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Peripherical thioester functionalization induces *J*-aggregation in bithiophene-DPP films and nanoparticles†

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In this work we demonstrated that the peripherical thioacetylation of a bithiophene–DPP molecule can greatly influence the solid-state properties triggering the formation of NIR emitting *J*-aggregates in both bithiophene–DPP films and nanoparticles. The morphology and the kinetic and thermal stability of the organic nanoparticles were also investigated.

The control of supramolecular aggregation of organic semiconductors is a critical and fundamental achievement since aggregation strongly affects the solid-state optic, electronic and optoelectronic properties of conjugated systems. Although processing conditions play a key role in the solid-state arrangement of organic semiconductors in thin film,1 chemists are attracted by the possibility to control the organization of conjugated molecules by the structural design.2-5 In this context, it is particularly interesting the case of diketopyrrolo [3,4.c]pyrrole (DPP) derivatives. DPPs are gaining a prominent position among organic dyes for optics and electronics thanks to their simple synthesis, intense color, high photostability, and good charge transport properties.^{6,7} Several studies have been reported on the control of the DPP aggregation varying the conjugation of the core or the N-substitution,8-11 while less explored is the impact on the DPP self-assembly given by the presence of terminal non-conjugated flexible chains. 12,13 In our previous works we observed that varying the length of terminal aliphatic side chains (C8 vs. C16) in a donor-acceptor-donor (DAD) triazole-thiophene-DDP derivative, it was possible to induce the formation of H- or J-aggregates in thin films and organic nanoparticles (ONPs).14,15 Especially, we found that, for both the C₈ and C₁₆ substituted molecules, H-aggregates were preferentially formed during the self-assembly, but in the case

of C₈ substituted species, the resulting ONPs were characterized by low colloidal stability with a remarkable tendency to form micrometric rod-like particles. This morphological evolution of ONPs was associated to an enhancement of the J-aggregate absorption, demonstrating the importance of terminal aliphatic chains in the supramolecular organization of the dye. The possibility to control the formation of H- or J-aggregates is a crucial aspect for the solid-state applications of DDPs. For instance, it has been reported that the increase in the DPP π – π stacking, with the consequent H-aggregation, improves carrier mobility in organic field-effect transistors¹⁶ while, in bulk heterojunction solar cells, DPPs that generate H-aggregates display distinctly lower performances than those forming J-aggregates.17 Furthermore, J-aggregates show NIR photoluminescence and they can be used as efficient solid state NIR emitters. 18-20 In this work we report the effect of introduction of thioacetyl groups at the end of long alkyl chains on the selfassembly behavior of a bithiophene-DPP derivative (DPP4T). The two compared molecules share the same conjugated core, N-substitution (2-ethylhexyl) and terminal aliphatic moieties (hexyl chains), which are unsubstituted for DPP4T-CH3, or functionalized with terminal thioacetyl groups for DPP4T- CH_2SAc (Fig. 1). DPP4T- CH_3 represents the reference system for this study since it leads to the formation of both H- and Jaggregates once in the solid state. 13,21,22 Hence, even small variations in the ratio between the two supramolecular species induced by the introduction of different substituents, can be easily detected and quantified. The interest about the correlations between the thioester functionalization and the supramolecular assembly was driven by recent findings on the organization of photoactive ternary blends in the presence of -SAc functionalized oligo(arylenethiophene)s.23 In fact, we showed that thioester polarizable group can modify the interactions with the solvent used for the processing, also impacting on the solid-state organization.

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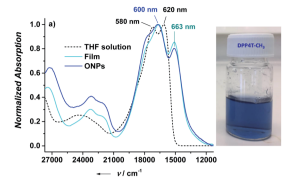
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[†] Electronic supplementary information (ESI) available: Synthesis, cyclic voltammetry, UV-vis and fluorescence characterization, film preparation and AFM characterization, ONPs preparation and DLS characterization. See DOI: 10.1039/d1ra01253c

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In the present study, in order to distinguish the two effects, DPP4T-CH₃ and DPP4T-CH₂SAc were processed with different techniques and in different conditions: as thin films by spincoating from a chloroform solution, or as ONPs obtained via re-precipitation method from a tetrahydrofuran (THF) solution in water. The two DPP derivatives were synthesized through a palladium-catalysed Suzuki-Miyaura coupling, a straightforward metal-catalyzed cross-coupling reaction.24,25 Synthetic details are provided in the ESI.† First, we performed the electrochemical and the optical characterization of the two dyes in solution. From the comparison between the cyclic voltammograms of the two species (Fig. S1†) it is clear that the peripherical thioacylation has negligible effects on the electrochemical bandgap (E_g) of the conjugated molecules. Both the oligomers show HOMO and LUMO energies of -5.4 and -3.5 eV respectively, with a resulting bandgap of 1.9 eV (Table S1†). Results are in perfect agreement with the literature data available for DPP4T-CH₃ ^{21,22,26} and with the UV-VIS characterization in different solvents (CHCl3, THF, toluene). In fact, from the comparison between the normalized UV-vis spectra of the two dyes in THF (black dotted line in Fig. 2a and b), only minor differences in the relative intensities between the vibronic peaks at 620 and 580 nm can be detected. This difference in the relative intensity between the two absorption maxima can be observed also in other solvents (Fig. S2a and b†), suggesting a different solvation behaviour for two molecules due to the presence of the -SAc groups. The fluorescence is peaked in the red region at around 650 nm (Fig. S2c and d†), with a Stoke's shift of ≈30 nm. The main optical properties in solution are listed in Table S2.† The solid-state aggregation properties were investigated on thin films of both species on quartz glasses (details in ESI†). As already reported by Pron and co-workers, DPP4T-CH₃ exhibits poor filming properties,²² and even testing different solvents it was not possible to obtain uniform depositions. Contrarily, DPP4T-CH₂SAc can be deposited as thin film independently on the solvent used, indicating an easy processability of thioester-functionalized organic semiconductors.23 The high uniformity of DPP4T-CH2SAc film was confirmed by the AFM analysis (Fig. 3a), that revealed a film thickness of \approx 137 nm and a surface roughness (R_g) of 2.42 \pm 0.1 nm. The absorption behaviour of the two molecules as thin film completely differs as can be seen from Fig. 2. DPP4T-CH₃ film presents an absorption maximum blue-shifted with respect to the maximum in solution and attributed to the formation of H-aggregates, and a further, red-shifted relative maximum, associated to the *J*-species. The blue-shifted absorption peak is at 600 nm with a H-shift of 540 cm⁻¹ and a shoulder at 566 nm,

Fig. 1 Molecular structure of DPP4T-CH₃ and of the thioacetate species DPP4T-CH₂SAc.



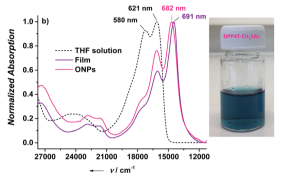


Fig. 2 Normalized absorption spectra for DPP4T-CH₃ (a) and DPP4T-CH₂SAc (b) in THF solution, as neat film and as ONP suspension in water. On the right are reported the picture of ONP suspensions of both species.

while the peak of J-aggregates is red-shifted of 1020 cm⁻¹ at 663 nm. For the DPP4T-CH₂SAc oligomer instead, we record a remarkable trend in the formation of J-aggregates, with a bathochromic shift of the absorption maximum of 1660 cm⁻¹. The spectrum presents a main peak at 691 nm, and two vibronic peaks at 623 and 570 nm ($\Delta E \approx 1550 \text{ cm}^{-1}$) with no clear evidence of H-species. The lower tendency to form stacked aggregates of acetylated derivative is supported also from the DSC analysis reported in Fig. 3b. No liquid crystal phase transition was apparently observed for any of the molecules. The melting $(T_{\rm m})$ and crystallization $(T_{\rm c})$ temperatures measured for the two dyes are lower with respect to those shown by other DPP compounds with unbranched chain substitution on the lactam

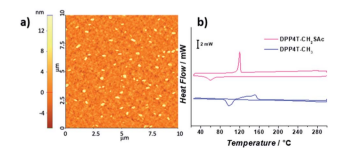


Fig. 3 (a) AFM $10 \times 10 \, \mu \text{m}^2$ topography of DPP4TS-CH₂SAc thin-film processed from a chloroform solution; (b) DSC thermograms of DPP4T-CH₃ (blue line) and DPP4T-CH₂SAc (pink line) recorded between 30 and 300 °C at a scan rate of 10 °C min⁻¹. Scale bar 2 mW.

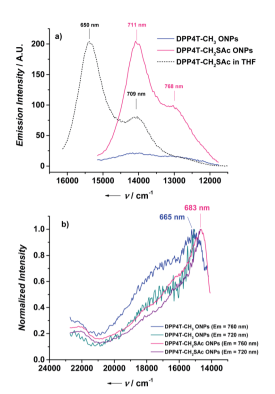


Fig. 4 (a) Emission spectra of DPP4T-CH $_3$ ONPs (excited at 600 nm, absorbance = 0.295), DPP4T-CH $_2$ SAc (excited at 620 nm, absorbance = 0.536), and the normalized emission spectrum of DPP4T-CH $_2$ SAc in THF (dashed black line); (b) normalized excitation spectra of DPP4T-CH $_3$ and DPP4T-CH $_2$ SAc ONPs recorded at different emission wavelength.

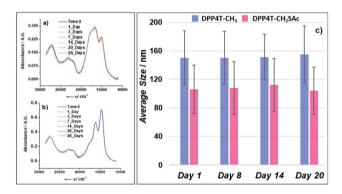


Fig. 5 (a) Absorption spectra of a DPP4T-CH₃ ONP suspension acquired during 28 days; (b) absorption spectra of a DPP4T-CH₂SAc ONP suspension acquired during 28 days; (c) evolution of the average size distribution of DPP4T-CH₃ and DPP4T-CH₂SAc ONPs during 20 days. The error bar represents the polydispersity index expressed in nm.

nitrogen. This has been attributed by other authors to the increase of intermolecular distance in aggregates produced by the 2-ethylhexyl substitution.²⁷ The **DPP4TS-CH₂SAc** thermogram displays sharp and narrow melting and crystallization transitions, centred at 120 and 60 °C respectively. The presence of a single and sharp peak is consistent with the formation of a single aggregate species, likely *J*-like as already observed in the

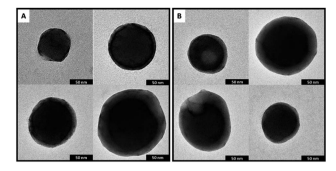


Fig. 6 (A) Representative TEM images of samples DPP4T-CH $_3$ and, (B) DPP4T-CH $_2$ SAc 1 day after the preparation.

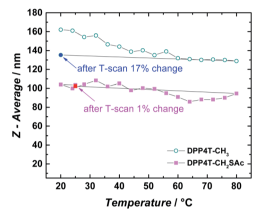


Fig. 7 Temperature dependence of the Z-average diameter of DPP4T-CH₃ and DPP4T-CH₂SAc ONP suspensions in water, varying the temperature in the range 20–80 °C. After the heating, suspensions were cooled at room temperature and the Z-average was measured again.

absorption spectra. Conversely, phase transitions for DPP4T-CH₃ look more complex, with a broad endothermic phase transition appearing as a bi-modal peak at 133 and 151 °C, and the exothermic phase transition centred at 98 °C. The broad and higher transition temperatures are consistent with the hypothesis of the oligomer DPP4T-CH₃ forming different aggregates with the highest melting energy required. The capability of the SAc-functionalization to trigger the formation of DPP4T Jaggregates independently on the processing technique was also tested on suspensions of organic nanoparticles (ONPs) in water. The normalized absorption spectra of the DPP4T-CH₃ and DPP4T-CH₂SAc ONPs are reported in Fig. 2. For both molecules, the ONP absorption spectra resemble those of the films. Actually, the major differences can be seen at higher frequency due to the scattering of particles. DPP4T-CH₃ ONPs spectrum is characterized by peaks related to H- and J-aggregates at 598 and 663 nm respectively, as in the case of film. The peak at 663 nm results less intense in ONPs, suggesting a lower tendency to form J-like species in these conditions. Even the spectrum of DPP4T-CH₂SAc retraces the film spectrum, with the maximum of absorption at 682 nm (691 nm for the film), and the same vibronic distribution. Contrarily to non-emissive DPP4T-CH₃

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ONPs, DPP4T-CH₂SAc ONPs gives photoluminescence in the NIR. As can be seen from Fig. 4a, the maximum of emission shifts from 650 to 711 nm ($\Delta E \approx 1300 \text{ cm}^{-1}$) moving from the THF solution to the ONPs suspension, with a shoulder at 768 nm ($\Delta E \approx 1050 \text{ cm}^{-1}$). The coexistence of different kinds of aggregates in DPP4T-CH₃ ONPs is confirmed from the analysis of the excitation spectra in Fig. 4b. In fact, the excitation spectra acquired close to the two maxima of emission (720 and 760 nm) have a different band-shape and they both differ from the absorption spectrum.9 In particular, the maximum of the excitation spectrum coincides with the maximum of absorption associated to the J-species at 663 nm. Instead, the excitation spectra measured at different wavelengths of DPP4T-CH₂SAc ONPs are quite similar, indicating that the luminescence at the two maxima is originated by the same species. Moreover, the excitation band-shape perfectly resembles the absorption spectrum, with a maximum at 683 nm and the same vibronic distribution. Combining the information derived from DSC and the excitation data it is clear that the peripherical SAc-groups inhibit the H-organization and strongly promote the formation of J-aggregates, which exhibit a dipole allowed transition in the NIR.13 This aspect, together with the low absorbance of biological tissues in this region, makes this system extremely interesting for the development of fluorescent biomarkers for in vitro and in vivo experiments. Finally, we studied the colloidal stability of the two ONP suspensions. In Fig. 5 are reported the absorption spectra on DPP4T-CH3 and DPP4T-CH2SAc ONPs acquired during four weeks. Both systems are characterized by a remarkable stability. All the spectra maintain unaltered the band-shape without showing any increasement of the scattering, that implies that no further agglomeration or precipitation occurred during this lapse of time. Results were confirmed by the dynamic light scattering (DLS) analysis (Fig. 5c). The DPP4T-CH₃ ONPs show a monomodal distribution (Fig. S3†) with an average diameter of 150 nm and a width of about 70 nm (corresponding to a polydispersity index PDI = 0.22). The introduction of the thioacetyl group generates smaller nanoparticles. In fact, for the DPP4T-CH₂SAc ONPs, the average size is 110 nm with a width of 70 nm (PDI = 0.40). The intensity distribution in this case exhibits a bi-modal distribution with two peaks at 15 and 110 mm (Fig. S4†). TEM images of samples, one day after their preparation, highlight the presence of nearly spherical objects in both DPP4T-CH3 and DPP4T-CH2SAc samples (Fig. 6). The statistical analysis of the nanoparticle size is in good agreement with DLS data. Indeed, the average diameter of sample **DPP4T-CH**₃ is 114 nm ($\sigma = 19\%$), while the mean diameter of sample **DPP4T-CH₂SAc** is 106 nm (σ = 46%). During 20 days, storing the samples at 4 °C, no variations in the size distribution of the two species occurred. In order to better characterize the two systems, we studied the effects of the temperature and ionic strength on their colloidal stability. DPP4T-CH3 e DPP4T-CH2SAc ONP suspensions in water were heated in the temperature range 20-80 °C and the Zaverage diameter was measured with a step of 5 °C. Finally, suspensions were cooled to room temperature and their size was measured again. As can be seen in Fig. 7, after this

CH₂SAc ONPs was observed, while the unsubstituted system exhibited a size reduction of 17%.

Interestingly, these temperature changes did not affect the supramolecular aggregation of the two oligomers in ONPs, since negligible variations in their absorption spectra were detected (Fig. S5†). Indeed, in the case of **DPP4T-CH₂SAc ONPs**, varying the temperature from 25 to 65 °C, just an increment of about 7% was observe for the absorption peak at 660 nm (Fig. S5†).

Regarding the impact of ionic strength, in both cases, the addition of 1 ml of phosphate buffered saline to 1 ml of ONP suspension (final salt concentration: phosphate buffer 0.01 M, KCl 0.0027 M, NaCl 0.137 M) did not induce any precipitation at least for one week. Contrarily with what we observed in our previous studies on triazole-thiophene-DPP ONPs, 14,15 where the J-aggregation was associated with a poor colloidal stability and with a remarkable tendency to aggregation, DPP4T-CH₂SAc **ONPs** possess a high colloidal stability (observed also for the other derivative). Moreover, the presence of -SAc terminal group induces the formation of NIR emitting *I*-aggregates, opening the way to a promising system for the in vitro and in vivo bioimaging. In conclusion we demonstrated that the simple introduction of a bulky and polarizable terminal thioacetyl group on alkyl chains attached to a thiophene-DPP core, is able to improve the processability of the molecule and to strongly condition the supramolecular aggregation with the formation of stable NIR emitting J-aggregates, independently on the processing technique and conditions.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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Notes and references

- D. Blasi, F. Viola, F. Modena, A. Luukkonen, E. MacChia, R. A. Picca, Z. Gounani, A. Tewari, R. Österbacka, M. Caironi, Z. M. Kovacs Vajna, G. Scamarcio, F. Torricelli and L. Torsi, *J. Mater. Chem. C*, 2020, 8, 15312–15321.
- Ku, D. Gao, J. Song, L. Shen, W. Chen, Y. Chen and S. Zhang, New J. Chem., 2015, 39, 5553-5560.
- 3 Y. Gao, G. Feng, T. Jiang, C. Goh, L. Ng, B. Liu, B. Li, L. Yang, J. Hua and H. Tian, *Adv. Funct. Mater.*, 2015, **25**, 2857–2866.
- 4 A. Punzi, M. A. M. Capozzi, V. Fino, C. Carlucci, M. Suriano, E. Mesto, E. Schingaro, E. Orgiu, S. Bonacchi, T. Leydecker, P. Samorì, R. Musio and G. M. Farinola, *J. Mater. Chem. C*, 2016, 4, 3138–3142.
- 5 M. A. M. Capozzi, A. Punzi, F. Babudri, R. Musio and G. M. Farinola, J. Org. Chem., 2018, 83, 14396–14405.

temperature scan, no variations in the average size of DPP4T-

- 6 M. Grzybowski and D. T. Gryko, *Adv. Opt. Mater.*, 2015, 3, 280–320.
- 7 Y. Patil and R. Misra, *J. Mater. Chem. C*, 2019, 7, 13020-13031.
- 8 N. Genevaz, P. Chávez, V. Untilova, A. Boeglin, C. Bailly, L. Karmazin and L. Biniek, *J. Mater. Chem. C*, 2018, **6**, 9140–9151.
- M. Kirkus, L. Wang, S. Mothy, D. Beljonne, J. Cornil,
 R. A. J. Janssen and S. C. J. Meskers, *J. Phys. Chem. A*, 2012,
 116, 7927–7936.
- 10 T. G. Hwang, J. Y. Kim, J. W. Namgoong, J. M. Lee, S. B. Yuk, S. H. Kim and J. P. Kim, *Photochem. Photobiol. Sci.*, 2019, 18, 1064–1074.
- 11 C. M. Mauck, P. E. Hartnett, E. A. Margulies, L. Ma, C. E. Miller, G. C. Schatz, T. J. Marks and M. R. Wasielewski, J. Am. Chem. Soc., 2016, 138, 11749–11761.
- 12 X. A. Jeanbourquin, A. Rahmanudin, A. Gasperini, E. Ripaud, X. Yu, M. Johnson, N. Guijarro and K. Sivula, *J. Mater. Chem. A*, 2017, 5, 10526–10536.
- 13 V. S. Gevaerts, E. M. Herzig, M. Kirkus, K. H. Hendriks, M. M. Wienk, J. Perlich, P. Müller-Buschbaum and R. A. J. Janssen, *Chem. Mater.*, 2014, 26, 916–926.
- 14 A. Punzi, E. Maiorano, F. Nicoletta, D. Blasi, A. Ardizzone, N. Ventosa, I. Ratera, J. Veciana and G. M. Farinola, *Eur. J. Org. Chem.*, 2016, 2016, 2617–2627.
- 15 A. Ardizzone, D. Blasi, D. Vona, A. Rosspeintner, A. Punzi, E. Altamura, N. Grimaldi, S. Sala, E. Vauthey, G. M. Farinola, I. Ratera, N. Ventosa and J. Veciana, *Chem. –Eur. J.*, 2018, 24, 11386–11392.

- 16 M. Shaker, B. Park, J.-H. Lee, W. Kim, C. K. Trinh, H.-J. Lee, J. woo Choi, H. Kim, K. Lee and J.-S. Lee, RSC Adv., 2017, 7, 16302–16310.
- 17 M. Más-Montoya and R. A. J. Janssen, *Adv. Funct. Mater.*, 2017, 27, 1605779.
- 18 J. Li, Y. Liu, Y. Xu, L. Li, Y. Sun and W. Huang, Coord. Chem. Rev., 2020, 415, 213318.
- 19 D. Blasi, D. M. Nikolaidou, F. Terenziani, I. Ratera and J. Veciana, *Phys. Chem. Chem. Phys.*, 2017, **19**, 9313–9319.
- 20 A. Barbieri, E. Bandini, F. Monti, V. K. Praveen and N. Armaroli, *Top. Curr. Chem.*, 2016, 374, 47.
- 21 Y.-F. Huang, S.-T. Chang, K.-Y. Wu, S.-L. Wu, G.-T. Ciou, C.-Y. Chen, C.-L. Liu and C.-L. Wang, ACS Appl. Mater. Interfaces, 2018, 10, 8869–8876.
- 22 G. Wiosna-Salyga, M. Gora, M. Zagorska, P. Toman, B. Luszczynska, J. Pfleger, I. Glowacki, J. Ulanski, J. Mieczkowski and A. Pron, RSC Adv., 2015, 5, 59616–59629.
- 23 A. Punzi, A. Operamolla, O. Hassan Omar, F. Brunetti, A. D. Scaccabarozzi, G. M. Farinola and N. Stingelin, *Chem. Mater.*, 2018, 30, 2213–2217.
- 24 A. Punzi, N. Zappimbulso and G. M. Farinola, *European J. Org. Chem.*, 2020, 2020, 3229–3234.
- 25 A. Operamolla, S. Colella, R. Musio, A. Loiudice, O. Hassan Omar, G. Melcarne, M. Mazzeo, G. Gigli, G. M. Farinola and F. Babudri, *Sol. Energy Mater. Sol. Cells*, 2011, 95, 3490–3503.
- 26 M. Gora, S. Pluczyk, P. Zassowski, W. Krzywiec, M. Zagorska, J. Mieczkowski, M. Lapkowski and A. Pron, *Synth. Met.*, 2016, 216, 75–82.
- 27 C. Kim, J. Liu, J. Lin, A. B. Tamayo, B. Walker, G. Wu and T.-Q. Nguyen, *Chem. Mater.*, 2012, 24, 1699–1709.