Spin Density Distribution in a Nitroxide Biradical Containing ¹³C-Enriched Acetylene Groups in the Bridge: DFT Calculations and EPR Investigation

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Abstract A specially synthesized nitroxide biradical $R_6^{-13}C \equiv C-p-C_6H_4-C \equiv {}^{13}C-R_6$, **B3**, where $R_6 = 1$ -oxyl-2,2,6,6-tetramethyl-1,2,3,6-tetrahydropyridine nitroxide, has been studied by electron paramagnetic resonance spectroscopy, and electron-nuclear double resonance (ENDOR). Spin density distribution and hyperfine splitting (hfs) constant on ${}^{13}C$ atoms calculations for biradical **B3** were carried out using B3LYP and PBE0 functionals and several different basis sets including N07 family and were compared with the experimental value of the hfs constant on ${}^{13}C$ atoms, measured from ENDOR spectra of **B3**. The mechanism of the intramolecular electron spin exchange in **B3** biradical is discussed.

1 Introduction

Scientists all over the world have been intrigued by the chemistry of stable free radicals since their discovery in 1900 [1]. Stable radicals of different types are presently used in a variety of fields [2], including spin labeling [3–12], the construction of organic magnetic materials [13], MRI contrast agents [14], redoxactive components of organic radical batteries [15], co-oxidants [16] and tools for

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controlled radical polymerization [17]. Some biradicals were found to be promising polarizing agents for dynamic nuclear polarization [18, 19]. Nevertheless, in many cases, biradical or polyradical structure and features, their magnetic properties are still the subject of investigation [20, 21].

Among numerous nitroxide biradicals, those containing acetylene groups in the bridge connecting two nitroxide rings present an interesting specific group [22–31]. In these biradicals, which are structurally rigid enough, spin exchange coupling between the unpaired electrons is realized by the indirect mechanism via the bridge of atoms and bonds connecting two paramagnetic nitroxide rings [27, 30–32]. The intramolecular dynamics and temperature behavior in such biradicals dissolved in various molecular and ionic liquid solvents is usually described well with a two-conformational model [32, 33]. In such biradicals, the intramolecular spin exchange coupling occurs by the indirect mechanism through the bridge of atoms and bonds.

It has been recently shown that such polyacetylene biradicals as $R_6-C \equiv C-R_6$ (B1), $R_6-C \equiv C-C \equiv C-R_6$ (**B2**), $R_6-C \equiv C-p-C_6H_4-C \equiv C-R_6$ (**B3**), where R_6 is 1-oxyl-2,2,6,6tetramethyl-1,2,3,6-tetrahydropyridine nitroxide [30, 31] with acetylene groups in the bridge are not "rigid" but "flexible", and biradical B2, due to fast rotations around its molecular axis, demonstrates average values of the exchange integral |J|. Biradical B3 is characterized by a higher rotation barrier in vacuum and is, therefore, more rigid. It was concluded from electron paramagnetic resonance (EPR) data that both biradicals exist in at least two different conformers with a slight influence of temperature on the |J/a| value. It decreases at high temperature, when faster rotation averages the configurations more effectively [30]. A short nitroxide biradical, **B1**, has been investigated by EPR spectroscopy supplemented with a single-crystal X-ray analysis and density functional theory (DFT) calculations [31]. Structural and EPR data were complemented with quantum chemical calculations, and conformational transitions in biradical B1 were also in a good agreement with quantum chemical computation. Nevertheless, understanding that in such chemically rigid polyacetylene nitroxide biradicals, intramolecular spin exchange can be realized only by the indirect mechanism, we could not measure the spin density distribution on atoms forming the bridge in biradicals and describe quantitatively spin delocalization and its influence on the |J| value.

In this work, we present our results on studying the spin density delocalization and the mechanism of the intramolecular spin exchange in the nitroxide biradical R_6 -¹³C \equiv C-*p*-C₆H₄-C \equiv ¹³C-R₆, **B3**, in which two carbon atoms are substituted by ¹³C isotopes (I = 1/2). Spin coupling in **B3** can be realized only via the bridge of atoms and bonds connecting two 1-oxyl-2,2,6,6-tetramethyl-1,2,3,6-tetrahydropyridine nitroxides, and it was investigated using EPR, electron-nuclear double resonance (ENDOR) spectroscopy as well as quantum chemical calculations.

2 Experimental

2.1 Synthesis of B3 Biradical

Methyl-¹³C-triphenylphosphonium iodide in anhydrous THF suspension was treated with butyl lithium solution. To this brown-colored solution was added the

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terephthalaldehyde (1), and after work-up, we got the 1,4-divinylbenzene, of which bromination with 2.2 equiv. Br₂ in CHCl₃ gave the tetrabromo product. Elimination with 6 equiv. KOt-Bu in THF generated the isotopically labeled 1,4-diethynyl benzene (2). The Sonogashira coupling with 4-iodo-2,2,6,6-tetramethyl-1,2,3,6tetrahydropyridin-1-yloxyl radical [36] in Et₃N and in the presence of CuI and Pd(PPh₃)₂Cl₂ biradical (3) was generated. According to our experience this synthetic methodology resulted no Glaser-coupled product formation was observed (while paramagnetic acetylenes reaction with 1,4-diiodobenzene generated two types of biradicals) (Fig. 1).

Reagents and conditions were: (i) 13 CH₃P(Ph)₃I (2.18 eq.), BuLi (2.18 eq.), THF, 2 h, 0 °C, then -78 °C, terephthalaldehyde (1.0 eq.) 15 min, rt, 3 h, 83 %, then CHCl₃, Br₂ (2.2 eq.), rt, 12 h, then KO*t*-Bu (6.0 eq.), THF, reflux, 3 h, 60 %. (ii) 4-iodo-2,2,6,6-tetramethyl-1,2,3,6-tetrahydropyridin-1-yloxyl radical (2.1 eq.), Et₃N, CuI (0.1 eq.), Pd(PPh₃)₂Cl₂ (0.06 eq.), 50 °C, 20 h, 25 %. Here, rt denotes room temperature.

Melting points were determined with a Boetius micro melting point apparatus and are uncorrected. Elemental analyses (C, H, N) were performed on Fisons EA 1110 CHNS elemental analyzer. Mass spectra were recorded on a Thermoquest Automass Multi. ¹H NMR and ¹³C NMR spectra were recorded with Bruker Avance 3 Ascend 500 spectrometer. Chemical shifts are referenced to Me₄Si. The paramagnetic compound was reduced with 5 equiv. pentafluorophenyl hydrazine/ radical. Measurements were run at 298 K probe temperature in CDCl₃ or CD₃OD solution. Flash column chromatography was performed on Merck Kieselgel 60 (0.040–0.063 mm). Qualitative TLC was carried out on commercially available plates ($20 \times 20 \times 0.02$ cm) coated with Merck Kieselgel GF254. It is essential that anhydr. THF can be used only, Et₃N was distilled from CaH₂ prior to use. Terephthalaldehyde and methyl-¹³C-triphenylphosphonium iodide and butyl lithium were purchased from Aldrich. 4-Iodo-2,2,6,6-tetramethyl-5,6-dihydropyridin-1yloxyl radical [36] was prepared according to published procedures, other reagents were purchased from Aldrich or Alfa Aesar.

1,4-Diethynylbenzene (2): methyl-¹³C-triphenylphosphonium iodide (4.86 g, 12 mmol) was suspended in dry THF (50 ml), cooled to 0 °C, and *n*-BuLi (4.8 ml, 12.0 mmol, 2.5 mol/l solution in hexane) was added. It was stirred for 2 h at the same temperature, then cooled to -78 °C, and the terephthalaldehyde (737 mg, 5.5 mmol) as solution in anhydrous THF was added dropwise. After



Fig. 1 Scheme of the synthesis of biradical B3

15 min of stirring at -78 °C, the mixture was warmed to room temperature and stirred for another 3 h. After stopping the reaction with water (50 ml), it was extracted with Et₂O (3 × 20 ml), and the combined organic phases were dried over MgSO₄, the pure olefin was obtained by column chromatography to give 1,4-divinylbenzene (600 mg, 83 %). The 1,4-divinylbenzene was dissolved in CHCl₃ (20 ml); to this solution, Br₂ (1.60 g, 10.0 mmol) was added, and the mixture was allowed to stay overnight. The solvent and bromine were evaporated off in vacuo, the residue was dissolved in THF (30 ml), then KOt-Bu (27.6 mmol, 3.10 g) was added in one portion, and the mixture was stirred and refluxed for 3 h. After cooling, the solvent was evaporated off, and the residue was partitioned between Et₂O (30 ml) and 5 % aq. H₂SO₄ (10 ml). The organic phase was separated, dried (MgSO₄), filtered and evaporated, and the residue was purified by flash column chromatography with (hexane/Et₂O, 100:1) to give a brownish-white solid (350 mg, 60 %), mp 93–94 °C, R_f 0.37 (hexane), MS (70 eV): m/z = 128 (M⁺, 100), 101 (6) 76 (33). Anal calculated for C₁⁸³C₂H₆: C 95.28; H 4.72, found C 95.20; H 4.59.

4,4'-(1,4-Phenylenebis(ethyne-2,1-diyl))bis(2,2,6,6-tetramethyl-1,2,3,6-tetrahy-dropyridin-1-yloxyl) biradical **B3.**

To a degassed solution of 4-iodo-2,2,6,6-tetramethyl-1,2,3,6-tetrahydropyridin-1yloxyl radical (1.40 g, 5.0 mmol) in anhydrous Et₃N (10 ml), CuI (47 mg, 0.25 mmol) and PdCl₂(PPh₃)₂ (112 mg, 0.16 mmol) were added, and the mixture was stirred for 15 min, followed by the addition of ¹³C containing 1,4-diethynylbenzene (300 mg, 2.34 mmol) and was stirred for 20 h at 50 °C under Ar, in a sealed tube. The reaction mixture was diluted with CHCl₃ (10 ml), filtered through Celite, and the solvent was evaporated. The residue was partitioned between CHCl₃ (30 ml) and aq. sat. NaCl solution (10 ml), the organic phase was separated, washed with 10 % aq. Na₂S₂O₃ (10 ml), water (10 ml), dried (MgSO₄) and to this mixture, activated MnO₂ (172 mg, 2.0 mmol) was added, and the mixture was set aside for overnight. Next, the mixture was filtered, evaporated and purified by flash column chromatography (hexane/CHCl₃) to give the title compound (254 mg, 25 %) as a deep-yellow solid, mp 233–235 °C, R_f 0.66 (hexane/EtOAc). MS (70 eV): *m/* z = 432 (M⁺, 42), 402 (29), 372(18), 357 (16), 257 (22), 42 (100). Anal calculated for C₂¹³₆C₂H₃₄N₂O₂: C 78.20; H 7.92; N 6.48; found C 78.15; H 7.80; N 6.42.

2.2 EPR Measurements

Toluene was selected as a solvent and was carefully purified as per the literature procedure [37]. Solutions were prepared, bubbled with nitrogen for 20–25 min. 0.5 ml of the solution was taken into a thin capillary and degassed by frieze-pump circle to remove oxygen, and sealed off under vacuum. Radical concentrations were sufficiently low ($\leq 4 \times 10^{-4} \text{ mol } 1^{-1}$) to eliminate intermolecular exchange broadening of EPR lines [38].

EPR spectra were recorded at X-band on a Bruker EMX-8 spectrometer with a modulation frequency of 100 kHz. Temperatures were controlled with an accuracy of ± 0.5 °C at temperatures between -20 and 75 °C by means of a JEOL JNM-VT-30 temperature control system. EPR parameters: the hyperfine splitting, hfs, constant on nitrogen ¹⁴N atom, *a*, and a value of the exchange integral |J/a|, which is

sensitive to any changes of the spin density distribution of the unpaired electron in the system. The exchange integral values |J/a| were calculated in accordance with [32, 39]. EPR spectra of biradicals were simulated with the computer program package created by Dr. A. A. Shubin (Boreskov Institute of Catalysis, Siberian Branch, Russian Academy of Sciences) and described in details in Ref. [39]. For our calculations, we used spin-Hamiltonian parameters of similar nitroxide radicals collected in [40].

In liquid solutions with rather low viscosity, the spin Hamiltonian \hat{H} comprises the isotropic hyperfine, the Zeeman interactions and the exchange coupling. In the case when both radical fragments are identical and one nucleus with a nonzero nuclear spin *I*, the following equation is valid [34, 35]:

$$\hat{H} = g\beta_{\rm e}H_0\left(S_z^{(1)} + S_z^{(2)}\right) + a\left(S_z^{(1)}I_z^{(1)} + S_z^{(2)}I_z^{(2)}\right) + JS^{(1)}S^{(2)}$$
(1)

The spin Hamiltonian is, here, written in frequency units; superscripts 1 and 2 denote different radical fragments; $S^{(k)}$ are electron spin operators; $S_z^{(k)}$ and $I_z^{(m)}$ are projections of the electron and nuclear spins to the Z-axis, respectively; g is the isotropic g-factor of the radical fragments; β_e is the Bohr magneton; H_0 is the external magnetic field; $a \approx 3 \times 10^8$ rad/s denotes the ¹⁴N isotropic hyperfine splitting (hfs) constant, and J is the exchange integral. In low-viscous solvents, the tensor of the dipole–dipole coupling is averaged to zero by fast rotation of the biradical molecule [32, 33]. For any individual conformation, the one value of |J| should correctly describe the position and integral intensities of all lines in the EPR spectrum. Values of |J| are usually measured in units of the hfs constant a, i.e., as |J/a|, with an accuracy of $\pm 2-3 \%$.

For the two-conformational model [32, 34, 35], the Arrhenius plot of |J/a| should be linear for any biradical, and this allows determining the differences in enthalpies, ΔH , and entropies, ΔS , of these two conformations:

$$\ln|J/a| = \Delta S/R - \Delta H/RT, \qquad (2)$$

The experimental observation of such plot confirms the validity of the twoconformational model for a biradical.

2.3 ENDOR Measurements

For measuring hyperfine splitting (hfs) constants of the interaction between the unpaired electron and ¹³C nucleus, the ENDOR method suggested by Mims has been used [41]. In this method, three microwave pulses are used, which form the signal of the stimulated echo. Between the second and the third microwave pulses (mw pulses), the radio-frequency pulse (rf pulse) is applied. The signal of the stimulated echo is recorded as a function of the rf pulse frequency (Fig. 2).

Measurements were carried out using a pulsed EPR spectrometer Elexsys E580 at X-band (10 GHz). A spectrometer is equipped with a standard commercial resonator EN 4118X-MD4W1, which was placed into cryostat CF935. Temperature control was carried out by the ITC503 unit. All measurements were done at 80 K. Two biradicals, **B3** and its analog **B3a** with a usual isotope ¹²C (I = 0), were





investigated. Both biradicals were dissolved in toluene at ~1 mmol/l concentration and degassed by pumping. ENDOR study was performed using 16-, 48- and 60-ns pulses as mw $\pi/2$ -pulse and 10 µs as rf one. It should be noted that no difference in ENDOR spectra using different lengths of the mw pulses was observed. Interpulse time τ was equal to 600 ns.

2.4 Calculation Details

Calculations were performed with ORCA 3.0.3 program package [42]. **B3** geometry was optimized on UKS/B3LYP/cc-pVTZ level and is in good agreement with previously reported results. Hfs constants were calculated using density functional theory with B3LYP and PBE0 functionals and variety of full electron basis sets such as EPR-II [43], cc-pVTZ, aug-cc-pVTZ [44], N07D and N07T family (including diffuse functions) [45]. Fine Lebedev 770 angular grid and 10^{-10} Eh SCF convergence tolerance were used. Solvent effects were simulated with COSMO model [46].

3 Results and Discussion

3.1 EPR Spectroscopy

Typical EPR spectrum of biradical **B3** at 298 K is shown in Fig. 3. The comparison of simulated and experimental spectra shows that they are in a very good agreement, and positions of all "exchange" lines in the magnetic field are practically coinciding (Fig. 3). Similar EPR spectra were described in Refs. [25, 30].

The exchange integral value I/a measured from spectra simulation at 298 K is equal to 3.4 ± 0.1 mT (a = 15.02 G), which coincides with those published in Refs. [25, 30]. One can see from Fig. 3 that amplitudes of the central and high-field simulated main "radical" lines are bigger than those of experimental ones. This is caused by the "stick" shape of the **B3** molecule, which is realized in the anisotropic rotation of **B3** performed in the experimental spectrum, while spectrum simulation



Fig. 3 Experimental (*solid line*) and calculated (*open circles*) EPR spectra of biradical B3 in toluene at 298 K

has been done for the case of the isotropic rotation [39]. Positions of all the lines for both spectra are coinciding.

Temperature dependences both of a and |J/a| values for **B3** dissolved in toluene slightly decrease with the increase of temperature and coincide in the error limits with reported [30]: $\Delta H = -0.5 \pm 0.2 \text{ kJ mol}^{-1}$ those in and $\Delta S = 6.8 \pm 0.7 \text{ J mol}^{-1} \text{ K}^{-1}$. Such small values were already explained in Ref. [30]: the barrier of rotation, E_a , of piperidine rings in biradicals with acetylene groups in the bridge is rather low as shown by results from the DFT calculations: it does not exceed 8.0 kJ mol⁻¹ for **B3**. Librations of the piperidine planes, relatively the p-C₆H₄ plane by up to 5°, have a value of E_a less than 0.1 kJ mol⁻¹, i.e., negligible. These results correlate well with the experimentally measured values of ΔH , which are equal to -0.5 kJ mol⁻¹ and close to -1.0 kJ mol⁻¹ in the case of B3 dissolved in toluene and ethanol, respectively. It should also be mentioned that such small values of E_a for biradical **B3** result in rather free intramolecular rotation of the nitroxide rings even at low temperatures, i.e., the EPR spectra and measured values of |J/a| characterize not individual conformations of **B3** but an averaged pattern with very fast transitions between several rotamers, and the measured value of |J/a| is averaged by all these conformations.

3.2 ENDOR Investigation

ENDOR spectra of biradicals **B3** and **B3a** are shown in Fig. 4. Signals of both samples are observed in the area of 3 and 5 MHz (and the same lines or two biradicals at the frequency region about 8–20 MHz, data are not shown), and we suppose that they reflect the interaction of the unpaired electron with a nuclear spin of nitrogen ¹⁴N. Besides these signals, in the ENDOR spectrum of **B3** enriched with ¹³C, two additional lines are observed in the area of 3.5-4.5 MHz. Subtracting the



Fig. 4 ENDOR spectra of biradicals B3a (a) and B3 (b). The spectrum (c) is a subtraction of a from b. The estimated value of the hfs constant with ¹³C is equal to ~ 0.48 MHz

spectrum of the usual biradical **B3a** from the spectrum of **B3** results in the signal (Fig. 4c), which contains only two peaks with a center at 3.7 MHz, what corresponds to a Larmor frequency of the carbon nucleus precession in the magnetic field, close to ~ 340 mT. The splitting between peaks is equal to 0.48 ± 0.14 MHz. These results clearly indicate that we observe a rather weak interaction between the unpaired electron and the ¹³C carbon atom. The hfs constant value in such case is equal to the splitting, i.e., to 0.48 MHz. It should be noted that we could not determine all values of the anisotropic hfs tensor from our study of biradical **B3** in frozen toluene solutions.

3.3 DFT Calculations

Important information about **B3** structure and features has been obtained from quantum chemical calculations. Calculated geometry of biradical **B3** is shown in Fig. 5, where the appropriate values of bond lengths, ¹³C and other isotropic hfs constants on different atoms in **B3** are listed for two cases of calculation methods: B3LYP/N07D and PBE0/N07D. The whole set of ¹³C isotropic hfs constant values calculated using B3LYP and PBE0 functionals at various basis sets is given in Table 1. The distance between centers of N–O bonds is equal to 17.78 Å.

It should be noted that while calculations of the dipolar components of the hfs tensor can be systematically tuned by going to a much larger basis set of common use, isotropic hfs constants are usually reproduced better using a specialized basis set described in [45]. According to this note, we should point out two important



Fig. 5 Calculated bond lengths and ¹³C isotropic hfs constants in biradical **B3**. *Numbers 1* and 2 indicate the method of calculation of ¹³C isotropic hfs constants: B3LYP/N07D and PBE0/N07D, respectively

Table 1 13 C isotropic hfs constant values calculated using B3LYP and PBE0 functionals at various basis sets

<i>a</i> (¹³ C), MHz
0.39
0.38
0.39
0.33
0.35
0.38
0.52
0.41
0.43
0.46
0.48

results: first, specialized basis sets, such as N07 family or EPR family, enable much better agreement with the experimental data than the conventional basis sets on far less computational prices even in this case of very small hfs coupling. Second, we point out that, for some reasons, PBE0 functional systematically gives better agreement of calculated ¹³C isotropic hfs constant with the experimental one in this particular case, while B3LYP functional, in turn, underestimates it, though both approaches yield results which are within the experimental error limits and agree well with the experimental value. We conclude that combination of PBE0/N07D or PBE0/EPR-II can be recommended as a cheap and precise way for computation of small hfs couplings in nitroxide biradicals. A more detailed comparison of the experimental results obtained in different solvents with calculated ones taking into account various polarities, we intend to do in the next paper.

4 Conclusion

A nitroxide biradical $R_6^{-13}C \equiv C$ -*p*- C_6H_4 - $C \equiv {}^{13}C$ - R_6 , **B3**, where $R_6 = 1$ -oxyl-2,2,6,6-tetramethyl–1,2,3,6-tetrahydropyridine nitroxide, has been investigated by X-band EPR and electron-nuclear double resonance (ENDOR) spectroscopy. Spin density distribution and hyperfine splitting (hfs) constant value on ${}^{13}C$ atoms in the bridge in biradical **B3** were calculated using B3LYP and PBE0 functionals and several different basis sets including N07 family. These results were compared with the experimental value of the hfs constant on ${}^{13}C$ atoms measured from ENDOR spectra of **B3**. A few recommendations concerning the calculation of electron spin density distribution and the isotropic hfs constant values in nitroxide biradicals are given. Results presented in this paper confirm the fact that intramolecular electron spin exchange in **B3** biradical is realized by the indirect mechanism.

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