

All-Polymer Microcavities for the Fluorescence Radiative Rate Modification of a Diketopyrrolopyrrole Derivative

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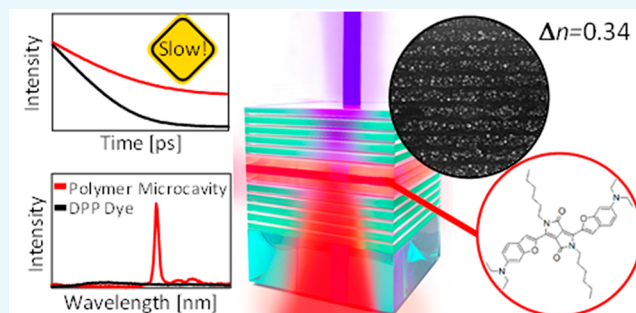
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Supporting Information

ABSTRACT: Controlling the radiative rate of emitters with macromolecular photonic structures promises flexible devices with enhanced performances that are easy to scale up. For instance, radiative rate enhancement empowers low-threshold lasers, while rate suppression affects recombination in photovoltaic and photochemical processes. However, claims of the Purcell effect with polymer structures are controversial, as the low dielectric contrast typical of suitable polymers is commonly not enough to provide the necessary confinement. Here we show all-polymer planar microcavities with photonic band gaps tuned to the photoluminescence of a diketopyrrolopyrrole derivative, which allows a change in the fluorescence lifetime. Radiative and

nonradiative rates were disentangled systematically by measuring the external quantum efficiencies and comparing the planar microcavities with a series of references designed to exclude any extrinsic effects. For the first time, this analysis shows unambiguously the dye radiative emission rate variations obtained with macromolecular dielectric mirrors. When different waveguides, chemical environments, and effective refractive index effects in the structure were accounted for, the change in the radiative lifetime was assigned to the Purcell effect. This was possible through the exploitation of photonic structures made of polyvinylcarbazole as a high-index material and the perfluorinated Aquivion as a low-index one, which produced the largest dielectric contrast ever obtained in planar polymer cavities. This characteristic induces the high confinement of the radiation electric field within the cavity layer, causing a record intensity enhancement and the steering the radiative rate. Current limits and requirements to achieve the full control of radiative rates with polymer planar microcavities are also addressed.



INTRODUCTION

Progress in the field of polymer photonics has quickly accelerated in the last decades due to their unique properties, including easy chemical tailoring, mechanical flexibility, and simple fabrication.¹ Devices employing all-polymer planar 1D photonic crystals such as light-emitting diodes (LEDs), photovoltaic cells, and sensors² are especially appealing as their fabrication is easy to scale up.³ However, the poor refractive index (n) contrast between transparent polymer materials limits light confinement³ and hinders the control of spontaneous emission rates, namely the Purcell effect.⁴ However, empowering the control of an emitter's radiative rate with polymer photonic crystals promises the easy large-area fabrication of either flexible low-threshold lasers⁵ and high-efficiency LEDs⁶ when the rate is increased or photo-

voltic and photocatalytic devices where an increase in the exciton lifetime (i.e., a reduction of the radiative rate) leads to a longer diffusion length and a higher device performance.⁷ Both radiative rate enhancement and radiative rate suppression have been well-demonstrated in metallic⁸ and inorganic⁹ structures. Indeed, the latter have been dominating

the photonics playground thanks to their low losses and optimal radiation confinement.¹⁰ Rate control has been achieved by employing inorganic optical resonators¹¹ and microcavities (MCs) of different typologies,¹² including planar ones,¹³ microdisks,¹⁴ micropillars,¹⁵ and photonic crystals.¹⁶ On the other hand, their fabrication requires severe conditions and is time and energy consuming. This aspect has hindered their adaptation for flexible devices, integration with organic and hybrid emitters, and large-area production. As such, achieving rate control with polymer structures would be a milestone for efficient solution-processable flexible photonics. Yet, the unambiguous observation of this effect in polymer structures has been disputed within the scientific community. Some claims of Purcell effect observation were made for hybrid

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61 silica/polystyrene systems,¹⁷ but possible extrinsic effects, such
 62 as changes in the refractive index of the effective medium,¹⁸
 63 and chemical effects, including exciton chemical traps,
 64 impurities, and local disorder affecting the structure, have
 65 been debated for these systems.¹⁹ In general, photolumines-
 66 cence (PL) lifetime (τ_{PL}) variations were reported for polymer
 67 synthetic opals²⁰ and 2D²¹ and planar MCs,²² but radiative
 68 rate variations were not investigated. Indeed, τ_{PL} measure-
 69 ments alone cannot disentangle radiative (Γ_{R}) and non-
 70 radiative (Γ_{NR}) decay rates without information on the
 71 photoluminescence external quantum efficiency (η , eq 1).²³
 72 Then, the conclusion that any changes in the fluorescence
 73 lifetime arise from a modification of the Γ_{R} value is valid only
 74 when η is taken into account.

$$\eta = \frac{\Gamma_{\text{R}}}{\Gamma_{\text{R}} + \Gamma_{\text{NR}}} \quad (1)$$

76 In this work, we unambiguously demonstrate radiative rate
 77 suppression within a planar polymer MC fabricated through
 78 the solution processing of a perfluorinated polymer, Aquivion
 79 (AQ, $n = 1.35$),² and polyvinylcarbazole (PVK, $n = 1.69$).^{3,24}
 80 The couple provides the highest dielectric contrast demon-
 81 strated for polymer planar microcavities ($\Delta n = 0.34$ in the UV-
 82 NIR range) so far.²⁵ The dye embedded in the cavity is a
 83 diketopyrrolopyrrole (DPP) derivative. Diketopyrrolopyrroles
 84 are some of the most studied organic dyes for electronics and
 85 photonics,²⁶ including in organic light-emitting diodes and
 86 solar cells,²⁷ due to their tailorable synthesis and high thermal-
 87 and photostabilities.²⁸ The same materials were also employed
 88 to fabricate a series of references. A systematic assessment of η
 89 and the emission decays allows us to unequivocally
 90 demonstrate an increase of the radiative lifetime (τ_{rad}) in
 91 opportunely tuned microcavities, ruling out any simpler
 92 alternative interpretations or extrinsic photophysical processes
 93 beside cavity Purcell effects.

94 ■ RESULTS

95 **Cavity Design and Properties.** The MCs were grown via
 96 spin-coating deposition and were formed by two dielectric
 97 mirrors, each of which consisted of 20 bilayers of AQ and PVK.
 98 The cavity between the mirrors contains two layers of cellulose
 99 acetate (CA) sandwiching a layer of polystyrene (PS) doped
 100 with a DPP derivative^{27,29} (DPP:PS), as sketched in Figure 1a.
 101 The same panel shows the chemical structure of the DPP dye,
 102 while the normalized PL and absorbance spectra of a thin film
 103 of the DPP:PS blend cast on a glass substrate are shown in
 104 Figure 1b. In the spectral range of interest, the DPP dye shows
 105 three distinct absorption maxima at $\lambda = 450, 600,$ and 655 nm
 106 (highest intensity). Upon excitation at 534 nm, the steady-state
 107 PL spectrum of the blend displays a Stokes shift of 22 nm as
 108 the maximum intensity appears at $\lambda = 677$ nm with a full-width
 109 at half-maximum (fwhm) of 40 nm. Additionally, the emission
 110 shows a broad shoulder at 745 nm.

111 The microcavity (MC_{tuned}) was engineered to tune the
 112 mirrors' photonic band gap (PBG) and cavity mode to the PL
 113 of the DPP dye using simple control of the spin-coating
 114 deposition parameters. Moreover, several reference samples
 115 were engineered and fabricated to compare the properties of
 116 the microcavity with those of the bare dye, particularly to
 117 exclude radiative rate variations due to extrinsic effects,
 118 including medium chemical effects, the polarity of the medium,
 119 the residual solvent diffusion among layers, waveguiding, out-

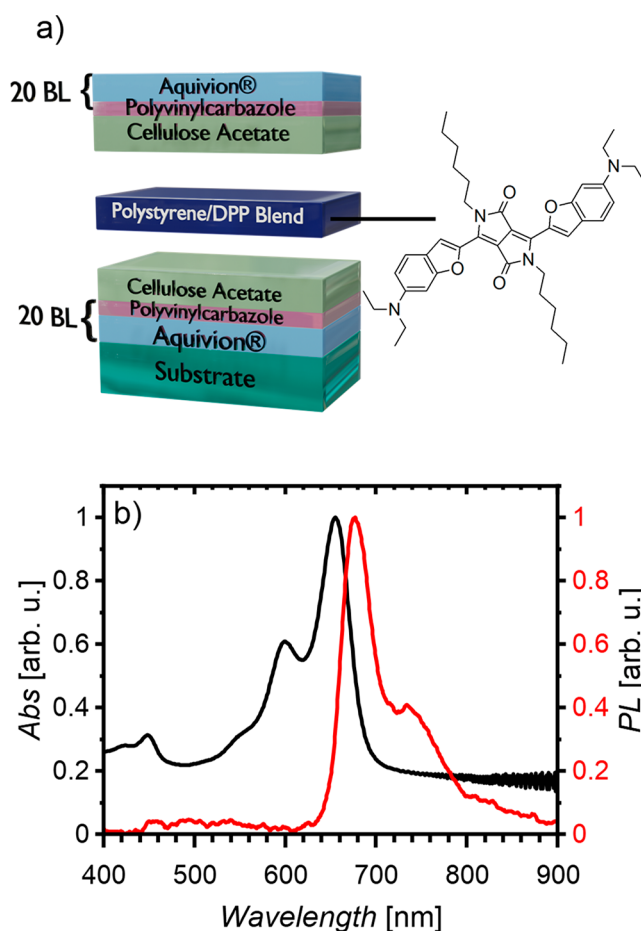


Figure 1. (a) Schematic of the MC structure, including the chemical structure of DPP. (b) Normalized absorbance and PL intensity spectra of the DPP:PS blend thin film.

coupling, and light extraction differences.²⁵ These references
 were a DPP:PS pristine blend film, a detuned microcavity
 (MC_{detuned}) with the PBG in the green region of the spectrum
 where the DPP:PS film does not show significant fluorescence,
 bilayer CA-DPP:PS (R1), and a more complex five-layer
 structure (R2) (see Supporting Figure S1 for the optical
 characterization of the thin films). These references serve to
 simulate possible defects that could be unintentionally inserted
 into the microcavity by the growth process as well as loss
 mechanisms such as waveguiding effects and self-absorption,
 which are known to affect light emission in polymer
 microcavities.²⁴

Figure 2a contrasts the reflectance spectra of both the tuned
 and detuned microcavities. Both structures display the
 characteristic features of planar MCs: an intense and wide
 reflectance band corresponding to the PBG of the photonic
 crystal mirror, with a sharp minimum assigned to the cavity
 mode, and a Fabry–Pérot interference pattern in the
 background. While the PBG and the cavity mode of MC_{tuned}
 strongly overlap the emission spectrum of the DPP blend,
 the shifted microcavity provides no spectral overlap; hence,
 any variation in the emission of the dye in this sample cannot
 be assigned to optical confinement effects provided by the
 microstructure.

For the further characterization of the MC_{tuned}, Figure 2b
 shows the transmittance spectrum of the microcavity, which,
 corresponding to the reflectance spectrum, shows a wide PBG

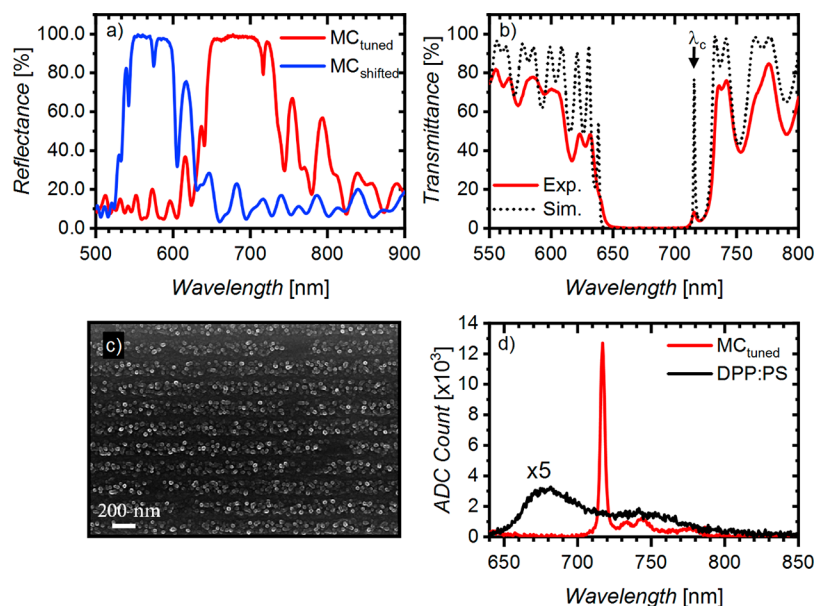


Figure 2. (a) Reflectance spectra of tuned microcavity (red line) and the detuned one (blue line). (b) Experimental (red full line) and simulated (black dotted line) transmittance of the tuned microcavity. (c) SEM image of a representative DBR forming the dielectric mirror in a tuned microcavity. (d) Photoluminescence spectra of the DPP:PS film (multiplied by 5, black full line) and the tuned microcavity (red full line).

147 with a maximum at $\lambda_c = 717$ nm assigned to the cavity mode.
 148 The PBG extends from 635 to 732 nm (fwhm of 250 meV), a
 149 larger value compared to other polymer planar microcavities
 150 with smaller dielectric contrasts.^{25,30} Moreover, due to the
 151 photonic band structure of the MC, the spectral position of the
 152 aforementioned features is strongly dependent on the angle of
 153 incidence and the polarization state of the incoming light beam
 154 (Supporting Figure S2),³¹ so the emission spectrum can
 155 possibly be tuned by changing the detection angle. It is worth
 156 noticing that the sample surface shows some minor spectral
 157 inhomogeneities due to small thickness variations across the
 158 sample, as reported in Supporting Figure S3a. Supporting
 159 Figure S3b shows a digital image of the strong PL from a
 160 microcavity cast on a flexible substrate under violet laser
 161 excitation.

162 The optical response of the structure was simulated using
 163 the complex refractive index dispersions of all the polymers
 164 employed that were previously reported in the literature^{2,3,24,32}
 165 and that measured for the DPP:PS film (Supporting Figure
 166 S4). Employing transfer matrix method (TMM) modeling,³
 167 the simulated transmittance reported in Figure 2a (dotted line)
 168 was best-fit to the experimental one, yielding the thicknesses of
 169 individual materials (74.5 nm for PVK, 159.5 nm for AQ, 184
 170 nm for DPP:PS, and 78 nm for the CA layers). The positions and
 171 widths of the interference fringes, as well as the PBG and λ_c in
 172 the experimental measurements, are all well-reproduced in the
 173 calculation. To obtain initial estimates of the layer thicknesses
 174 for simulations, SEM measurements were performed. The
 175 SEM micrograph in Figure 2c (and Supporting Figure S5)
 176 shows the layering of the DBR after the microcavity was freeze-
 177 cracked, which caused uneven fracture artifacts in the image.
 178 The AQ layers are distinguishable thanks to their spherical
 179 aggregates, which have been previously reported in the
 180 literature for perfluorosulfonic acid ionomer dispersions.³³
 181 From the SEM images, the uniformity is clear. As described in
 182 detail in Supporting Figure S6, the average thickness of the AQ
 183 layers was estimated to be 108 ± 8 nm, and that of PVK was
 184 estimated to be 65 ± 6 nm. We noticed that the freeze-

185 cracking process induced delamination that prevented the
 186 direct observation of the cavity layers, whose thicknesses were
 187 then estimated through AFM measurements to be 88 ± 33 nm
 188 for the CA layers and 113 ± 67 for the DPP:PS layer (see
 189 Supporting Figure S7 for details). We observed good
 190 agreement within the experimental uncertainty between
 191 thickness derived from SEM/AFM for the PVK, CA, and
 192 DPP:PS layers and those derived from optical simulations,
 193 while some discrepancy was observed for the AQ layers.
 194 However, this is expected as AQ, being a material sensitive to
 195 its chemical environment, is strongly perturbed by the abrupt
 196 change induced by the freeze-cracking process and the
 197 subsequent vacuum environment of the SEM chamber.²

198 Figure 2d compares the PL spectra for a DPP:PS thin film
 199 and MC_{tuned}. Both the DPP:PS layer in the microcavity and
 200 that in the reference DPP:PS film were cast under identical
 201 conditions. Due to the spectral overlap of the dye PL with the
 202 PBG and the very sharp cavity mode (fwhm of ~ 3.5 nm), the
 203 local photonic density of states (LPDOS) strongly modulates
 204 the fluorescence line-shape.^{3,34} Indeed, the LPDOS is very low
 205 at the PBG and is strongly enhanced at the cavity mode,
 206 channeling the emission into the latter. At near-normal
 207 incidence, the enhanced LPDOS at the cavity mode produces
 208 a 40-fold intensity enhancement of the DPP emission at $\lambda_c =$
 209 717 nm as compared to the bare emitter film (see Supporting
 210 Figure S8 for a comparison with other spectral regions), a
 211 record value among polymer microcavities (the highest being
 212 $8.9\times$ to the best of our knowledge).^{22b} As the collection angle
 213 increases, the PBG and λ_c shift to shorter wavelengths,
 214 changing the spectral regions that are either enhanced or
 215 suppressed (see Supporting Figures S9 and S10 for details).
 216 The finesse of MC_{tuned} is quantified through the quality factor
 217 ($Q = \lambda_c / \Delta\lambda_c$) of approximately 205, a relatively high value
 218 compared to the highest reported so far for all-polymer
 219 microcavities ($Q = 255$).³⁰ We also notice three additional
 220 weak emission peaks at approximately 733, 743, and 775 nm in
 221 the cavity PL spectra that corresponded to the local minima in
 222 the Fabry–Pérot interference pattern (Figure 2a). Unsurpris-

ingly, the PL is strongly suppressed at the PBG wavelengths where the LPDOS is lower. As expected from the angular dispersion of transmittance in the microcavities³⁴ (Supporting Figure S2), the fluorescence from MC_{tuned} is highly directional compared to the Lambertian emission of the DPP:PS film (Supporting Figure S9).

Moreover, we consider the overall microcavity effect on the fluorescence intensity, where only part of the dye's fluorescence spectrum overlaps the PBG and the microcavity mode. Supporting Figure S10 displays the angle-resolved PL intensity data at the cavity mode wavelength as well as those integrated for all wavelengths for the microcavity, the standalone DPP:PS film, and the reference R2. Then the total emission enhancement factor ($G_{\text{tot}}^{\text{exp}}$) according to eq 2³⁵ can be calculated as ~ 1.4 with respect to the dye and ~ 1.3 with respect to R2, indicating an overall enhancement in the PL intensity. The relevant calculations are more thoroughly discussed in the Supporting Information.

$$G_{\text{tot}}^{\text{exp}} = \frac{\iint \text{PL}_{\text{cav}}(\lambda, \theta) d\lambda d\theta}{\iint \text{PL}_{\text{ref}}(\lambda, \theta) d\lambda d\theta} \quad (2)$$

Cavity Effects and Radiative Rate Modification. Figure 3 compares the fluorescence decay upon excitation with a

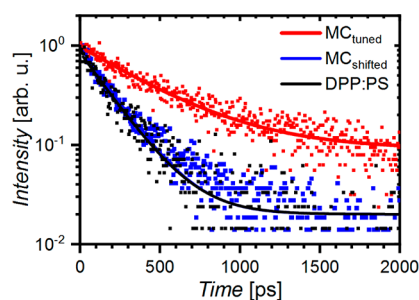


Figure 3. Photoluminescence decay (squares) and fitted data (lines) for the tuned microcavity (red) and for two references: the DPP:PS film (black) and a detuned microcavity (blue, MC_{detuned}) around λ_c .

pulsed laser ($\lambda = 405$ nm) for the standalone DPP:PS film (black squares), MC_{detuned} (blue squares), and MC_{tuned} (red squares), which was evaluated from the PL intensity at $\lambda_{\text{MCtuned}} \pm 10$ nm. As the reference intensities are lower than the tuned microcavity due to the previously discussed enhancement effect (see Figure 2c), their decay signals have a lower signal-to-noise ratio. Notwithstanding, a strong difference between the lifetimes of MC_{tuned} and the two references can clearly be observed, even without the need for fitting.

τ_{PL} , which is related to the radiative and nonradiative decay rates ($\frac{1}{\tau_{\text{PL}}} = \Gamma_{\text{R}} + \Gamma_{\text{NR}} = \frac{1}{\tau_{\text{R}}} + \frac{1}{\tau_{\text{NR}}}$), was retrieved from the single-exponential fitting of the decays (Figure 3, Supporting Figure S11, and Table S1). Then, the radiative lifetimes for all

the samples (τ_{rad}) were calculated as the ratio between the best-fitted τ_{PL} and the quantum efficiency η measured for all samples (Table S1). We would like to stress the role of the external quantum efficiency (eq 1) in disentangling the radiative and nonradiative (τ_{NR}) lifetimes and quantifying the Purcell effect. As reported in eq 3, the radiative rate can only be calculated from the quantum efficiency and the overall PL lifetime. Unfortunately, as η measurements are rarely reported, making estimations of the actual radiative rate and its variations is highly speculative.³⁶

$$\tau_{\text{R}} = \frac{\tau_{\text{PL}}}{\eta} \quad (3)$$

The fits for the references are almost superimposable, as seen in Figure 3. The results of such calculations are summarized in Table 1, showing that the two references have similar τ_{PL} values (190 ps for the DPP:PS layer and 185 ps for MC_{detuned}). Comparable lifetimes were also observed for other references that emulated the boundaries of the defect layer (as reported in Table S1 of the Supporting Information), while the value more than doubles (~ 416 ps) for the tuned microcavity. Regarding η instead, the value for the bare DPP:PS film is $\sim 6\%$, which decreases by half for the MC_{detuned} reference ($\sim 3\%$) and to a sixth for MC_{tuned} ($\sim 1\%$). Astonishingly, with respect to the DPP:PS film ($\tau_{\text{rad}} = 3.2$ ns), the microcavity shows a 10-fold increase in the radiative lifetime ($\tau_{\text{rad}} \sim 42$ ns) as compared to a slight increase of that for MC_{detuned} ($\tau_{\text{rad}} = 6.2$ ns). Corresponding changes in the radiative rates were derived ($\Gamma_{\text{rad}} = \frac{1}{\tau_{\text{rad}}}$) and are reported in

Table 1.

From data in Table 1, we derived the ratio between the radiative decay rate of the emitter modified by the environment ($\Gamma_{\text{rad}}^{\text{mod}}$) and its decay in vacuum $\Gamma_{\text{rad}}(P)$, usually called the Purcell factor. In our case, the error $0.03 \leq P \leq 0.24$ (for the average values, $P = 0.08$) is included when considering the DPP:PS film as a reference and the error $0.06 \leq P \leq 0.47$ ($P = 0.15$) is included when considering the detuned microcavity as a reference. The latter accounts for enhanced self-absorption due to the small Stokes shift, the longer photon dwell time in the cavity, out-coupling effects, and any variations due to chemical effects or the effective dielectric environment. Notwithstanding the significant error propagating from uncertainty in the quantum yield measurements, the present P values are consistent with strong radiative rate suppression. If, however, the overall PL rates were considered only, i.e., neglecting η as often occurs,²³ P would have much larger values (0.44–0.46), falsely indicating a smaller and speculative rate suppression. The PL decay and η for all other references used (Table S1) show very similar values to those for references reported in Table 1. Finally, we notice that the effective refractive indices of all our references (also reported in) show a variation below 10%.

Table 1. Radiative Decays: Photoluminescence Lifetime τ_{PL} , Quantum Efficiency η , and the Calculated Radiative τ_{rad} and Non-Radiative τ_{NR} Lifetimes for the Microcavities and the DPP:PS Film^a

sample	τ_{PL} [ps]	η [%]	τ_{rad} [ns]	τ_{NR} [ps]	Γ_{rad} [ns ⁻¹]	Γ_{NR} [ns ⁻¹]
DPP:PS	190 ± 20	6 ± 2	3.2 ± 1.4	202 ± 89	0.32	4.9
MC _{detuned}	185 ± 20	3 ± 1	6.2 ± 2.7	191 ± 84	0.16	5.2
MC _{tuned}	416 ± 20	1 ± 0.5	41.6 ± 22.8	420 ± 230	0.02	2.4

^aSee Table S1 for more details.

307 ■ DISCUSSION

308 To date, this is the first case where the Purcell effect is
 309 unambiguously confirmed in all-polymer planar microcavities.
 310 It is then useful to provide a deeper investigation of the
 311 phenomena behind it and explain the reasons that prevented
 312 its observation so far.

313 The puzzling results summarized in Table 1 concern the
 314 simultaneous reduction in η and the increase in τ_{PL} for the
 315 tuned microcavity with respect to all references in both Table 1
 316 and Table S1. Indeed, in contrast with our data (Table 1)
 317 where Γ_{NR} is almost halved, a decrease in η usually causes a
 318 decrease in τ_{PL} due to enhanced nonradiative recombination
 319 rates.³⁷ Moreover, the radiative rate is known to be strongly
 320 dependent on the effective refractive index surrounding the
 321 emitters, as demonstrated in planar silicon slot waveguides.^{19d}
 322 The variations in the value of τ_{rad} between the tuned cavity and
 323 the references show remarkable dissimilarity and do not
 324 comply with this interpretation because the references have
 325 effective refractive indices similar (within 10%, see Table S1)
 326 to that of MC_{tuned} , making this alternative explanation
 327 unsuitable for the radiative rate change observed for our
 328 plastic microcavities.

329 To explain the significant extension of the radiative rate for
 330 the MC_{tuned} , we must invoke an unusual change in the light–
 331 matter interaction: the Purcell-effect, i.e., the modification of
 332 the spontaneous emission rate of a quantum system. According
 333 to theory, both radiative rate suppression and radiative rate
 334 enhancement should be observable in planar microcavities.^{13,38}

335 Ideally, rate enhancement is achieved when three conditions
 336 are satisfied: (i) the electromagnetic field is strongly confined,
 337 (ii) the emission intensity is spectrally sharper than the cavity
 338 mode and tuned to the region where the LPDOS is at a
 339 maximum, and (iii) the emitter is placed at an antinode of the
 340 microcavity electric field standing wave where its intensity is at
 341 a maximum. Conversely, if one or more of these conditions is
 342 not satisfied, rate suppression should occur.³⁹

343 To discuss the role of these three requirements, it is useful to
 344 recall the relevant theoretical framework. According to the
 345 Wigner–Weisskopf approximation,⁴⁰ the modified emission
 346 rate is directly proportional to the LPDOS,^{31,41} which is a
 347 function of the angular frequency (ω) and of the emitter
 348 position (\mathbf{r}) in the microcavity as described by a modification
 349 of Fermi's golden rule ($\Gamma_{\text{rad}}^{\text{mod}} = \frac{2\pi}{\hbar^2} \text{LPDOS}(\omega, \mathbf{r})$).³⁴ Then,
 350 when LPDOS is at a minimum at the band gap, the radiative
 351 rate is suppressed, while at the cavity mode (where it is
 352 maximum) the radiative rate can be enhanced. Furthermore, if
 353 the emitter is placed at an antinode of the electric field
 354 amplitude inside the cavity, the emission is enhanced, and vice
 355 versa for the positioning at a node. As DPP is a broad-
 356 spectrum emitter, all off-resonance photons are expected to
 357 experience suppression due to the low density of states outside
 358 the cavity mode. This again is assigned to the enhanced
 359 dielectric contrast, which is comparable to that achieved in
 360 some inorganic dielectric microcavities that exhibited a PL rate
 361 enhancement.³⁸

362 Thus, far, the achieved result is promising for applications
 363 where the suppression of radiative rate and the enhancement
 364 of the lifetime are desired effects, such as light-harvesting
 365 devices, where the efficiency of the devices is limited by the
 366 diffusion length of the excitons and thus by the radiative
 367 lifetime.⁷ On the other hand, radiative rate enhancement is

desired for all light-emission applications, most importantly
 low-threshold lasers.

Theoretically, the maximum achievable Purcell factor in a
 cavity (P_{max} , eq 4)¹⁰ is dictated by the quality factor Q and the
 effective cavity volume (V_{eff}), which represents the electro-
 magnetic field confinement in all the directions.

$$P_{\text{max}} = \frac{3}{4\pi^2} \left(\frac{\lambda}{n} \right)^3 \frac{Q}{V_{\text{eff}}} \geq \frac{\Gamma_{\text{rad}}^{\text{mod}}}{\Gamma_{\text{rad}}} \quad (4)$$

In our case, there is no lateral confinement in the plane of
 the cavity layer. However, we can estimate the relative
 confinement along the periodicity direction from the
 penetration depth (L_{eff}) of the electromagnetic field into the
 dielectric mirrors (eq 5). The penetration depth depends on
 the dielectric contrast ($\Delta n = n_{\text{H}} - n_{\text{L}}$), the geometric length of
 the defect layers (L_{MC}), the optical lengths within the dielectric
 mirrors (L_{DBR}), the periodicity of the structure (D), and the
 effective refractive index (n_{eff}).²⁵

$$L_{\text{eff}} = L_{\text{MC}} + 2L_{\text{DBR}} = L_{\text{MC}} + 4D \frac{n_{\text{eff}}}{|n_{\text{H}} - n_{\text{L}}|} \quad (5)$$

In the MC_{tuned} case, L_{eff} is approximately 4.6 μm , which is
 much smaller than the overall geometrical length of 9.7 μm .
 Then, the system shows a stronger confinement with respect to
 the previously investigated all-polymer microcavities, where the
 lower refractive index contrast causes L_{eff} to be comparable or
 even longer than the photonic structure itself (Table S2). This
 simple characteristic should explain why radiative rate
 variations have not been confirmed so far. In fact, $\Delta n = 0.34$
 represents at least a 50% increase from the highest value
 reported in the literature for polymer microcavities so far.^{22b,42}
 On the other hand, most of the emission of broad emitters is
 suppressed by the PBG or is leaked outside the cavity mode,
 hence indicating the need to integrate very narrow emitters (PL
 full-width half-maximum below the cavity mode width) in the
 microcavities.

Theoretical and experimental observations indicate that the
 maximum enhancement or suppression for a narrow emitter
 placed at the antinode of the standing wave in a planar
 dielectric microcavity is around 30%.⁴³ Hybrid planar systems
 using polymer emitters or spacers in dielectric and metallic
 systems a report similar enhancement in the overall decay rate,
 including nonradiative decay.^{36,44} However, if micropillars or
 microbeams with micrometer-scale diameters were fabricated
 from the planar microcavity, the Purcell factor could reach
 10.⁴⁵ At the state of the art, lateral nanostructuring remains the
 most reliable approach to radiative rate enhancement.¹⁰

From the data reported in Table 1, we noticed that a change
 in the nonradiative rate was also observed. We suggest here a
 possible explanation for the effect, even though this is not the
 main focus of this work. Nonradiative decays are known to be
 activated after photoexcitation by the photon field.⁴⁶ The
 microcavity deeply changes the environment of the dye and
 induces the strong localization of the electrical field within the
 layers driven by the dielectric contrast. In Figure S12, we
 report the square modulus of the absolute electric field
 amplitude ($|E|^2$) and its maximum value in the cavity as
 calculated using TMM;⁴⁷ the DPP:PS layer position is
 highlighted in red. As per the calculations, a strong resonance
 enhancement of the electric field intensity exceeding 40 \times the
 unmodulated intensity can be theoretically expected at a
 frequency of the photon field resonant with molecular

426 electronic transitions. According to Siebrand,⁴⁶ this enhanced
427 field modifies the Hamiltonian describing the process and thus
428 is a likely explanation for the reduction of the nonradiative rate
429 observed. In agreement with this conclusion, we noticed that
430 no changes of the nonradiative rate were observed for the
431 detuned microcavity, where field enhancement still occurs but
432 is at a frequency nonresonant with the molecular fluorescence.

433 ■ CONCLUSION

434 In conclusion, we demonstrated radiative rate suppression with
435 a 10-fold radiative lifetime increase and record PL enhance-
436 ment in fully solution-processed polymer planar microcavities,
437 which could open up new perspectives for flexible devices. The
438 larger dielectric contrast employed for the microcavity growth
439 allowed the unambiguous radiative emission rate variation to
440 be observed for the first time thanks to better confinement
441 within the sample thickness. Nevertheless, smaller mode
442 volume as well as the spatial and spectral optimization of
443 fluorophores are still necessary to achieve radiative rate
444 enhancement. The careful synergy of polymer refractive
445 index engineering, advanced dye synthesis, and solution-
446 fabricated flexible structures provides novel perspectives to
447 polymer photonics.

448 ■ METHODS

449 **Dye Synthesis.** DPP was synthesized following the
450 literature procedure.^{29a} The dye shows a fluorescence quantum
451 efficiency of 0.55 in toluene solutions (the solvent used to
452 prepare the cavity layer) and essentially the same values in
453 tetrahydrofuran and dichloromethane.

454 **Microcavity Fabrication.** All samples were grown via the
455 alternating spin coating of 100 μL of the polymer solutions on
456 $25 \times 25 \text{ mm}^2$ glass substrates at 175 rps for the dielectric
457 mirrors and 75 rps for the PS-dye solution. The mirrors with
458 20 bilayers were cast by alternating the deposition of the
459 Aquivion D79-25BS water/ethanol dispersion and the PVK
460 solution in toluene (40 mg/mL). The DPP:PS layer was
461 sandwiched between two layers of CA (30 mg/mL in
462 diacetone alcohol). The top layer serves to prevent water
463 percolation during the subsequent Aquivion deposition, and
464 the bottom one instead serves to maintain the symmetry of the
465 structure. The emitter layer was obtained by casting a solution
466 of the dye in PS/toluene (1 mg/mL DPP and 30 mg/mL PS).

467 **Optical Characterization.** Transmittance measurements
468 were performed with a setup consisting of deuterium and
469 tungsten-halogen sources (spectral range of 230–2500 nm)
470 using an AvaSpec-ULS4096CL-EVO CMOS (spectral range of
471 200–1100 nm and resolution of 1.4 nm) spectrometer. Angle-
472 resolved spectra were recorded using a homemade setup with
473 an angular resolution $\leq 1^\circ$. Steady-state PL measurements were
474 performed by exciting the samples with an Oxxius 405 nm CW
475 laser focused on a 1 mm^2 spot. The fluorescence spectra were
476 collected with the same spectrometer. The collection setup
477 allowed the the transmittance and the PL to be measured on
478 the same sample spot.

479 **SEM Measurements.** SEM measurements were performed
480 using the FE-SEM Zeiss SUPRA 40 VP instrument (Carl Zeiss,
481 Oberkochen, Germany) at an acceleration voltage of 10 kV.
482 The microcavity sample was frozen in liquid nitrogen and
483 fractured to reveal the cross section upon which a thin carbon
484 layer was deposited using a high-vacuum evaporator (Polaron
485 6700).

Time-Resolved PL Measurements. TRPL measurements 486
were carried out using a femtosecond tunable Ti:sapphire laser 487
(Coherent Chameleon Ultra II) and a streak camera detection 488
system. Type I phase-matched second harmonic generation 489
was performed using a β -barium borate crystal, leading to 490
pulses with central wavelengths of 405 nm and spot diameters 491
of 6–8 μm at the sample. The emission was collected at 30° 492
from normal incidence and analyzed by a spectrograph 493
(Princeton Instruments Acton SP2300) coupled to a streak 494
camera (Hamamatsu C5680), resulting in a spectral resolution 495
around 1 nm and a temporal resolution of 20 ps. 496

Quantum Efficiency. External PL quantum efficiencies for 497
microcavities and references were measured by the widely 498
utilized method from de Mello et al.⁴⁸ using an integrating 499
sphere (Avantes AvaSphere-50) fiber-coupled with a 405 nm 500
LDH-P-C-405 laser and an Avantes AvaSpec-2048 calibrated 501
spectrometer (200–1150 nm resolution). Typical uncertainty 502
in the quantum efficiency measurements for low values 503
($<10\%$) can be in the range of 30–50%.⁴⁹ 504

Refractive Index Measurements. A VASE instrument (J. 505
A. Woollam Co., Lincoln, NE) in the range 250–2500 nm was 506
used for spectroscopic ellipsometry measurements at different 507
incidence angles from 55° to 75° on films on both fused silica 508
and silicon substrates. Varian Cary 6000i spectrometer in the 509
spectral range of 200–1800 nm was used to measure the 510
reflectance and transmittance at normal incidence. Then, the 511
complex refractive index was evaluated using WVASE32 512
software (J. A. Woollam, ver. 3.774, Lincoln, NE), adopting 513
oscillator models to guarantee a Kramers–Kronig consistency. 514

515 ■ ASSOCIATED CONTENT

516 ■ Supporting Information

The Supporting Information is available free of charge at 517
<https://pubs.acs.org/doi/10.1021/acsomega.2c00167>. 518

Further optical characterization of reference samples, 519
complex refractive index of the polymer/dye blend, 520
microscopic characterization of the samples, detailed PL 521
behavior of the tuned microcavity and references, further 522
time-resolved PL decay analysis, comparison among 523
microcavities reported in the literature, and the 524
calculated electric field within the tuned microcavity 525
(PDF) 526

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559 Author Contributions

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577 ■ REFERENCES

578 (1) Annadhasan, M.; Basak, S.; Chandrasekhar, N.; Chandrasekar, R.
579 Next-Generation Organic Photonics: The Emergence of Flexible
580 Crystal Optical Waveguides. *Adv. Opt. Mater.* **2020**, *8* (21), 2000959.
581 (2) Megahd, H.; Oldani, C.; Radice, S.; Lanfranchi, A.; Patrini, M.;
582 Lova, P.; Comoretto, D. Aquivion–Poly(N-vinylcarbazole) Holistic
583 Flory–Huggins Photonic Vapor Sensors. *Adv. Opt. Mater.* **2021**, *9*
584 (5), 2170017.
585 (3) Lova, P.; Manfredi, G.; Comoretto, D. Advances in functional
586 solution processed planar one-dimensional photonic crystals. *Adv.*
587 *Opt. Mater.* **2018**, *6* (24), 1800730–26.
588 (4) Purcell, E. M.; Torrey, H. C.; Pound, R. V. Resonance
589 Absorption by Nuclear Magnetic Moments in a Solid. *Phys. Rev.* **1946**,
590 *69* (1–2), 37–38.
591 (5) Prieto, I.; Llorens, J. M.; Muñoz-Camúñez, L. E.; Taboada, A.
592 G.; Canet-Ferrer, J.; Ripalda, J. M.; Robles, C.; Muñoz-Matutano, G.;
593 Martínez-Pastor, J. P.; Postigo, P. A. Near thresholdless laser
594 operation at room temperature. *Optica* **2015**, *2* (1), 66–69.
595 (6) (a) Shambat, G.; Ellis, B.; Majumdar, A.; Petykiewicz, J.; Mayer,
596 M. A.; Sarmiento, T.; Harris, J.; Haller, E. E.; Vučković, J. Ultrafast
597 direct modulation of a single-mode photonic crystal nanocavity light-
598 emitting diode. *Nat. Commun.* **2011**, *2*, 539. (b) Cho, H.; Chung, J.;
599 Song, J.; Lee, J.; Lee, H.; Lee, J.; Moon, J.; Yoo, S.; Cho, N. S.
600 Importance of Purcell factor for optimizing structure of organic light-
601 emitting diodes. *Opt. Express* **2019**, *27* (8), 11057–11068.
602 (7) Vuong, L. T.; Kozyreff, G.; Betancur, R.; Martorell, J. Cavity-
603 controlled radiative recombination of excitons in thin-film solar cells.
604 *Appl. Phys. Lett.* **2009**, *95* (23), 233106.

(8) Murataj, I.; Channab, M.; Cara, E.; Pirri, C. F.; Boarino, L.; 605
Angelini, A.; Ferrarese Lupi, F. Hyperbolic Metamaterials via 606
Hierarchical Block Copolymer Nanostructures. *Adv. Opt. Mater.* 607
2021, *9* (7), 2001933. 608
(9) Gevaux, D. G.; Bennett, A. J.; Stevenson, R. M.; Shields, A. J.; 609
Atkinson, P.; Griffiths, J.; Anderson, D.; Jones, G. A. C.; Ritchie, D. A. 610
Enhancement and suppression of spontaneous emission by temper- 611
ature tuning InAs quantum dots to photonic crystal cavities. *Appl.* 612
Phys. Lett. **2006**, *88* (13), 131101. 613
(10) Vahala, K. J. Optical microcavities. *Nature* **2003**, *424* (6950), 614
839–846. 615
(11) Jacob, Z.; Smolyaninov, I. I.; Narimanov, E. E. Broadband 616
Purcell effect: Radiative decay engineering with metamaterials. *Appl.* 617
Phys. Lett. **2012**, *100* (18), 181105. 618
(12) Megahd, H.; Comoretto, D.; Lova, P. Planar microcavities: 619
Materials and processing for light control. *Optical Materials: X* **2022**, 620
13, 100130. 621
(13) Vredenberg, A. M.; Hunt, N. E. J.; Schubert, E. F.; Jacobson, D. 622
C.; Poate, J. M.; Zydzik, G. J. Controlled atomic spontaneous 623
emission from Er³⁺ in a transparent Si/SiO₂ microcavity. *Phys. Rev.* 624
Lett. **1993**, *71* (4), 517–520. 625
(14) Xie, Z. G.; Götzinger, S.; Fang, W.; Cao, H.; Solomon, G. S. 626
Influence of a Single Quantum Dot State on the Characteristics of a 627
Microdisk Laser. *Phys. Rev. Lett.* **2007**, *98* (11), 117401. 628
(15) Ding, X.; He, Y.; Duan, Z. C.; Gregersen, N.; Chen, M. C.; 629
Unsleber, S.; Maier, S.; Schneider, C.; Kamp, M.; Höfling, S.; Lu, C.- 630
Y.; Pan, J.-W. On-Demand Single Photons with High Extraction 631
Efficiency and Near-Unity Indistinguishability from a Resonantly 632
Driven Quantum Dot in a Micropillar. *Phys. Rev. Lett.* **2016**, *116* (2), 633
020401. 634
(16) Noda, S.; Fujita, M.; Asano, T. Spontaneous-emission control 635
by photonic crystals and nanocavities. *Nat. Photonics* **2007**, *1* (8), 636
449–458. 637
(17) (a) Rout, D.; Kumar, G.; Vijaya, R. Amplified emission and 638
modified spectral features in an opal hetero-structure mediated by 639
passive defect mode localization. *J. Phys. D: Appl. Phys.* **2018**, *51* (1), 640
015112. (b) Petrov, E. P.; Bogomolov, V. N.; Kalosha, I. I.; 641
Gaponenko, S. V. Spontaneous Emission of Organic Molecules 642
Embedded in a Photonic Crystal. *Phys. Rev. Lett.* **1998**, *81* (1), 77–80. 643
(18) Zhu, Y.; Xu, W.; Zhang, H.; Wang, W.; Tong, L.; Xu, S.; Sun, 644
Z.; Song, H. Highly modified spontaneous emissions in YVO₄:Eu³⁺ 645
inverse opal and refractive index sensing application. *Appl. Phys. Lett.* 646
2012, *100* (8), 081104. 647
(19) (a) Megens, M.; Schriemer, H. P.; Lagendijk, A.; Vos, W. L. 648
Comment on “Spontaneous Emission of Organic Molecules 649
Embedded in a Photonic Crystal. *Phys. Rev. Lett.* **1999**, *83* (25), 650
5401–5401. (b) Wang, W.; Song, H.; Bai, X.; Liu, Q.; Zhu, Y. 651
Modified spontaneous emissions of europium complex in weak 652
PMMA opals. *Phys. Chem. Chem. Phys.* **2011**, *13* (40), 18023–18030. 653
(c) Petrov, E. P.; Bogomolov, V. N.; Kalosha, I. I.; Gaponenko, S. V.; 654
et al. Petrov et al. Reply: *Reply. Phys. Rev. Lett.* **1999**, *83* (25), 5402. 655
(d) Creatore, C.; Andreani, L. C.; Miritello, M.; Lo Savio, R.; Priolo, 656
F. Modification of erbium radiative lifetime in planar silicon slot 657
waveguides. *Appl. Phys. Lett.* **2009**, *94* (10), 103112. 658
(20) Priya; Schöps, O.; Woggon, U.; Nair, R. V. Inhibited 659
spontaneous emission using gaplike resonance in disordered photonic 660
structures. *Phys. Rev. A* **2018**, *98* (4), 043835. 661
(21) Gan, X.; Clevenson, H.; Tsai, C.-C.; Li, L.; Englund, D. 662
Nanophotonic filters and integrated networks in flexible 2D polymer 663
photonic crystals. *Sci. Rep.* **2013**, *3*, 2145. 664
(22) (a) Lova, P.; Olivieri, M.; Surace, A.; Topcu, G.; Emirdag- 665
Eanes, M.; Demir, M. M.; Comoretto, D. Polymeric Planar 666
Microcavities Doped with a Europium Complex. *Crystals* **2020**, *10* 667
(4), 287. (b) Athanasiou, M.; Papagiorgis, P.; Manoli, A.; Bernasconi, 668
C.; Bodnarchuk, M. I.; Kovalenko, M. V.; Itskos, G. Efficient 669
Amplified Spontaneous Emission from Solution-Processed CsPbBr₃ 670
Nanocrystal Microcavities under Continuous Wave Excitation. *ACS* 671
Photonics **2021**, *8* (7), 2120–2129. 672

- 673 (23) Pelton, M. Modified spontaneous emission in nanophotonic
674 structures. *Nat. Photonics* **2015**, *9* (7), 427–435.
- 675 (24) Lova, P.; Grande, V.; Manfredi, G.; Patrini, M.; Herbst, S.;
676 Würthner, F.; Comoretto, D. All-polymer photonic microcavities
677 doped with perylene bisimide j-aggregates. *Adv. Opt. Mater.* **2017**, *5*
678 (21), 1700523.
- 679 (25) Lova, P.; Megahd, H.; Stagnaro, P.; Alloisio, M.; Patrini, M.;
680 Comoretto, D. Strategies for dielectric contrast enhancement in 1D
681 planar polymeric photonic crystals. *Appl. Sci.* **2020**, *10* (12), 4122.
- 682 (26) Skonieczny, K.; Papadopoulos, I.; Thiel, D.; Gutkowski, K.;
683 Haines, P.; McCosker, P. M.; Laurent, A. D.; Keller, P. A.; Clark, T.;
684 Jacquemin, D.; Guldi, D. M.; Gryko, D. T. How To Make
685 Nitroaromatic Compounds Glow: Next-Generation Large X-Shaped,
686 Centrosymmetric Diketopyrrolopyrroles. *Angew. Chem., Int. Ed.* **2020**,
687 *59* (37), 16104–16113.
- 688 (27) Grzybowski, M.; Gryko, D. T. Diketopyrrolopyrroles: Syn-
689 thesis, Reactivity, and Optical Properties. *Adv. Opt. Mater.* **2015**, *3*
690 (3), 280–320.
- 691 (28) Bao, W. W.; Li, R.; Dai, Z. C.; Tang, J.; Shi, X.; Geng, J. T.;
692 Deng, Z. F.; Hua, J. Diketopyrrolopyrrole (DPP)-Based Materials and
693 Its Applications: A Review. *Front. Chem.* **2020**, *8*, 679.
- 694 (29) (a) Purc, A.; Koszarna, B.; Iachina, I.; Friese, D. H.; Tasiar, M.;
695 Sobczyk, K.; Pędziński, T.; Brewer, J.; Gryko, D. T. The impact of
696 interplay between electronic and steric effects on the synthesis and the
697 linear and non-linear optical properties of diketopyrrolopyrrole
698 bearing benzofuran moieties. *Organic Chemistry Frontiers* **2017**, *4*
699 (5), 724–736. (b) Hupfer, M. L.; Koszarna, B.; Ghosh, S.; Gryko, D.
700 T.; Presselt, M. Langmuir–Blodgett Films of Diketopyrrolopyrroles
701 with Tunable Amphiphilicity. *Langmuir* **2021**, *37* (34), 10272–10278.
- 702 (30) Manfredi, G.; Lova, P.; Di Stasio, F.; Krahne, R.; Comoretto, D.
703 Directional Fluorescence Spectral Narrowing in All-Polymer Micro-
704 cavities Doped with CdSe/CdS Dot-in-rod Nanocrystals. *ACS*
705 *Photonics* **2017**, *4* (7), 1761–1769.
- 706 (31) *Organic and Hybrid Photonic Crystals*; Comoretto, D., Ed.;
707 Springer International Publishing: Basel, Switzerland, 2015.
- 708 (32) Sultanova, N.; Kasarova, S.; Nikolov, I. Dispersion Properties of
709 Optical Polymers. *Acta Phys. Polym., A* **2009**, *116*, 585.
- 710 (33) da Silva, J. S.; Carvalho, S. G. M.; da Silva, R. P.; Tavares, A. C.;
711 Schade, U.; Puskar, L.; Fonseca, F. C.; Matos, B. R. SAXS signature of
712 the lamellar ordering of ionic domains of perfluorinated sulfonic-acid
713 ionomers by electric and magnetic field-assisted casting. *Phys. Chem.*
714 *Chem. Phys.* **2020**, *22* (24), 13764–13779.
- 715 (34) Barth, M.; Gruber, A.; Cichos, F. Spectral and angular
716 redistribution of photoluminescence near a photonic stop band. *Phys.*
717 *Rev. B* **2005**, *72* (8), 085129.
- 718 (35) Schubert, E. F.; Hunt, N. E. J.; Micovic, M.; Malik, R. J.; Sivco,
719 D. L.; Cho, A. Y.; Zydzik, G. J. Highly Efficient Light-Emitting Diodes
720 with Microcavities. *Science* **1994**, *265* (5174), 943–945.
- 721 (36) Wang, J.; Cao, R.; Da, P.; Wang, Y.; Hu, T.; Wu, L.; Lu, J.;
722 Shen, X.; Xu, F.; Zheng, G.; Chen, Z. Purcell effect in an organic-
723 inorganic halide perovskite semiconductor microcavity system. *Appl.*
724 *Phys. Lett.* **2016**, *108* (2), 022103.
- 725 (37) *Excited States and Photochemistry of Organic Molecules*;
726 Klessinger, M.; Michl, J., Eds.; VCH: New York, NY, 1995.
- 727 (38) Goldberg, D.; Menon, V. M. Enhanced amplified spontaneous
728 emission from colloidal quantum dots in all-dielectric monolithic
729 microcavities. *Appl. Phys. Lett.* **2013**, *102* (8), 081119.
- 730 (39) (a) Yamamoto, Y.; Machida, S.; Björk, G. Micro-cavity
731 semiconductor lasers with controlled spontaneous emission. *Opt.*
732 *Quantum Electron.* **1992**, *24* (2), S215–S243. (b) Björk, G. On the
733 spontaneous lifetime change in an ideal planar microcavity-transition
734 from a mode continuum to quantized modes. *IEEE J. Quantum*
735 *Electron.* **1994**, *30* (10), 2314–2318. (c) Björk, G.; Machida, S.;
736 Yamamoto, Y.; Igeta, K. Modification of spontaneous emission rate in
737 planar dielectric microcavity structures. *Phys. Rev. A* **1991**, *44* (1),
738 669–681.
- 739 (40) Scully, M. O.; Zubairy, M. S. *Quantum Optics*; Cambridge
740 University Press: Cambridge, U.K., 1997.
- (41) Frezza, L.; Patrini, M.; Liscidini, M.; Comoretto, D. Directional
741 enhancement of spontaneous emission in polymer flexible micro-
742 cavities. *J. Phys. Chem. C* **2011**, *115* (40), 19939–19946.
- (42) Menon, V. M.; Luberto, M.; Valappil, N. V.; Chatterjee, S.
743 Lasing from InGaP quantum dots in a spin-coated flexible
744 microcavity. *Opt. Express* **2008**, *16* (24), 19535–19540.
- (43) Tanaka, K.; Nakamura, T.; Takamatsu, W.; Yamanishi, M.; Lee,
745 Y.; Ishihara, T. Cavity-Induced Changes of Spontaneous Emission
746 Lifetime in One-Dimensional Semiconductor Microcavities. *Phys. Rev.*
747 *Lett.* **1995**, *74* (17), 3380–3383.
- (44) (a) Chebykin, A. V.; Orlov, A. A.; Shalin, A. S.; Poddubny, A.
751 N.; Belov, P. A. Strong Purcell effect in anisotropic ϵ -near-zero
752 metamaterials. *Phys. Rev. B* **2015**, *91* (20), 205126. (b) Lemmer, U.;
753 Hennig, R.; Guss, W.; Ochse, A.; Pommerehne, J.; Sander, R.;
754 Greiner, A.; Mahrt, R. F.; Bässler, H.; Feldmann, J.; Göbel, E. O.
755 Microcavity effects in a spin-coated polymer two-layer system. *Appl.*
756 *Phys. Lett.* **1995**, *66* (11), 1301–1303.
- (45) Böckler, C.; Reitzenstein, S.; Kistner, C.; Debusmann, R.;
758 Löffler, A.; Kida, T.; Höfling, S.; Forchel, A.; Grenouillet, L.; Claudon,
759 J.; Gérard, J. M. Electrically driven high-Q quantum dot-micropillar
760 cavities. *Appl. Phys. Lett.* **2008**, *92* (9), 091107.
- (46) Siebrand, W., Nonradiative Processes in Molecular Systems. In
762 *Dynamics of Molecular Collisions*, Part A; Miller, W. H., Ed.; Springer:
763 Boston, MA, 1976; pp 249–302.
- (47) Sukhoivanov, I. A.; Guryev, I. V. *Photonic crystals: Physics and*
765 *Practical Modeling*; Springer Series in Optical Sciences, Vol. 152;
766 Springer Verlag: Berlin, Germany, 2009.
- (48) de Mello, J. C.; Wittmann, H. F.; Friend, R. H. An improved
768 experimental determination of external photoluminescence quantum
769 efficiency. *Adv. Mater.* **1997**, *9* (3), 230–232.
- (49) (a) Theander, M.; Inanäs, O.; Mammo, W.; Olinga, T.;
771 Svensson, M.; Andersson, M. R. Photophysics of Substituted
772 Polythiophenes. *J. Phys. Chem. B* **1999**, *103* (37), 7771–7780.
- (b) Valenta, J. Determination of absolute quantum yields of
774 luminescing nanomaterials over a broad spectral range: from the
775 integrating sphere theory to the correct methodology. *Nanoscience*
776 *Methods* **2014**, *3* (1), 11–27.