Conductivity in organic semiconductors
hybridized with the vacuum field

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Much effort over the past decades has been focused on improving carrier mobility in organic thin-film transistors by optimizing the organization of the material or the device architecture. Here we take a different path to solving this problem, by injecting carriers into states that are hybridized to the vacuum electromagnetic field. To test this idea, organic semiconductors were strongly coupled to plasmonic modes to form coherent states that can extend over as many as $10^5$ molecules and should thereby favour conductivity. Experiments show that indeed the current does increase by an order of magnitude at resonance in the coupled state, reflecting mostly a change in field-effect mobility. A theoretical quantum model confirms the delocalization of the wavefunctions of the hybridized states and its effect on the conductivity. Our findings illustrate the potential of engineering the vacuum electromagnetic environment to modify and to improve properties of materials.

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Light and matter can enter into the strong-coupling regime by exchanging photons faster than any competing dissipation processes. This is normally achieved by placing the material in a confined electromagnetic environment, such as a Fabry–Perot cavity composed of two parallel mirrors that is resonant with an electronic transition in the material. Alternatively, one can use surface plasmon resonances, as in this study. Strong coupling leads to the formation of two hybridized light–matter polaritonic states, $P^+$ and $P^-$, separated by the so-called Rabi splitting, as illustrated in Fig. 1a. According to quantum electrodynamics, in the absence of dissipation, the Rabi splitting for a single molecule is given by

$$h\Omega_R = 2\sqrt{\frac{h\omega_0}{2\varepsilon_0
\nu}} \cdot \sqrt{n_{ph} + 1}$$

where $h\omega_0$ is the cavity resonance or transition energy ($h$ the reduced Planck constant), $\varepsilon_0$ the vacuum permittivity, $\nu$ the mode volume, $d$ the transition dipole moment of the material and $n_{ph}$ the number of photons present in the system. The last term in equation (1) implies that, even in the dark, the Rabi splitting has a finite value owing to the interaction with the vacuum electromagnetic field. This vacuum Rabi splitting can be further increased by coupling a large number $N$ of oscillators to the electromagnetic mode, as $h\Omega_R^2 \propto \sqrt{N}$ (ref. 1). In this ensemble coupling, in addition to $P^+$ and $P^-$, there are many other combinations of states, known as dark states$^2$, that are located in the middle of the Rabi splitting, as illustrated in Fig. 1a.

Polaritons have been extensively studied in inorganic and organic materials$^3$–$^{25}$. Vacuum Rabi splittings as large as 1 eV have been reported for strongly coupled molecules$^3$, thereby significantly modifying the electronic structure of the molecular material, as can be seen in the work function, the ground state energy shift, and the chemical reactivity$^8$–$^{10}$. In such situations, $P^+$ and $P^-$ are coherent collective states (Fig. 1b) that extend beyond the mode volume and may involve $\sim 10^5$ oscillators, as in the present systems.

This collective nature has been demonstrated by the coherent fluorescence of $P^-$ over distances of micrometres in strongly coupled molecular states$^{11,25}$.

Organic semiconductors have generated considerable interest due to their potential to create inexpensive and flexible devices that are easily processed on a large scale$^{26}$–$^{36}$. However, technological applications are at present limited by the low mobility of the charge carriers, which is associated with the disorder in these materials$^{30}$–$^{33}$. Hence, charge carrier mobility in organic semiconductors should also benefit from the extended coherence associated with light-hybridized states as compared to the normal carrier mobilities limited by the hopping between molecules and by scattering induced by molecular disorder.

Although electron-injected polariton LEDs and lasers have been reported in recent years$^{39}$–$^{41}$, conductivity mediated by light-hybridized materials has not been considered. One of the challenges in those studies has been to inject electrons into $P^+$ or $P^-$, as these states have not just well-defined energies but also well-defined momenta associated with their dispersion curves. In this regard, molecular materials have the advantage that the energies of the states are inhomogeneously broadened and fluctuating in time, resulting in spectral widths that facilitate polariton creation. Indeed the very first study of polariton electroluminescence was carried out on organic materials$^{42}$.

In this study, an initial screening was carried out on different organic semiconductors known to be air stable and exhibit good field-effect mobilities in solution-processed devices ($10^{-2}$–$10^{-1}$ cm$^2$ V$^{-1}$ s$^{-1}$ for optimized field-effect geometries). We focus here on three members of the aromatic diimide family, an example of which, PD12EH-CN$_x$, is shown in Fig. 1c (refs 34–36). The synthesis of this molecule is described in Supplementary Section I. The other molecules are the brominated version of the former, PD12EH-Br$_x$, and the diimide naphthalene polymer, P(NDI2OD-T2) (ActivInk N2200 from Polyera). Note that we

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Figure 1 | Light–matter strong coupling with organic semiconductors. a, Illustration of the strong coupling between a molecular electronic transition ($S_0$ to $S_1$) and an optical cavity mode of the same energy ($\hbar\omega_C$). Two new hybrid light–matter states $P^+$ and $P^-$ are formed, separated by the Rabi splitting $\hbar\Omega$. The energy level of the so-called dark states (DS) is also indicated. b, Illustration of the collective nature $P^+$ and $P^-$, forming a coherent state involving many molecules or molecular units ($M$). c, Molecular structure of PDI2EH-CN$_2$. d, Absorbance spectra of a thin film of PDI2EH-CN$_2$ (dashed blue) and examples of resonances of two hole arrays whose resonance matches the centre of the molecular absorbance. e, Electron microscope image of a hexagonal array milled in a Al metal film which provides the surface plasmon resonance that is strongly coupled to the molecule. f, g, Dispersion of the surface plasmon modes in the absence (f) and presence (g) of PDI2EH-CN$_2$ as a function of the period $P$ every 20 nm between 240 nm (top curve) and 460 nm (bottom curve). In g, the dashed lines are guides to the eye, with the vertical lines indicating the position of the PDI2EH-CN$_2$ absorption peaks shown in d and the curved lines indicating the shift in the plasmon resonances.

have chosen these organic materials in view of their very different charge transport capacities, but also because they all possess a lowest unoccupied molecular orbital (LUMO) energy level at $\sim 4$ eV (see Supplementary Section IV). This last observation rules out the injection capacity as a main parameter for the interpretation of the results appearing later in the manuscript when comparing their relative response. Instead, as we will see, the differences that are important are the intrinsic charge transport capacity of each molecular system, influenced by the film disorder. These compounds possess intense absorption bands (Fig. 1d) that can be used to strongly couple them to surface plasmon resonances. For this we used a hexagonal array of holes (Fig. 1e), in either Al or Ag films, which give rise to well-defined surface plasmon resonances,$^{37,38}$ also shown in Fig. 1d. The low work function of an Al electrode is generally favourable for electron injection in n-type semiconductors. In the case of a Ag electrode, we still have
good electron injection, as confirmed in our experiments (see Supplementary Section IV, Supplementary Fig. 6). Such efficient electron injection from metals with a high work function into PDI derivatives has been previously reported.\textsuperscript{15,36}

Figure 1f,g shows, respectively, the dispersion of surface plasmon resonances of the Ag hole array as a function of hole period in the bare state and when the array is covered with a thin film of PDI2EH-CN\textsubscript{2}. Unlike with Fabry–Perot cavities, in the case of strong coupling with surface plasmons on hole arrays the upper polariton is less visible owing to Fano-type interference.\textsuperscript{6} However, a Rabi splitting of \(\sim 0.7\) eV can be estimated, corresponding to 30\% of the transition energy, typical of the ultra-strong-coupling regime.\textsuperscript{67}

To assess the conductivity of strongly coupled organic semiconductors, we compare it to that of the bare molecular material, by recording current–voltage (I–V) curves for thin films of the semiconductors between two electrodes, as illustrated in Fig. 2a. First, a 100-nm-thick layer of Al or Ag was evaporated on a glass substrate through a shadow mask to yield a long 80-\(\mu\)m-wide strip. Then hexagonal arrays with period \(P\) were milled over a 55-\(\mu\)m-long area on the metal strip. The active channel, 50-\(\mu\)m long, was defined by milling gaps \(g\) of 200 nm separating this area from the source and drain electrodes (Fig. 2a). The width of these gaps was found to be very important, because a distance larger than the decay of the evanescent field (\(\sim\) the wavelength) of the surface plasmon would result in the current flowing through an area where the material would not be hybridized. Note that the array continues for a few micrometres on both electrodes to ensure that the current is already injected into the strongly coupled system before reaching the active channel (Fig. 2a).

The I–V curves were then recorded for different array periods and compared to that of an unstructured metal film also covered with the organic semiconductors (the reference). An example is shown for selected periods in Fig. 2b in the case of the PDI2EH-CN\textsubscript{2}. Similar experiments were carried out for the other two compounds. By plotting the current average at \(\pm 15\) V versus \(2\pi/P\), with \(P\) being the period of the array, two current resonances are clearly visible, as can be seen in Fig. 2c for both PDI2EH-CN\textsubscript{2} and P(NDI2OD-T2). Interestingly, the current peaks occur for different periods. These correspond exactly to the intersections of the molecular absorption peaks and the surface plasmon modes (1,0) and (1,1) of the substrate side of the hole array where strong coupling occurs (Fig. 2d). In other words, the current is boosted when the organic...
Figure 3 | Conductivity under strong coupling on Al arrays and gating. **a**, Spectra of Al hole arrays with and without PDI2EH-CN$_2$ as indicated. **b**, I–V curves for selected periods versus reference in the same geometry as in Fig. 2a. **c**, Three-terminal gating device geometry, composed of: Al (40 nm) acting as a top gate; ∼600 nm of Cytop, a polymer dielectric acting as a gate insulator ($\varepsilon_r = 2.1$); 100 nm of PDI2EH-CN$_2$ as the semiconductor layer; Al electrodes and nanohole array structure ($W = 50 \mu m$, $L = 50 \mu m$, $V_D = +60 V$). Note that the surface plasmon modes are only slightly modified by the presence of the dielectric, as most of the evanescent plasmonic field is confined within the organic semiconductor layer. **d**, **e**, Typical transfer characteristics ($I_D$–$V_G$) of top-gate bottom-contact FETs realized on the structure proposed in Fig. 3c. In particular the graphs show the difference between a gated structure with hole arrays of periodicity $P = 280$ nm (**d**) versus a reference structure with the same geometry but without nanohole arrays (**e**). The ratio of the mobilities is $\mu_{FET}(P=280\text{nm})/\mu_{FET(\text{reference})} \sim 10$, which corresponds essentially to the ratio of the current measured in the I–V curves.

The semiconductor is strongly hybridized with the surface plasmon modes. The current increases by over an order of magnitude in this regime. The third compound, PDI2EH-Br$_2$, showed no detectable change in current with conductivity in the pA range (see Supplementary Fig. 5). The differences between these compounds will be discussed below. The widths of the conductivity resonances observed in Fig. 2c reflect the widths of both the absorbance of the semiconductor and the plasmon resonance. For that reason, a true off-resonance with periodic arrays is hard to achieve, weakening the contrast between the conductivity minima and maxima with respect to what would be observed if the semiconductor and plasmon resonances were sharper. Please note that, in Fig. 2c, Ag and Al were chosen as the metal electrodes because, in each case, the charge injection is optimal for each electrode/semiconductor combination (see Supplementary Section IV). In addition, the injection from different metals was tested for PDI2EH-CN$_2$ and P(NDI2OD-T2), as reported in the Supplementary Fig. 2 for PDI2EH-CN$_2$ and Supplementary Fig. 6 for P(NDI2OD-T2).

The reference in the above data is always the unstructured metal film covered with the organic semiconductors. This reference shows a current larger than just the semiconductor, enhanced by the metallic substrate, but not shorted by the metal. Indeed, the
measured field-effect mobility results from the intrinsic organic semiconducting properties. We have also checked random arrays of holes, which show no current increase (Fig. 2b), in agreement with expectations, as there are no well-defined plasmon resonances in such systems.

In the case of PDI2EH-CN12, the same experiment was repeated on hexagonal hole arrays milled in an Al film with the spectra shown in Fig. 3a. The system shows (Fig. 3b) the strongest current resonance for a period of 280 nm, which again corresponds to the strongly coupled condition. The current increases by roughly an order of magnitude as compared to the reference consisting of PDI2EH-CN12 on a flat Al film. To further assess the electronic properties of the coupled system, a three-terminal gated configuration was prepared, as illustrated schematically in Fig. 3c. Figure 3d,e shows the current versus the gating voltage for the on-resonance sample in Al and the reference, respectively. The mobility extracted from this data shows an increase from 2.0 × 10−3 cm2 V−1 s−1—a value in agreement with the values extracted in a standard geometry (see Supplementary Table 1)—to 1.7 × 10−2 cm2 V−1 s−1. In other words the increased carrier mobility in the coupled system is responsible for most of the enhanced current.

To explain the above results, we introduced a simple theoretical model where each molecule is represented by two states forming the conduction and valence bands of the semiconductor. The molecular states are resonantly coupled by the vacuum field with the relevant Hamiltonian

\[
H = \left( -\sum_{j=1}^{M-1} J_j (\epsilon_j^c c_j^+ + \epsilon_j^h c_j^+) + \sum_{j=1}^{M-1} J_j (\epsilon_j^h c_j^+ + \epsilon_j^c c_j^+) - \sum_{j=1}^{M} \sum_{j'=1}^{M} (\epsilon_j^c n_j^e + \epsilon_j^h n_j^h) \right) + \sum_{j=1}^{M} \sum_{j'=1}^{M} (\epsilon_j^e h^c + \epsilon_j^h h^h + h.c.
\]

Here, in a second-quantized formalism, \( \epsilon_j^c (\epsilon_j^h) \) and \( \epsilon_j^e (\epsilon_j^h) \) correspond to the destruction (creation) of an electron in the conduction band and of a hole in the valence band, respectively, with \( n_j^e = \epsilon_j^c c_j^+ (n_j^h = \epsilon_j^h c_j^+) \) the local electron (hole) occupation, and \( j \) the site index (that is, the molecule location) for a system of size \( M \), where \( M \) corresponds to the number of molecules. The local coupling \( g \propto d_{a} \exp(-ik\ell) \) (with strength \( d_{a} \)) is mediated by the surface plasmon field (with a delocalized surface plasmon wavevector \( k \)) resonant with the interband transition \( \epsilon_{i}^{e,h} \), here represented by a destruction (creation) operator \( a_{i}^c (a_{i}^h) \). \( \ell \) is the intermolecular spacing, in the nanometre range, and thus \( k\ell \ll 1 \). The third term represents processes where a surface plasmon is coherently created when an electron is transferred from the conduction to the valence band \( (a_{i}^c c_{j}^h) \) and, vice versa, where a surface plasmon is coherently absorbed when an electron is promoted from the valence to the conduction band \( (c_{i}^e a_{j}^h) \). The last term represents possible local shifts of the energy \( \epsilon_{i}^c (\epsilon_{i}^h) \) for the electrons (holes), arising, for example, because of disorder.

The two-band Hamiltonian equation (2) is reminiscent of the well-known Jaynes–Cummings model; however it takes into account the dynamics of electrons and holes. The terms \( J_j^e \) and \( J_j^h \) represent the quantum mechanical tunnelling rates of the charged particles between neighbouring sites. In the absence of disorder \( (J_j^e = J, \epsilon_j^c = \epsilon_e \) and \( J_j^h = J, \epsilon_j^h = \epsilon_h \) for all \( j \)) and for \( g \ll g = 0 \), the first two terms in equation (2) imply extended Bloch-type wavefunctions for electrons and holes, which can in principle lead to a finite conductivity even at zero temperature. However, disorder in semiconductors in one dimension is known to invariably lead to wavefunction localization and vanishing conductivity. Indeed, this is shown in Fig. 4a for \( g = 0 \), where the probability density of a few chosen electronic states is shown to be exponentially localized for a situation where we have chosen disordered hopping rates \( J_j^{eh} = J + \delta J_j^{eh} \) and energy shifts \( \epsilon_j^{eh} = \epsilon + \delta \epsilon_j^{eh} \), with \( \delta J_j^{eh} \) and \( \delta \epsilon_j^{eh} \) sets of random values (see Supplementary Section VI).
On coupling the system to the vacuum electromagnetic field, it is expected that polaritonic states delocalized throughout the mode volume should appear\textsuperscript{42,43}. This is shown in Fig. 4b (black curve) for \( g = 1 \), where, remarkably, the electronic component of the polaritonic density is shown to be delocalized despite the strong disorder. This value for \( g \) was chosen so that the collective coupling \( g \sqrt{N} \) is larger than all other coherent and incoherent rates, allowing the strong-coupling regime to be reached (see Supplementary Section VI). Surprisingly, in addition to the polaritonic state we find that under conditions of strong coupling also other electronic states in the conduction band of the model are modified by the hybridization with the vacuum field and can become delocalized over the same length of the sample, as shown in Fig. 4b for three eigenstates. This should have a considerable impact on conductivity, which we investigate next.

To probe the dynamics, we solve the master equation for the full density matrix of the system (see Supplementary Section VI) in the limit of low charge density. To mimic experiments, the master equation includes dissipative processes such as spontaneous decay of the plasmon mode and dephasing of the electron and hole wavefunctions—arising, for example, from local molecular vibrations. Electrons are incoherently pumped into the system at a rate \( \gamma_P \), similar to the effects of an applied voltage. Figure 4c shows the equivalent of the \( I--V \) curves derived from this model for \( M = 30 \) sites and both \( g = 0 \) and \( g = 1 \). As can be seen, the current increases significantly in the presence of strong coupling. In Supplementary Section VI we show that in fact significant enhancement is obtained only in the strong-coupling regime, where the electronic wavefunctions hybridized with light are delocalized.

Despite its simplicity, the model clearly shows that the origin of the enhanced mobility under strong coupling can be explained by the extended coherence of the states induced by the hybridization with the vacuum field. This occurs over lengths that correspond to the mode volume (on the scale of the wavelength of the resonance), in contrast with the normal situation where the carrier is mostly confined to the molecular (or a few molecules) scale, in particular at room temperature.

It is important to note that, unlike for all conductors, disorder can strongly hinder charge transport whatever the transport mechanism. This is clearly shown in the case of PDIEH-BT\textsubscript{r}, which has a similar molecular structure to PDIEH-CN\textsubscript{r}, but a much lower mobility\textsuperscript{16} as a result of disorder. We found extremely weak resonant enhancement, if any, despite the strong coupling (see Supplementary Fig. 5). The P(ND2OD-T2) polymer presents an intermediate case where, despite the lower conductivity, the latter is enhanced by strong coupling. This is also confirmed by the model in the presence of large disorder, where, although the relative improvement of conductivity increases with disorder strength for a fixed light coupling \( g \) over the case \( g = 0 \), the overall conductivity tends to vanish.

The enhanced currents and the improved mobilities reported here demonstrate the potential of strong coupling for boosting conductivity in organic semiconductors. It is worth noting that at the same time the semiconducting properties are preserved. Although we used surface plasmon resonances to couple to the organic semiconductors, metal micro-cavities or dielectric resonators could also be used provided that the mode volumes are sufficiently small such that the strong-coupling condition can be met. In particular metallic Fabry–Perot cavities lead to even larger Rabi splittings than plasmonic arrays, which is important to increase the fraction of coupled oscillators in the sample, as we have shown elsewhere\textsuperscript{16}. The challenge, we have found, is to ensure that there is no leakage via the top mirror when measuring the current parallel to the cavity.

Conductivity of strongly coupled systems can be implemented with ease in a variety of designs and configurations, including in plastics, depending on the application. Finally, our results show the importance of engineering the electromagnetic environment of a material, not just for optical purposes, but also as a means to modify its properties through the formation of hybrid light–matter states. This opens new prospects for materials science as well as device technology.

Note added in proof: Since the submission of this work, two theoretical works have addressed the question of suppression of backscattering of conduction electrons for a two-dimensional electron gas off-resonantly coupled to a high-frequency classical electromagnetic field\textsuperscript{44,45}.

Methods

Methods and any associated references are available in the online version of the paper.

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Author contributions

T.W.E. conceived the idea and supervised the project. T.W.E. and E.O. designed the device experiments. J.G., J.A.H. and J.F.D. undertook the spectroscopic experiments. E.D., E.O., J.G., J.A.H. and J.F.D. fabricated and performed the device experiments. B.D., P.S., C. Genet and E.O. helped with the interpretation of the experimental data. G.P., C. Genet and J.S. contributed to the discussions and the preparation of the manuscript.

Additional information

Supplementary information is available in the online version of the paper. Reprints and permissions information is available online at www.nature.com/reprints. Correspondence and requests for materials should be addressed to T.W.E.

Competing financial interests

The authors declare no competing financial interests.
Methods

Two- and three-terminal devices including nanohole arrays. All the electrodes were realized on a glass substrate BK7 (25 × 25 mm) after standard cleaning procedures with sonication in Alconox solution (0.8% solution in milliQ water), then rinsed with water and sonicated for 1 h in (spectroscopically pure) ethanol. The glass substrates were then dried in a oven overnight. Ag and Al electrodes (100 nm) were fabricated using an electron beam evaporator (Plassys ME 300) at optimized working pressure (~10⁻⁶ mbar) and deposition rates (~0.4 nm s⁻¹). 

Plasmonic hole arrays were generated by the NPVE software program and milled using a Carl Zeiss Auriga FIB system. 0.5 wt% of PDI2EH-Br or 0.5 wt% of PDI2EH-CN, or 1 wt% P(NDI2OD-T2) solutions were prepared by dissolving the molecules in spectroscopic grade anhydrous chloroform in an inert atmosphere and spin-coating onto the electrodes at 750 r.p.m., 1,000 r.p.m. and 4,000 r.p.m., respectively, to achieve 100-nm-thick films.

Regarding the fabrication of field-effect transistor devices starting from electrode pairs bearing nanohole arrays, the two-terminal structures described above were completed by spin-coating of a commercial fluoropolymer (Cytop-809-M) at 2,500 r.p.m., leading to 700-nm-thick dielectric layers. The dielectric layer, processed from an orthogonal solvent which prevents the underlying PDI semiconductor films were prepared by spin-coating onto a conductive Au/glass substrate, which forms a film exceeding 10 nm and therefore sufficient to ensure that only the top material was probed. The metal Al and Ag films on glass substrates were prepared as previously described and measured over time to determine whether the exposure to air would affect their respective work functions. The exposure of all metal films employed as electrodes in the electrical characterization measurements is minimized, amounting to some minutes in total, including the transfer time from the evaporating chamber into the FIB chamber (ultrahigh vacuum at 10⁻⁶ mbar) and then into the glovebox (nitrogen). The ionization energy of P(NDI2OD-T2) amounted to 5.55 ± 0.05 eV, in perfect accordance with previous measurements. By considering an optical band gap of 1.55 eV, as reported in ref. 47, it was possible to estimate the energy of the LUMO level of P(NDI2OD-T2) as 4.00 ± 0.05 eV.

The ionization energy measured on Al and Ag films corresponds to their absolute work function. For Al films, the work function was found to be constant with time with a value of 4.2 ± 0.05 eV. For Ag films, the variation of the work function over a period of time up to four days was monitored. The work function changed from 4.72 eV (fresh sample) up to 4.81 eV (3 days 14 h) which, including the experimental error (normally about 50 meV), gives a net variation of approximately 50 meV. The Ag work function was found to be 4.75 eV after some minutes of exposure to air, which most probably corresponds to the value the Ag work function during the device experiments.

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