics & Astronomy, University of Southampton, SO17 1BJ, United Kingdom
2Department of Materials Science and Technology, University of Crete, and FORTH, P.O. 1527, 71110 Heraklion, Greece
3Department of Physics and Astronomy, University of Sheffield, Sheffield S3 7RH, United Kingdom

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Time and angle-resolved measurements reveal ultrafast dynamics of excitations in organic microcavities leading to dramatic modulation of probe transmission (∼30%). We find that the induced changes have both fast and long-lived components. Fast response times are defined by vibronic relaxation and intersystem crossing (S1→T1), whereas long-lived changes are attributed to a buildup of carriers in the nonradiative triplet state whose lifetime is longer than the repetition rate of the laser pulses, resulting thus in incomplete recovery of the ground state. Blueshifts of the lower polariton branch in the presence of the pump pulse indicate the presence of nonlinear interactions in the sample. However, there is no evidence yet for the pair-type scattering processes in porphyrin microcavities.

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The modified dispersion that accompanies strong coupling of embedded resonant absorbers in an optical microcavity1 has been shown to introduce completely new and extremely strong nonlinear optical interactions.2–5 This has been shown in a wide variety of ways in conventional III-V heterostructure microcavities grown using traditional semiconductor fabrication. In particular, the polaritons on the lower branch of the dispersion relation can undergo mutual pair scattering allowed by energy and momentum conservation.6 The polariton-polariton interactions are made much stronger by strong nonlinear optical interactions.2–5 This has been shown to introduce completely new and extremely strong nonlinear optical interactions in the sample. However, there is no evidence yet for the pair-type scattering processes in porphyrin microcavities.

In this work, we report on the initial study of exciton-polariton dynamics in strongly coupled organic microcavities filled with porphyrin based dye. Ultrafast changes in the transmitted probe pulse are observed when resonantly injecting polaritons at different points of the lower polariton branch with short pump pulses. The changes in transmission have both short- and long-lived components. We attribute the short-lived component to fast vibronic relaxation (τS2 ~ few ps) of the excited S2 state into the lower-lying T1 state, then followed by intersystem crossing (ISC) S1→T1 into triplet state (τISC ~ 100 ps), Fig. 1. Notably ISC dominates over ground-state S1→S0 relaxation because of typically long radiative lifetimes of the S1 state of the order of τS1 ~ few ns. Accumulation of carriers in the very slowly decaying (τT1 ~ 10 μs) nonradiative triplet states is responsible for the observed long-lived response. Contrary to demonstrations of polariton amplification in III-V microcavities, our experiments show reduction in the transmitted probe power indicating that no parametric scattering is observed in these samples. Our experiments clearly suggest that new organic microcavity systems that do not have fast deactivation pathways to lower-lying states are required in order for polariton-polariton interaction (parametric scattering of polaritons) to take place.

The microcavity (MC) under study consists of 125-nm-thick polystyrene film doped with the molecular dye tetraphenyl-porphyrin zinc (ZnTPP) and positioned between two dielectric distributed Bragg reflector (DBR) mirrors.14 Organic film was spin-cast onto the PECVD grown DBR mirror composed of 11 λ/4 pairs of SiN/SiO2 on a quartz substrate. The top DBR mirror consisting of 12 λ/4 layers of TeO2 and LiF was thermally evaporated onto the organic film to avoid damage to the film. The resulting cavity finesse was in excess of Q>400. The absorption spectra of the dye

![Diagram](https://example.com/diagram.png)

**FIG. 1.** Absorption spectra of zinc porphyrin dye and corresponding energy states S0, singlets, Tn, triplets. Microcavity is formed by positioning the organic film between two dielectric mirrors. Two new states LP, UP separated by Rabi energy are formed through resonant light–matter interaction.
is plotted in Fig. 1 showing peak absorbance at 2.76 eV which corresponds to the $S_2$ singlet excited state. Resonantly coupling this transition to the MC confined photon produces two new eigenstates of the system, namely the lower and upper polariton modes, Fig. 1 (dashed). Continuous wave (CW) characterization of these microcavities has been reported by Connolly et al.\textsuperscript{14} Here we map the microcavity dispersion relations by looking at transmission spectra at different angles using white light continuum from an optical parametric amplifier (OPA). In Fig. 2(a), both lower and upper polariton branches are clearly resolved, split by a Rabi energy of 135 meV.

Although it is possible to explore the nonlinearities using CW experiments, it has proved much more fruitful and clear to examine the nonlinear properties in the ultrafast regime, before combinations of different scatterings muddy the clear identification of the processes that occur. We have investigated the nonlinear response and the dynamics of the cavity polaritons, by employing angle resolved pump-probe experiments. This requires the use of two-color pump-probe ultrafast spectroscopy with the ability to tune the wavelength of each pulse independently in the 2.7–2.9-eV (460–430-nm) spectral region. These are derived from an optical parametric amplifier (OPA) pumped by a regeneratively amplified Ti:sapphire system (RegA) producing 5 µJ, 150-fs pulses at 50–250-kHz rate. The frequency doubled pump is used to resonantly pump the lower polariton branch, whereas the probe pulse is produced by mixing the residual pump pulse at 800–860-nm with the amplified idler pulse (around 950 nm) from the OPA, to generate tuneable pulses from 420–440 nm. The tuning ranges were a little restricted by the available mixing processes. Despite this, they allowed us to investigate many of the expected parametric scattering regimes.

A strong 150-fs pump pulse excites the sample at the chosen angle of incidence tuned in wavelength to resonantly excite lower branch polaritons. The resonance condition is easily observed from an increase in the transmission through the sample, and a spectral hole eaten out of the reflection spectrum (corresponding to the lower polariton). A weaker probe pulse arrives at the sample at a different angle of incidence (using our ultrafast goniometer\textsuperscript{6}), again resonantly tuned to the lower polariton, and the transmission of this probe pulse is measured in the presence and absence of the pump. The signature for polariton scattering is a pump-induced change in probe transmission, which depends on the energy momentum $(E,k)$ of the initial and final polariton states.

We first present data with the pump pulse set to $\theta$=40° and the probe pulse set to $\theta$=30°, injecting polaritons at a position from which pair scattering of pump polaritons would be possible. The polariton dispersion curve is shown in Fig. 2(a), together with spectra of transmitted pump and probe pulses at angles of 40° and 30°, respectively. This shows the high quality of the sample polariton dispersion, and the independently tuned pump and probe pulses. Typical time-resolved data are shown in Fig. 2(b).

Injecting pump polaritons results in $\sim$30% reduction in probe transmission in comparison to the probe transmission in the absence of the pump, even for large negative time delays. This dramatic reduction contrasts with experiments on III-V microcavities which show an induced increase in probe transmission (as energy is transferred from pump to probe beams) at zero delay. Typical pump powers of sub-1 W/cm$^2$ were used, however, the changes here were linear in pump power with no threshold effect. The change occurs across the whole spectral bandwidth, and no clear spectral shift of the peak position is observed (indicating that heating of the sample is not severe). The time-resolved change in the probe transmission (inset) shows that the reduction in transmission reaches maximum modulation $\Delta T$ of $\sim$5% within first 2 ps, decays slightly over 10 ps, and then persists for long times ($\mu$s). Notably, comparable modulation amplitudes have been reported for strongly coupled organic microcavities based on $J$ aggregates of cyanine dye,\textsuperscript{15} in which modulation amplitude was enhanced by more than one order of magnitude compared to bare organic film (no microcavity present). The presence of a long-lived signal is characteristic in metalloporphyrins because of the long lifetime of the first excited state and significant triplet intersystem crossing. This typically manifests itself as an incomplete recovery of the ground state because the lowest-lying triplet state becomes populated.\textsuperscript{16} The lifetime of these nonradiative triplet states is quite long (few tens of $\mu$s),\textsuperscript{17–19} much longer than the repetition rate of the laser pulses (40 kHz).

Similar results were obtained over a range of pump and probe angles. They suggest that parametric scattering is not observed in these samples. On the other hand, the observed changes indicate a rapid bleaching of the $S_2$ state after exci-
and are available to scatter to lower energy states. We also observe an ultrafast blueshift of the resonant polariton energy when the pump pulse is present, due to nonlinear interactions in the sample.\textsuperscript{21} Notably, the increased transmission around 2.8 eV (450 nm) is associated with the photobleaching of the Soret transition ($S_0 \rightarrow S_2$) also observed by O’Keefe et al.\textsuperscript{20} Again there is an ultrafast turn-on of the induced change, which decays on the same time scales noted previously. This leads us to believe that we excite real excitations in the molecules, which decay from the $S_2$ state into a long-lived state over a relatively short time scale ($\approx 100$ ps), most likely the $T_1$ state which is metastable. The occupation of this state causes a change in the oscillator strength and energy position of the $S_2$ state, leading to the induced changes observed.

In summary, there is no evidence yet for pair-type scattering processes in porphyrin strongly coupled microcavities at 300 K. Such processes are driven by exciton-exciton exchange interactions, for which there is preliminary evidence in our experiments. However, a small size of the excitons in these organic semiconductors reduces exciton interactions substantially leading to higher thresholds for pair scattering. It is also possible that due to low-lying metastable states which compete rapidly for the excitations injected, the pair scattering appears to be masked. On the other hand, other systems such as strong-coupled $J$-aggregate microcavities are much more promising for observation of stimulated scattering, since relaxation to lower-lying states in this material is strongly suppressed. The effect of injecting excitations into these molecules is as yet substantially unclear.

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\begin{figure}
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\includegraphics[width=\textwidth]{fig3.png}
\caption{(a) Pumping away from the magic angle into the excitons, which recovers slightly. This may be due to transfer to the $T_1$, $S_0$ states on a 10–50-ps time scale. Further information can be gained from exciting the sample at higher energy into the bottleneck region of polariton dispersion (almost resonant to the $S_2$ exciton) as in Fig. 3(a).
In this case, we see similar results, with a decrease in probe transmission, which is larger in magnitude than when pumping at lower angles and energies. This may be due to excited excitons that remain for a longer time in the sample and are available to scatter to lower energy states. We also observe an ultrafast blueshift of the resonant polariton energy when the pump pulse is present, due to nonlinear interactions in the sample.\textsuperscript{21} Notably, the increased transmission around 2.8 eV (450 nm) is associated with the photobleaching of the Soret transition ($S_0 \rightarrow S_2$) also observed by O’Keefe et al.\textsuperscript{20} Again there is an ultrafast turn-on of the induced change, which decays on the same time scales noted previously. This leads us to believe that we excite real excitations in the molecules, which decay from the $S_2$ state into a long-lived state over a relatively short time scale ($\approx 100$ ps), most likely the $T_1$ state which is metastable. The occupation of this state causes a change in the oscillator strength and energy position of the $S_2$ state, leading to the induced changes observed.}
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