High-speed flow of interacting organic polaritons

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The overwhelming majority of the previous research has focused on adopting inorganic-based systems (typically semiconductor quantum wells) operating at cryogenic temperatures; however, this limits their potential technological applications. Due to their ease of fabrication, low costs and high binding energies24,25, organic polaritons are an emerging field with an outstanding potential, as demonstrated by the recent observation of room temperature condensation26,27. Conventional polaritonic structures are planar microcavities where the optical active layer is embedded between two Distributed Bragg Reflectors (DBRs). However, the high dissipation rate of organic planar microcavities (short lifetimes) together with their relatively low group velocity (typically 1–2 μm ps⁻¹) has so far prevented the observation of the fascinating physics related to the flows of light–matter particles as well as any possible applications in cascadable-on-chip technologies that have been intensively investigated in inorganic systems28–31. Here, we show organic polariton propagation for distances longer than 120 microns and with group velocities of ~ 50% of the speed of light in a single DBR structure operating at room temperature.

To obtain such fast velocities, an organic exciton is coupled with a high-quality Bloch surface wave (BSW). The BSW is an evanescent, lossless and propagating optical mode with very high group velocity32–34, that differs from other surface modes such as the Tamm states or surface plasmons because it does not undergo dissipation caused by metallic losses35. Due to their high-quality factor, BSWs are widely studied for sensing applications, and their remarkable potential for the field of on-chip optical manipulation has only recently been explored36–39. Unlike the bare optical BSW investigated in these works, the excitonic component of the Bloch surface wave polariton (BSWP) allows the exploitation of nonlinear interactions. Here, we demonstrate the blueshift of the polariton resonance with increasing particle densities, enabling the local dynamic manipulation of the BSWP dispersion by external laser pulses and also laying the groundwork for polariton devices with logic functionalities that operate at room temperature.

MATERIALS AND METHODS

The DBR fabrication process consists of the deposition of seven TiO2/SiO2 pairs (dTiO2 = 85 nm and dSiO2 = 120 nm) on a 130-μm thick glass substrate using an e-beam evaporator.
A 35-nm thick layer of a perylene derivative (Lumogen Red F305), a small molecule with the absorption and emission spectra shown in Figure 1a, is thermally evaporated onto the SiO2 top layer of the DBR, with the base pressure of $\sim 10^{-7}$ mbar, and at the deposition rate of $\sim 1.0 \text{ Å s}^{-1}$.

The resulting structure sustains only high-quality TE polarized BSWP modes. The extraction of the signals lying beyond the critical angle is obtained through a high numerical aperture (NA 1.49) oil immersion microscope objective positioned at the glass substrate side, obviating the need to use additional optical elements[6,41]. The spectrometer and the CCD imaging camera (Princeton Instruments, Inc., Trenton, NJ, USA) allow the energy resolution of the BSWP signal. In the non-resonant configuration (upper part of Figure 1c), a TM-polarized continuous wave laser at 2.33 eV is focused (2 μm spot radius) through a second objective (NA 0.45) from the organic deposition side to prevent any possible coupling of the excitation laser to the surface propagating modes. The laser intensity on the sample surface is limited to 10 W cm$^{-2}$, preventing damage to the organic layer. Path A in Figure 1c shows the optical configuration used for the space–energy and space–energy–image of the emission: the back focal plane (Fourier plane) of the detection objective is recreated with S1 selection, with critical angle and BSWP indicated by dashed blue and red lines, respectively. In the bottom part, a sketch of the resonant configuration, sketched in the bottom part of Figure 1c, is used for the resonant injection of the polaritons. To match the BSWP dispersion (both in energy and momentum), the oil immersion objective is used for both the excitation and detection of the polariton mode, that is, operating in the reflection configuration. The slit S2 is positioned at the image plane of the sample and allows the selection of different areas in the field of view. A 300-mm spectrometer with a 1200-μm width grating blazed at 600 nm and a high-sensitivity CCD imaging camera are used for collecting the energy-resolved images with an overall resolution of 0.4 nm. In the non-resonant configuration, a low-pass filter at 600 nm is used for the energy-resolved propagation measurements to cut-off the residual laser signal passing through the sample.

The experimental data are supported by transfer matrix (TMM) calculations; the thicknesses and refractive indexes of the layers constituting the DBR and of the organic material are evaluated by ellipsometric measurements (Supplementary Information).

**RESULTS AND DISCUSSION**

The calculated dispersion by TMM and the experimental data of the BSWP emission under non-resonant excitation are shown in Figure 2a in the top and bottom panels, respectively. The experimental emission perfectly matches the simulated polaritonic mode (dashed orange line on the experimental emission in Figure 2a) that deviates from the bare optical BSW (blue line) when approaching the exciton energy (yellow line at 2.13 eV), showing the anti-crossing behavior typical of strongly coupled systems with a Rabi splitting of 186 meV. Short-propagating evanescent modes attributed to the limit of the DBR optical band-gap (sideband modes) are also visible in both the simulated and experimental results at low energies and high momenta[42].

The BSWP is highly sensitive to small changes of the optical index at the surface of the device[43], and any spatial inhomogeneity of the

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**Figure 1** Material, sample structure and optical setup. (a) Absorption and emission spectra of Lumogen Red F305 in the solid state (35-nm thick thermally evaporated thin film). (b) Illustration of the sample structure. The organic layer is deposited on a DBR composed of seven pairs of dielectric oxides ($\text{TiO}_2/\text{SiO}_2$). (c) In the upper part, a sketch of the leakage radiation microscope setup in a non-resonant configuration is shown. The excitation source (at 2.33 eV) is focused on the sample from the organic deposition side. The emission is collected through a microscope objective with $\text{NA} = 1.49$, extracting the signals lying beyond the critical angle. The use of two detection arms together with a spectrometer and a CCD imaging camera enables visualization of the energy-resolved space and momentum maps. The slit (S1) are used for the selection of specific directions of propagations. Inset: experimental 2D momentum space with S1 selection, with critical angle and BSWP indicated by dashed blue and red lines, respectively. In the bottom part, a sketch of the resonant configuration is shown. The excitation laser (100-fs laser pulse) is an energetically matched with the BSWP mode and focused through the oil immersion objective. The dispersion map of the reflected light is visualized on the CCD camera. The slit (S2) are used for the selection of the excitation spot area to cutoff the propagating signal and reveal the BSWP as a dip in the reflectance map.
Figure 2 Dispersion and space propagation of the BSWP. (a) Top panel: TMM calculation of the electromagnetic mode dispersions. The light cone is delimited by the red line, and the blue line indicates the bare optical BSW and the yellow line indicates the exciton energy (2.13 eV). The sharp signal in the center of the image is the BSWP mode, and the signal at higher momenta corresponds to the sideband modes of the DBR stop band. Bottom panel: Experimental measurement of the emission dispersion. Theoretical BSWP (dashed orange line), obtained with all the parameters extracted from the materials constituting the as-grown structure (Supplementary Information), fits the experimental results exactly. A weaker signal from the sideband modes is also detected. (b) Space map of the non-resonant excited polariton emission; the white arrows are a guide for the eye indicating the radial directions starting from the excitation spot, that is, the BSWP directions of propagation. (c) The emission intensity profile at 1.925 eV as a function of the detection angle. In the inset, the experimental data (blue dots) are fitted with a Lorentzian function with a full width at half maximum of 0.4° (red line).

Figure 3 Polariton propagation. (a) Space propagation of the BSWP after filtering in momentum space and with a low-pass filter at 600 nm. (b) Space propagation in the energy domain (associated to a). The red dashed line indicates the position of the excitation spot. (c) Propagation lengths versus energy and exciton fraction. (d) Intensity profiles (logarithmic scale) of the energy-resolved polariton propagation at different exciton fractions. At 1.97 eV (blue line), with the excitonic component >20%, the propagation distance is ~30 μm; at 1.92 eV (red line), the exciton fraction is 12.7% and the propagation length is 120 nm. At lower energies (1.88 eV, black line), an initial rise of the intensity is observed, followed by a long decay of 200 μm. (e) Polariton lifetime, versus in-plane wavevector and exciton fraction, evaluated from propagation lengths and group velocity. The red line is the TMM calculation of the bare BSW lifetimes.
sample deposition results in a broader BSWP dispersion\textsuperscript{14}. Therefore, the sharpness of the polariton mode that is clearly visible in the emission profile shown in Figure 2c, is a first hint of the high local homogeneity of the organic deposition.

The propagation of the polariton fluid on the sample surface is clearly visible in Figure 2b, where a few defects in the structure act as small perturbations on the flow trajectory. This also demonstrates that BSWP is an ideal workbench for the study of transport properties of dipole-like excitations strongly coupled with an electromagnetic mode; these are currently a focus of intense investigations and have recently been proposed to be strongly enhanced in the case of molecules and atoms placed inside an optical cavity\textsuperscript{44-47}.

Figure 3a shows the polariton flow along a given direction, obtained by filtering the signal in the back focal plane of the detection objective (slit S1 in the inset of Figure 1c). A low-energy pass filter at 600 nm is used to filter out the non-resonant laser light, resulting in the energy-resolved emission spectrum presented in Figure 3b. The residual emission of uncoupled excitons is detected close to the excitation beam position (red dashed line).

Sideband modes and short-living polaritons at higher energies populated through non-resonant pumping can also decay into the lower polariton branch; however, owing to their weak-propagating nature, they can only affect polariton formation very close to the excitation spot (less than 10 μm)\textsuperscript{48-50}. Although this ‘filling effect’ is small, it produces the initial rise of the signal at lower energies, as shown in Figure 3b and 3d (black arrow). At longer distances, the emission is attributed only to the long propagating BSWP and follows an exponential decay. The propagation length values at different energies, corresponding to different excitonic fractions, are shown in Figure 3c and exhibit ranges up to 300 μm (8% of exciton content), with a mean propagation length of ~120 μm and 13% of excitonic content.

The polariton lifetimes (τ\textsubscript{pol}) can be estimated from the experimental propagation length and group velocity (v\textsubscript{g})\textsuperscript{51}. The flow speed as a function of energy (E) is extracted directly from the BSWP dispersion as v\textsubscript{g} = \frac{∂E}{∂k\textsubscript{m}}. Group velocities ranging from 120 to 250 μm ps\textsuperscript{-1} (Supplementary Information) are obtained, ~100 times faster than in standard planar microcavities. At the energy of the maximum emission intensity, the polariton lifetime is ~1 ps (see data in Figure 3e) and the dissipative character of the BSWP is attributed mainly to the excitonic component.

Indeed, the DBR mode quality (evaluated from the ellipsometric data of the DBR structure) is estimated to be close to E/ΔE = 7000, giving a corresponding photon lifetime of ~3 ps (red line in Figure 3e). Although this value is about one order of magnitude higher than that obtained in the state-of-the-art organic microcavities, it is not as high as the lifetimes reported for inorganic planar microcavities, where lifetimes of some hundreds of ps have been obtained\textsuperscript{51,52}. However, thanks to the high speed of BSWP, the propagation lengths obtained in the present structure are comparable to the ones achieved with the best inorganic-based planar microcavities\textsuperscript{51,52}.

To fully exploit the potential of this system, we resonantly injected polaritons using the configuration depicted in the bottom panel of Figure 1c. Resonant excitation below the absorption energy of the exciton is a favorable configuration for the observation of the blueshift of the BSWP induced by polariton-polariton interactions only, avoiding the formation of a large reservoir of uncoupled excitons and reducing the heating of the sample and its degradation.

By selecting the sample area under the excitation spot through the slit S2 (Figure 1c), the propagating signal can be subtracted from the energy-momentum map, and the BSWP appeared as a dip in the reflectance dispersion, as shown in Figure 4a for the low-energy density signal of 150 μJ cm\textsuperscript{-2} (measured before the pump enters the microscope objective).

When the pumping energy density is increased to ρ\textsubscript{E} = 10 mJ cm\textsuperscript{-2}, the BSWP mode energy blueshifts. The blue (red) dots in Figure 4c represent the experimental peaks at different energies for the low (high) pumping power. The experimental dispersion at low power perfectly matches the results of the TMM calculation with the exciton peak at 2.13 meV (blue line in Figure 4c), whereas at ρ\textsubscript{E} = 10 mJ cm\textsuperscript{-2} the TMM reproduces the experimental dispersion resulting from an exciton energy blueshift of ~8 meV (red line in Figure 4c). Despite the reflectance spectra that are inevitably broader than the actual polariton dispersion due to the setup angular resolution (0.1°) that affects the momentum resolution (0.018 μm\textsuperscript{-1}), the shift of the polariton energy is larger than the mode line width and can be clearly resolved in our experiments, as shown in Figure 4b. This demonstrates the possibility of using polariton nonlinearities to tune a mode in and out of resonance with an external optical beam.

The wide range of energies covered by the laser pulse makes it possible to observe the trend of increasing polariton blueshift for energies approaching the exciton resonance (Figure 4d), as expected when increasing the exciton fraction of the BSWP mode.

Based on the BSWP depth in the energy–momentum map, it is reasonable to consider that a fraction of ~1% of the pumping power is transferred to the BSWP mode. From the shift of the exciton

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**Figure 4** BSWP nonlinearities. (a) Bare experimental data of the BSWP dispersion in reflectance configuration (i.e., with resonant excitation of polaritons) at 150-μJ cm\textsuperscript{-2} pumping energy. (b) Energy-resolved signal at in-plane wavevector k = 10.34 μm\textsuperscript{-1} under 150-μJ cm\textsuperscript{-2} and 10-mJ cm\textsuperscript{-2} excitation energy densities (blue and red dots, respectively, with Gaussian fit as solid lines), showing the blueshift of the BSWP resonance. (c) Dots are the experimental BSWP dispersions with 150-μJ cm\textsuperscript{-2} (blue) and 10-mJ cm\textsuperscript{-2} (red) resonant pump pulse. TMM calculations (solid lines) of the BSWP dispersion perfectly match the experimental values when considering an exciton energy of 2.13 eV (blue) and a blue-shifted (8 meV) exciton (red). The green ellipse depicts the full width at half maximum of the laser spot in energy–momentum space. (d) The expected blueshift as a function of the excitonic fraction (black dashed line) fits the experimental results (red dots) for an exciton blueshift of 8 meV.
transition, we can therefore estimate the interaction constant to be of the order of $10^{-3}$ eV $\mu$m$^2$, in accordance with the value indirectly extracted via non-resonant excitation of organic polaritons in a planar microcavity[26]. Thus, the strong coupling with organic molecules contributes to the enhancement of the nonlinear behavior of the BSW; nevertheless, despite the cryogenic temperatures, the nonlinearities offered by inorganic polaritons, are still between three and four orders of magnitude higher\textsuperscript{13,34}.

CONCLUSIONS

Room temperature ballistic propagation of light–matter excitations in an organic semiconductor for distances well beyond one hundred microns is demonstrated thanks to the adoption of a BSW model. Indeed, the BSW speed exceeds 150 ps $^{-1}$, which is two orders of magnitude higher than typical polariton velocities in standard planar microcavities. Moreover, through the resonant injection of a traveling polariton wave-packet, the exciton component of the BSWP manifests in the density-dependent self-energy of the system, showing blueshifts of the polariton resonance for increasing pump powers. This is the first direct measurement of this term, which arises from the excitonic components of polaritons, that enables tuning of the polariton energy and demonstrates a fundamental element for the implementation of polariton on-chip devices.

Although the interaction constant values reported for inorganic microcavities are higher than those observed in this work with organic semiconductors, these results are, however, very promising and can be further improved with the higher confinement of the electromagnetic field. In this respect, one of the most promising aspects of BSWP is that the surface can be easily patterned to localize the field and enhance the nonlinear optical response of organic materials, paving the way for the realization of ultrafast, low-loss polariton devices operating at room temperature.

CONFLICT OF INTEREST

The authors declare no conflict of interest.

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