

Kinetically Controlled Coassembly of Multichromophoric Peptide Hydrogelators and the Impacts on Energy Transport

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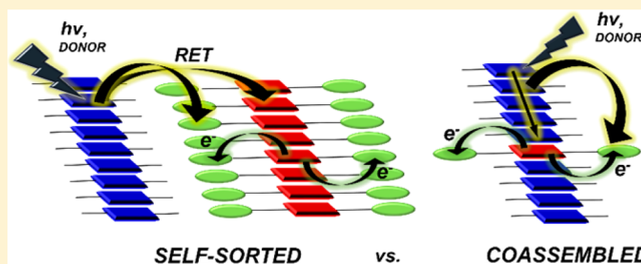
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Supporting Information

ABSTRACT: We report a peptide-based multichromophoric hydrogelator system, wherein π -electron units with different inherent spectral energies are spatially controlled within peptidic 1-D nanostructures to create localized energy gradients in aqueous environments. This is accomplished by mixing different π -conjugated peptides prior to initiating self-assembly through solution acidification. We can vary the kinetics of the assembly and the degree of self-sorting through the choice of the assembly trigger, which changes the kinetics of acidification. The hydrolysis of glucono- δ -lactone (GdL) provides a slow pH drop that allows for stepwise triggering of peptide components into essentially self-sorted nanostructures based on subtle pK_a differences, whereas HCl addition leads to a rapid formation of mixed components within a nanostructure. Using ¹H NMR spectroscopy and fiber X-ray diffraction, we determine the conditions and peptide mixtures that favor self-sorting or intimate comixing. Photophysical investigations in the solution phase provide insight into the correlation of energy-transport processes occurring within the assemblies to the structural organization of π -systems.



INTRODUCTION

Supramolecular systems can be engineered with dynamic properties as a result of the noncovalent nature of the intermolecular interactions that drive their assembly, so they offer promising materials for mimicking the dynamic nature of physiological environments.^{1–3} Multicomponent supramolecular assemblies have been used toward creating artificial light-harvesting complexes, whereby the energy migration from an excited donor to an acceptor occurs within an array of donor–acceptor units.^{4–8} There are several examples of multichromophoric assemblies aimed toward applications not only for photosynthetic mimicry but also for excitonic solar cells,⁹ light-emitting assemblies,^{10–13} and as biophysical molecular rulers.¹⁴ The synthesis details and characterization of energy-transfer events are both equally important for the development of multichromophoric arrays in such applications. For example, Friend, Rowan, and Nolte and co-workers have developed Pt-porphyrin and perylene-bis(dicarboximide) units in polyisocyanopeptides and investigated the sequential energy and electron transfer between the donor and acceptor units.¹⁵ Hodgkiss and Thordarson and co-workers reported a highly efficient multichromophore array based on perylene systems, whereby efficient incoherent energy transfer was reported.¹⁶ Basché and

Müllen and co-workers reported terrylene and perylene diimide dyad systems and quantified singlet–singlet annihilation.^{17,18} Ajayaghosh and co-workers extensively investigated a library of oligo(*p*-phenylenevinylene) organogelators that are capable of excitation energy transfer between specific donor–acceptor systems.^{19–24} A system reported on C₃-symmetric self-sorted units of oligo(*p*-phenylenevinylene) and perylene bisimide was even utilized as a sensing platform for volatile aromatic compounds.²⁵

Among the biologically relevant supramolecular structures, peptide-based assemblies appended with π -electron systems, both in solution phase and in xerogels, hold promise for potential applications toward energy-harvesting materials under aqueous conditions²⁶ as well as solution-processable active layers for device applications such as in field-effect transistors^{27–29} and biointerfacing^{30–34} applications. Tovar and co-workers previously presented a two-component peptide scaffold bearing oligo(*p*-phenylenevinylene) and quaterthiophene π -electron units that demonstrated excitonic energy transfer.³⁵ They also demonstrated that quaterthiophene and

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70 naphthalenediimide chromophores embedded in the same
 71 peptide supported photoinduced electron transfer also under
 72 aqueous conditions.³⁶ Adams and co-workers reported self-
 73 sorting among perylene bisimide and stilbene gelators, creating
 74 heterojunction arrays with self-sorted fiber networks that led to
 75 a photoconductive material.³⁷ Side-chain functionalization of
 76 lysine units with tetraphenylporphyrin and naphthalene diimide
 77 was also shown to demonstrate a switch between the H- or J-
 78 type of aggregation depending on the protonation state of the
 79 porphyrin.³⁸ In a study that utilized chiral cholesterol moieties,
 80 the formation of p-n heterojunctions among self-sorted
 81 quaterthiophene- and perylene diimide organogelators was
 82 achieved via a heat-cool method, whereby the sequential
 83 dissociation and reassembly was governed by the different
 84 dissociation constants of each component.³⁹ We combine three
 85 distinct π -electron units (oligo(*p*-phenylenevinylene), quater-
 86 thiophene, and naphthalenediimide) within peptidic moieties
 87 that form assemblies in aqueous environments under acidic
 88 conditions. The combination of photonic donor-acceptor pairs
 89 and electron donor-acceptor pairs within one nanostructure
 90 enables potentially simultaneous and/or sequential energy- and
 91 electron-transfer events and allows for the creation of localized
 92 electric fields between the donor and acceptor units that result
 93 from charge separation. Moreover, by utilizing different
 94 acidification triggers and controlling the kinetics of self-
 95 assembly, we aim to control the spatial ordering of the
 96 chromophores within the peptidic nanostructure and system-
 97 atically impact the unique energy-transport processes fostered
 98 within these nanostructures.

99 ■ RESULTS AND DISCUSSION

100 **Design Considerations.** We compare the photophysical
 101 responses of self-sorted and randomly coassembled binary
 102 mixtures of peptides bearing different π -electron systems
 103 (Figure 1). To achieve such control over the spatial ordering
 104 of the binary peptide mixtures, we modulated the rate of

acidification. Although there are several studies conducted on
 105 controlling the supramolecular polymerization of multicompo-
 106 nent systems,⁴⁰ accurate control over the assembly dynamics
 107 and spatial ordering is rarely discussed in the context of
 108 engineering the optoelectronic function of a material. A
 109 previous report on the nonequilibrium hydrogelation of a
 110 quaterthiophene⁴¹ demonstrated that the nanoscale morphol-
 111 ogies vary depending on the rate of acid diffusion into the
 112 peptide solution: the rapid assembly leads to random but dense
 113 matted fiber networks, whereas the relatively slower assembly
 114 results in a more homogeneous and anisotropic fiber network.
 115

Here, we use different assembly triggers that lower the pH of
 116 the peptide solutions at different rates in order to have control
 117 over the kinetics of assembly and the molecular ordering of
 118 binary peptide mixtures. Self-sorting was induced via chemical
 119 programming, utilizing the difference in the pK_a of the two
 120 components as the pH is gradually decreased upon the slow
 121 hydrolysis of glucono- δ -lactone (GdL).^{42,43} Thus, as the pH
 122 slowly decreases, we expect a greater extent of protonation/
 123 charge screening for the higher pK_a peptide, triggering it to
 124 preferentially assemble first. This should favor the formation of
 125 essentially self-sorted structures (Figure 1b). To form the
 126 randomly coassembled structures, acid was rapidly added to
 127 solutions of essentially dissolved peptide units, whereby global
 128 protonation/charge screening of all peptides occurs regardless
 129 of the pK_a (Figure 1c). The Asp-Val-Val and Lys-Ala-Ala
 130 peptide sequences were chosen for the oligo(*p*-phenylene-
 131 enevinylene)- and quaterthiophene-appended peptides
 132 (OPV3 and OT4-NDI), respectively, to establish a difference
 133 in the pK_a of each component (see below). The 1,4-
 134 distyrylbenzene, an oligo(*p*-phenylenevinylene), serves as an
 135 energy donor that upon photoexcitation can funnel the energy
 136 to a quaterthiophene unit acceptor. Similarly, the quaterthio-
 137 phene serves as an excited-state electron donor to the
 138 naphthalenediimide in OT4-NDI. A control molecule was
 139 also synthesized in which the side-chain amine was acylated
 140 instead of bearing naphthalenediimide groups (OT4-Ac). This
 141 control molecule allows us to observe photonic energy-transfer
 142 events without the associated electron transfer. We investigate
 143 the occurrence of energy-transport processes under aqueous
 144 environments in the solution phase and whether the
 145 components are self-sorted because of the slow GdL hydrolysis
 146 or kinetically coassembled by rapid acidification.
 147

Characterization of Assembly Behavior. The apparent
 148 pK_a s of the OPV3 donor, the OT4-NDI acceptor, and the OT4-
 149 Ac control acceptor were observed as pH 6.2, 6.5, and 5.4,
 150 respectively (Figure S10). The OT4-NDI-containing samples
 151 showed buffering regions around pH 12. The gradual titration
 152 with aqueous HCl for 1:1 binary mixtures of OPV3/ OT4-NDI
 153 and OPV3/ OT4-Ac only showed one apparent pK_a rather
 154 than a distinct pK_a for each component. To initiate the self-
 155 sorting behavior within the two-component peptide hydrogels,
 156 30 mg/mL of GdL was added to the peptide solutions that
 157 contained 5 mg/mL of each component. In previous studies
 158 with related peptides appended with π -systems, only 5 to 10
 159 mg/mL of GdL was necessary to form hydrogels from 5 mg/
 160 mL one-component or binary-peptide gelator solutions.^{42,43}
 161 The necessary higher GdL concentration for the hydrogelation
 162 of the systems reported herein can be possibly attributed to the
 163 multiple carboxylates (total of four, two from the backbone
 164 termini and two from the aspartic acid side chains, for the
 165 OPV3 donor units) that need to be protonated to favor the
 166 self-assembly process. To further monitor the self-assembly
 167

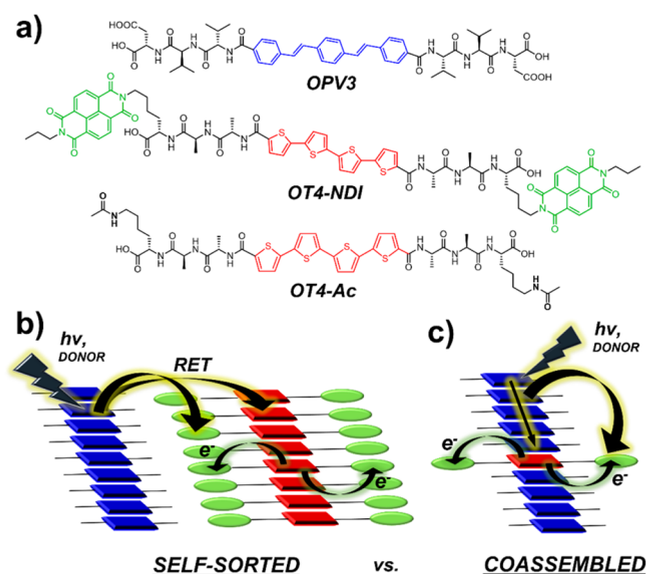


Figure 1. (a) Molecular structures of the peptides studied herein and (b),(c) diagrams of the potential energy (e.g., resonance-energy transfer (RET)) and electron-transfer events occurring within a two-component peptidic nanostructure with three π -electron units for (b) self-sorted and (c) randomly coassembled systems.

168 process with GdL, we followed the disappearance of ^1H NMR
 169 peaks corresponding to the assembly of each gelator
 170 component together with the in situ measurement of pH.⁴⁴
 171 The assembly is correlated to the disappearance of unique
 172 NMR peaks of each component over time, and the results
 173 showed a sequential assembly for the 1:1 OPV3 and OT4-Ac
 174 mixture (Figure 2a), whereby OPV3 assembles first. Alter-

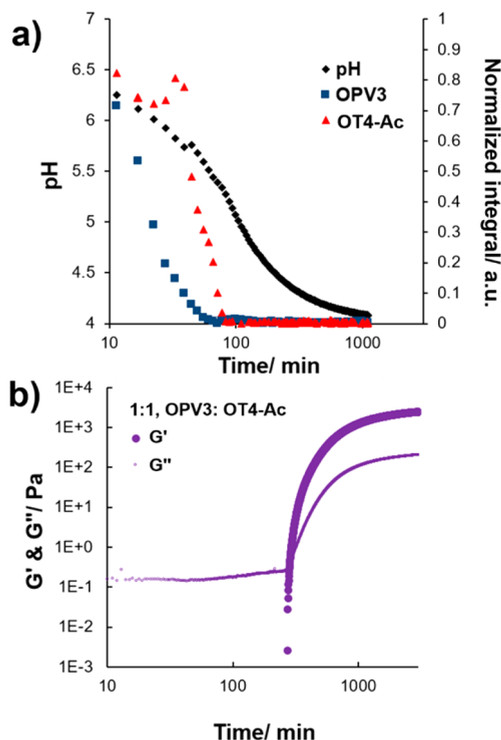


Figure 2. Monitoring of peptide assembly and hydrogel formation for 1:1 OPV3 and OT4-Ac solutions prepared with 30 mg/mL GdL via (a) ^1H NMR and (b) rheology (G' = solid circles; G'' = smaller circles). Each peptide component is at 5 mg/mL (total of 10 mg/mL peptides for the 1:1 mixed sample).

175 natively, the 1:1 OPV3 and OT4-NDI mixture (Figure S11)
 176 did not show any peaks for OT4-NDI even at the beginning of
 177 the measurement (before the addition of GdL), suggesting that
 178 the assembly might have already started for this component
 179 even at higher pH conditions (ca. pH 8). This observation of
 180 assembly formation at higher pH was analogous to a previously
 181 reported hydrophobic oligo(*p*-phenylenevinylene) dipeptide,
 182 which favors the formation of wormlike micelles.⁴⁵ All
 183 homoassemblies (one-component samples) and 1:1 mixed
 184 assemblies successfully formed self-supporting gels, as con-
 185 firmed by the rheology measurements (Figure 2b and S14; the
 186 storage modulus G' was an order of magnitude larger than the
 187 loss modulus G'' at the end of the time-sweep measurements).
 188 The storage moduli values for these hydrogels spanned from
 189 0.02 to 2.5 kPa, with the 1:1 OPV3 and OT4-NDI mixed
 190 assemblies having the lowest G' values and the 1:1 OPV3 and
 191 OT4 mixed assemblies with the assemblies having the highest
 192 G' values. For the 1:1 OPV3 and OT4-NDI mixed assembled
 193 hydrogel (with a total of 10 mg/mL peptide content) the G' is
 194 even lower than its individual components at 5 mg/mL, which
 195 demonstrates the complex and nonlinear nature of the
 196 mechanical properties of two-component supramolecular
 197 systems. These mechanical properties were also observed in

another related binary system composed of dipeptide
 198 components appended with naphthalene units.⁴⁶ Considering
 199 the apparent pK_a values from the titration experiments and the
 200 spectral information from time-resolved NMR measurements,
 201 OT4-NDI assembles first, followed by OPV3 and then OT4-
 202 Ac. The rheology plots show that the OT4-Ac forms the
 203 hydrogel the fastest, followed by OPV3 and then OT4-NDI
 204 (Figure S14). Correlating the ^1H NMR results, which probe
 205 molecular-aggregation events, to the time frame of gelation
 206 observed from rheological measurements, it is observed that the
 207 aggregation time does not coincide with the amount of time
 208 required to observe the inflection points for G' and G'' plots for
 209 this peptidic system. These plots suggest that for this material
 210 and the conditions imposed upon assembly the kinetics of
 211 aggregation of the individual components do not directly
 212 correlate to the rate at which the fibrillar network reaches the
 213 critical fiber-network density to form a hydrogel. Furthermore,
 214 because the fiber network density is linked to the bulk
 215 mechanical property of hydrogels, there is also no direct
 216 correlation that can be made between the kinetics of assembly
 217 formation and hydrogel stiffness for these systems. 218

Further monitoring of the GdL hydrolysis-triggered gelation
 219 process via fiber X-ray diffraction supports the occurrence of
 220 self-sorting based on the presence of distinct signals from the
 221 individual components (Figure 3). No diffraction pattern was
 222 3

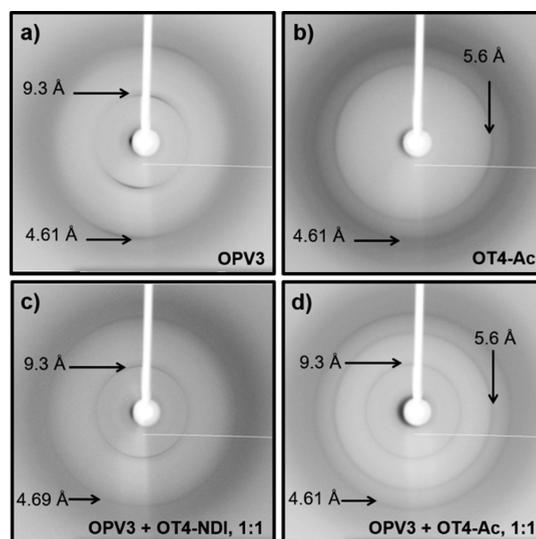


Figure 3. Fiber X-ray diffraction data for hydrogels prepared with 30 mg/mL GdL, composed of one peptide component ((a) OPV3 only; (b) OT4-Ac only) and two peptide components ((c) OPV3/OT4-NDI; (d) OPV3/OT4-Ac; both 1:1). Each peptide component is at 5 mg/mL (total of 10 mg/mL peptides for the 1:1 mixed samples).

observed for OT4-NDI as a result of the difficulty in aligning
 223 the weaker gels ($G' < 100$ Pa) between the capillaries used for
 224 the measurements. The persistence of diffraction signals from
 225 pure OPV3 (9.3 Å and 4.61/4.7 Å, Figure 3a) and pure OT4-
 226 Ac (5.6 and 4.61 Å, Figure 3b) in the mixed-peptide assemblies
 227 prepared with GdL indicates the self-sorted nature of the
 228 nanostructures (Figure 3c,d).⁴² From these characterization
 229 techniques, we determined the chronology of aggregation and
 230 the onset of peptide gelation, and we established that self-
 231 sorting occurs when hydrogels are prepared with GdL for both
 232 OPV3/OT4-Ac and OPV3/OT4-NDI systems. 233

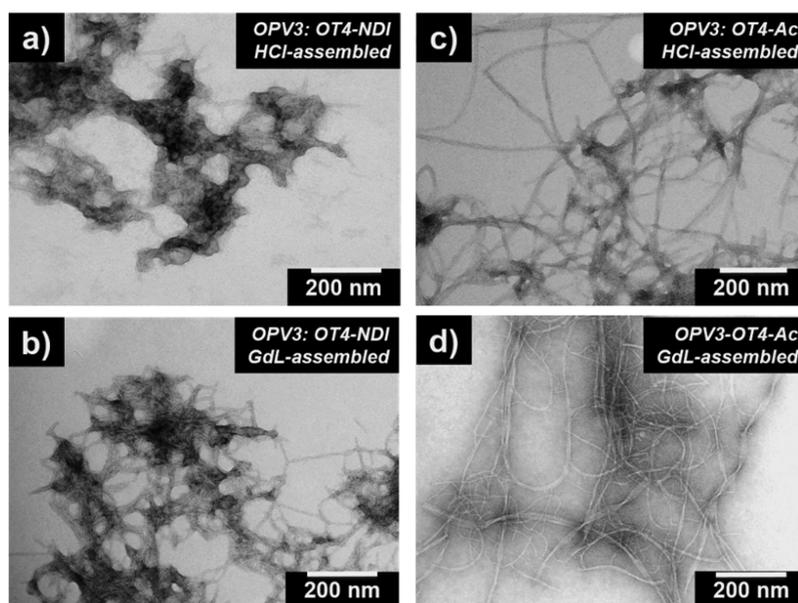


Figure 4. TEM images of 1:1 mixed assemblies ($3 \mu\text{M}$ peptide concentration for each component) of OPV3 with (a),(b) OT4-NDI or (c),(d) OT4-Ac (1:1) assembled via (a),(c) HCl addition and (b),(d) GdL (10 mg/mL) hydrolysis.

234 TEM was used to visualize the nanostructures resulting from
 235 GdL or HCl addition (ca. $3 \mu\text{M}$ peptide concentration for each
 236 component). Under such dilute conditions, 10 mg/mL GdL
 237 was enough to observe nanostructure formation. For the OPV3
 238 and OT4-Ac homoassemblies (Figure S15), both methods of
 239 triggering assembly resulted in the same type of 1-D
 240 morphologies. For the OT4-NDI peptides, rapid acidification
 241 with HCl shows some evidence of micellar-shaped random
 242 aggregates, whereas the addition of GdL results in structures
 243 with 1-D morphology coexisting with these micellar structures.
 244 The predominantly micellar nature of nanostructures formed
 245 by OT4-NDI with GdL and HCl further supports the
 246 formation of relatively weaker gels than for the other peptides
 247 because such low-aspect-ratio structures generally have a lower
 248 propensity to form cross-links among the nanostructures. For
 249 the OPV3 and OT4-NDI mixed assemblies, both the rapid
 250 acidification with HCl and the addition of GdL result in the
 251 formation of micellar structures along with 1-D nanostructures
 252 (Figure 4a,b). In contrast, the mixed assemblies of OPV3 and
 253 OT4-Ac (Figure 4c,d) both show 1-D nanostructures with high
 254 aspect ratios.

255 **Photophysical Characterization.** We utilized the π -
 256 electron units embedded in the peptidic moieties, each with
 257 unique spectroscopic signatures, as reporters to obtain more
 258 information about the spatial organization occurring within the
 259 nanostructures under HCl or GdL assembly conditions. The
 260 steady-state photophysical spectra of the homoassemblies and
 261 mixed assemblies triggered to assemble via HCl and GdL in the
 262 aqueous solution phase (Figures 5, 6, and S16) consistently
 263 showed trends for H-like aggregation. Namely, this included
 264 blue-shifted, quenched absorption and red-shifted, quenched
 265 photoluminescence (PL) of the assembled samples as
 266 compared to their molecularly dissolved states. The recorded
 267 spectra from these aqueous solutions should be treated as
 268 ensemble averages of the response from polydisperse solutions
 269 of peptide assemblies. The spectra for HCl-triggered solutions
 270 were recorded immediately after acidification (i.e., within 1 h)
 271 whereas the GdL-triggered samples were measured at 20 h to
 272 allow enough time for the lactone to hydrolyze and produce

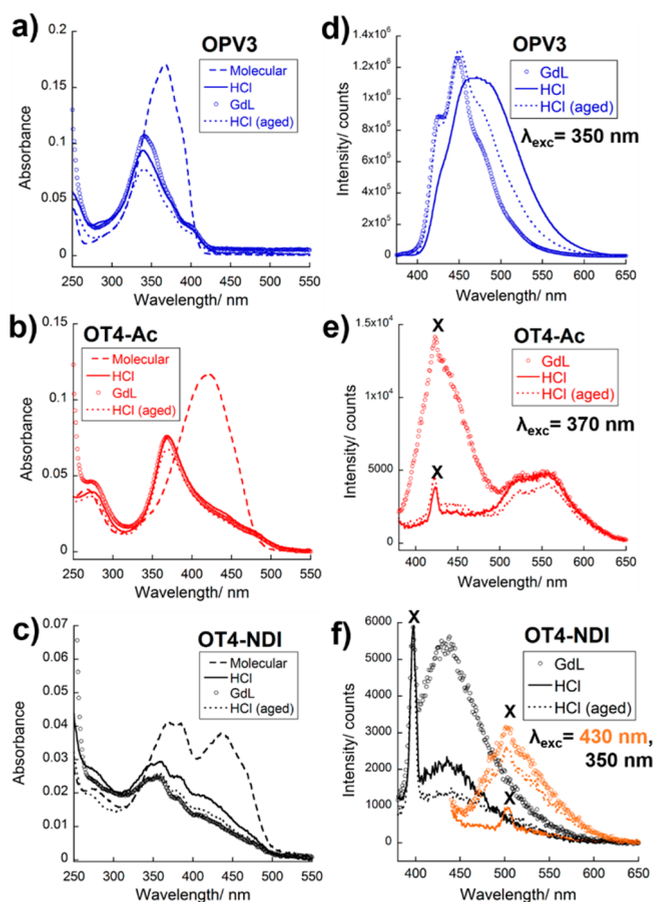


Figure 5. Solution-phase UV-vis absorption (a–c) and steady-state emission (d–f) spectra for GdL- (10 mg/mL) or HCl-assembled peptide homoassemblies of (a) OPV3 (blue), (b) OT4-Ac (red), and (c) OT4-NDI (black); solutions measured under different conditions: molecularly dissolved samples (---), (a),(d) with GdL (empty circles), (b),(e) with HCl (—), and (c),(f) with HCl, aged (...); [peptide] = $3 \mu\text{M}$; X = Raman scatter of water.

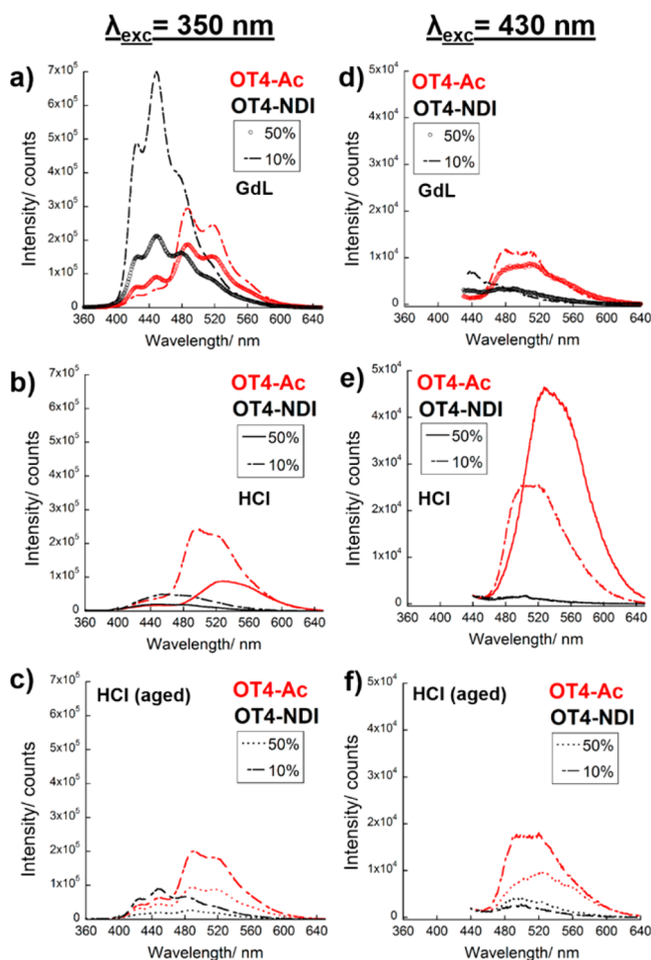


Figure 6. Solution-phase steady-state PL spectra for mixed peptide structures of OPV3 with OT4-Ac (red) and with OT4-NDI (black) prepared with GdL (10 mg/mL) or HCl, measured under different conditions: (a–c) $\lambda_{\text{exc}} = 350$ nm; (d–f) $\lambda_{\text{exc}} = 430$ nm; (a) with GdL (empty circles), (b) with HCl (—), and (c) with HCl, aged (...); [OPV3] = 3 μ M.

273 enough acid to achieve the same pH as the HCl-triggered
274 solutions. The HCl-assembled samples were then aged for the
275 same amount of time that the GdL was allowed to hydrolyze,
276 after which the absorption and PL spectra were then
277 remeasured. For this multichromophoric system, the spectral
278 overlap between the emission of the oligo(*p*-phenylenevinyl-
279 ene) donor with the absorption of quaterthiophene or
280 naphthalenediimide acceptors encourages energy transfer via
281 exciton migration or resonance-energy transfer (RET) from the
282 donor to these acceptors, with a better spectral overlap with
283 quaterthiophene. For quaterthiophene and naphthalenediimide,
284 their relative HOMO and LUMO levels encourage excited-state
285 electron transfer and inhibit energy transfer upon photo-
286 excitation as previously reported.³⁶ As a point of reference, the
287 spectral properties of the homoassemblies prepared via GdL-
288 and HCl-mediated assembly showed no apparent differences in
289 the maximum absorption peaks except for some slight intensity
290 variations (Figure 5a–c). The aged HCl-assembled samples
291 more closely resemble the absorption spectra of the GdL-
292 assembled samples, suggesting the presence of kinetically
293 trapped structures formed by rapid acidification. They also
294 suggest the dynamic nature of these assemblies, which allow
295 some extent of reorganization over time.

The PL spectra for the homoassemblies of OPV3, OT4-Ac, 296
and OT4-NDI all showed different features when assembled via 297
GdL hydrolysis or via rapid HCl addition (Figure 5d–f). The 298
HCl-assembled OPV3 shows a broad emission with $\lambda_{\text{max}} = 470$ 299
nm, whereas the aged HCl- and GdL-triggered assemblies show 300
blue-shifted peaks with more resolved vibronic bands at 420, 301
440, and 470 nm, suggesting a different packing pattern than 302
for the initially trapped HCl assemblies. These bands can be 303
assigned to strong exciton coupling and are consistent with 304
those that we have observed for the thermally annealed 305
structures of other OPV3 tetrapeptides.³⁵ These results support 306
that the slower hydrolysis of GdL could be achieving a more 307
thermodynamically favored structure that is accessed after a 308
dynamic rearrangement due to heating or aging of the 309
kinetically trapped structures formed after rapid acidifica- 310
tion.^{47–49} For OT4-Ac and OT4-NDI excited at 370 and 350 311
nm (maximum absorption for assembled OT4-Ac and OPV3, 312
respectively; naphthalenediimide has minimal absorption at 350 313
nm), the PL peak of the aged HCl-assembled solutions does 314
not vary from the early time-point measurements. The OT4- 315
NDI peptide excited at 350 nm shows an emission profile that 316
corresponds to minimal PL contributions from naphthalene- 317
diimide. However, for excitation at 430 nm, which corresponds 318
to the quaterthiophene component of OT4-NDI, the GdL- 319
assembled and aged HCl samples both show a very weak 510 320
nm peak similar to the maximum PL peak for a monomeric 321
quaterthiophene peptide. The aged HCl-assembled sample 322
shows a completely quenched spectrum (430 nm excitation), 323
with the Raman peak from water at 500 nm as the only 324
apparent feature. Figure 5e,f show signals with very weak 325
intensities and apparent Raman scattering peaks for water. The 326
weak signals are expected from the weakly fluorescent 327
quaterthiophene assemblies but can also be a consequence of 328
the electron transfer from quaterthiophene to naphthalenedi- 329
imide units. 330

For the mixed assemblies, analogous to the observations in 331
our previous study for an oligo(*p*-phenylenevinylene)-based 332
donor–acceptor system,⁵⁰ it is expected that the photophysical 333
behavior will reflect the spatial organization of the π -electron 334
units, whether it is self-sorted via GdL-triggered assembly or 335
randomly coassembled via HCl-triggered assembly. The 336
absorption profiles are dominated by the OPV3 signals, 337
which have the highest extinction coefficient among the three 338
chromophores studied, even at 50 mol % of the acceptor units 339
(Figure S16). At 10 and 50 mol %, higher-intensity shoulders at 340
450 nm are more evident for the HCl-assembled than for the 341
GdL-assembled comixtures. For the PL spectra of mixed 342
assemblies, the samples were excited at 350 nm (where oligo(*p*- 343
phenylenevinylene) has the strongest absorption among the 344
three chromophores) and at 430 nm (where only the 345
quaterthiophene component absorbs). The 10 and 50 mol % 346
OT4-Ac in OPV3 and OT4-NDI in OPV3 are compared 347
under different assembly conditions. In all cases, the PL peak 348
intensities of these mixed-peptide solutions under acidic 349
conditions are lower than that of the OPV3 homoassemblies 350
as shown in Figure 5d. The 10 and 50 mol % OT4-Ac in OPV3 351
GdL self-sorted assemblies excited at 350 nm (Figure 6a) both 352
show quenched emission peaks in the OPV3 region with 353
bimodal peaks around 490 and 520 nm resembling assembled 354
oligothiophene emission in more hydrophobic environ- 355
ments,^{51–54} demonstrating some extent of energy transfer. 356
The self-sorted OT4-NDI and OPV3 GdL-assembled samples 357
showed less OPV3 peak quenching than did the OT4-Ac and 358

359 **OPV3** self-sorted assemblies but maintained the **OPV3**
360 vibronic progression. The HCl-assembled samples (Figure
361 6b) showed a similar trend for quaterthiophene PL signatures
362 but with less-resolved spectral features. Compared to the GdL-
363 assembled samples, the **OPV3** emission region was completely
364 quenched, and the peak was further red-shifted to 540 nm at 50
365 mol %, showing a higher efficiency of energy transfer, which is
366 expected from a more intimate comixing for HCl-assembled
367 samples. Upon aging (Figure 6c), the HCl-assembled samples
368 show more vibronic features in the **OPV3** region but are still
369 more quenched than the self-sorted assemblies prepared with
370 GdL. Furthermore, comparing the **OT4-Ac** and **OT4-NDI**
371 coassemblies with **OPV3** prepared with HCl, the intensities
372 (i.e., relative quantum yields) of the **OT4-NDI** coassemblies
373 are much lower than those of the **OT4-Ac** coassemblies with
374 **OPV3**, verifying the complete energy funneling from oligo(*p*-
375 phenylenevinylene) to quaterthiophene, to relatively spectrally
376 dark naphthalenediimide in the **OT4-NDI/OPV3** coassem-
377 blies. This is indicative that energy transfer is still more efficient
378 in the aged HCl-assembled samples than in the GdL-assembled
379 samples, showing that the reorganization that occurs in the
380 kinetically trapped structures formed upon rapid acidification
381 does not achieve the complete self-sorting that is induced with
382 GdL hydrolysis.

383 For the mixed assemblies excited at 430 nm (Figure 6d–f),
384 the emission spectra provide information on any energy-
385 transfer events that occur from quaterthiophene. The PL signals
386 for self-sorted **OT4-Ac** and **OPV3** mixed assemblies prepared
387 with GdL coincide with the emission of assembled **OT4-Ac**
388 homoassemblies, which is consistent with self-sorting. For the
389 HCl-assemblies at 50 mol % **OT4-Ac**, the PL spectrum has a
390 maximum emission peak at ca. 540 nm, which coincides with
391 what was previously reported for quaterthiophene peptides,^{29,51}
392 whereas the maximum emission peak for 10 mol % **OT4-Ac** at
393 510 nm is reminiscent of molecularly dissolved samples
394 prepared under basic conditions. At 10 mol %, where **OT4-**
395 **Ac** is a minority component in the comixture, the similarity of
396 the PL profile to molecularly dissolved samples suggests that
397 most of the **OT4-Ac** units are isolated within 90 mol % **OPV3**
398 1-D nanostructures. Upon aging of the HCl-assembled samples,
399 the 50 mol % **OT4-Ac** sample showed a similar bimodal peak
400 to that observed in GdL-assembled samples, whereas the 10
401 mol % sample showed a broader quenched peak. For the **OT4-**
402 **NDI/OPV3** HCl-triggered mixed assemblies, the quaterthio-
403 phene PL region showed a significant quenching as compared
404 to the **OT4-Ac** and **OPV3** samples, which are prepared with
405 the same concentration of quaterthiophene units.

406 These results support the idea that the selective electron
407 transfer from quaterthiophene to naphthalenediimide upon
408 excitation at 430 nm efficiently occurs without preference as to
409 the assembly method used. For the mixed assemblies excited at
410 350 nm, the steady-state emission behavior supports that the
411 sequential and/or concerted energy/electron transfer events
412 from **OPV3** to **OT4-Ac/OT4-NDI** occur more efficiently
413 when the components are randomly mixed (prepared with
414 HCl) than self-sorted (prepared with GdL). These data
415 demonstrate the dependence of the photophysical behavior
416 on the choice of assembly trigger (GdL vs HCl) for solution-
417 phase assembly. Hence, the spatial organization of the
418 chromophores within the peptide nanostructures coincides
419 with the structural information obtained for these samples in
420 the gel phase via ¹H NMR and fiber XRD (Figures 2a and 3).

Circular dichroism (CD) measurements confirm that both
421 the HCl and GdL-assembled peptide nanostructures maintain
422 similar chirality in both pure and mixed assemblies, except for
423 **OT4-NDI** (Figure S17). Low-intensity CD signals were
424 recorded for **OT4-NDI** homoassemblies corresponding to the
425 naphthalenediimide region of absorption but not for
426 quaterthiophene. Bisignate signals within the region of
427 chromophore absorption corresponding to an exciton-coupled
428 Cotton band were observed for all other peptide nanostructures
429 prepared under both assembly conditions, which is indicative of
430 the local chiral environment of the π -systems undergoing
431 exciton coupling. The CD signals within the high-energy region
432 that distinguish the structural motifs for conventional peptides
433 were not observed for GdL-assembled samples because of a
434 high intensity signal from chiral GdL and gluconic acid as well
435 as interference from sample scattering (Figure S18). For the
436 HCl-assembled samples, the broad peak with minima at ~250
437 nm corresponds to a red-shifted β -sheet signal, which is
438 correlated to local twisting within individual assembly units.^{55,56} 439

CONCLUSIONS

440 A two-component peptide hydrogelator system with multi-
441 chromophoric units can be spatially engineered on the basis of
442 the choice of the assembly trigger. The slow hydrolysis of GdL
443 provides access to self-sorted structures if each of the
444 components has a different pK_a . However, rapid addition of
445 HCl to the peptide solutions favors the random comixing of
446 two components within the same nanostructure, which can
447 reorganize upon aging. Both methods of assembly led to the
448 formation of self-supporting hydrogels. ¹H NMR spectroscopy
449 and fiber XRD support that self-sorting behavior is induced
450 when GdL is added to the binary peptide mixtures. Correlating
451 the ¹H NMR spectra and rheology profiles, aggregation occurs
452 the fastest for **OT4-NDI**, but the fastest gelation was achieved
453 with **OT4-Ac**. Thus, aggregation can start earlier in one
454 component, but the propagation of the supramolecular
455 polymerization leading to entangled networks that can support
456 a hydrogel structure can occur at a different rate. Whether the
457 two components were randomly coassembled or self-sorted, the
458 chirality within the nanostructures was maintained as evidenced
459 by the exciton-coupled CD bands within the region of the
460 peptide-embedded chromophore absorptions. Both randomly
461 coassembled and self-sorted peptide nanostructures showed
462 evidence for energy and electron transfer. Selectively exciting
463 **OPV3** initiates exciton migration and resonance energy
464 transfer, which both occur more efficiently if the chromophores
465 are randomly coassembled within one nanostructure. However,
466 the excited-state electron transfer from quaterthiophene to
467 naphthalenediimide does not require any specific comixing. This
468 work provides new insights into the photophysical behavior of
469 multichromophoric peptide assemblies whether the compo-
470 nents are self-sorted, which is useful for establishing p-n
471 heterojunctions, or randomly comixed, which is useful for
472 creating photosynthetic mimics that require high energy-
473 transfer efficiency. The observation of multiple energy-trans-
474 port processes in completely aqueous environments, along with
475 the ability to spatially control the distribution of π -electron
476 units within peptidic nanostructures, is an important step in the
477 continued engineering of functional bioelectronic materials. 478

479 ■ ASSOCIATED CONTENT

480 ● Supporting Information

481 The Supporting Information is available free of charge on the
482 ACS Publications website at DOI: 10.1021/jacs.7b04006.

483 General synthesis procedures and experimental condi-
484 tions; characterization data for peptides (¹H NMR,
485 ESI-MS, HPLC traces); pH titration curves; ¹NMR
486 stacked plots and data for monitoring OPV3 and OT4-
487 NDI assembly with GdL; supplementary rheology data,
488 TEM images, UV-vis absorption spectra, and CD
489 spectra (PDF)

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499 Notes

500 The authors declare no competing financial interest.

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