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Introduction

Discovered in the early $1970s¹$, diketopyrrolopyrroles $(2,5$ -dihydropyrrolo[4,3-c]pyrrolo-1,4-diones, DPPs) remain one of the most widely used dyes, finding application in many areas of high technology. Numerous studies have been devoted to the development of DPP derivatives as high-performance materials for use in various applications such as high-performance pigments, organic field-effect transistors, bulk-heterojunction solar cells, dye-sensitized solar cells, organic light-emitting

Synthesis and photoinduced behavior of DPPanchored nitronyl nitroxides: a multifaceted approach†

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Understanding and controlling spin dynamics in organic dyes is of significant scientific and technological interest. The investigation of 2,5-dihydropyrrolo[4,3-c]pyrrolo-1,4-dione derivatives (DPPs), one of the most widely used dyes in many fields, has so far been limited to closed-shell molecules. We present a comprehensive joint experimental and computational study of DPP derivatives covalently linked to two nitronyl nitroxide radicals ($DPP^{Th}-NN_2$). Synthesis, single crystal X-ray diffraction study, photophysical properties, magnetic properties established using steady-state and pulse EPR, fast spin dynamics, and computational modelling using density functional theory and ab initio methods of electronic structure and spectroscopic properties of DPP^{Th} -NN₂ are presented. The single-crystal X-ray diffraction analysis of DPP^{Th} -NN2 and computational modeling of its electronic structure suggest that effective conjugation along the backbone leads to noticeable spin-polarization transfer. Calculations using ab initio methods predict a weak exchange interaction of radical centers through a singlet ground state of DPPTh with a small singlet–triplet splitting (Δ E_{ST}) of about 25 cm⁻¹ (~0.07 kcal mol⁻¹). In turn, a strong ferromagnetic exchange interaction between the triplet state of DPPTh chromophore and nitronyl nitroxides (with $J \sim 250$ cm⁻¹) was predicted. **PAPER**
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diodes, fluorescence imaging, and many other fields.² Despite the significant progress achieved by researchers in the exploration of DPP-based compounds, only recently have their conjugated derivatives substituted with stable radical groups been obtained. Single crystal X-ray data on a DPP derivative bearing radical groups, namely, bis(thiophen-2-yl)-2,5-dihydropyrrolo[3,4-c]pyrrole-1,4-dione with terminal nitronyl nitroxide groups $(DPPTh-NN₂)$ were first reported in 2020.³

Later, nitronyl-nitroxide diradicals with extended DPPcontaining linkers ($DPPTh-Ph-NN₂$, $DPP^{Fu}-Ph-NN₂$, and DPP^{Ph} -Ph-NN₂, Fig. 1) were obtained and partially characterized.⁴ In addition to the listed diradicals (Fig. 1), perylene bisimide (PBI) and iso-indigo (IIn) conjugated diradicals were also synthesized.⁴ Although all diradicals retain the intrinsic optical properties of the dyes or, more specifically, pi-conjugated chromophore, at the same time they exhibit an indirect spin coupling between two distant paramagnetic centres. Room temperature EPR data⁴ provided only nearly identical 9-line spectra, demonstrating that the exchange coupling parameter *J* is much larger than twice the hyperfine coupling constant. The distances between the radical units were very large $(C2-C2)$ distance in the range 2.2–2.4 nm for DPP-based diradicals) due to the additional phenyl rings connecting the NN moieties to the DPP core.⁴ To date, comprehensive temperature-dependent studies of liquid and frozen solutions via EPR spectroscopy, as well as magnetization measurements of polycrystalline powder,

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[†] Electronic supplementary information (ESI) available: Electronic absorption and FTIR spectra, CV profiles versus the Fc⁺/Fc redox couple, TGA and DSC curves of DPPTh-NN₂, details of DFT and ab initio calculations were presented. CCDC 1908216 for DPPTh-NN₂. For ESI and crystallographic data in CIF or other electronic format see DOI: <https://doi.org/10.1039/d4ra00916a>

Fig. 1 Chemical structures of DPP-linked nitronyl-nitroxide diradicals: DPPTh-NN₂, DPPTh-Ph-NN₂, DPP^{Fu}-Ph-NN₂, and DPP^{Ph}-Ph-NN₂.

have not been conducted. These analyses are crucial for a deeper understanding and clear differentiation of the magnetic properties of such diradicals. Earlier, Matsuda et al. demonstrated the exponential distance dependence of the exchange interaction parameter over extended phenylacetylene bridged NN biradicals.⁵ Their findings indicate that the EPR spectra of the examined diradicals in liquid phase will display a consistent nine-line pattern (indicative of strong exchange) when the distance between the C2 atoms of the NN moieties ranges from 0.7 to 2.7 nm. This observation is coupled with significant variations (spanning orders of magnitude) in the exchange coupling parameters, highlighting the sensitivity of magnetic interactions to molecular spacing within these systems.

Thus, the magnetic properties of DPP-based diradicals require much more in-depth studies. Since DPP derivatives have found applications in the solar cells, OLEDs, fluorescence imaging, etc.,^{2d-g} information available about their electronic absorption spectra is insufficient to understand their excitation relaxation dynamics which is very important to broaden the scope of these molecules in respective fields. Our study was undertaken with the aim of quantitatively elucidating the magnetic properties of the DPP^{Th} -NN₂ diradical. Continuouswave and time-resolved EPR spectra were recorded, and exchange and dipole–dipole interactions were theoretically assessed not only in the ground state of DPP linker, but also in the case of its triplet excitation. Furthermore, we tried to resolve the issue of possible relaxation pathways for the excitation of the DPP chromophore. To do this, we performed both DFT and high-level calculations of the electronic structure of the DPPTh-NN2 diradical. In this paper, we described in detail the synthesis of $DPPTh-NN₂$, single crystal analysis, insights into electronic structure, and the results on the continuous wave, echodetected and time-resolved EPR study.

Results and discussion

Synthesis and structure of the DPP^{Th} -NN₂ diradical

The synthetic route to the $DPP^{Th}-NN_2$ diradical is shown in Scheme 1. The diradical precursor, the corresponding dibromo-

substituted $DPP^{Th} - Br_2$, was synthesized according to the literature-reported procedure⁶ with necessary modifications. To introduce two nitronyl nitroxide (NN) groups into the DPPTh core, we chose a palladium catalysed cross-coupling of bromoderivative of DDP with the (nitronyl nitroxide-2 ide)(triphenylphosphine)gold complex. This method, originally proposed by Keiji Okada,⁷ with the recent significant improvements^{8,9} has become an efficient protocol for the directed synthesis of nitronyl nitroxides.¹⁰⁻¹² Thus, a new path has been opened to various in-demand nitronyl nitroxides for a variety of research fields, including radical chemistry, the creation of high-spin systems and rechargeable batteries, the design of molecule-based magnets and molecular units for spintronics.^{13,14} Naturally, using the described approach, the $DPPTh-NN₂$ diradical was successfully obtained in a moderate yield of \sim 72% when using Pd(PPh₃)₄ catalyst (Scheme 1).

In chemistry of diradicals, X-ray diffraction analysis is of paramount importance because it unveils valuable information about molecular organization and interactions in the solid state. We succeeded in growing single crystals of $DPP^{Th}-NN_2$ by slow diffusion of methanol into its solution in CH_2Cl_2 at 5 °C. The crystalline product appeared to be stable at ambient conditions and no phase transitions were observed in the DSC analysis in the temperature range from 20 °C to the melting point of 216 °C (see ESI†).

At room temperature, attempts to perform an X-ray diffraction analysis of DPP^{Th} -NN₂ single-crystal samples failed. Cooling the single-crystal sample at 193 K led to the freezing of the alkyl-chain movement, which made it possible to solve the molecular and crystal structure of the $DPP^{Th}-NN_2$ diradical. According to XRD, the crystal structure of $DPP^{Th}-NN_2$ is centrosymmetric (space group $\overline{P_1}$) and contains two crystallographically independent molecules in an asymmetric unit (hereafter called as A and B). The molecular structures of A and B DPPTh-NN₂ molecules are depicted in Fig. 2. Bond lengths and angles in both forms of the molecule are similar (Table S1†) and comparable to typical values for similar molecular fragments, as indicated by a search in the Cambridge Structural Database using the Mogul program.¹⁵

Scheme 1 Synthesis of the $DPP^{Th}-NN_2$ nitronyl nitroxide diradical

Fig. 2 Structures of crystallographically independent molecules A and B (with solvate CH_2Cl_2) in crystals of the DPPTh-NN₂ diradical.

In the DPPTh-NN₂ diradical, the π -conjugated system, consisting of the DPP core, thiophene rings, and nitronyl nitroxide moieties, presents high planarity with small interplanar twist angles. In molecules A and B, dihedral torsion angles between planes of the nitronyl nitroxide groups and planes of the nearest side thiophene rings are 2.90 and 13.24° respectively. In turn, the two thiophene rings are imposed in anti-orientation with respect to each other and twisted by 9.16° (in molecule A) and 20.69 \degree (in molecule B) relative to the mean plane of DPPTh. The observed planarity of the thiophene–nitronyl nitroxide system leads to short intramolecular H-bonds O \cdots H–C (2.41 and 2.45 Å, dotted lines in Fig. 2), as well as extremely short contacts between O_{NO} and S atoms (2.74 and 2.79 Å) compared to the sum of the van der Waals radii of the S and O atoms (3.32 Å, dashed purple lines in Fig. 2). Small torsion angles provide efficient conjugation along the backbone, thereby facilitating the noticeable transfer of spin-polarization from the NN radical units to the DPPTh chromophore, as observed in the EPR spectra and demonstrated by DFT calculations (Fig. S6, ESI†).

The crystal packing shows that the molecules adopt weak intermolecular π – π stacking separated by distance of 3.74 Å between centroids of the DPPTh moieties (Fig. 3). Inside such stacks, short intermolecular contacts C-H \cdots O_{NO} (2.36, 2.45, 2.56, and 2.64 Å) are realized between the oxygen atoms of the nitroxide fragments and the hydrogen atoms of the methyl groups (Fig. 4a). Similar contacts are often observed in crystals of nitronyl nitroxides, and they are characterized by a rather high binding energy.¹⁶ Because of this, interactions of this type have a significant effect on the motif of the crystal packing of nitroxide radicals. The most important consequence of C-H \cdots O_{NO} interactions is the geometry of the mutual arrangement of nitroxide groups and, above all, the distances between the oxygen atoms of paramagnetic centers, since the latter

Fig. 3 (a) Solid state molecular packing of DPPTh-NN₂ with $\pi-\pi$ -interactions; (b) side view of DPPTh-NN₂ with $\pi-\pi$ -distances between centroids of the DPPTh moieties.

predetermine the magnetic properties of paramagnetic samples. Moreover, C-H \cdots O_{NO} interactions predetermine the relative rotation angle of molecules in the stacks, as well as the shortest intermolecular distances between nitroxide oxygen atoms (3.617 and 3.887 Å). In addition, one can see multiple short intermolecular C–H \cdots O_{NO} contacts (2.40, 2.55, and 2.64 Å) between the $DPP^{Th}-NN_2$ diradicals belonging to neighbouring stacks (Fig. 4b). These interactions lead to the shortest intermolecular distances between the nitroxide oxygen atoms, equal to 3.653 Å. On the whole, in the solid phase of $DPP^{Th}-NN_2$, the shortest $O \cdots O$ distances between O atoms of nitroxide groups exceed 3.617 Å, which is much greater than the sum of the van der Waals radii of O atoms (3.04 Å). Therefore, in the crystalline phase of DPPTh-NN₂ diradicals, both intra- and intermolecular exchange interactions should be weak.

DFT and ab initio calculations of the electronic structure and magnetic properties of the $DPPTh-NN₂$ diradical

To better understand the magnetic and spectroscopic properties of the $DPPTh-NN₂$ diradical, we performed a series of DFT and ab initio calculations. All calculations were performed for model geometries that differ from the geometry of the DPPTh- $NN₂$ by replacing the long *n*-hexyl substituents with methyl groups. In addition, two types of model geometry were used in the calculations: one was based on the XRD analysis, and the other was optimized in a toluene solution.

Calculations of intra- and intermolecular exchange interactions. Since the DPP^{Th} -NN₂ diradical has SOMOs of a disjoint type (Fig. S7, S8, ESI and discussion ibid†), the parameters of the intramolecular exchange interaction between the NN fragments were calculated at the SA-CASSCF/NEVPT2 level of theory.¹⁴ The largest active space used in these calculations consisted of 14 electrons on 13 MOs (Fig. S8, ESI†). For the model geometry based on XRD analysis, the parameter J was calculated to be -11.3 cm^{-1} (for molecule A) at the highest level of calculations. As expected, the BS-DFT calculations overestimated absolute value of J (−60, −53 and −79 cm⁻¹ with B3LYP, M06 and M06-2X functionals, respectively). Thus, the intramolecular exchange interaction in the diradical is indeed very weak and antiferromagnetic, and the ground state of the diradical is diamagnetic.

Fig. 4 (a) Short C–H \cdots O_{NO} contacts and the shortest O \cdots O distances between neighbouring DPPTh-NN₂ molecules in a stack and (b) between DPPTh-NN₂ molecules belonging to different stacks.

Fig. 5 Normalized electronic absorption spectra of the DPP^{Th} precursor (black curve) and DPPTh-NN2 diradical (red curve) measured in toluene at room temperature, as well as the calculated positions and oscillator strengths (f, right axis) of electronic transitions depicted as black bars for DPPTh and red bars for DPPTh-NN₂

We also calculated the parameters of intermolecular exchange interactions of neighboring radical fragments in the stack. Interestingly, for a pair of diradicals, J parameters of different sign (-5.6 and 6.4 cm^{-1}) were predicted for two pairs of adjacent radical fragments (Fig. S9, ESI†). An antiferromagnetic exchange interaction was predicted in the case when ONCNO fragments of neighboring radical fragments are almost parallel, and a ferromagnetic one, in the case of a significant deviation from parallel arrangement of the corresponding fragments.

Electronic absorption spectra of DPP^{Th} and $DPP^{Th}-NN_2$. Before proceeding to the calculation and analysis of exchange interactions in the excited states of the DPP^{Th} -NN₂ diradical, it is reasonable to interpret and analyze its electronic absorption spectrum and compare it with the spectrum of the parent DPPTh. Fig. 5 displays the electronic absorption spectra of both the DPP^{Th} and $DPP^{Th}~NN_2$ in toluene solution at room

temperature. The DPP^{Th} shows an intense absorption band in the visible region with two maxima; this pattern is characteristic of diaryl-substituted DPP core.^{2f,g} According to the TD-B3LYP calculation, this structured band corresponds to a single electronic excitation (507 nm, $f = 0.48$, Fig. 5), in which the electron promotion from the highest occupied molecular orbital (HOMO) to the lowest unoccupied molecular orbital (LUMO) dominates (Fig. 6a). The next transition with a much smaller intensity is predicted at 360 nm $(f = 0.0034)$. Thus, the maxima at 550 and 512 nm and the shoulder at about 480 nm represent the vibrational structure of the long wavelength band (with $\nu \sim$ 1350 cm−¹).

The electronic absorption spectrum of the $DPP^{Th}-NN_2$ diradical contains a very similar intense absorption band with the same structure as in the DPP^{Th} spectrum, but noticeably redshifted. Note that the DPPTh-NN₂ spectrum was calculated for the high-spin triplet state of the diradical. The wavefunction of the ground open-shell singlet cannot be correctly described in the one-determinant approximation, and thus its UV-Vis spectrum cannot be calculated at the TD-DFT level. Fig. S7 (ESI†) represents the MO diagram with a series of α - and β -type MOs involved in electronic excitations of the $DPPTh-NN₂$ diradical. It can be seen that highest occupied and lowest unoccupied MOs of both α - and β -types are similar to the HOMO and LUMO of the DPPTh (Fig. 6a) and differ from them only in some delocalization to NN fragments. According to the TD-UB3LYP calculation, the intense structured band in the spectrum of diradical also corresponds to a single intense transition (563 nm, $f = 0.96$, Fig. 5), which in turn corresponds to the HOMO \rightarrow LUMO promotions for α - and β -electrons (Fig. 6b). Since these HOMO and LUMO are similar to those of the parent $DPPTh$ (except for a slight delocalization), the intense long wavelength bands of the precursor and diradical have very similar shape and intensity, but the diradical band is redshifted. PSC Advances Fourier and the computer of the computation of the system is the common and the common attention is the common and the common attenti

In addition, the diradical spectrum shows a weak feature (around 680 nm) at the tail of the absorption band discussed above. According to the calculations, the $DPP^{Th}-NN_2$ has two electronic transitions in this region: at 672 nm $(f = 0.0037)$ and 673 nm ($f = \sim 10^{-8}$). In turn, the MO diagram (Fig. S7, ESI†) shows that two SOMOs (MO-166a and MO-167a) localized on the NN fragments are practically degenerate and lie below the

Fig. 6 Molecular orbitals (MO) involved in the long wavelength transition in the DPPTh spectrum (a), in the intense long wavelength transition in the DPPTh-NN₂ spectrum (b), and in the weak transition at 672 nm (c).

Fig. 7 Energy diagram of the six lowest energy states of the DPP^{Th} -NN2 diradical. The energies were calculated at the CASSCF(14,13)/ NEVPT2/def2-TZVP level for the optimized geometry. The splitting of the lowest excited quintet state is calculated at the CAS(10,10)/QDPT level.

HOMO by about 0.2 eV. The corresponding unoccupied MO-168 β and MO-169 β localized on the NN fragments (partners of the α -SOMOs) are also practically degenerate and lie above LUMO by about 0.2 eV. It is these MOs, along with HOMO and LUMO, that are involved in long wavelength low intensity transitions. Both transitions consist mainly of two contributions: the promotion of an a-electron from the SOMO to the LUMO and the promotion of a β -electron from the HOMO to the SOMO partner (Fig. 6c). These transitions are more complex than those previously calculated for a number of nitronyl nitroxide radicals.¹⁴

Analysis of the spin-Hamiltonian parameters in the excited states. In the excited states of the $DPPTh-NN₂$ diradical, a much stronger exchange interaction is expected. For example, the exchange interaction of the local excited triplet state of the DPPTh core with the doublet states of the NN fragments should lead to four excited states: a singlet, two triplets, and a quintet. Knowledge of the sequence and splitting of these multiplets, as well as zero-field splitting of triplet and quintet states, is very important for the interpretation of time-resolved and echodetected EPR spectra, and the phase relaxation kinetics in the presence and absence of laser radiation.

The results of the high-level calculations for the six lowenergy DPP^{Th} -NN₂ multiplets are presented in Fig. 7 and S10 (ESI†), as well as in Table 1. Calculations were carried out both for the optimized and XRD-based geometries. An analysis of the CASSCF wave functions demonstrates that the Q_1 state, as well as T2, T3, and S1 states, actually arose as a result of the exchange interaction of the local triplet state corresponding to the HOMO \rightarrow LUMO excitation (Fig. 6a) with NN radical fragments (for details, see ESI, Section 3†). The energy splitting between these multiplets corresponds to the exchange interaction of the local triplet state with doublets of radical fragments (with parameter J_1) and between radical fragments (with J_2), respectively (Table 1). Results of Table 1 show that the exchange interaction of the radical fragments is weak and antiferromagnetic for both the ground and triplet excited states. In turn, the exchange interaction between the local triplet state and NN fragment is strong and ferromagnetic. The latter can be explained by the McConnell I mechanism,¹⁷ as carbon atom of ON-C-NO fragment has high negative spin population and is bound with carbon atom of thiophen ring with high positive spin population (Fig. S11†). Paper
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For the parent DPP^{Th} , previous femtosecond transient absorption spectroscopy and time-correlated single-photon counting studies¹⁸ have demonstrated that the singlet excited S_1 state of this compound relaxes directly to the ground state via internal conversion and fluorescence, bypassing the triplet state. For the $DPP^{Th}-NN_2$ the energy diagram is much more complicated. Excitation of the diradical by the 532 nm laser

Table 1 Results of the CASSCF(14,13)/NEVPT2/def2-TZVP calculations of the low-energy spectrum of the DPPTh-NN₂, consisting two singlet, three triplet and one quintet states, as well as the parameters of corresponding exchange interactions^a and the zero-field splitting (ZFS) parameters (D, E/D)

 a Parameters J_1 and J_2 of the exchange interaction correspond to the spin Hamiltonian of the form $\hat{H} = -2J_1(\vec{S}_{\mathrm{DPP^{Th}}} \cdot \vec{S}_1 + \vec{S}_{\mathrm{DPP^{Th}}} \cdot \vec{S}_2) - 2J_2(\vec{S}_1 \cdot \vec{S}_2),$ where $\vec{S}_{\text{DPP}^{\text{Th}}}$ is the spin of the DPPTh core triplet state, and \vec{S}_1 and \vec{S}_2 correspond to the NN radical fragments. b ZFS parameters estimated at the CASSCF(10,10)/QDPT level. c ZFS parameters estimated at the RO-B3LYP level.

pulse leads to both singlet and triplet states arising from the exchange interactions of NN fragments through the excited singlet state localized mainly on the $DPPTh$ core. However, as was discussed previously, slightly below this pair of states, there are also states responsible for a weak feature in the UV-Vis spectrum (Fig. 5) (4 states in total - two triplets and two singlets). Finally, as shown by calculations, this sequence of excited states is closed by singlet, triplet, and quintet states presented in Fig. 7.

Thus, in the case of the $DPP^{Th}-NN_2$ diradical, between the state populated upon excitation and the ground state there are a large number of states, the energy gaps between which are small. In addition, among these states there are pairs of very close singlet and triplet levels that arise due to the exchange interaction of NN fragments. Therefore, one might expect a very fast conversion of the excited state (singlet or triplet), which occurs immediately after the excitation, into the Q_1 state. At room temperature, the Q_1 state is expected to be short-lived due to thermally activated relaxation through the T_2 state. However, such a process is absent at cryogenic temperatures, and one can expect a sufficiently long-lived Q_1 state at these temperatures.

The steady-state and echo-detected EPR spectra of the **DPPTh-NN₂.** Although the ground state of the DPPTh-NN₂ diradical is singlet and diamagnetic, its triplet state at room temperature is substantially populated (about 70% using predicted *J* value). Thus, the EPR spectrum of $DPPTh-NN₂$ in a toluene solution at ambient temperature (Fig. 8a) was recorded, and it is characteristic of bis(nitronyl-nitroxide) systems with an intramolecular exchange interaction significantly exceeding the hyperfine coupling $(|J| \gg A_N)$, which is also consistent with the results of our calculations ($|J| = 3.7 \times 10^3$ mT, Table 1). The spectrum has a center at $g_{\text{iso}} = 2.0065$ and contains nine lines due to the coupling of two unpaired electrons with four equivalent 14 N nuclei with $|A_N|/2$ ∼ 0.374 mT, which is experimental evidence of the diradical nature of $DPPTh-NN₂$.

The EPR spectrum of the DPP^{Th} -NN₂ diradical in a frozen toluene solution at 135 K (Fig. 8b) is difficult to interpret, since the symmetry is partly lost and some anisotropic components may overlap in $|\Delta m_S| = 1$ region. This results in a different number of shoulders in the outermost regions of the spectrum. Therefore, the zero-field splitting parameters can be estimated roughly from the low temperature EPR experiment. However, clear evidence of a diradical character comes from the observed $|\Delta m_{\rm s}| = 2$ transition. This signal has a low signal-to-noise ratio (S/N), because the average value of the $N \cdot \cdot N$, O $\cdot \cdot \cdot O$ and $N \cdot \cdot \cdot O$ distances is large and estimated at 15.3 Å (for both the XRD and optimized structures), which implies a small $|D|$ and, accordingly, a low probability of this transition. In the point dipole approximation, a distance of 15.3 Å leads to $|D| = 0.81$ mT (7.6 × 10⁻⁴ cm⁻¹). Note that this value cannot explain the rather large width of the EPR spectrum (Fig. 8b). However, the calculation of the spin–spin contribution at the DFT level gave an even smaller but close value: $D = -0.73$ mT (–6.8 \times 10⁻⁴ cm⁻¹) and $E/D = 0.014$. RSC Advances Continued and interferious article for the method interferious Continued Continued and the common and the proposition of the creation of the creation of the set of the common and the common and the set of the

Fig. 9a shows the echo-detected (ED) EPR spectra of the DPPTh-NN2 diradical recorded at 10 K under the same conditions in the presence and absence of pulse laser excitation. The spectrum recorded in the absence of irradiation features one slightly asymmetric EPR line, the maximum of which corresponds to $g =$ 2.008. This signal apparently corresponds to the lowest triplet state of the DPPTh-NN₂ diradical (T₁), and, according to calculations, its population at 10 K is low, although it cannot be ruled out that $E(T_1)$ may be substantially overestimated.

In the presence of laser radiation, the observed EPR signal retains its shape, but becomes less intense. Based on the quantum chemical calculations and discussion presented above, one should expect that the lowest excited quintet state (Q_1) may be populated upon excitation, thus being paramagnetic. However, still we do not observe any reliable spectral manifestations of such state. In addition, the transverse relaxation curves in the presence and absence of photoexcitation are

Fig. 8 Experimental (black) and simulated (red) continuous-wave (CW) X-band EPR spectra of DPPTh-NN₂ in toluene at room temperature (a) and in frozen toluene at 135 K (b) $(C \approx 0.1 \text{ mM})$

Fig. 9 Echo-detected EPR spectra (a) and phase relaxation kinetics (two-pulse echo decay vs. interpulse delay) (b) the DPPTh-NN₂ diradical in toluene glass at $T = 10$ K in the presence/absence of photoexcitation. Kinetic curves are normalized to the maximum.

remarkably similar (Fig. 9b), indicating that they refer to the same paramagnetic species.

We also applied time-resolved (TR) EPR technique to probe the intermediate species formed upon photoexcitation of $DPPTh-NN₂$ in toluene glass at 80 K. TR EPR is based on continuous wave detection and typically is more sensitive to photoexcited paramagnetic species compared to pulse EPR. However, no TR EPR signals were observed. In addition, the sample with pure $DPPTh$ moiety (without radicals attached) has been investigated with the same zero TR EPR signal. The failure to detect the TR EPR signal in the case of DPP^{Th} is consistent with the relaxation of the S_1 state directly to the ground state, as discussed above. The failure in the case of the diradical may be due to the short lifetimes of T_2 , T_3 and Q_1 states, if they are actually formed.

The observed decrease of the echo signal under light can be assigned to a slight heating of the sample. Indeed, an insignificant heating by a few degrees $(3-5 K)$ can noticeably modify EPR signal at 10 K, which is proportional to $1/T$ (see, *e.g.* ref. 19 and 20). At the same time, the opposite effect of heating should be present due to the higher population of T_1 state at higher temperatures. Apparently, the former mechanism dominates, causing spectral changes shown in Fig. 9a. Additional perturbation of resonator under laser radiation can also contribute to the observed signal decrease.

Conclusions

In summary, we have synthesized a DPP-derivative bearing two radical groups, i.e. bis(thiophen-2-yl)-2,5-dihydropyrrolo[3,4-c] pyrrole-1,4-dione terminally capped with nitronyl nitroxide groups ($DPP^{Th}-NN_2$). Notably, we used palladium-catalyzed cross-coupling reaction of the corresponding dibromoderivative with the (nitronyl nitroxide-2ide)(triphenylphosphine)gold complex as a convergent step to synthesize $DPP^{Th}-NN_2$ diradical. The incorporation of two

radicals resulted in interesting spectroscopic, photophysical, and magnetic properties, and the expected unusual excitedstate dynamics. A computational study predicted that the $DPPTh-NN₂$ diradical has singlet ground state with a small singlet–triplet energy gap ΔE_{ST} of about 0.2 kcal mol⁻¹ in solution. This value is consistent with strong on the EPR scale exchange interactions observed in the EPR. The inclusion of two NN radicals leads to an increase in the number of excited states both due to excitations involving orbitals localized on NN fragments and due to the exchange interaction of these fragments. Consequently, the density of states increases and, in turn, the energy gaps between them decrease. This effect, combined with the removal of spin prohibition for certain transitions, is expected to signicantly speed up the relaxation process of the excitation initially localized on the DPPTh core. Although we were unable to experimentally detect the lowenergy quintet state, likely due to its brief lifetime or low yield, future studies using ultrafast techniques, such as femtosecond spectroscopy, could potentially reveal its properties.

Experimental and computational details

Materials and instrumentation

All chemicals and reagents were purchased from commercial suppliers and used without further purification. Solvents used for spectroscopic measurements were spectral grade quality. 2,5-Dihydro-1,4-dioxo-3,6-dithienylpyrrolo[3,4-c]-pyrrole (2) was synthesized by adapting reported procedure. 21 All reactions were monitored by thin-layer chromatography (TLC) carried out on silica gel plates. Preparative separations were performed by column chromatography on silica gel grade 60 (0.040–0.063 mm) from Merck.

¹H and ¹³C NMR spectra were recorded on Bruker Avance 300 (300 MHz) spectrometer. The chemical shifts were reported in ppm and referenced to the residual solvent peak. $s = singlet$, $d =$ doublet, $t =$ triplet, $m =$ multiplet, $b =$ broad. Infrared spectroscopy measurements were conceded on Nicolet 730 FTIR spectrometer equipped with an attenuated total reflection (ATR) setup. The UV/Vis spectra were recorded at 298 K with a PerkinElmer Lambda 900 spectrophotometer. Matrix-assisted laser desorption ionization time-of-flight (MALDI-TOF) mass spectra were acquired with a Bruker Reflex II MALDI-TOF mass spectrometer, calibrated against a mixture of C_{60}/C_{70} . The X-ray crystallographic data for the molecules were collected on a STOE IPDS 2T diffractometer using Cu-K α IµS source. Crystal structure contains two independent molecules of NN_2 -DPP (A and B) both of them are located on an centre of inversion which result in a C_i symmetry. The alkyl chains are disordered. One solvent molecule (CH_2Cl_2) complete the unit cell.

Computational details

All quantum chemical calculations were performed for model structures that differ from the structures of the compounds under study by replacing the long n -butyl substituents with methyl groups. This substitution should only slightly affect the electronic and spectral properties of the parent $DPPTh$ and the DPPTh-NN₂ diradical.

To analyze magnetic properties of polycrystalline samples, all quantum chemical calculations were performed for the model geometry of $DPP^{Th}-NN_2$ diradicals and their pairs obtained from XRD analysis. The parameters of the intramolecular exchange interaction $(H = -2J\vec{S}_1\vec{S}_2)$ were computed using the accurate ab initio CASSCF²² and CASSCF/NEVPT2 (ref. 23 and 24) procedures. To calculate the J parameters for intermolecular exchange interactions, the spin-unrestricted brokensymmetry (BS) approach²⁵ at the BS-B3LYP/def2-TZVP level of theory²⁶⁻²⁸ using the Yamaguchi formula²⁹ was utilized

$$
J=-\frac{E^\mathrm{HS}-E^\mathrm{LS}_\mathrm{BS}}{\left\langle S^2 \right\rangle^\mathrm{HS}-\left\langle S^2 \right\rangle_\mathrm{BS}^\mathrm{LS}}
$$

:

To rationalize the results of experiments performed in solutions, calculations were carried out using the model geometry optimized in toluene solution at the UB3LYP/def2-TZVP and UM06-2X/def2-TZVP levels of theory;³⁰ the solvent was taken into account according to the CPCM model.³¹

The energies of electronic transitions and their oscillator strengths in the electronic absorption spectra of $DPPTh$ and $DPPTh-NN₂$ were calculated using the time-dependent $DFT³²$ at the TD-B3LYP/def2-TZVP level. In the case of $DPPTh-NN₂$, the calculations were performed for the high-spin triplet state of diradical. The zero-field splitting parameters (ZFS, D and E/D) for the lowest-energy triplet state of the $DPPTh-NN₂$ diradical was calculated at the RO-BP86/def2-TZVP level,^{33,34} while the g-tensor and hyperfine splitting tensors at the B1LYP/def2-TZVP level.³⁵

The energies of low-lying states (2 singlets, 3 triplets and 1 quintet) were also calculated at the CASSCF(14,13)/NEVPT2/ def2-TZVP level. Moreover, the splittings of the triplet and quintet multiplets (ZFS) due to the dipolar spin–spin interaction and spin–orbit coupling (SOC) were evaluated using CASSCF(10,10) wavefunctions and quasi-degenerate perturbation theory $(QDPT)$,³⁶ as realized in the CASSCF and MRCI modules of the ORCA 5.0.5 software package.³⁷ The contribution of SOC was calculated using the mean-field formalism $(SOMF(1X))$ ³⁸ and found to be negligible. To the best of our knowledge, this work is the first to provide high-level calculations of the low-energy spectrum of a dye substituted with two radical fragments.

The same software package was used for all other calculations. The molecular orbitals were visualized using the Chemcraft software³⁹

Synthetic methods and characterization

3,6-Di(thiophen-2-yl)-2,5-dihydropyrrolo[3,4-c]pyrrole-1,4 dione (2).²¹ To a 250 mL two-neck round bottom flask equipped with magnetic stirrer, potassium t -butoxide (6.5 g, 57.9 mmol), 2-thiophenecarbonitrile (1) $(5.0 \text{ g}, 45.9 \text{ mmol})$ and tert-amyl alcohol (30 mL) were added and the mixture was heated to 100 ° C under a nitrogen atmosphere. At this temperature, a solution of di-ethyl succinate (3.85 g, 22.1 mmol) and tert-amyl alcohol (5 mL) was added to the reaction mixture over 1 h using a dropping funnel. The reaction mixture was stirred at 100 °C for 20 h, and then cooled to room temperature, neutralized with glacial acetic acid (35 mL), and gently reflux temperature for 1 h. The resulting pigment suspension was suspended in water–methanol mixture $(1:1, 50 \text{ mL})$ and filtered to get pigment cake, which was washed with water–methanol mixture until no color found in washings. The crude compound was dried at 100 $\mathrm{^{\circ}C}$ in vacuo, and obtained 4.0 g of compound 2. The product was used in the next step without further purification. RSC Advances

in pays and referenced to the residual solvent peak, a singlet, CoSSCP, 0.13 Terms are determined on 19 February 10:01 PM. This contract the material one of SOC was elected unit the CoNTEXP and West Tend Wes

2,5-Dihexyl-3,6-di(thiophen-2-yl)-2,5-dihydropyrrolo[3,4-c] pyrrole-1,4-dione (DPPTh).^{40,41} To a 250 mL two-neck round bottom flask equipped with magnetic stirrer, compound 2 $(1.00 \text{ g}, 3.33 \text{ mmol})$ and anhydrous K_2CO_3 $(1.4 \text{ g}, 10.1 \text{ mmol})$ were added in 100 mL of anhydrous N,N-dimethylformamide (DMF). The reaction mixture was heated to 120 °C under nitrogen for 1 h. Alkyl bromide (8.0 mmol) was then added dropwise, and the reaction mixture was further stirred and heated 12 h at 130 °C. The reaction mixture was cooled to room temperature; then it was poured into 100 mL of water, and the resulting suspension was stirred at room temperature for 1 h. The solid was collected by vacuum filtration, washed with copious amount of water, followed by methanol, and then airdried. The crude product was checked on TLC and ¹H NMR to confirm the isomer presence and it is found to be around $45:55$ ratio based on ¹H NMR spectra. The crude mixture was purified on chromatography using 3% ethyl acetate in hexane as an eluent, and the solvent was removed under vacuum to obtain a pure product DPP $^{\rm Th}$ in 40% yield. $^{\rm 1}{\rm H}$ NMR (300 MHz, CDCl $_{\rm 3}$, ppm) $\delta = 8.92$ (d, 2H Th–H), 7.64 (d, 2H, Th–H), 7.27 (m, 2H, Th–H), 4.07 (d, 4H, N–CH₂), 1.74 (m, 4H, CH₂–CH₂), 1.41–1.28 (m, 12H), 0.87 (t, 12H). ¹³C NMR (125 MHz, CDCl₃, ppm) δ = 161.37, 140.02, 135.21, 130.63, 129.77, 128.58, 107.70, 42.21, 31.39, 29.89, 26.53, 22.53, 13.98. MALDI-TOF, Mw 468.9518. Anal. calcd for $C_{26}H_{32}N_2O_2S_2$: C, 66.63; H, 6.88; N, 5.98; S, 13.68. Found: C, 66.83; H, 7.01; N, 6.16; S, 13.42.

3,6-Bis(5-bromothiophen-2-yl)-2,5-dihexyl-2,5-dihy-

dropyrrolo[3,4-c]pyrrole-1,4-dione (DPPTh-Br₂). Compound DPPTh-Br was synthesized by following a modified procedure reported. 41 To a 100 mL two-neck round bottom flask, compound DPP^{Th} (0.60 g, 1.28 mmol) was added and dissolved in to dry DMF (10 mL) under nitrogen atmosphere. The reaction mixture was cooled down to 0 °C in ice bath and N-bromosuccinimide (NBS) (0.46 g, 2.6 mmol) was added portion wise. It was stirred at room temperature for 12 h. This mixture was poured into water (100 mL) and stirred for 1 h. Solid was vacuum filtered and washed with MeOH and $H₂O$ to remove excess NBS. The crude solid was purified through column chromatography using hexane : DCM $(1:1)$ as an eluent. Compound $DPPTh-Br$ was obtained as a dark purple solid (0.58 g, yield 83%). ¹H NMR (300 MHz, CD₂Cl₂, ppm) δ = 8.67 (d, 2H Th–H), 7.27 (d, 2H, Th–H), 3.98 (d, 4H, N–CH2), 1.64–1.74 (m, 4H), 1.30-8-1.40 (m, 12H), 0.89 (t, 6H). 13C NMR (75 MHz, CDCl₃, ppm) δ = 160.95, 139.01, 135.32, 131.62, 131.12, 119.10, 107.90, 42.30, 31.90, 29.95, 26.80, 22.63, 14.07. UV/Vis (toluene): λ_{max} : 550 nm. MALDI-TOF, Mw 626.47 (626.55). Anal. calcd for

 $C_{6}H_{30}Br_{2}N_{2}O_{2}S_{2}$: C, 49.85; H, 4.83; Br, 25.51; N, 4.47; O, 5.11; S, 10.24. Found: C, 49.76; H, 4.75; Br, 26.05; N, 4.55; S, 9.89. 3,6-

Bis(5-(4,4,5,5-tetramethyl-4,5-dihydro-3-oxid-1-oxyl-1H-imidazole-2-yl)thiophene-2yl)-2,5-dihexyl-2,5-dihydropyrrolo[3,4-c] pyrrole-1,4-dione (DPPTh-NN₂). A solution of dibromide Br₂-DPP (100 mg, 0.16 mmol), nitronyl nitroxide gold complex $(218 \text{ mg}, 0.35 \text{ mmol})$, and $[\text{Pd}(PPh_3)_4](20 \text{ mg})$ in THF (15 mL) was stirred at 65 °C in an argon atmosphere for 16 h. The reaction mixture was cooled to room temperature and solvent was evaporated. The residue was purified by column chromatography on silica gel by using $CH_2Cl_2/ethyl$ acetate $(10:1)$ solvent mixture as an eluent. The resulted product was recrystallized two times from a mixture of CH_2Cl_2 with MeOH to obtain diradical NN₂-DPP (90 mg, 72 %). Compound DPPTh-NN₂ was eluted with $R_f = 0.7$ (10 : 1 mixture of CH₂Cl₂/ethyl acetate); IR: ν = 2956, 2927, 2856, 1662, 1552, 1436, 1400, 1367, 1367, 1211, 1176, 1134, 1095, 870, 827, 729, 620, 540 cm^{-1} ; UV/Vis (toluene): λ_{max} : two maxima at 573 and 624 nm and a shoulder at ∼680 nm; MALDI-TOF MS: m/z calcd. (%) for $C_{52}H_{76}N_4O_4S_4$: 778.35 (100) $[M]^+$, 779.36 (43) [M + 1]⁺; found: 778 (average $M_{\rm W}$); elemental analysis calcd: C 61.67, H

6.99, N 10.79, S 8.23; found: C 61.80, H 7.1, N 10.3, S 8.05. Melting point, 216 °C.

Single-crystal XRD analysis

Single crystal X-ray diffraction data for $DPP^{Th}-NN_2$ were collected at 193 K on a STOE IPDS 2T diffractometer with Cu-K_a $I\mu S$ mirror system (Table 2). The structure was solved using direct methods, expanded with Fourier techniques and refined with the SHELXT software package. All non-hydrogen atoms were refined anisotropically. Hydrogen atoms were included in the structure factor calculation on geometrically idealized positions. Crystallographic data have been deposited with the Cambridge Crystallographic Data Centre as supplementary publication no. CCDC 1908216.†

Photoinduced EPR experiments

Pulse EPR experiments were conducted at Bruker Elexsys E580 spectrometer at the Center of Collective Use ''Mass spectrometric investigations'' SB RAS at X-band (9 GHz). Spectrometer is equipped with flow helium cryostat (Oxford Instruments) and thermocontroller Lakeshore and allows measurements at $T = 4$ 300 K. In order to measure echo-detected (ED) EPR spectra and transverse (phase) relaxation times (T_2) we used standard twopulse sequence with the pulse lengths being 10 and 20 ns for $\pi/2$ and π pulses, respectively. In case of T_2 measurements, interpulse delay was incremented. Time-Resolved (TR) EPR experiments were performed using homemade spectrometer based on commercial Bruker EMX microwave bridge. **PSC Advances**
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In all cases the studied compounds were dissolved in toluene in concentrations ca. 0.5 mM and placed into quartz sample tubes with outer diameter of 2.8 mm. The samples were degassed by a few freeze–pump–thaw cycles and then sealed off in vacuo. EPR studies were performed for two samples: DPP^{Th} - NN_2 biradical and its photosensitive moiety DPPTh free of radical fragments.

Laser irradiation was provided by Nd:YaG LOTIS-TII system at 532 nm, power ∼ 20 mJ per pulse, 10 Hz repetition rate. In case of pulse EPR detection laser was synchronized with EPR spectrometer; laser pulse proceeded the first microwave pulse by 700 ns. In TR EPR experiments the detection was also synchronous with laser pulsing; microwave absorption was detected starting from 100 ns till 9 μ s after the laser pulse.

Conflicts of interest

There are no conflicts to declare.

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