

# Electronic Supplementary Information

for

## Photochemical synthesis of carbazole-fused Blatter radicals: Effective spin injection to the carbazole system

Paulina Bartos,<sup>a\*</sup> Patrycja Szamweber,<sup>b</sup> Bruno Camargo,<sup>c</sup> Anna Pietrzak,<sup>d</sup> Piotr Kaszyński<sup>a,b,e\*</sup>

<sup>a</sup> Faculty of Chemistry, University of Łódź, 91-403 Łódź, Poland

<sup>b</sup> Centre of Molecular and Macromolecular Studies, Polish Academy of Sciences, 90-363 Łódź, Poland

<sup>c</sup> Institute of Experimental Physics, Faculty of Physics, University of Warsaw, 02-093 Warsaw, Poland

<sup>d</sup> Faculty of Chemistry, Łódź University of Technology, 90-924 Łódź, Poland

<sup>e</sup> Department of Chemistry, Middle Tennessee State University, Murfreesboro, TN, 37132

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## 1. Synthetic details

**General.** Reagents and solvents were obtained commercially. For separation of radicals, silica gel 60 (35-70 micron) passivated with Et<sub>3</sub>N or high-purity grade silica gel (w/ Ca, ~0.1%; purchased from Sigma-Aldrich, product no. 60752-1KG) were used. NMR spectra were obtained at 600 and 400 MHz (<sup>1</sup>H), and 151 and 101 MHz (<sup>13</sup>C) in CDCl<sub>3</sub> and referenced to the solvent ( $\delta = 7.26$  ppm for <sup>1</sup>H and  $\delta = 77.16$  ppm for <sup>13</sup>C).<sup>1</sup> IR spectra were recorded using Nexus FT-IR Thermo Nicolet IR spectrometer in KBr tablets or an Agilent Cary 630 FTIR spectrometer, in neat. UV spectra were measured in CH<sub>2</sub>Cl<sub>2</sub> on PerkinElmer Lambda 45 spectrophotometer. Melting points were determined on a Stuart SMP30 Advanced Digital Melting Point Apparatus and are uncorrected. High-resolution mass spectrometry (HRMS) measurements were performed using SYNAPT G2-Si High Resolution Mass Spectrometry equipped with an ESI or APCI source and Quantitative Time-of-Flight (QuanTof) mass analyzer. Irradiations were conducted with a 300 W halogen lamp ("Portable halogen Work Lamp" without the protecting front glass window) equipped with a T3 double-ended RSC base J118 light bulb.

### Photochemical preparation of radicals 1. General procedure.

A solution of precursor **2** (0.10 mmol, see the ESI for synthesis) in CH<sub>2</sub>Cl<sub>2</sub>, EtOAc or EtOH (100 mL) was placed in a 250 mL borosilicate round bottom flask fitted with a reflux condenser. The solution was stirred and irradiated with a 300 W halogen lamp, which was set about 30 cm from the flask. The reaction mixture was allowed to warm up to 30–35°C. Progress of the reaction was monitored by TLC (20% EtOAc/petroleum ether) and the irradiation was stopped after 72 h. The solvent was evaporated, the residue was adsorbed onto silica gel (w/Ca content, ~0.1%) and the products were separated from unreacted starting **2** (if present) using a short chromatographic column (silica gel with "Ca", ~0.1%) using EtOAc in pet. ether gradient 5–10%. The obtained radical was recrystallized.

*6-Phenyl-7H-indolo[3,2,1-de][1,2,4]triazino[5,6,1-kl]phenazin-7-yl (1c).* Obtained from **2c** (37.2 mg, 0.1 mmol) in CH<sub>2</sub>Cl<sub>2</sub>: 14.1 mg (37% yield); in EtOAc: 18.4 mg (45% yield); in EtOH; 3.4 mg (9% yield); black brown microcrystals: mp 264–266 °C (hexane/ EtOAc 4:1); IR (KBr)  $\nu$  1587, 1483, 1449, 1394, 1353, 1340, 1322, 771, 736, 688 cm<sup>-1</sup>; UV (CH<sub>2</sub>Cl<sub>2</sub>)  $\lambda_{\max}$  (log  $\epsilon$ ) 852.0 sh (2.63), 763.0 (3.48), 693.5 (3.44), 636.0 (3.19), 531.0 (3.31), 450.0 (3.50), 355.0 (4.00), 285.5 (4.35), 244.0 (4.56) nm; HRMS

(ESI-TOF)  $[M+H]^+$   $m/z$  calcd for  $C_{25}H_{16}N_3$ : 372.1375; found: 372.1360. Anal. Calcd for  $C_{25}H_{15}N_4$ : C, 80.84; H, 4.07; N, 15.08. Found: C, 80.82; H, 4.08; N, 15.14%.

*1-Oxo-6-phenyl-1H-indolo[3,2,1-de][1,2,4]triazino[5,6,1-kl]phenazin-4-ium-5-ide (1c-oxo)*. Obtained as a highly polar byproduct (AcOEt eluent) in photocyclization of **2c** (37.2 mg, 0.1 mmol) in  $CH_2Cl_2$ : 5.6 mg (12% yield); in EtOAc: 5.0 mg (13% yield); dark blue micro-crystals: mp 268–270 °C;  $^1H$  NMR (400 MHz,  $CDCl_3$ )  $\delta$  8.54 (d,  $J$  = 8.1 Hz, 1H), 8.38–8.35 (m, 2H), 8.24 (d,  $J$  = 9.9 Hz, 1H), 8.16 (d,  $J$  = 6.8 Hz, 1H), 7.91–7.83 (m, 2H), 7.59–7.52 (m, 5H), 7.44 (d,  $J$  = 7.7 Hz, 1H), 6.82 (d,  $J$  = 10.1 Hz, 1H);  $^{13}C\{^1H\}$  NMR (101 MHz,  $CDCl_3$ )  $\delta$  179.7, 161.5, 148.3, 134.9, 134.7, 133.1, 132.2, 131.8, 130.4, 128.7, 128.4, 128.0, 126.0, 125.5, 123.2, 120.7, 119.9, 119.1, 113.9, 111.2, 108.7; IR (neat)  $\nu$  1682, 1571, 1536, 1506, 1491, 1435, 1380, 1330, 1179, 1125, 1043, 971, 781, 746, 693  $cm^{-1}$ ; UV ( $CH_2Cl_2$ )  $\lambda_{max}$  (log  $\epsilon$ ) 702.0 (2.76), 612.0 (3.96), 566.0 (3.85), 526.5 (3.57), 312.5 (4.12), 276.0 (4.20) nm; HRMS (AP-TOF)  $m/z$   $[M+H]^+$  Calcd for  $C_{25}H_{15}N_4O$ : 387.1246; found: 387.1253. Anal. Calcd for  $C_{25}H_{14}N_4O$ : C, 77.71; H, 3.65; N, 14.50; O, 4.14. Found: C, 77.42; H, 3.78; N, 14.61%.

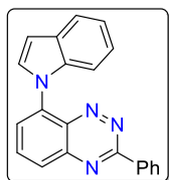
*2,14-Di-tert-butyl-6-phenyl-7H-indolo[3,2,1-de][1,2,4]triazino[5,6,1-kl]phenazin-7-yl (1d)*. Obtained from **2d** (48.5 mg, 0.1 mmol) in  $CH_2Cl_2$ : 1.1 mg (~2% yield); in EtOAc: 33.9 mg (70% yield); in EtOH: 0.5 mg (~1% yield); black brown microcrystals: mp 294.0–295.0 °C (MeCN/ $CH_2Cl_2$ ); IR (neat)  $\nu$  2954, 1578, 1487, 1444, 1389, 1264, 1140, 851, 770, 691  $cm^{-1}$ ; UV ( $CH_2Cl_2$ )  $\lambda_{max}$  (log  $\epsilon$ ) 852.0 sh (2.62), 781.0 (3.78), 708.0 (3.69), 650.0 sh (3.40), 535.5 (3.45), 456.5 (3.69), 357.5 (4.16), 288.5 (4.52), 245.0 (4.72) nm; HRMS (ESI-TOF)  $[M+H]^+$   $m/z$  calcd for  $C_{33}H_{32}N_4$ : 484.2627; found: 484.2604. Anal. Calcd for  $C_{33}H_{31}N_4$ : C, 81.95; H, 6.46; N, 11.58. Found: C, 81.65; H, 6.87; N, 11.55%.

### Preparation of 8-substituted 3-phenylbenzo[e][1,2,4]triazines **2**. General procedure.

A mixture of 8-fluoro-3-phenylbenzo[e][1,2,4]triazine<sup>2</sup> (**3**, 225.0 mg, 1.0 mmol), appropriate heterocycle **4** (1.1 mmol), and 60% NaH (45.0 mg, 1.1 mmol) in dry DMSO (6 mL) were stirred at 80 °C for 3 h under Ar. After cooling,  $CH_2Cl_2$  (30 mL) was added, and the organic layer was washed well with  $H_2O$  (3  $\times$  25 mL) and brine (25 mL). The organic layer was dried ( $Na_2SO_4$ ), and the solvent was removed in *vacuo*. The resulting solid residue was absorbed

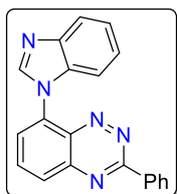
onto silica and purified by column chromatography using EtOAc in pet. ether gradient 5–10%. The solvent was evaporated, and the product was recrystallized.

### 8-(1*H*-Indol-1-yl)-3-phenylbenzo[*e*][1,2,4]triazine (2a).



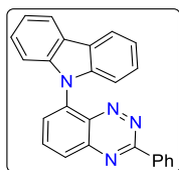
Obtained 280.0 mg (87% yield) as yellow crystals: mp 198–199 °C (EtOAc); <sup>1</sup>H NMR (600 MHz, CDCl<sub>3</sub>) δ 8.82–8.77 (m, 2H), 8.12 (dd, *J*<sub>1</sub> = 8.6 Hz, *J*<sub>2</sub> = 1.5 Hz, 1H), 8.09 (t, *J* = 7.8 Hz, 1H), 7.99 (d, *J* = 7.1 Hz, 1H), 7.78 (d, *J* = 3.0 Hz, 1H), 7.77–7.74 (m, 1H), 7.64–7.58 (m, 3H), 7.50–7.46 (m, 1H), 7.26–7.21 (m, 2H), 6.85 (d, *J* = 3.2 Hz, 1H), <sup>13</sup>C{<sup>1</sup>H} NMR (151MHz, CDCl<sub>3</sub>) δ 160.0, 142.2, 142.1, 137.7, 137.2, 135.5, 135.3, 132.0, 131.2, 129.8, 129.2, 129.1, 127.7, 125.7, 122.8, 121.4, 121.2, 110.6, 104.9; IR (KBr) ν 1601, 1563, 1509, 1457, 1389, 1355, 1318, 1244, 1211, 1149, 1010, 799, 745, 703 cm<sup>-1</sup>; UV (CH<sub>2</sub>Cl<sub>2</sub>) λ<sub>max</sub> (log ε) 427.5 (3.40), 352.0 (3.62), 306.0 (4.13), 261.5 (4.63) nm; HRMS (ESI-TOF) [M+H]<sup>+</sup> calcd for C<sub>21</sub>H<sub>15</sub>N<sub>4</sub>: 323.1297; found: 323.1299. Anal. Calcd for C<sub>21</sub>H<sub>14</sub>N<sub>4</sub>: C, 78.24; H, 4.38; N, 17.38. Found: C, 78.25; H, 4.34; N, 17.37%.

### 8-(1*H*-Benzo[*d*]imidazol-1-yl)-3-phenylbenzo[*e*][1,2,4]triazine (2b).



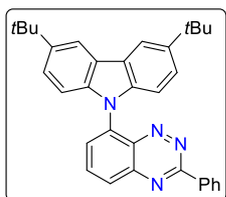
Obtained 301.0 mg (93% yield) as yellow crystals: mp 200–201 °C (EtOAc); <sup>1</sup>H NMR (600 MHz, CDCl<sub>3</sub>) δ 8.81–8.77 (m, 2H), 8.56 (s, 1H), 8.23 (d, *J* = 8.5 Hz, 1H), 8.16 (t, *J* = 8.0 Hz, 1H), 8.02 (d, *J* = 7.5 Hz, 1H), 7.97 (d, *J*<sub>1</sub> = 8.0 Hz, 1H), 7.64–7.60 (m, 3H), 7.49 (d, *J* = 7.9 Hz, 1H), 7.40 (t, *J* = 7.5 Hz, 1H), 7.36 (t, *J* = 7.5 Hz, 1H); <sup>13</sup>C{<sup>1</sup>H} NMR (151MHz, CDCl<sub>3</sub>) δ 160.6, 144.8, 144.0, 142.2, 141.4, 135.5, 135.0, 134.6, 134.5, 132.3, 129.31, 129.28, 129.2, 125.3, 124.2, 123.5, 121.0, 110.6; IR (neat) ν 1605, 1563, 1502, 1449, 1385, 1327, 1242, 1198, 1010, 803, 764, 737, 705 cm<sup>-1</sup>; UV (CH<sub>2</sub>Cl<sub>2</sub>) λ<sub>max</sub> (log ε) 352.5 (3.78), 295.0 sh (4.29), 260 (4.59) nm; HRMS (ESI-TOF) [M+H]<sup>+</sup> calcd for C<sub>20</sub>H<sub>14</sub>N<sub>5</sub>: 324.1249; found: 324.1243. Anal. Calcd for C<sub>20</sub>H<sub>13</sub>N<sub>5</sub>: C, 74.29; H, 4.05; N, 21.66. Found: C, 74.30; H, 4.11; N, 21.45%.

### 9-(3-Phenylbenzo[e][1,2,4]triazin-8-yl)-9H-carbazole (2c).



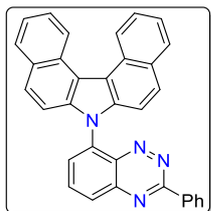
Obtained 276.5–281.0 mg (73–75% yield) as yellow crystals: mp 268.0–269.0 °C (EtOAc);  $^1\text{H}$  NMR (400 MHz,  $\text{CDCl}_3$ )  $\delta$  8.79–8.75 (m, 2H), 8.28 (dd,  $J_1 = 8.6$  Hz,  $J_2 = 1.0$  Hz, 1H), 8.22–8.17 (m, 3H), 8.02 (dd,  $J_1 = 7.2$  Hz,  $J_2 = 1.1$  Hz, 1H), 7.60–7.57 (m, 3H), 7.36 (td,  $J_1 = 7.5$  Hz,  $J_2 = 1.2$  Hz, 2H), 7.32 (td,  $J_1 = 7.1$  Hz,  $J_2 = 0.7$  Hz, 2H), 7.12 (d,  $J = 8.0$  Hz, 2H);  $^{13}\text{C}\{^1\text{H}\}$  NMR (101 MHz,  $\text{CDCl}_3$ )  $\delta$  160.2, 143.0, 142.4, 142.3, 136.6, 135.8, 135.3, 132.0, 129.7, 129.6, 129.2, 129.1, 126.2, 124.1, 120.6, 120.5, 110.2; IR (KBr)  $\nu$  1606, 1563, 1504, 1451, 1371, 1324, 1276, 1232, 1004, 803, 750, 706  $\text{cm}^{-1}$ ; UV ( $\text{CH}_2\text{Cl}_2$ )  $\lambda_{\text{max}}$  (log  $\epsilon$ ) 444.5 (3.17), 354.5 (3.65), 333.5 (4.05), 319.5 (4.09), 270.5 (4.63), 258.5 (4.65), 248.5 (4.66), 239.0 (4.78) nm; HRMS (ESI-TOF)  $[\text{M}+\text{H}]^+$  calcd for  $\text{C}_{25}\text{H}_{17}\text{N}_4$ : 373.1453; found: 373.1458. Anal. Calcd for  $\text{C}_{25}\text{H}_{16}\text{N}_4$ : C, 80.63; H, 4.33; N, 15.04. Found: C, 80.59; H, 4.27; N, 15.06%.

### 3,6-Di-*tert*-butyl-9-(3-phenylbenzo[e][1,2,4]triazin-8-yl)-9H-carbazole (2d).



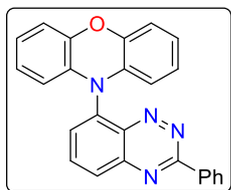
Obtained 448.0 mg (92% yield) as yellow crystals: mp 301–303 °C (EtOAc);  $^1\text{H}$  NMR (400 MHz,  $\text{CDCl}_3$ )  $\delta$  8.78 – 8.76 (m, 2H), 8.23 (dd,  $J_1 = 8.6$ ,  $J_2 = 1.3$  Hz, 1H), 8.18 – 8.17 (m, 2H), 8.15 (dd,  $J_1 = 8.6$  Hz,  $J_2 = 7.3$  Hz, 1H), 7.99 (dd,  $J_1 = 7.3$ ,  $J_2 = 1.3$  Hz, 1H), 7.61 – 7.57 (m, 3H), 7.40 (dd,  $J_1 = 8.6$  Hz,  $J_2 = 1.9$  Hz, 2H), 7.05 (d,  $J = 8.6$  Hz, 2H), 1.46 (s, 18H);  $^{13}\text{C}\{^1\text{H}\}$  NMR (101 MHz,  $\text{CDCl}_3$ )  $\delta$  160.1, 143.5, 143.0, 142.4, 140.8, 137.2, 135.8, 135.4, 131.9, 129.2, 129.15, 129.1, 129.0, 124.2, 123.9, 116.5, 109.7, 34.9, 32.2; IR (neat)  $\nu$  2955, 1609, 1564, 1482, 1360, 1290, 1236, 1176, 1102, 1057, 1009, 882, 797, 699  $\text{cm}^{-1}$ ; UV ( $\text{CH}_2\text{Cl}_2$ )  $\lambda_{\text{max}}$  (log  $\epsilon$ ) 468.5 (3.23), 363.5 sh (3.59), 339.0 (4.06), 326.5 (4.07), 293.0 (4.50), 272.5 (4.62), 252.0 (4.69), 242.0 (4.77) nm; HRMS (ESI-TOF)  $[\text{M}+\text{H}]^+$  calcd for  $\text{C}_{33}\text{H}_{33}\text{N}_4$ : 485.2705; found: 485.2705. Anal. Calcd for  $\text{C}_{33}\text{H}_{32}\text{N}_4$ : C, 81.78; H, 6.66; N, 11.56. Found: C, 81.72; H, 6.73; N, 11.54%.

### 7-(3-Phenylbenzo[e][1,2,4]triazin-8-yl)-7H-dibenzo[c,g]carbazole (2e).



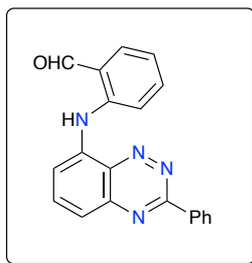
Obtained 417.0 mg (88% yield) as yellow crystals: mp 295.0–297.0 °C (EtOAc);  $^1\text{H}$  NMR (400 MHz,  $\text{CDCl}_3$ )  $\delta$  9.31 (d,  $J = 8.5$  Hz, 2H), 8.76–8.74 (m, 2H), 8.36 (dd,  $J_1 = 8.6$  Hz,  $J_2 = 1.2$  Hz, 1H), 8.23 (dd,  $J_1 = 8.7$  Hz,  $J_2 = 7.3$  Hz, 1H), 8.07 (dd,  $J_1 = 7.2$  Hz,  $J_2 = 1.2$  Hz, 1H), 8.03 (dd,  $J_1 = 8.2$  Hz,  $J_2 = 1.4$  Hz, 2H), 7.80 (d,  $J = 8.8$  Hz, 2H), 7.74 (ddd,  $J_1 = 8.4$  Hz,  $J_2 = 7.7$  Hz,  $J_3 = 1.4$  Hz, 2H), 7.60–7.53 (m, 5H), 7.30 (d,  $J = 8.8$  Hz, 2H);  $^{13}\text{C}\{^1\text{H}\}$  NMR (101 MHz,  $\text{CDCl}_3$ )  $\delta$  160.5, 143.3, 142.3, 139.5, 135.8, 135.6, 135.2, 132.1, 130.7, 130.6, 130.56, 129.3, 129.28, 129.2, 129.15, 127.2, 125.71, 125.67, 123.7, 118.6, 111.8; IR (neat)  $\nu$  1609, 1562, 1502, 1381, 1322, 1175, 1108, 1059, 1009, 919, 791, 702  $\text{cm}^{-1}$ ; UV ( $\text{CH}_2\text{Cl}_2$ )  $\lambda_{\text{max}}$  (log  $\epsilon$ ) 429.5 (3.16), 364.5 (4.48), 346.5 (4.37), 303.0 (4.50), 378.0 (4.84) nm; HRMS (ESI-TOF)  $[\text{M}+\text{H}]^+$  Calcd for  $\text{C}_{33}\text{H}_{21}\text{N}_4$ : 473.1766; found: 473.1772. Anal. Calcd for  $\text{C}_{33}\text{H}_{20}\text{N}_4$ : C, 83.88; H, 4.27; N, 11.86. Found: C, 83.72; H, 4.48; N, 11.93%.

### 10-(3-Phenylbenzo[e][1,2,4]triazin-8-yl)-10H-phenoxazine (2f).



Obtained 341.8 mg (88% yield) as yellow crystals: mp 252.5–253.5 °C (EtOAc);  $^1\text{H}$  NMR (600 MHz,  $\text{CDCl}_3$ )  $\delta$  8.78–8.74 (m, 2H), 8.27 (dd,  $J_1 = 8.6$  Hz,  $J_2 = 1.0$  Hz, 1H), 8.17 (dd,  $J_1 = 8.6$  Hz,  $J_2 = 7.2$  Hz, 1H), 7.91 (dd,  $J_1 = 7.2$  Hz,  $J_2 = 1.1$  Hz, 1H), 7.60–7.57 (m, 3H), 6.76 (dd,  $J_1 = 8.0$  Hz,  $J_2 = 1.1$  Hz, 2H), 6.66 (td,  $J_1 = 7.9$  Hz,  $J_2 = 1.5$  Hz, 2H), 6.49 (td,  $J_1 = 7.7$  Hz,  $J_2 = 1.3$  Hz, 2H), 5.73 (dd,  $J_1 = 7.9$  Hz,  $J_2 = 1.1$  Hz, 2H);  $^{13}\text{C}\{^1\text{H}\}$  NMR (151 MHz,  $\text{CDCl}_3$ )  $\delta$  160.3, 144.1, 143.7, 143.6, 137.6, 136.5, 135.2, 134.7, 134.1, 132.0, 130.9, 129.2, 129.1, 123.3, 122.0, 116.0, 113.4; IR (neat)  $\nu$  1589, 1563, 1490, 1384, 1338, 1271, 1080, 1044, 1001, 798, 738, 709  $\text{cm}^{-1}$ ; UV ( $\text{CH}_2\text{Cl}_2$ )  $\lambda_{\text{max}}$  (log  $\epsilon$ ) 460.0 (2.66), 320.0 sh (4.05), 276.0 (4.45), 260.0 (4.48), 239.5 (4.71) nm; HRMS (ESI-TOF)  $m/z$   $[\text{M}+\text{H}]^+$  calcd for  $\text{C}_{25}\text{H}_{17}\text{N}_4\text{O}$ : 389.1402; found: 389.1399. Anal. Calcd for  $\text{C}_{25}\text{H}_{16}\text{N}_4\text{O}$ : C, 77.30; H, 4.15; N, 14.42. Found: C, 77.12; H, 4.49; N, 14.21%.

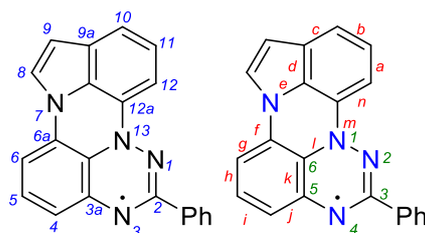
## 2-((3-Phenylbenzo[e][1,2,4]triazin-8-yl)amino)benzaldehyde (**5**).



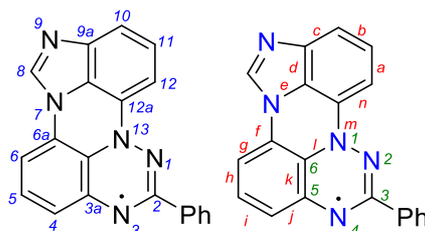
A solution of precursor **2a** (32.6 mg, 0.1 mmol) in EtOH (100 mL) was irradiated for 72 h. The reaction mixture was separated by column chromatography using EtOAc in pet. ether gradient 10-20% giving 17.4 mg (53% yield) of aldehyde **5** as orange crystals: mp 143-145 °C (EtOAc);  $^1\text{H}$  NMR (600 MHz,  $\text{CDCl}_3$ )  $\delta$  11.88 (s, 1H), 10.08 (s, 1H), 8.80-8.77 (m, 2H), 7.88 (d,  $J = 8.0$  Hz, 1H), 7.85 (d,  $J = 8.5$  Hz, 1H), 7.81 (d,  $J = 7.6$  Hz, 1H), 7.77 (dd,  $J_1 = 7.7$  Hz,  $J_2 = 1.5$  Hz, 1H), 7.62-7.57 (m, 5H), 7.13 (t,  $J = 7.1$  Hz, 1H);  $^{13}\text{C}$   $\{^1\text{H}\}$  NMR (176 MHz,  $\text{CDCl}_3$ )  $\delta$  194.1, 160.7, 143.8, 142.5, 139.7, 139.5, 137.0, 136.9, 135.8, 135.4, 131.6, 129.1, 129.0, 123.2, 120.6, 120.0, 115.9, 112.3; IR (neat)  $\nu$  2922, 2851, 2734, 1665, 1579, 1456, 1371, 1316, 1166, 877, 760, 686  $\text{cm}^{-1}$ ; HRMS (ESI-TOF)  $m/z$   $[\text{M}+\text{H}]^+$  Calcd for  $\text{C}_{20}\text{H}_{15}\text{N}_4\text{O}$ : 327.1240; found: 327.1241.

## 2. Nomenclature of radicals **1** according to IUPAC rules

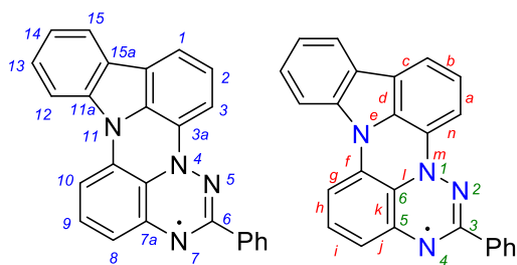
In blue: numbering of the main structure. In green: numbering of the [1,2,4]triazinyl fused ring. In red: assigned letters to the phenazine core.



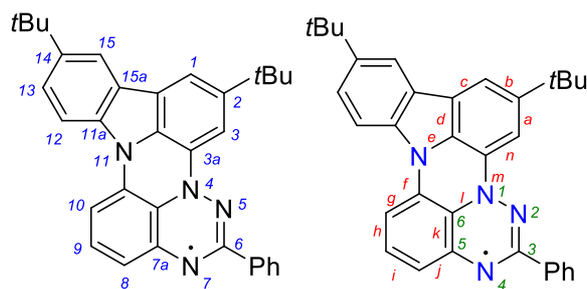
2-Phenyl-3*H*-pyrrolo[3,2,1-*de*][1,2,4]triazino[5,6,1-*kl*]phenazin-3-yl (**1a**).



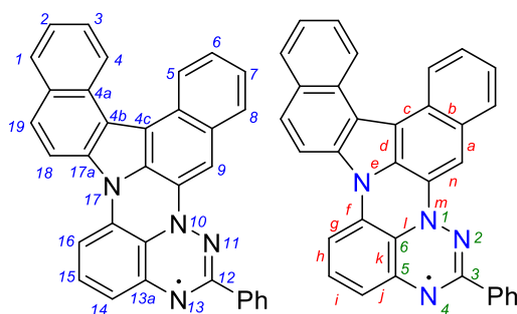
2-Phenyl-3*H*-imidazo[4,5,1-*de*][1,2,4]triazino[5,6,1-*kl*]phenazin-3-yl (**1b**).



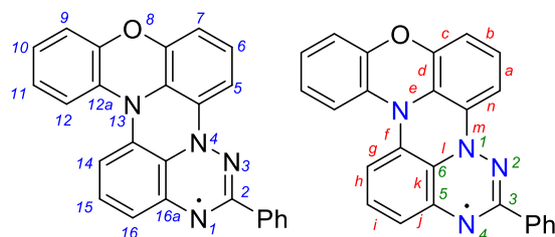
6-Phenyl-7*H*-indolo[3,2,1-*de*][1,2,4]triazino[5,6,1-*kl*]phenazin-7-yl (**1c**).



2,14-Di-*tert*-butyl-6-phenyl-7*H*-indolo[3,2,1-*de*][1,2,4]triazino[5,6,1-*kl*]phenazin-7-yl (**1d**).

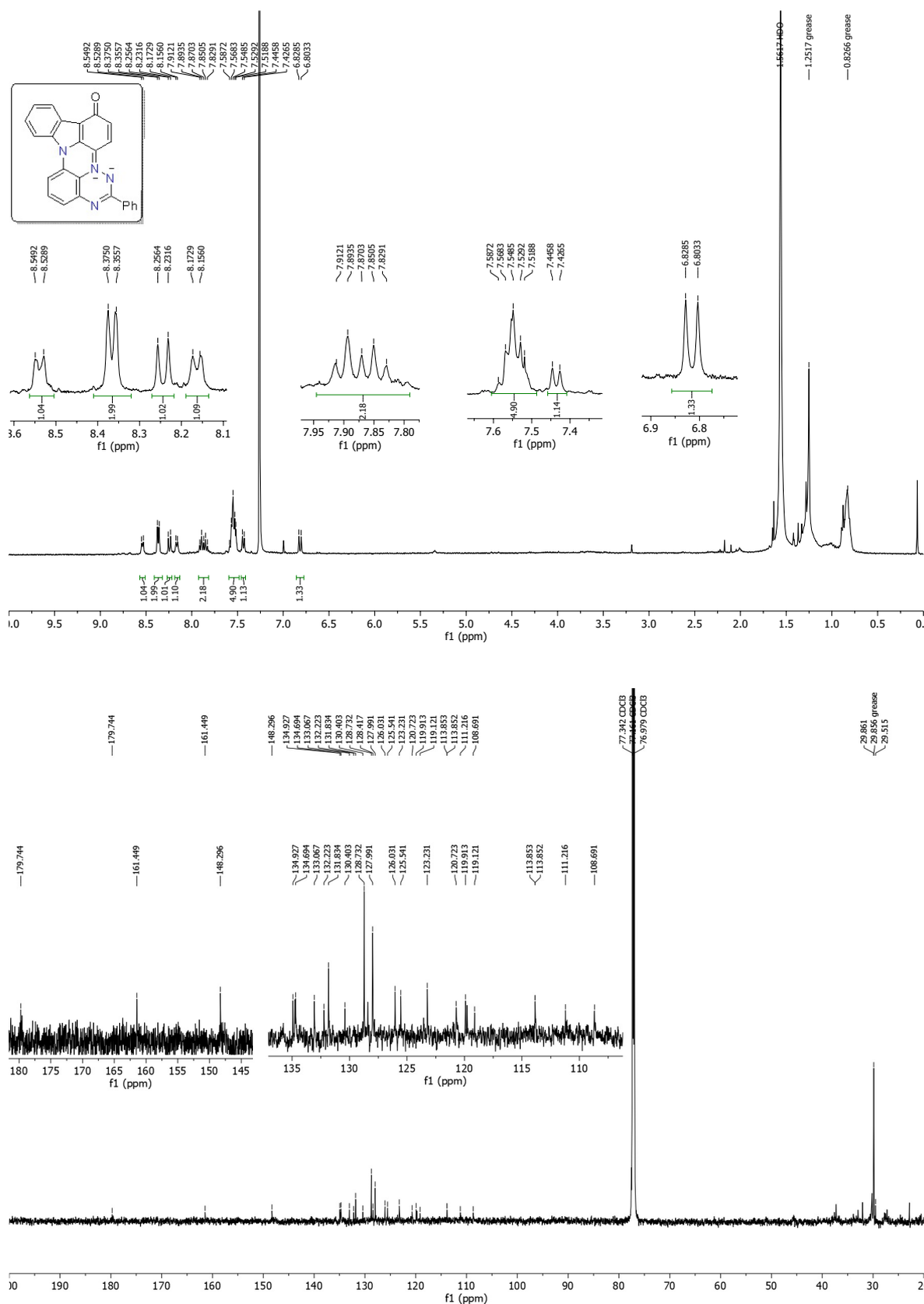


12-Phenyl-13*H*-benzo[*b*]benzo[4,5]indolo[3,2,1-*de*][1,2,4]triazino[5,6,1-*kl*]phenazin-13-yl (**1e**).

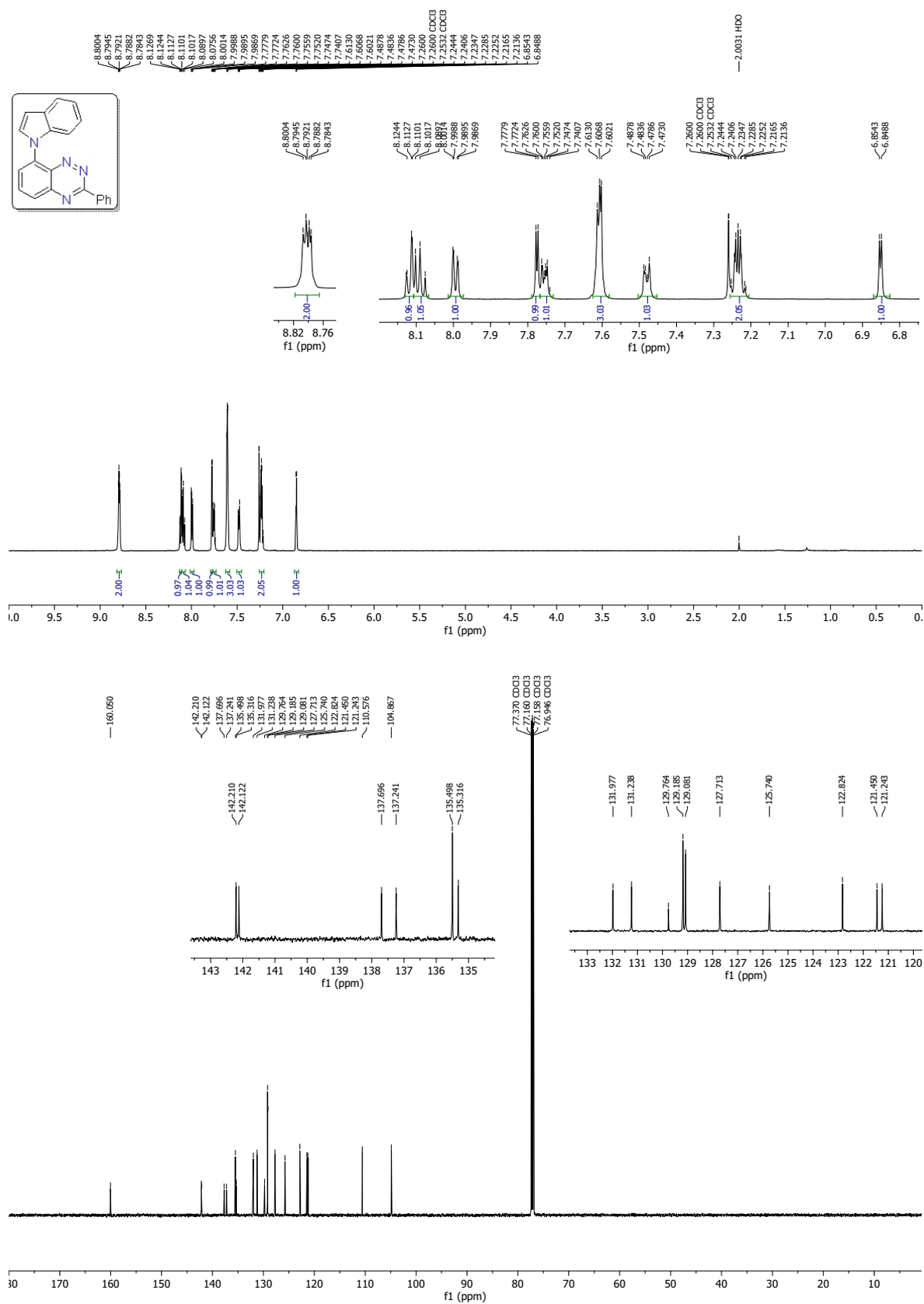


2-Phenyl-1*H*-benzo[5,6][1,4]oxazino[2,3,4-*de*][1,2,4]triazino[5,6,1-*kl*]phenazin-1-yl (**1f**).

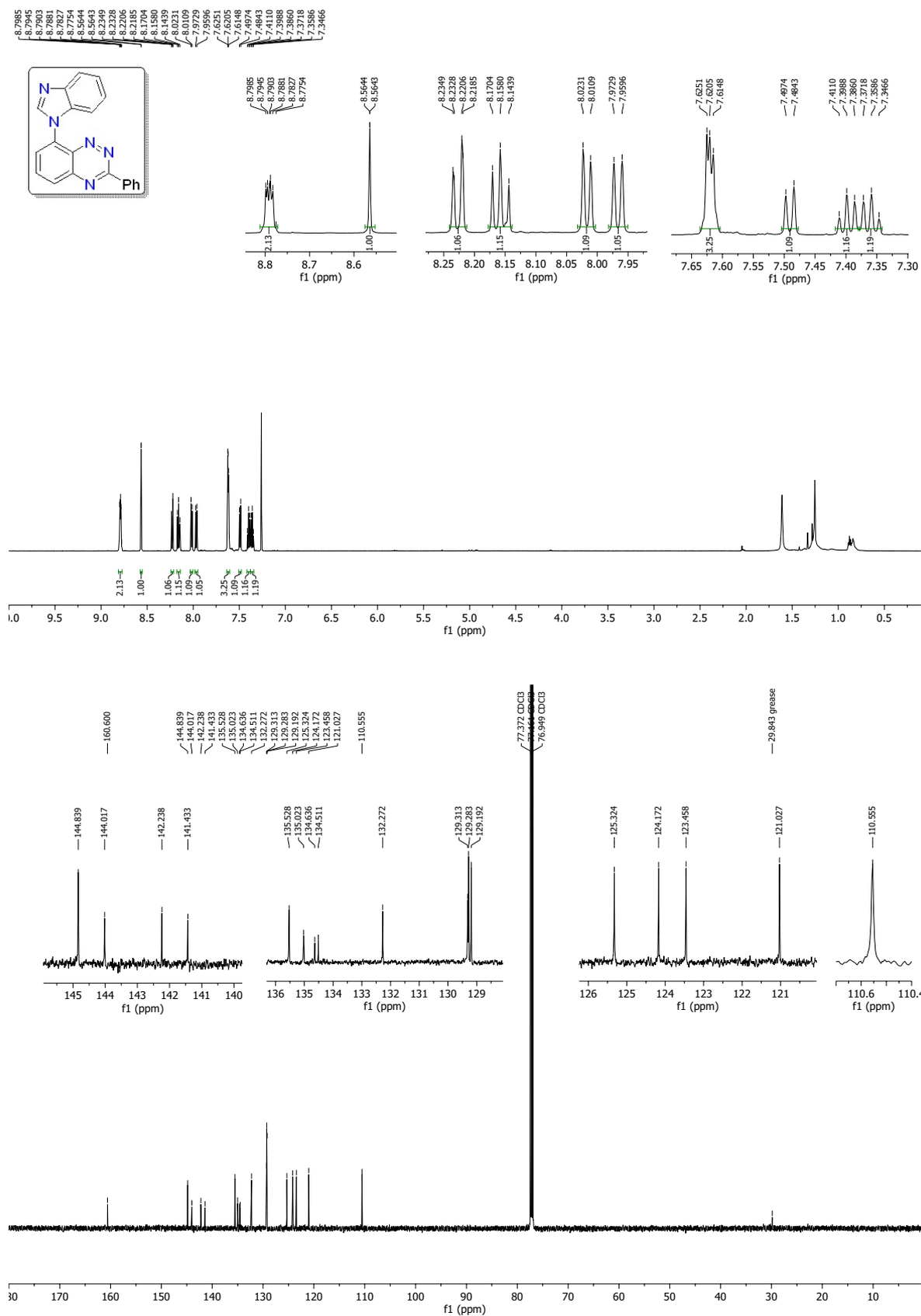
### 3. NMR spectra



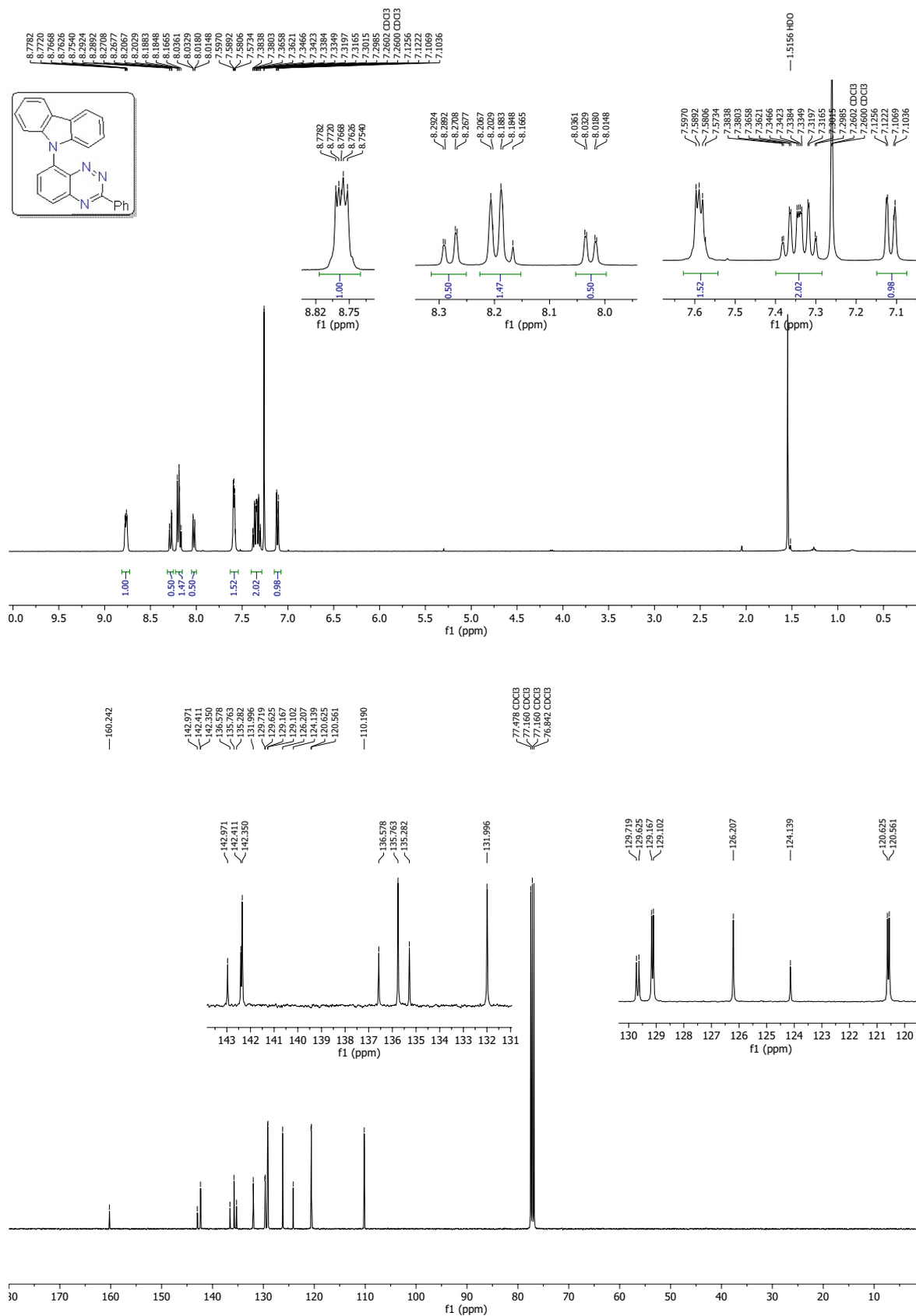
**Figure S1.** <sup>1</sup>H NMR and <sup>13</sup>C NMR spectra of 1-oxo-6-phenyl-1H-indolo[3,2,1-de][1,2,4]triazino[5,6,1-k]phenazin-4-ium-5-ide (**1c-oxo**) recorded in CDCl<sub>3</sub> at 400 and 101 MHz, respectively.



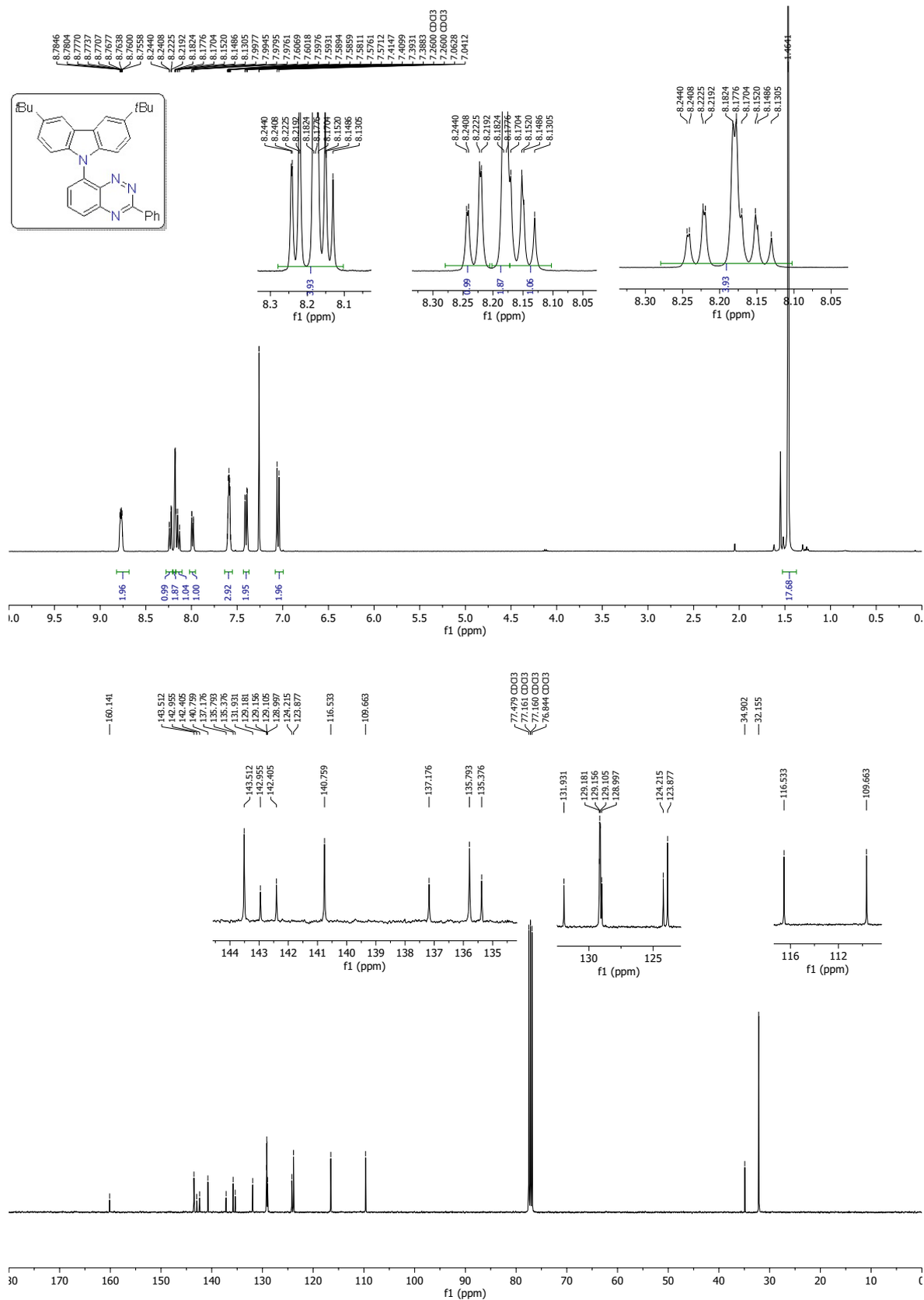
**Figure S2.** <sup>1</sup>H NMR and <sup>13</sup>C NMR spectra of 8-(1*H*-indol-1-yl)-3-phenylbenzo[*e*][1,2,4]triazine (**2a**) recorded in CDCl<sub>3</sub> at 600 and 151 MHz, respectively.



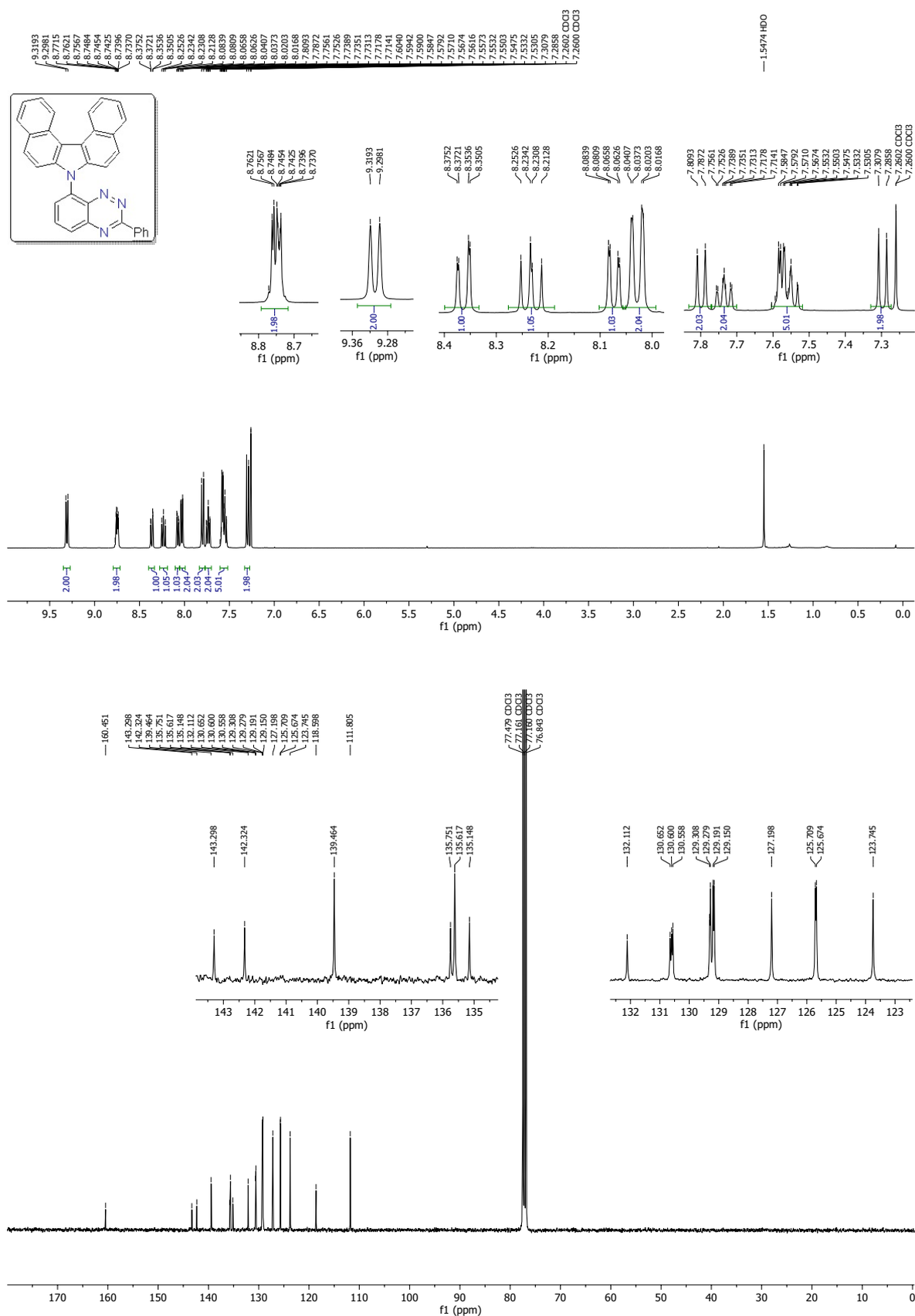
**Figure S3.** <sup>1</sup>H NMR and <sup>13</sup>C NMR spectra of 8-(1H-benzo[d]imidazol-1-yl)-3-phenylbenzo[e][1,2,4]triazine (**2b**) recorded in CDCl<sub>3</sub> at 600 and 151 MHz, respectively.



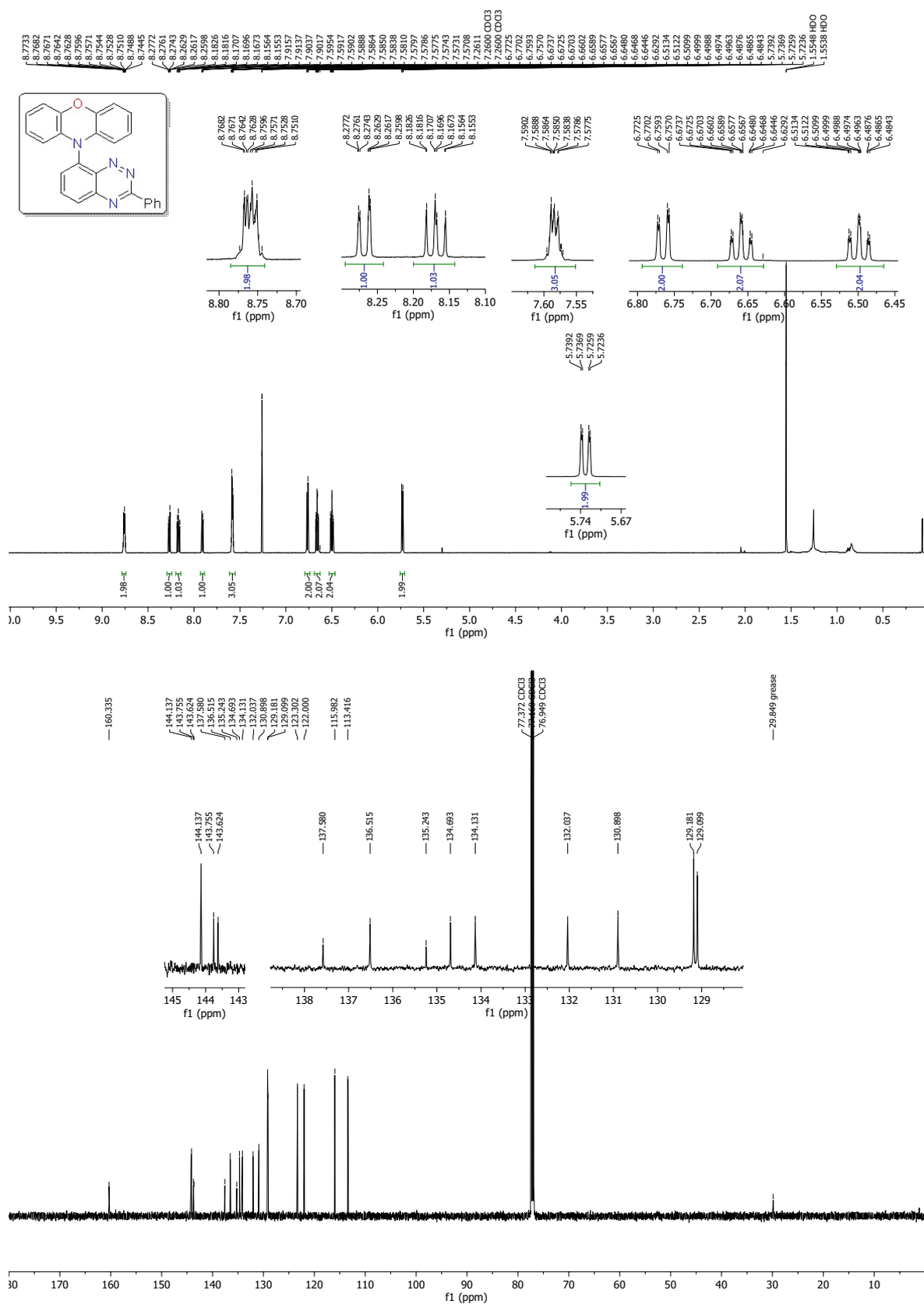
**Figure S4.** <sup>1</sup>H NMR and <sup>13</sup>C NMR spectra of 9-(3-phenylbenzo[e][1,2,4]triazin-8-yl)-9H-carbazole (**2c**) recorded in CDCl<sub>3</sub> at 400 and 101 MHz, respectively.

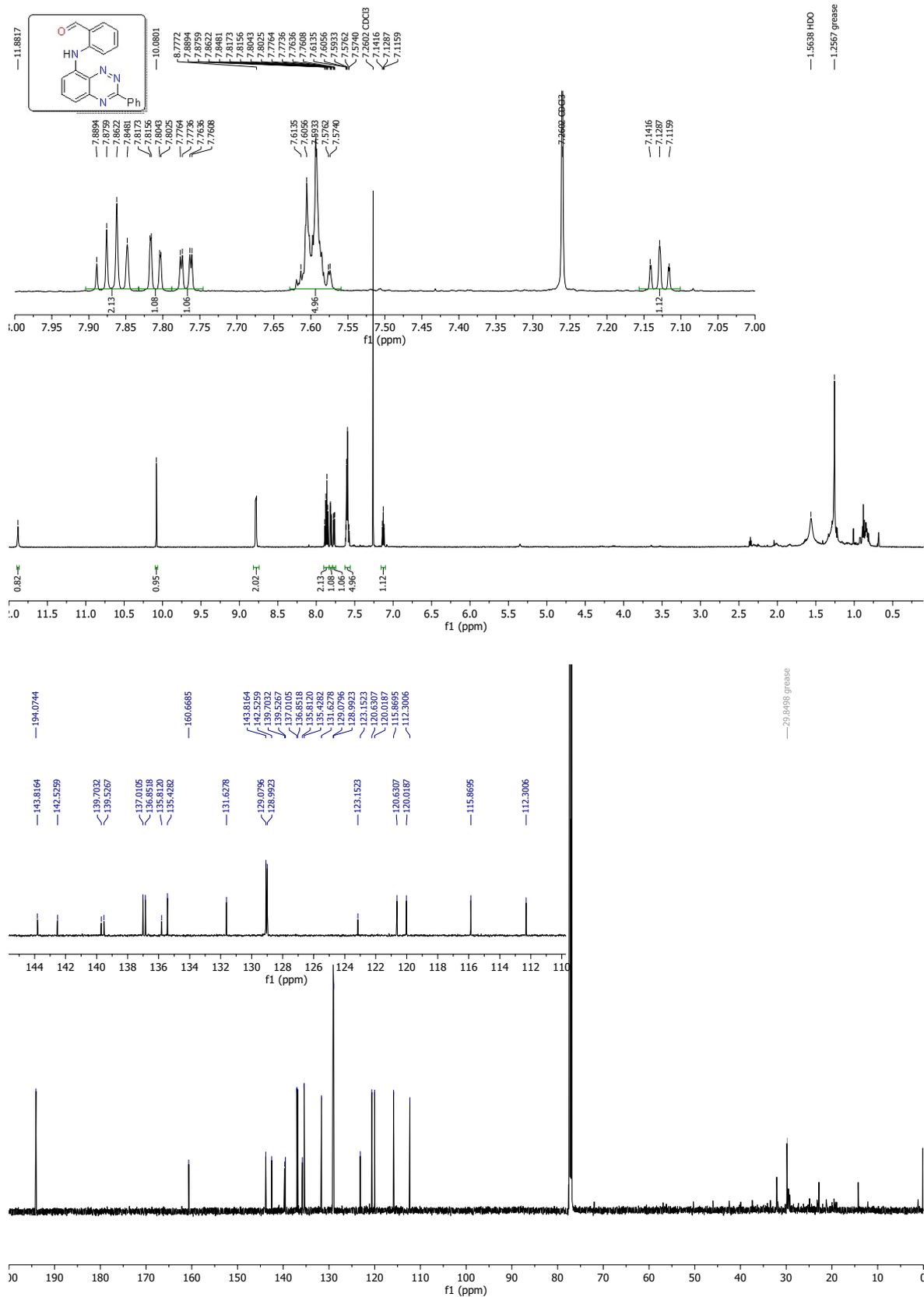


**Figure S5.** <sup>1</sup>H NMR and <sup>13</sup>C NMR spectra of 3,6-di-*tert*-butyl-9-(3-phenylbenzo[*e*][1,2,4]triazin-8-yl)-9H-carbazole (**2d**) recorded in CDCl<sub>3</sub> at 400 and 101 MHz, respectively.



**Figure S6.** <sup>1</sup>H NMR and <sup>13</sup>C NMR spectra of 7-(3-phenylbenzo[e][1,2,4]triazin-8-yl)-7H-dibenzo[c,g]carbazole (**2e**) recorded in CDCl<sub>3</sub> at 400 and 101 MHz, respectively.





**Figure S8.** <sup>1</sup>H NMR and <sup>13</sup>C NMR spectra of 2-((3-phenylbenzo[e][1,2,4]triazin-8-yl)amino)benzaldehyde (**5**) recorded in CDCl<sub>3</sub> at 600 and 176 MHz, respectively.

#### 4. XRD data collection and refinement for **1d** and **1c-oxo**

Single-crystal XRD measurements for **1d** and **1c-oxo** were performed with a Rigaku XtaLAB Synergy, Pilatus 300K diffractometer. The measurements were conducted at 100.0(5) K using the  $\text{CuK}\alpha$  radiation ( $\lambda=1.54184$  Å). The data was integrated using CrysAlisPro program.<sup>3</sup> Intensities for absorption were corrected using multi-scan method as in SCALE3 ABSPACK scaling algorithm implemented in CrysAlisPro program. Additional crystal and refinement information are listed in Table S1. Selected geometrical parameters are provided in Table S2 and compared to those of **A(X=H)** and **B(X=H)**.<sup>2</sup> Selected close contacts for **1d** are presented in Figure S9 and in Table S3. All dimensions and s.u. values are copied as they appear in the CIF.

CCDC: 2327427 and 2327428 contain the supplementary crystallographic data for this paper. The data is provided free of charge by The Cambridge Crystallographic Data Center.

**Table S1.** Selected Structural Data for **1d** and **1c-oxo**.

	<b>1d</b> CCDC: 2327427	<b>1c-oxo</b> CCDC: 2327428
Formula	$\text{C}_{33}\text{H}_{31}\text{N}_4$	$\text{C}_{25}\text{H}_{14}\text{N}_4\text{O}$
Formula Weight	483.62	386.40
Crystal System	Monoclinic	Monoclinic
Space Group	$C2/c$	$P2_1/n$
$a/\text{Å}$	22.1999(3)	7.4588(4)
$b/\text{Å}$	10.4920(1)	23.524(1)
$c/\text{Å}$	21.9787(3)	19.4019(1)
$\alpha/^\circ$	90	90
$\beta/^\circ$	98.613(1)	92.652(4)
$\gamma/^\circ$	90	90
Volume/ $\text{Å}^3$	5061.6(1)	3400.6(3)
Z	8	8
$2\theta$ range for data collection/ $^\circ$	8.056 to 157.474	7.516 to 134.16
Index ranges	$-26 \leq h \leq 26, -13 \leq k \leq 9, -27 \leq l \leq 27$	$-8 \leq h \leq 8, -28 \leq k \leq 28, -20 \leq l \leq 22$
No. of measured, independent, and observed [ $I > 2\sigma(I)$ ] reflections	33089, 5125, 4376	34713, 5961, 4006
$R_{\text{int}}$	0.0298	0.1146
Goodness-of-fit on $F^2$	1.055	1.060
Final $R$ indexes [ $F^2 > 2\sigma(F^2)$ ]	$R_1=0.0384, wR2=0.0966$	$R_1=0.0616, wR2=0.1542$
Final $R$ indexes [all data]	$R_1=0.0461, wR2=0.1012$	$R_1=0.0948, wR2=0.1756$
Data/restraints/parameters	5125/126/396	5961/0/541
Largest diff. peak/hole $\text{Å}^{-3}$	0.27/-0.32	0.32/-0.24

**Table S2.** Pertinent geometrical parameters for selected structures.<sup>a</sup>

	<b>1d</b>	<b>1c-oxo</b>	<b>A(X=H)<sup>b</sup></b>	<b>B(X=H)<sup>b</sup></b>		
CCDC #	2327427	2327428	1483990	1483989		
		A	B			
<i>d</i> N5–N4	1.3652(13)	1.366(3)	1.357(3)	<i>d</i> N1–N12	1.35(1)	1.368(3)
<i>d</i> N5–C6	1.3413(15)	1.333(4)	1.346(4)	<i>d</i> N1–C2	1.32(1)	1.331(3)
<i>d</i> C6–N7	1.3363(15)	1.332(4)	1.317(4)	<i>d</i> C2–N3	1.364(9)	1.340(4)
<i>d</i> N7–C7a	1.3772(15)	1.383(4)	1.375(4)	<i>d</i> N3–C3a	1.38(1)	1.372(4)
<i>d</i> C7a–C7a'	1.4108(16)	1.397(4)	1.422(4)	<i>d</i> C3a–C3a'	1.38(1)	1.418(4)
<i>d</i> C7a'–C10a	1.4185(16)	1.424(4)	1.420(4)	<i>d</i> C3a'–C6a	1.36(1)	1.401(4)
<i>d</i> C10a–N11	1.4035(15)	1.387(4)	1.412(3)	<i>d</i> C6a–X7	1.39(1)	1.757(3)
<i>d</i> N11–C3a'	1.3834(15)	1.385(4)	1.373(4)	<i>d</i> X7–C7a	1.381(1)	1.757(3)
<i>d</i> N4–C3a	1.3996(15)	1.335(4)	1.360(3)	<i>d</i> N12–C11a	1.42(1)	1.420(3)
<i>d</i> C6–Ph	1.4904(16)	1.496(4)	1.488(4)	<i>d</i> C2–Ph	1.50(1)	1.481(4)
$\alpha$ N4–N5–C6	115.13(10)	116.3(2)	115.7(2)	$\alpha$ N12–N1–C2	114.1(6)	116.3(2)
$\alpha$ N5–C6–N7	128.11(11)	128.4(3)	128.2(3)	$\alpha$ N1–C2–N3	130.2(7)	128.1(3)
$\alpha$ C3a'–N11–C10a	118.53(10)	118.9(2)	119.0(2)	$\alpha$ C7a–X7–C6a	118.5(6)	101.53(13)
$\alpha$ C7a'–N4–C3a	119.76(10)	120.4(2)	120.4(2)	$\alpha$ C3a'–N12–C11a	118.6(6)	123.2(2)
$\theta$ N4–N5–C6–N7	0.12(17)	0.7(4)	4.4(4)	$\theta$ N12–N1–C2–N3	1(1)	-0.8(4)
$\theta$ C7a'–C10a–N11–C3a'	-6.52(15)	-0.7(4)	2.0(4)	$\theta$ C3a'–C6a–X7–C7a	0(1)	17.5(3)
$\beta$ triazine-(N12)Ph	5.7	5.7	7.2	$\beta$ triazine-(N12)Ph	3.5	14.3
$\beta$ triazine-(C2)Ph	2.9	5.9	22.4	$\beta$ triazine-(C2)Ph	2.9	30.2

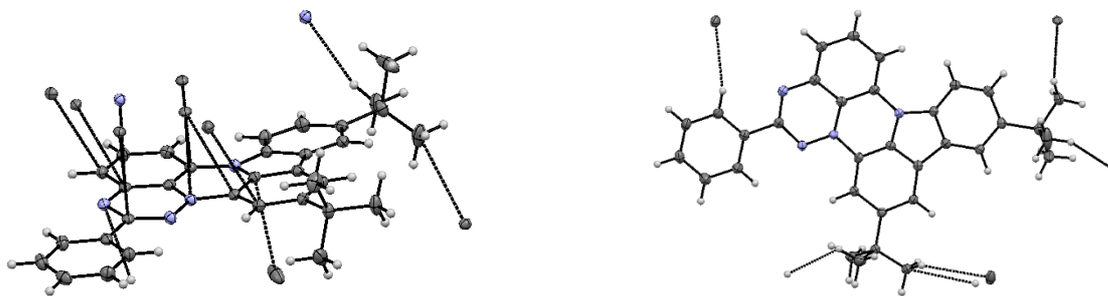
<sup>a</sup> For each structure appropriate systematic numbering scheme is used as shown in table head. *d* - interatomic distance,  $\alpha$  - interatomic angle,  $\theta$  - dihedral angle,  $\beta$  - inter-ring angle. <sup>b</sup> Ref<sup>2</sup>.

**Table S3.** Selected close contacts in the solid-state structure of **1d**.<sup>a</sup>

Contact X...Y	<i>d</i> <sub>X...Y</sub> /Å	<i>d</i> <sub>X-Y-vdW</sub> /Å	Calculated spin densities	Type of interaction <sup>b</sup>
C(6)···C(10a) <sup>c</sup>	3.288(2)	-0.112 <sup>e</sup>	+0.021...-0.057	F
C(3)···C(8) <sup>c</sup>	3.322(2)	-0.078 <sup>e</sup>	+0.094...-0.001	F
C(3a')-CH <sub>3</sub> (tBu) <sup>d</sup>	3.286(2)	-0.114 <sup>e</sup>	+0.053...0.001	A

<sup>a</sup> The systematic numbering scheme of the indolo[3,2,1-*de*][1,2,4]triazino[5,6,1-*kl*]phenazine skeleton for **1d** is used; see pg S8. <sup>b</sup> F – ferromagnetic, A –

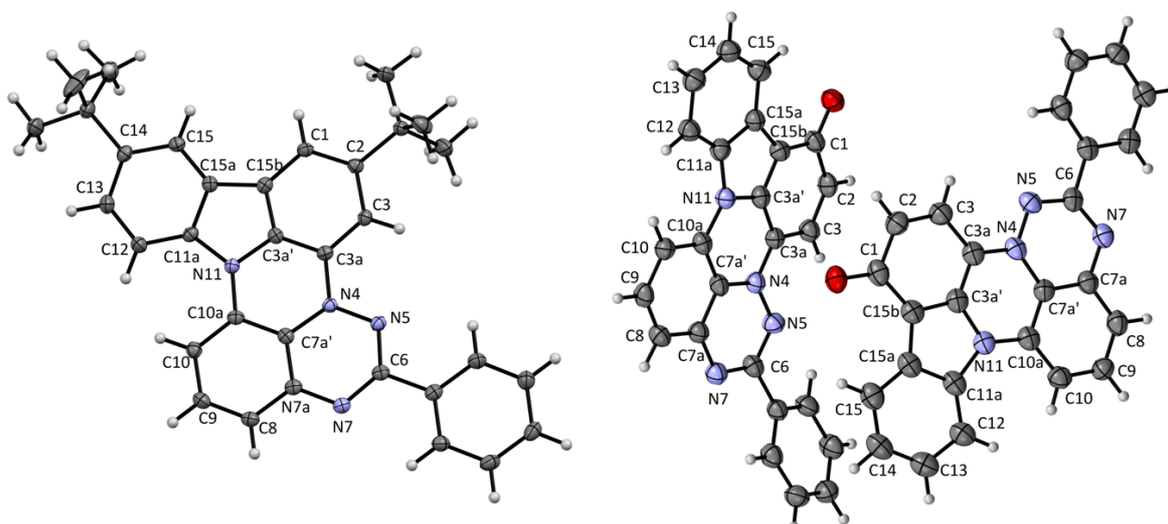
antiferromagnetic. <sup>c</sup> Symmetry code:  $-x + \frac{1}{2}, -y + \frac{3}{2}, -z + 1$ . <sup>d</sup> Symmetry code:  $-x + 1, y, -z + \frac{3}{2}$ .  
<sup>e</sup> Assuming 1.70 Å for the van der Waals radius of the carbon atom.



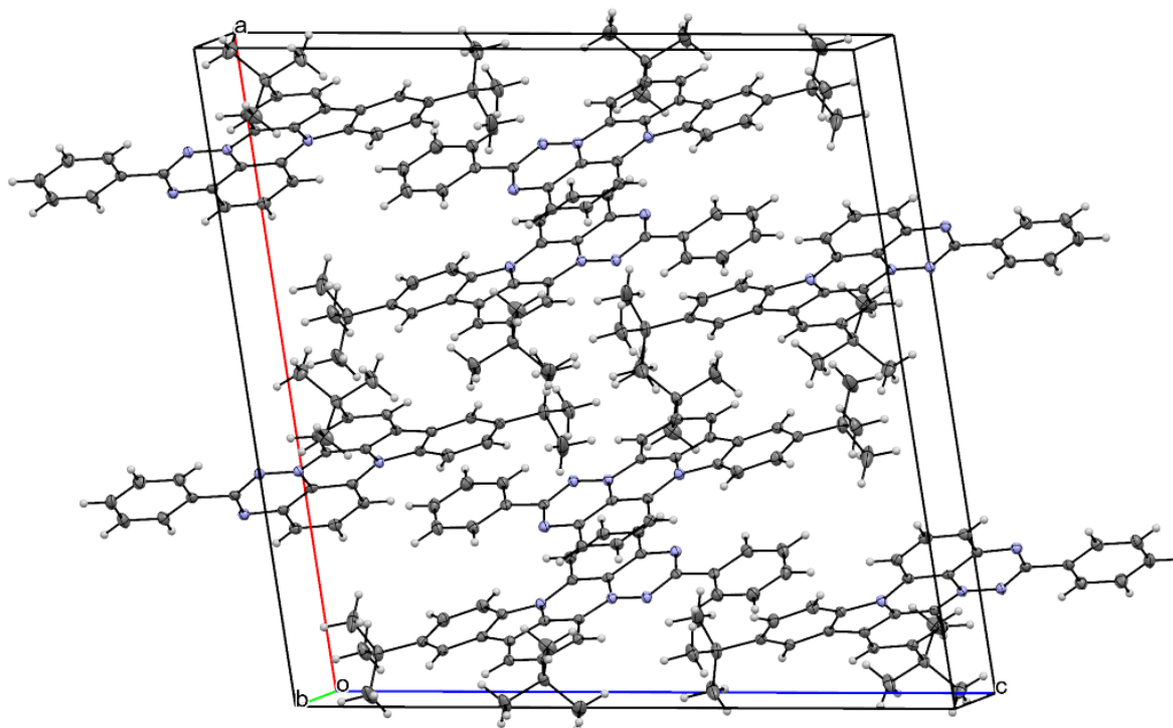
**Figure S9.** Close contacts in the crystal of **1d**: intradimer (left) and interdimer (right).

### **Structure solution and refinement**

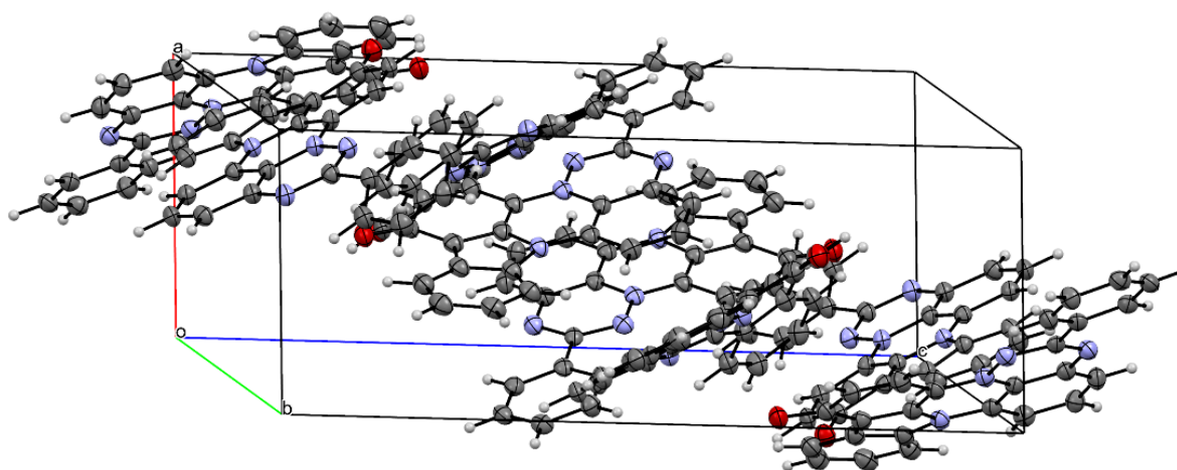
The structures were solved with the ShelXT<sup>4</sup> structure solution program using Intrinsic Phasing and refined by the full-matrix least-squares minimization on  $F^2$  was performed with the ShelXLrefinement package.<sup>5</sup> All non-hydrogen atoms were refined anisotropically and C–H hydrogens were generated geometrically using the HFIX command as in ShelXL. Hydrogen atoms were refined isotropically and constrained to ride on their parent atoms. Two *tert*-butyl groups in **1d** are rotationally disordered over two orientations with occupancy ratios 0.944(2):0.056(2) and 0.861(2):0.139(2). Sums of occupancies of relevant sites were set equal to 1 and refined using free variables. PART instruction was applied to exclude bonding between equivalent disordered atoms. Anisotropic displacement parameters of neighboring disordered atoms were restrained using SIMU and RIGU procedures as in ShelXL. The crystal data and structure refinement descriptors are presented in Table S1. Partial packing diagrams are shown in Figures S10 – S12.



**Figure S10.** Molecular structures of **1d** and **1c-oxo** with the numbering scheme of the indolo[3,2,1-*de*][1,2,4]triazino[5,6,1-*kl*]phenazine skeleton. The structure of **1c-oxo** contains two molecules in the asymmetric unit. Atomic displacement ellipsoids are drawn at 50% probability level.



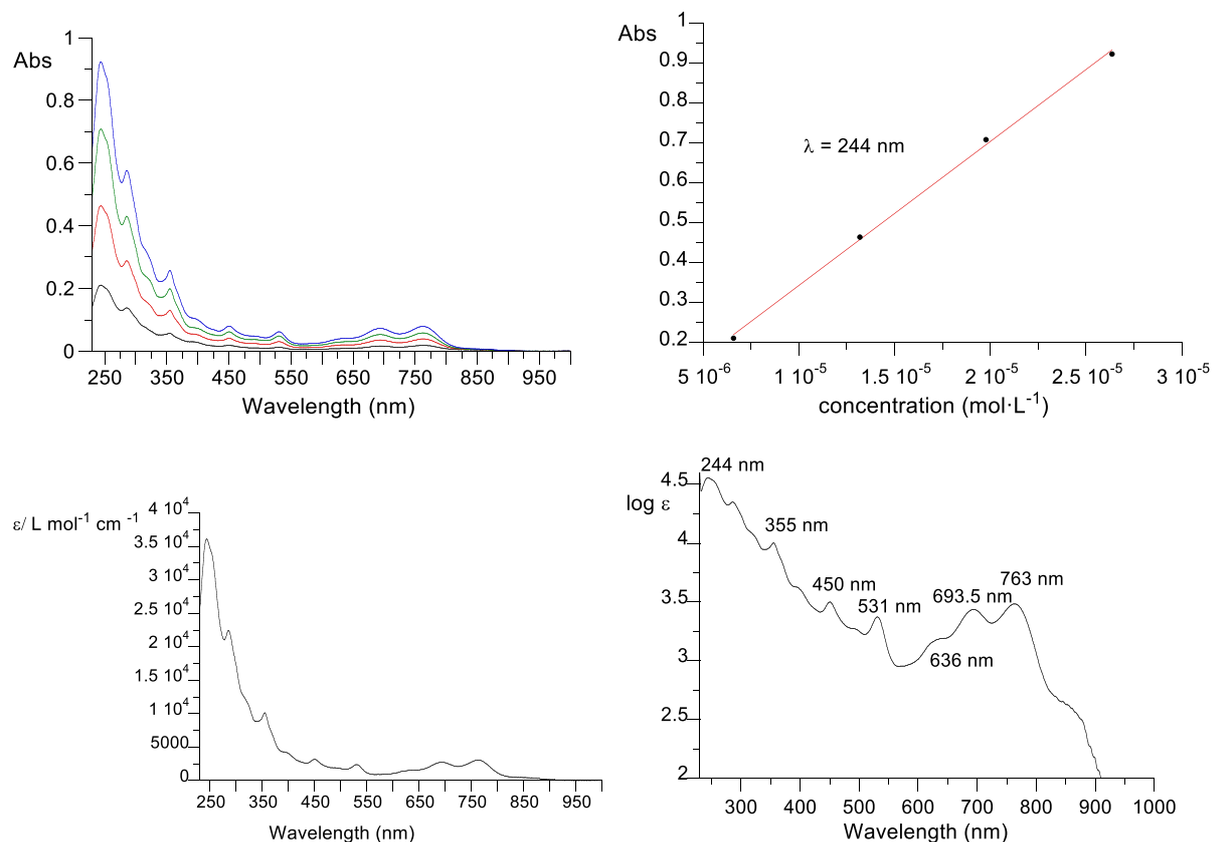
**Figure S11.** Packing diagram for the unit cell of **1d**.



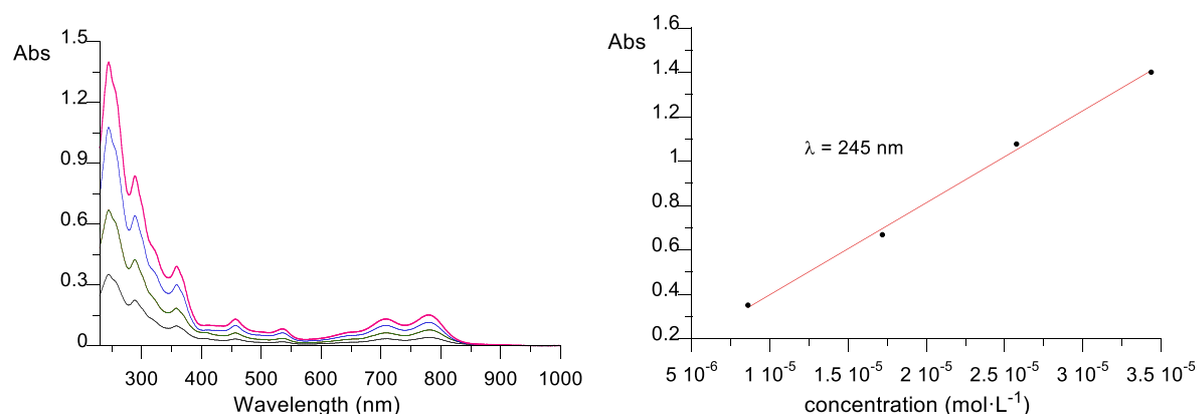
**Figure S12.** Packing diagram for the unit cell of **1c-oxo**.

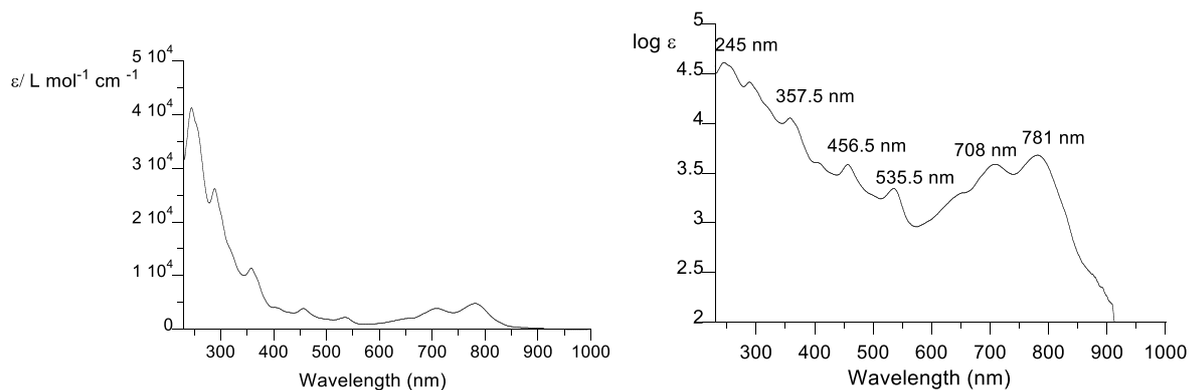
## 5. Electronic absorption spectroscopy

Electronic absorption spectra for radicals **1** and their precursors **2** were recorded in spectroscopic grade  $\text{CH}_2\text{Cl}_2$  at concentrations in a range  $1\text{--}10 \times 10^{-5}$  M and fitted to the Beer–Lambert law. Results for radicals **1c** and **1d** are shown in Figures S13 and S14. Figure S15 shows electronic absorption spectra for **1c-oxo**. Spectra of precursors **2** are shown in Figures S16–S21.

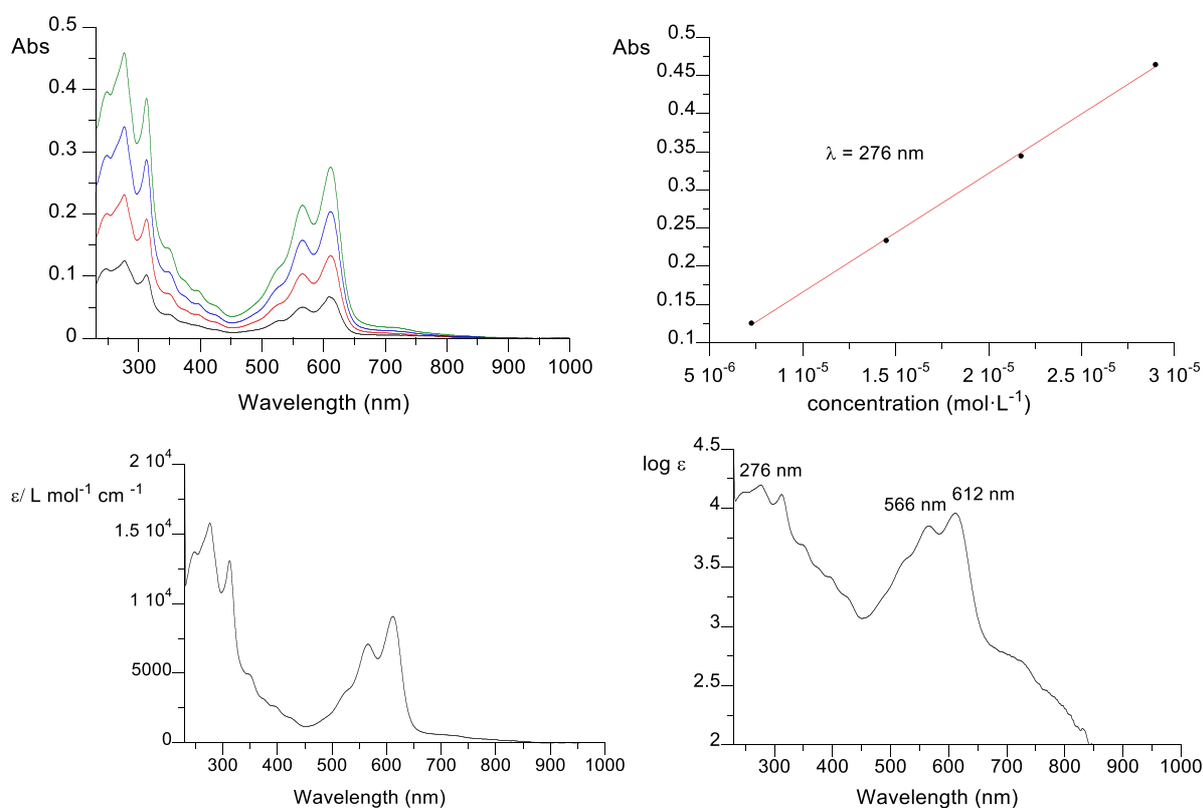


**Figure S13.** Clockwise: electronic absorption spectra for **1c** in  $\text{CH}_2\text{Cl}_2$  for four concentrations, determination of molar extinction coefficient  $\epsilon$  at  $\lambda = 244.0$  nm (best fit function:  $\epsilon = 35169(402) \times \text{conc}$ ,  $r^2 = 0.998$ ), molar extinction  $\log(\epsilon)$  plot.

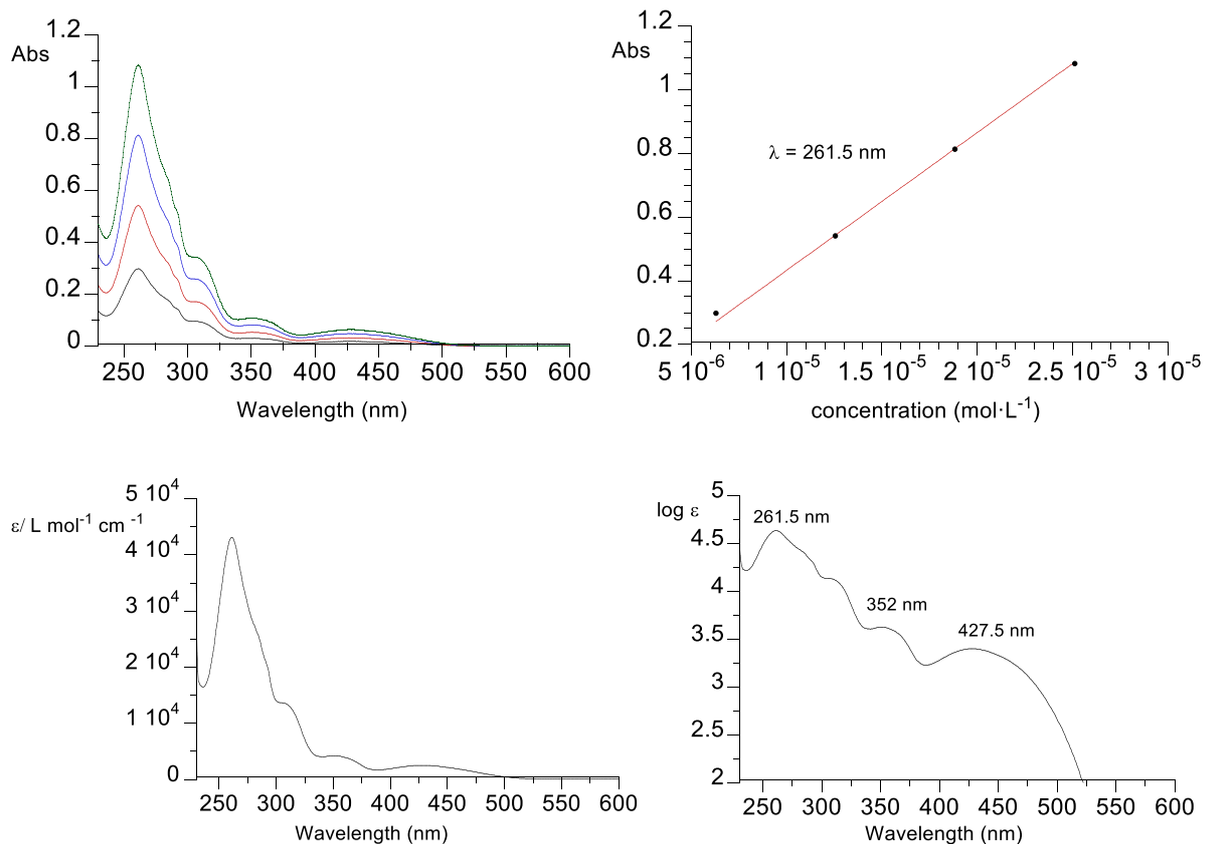




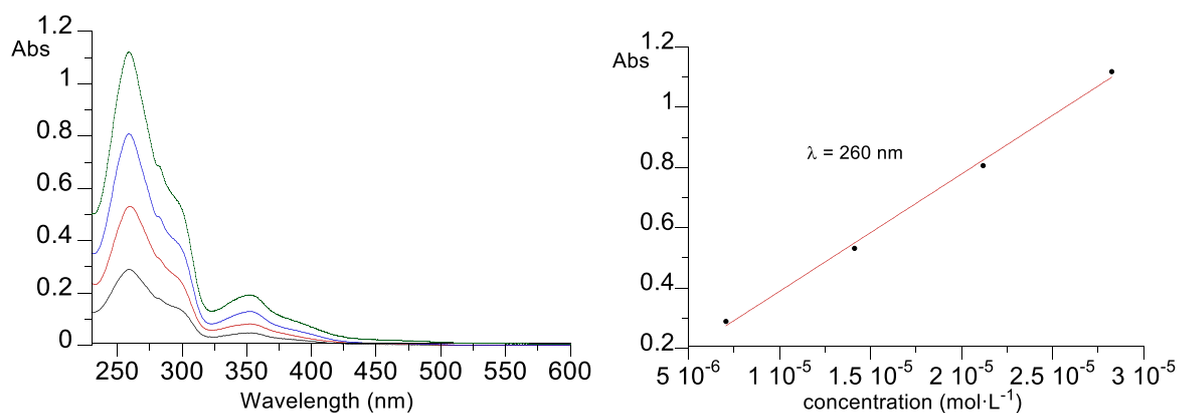
**Figure S14.** Clockwise: electronic absorption spectra for **1d** in  $\text{CH}_2\text{Cl}_2$  for four concentrations, determination of molar extinction coefficient  $\epsilon$  at  $\lambda = 245.0$  nm (best fit function:  $\epsilon = 40831(504) \times \text{conc}$ ,  $r^2 = 0.997$ ), molar extinction  $\log(\epsilon)$  plot.

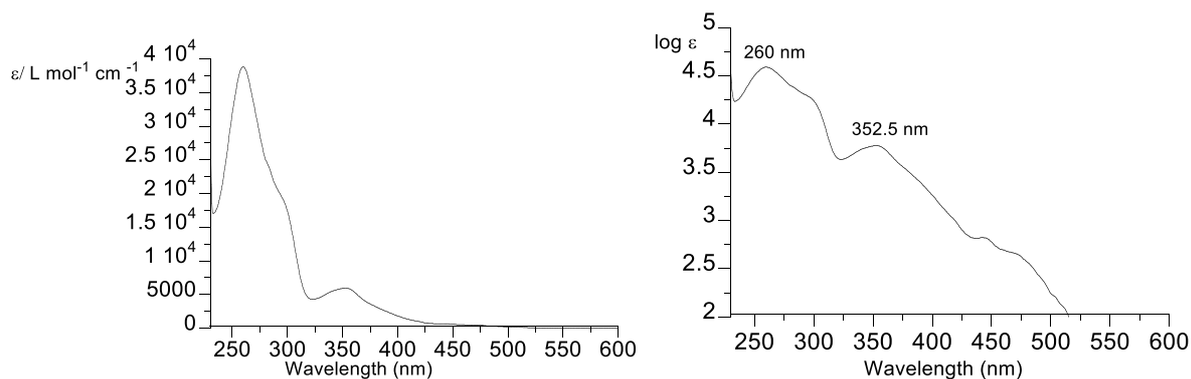


**Figure S15.** Clockwise: electronic absorption spectra for **1c-oxo** in  $\text{CH}_2\text{Cl}_2$  for four concentrations, determination of molar extinction coefficient  $\epsilon$  at  $\lambda = 276.0$  nm (best fit function:  $\epsilon = 16030(148) \times \text{conc}$ ,  $r^2 = 0.998$ ), molar extinction  $\log(\epsilon)$  plot.

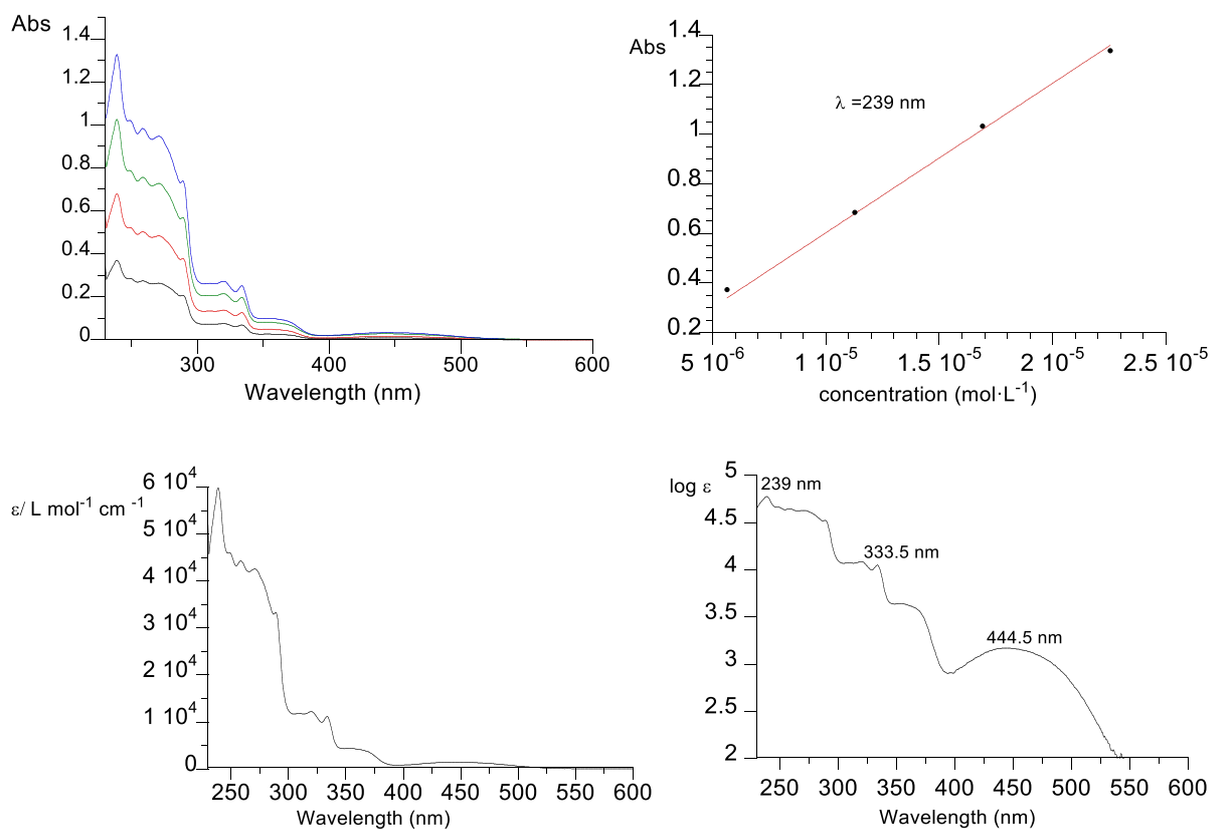


**Figure S16.** Clockwise: electronic absorption spectra for **2a** in  $\text{CH}_2\text{Cl}_2$  for four concentrations, determination of molar extinction coefficient  $\epsilon$  at  $\lambda = 261.5 \text{ nm}$  (best fit function:  $\epsilon = 43285(445) \times \text{conc}$ ,  $r^2 = 0.999$ ), molar extinction  $\log(\epsilon)$  plot.

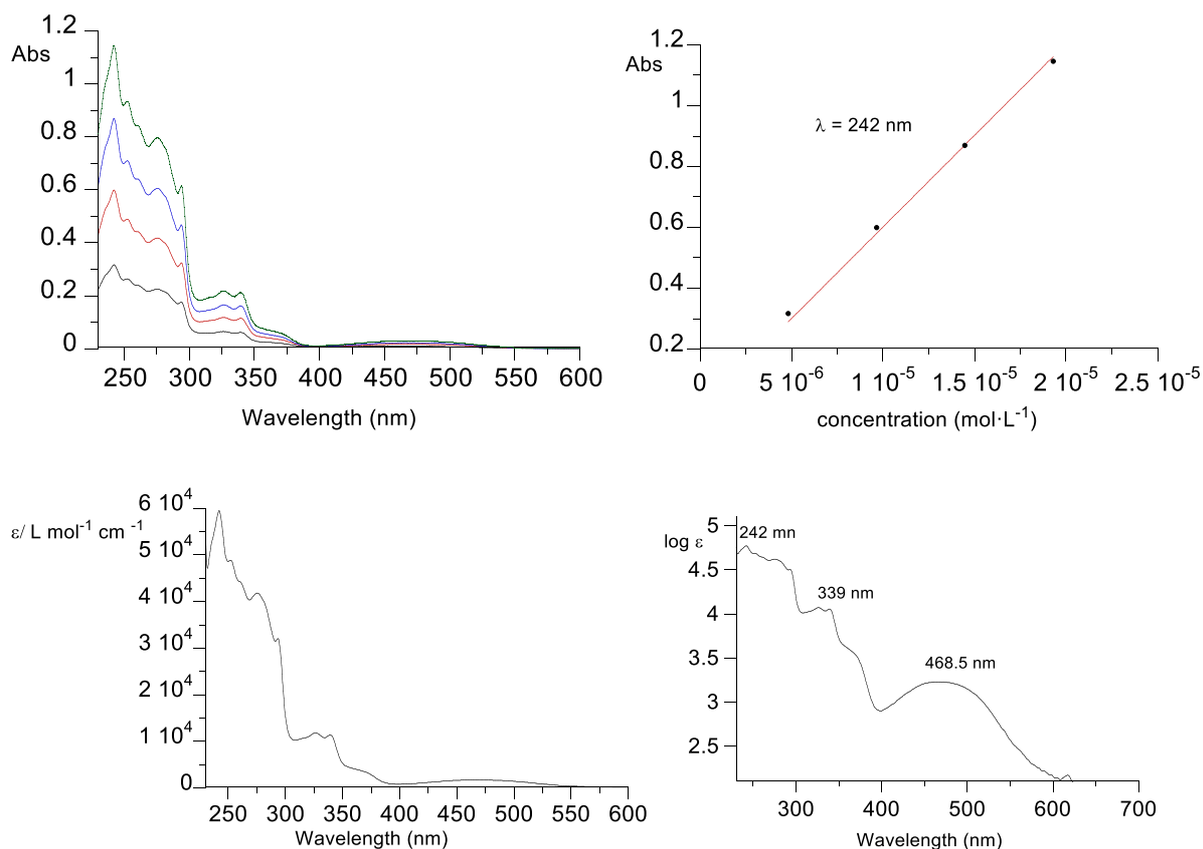




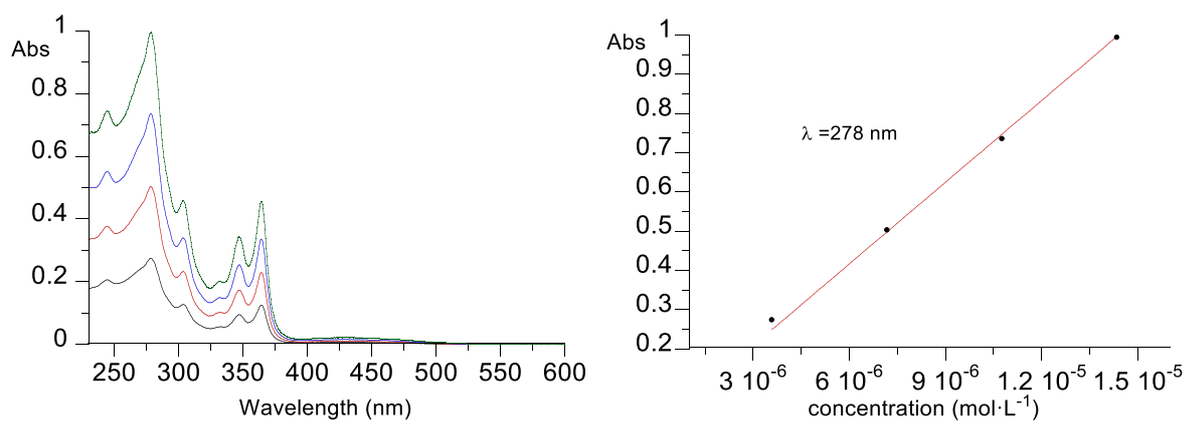
**Figure S17.** Clockwise: electronic absorption spectra for **2b** in  $\text{CH}_2\text{Cl}_2$  for four concentrations, determination of molar extinction coefficient  $\epsilon$  at  $\lambda = 260.0$  nm (best fit function:  $\epsilon = 38896(529) \times \text{conc}$ ,  $r^2 = 0.9984$ ), molar extinction  $\log(\epsilon)$  plot.

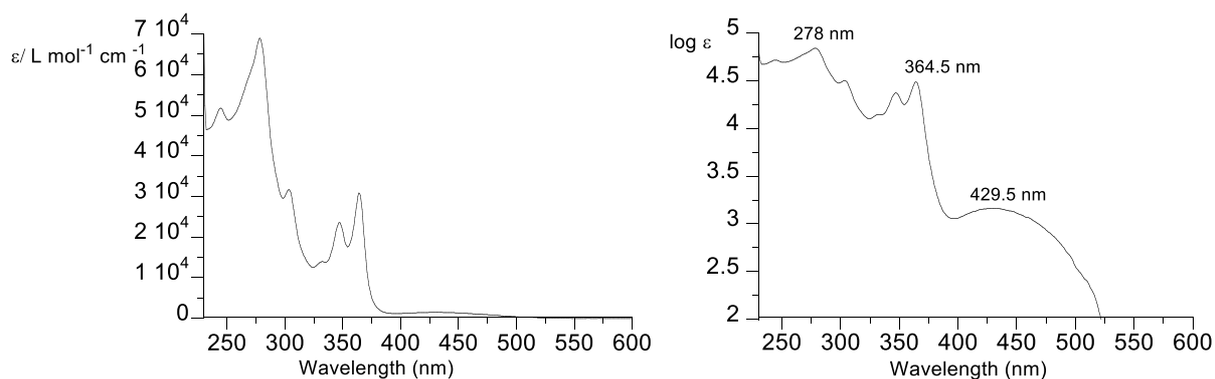


**Figure S18.** Clockwise: electronic absorption spectra for **2c** in  $\text{CH}_2\text{Cl}_2$  for four concentrations, determination of molar extinction coefficient  $\epsilon$  at  $\lambda = 239.0$  nm (best fit function:  $\epsilon = 60247(792) \times \text{conc}$ ,  $r^2 = 0.9983$ ), molar extinction  $\log(\epsilon)$  plot.

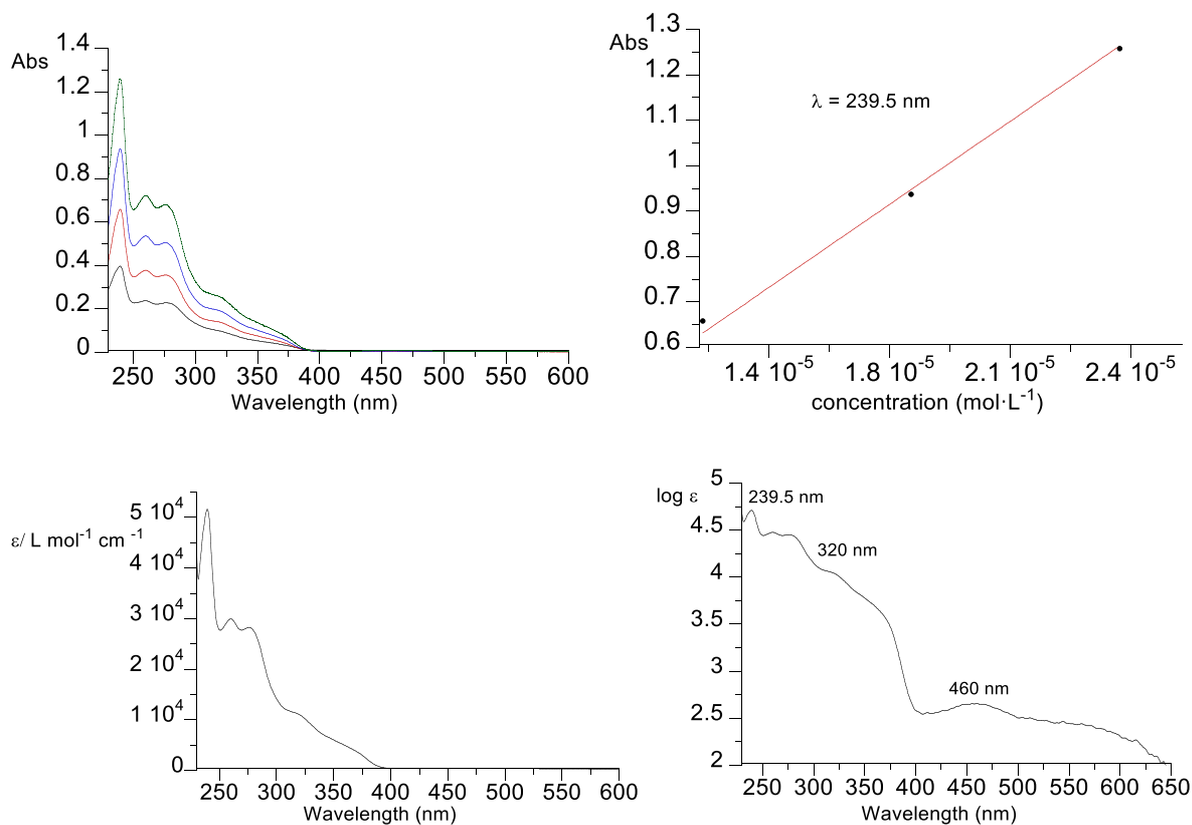


**Figure S19.** Clockwise: electronic absorption spectra for **2d** in  $\text{CH}_2\text{Cl}_2$  for four concentrations, determination of molar extinction coefficient  $\epsilon$  at  $\lambda = 242.0 \text{ nm}$  (best fit function:  $\epsilon = 60138(794) \times \text{conc}$ ,  $r^2 = 0.998$ ), molar extinction  $\log(\epsilon)$  plot.





**Figure S20.** Clockwise: electronic absorption spectra for **2e** in  $\text{CH}_2\text{Cl}_2$  for four concentrations, determination of molar extinction coefficient  $\epsilon$  at  $\lambda = 278.0$  nm (best fit function:  $\epsilon = 69413(830) \times \text{conc}$ ,  $r^2 = 0.9986$ ), molar extinction  $\log(\epsilon)$  plot.

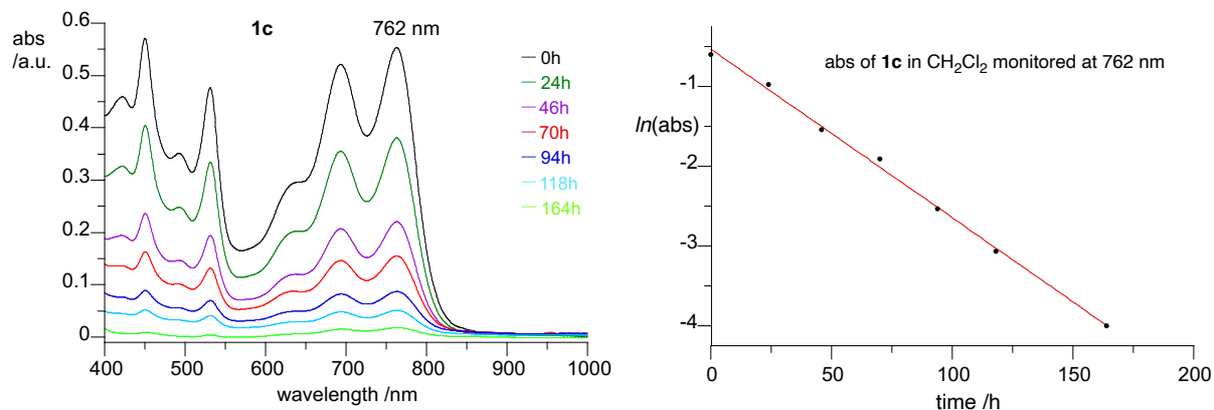


**Figure S21.** Clockwise: electronic absorption spectra for **2f** in  $\text{CH}_2\text{Cl}_2$  for four concentrations, determination of molar extinction coefficient  $\epsilon$  at  $\lambda = 239.5$  nm (best fit function:  $\epsilon = 52258(636) \times \text{conc}$ ,  $r^2 = 0.998$ ), molar extinction  $\log(\epsilon)$  plot.

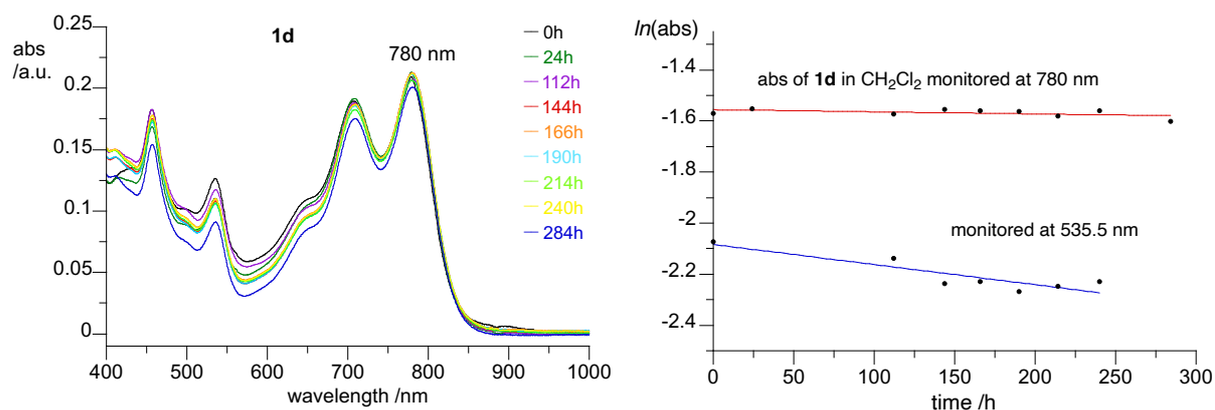
## 6. Stability of radicals **1c** and **1d** towards atmospheric oxygen

Oxidative stability of radicals **1c** and **1d** was investigated in  $\text{CH}_2\text{Cl}_2$  solutions in a quartz cuvette and the absorbance at 762 nm (for **1c**) and 780 nm (for **1d**) was measured periodically. The cuvettes were filled up to 2/3 of the volume with the solution and open daily

to exchange air atmosphere. If needed, fresh solvent was added to maintain the original volume of the solution before each measurement. Results are shown in Figures S22 and S23.



**Figure S22.** Left: electronic absorption spectra measured for radical **1c** in  $\text{CH}_2\text{Cl}_2$  solution exposed to air at several time intervals at 273 K. Right: first-order decay of the absorption peak at 762 nm. Best fit line:  $\ln(\text{abs}) = -0.53(4) - 0.0211(5) \times t$ ,  $r^2 = 0.997$ .

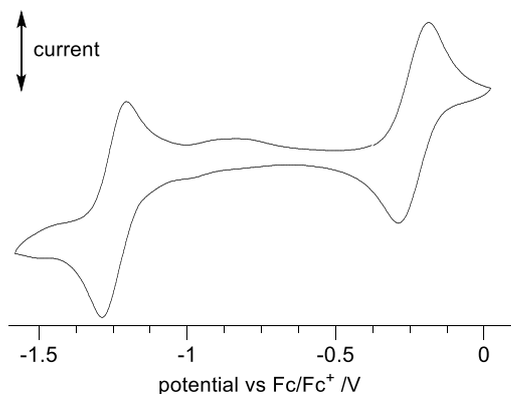


**Figure S23.** Left: electronic absorption spectra measured for radical **1d** in  $\text{CH}_2\text{Cl}_2$  solution exposed to air at several time intervals at 273 K. Right: first-order decay of the absorption peak at 780 nm (red) and 535.5 nm (blue). Best fit lines:  $\ln(\text{abs}_{780}) = -1.556 - 8.38 \times 10^{-5} \times t$ ,  $r^2 = 0.28$ , and  $\ln(\text{abs}_{535}) = -2.08 - 7.9 \times 10^{-4} \times t$ ,  $r^2 = 0.79$ .

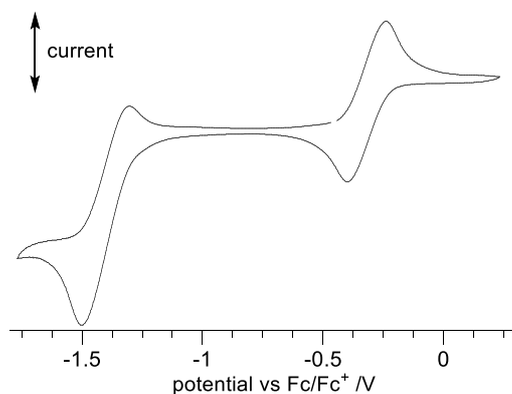
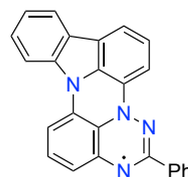
## 7. Electrochemical results

The electrochemical characterization of selected radicals was conducted using Autolab PGSTAT128N potentiostat/galvanostat instrument in dry and degassed  $\text{CH}_2\text{Cl}_2$  (concentration 0.5 mM) in the presence of  $[n\text{-Bu}_4\text{N}]^+[\text{PF}_6]^-$  as an electrolyte (concentration 50 mM) using glassy carbon as the working electrode and Ag/AgCl as the reference electrode with a scan rate of  $50 \text{ mV s}^{-1}$  at *ca.* 20 °C. In the end of each measurement ferrocene was added and the peak potentials were referenced to the  $\text{Fc}/\text{Fc}^+$  couple (0.46 V vs SCE).<sup>6</sup>

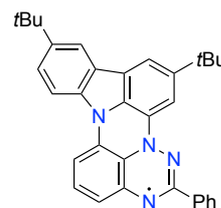
Cyclic voltammetry (CV) plots are shown in Figures S24 and S25 and numerical result are shown in Table S4. A correlation of  $E_{1/2}^{0/+1}$  with Hammett parameters<sup>7</sup> for model substituents is shown in Figure S26.



**Figure S24.** Cyclic voltammogram for **1c**.



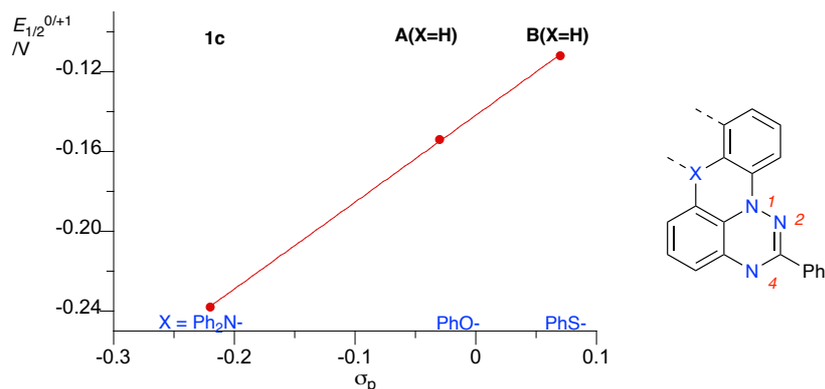
**Figure S25.** Cyclic voltammogram for **1d**.



**Table S4.** Cyclic voltammetry data for selected radicals in  $\text{CH}_2\text{Cl}_2$ .<sup>a</sup>

Radical	$E_{1/2}^{-1/0}$ /V	$E_{1/2}^{0/+1}$ /V	$E_{\text{cell}}$ /V <sup>a</sup>
<b>A(X=H)</b> <sup>b</sup>	-1.317	-0.154	1.163
<b>B(X=H)</b> <sup>b</sup>	-1.202	-0.112	1.090
<b>1c</b>	-1.243	-0.238	1.005
<b>1d</b>	-1.404	-0.320	1.085

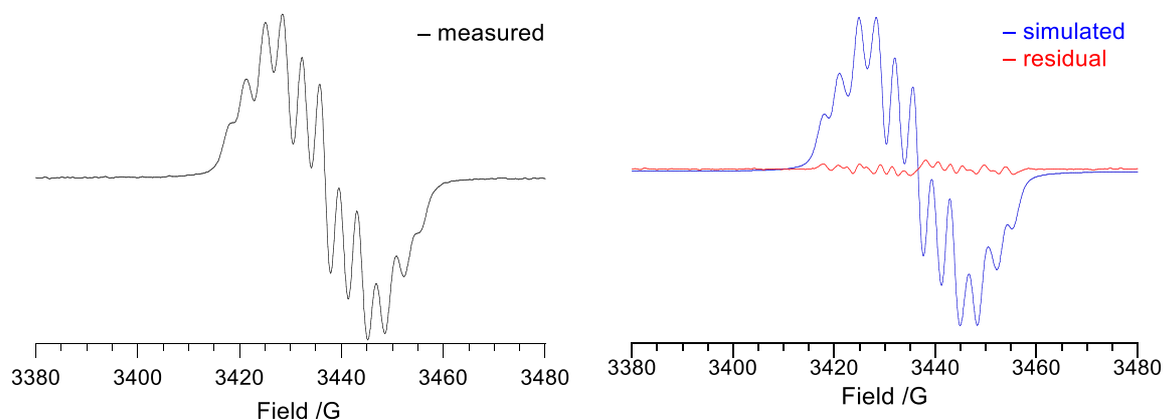
<sup>a</sup> Potential vs the Fc/Fc<sup>+</sup> couple. <sup>b</sup>Ref.<sup>8</sup>



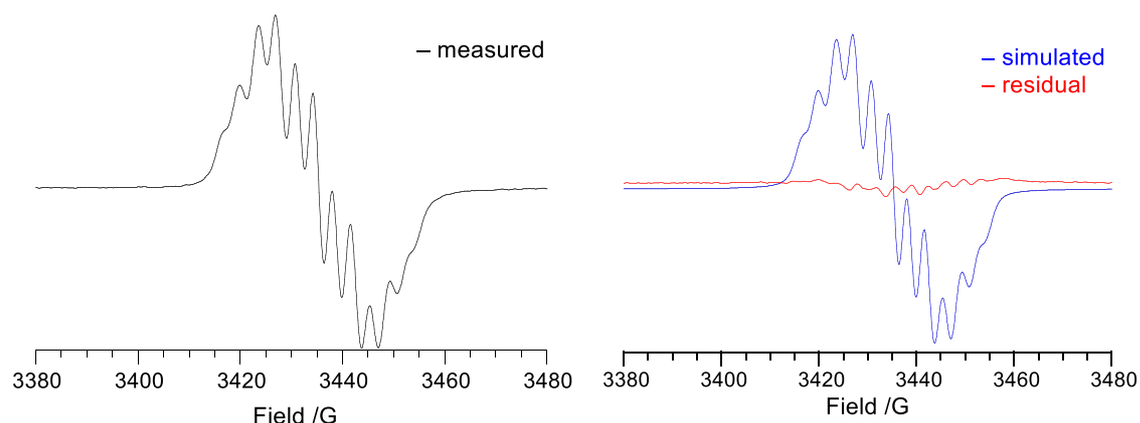
**Figure S26.** Correlation of  $E_{1/2}^{0/+1}$  with Hammett parameter  $\sigma_p$  for model substituents X (blue). Best fitting functions:  $a_{N1} = 0.435(6) \times E_{1/2}^{0/+1} - 0.142(1)$ ,  $r^2 = 0.999$ .

## 8. EPR spectroscopy

EPR spectra for radicals **1c** and **1d** were recorded on an X-band EMX-Nano EPR spectrometer at room temperature in dilute and degassed solutions in benzene. The microwave power was set using the Power Sweep program below the saturation of the signal, modulation frequency of 100 kHz, modulation amplitude of 0.5 G<sub>pp</sub> and spectral width of 100 G. Accurate g-values were obtained using TEMPO as EMX-Nano internal standard. Simulations of the spectra were performed with *Easy Spin* (Matlab) using DFT results (*vide infra*) as the starting point including all nitrogen and 6 hydrogen atoms. The resulting *hfcc* values were perturbed several times until a global minimum for the fit was achieved. For consistency, the previously reported<sup>9</sup> spectrum for **A(X=H)** and a newly recorded spectrum of **B(X=H)** were resimulated using the same parameters. Experimental and simulated spectra for **1c** and **1d** are shown in Figures S27 and S28 and resulting *hfcc* are listed in Table S5.



**Figure S27.** Experimental (black, left), simulated (blue, right) and difference (red, right) spectra for **1c** recorded in benzene at *ca* 20 °C.



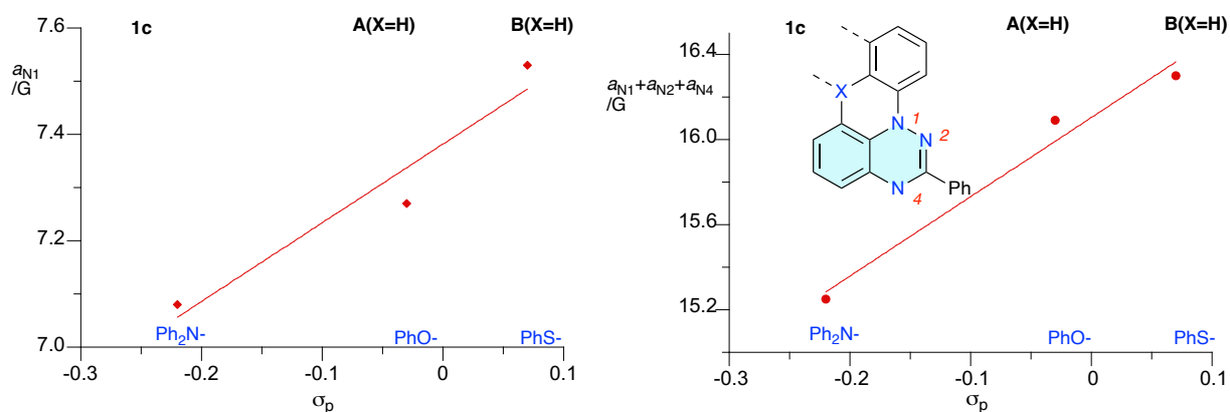
**Figure S28.** Experimental (black, left), simulated (blue, right) and difference (red, right) spectra for **1d** recorded in benzene at *ca* 20 °C.

**Table S5.** Experimental hyperfine coupling constants (G) for selected radicals in benzene at *ca.* 20 °C.

atom	A (X=H) <sup>a</sup>	B (X=H)	1c	1d
$a_{N(11)}$	7.53	7.27	7.08	7.14
$a_{N(1)}$	4.35	4.39	4.15	4.00
$a_{N(3)}$	4.42	4.43	4.02	4.00
$a_N$	-	-	0.35	0.36
$a_H$	1.88	1.39	2.77	2.79
$a_H$	1.68	1.26	2.31	2.47
$a_H$	1.09	1.21	0.34	0.36
$a_H$	1.07	1.10	0.32	0.35
$a_H$	1.07	1.10	0.29	0.34
$a_H$	1.07	0.75	0.29	0.34
$g$	2.0026	2.0040	2.0036	2.0044

<sup>a</sup> Experimental data for simulation taken from ref<sup>9</sup>

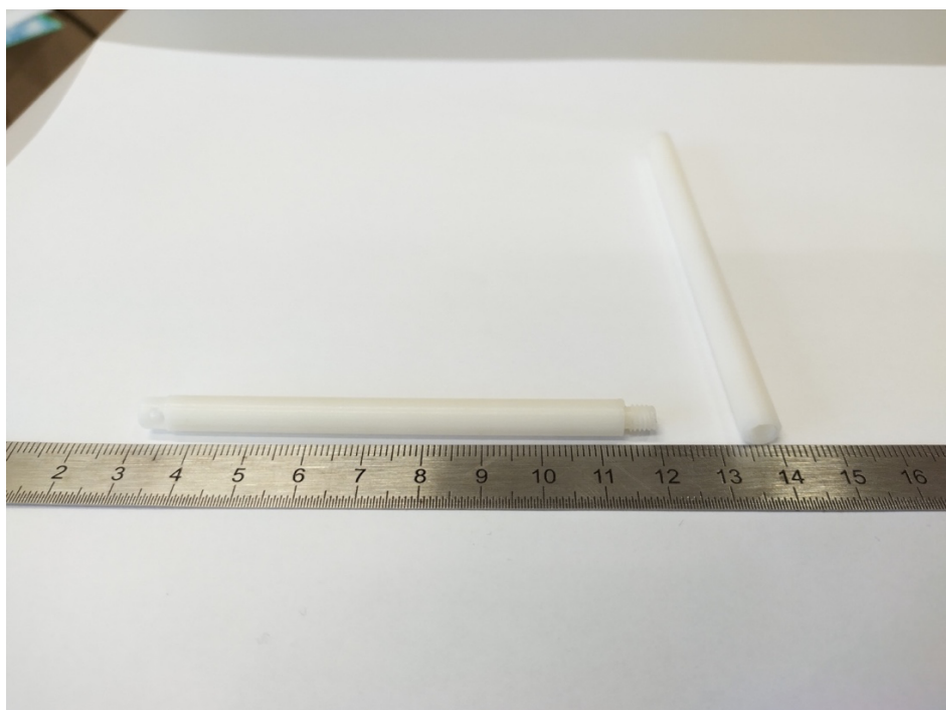
Figure S29 shows correlations of selected hfcc with Hammett parameters<sup>7</sup> for model substituents.



**Figure S29.** Correlation of  $a_{N1}$  hfcc (left) and sum of  $a_N$  (right) with Hammett parameter  $\sigma_p$  for model substituents (blue). Best fitting functions:  $a_{N1} = 1.5(4) \times \sigma_p + 7.38(5)$ ,  $r^2 = 0.930$ ;  $\sum a_N = 3.7(6) \times \sigma_p + 16.10(8)$ ,  $r^2 = 0.976$ .

## 9. SQUID magnetometry

Magnetic susceptibility of polycrystalline sample of radical **1d** was measured in a paramagnetic cavity of known background (a two-part Delrin holder, Figure S30) as a function of temperature in the cooling mode (300 K  $\rightarrow$  2 K) with a sweep rate of 0.5 K min<sup>-1</sup> at 0.6 T, using a SQUID magnetometer (Quantum Design MPMS-XL-7T). Measurements of magnetization  $M$  vs  $H$  for **1d** were conducted at 2 K for several magnetic field strengths (0.05–7 T).



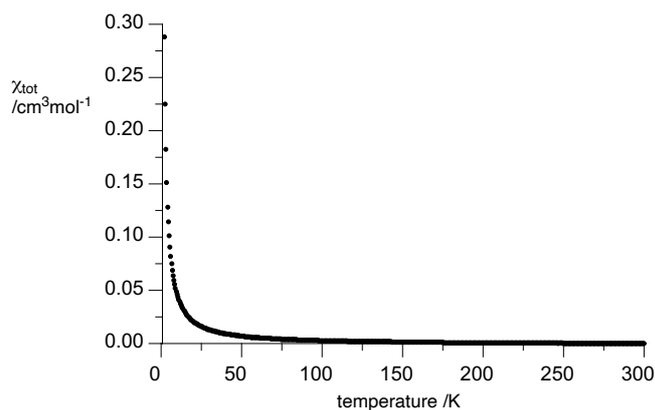
**Figure S30.** Delrin sample holder used for SQUID measurements.

### Raw data workup

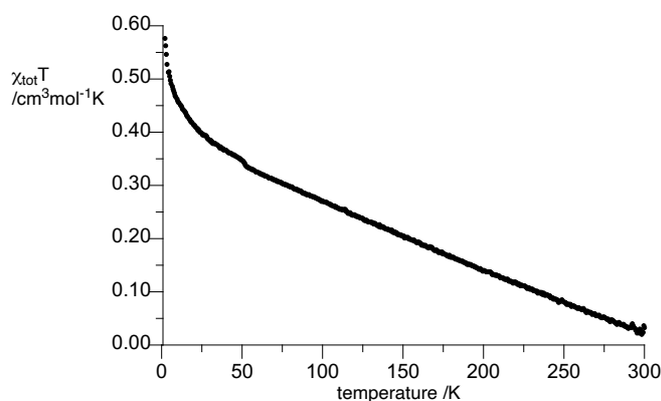
The magnetic contribution of the sample holder was removed numerically from the raw data (voltage vs sample position,  $U(x)$ ) by subtracting the  $U(x)$  contribution for the empty holder in the entire temperature range. The holder brings a large, temperature-constant paramagnetic component.

### Analysis for mono-radical **1d**

A microcrystalline sample of derivative **1d** ( $m = 8.53$  mg,  $1.764 \times 10^{-5}$  mol,  $M_w = 483.64$  g mol<sup>-1</sup>) was analyzed at 0.6 T in the 300–2 K temperature range. Total molar magnetic susceptibility  $\chi_{\text{tot}}(T)$  and  $\chi_{\text{tot}}T(T)$  plots are shown in Figures S31 and S32, respectively.



**Figure S31.**  $\chi_{\text{tot}}$  vs  $T$  plot for **1d**.



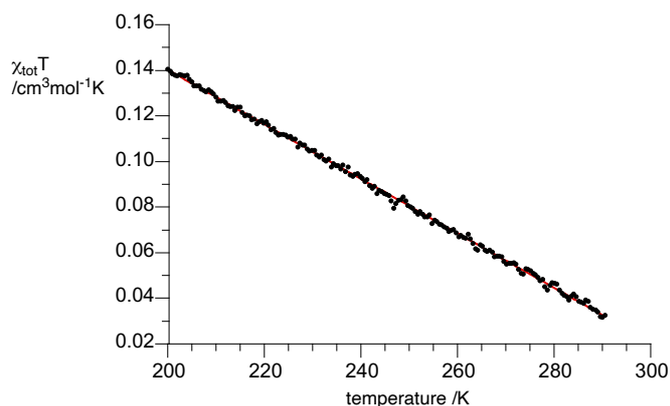
**Figure S32.**  $\chi_{\text{tot}}T$  vs  $T$  plot for **1d**.

The diamagnetic correction for the sample was estimated from the linear portion of high temperature plot  $\chi_{\text{tot}} \cdot T$  vs  $T$  assuming ideal paramagnetic behavior of the sample using the Curie law (eq S1).

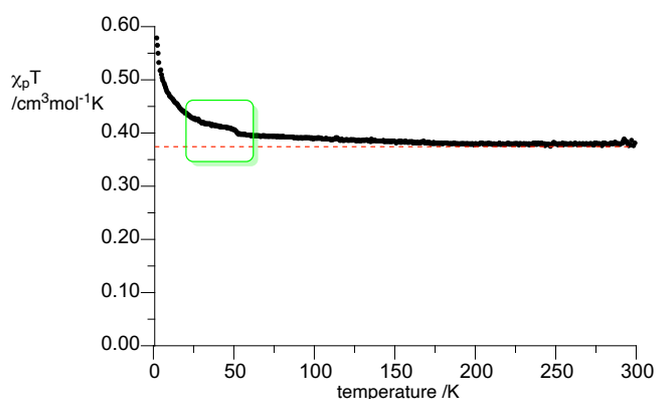
$$\chi_{\text{tot}} \cdot T = (\chi_p + \chi_{\text{dia}}) \cdot T = C + \chi_{\text{dia}} \cdot T \quad (\text{eq S1})$$

where  $C = 0.375 \text{ cm}^3 \text{ mol}^{-1} \text{ K}$  for an ideal paramagnet ( $S = 1/2$ ).

Fitting the high temperature portion of the data gave the diamagnetic susceptibility  $\chi_d$  of  $-11.95(3) \times 10^{-4} \text{ cm}^3 \text{ mol}^{-1}$  (Figure S33). The total magnetic susceptibility  $\chi_{\text{tot}}$  was corrected for  $\chi_d$  and the resulting paramagnetic susceptibility is plotted as  $\chi_p T(T)$  in Figure S34. The observed bump on the  $\chi_p T(T)$  plot at about 50 K is attributed to the presence of small amounts of oxygen (air) in the holder's closed cavity.



**Figure S33.** High temperature portion of the  $\chi_{\text{tot}}T$  vs  $T$  plot for **1d**. The best fitting function:  $\chi_{\text{tot}}T = -0.001196(3) \times T + 0.380(1)$ ,  $r^2 = 0.999$ .



**Figure S34.**  $\chi_p T$  vs  $T$  plot for **1d** after diamagnetic correction  $\chi_d$  of  $-11.96 \times 10^{-4} \text{ cm}^3\text{mol}^{-1}$ . The horizontal red line marks the Curie value of  $0.375 \text{ Kcm}^3\text{mol}^{-1}$  for an ideal paramagnet. The green box marks the feature related to traces of molecular oxygen trapped in the holder cavity with the sample.

The feature in the range 30–50 K related to molecular oxygen was removed from further analysis.

### Analysis for radical dimer **1d**

The same experimental magnetization data was analyzed as dimer of two spins. Thus, a microcrystalline sample of derivative **1d** ( $m = 8.53 \text{ mg}$ ,  $0.882 \times 10^{-5} \text{ mol}$ ,  $M_w = 967.28 \text{ g mol}^{-1}$  per dimer) was analyzed at 0.6 T in the 300–2 K temperature range at 0.6 T. The total molar magnetic susceptibility  $\chi_{\text{tot}}(T)$  and  $\chi_{\text{tot}}T(T)$  plots are shown in Figures S31 and S32, respectively (*vide supra*).

Considering the discrete dimer structure of **1d** in the solid-state, the magnetic data was

analyzed using the Bleaney-Bowers model<sup>10</sup> (the BB model) and the BB model with a mean field approximation based on the Hamiltonian:  $\mathcal{H} = -2JS_1S_2$ ,<sup>11</sup> for two 1/2 spins.

The Bleaney-Bowers model for two interacting spins is shown in eq S2.

$$\chi_{BB}(T) = \frac{N_A g \mu_B}{B} \left( \frac{\sinh\left(\frac{g \mu_B B}{k_B T}\right)}{1 + \exp\left(-\frac{2J}{k_B T}\right) + 2 \cosh\left(\frac{g \mu_B B}{k_B T}\right)} \right)$$

For low fields it reduces to:  $\chi_{BB}(T) = \frac{N_A g^2 \mu_B^2}{k_B T \left(3 + \exp\left(-\frac{2J}{k_B T}\right)\right)}$  (eq S2)

where  $N_A$ -Avogadro number,  $g$ -electron g-factor,  $\mu_B$ -Bohr magneton (in CGS units),  $k_B$  – Boltzmann constant,  $S$  – electron spin,  $B$  – magnetic field (in T),  $K$  – temperature (K),  $J$  – exchange integral.

Since  $N_A g^2 \mu_B^2 / k_B = 1.504 \text{ K cm}^3/\text{mol}$ , eq S2 gives eq S3 after substitution:

$$\chi_{BB} = \frac{1.504}{T \left(3 + \exp\left(-\frac{2J}{k_B T}\right)\right)} \text{ or } \chi_{BB} T = \frac{1.504}{\left(3 + \exp\left(-\frac{2J}{k_B T}\right)\right)} \quad (\text{eq S3})$$

and with the diamagnetic component it gives eq S4:

$$\chi_{tot} T = \chi_{BB} T + \chi_{dia} T = \frac{1.504}{\left(3 + \exp\left(-\frac{2J}{k_B T}\right)\right)} + \chi_{dia} T \quad (\text{eq S4})$$

The correction for a mean field (the  $zJ'$  model),<sup>11</sup> which accounts for interdimer interactions gives eq S5:

$$\chi' T = \frac{\chi T}{1 - \left(\frac{zJ'}{N_A g^2 \mu_B^2}\right) \chi} = \frac{\chi T}{1 - \left(\frac{zJ'}{k_B}\right) \left(\frac{k_B}{N_A g^2 \mu_B^2}\right) \chi} \quad (\text{eq S5})$$

Where  $\chi$ -molar susceptibility for a given model,  $zJ'/k_B$ - interaction parameter between the nearest neighbor magnetic species in K, and  $N_A g^2 \mu_B^2 / k_B = 1.504 \text{ K cm}^3/\text{mol}$ . For the Bleaney-Bowers model<sup>10</sup> (BB model) the mean field approximation takes the form of equation S6:

$$\chi' T = \frac{\chi_{BB} T}{1 - \left(\frac{zJ'}{1.504 \times k_B}\right) \chi_{BB}} \quad (\text{eq S6})$$

After inclusion of the diamagnetic correction,  $\chi_{dia}$ , the total magnetization  $\chi_{tot}$  can be expressed as eq S7:

$$\chi'_{tot} T = \frac{\chi_{BB} T}{1 - \left(\frac{zJ'}{1.504 \times k_B}\right) \chi_{BB}} + \chi_{dia} T \quad (\text{eq S7})$$

Which gives the fitting function eq S8:

$$\chi'_{tot}T = \frac{1.504}{3 + \exp\left(\frac{-m_1}{M_0}\right)} \times \frac{1}{1 - \left(\frac{m_2}{1.504}\right) \times 1.504 / (3 + \exp\left(\frac{-m_1}{M_0}\right))} + m_3T \quad (\text{eq S8})$$

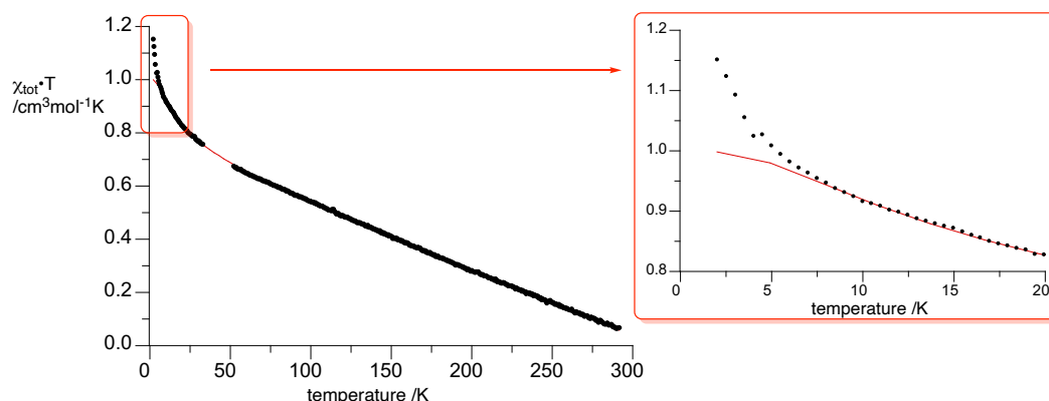
where

$$m_1 = 2J/k_B$$

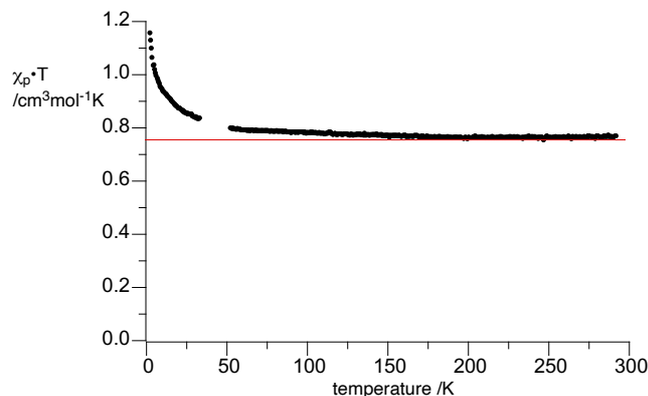
$$m_2 = zJ'/k_B$$

$$m_3 = \chi_{\text{dia}}$$

The total molar magnetic susceptibility data,  $\chi_{\text{tot}}$ , was fitted to the pure BB model containing the diamagnetic correction,  $\chi_{\text{dia}}$  (eq S4, Figure S35). Expansion of the low temperature section of the plot shows discrepancy between the data and the model. The resulting  $\chi_{\text{dia}}$  was used to obtain the paramagnetic susceptibility  $\chi_p$ , and the  $\chi_p T$  vs  $T$  plot is shown in Figure S36.

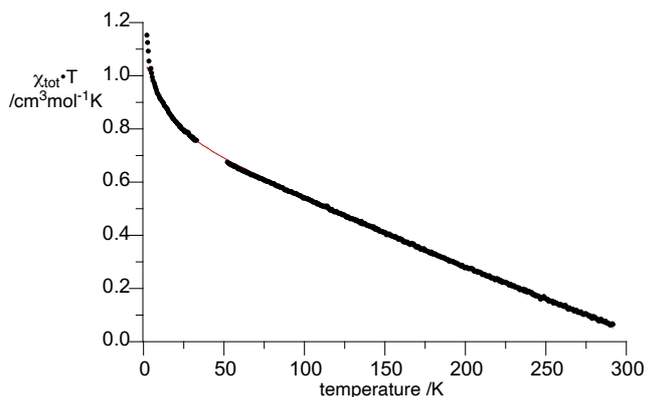


**Figure S35.** Left:  $\chi_{\text{tot}}T$  vs  $T$  plot for dimeric **1d** and fitting to the Bleaney-Bowers model (eq. S4). Fitting parameters:  $2J/k_B = 16.6(2)$  K and  $\chi_{\text{dia}} = -0.002418(3)$   $\text{cm}^3\text{mol}^{-1}$ ;  $r^2 = 0.998$ . Right: expanded low temperature portion of the plot.



**Figure S36.**  $\chi_p T$  vs  $T$  plot for dimer **1d** plot after applying the diamagnetic correction  $\chi_{\text{dia}} = -0.002418(3)$   $\text{cm}^3\text{mol}^{-1}$ . The horizontal red line marks the Curie value of  $0.75$   $\text{Kcm}^3\text{mol}^{-1}$  for two ideal spins  $S = 1/2$ .

Fitting the  $\chi_{\text{tot}}$  data to the BB model with mean field approximation (eq S8) is shown in Figure S37.



**Figure S37.**  $\chi_{\text{tot}}T$  vs  $T$  plot for dimer **1d** plot and fitting to the Bleaney-Bowers model with mean field approximation (eq. S8). Fitting parameters:  $2J/k_B = 12.0(4)$  K,  $zJ'/k_B = 0.10(1)$  K, and  $\chi_{\text{dia}} = 0.002496(7)$   $\text{cm}^3 \text{mol}^{-1}$ ;  $r^2 = 0.998$ .

Results demonstrate that the pure BB model describes well magnetic behavior of the analyzed solid-state sample of **1d**, which indicates that the sole exchange interaction is between molecules in the dimer and essentially negligible between the dimers above 8 K.

#### Magnetization $M$ vs magnetic field $B$ for monomeric radical **1d**

Paramagnetic susceptibility is described by the Brillouin model (the B model, eq S9):

$$\chi_p(T, B) = n_s \frac{N_A g \mu_B}{B} \left( \frac{(S+0.5)}{\tanh((S+0.5)g\mu_B B/k_B T)} - \frac{0.5}{\tanh(0.5g\mu_B B/k_B T)} \right) \quad \text{eq S9}$$

where  $N_A$ -Avogadro number,  $g$ -electron g-factor,  $\mu_B$ -Bohr magneton (in CGS units),  $k_B$ -Boltzmann constant,  $S$ -electron spin,  $B$  – magnetic field (in T),  $T$  – temperature (K), and  $n_s$  – fraction of units with spin  $S$  in 1 mol.

Combined constants give:

$$\frac{g\mu_B}{k_B} = 1.3450 \text{ K/T} \quad \text{and} \quad \mu_B N_A g = 11183.45 \text{ cm}^3 \text{ G/mol (or emu/mol)}$$

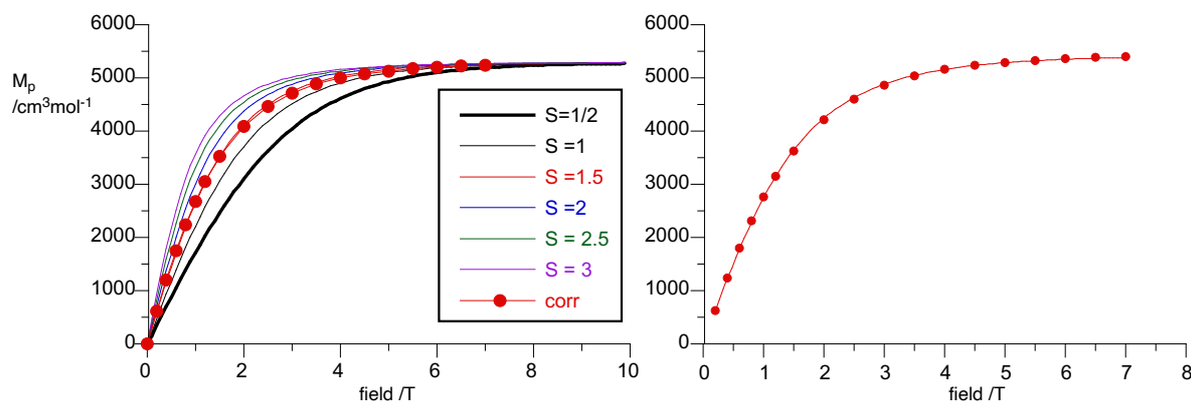
Considering that  $M_p = \chi_p \times B$  substitution gives eq S10:

$$M_p = n_s \times 11183.45 \times \left( \frac{(S+0.5)}{\tanh\left(\frac{(S+0.5) \cdot 1.3450 \cdot B}{T}\right)} - \frac{0.5}{\tanh\left(\frac{0.67248 \cdot B}{T}\right)} \right) \quad \text{(eq S10)}$$

For  $n_s$  to express the number of active spins in the material per 1 mol, eq S9 is divided by the number of spins in the fundamental spin unit  $S$ , which leads to eq S10:

$$M_p = \frac{n_s}{2S} \times 11183.45 \times \left( \frac{(S+0.5)}{\tanh\left(\frac{(S+0.5) \cdot 1.3450 \cdot B}{T}\right)} - \frac{0.5}{\tanh\left(\frac{0.67248 \cdot B}{T}\right)} \right) \quad \text{(eq S11)}$$

The fitting of the experimental data to equation S11 is shown in Figure S38.



**Figure S38.** Left:  $M_p$  vs  $B$  experimental data for monoradical **1d** (red dots) and Brillouin function plots for several  $S$  values. Right:  $M_p$  vs  $B$  plot for monoradical **1d** (red dots). Numerical fitting to eq S11 gave  $S = 1.56(1)$ ,  $n_s = 0.968(1)$ ,  $r^2 = 0.9999$ . Measurement performed at  $T = 2$  K.

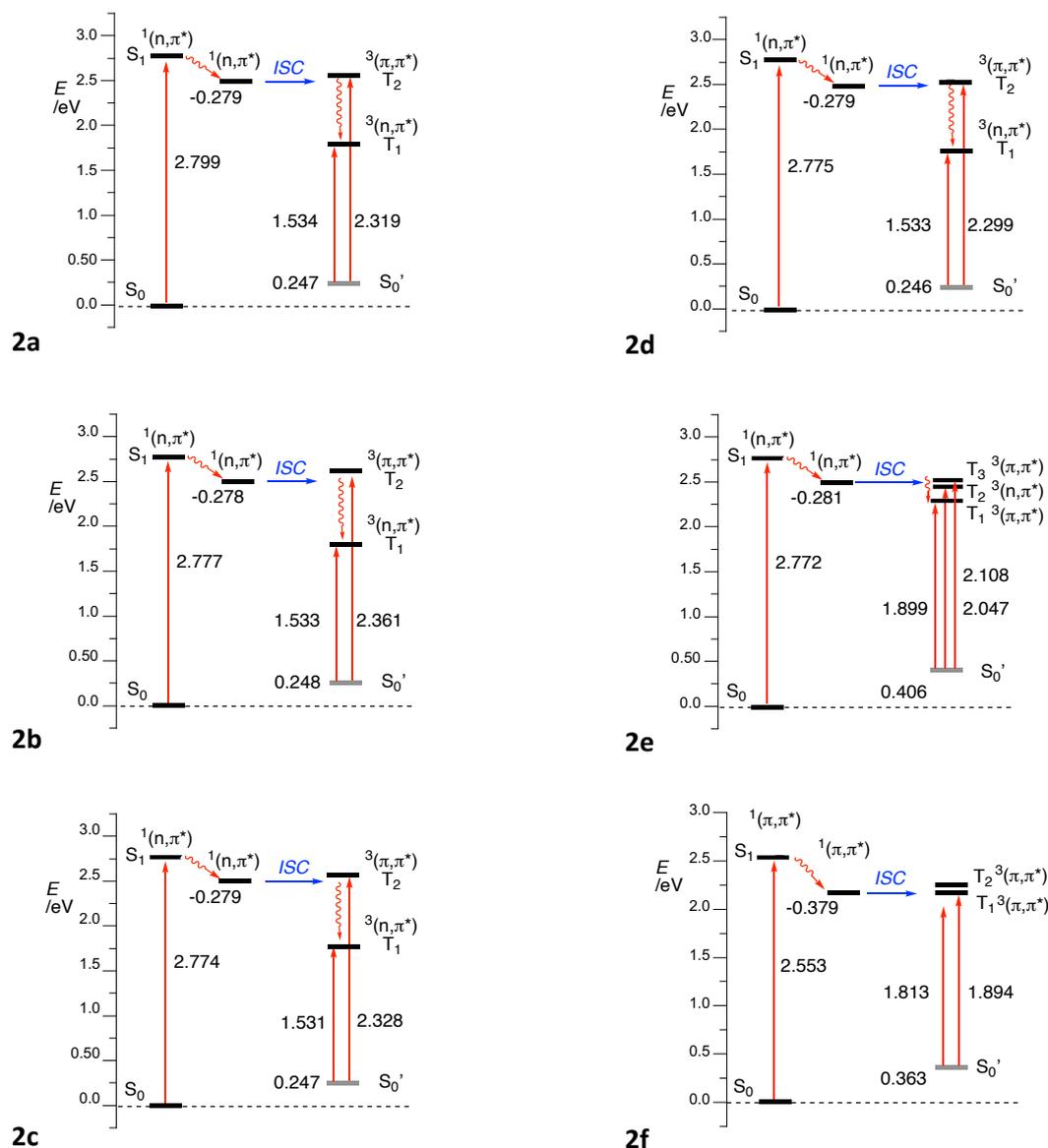
## 10. Computational details and results

Quantum-mechanical calculations were carried out using Gaussian 16 suite of programs.<sup>12</sup> Geometry optimizations of the precursors **2** were conducted at the CAM-B3LYP/6-311G(d,p) level of theory in ethyl acetate dielectric medium (PCM model<sup>13</sup>) requested with the SCRF(Solvent=EthylEthanoate) keyword and using tight convergence limits and without symmetry constrains. Geometries of radicals **1** and reference radicals **A(X=H)** and **B(X=H)** were optimized at the UB3LYP/6-311G(d,p) level of theory in vacuum using tight convergence limits and without symmetry constrains.

### *a) mechanistic investigation of photocyclization of 2*

Mechanistic investigation of photocyclization of model **2'** (in which t-Bu was approximated with Me) was conducted at the CAM-B3LYP/6-311G(d,p) level of theory in EtOAc dielectric medium (PCM model<sup>13</sup>) requested with the SCRF(Solvent=EthylEthanoate) keyword and tight convergence limits. Excitation calculations of **2'** were conducted using the TD-DFT method for closed-shell systems. Geometry optimization in the  $S_1$  state was performed using Fopt with default convergent limits and TD=(singlets, root=1, NStates=3) keywords in AcOEt dielectric medium (PCM model<sup>13</sup>).

The triplet state geometries of precursors **2** were obtained using the UCAM-B3LYP/6-311G(d,p) method and starting with GS geometries of **2**. TD-DFT calculations for closed-shell singlet at the triplet geometries using CAM-B3LYP/6-311G(d,p) method and TD=(triplets, root=1, NStates=12) keyword gave the forbidden  $S_0 \rightarrow T_n$  transitions. Partial Jablonski diagrams for **2a–2f** are shown in Figure S39.



**Figure S39.** Partial Jablonski diagrams for precursors **2a–2f**.

Natural charges for radical ions  $2^{+}$  and  $2^{-}$  were obtained by population analysis (pop=NBO) of the UCAM-B3LYP/6-311G+(d,p) wavefunction obtained by single point calculation of the radical ion generated by adding or subtracting an electron from precursors **2** at the optimized GS geometry (CAM-B3LYP/6-311G(d,p) level of theory in AcOEt dielectric medium) in AcOEt dielectric medium (PCM model).<sup>13</sup>

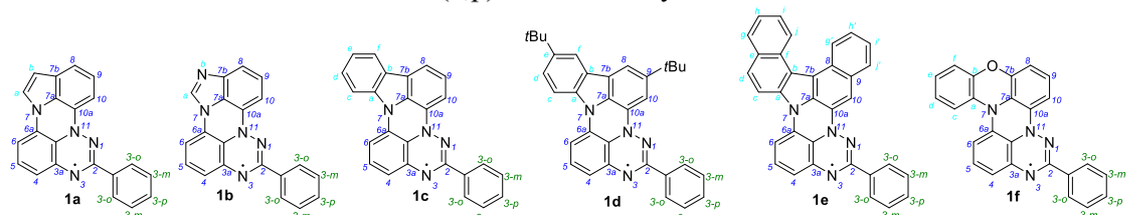
**b) isotropic Fermi contact coupling constants (hfcc) and spin densities for radicals**

Isotropic Fermi contact coupling constants for radicals **A**, **B**, **1** and carbazole substituted reference radicals were calculated using the UCAM-B3LYP/EPR-III // UB3LYP/6-311G(d,p) method in benzene dielectric medium requested with the SCRf(Solvent=Benzene) keywords (PCM model).<sup>13</sup> Fermi contact parameters for **B(X=H)** were obtained with the EPR-III basis

set for all atoms except for the sulfur atom, for which 6-311G+(2df) basis set was used implemented with the “gen” keyword. The resulting *hfcc* values are shown in Table S6 and spin densities are listed in Table S7 and S8.

Spin densities calculated for the triazine, benzo[*e*][1,2,4]triazine with the Ph ring at the N(1) position of the benzo[*e*][1,2,4]triazine ring were correlated with Hammett parameters<sup>7</sup> for model substituents and results are shown in Figure S40.

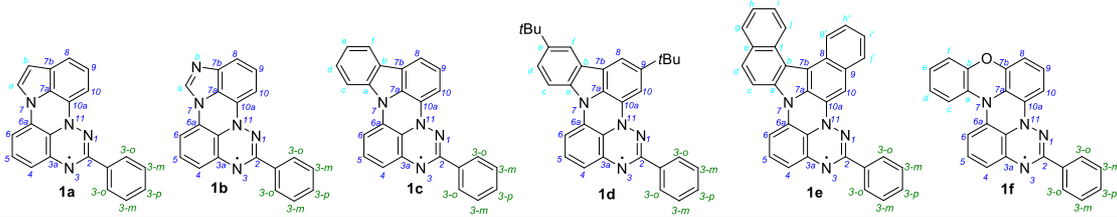
**Table S6.** DFT calculated hyperfine coupling constants (G) of radicals **1** in benzene at the UCAM-B3LYP/EPR-III // UB3LYP/6-311G(d,p) level of theory.<sup>a</sup>



<i>hfcc</i> /G	<b>1a</b>	<b>1b</b>	<b>1c</b>	<b>1d</b>	<b>1e</b>	<b>1f</b>
<i>a</i> <sub>N(1)</sub>	3.81	4.05	3.78	3.71	3.94	3.87
<i>a</i> <sub>N(3)</sub>	4.03	4.16	4.04	4.01	4.09	4.09
<i>a</i> <sub>N(11)</sub>	6.04	5.88	6.08	6.18	5.62	6.32
<i>a</i> <sub>N(7)</sub>	0.07	-0.09	0.39	0.41	0.04	0.77
<i>a</i> <sub>N(b)</sub>	-	0.29	-	-	-	-
<i>a</i> <sub>H(4)</sub>	-0.55	-0.89	-0.43	-0.37	-0.75	-0.31
<i>a</i> <sub>H(5)</sub>	-0.74	-0.53	-0.82	-0.88	-0.73	-0.89
<i>a</i> <sub>H(6)</sub>	-0.84	-1.26	-0.73	-0.65	-1.12	-0.65
<i>a</i> <sub>H(8)</sub>	-3.32	-3.10	-2.94	-2.92	-	-2.01
<i>a</i> <sub>H(9)</sub>	1.33	1.29	1.17	-	-	1.09
<i>a</i> <sub>H(10)</sub>	-3.62	-3.32	-3.15	-3.17	-4.50	-2.11
<i>a</i> <sub>H(a)</sub>	0.49	-0.03	-	-	-	-
<i>a</i> <sub>H(b)</sub>	-0.91	-	-	-	-	-
<i>a</i> <sub>H(c)</sub>	-	-	-0.29	-0.29	-0.18	-0.29
<i>a</i> <sub>H(d)</sub>	-	-	0.29	0.32	0.32	0.18
<i>a</i> <sub>H(e)</sub>	-	-	-0.32	-	-	-0.29
<i>a</i> <sub>H(f)</sub>	-	-	0.28	0.26	-	0.14
<i>a</i> <sub>H(g)</sub>	-	-	-	-	0.11	-
<i>a</i> <sub>H(g')</sub>	-	-	-	-	1.03	-
<i>a</i> <sub>H(h)</sub>	-	-	-	-	-0.12	-
<i>a</i> <sub>H(h')</sub>	-	-	-	-	-1.68	-
<i>a</i> <sub>H(i)</sub>	-	-	-	-	0.11	-
<i>a</i> <sub>H(i')</sub>	-	-	-	-	0.89	-
<i>a</i> <sub>H(j)</sub>	-	-	-	-	-0.10	-
<i>a</i> <sub>H(j')</sub>	-	-	-	-	-1.48	-
<i>a</i> <sub>H(3-o) avg</sub>	0.50	0.52	0.50	0.49	0.54	0.50
<i>a</i> <sub>H(3-m) avg</sub>	-0.29	-0.31	-0.30	-0.29	-0.32	-0.29
<i>a</i> <sub>H(3-p)</sub>	0.41	0.44	0.41	0.40	0.46	0.42

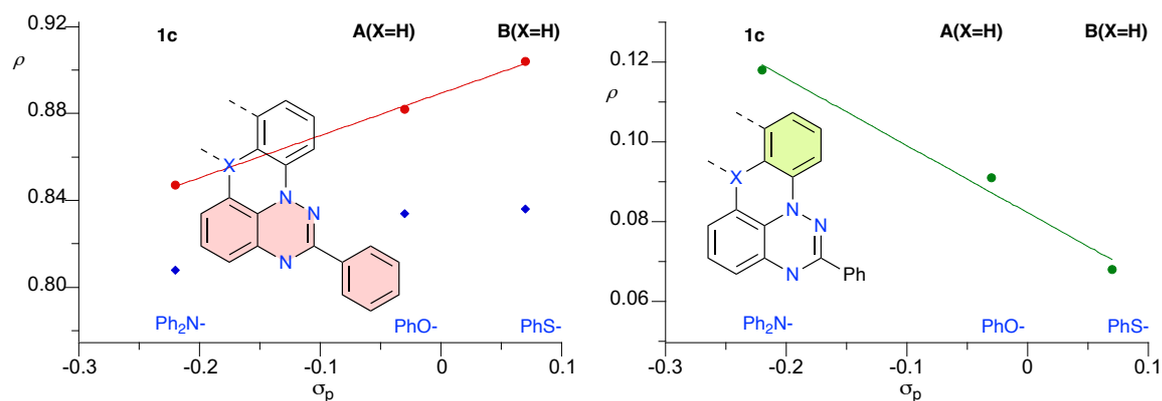
<sup>a</sup> For comparative purposes, the numbering of the systems is according to radical **1a**.

**Table S7.** DFT calculated spin densities of radicals **1** in benzene at the UCAM-B3LYP/EPR-III//UB3LYP/6-311G(d,p) level of theory.<sup>a</sup>



Spin density	1a	1b	1c	1d	1e	1f
$\rho_{N(1)}$	0.270	0.281	0.266	0.266	0.280	0.269
$\rho_{N(3)}$	0.271	0.280	0.272	0.268	0.275	0.275
$\rho_{N(11)}$	0.235	0.232	0.235	0.236	0.219	0.243
$\rho_{N(7)}$	0.016	0.010	0.026	0.027	0.011	0.044
$\rho_{C(2)}$	-0.058	-0.067	-0.058	-0.057	-0.062	-0.063
$\rho_{C(3a)}$	0.014	0.010	0.021	0.022	0.017	0.014
$\rho_{C(3b)}$	0.066	0.064	0.071	0.070	0.063	0.076
$\rho_{C(4)}$	0.006	0.020	0.002	-0.001	0.014	-0.002
$\rho_{C(5)}$	0.023	0.014	0.025	0.026	0.020	0.030
$\rho_{C(6)}$	0.028	0.044	0.022	0.021	0.040	0.020
$\rho_{C(6a)}$	0.022	0.014	0.019	0.021	0.014	0.027
$\rho_{C(7a)}$	0.053	0.053	0.021	0.053	0.037	0.090
$\rho_{C(7b)}$	-0.042	-0.040	-0.041	-0.028	-0.024	-0.043
$\rho_{C(8)}$	0.125	0.114	0.112	0.103	0.063	0.075
$\rho_{C(9)}$	-0.062	-0.058	-0.054	-0.049	-0.045	-0.043
$\rho_{C(10)}$	0.128	0.116	0.107	0.094	0.149	0.069
$\rho_{C(10a)}$	-0.079	-0.073	-0.069	-0.053	-0.075	-0.062
$\rho_{C(a)}$	-0.022	-0.001	-0.011	-0.013	-0.013	-0.009
$\rho_{X(b)}$	0.037	0.019	0.014	0.015	0.015	0.012
$\rho_{C(c)}$	-	-	0.013	0.012	0.010	0.011
$\rho_{C(d)}$	-	-	-0.011	-0.012	-0.011	-0.007
$\rho_{C(e)}$	-	-	0.013	0.012	0.005	0.010
$\rho_{C(f)}$	-	-	-0.008	-0.007	-0.006	-0.008
$\rho_{C(g)}$	-	-	-	-	-0.004	-
$\rho_{C(g')}$	-	-	-	-	-0.047	-
$\rho_{C(h)}$	-	-	-	-	0.004	-
$\rho_{C(h')}$	-	-	-	-	-0.036	-
$\rho_{C(i)}$	-	-	-	-	-0.004	-
$\rho_{C(i')}$	-	-	-	-	0.064	-
$\rho_{C(j)}$	-	-	-	-	0.004	-
$\rho_{C(j')}$	-	-	-	-	0.057	-
$\rho_{C(3-ipso)}$	0.005	0.006	0.003	0.003	0.006	0.020
$\rho_{C(3-o)}$ avg	-0.021	-0.022	-0.020	-0.020	-0.022	-0.022
$\rho_{C(3-m)}$ avg	0.011	0.012	0.012	0.012	0.013	0.011
$\rho_{C(3-p)}$	-0.017	-0.018	-0.017	-0.016	-0.018	-0.017

<sup>a</sup> For comparative purposes, the numbering of the systems is according to radical **1a**.



**Figure S40.** Left: Correlation of DFT spin densities for the triazine ring (blue diamonds) and benzo[*e*][1,2,4]triazine containing the phenyl ring (red) with Hammett parameter  $\sigma_p$  for model substituents (blue). Best fitting function:  $\rho = 0.195(9)\times\sigma_p + 0.889(1)$ ,  $r^2 = 0.998$ .

Right: A similar correlation of DFT spin density on the N-Ph ring (green) with  $\sigma_p$ . Best fitting function:  $\rho = -0.17(2)\times\sigma_p + 0.082(3)$ ,  $r^2 = 0.983$ .

**Table S8.** DFT calculated spin densities of substituted carbazole radicals in benzene at the UCAM-B3LYP/EPR-III//UB3LYP/6-311G(d,p) level of theory.<sup>a</sup>

	nitroxide	carb-N-CPh <sub>2</sub>	N-oxide
Spin density	nitroxide	carb-N-CPh <sub>2</sub>	N-oxide
$\rho_{C(1)}$	-0.031	0.009	0.103
$\rho_{C(2)}$	0.053	-0.005	-0.048
$\rho_{C(3)}$	-0.040	0.009	0.095
$\rho_{C(4)}$	0.063	-0.006	-0.048
$\rho_{C(4a)}$	-0.022	0.009	0.083
$\rho_{C(4b)}$	0.005	0.009	0.083
$\rho_{C(5)}$	-0.005	-0.006	-0.048
$\rho_{C(6)}$	0.004	0.009	0.095
$\rho_{C(7)}$	-0.005	-0.005	-0.048
$\rho_{C(8)}$	0.005	0.009	0.103
$\rho_{C(8a)}$	-0.004	0.011	-0.077
$\rho_{N(9)}$	0.005	0.004	0.287
$\rho_{C(9a)}$	0.039	0.011	-0.077
<b>carbazole total</b>	<b>0.066</b>	<b>0.061</b>	<b>0.505</b>
$\rho_N$	0.418	—	—
$\rho_O$	0.507	—	0.495
$\rho_{tBu}$	0.009	—	—
$\rho_C$	—	0.592	—
$\rho_{PhX2}$	—	0.347	—

**c) spin delocalization in radicals in benzene dielectric medium**

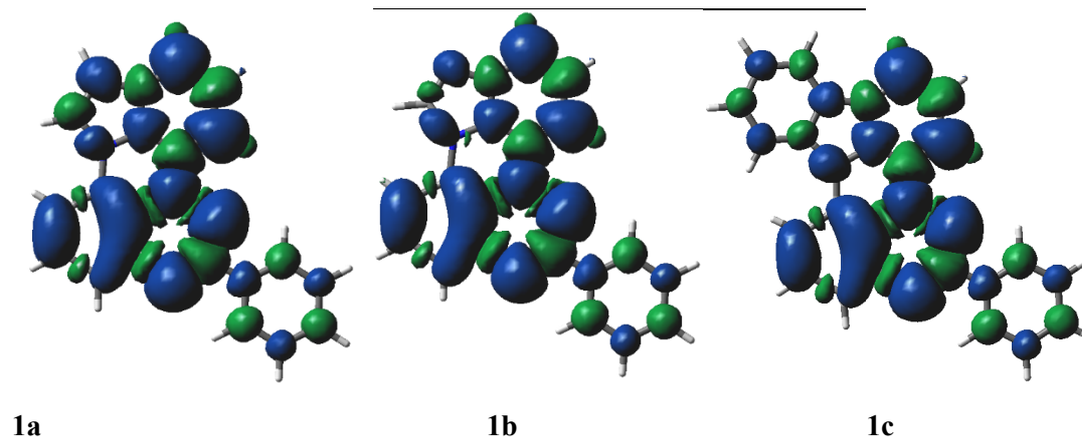
Spin delocalization parameter RDV (Radical Delocalization Value)<sup>14</sup> was calculated according to the formula (eq S12):

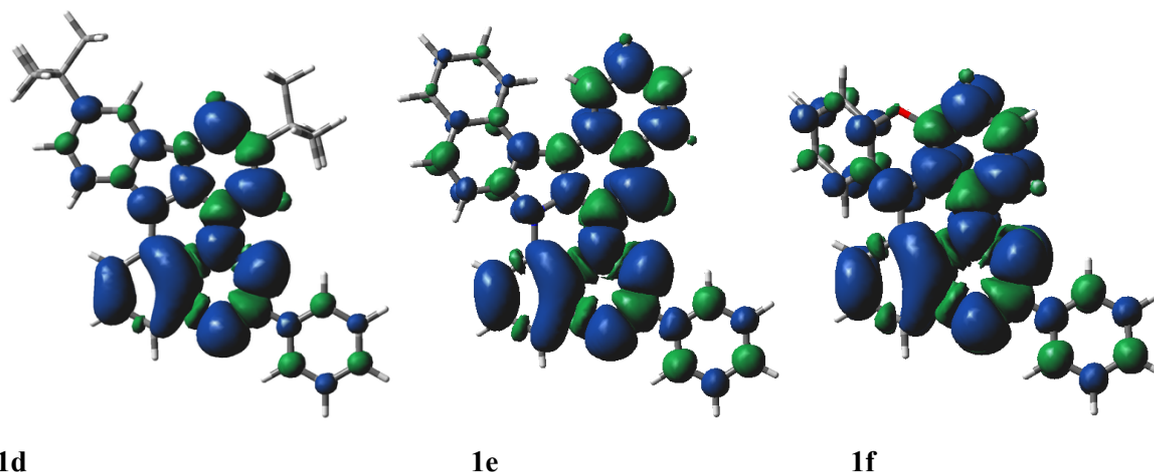
$$RDV = \sum_{i=1}^n (\rho_i)^2 \quad \text{eq S12}$$

where spin concentration  $\rho_i$  on heavy atoms  $i$  (hydrogen atoms summed up to heavy atoms) is obtained with the UCAM-B3LYP/EPR-III // UB3LYP/6-311G(d,p) method in benzene dielectric mediums using the PCM model.<sup>13</sup> For the purpose of this work, an inverse is reported:  $RDV^{-1} = 1/RDV$ , since now larger values corresponds to greater delocalization. Results are shown in Table S9 and in Figure S41. Figure S42 shows correlations  $RDV^{-1}$  with Hammett parameters for model substituents.

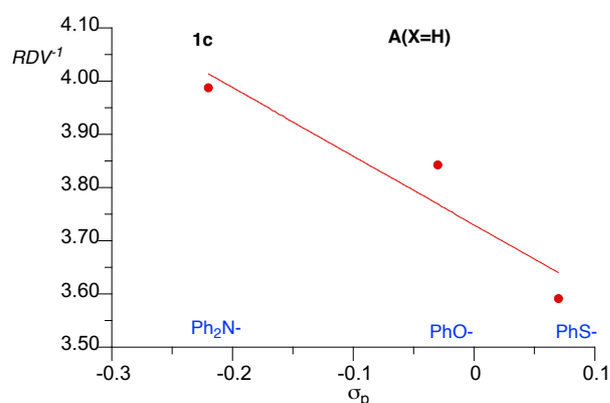
**Table S9.** Radical delocalization value ( $RDV^{-1}$ ) for radicals **1**.

Radical	$RDV^{-1}$ (in benzene)
<b>A (X=H)</b>	3.843
<b>B (X=H)</b>	3.591
<b>1a</b>	3.826
<b>1b</b>	3.782
<b>1c</b>	3.988
<b>1d</b>	4.173
<b>1e</b>	3.822
<b>1f</b>	4.017





**Figure S41.** Spin densities of radicals **1** at the UCAM-B3LYP/EPR-III // UB3LYP/6-311G(d,p) level of theory in benzene dielectric medium. Isovalue: MO = 0.020, Density = 0.0004.



**Figure S42.** Correlation of  $RDV^{-1}$  with Hammett parameters  $\sigma_p$  for model substituents (blue). Best fitting functions:  $a_{N1} = -1.3(4) \times \sigma_p + 3.73(6)$ ,  $r^2 = 0.89$ .

#### **d) electronic excitations**

Electronic excitation energies in  $\text{CH}_2\text{Cl}_2$  dielectric medium were obtained at the UB3LYP/6-311++G(d,p) // UB3LYP/6-311G(d,p) level of theory using the time-dependent TD-DFT method<sup>15</sup> supplied in the Gaussian 16 package. Solvation models in calculations were implemented with the PCM model<sup>13</sup> using the SCRF(solvent= $\text{CH}_2\text{Cl}_2$ ) keyword. Three lowest excitation energies, classified as  $\pi \rightarrow \pi^*$  transitions are listed in Table S10.

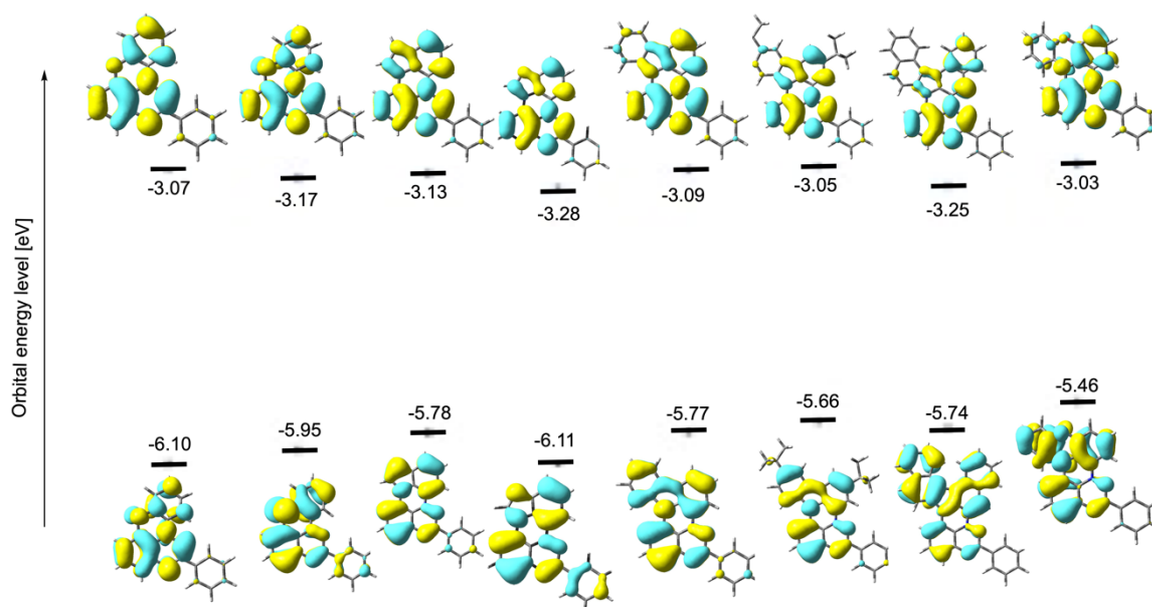
Energies of FMOs involved in the low energy transitions are listed in Table S11. The calculated energies of HOMO and LUMO orbitals together with their visualizations are shown in Figure 43.

**Table S10.** Electronic transition energies and oscillator strength values with the indicated main electronic transition obtained at the TD-UB3LYP/6-311++G(d,p) // UB3LYP/6-311G(d,p) level of theory in CH<sub>2</sub>Cl<sub>2</sub> dielectric medium.

Radical	$\pi \rightarrow \pi^*$ $\beta$ -HOMO $\rightarrow$ $\beta$ -LUMO /nm ( <i>f</i> )	$\pi \rightarrow \pi^*$ $\alpha$ -HOMO $\rightarrow$ $\alpha$ -LUMO /nm ( <i>f</i> )	$\pi \rightarrow \pi^*$ $\beta$ -HOMO-1 $\rightarrow$ $\beta$ -LUMO /nm ( <i>f</i> )	$\pi \rightarrow \pi^*$ $\alpha$ -HOMO $\rightarrow$ $\alpha$ -LUMO+1 /nm ( <i>f</i> )
<b>A(X=H)</b>	586.6 (0.053)	614.8 (0.002)	389.7 (0.040)	452.5 (0.041)
<b>1a</b>	679.2 (0.080)	614.0 (0.005)	476.6 (0.003)	453.0 (0.008)
<b>1b</b>	627.7 (0.067)	597.8 (0.015)	447.7 (0.004)	453.6 (0.010)
<b>1c</b>	662.4 (0.084)	626.6 (0.007)	465.4 (0.019)	494.4 (0.047)
<b>1d</b>	679.0 (0.099)	632.8 (0.007)	504.5 (0.016)	440.4 (0.021)
<b>1e</b>	707.8 (0.063)	605.5 (0.014)	638.6 (0.044)	499.7 (0.116)
<b>1f</b>	754.1 (0.073)	621.2 (0.004)	468.0 (0.025)	488.8 (0.011)

**Table S11.** Energies of MO involved in low energy transitions obtained from the UB3LYP/6-311++G(d,p) // UB3LYP/6-311G(d,p) method in CH<sub>2</sub>Cl<sub>2</sub> dielectric medium.

Radical	$\alpha$ -HOMO $\pi$ /eV	$\alpha$ -LUMO $\pi^*$ /eV	$\alpha$ -LUMO+1 $\pi^*$ /eV	$\beta$ -HOMO-1 $\pi$ /eV	$\beta$ -HOMO $\pi^*$ /eV	$\beta$ -LUMO $\pi^*$ /eV
<b>A(X=H)</b>	-4.933	-1.870	-1.195	-6.975	-6.095	-3.066
<b>B(X=H)</b>	-5.019	-1.850	-1.337	-6.874	-5.953	-3.170
<b>1a</b>	-4.866	-1.834	-1.182	-6.598	-5.782	-3.129
<b>1b</b>	-5.036	-1.929	-1.398	-6.927	-6.112	-3.276
<b>1c</b>	-4.823	-1.840	-1.466	-6.487	-5.771	-3.090
<b>1d</b>	-4.762	-1.796	-1.341	-6.299	-5.660	-3.048
<b>1e</b>	-4.939	-1.927	-1.784	-5.908	-5.741	-3.250
<b>1f</b>	-4.806	-1.804	-1.306	-6.513	-5.461	-3.032



**Figure S43.** The contours and energies of  $\beta$ -HOMO (lower) and  $\beta$ -LUMO (upper) orbitals calculated at the B3LYP/6-311++G(d,p)//B3LYP/6-311G(d,p) level of theory in  $\text{CH}_2\text{Cl}_2$  dielectric medium.

***e) partial output data from TD-DFT calculations for radicals 1***

Method: UB3LYP/6-311++G(d,p)// UB3LYP/6-311G(d,p)

Keywords: TD(nstates=20, root=1) SCRF(solvent=CH2CL2) SCF=tight

**A (X=H)**

```
Excited State 1: 2.048-A' 2.0167 eV 614.79 nm f=0.0019 <S**2>=0.799
  78A -> 79A 0.94989
  76B -> 78B -0.10713
  77B -> 78B -0.20245
This state for optimization and/or second-order correction.
Total Energy, E(TD-HF/TD-DFT) = -970.866917142
Copying the excited state density for this state as the 1-particle RhoCI
density.

Excited State 2: 2.078-A' 2.1136 eV 586.60 nm f=0.0529 <S**2>=0.830
  78A -> 79A 0.19398
  78A -> 80A -0.10304
  77B -> 78B 0.95413

Excited State 3: 2.191-A' 2.7399 eV 452.51 nm f=0.0407 <S**2>=0.950
  77A -> 80A 0.12804
  78A -> 80A 0.94501
  72B -> 78B 0.13306
  77B -> 80B -0.10501

Excited State 4: 2.047-A'' 2.9565 eV 419.35 nm f=0.0011 <S**2>=0.798
  73B -> 78B 0.98747

Excited State 5: 2.397-A' 3.1101 eV 398.66 nm f=0.0398 <S**2>=1.186
  76A -> 79A 0.12731
```

77A -> 79A 0.29700  
78A -> 79A 0.12701  
76B -> 78B 0.87466  
76B -> 79B -0.12656  
77B -> 79B -0.17488

### 1a

Excited State 1: 2.253-A' 2.0194 eV 613.96 nm f=0.1017 <S\*\*2>=1.019

83A -> 86A -0.11775  
84A -> 87A -0.11051  
83B -> 84B 0.94510

This state for optimization and/or second-order correction.

Total Energy, E(TD-HF/TD-DFT) = -1026.50058751

Copying the excited state density for this state as the 1-particle RhoCI density.

Excited State 2: 2.166-A' 2.3691 eV 523.34 nm f=0.0052 <S\*\*2>=0.923

83A -> 85A 0.26477  
84A -> 85A 0.85893  
84A -> 87A 0.18849  
84A -> 95A 0.15933  
81B -> 84B 0.10106

Excited State 3: 2.767-A' 2.9184 eV 424.84 nm f=0.0192 <S\*\*2>=1.664

81A -> 85A -0.11226  
82A -> 86A 0.16170  
82A -> 87A -0.10915  
83A -> 87A -0.20464  
84A -> 85A -0.20513  
84A -> 86A 0.12892  
84A -> 87A 0.45994  
84A -> 95A -0.13869  
79B -> 84B 0.18215  
81B -> 84B -0.10938  
81B -> 85B 0.12765  
82B -> 84B 0.55750  
82B -> 88B 0.16317  
83B -> 86B -0.21225  
83B -> 88B -0.18785

Excited State 4: 2.362-A' 3.1631 eV 391.97 nm f=0.0031 <S\*\*2>=1.145

84A -> 86A 0.72851  
84A -> 87A 0.31048  
82B -> 84B -0.46102  
83B -> 85B -0.13668  
83B -> 86B -0.10248

### 1b

Excited State 1: 2.109-A' 1.9751 eV 627.72 nm f=0.0666 <S\*\*2>=0.862

84A -> 85A -0.20493  
84A -> 87A -0.13510  
83B -> 84B 0.94996

This state for optimization and/or second-order correction.

Total Energy, E(TD-HF/TD-DFT) = -1043.28180792

Copying the excited state density for this state as the 1-particle RhoCI density.

Excited State 2: 2.035-A' 2.0742 eV 597.76 nm f=0.0152 <S\*\*2>=0.785

84A -> 85A	0.94166					
84A -> 87A	-0.11511					
81B -> 84B	-0.10339					
83B -> 84B	0.19847					
Excited State 3:	2.089-A'	2.7342 eV	453.46 nm	f=0.0099	<S**2>=0.841	
84A -> 86A	0.92726					
84A -> 87A	0.25105					
82B -> 84B	-0.19683					
Excited State 4:	2.230-A'	2.7696 eV	447.66 nm	f=0.0038	<S**2>=0.993	
82A -> 87A	-0.10477					
84A -> 85A	0.10309					
84A -> 87A	0.39042					
81B -> 84B	-0.22890					
82B -> 84B	0.82190					
Excited State 5:	2.045-A''	2.8632 eV	433.02 nm	f=0.0011	<S**2>=0.795	
79B -> 84B	0.98300					
79B -> 87B	-0.10458					
Excited State 6:	2.220-A'	2.9983 eV	413.52 nm	f=0.0320	<S**2>=0.982	
83A -> 85A	0.12461					
84A -> 85A	0.13074					
84A -> 86A	-0.10269					
84A -> 87A	0.47322					
78B -> 84B	0.13388					
81B -> 84B	0.79117					
Excited State 7:	2.157-A'	3.0221 eV	410.26 nm	f=0.1628	<S**2>=0.913	
83A -> 86A	0.11670					
84A -> 86A	-0.27813					
84A -> 87A	0.56728					
78B -> 84B	0.22875					
81B -> 84B	-0.48728					
82B -> 84B	-0.45850					
83B -> 85B	-0.12236					
83B -> 86B	-0.10159					

### 1c

Excited State 1: 2.078-A 1.8719 eV 662.35 nm f=0.0835 <S\*\*2>=0.830

97A -> 99A -0.13791

96B -> 97B 0.97392

This state for optimization and/or second-order correction.

Total Energy, E(TD-HF/TD-DFT) = -1180.91934695

Copying the excited state density for this state as the 1-particle RhoCI density.

Excited State 2: 2.040-A 1.9788 eV 626.57 nm f=0.0072 <S\*\*2>=0.790

96A -> 98A -0.12894

97A -> 98A 0.95847

97A -> 99A -0.12347

94B -> 97B -0.11344

Excited State 3: 2.219-A 2.5080 eV 494.36 nm f=0.0472 <S\*\*2>=0.981

97A -> 98A 0.11540

97A -> 99A 0.70330

97A -> 100A 0.24740

95B -> 97B -0.57360

96B -> 97B 0.11355

Excited State 4: 2.136-A 2.6639 eV 465.42 nm f=0.0190 <S\*\*2>=0.890  
 97A -> 99A 0.24337  
 97A ->100A 0.70774  
 95B -> 97B 0.61888

Excited State 5: 2.251-A 2.7581 eV 449.53 nm f=0.0483 <S\*\*2>=1.016  
 95A -> 99A 0.11688  
 97A -> 99A -0.56112  
 97A ->100A 0.60484  
 95B -> 97B -0.44774  
 96B -> 98B 0.15889

Excited State 6: 2.044-A 2.9473 eV 420.68 nm f=0.0011 <S\*\*2>=0.795  
 92B -> 97B 0.98139

Excited State 7: 2.223-A 2.9905 eV 414.59 nm f=0.0630 <S\*\*2>=0.986  
 94A -> 98A 0.10939  
 96A -> 98A 0.23972  
 97A -> 98A 0.13365  
 97A -> 99A -0.10856  
 94B -> 97B 0.90862  
 94B -> 98B -0.10142

**1d**

Excited State 1: 2.074-A 1.8260 eV 678.98 nm f=0.0987 <S\*\*2>=0.825  
 129A -> 131A 0.12438  
 128B -> 129B 0.97648

This state for optimization and/or second-order correction.

Total Energy, E(TD-HF/TD-DFT) = -1495.51217571

Copying the excited state density for this state as the 1-particle RhoCI density.

Excited State 2: 2.040-A 1.9593 eV 632.79 nm f=0.0069 <S\*\*2>=0.791  
 128A -> 130A -0.14351  
 129A -> 130A 0.96025  
 126B -> 129B -0.10457

Excited State 3: 2.189-A 2.4578 eV 504.45 nm f=0.0161 <S\*\*2>=0.948  
 127A -> 131A 0.13381  
 129A -> 131A 0.39244  
 127B -> 129B 0.87195  
 127B -> 131B -0.10107

Excited State 4: 2.188-A 2.6519 eV 467.53 nm f=0.0430 <S\*\*2>=0.946  
 129A -> 130A 0.10897  
 129A -> 131A -0.58068  
 129A -> 132A 0.66865  
 122B -> 129B -0.11005  
 127B -> 129B 0.35754  
 128B -> 131B 0.10689

Excited State 5: 2.242-A 2.7575 eV 449.63 nm f=0.0531 <S\*\*2>=1.007  
 128A -> 131A 0.10505  
 129A -> 131A 0.60358  
 129A -> 132A 0.68232  
 126B -> 129B -0.11020  
 127B -> 129B -0.20748  
 128B -> 130B 0.19854

**1e**

Excited State 1: 2.143-A 1.7516 eV 707.84 nm f=0.0634 <S\*\*2>=0.899

123A ->124A 0.16408  
123A ->125A -0.13404  
120B ->123B -0.11867  
121B ->123B -0.17298  
122B ->123B 0.92740

This state for optimization and/or second-order correction.

Total Energy, E(TD-HF/TD-DFT) = -1488.27105042

Copying the excited state density for this state as the 1-particle RhoCI density.

Excited State 2: 2.141-A 1.9416 eV 638.56 nm f=0.0438 <S\*\*2>=0.896

121B ->123B 0.96373  
122B ->123B 0.15525

Excited State 3: 2.042-A 2.0478 eV 605.46 nm f=0.0141 <S\*\*2>=0.793

123A ->124A 0.85894  
123A ->125A 0.41401  
119B ->123B 0.10439

Excited State 4: 2.239-A 2.4811 eV 499.72 nm f=0.1163 <S\*\*2>=1.003

122A ->125A 0.12254  
123A ->124A -0.38969  
123A ->125A 0.79567  
123A ->126A -0.13064  
123A ->127A 0.10499  
119B ->123B 0.11090  
120B ->123B 0.14770  
121B ->124B 0.10370  
122B ->123B 0.19988

Excited State 5: 3.211-A 2.6444 eV 468.85 nm f=0.0234 <S\*\*2>=2.328

120A ->125A 0.13014  
122A ->124A -0.31029  
122A ->125A 0.44240  
122A ->126A 0.14443  
123A ->124A 0.14201  
123A ->125A -0.18742  
120B ->123B 0.35574  
120B ->125B 0.15802  
121B ->125B 0.17234  
122B ->123B -0.13281  
122B ->124B 0.24811  
122B ->125B 0.45589

**1f**

Excited State 1: 2.063-A 1.6443 eV 754.05 nm f=0.0725 <S\*\*2>=0.814

100B ->101B 0.98432

This state for optimization and/or second-order correction.

Total Energy, E(TD-HF/TD-DFT) = -1256.14705961

Copying the excited state density for this state as the 1-particle RhoCI density.

Excited State 2: 2.043-A 1.9957 eV 621.25 nm f=0.0037 <S\*\*2>=0.793

100A ->102A 0.15877  
101A ->102A 0.95124  
101A ->103A 0.11748

Excited State 3: 2.196-A 2.5366 eV 488.78 nm f=0.0105 <S\*\*2>=0.956

101A ->102A -0.11604  
101A ->103A 0.93773

101A ->104A -0.13897  
 101A ->105A 0.12040  
 99B ->101B 0.11137

Excited State 4: 2.208-A 2.6493 eV 467.98 nm f=0.0246 <S\*\*2>=0.969  
 100A ->102A -0.13768  
 100A ->103A 0.11633  
 101A ->102A 0.12778  
 101A ->103A -0.10219  
 95B ->101B -0.12006  
 99B ->101B 0.92079  
 100B ->102B -0.11260

Excited State 5: 3.117-A 2.8509 eV 434.89 nm f=0.0153 <S\*\*2>=2.179  
 100A ->102A 0.43767  
 100A ->104A -0.20391  
 101A ->102A -0.12060  
 101A ->104A 0.51509  
 99B ->101B 0.18848  
 100B ->102B 0.54778  
 100B ->104B -0.21668

### Mechanistic studies

CAM-B3LYP/6-311G(d,p)//CAM-B3LYP/6-311G(d,p), SCRF(Solvent=EthylEthanoate)

#### 2a,

Excited State 1: Singlet-A 2.7791 eV 446.13 nm f=0.0031 <S\*\*2>=0.000  
 81 -> 85 0.65592  
 81 -> 90 0.10214  
 82 -> 85 0.11243  
 83 -> 85 0.12334

This state for optimization and/or second-order correction.

Total Energy, E(TD-HF/TD-DFT) = -1027.23335250

Copying the excited state density for this state as the 1-particle RhoCI density.

Excited State 2: Singlet-A 3.3572 eV 369.31 nm f=0.0651  
 <S\*\*2>=0.000  
 79 -> 85 -0.11450  
 83 -> 85 -0.14677  
 84 -> 85 0.66657

Excited State 3: Singlet-A 3.9529 eV 313.65 nm f=0.0150 <S\*\*2>=0.000  
 82 -> 85 -0.19600  
 83 -> 85 0.63391  
 84 -> 85 0.17816

Excited State 4: Singlet-A 4.0176 eV 308.60 nm f=0.1545  
 <S\*\*2>=0.000  
 77 -> 85 -0.14536  
 79 -> 86 0.10852  
 81 -> 85 -0.13847  
 82 -> 85 0.61395  
 83 -> 85 0.19123

#### 2a, S1 opt

Excited State 1: Singlet-A 2.1744 eV 570.21 nm f=0.0026 <S\*\*2>=0.000  
 82 -> 85 0.55412  
 82 -> 90 0.10724  
 83 -> 85 0.38444

This state for optimization and/or second-order correction.

Total Energy, E(TD-HF/TD-DFT) = -1027.24358869

Copying the excited state density for this state as the 1-particle RhoCI density.

Excited State 2: Singlet-A 3.3270 eV 372.66 nm f=0.0855 <S\*\*2>=0.000  
 79 -> 85 -0.11205  
 82 -> 85 0.11472  
 83 -> 85 -0.10173  
 84 -> 85 0.66215

Excited State 3: Singlet-A 3.6059 eV 343.83 nm f=0.0006 <S\*\*2>=0.000  
 82 -> 85 0.10111  
 82 -> 86 0.55280  
 83 -> 86 0.38464

**2a, at T geom**

Excited State 1: Triplet-A 1.5340 eV 808.26 nm f=0.0000 <S\*\*2>=2.000  
 82 -> 85 0.59010  
 82 -> 86 -0.15730  
 82 -> 90 0.15169  
 83 -> 85 -0.26020

This state for optimization and/or second-order correction.

Total Energy, E(TD-HF/TD-DFT) = -1027.27003263

Copying the excited state density for this state as the 1-particle RhoCI density.

Excited State 2: Triplet-A 2.3185 eV 534.75 nm f=0.0000 <S\*\*2>=2.000  
 78 -> 85 0.13654  
 79 -> 85 -0.28365  
 79 -> 86 0.11744  
 81 -> 85 0.43756  
 81 -> 86 0.13353  
 82 -> 85 0.11613  
 83 -> 85 0.14880  
 84 -> 85 0.29719

Excited State 3: Triplet-A 3.1418 eV 394.63 nm f=0.0000 <S\*\*2>=2.000  
 79 -> 85 0.28118  
 80 -> 88 -0.12424  
 80 -> 89 0.15578  
 81 -> 85 0.34816  
 81 -> 86 0.23957  
 81 -> 88 0.10548  
 83 -> 85 -0.10687  
 84 -> 85 -0.29857

**2b,**

Excited State 1: Singlet-A 2.7770 eV 446.46 nm f=0.0031 <S\*\*2>=0.000  
 81 -> 85 0.65692  
 83 -> 85 0.11846

This state for optimization and/or second-order correction.

Total Energy, E(TD-HF/TD-DFT) = -1043.29067015

Copying the excited state density for this state as the 1-particle RhoCI density.

Excited State 2: Singlet-A 3.7814 eV 327.88 nm f=0.0710  
 <S\*\*2>=0.000  
 79 -> 85 -0.16037  
 82 -> 85 0.23504  
 83 -> 85 -0.15080  
 84 -> 85 0.60724

Excited State 3: Singlet-A 4.0396 eV 306.92 nm f=0.1865  
 <S\*\*2>=0.000

77 -> 85	-0.13460
81 -> 85	-0.13609
82 -> 85	0.60096
83 -> 85	0.16056
84 -> 85	-0.17431

**2b, S1 opt**

Excited State 1: Singlet-A 2.1737 eV 570.39 nm f=0.0027 <S\*\*2>=0.000  
 82 -> 85 0.31135  
 83 -> 85 0.42143  
 84 -> 85 0.42622

This state for optimization and/or second-order correction.

Total Energy, E(TD-HF/TD-DFT) = -1043.30089767

Copying the excited state density for this state as the 1-particle RhoCI density.

Excited State 2: Singlet-A 3.6127 eV 343.19 nm f=0.0008  
 <S\*\*2>=0.000  
 82 -> 86 -0.31176  
 83 -> 86 -0.42243  
 84 -> 86 -0.42455

Excited State 3: Singlet-A 3.7347 eV 331.98 nm f=0.1300  
 <S\*\*2>=0.000  
 79 -> 85 -0.14589  
 81 -> 85 0.24618  
 82 -> 85 0.26817  
 83 -> 85 0.31851  
 84 -> 85 -0.46884

**2b, at T geom**

Excited State 1: Triplet-A 1.5333 eV 808.64 nm f=0.0000 <S\*\*2>=2.000  
 82 -> 85 0.45417  
 82 -> 86 -0.11708  
 82 -> 90 0.11380  
 83 -> 85 0.38490  
 84 -> 85 0.24908

This state for optimization and/or second-order correction.

Total Energy, E(TD-HF/TD-DFT) = -1043.32728476

Copying the excited state density for this state as the 1-particle RhoCI density.

Excited State 2: Triplet-A 2.3607 eV 525.19 nm f=0.0000 <S\*\*2>=2.000  
 78 -> 85 -0.10217  
 79 -> 85 -0.30953  
 79 -> 86 0.12052  
 81 -> 85 0.43316  
 81 -> 86 0.14984  
 82 -> 85 0.17249  
 84 -> 85 -0.27613

Excited State 3: Triplet-A 3.2031 eV 387.08 nm f=0.0000 <S\*\*2>=2.000  
 77 -> 86 0.13253  
 79 -> 85 0.36871  
 80 -> 88 -0.11763  
 80 -> 89 0.18988  
 81 -> 85 0.31660  
 81 -> 86 0.24175  
 81 -> 88 0.14491  
 84 -> 85 0.16193

**2c,**

Excited State 1: Singlet-A 2.7739 eV 446.96 nm f=0.0027 <S\*\*2>=0.000  
 92 -> 98 0.14719  
 94 -> 98 0.65629

This state for optimization and/or second-order correction.

Total Energy, E(TD-HF/TD-DFT) = -1180.83162975

Copying the excited state density for this state as the 1-particle RhoCI density.

Excited State 2: Singlet-A 3.2178 eV 385.30 nm f=0.0385  
 <S\*\*2>=0.000  
 97 -> 98 0.68592

Excited State 3: Singlet-A 3.9275 eV 315.69 nm f=0.0242  
 <S\*\*2>=0.000  
 95 -> 98 0.25948  
 96 -> 98 0.63835

Excited State 4: Singlet-A 4.0002 eV 309.94 nm f=0.1335 <S\*\*2>=0.000  
 89 -> 98 -0.13149  
 91 -> 99 -0.11177  
 95 -> 98 0.60076  
 96 -> 98 -0.27905

**2c, S1 opt**

Excited State 1: Singlet-A 2.1686 eV 571.73 nm f=0.0025  
 <S\*\*2>=0.000  
 95 -> 98 0.66897  
 95 -> 99 -0.11518  
 95 ->104 0.12679

This state for optimization and/or second-order correction.

Total Energy, E(TD-HF/TD-DFT) = -1180.84188883

Copying the excited state density for this state as the 1-particle RhoCI density.

Excited State 2: Singlet-A 3.1945 eV 388.11 nm f=0.0528 <S\*\*2>=0.000  
 97 -> 98 0.68040

Excited State 3: Singlet-A 3.6105 eV 343.40 nm f=0.0005 <S\*\*2>=0.000  
 95 -> 98 0.11841  
 95 -> 99 0.66777  
 95 ->113 0.12332

**2c, at T geom**

Excited State 1: Triplet-A 1.5306 eV 810.03 nm f=0.0000 <S\*\*2>=2.000  
 95 -> 98 0.64180  
 95 -> 99 -0.16943  
 95 ->104 0.16479

This state for optimization and/or second-order correction.

Total Energy, E(TD-HF/TD-DFT) = -1180.86825041

Copying the excited state density for this state as the 1-particle RhoCI density.

Excited State 2: Triplet-A 2.3276 eV 532.67 nm f=0.0000  
 <S\*\*2>=2.000  
 91 -> 98 -0.33464  
 91 -> 99 0.12478  
 91 ->101 0.10886  
 94 -> 98 0.45876  
 94 -> 99 0.13826  
 97 -> 98 0.29601

Excited State 3: Triplet-A 3.1072 eV 399.03 nm f=0.0000  
 <S\*\*2>=2.000  
 91 -> 98 -0.21829  
 92 ->101 0.10225  
 92 ->102 -0.14168  
 94 -> 98 -0.33338  
 94 -> 99 -0.21054  
 94 ->101 -0.12006  
 97 -> 98 0.41138

**2d,**

Excited State 1: Singlet-A 2.7749 eV 446.80 nm f=0.0026 <S\*\*2>=0.000  
 101 ->106 -0.35156  
 102 ->106 0.56694  
 103 ->106 0.10597

This state for optimization and/or second-order correction.

Total Energy, E(TD-HF/TD-DFT) = -1259.43484969

Copying the excited state density for this state as the 1-particle RhoCI density.

Excited State 2: Singlet-A 3.0766 eV 402.99 nm f=0.0489 <S\*\*2>=0.000  
 105 ->106 0.68465

Excited State 3: Singlet-A 3.7960 eV 326.62 nm f=0.0042  
 <S\*\*2>=0.000  
 104 ->106 0.68788

Excited State 4: Singlet-A 3.9908 eV 310.67 nm f=0.1344  
 <S\*\*2>=0.000  
 97 ->106 -0.14195  
 99 ->107 -0.12465  
 102 ->106 -0.13187  
 103 ->106 0.62717  
 104 ->106 -0.10571  
 105 ->107 0.13349

**2d, S1 opt**

Excited State 1: Singlet-A 2.1712 eV 571.05 nm f=0.0024  
 <S\*\*2>=0.000  
 103 ->106 0.66641  
 103 ->107 0.11578  
 103 ->112 -0.11960

This state for optimization and/or second-order correction.

Total Energy, E(TD-HF/TD-DFT) = -1259.44509320

Copying the excited state density for this state as the 1-particle RhoCI density.

Excited State 2: Singlet-A 3.0515 eV 406.30 nm f=0.0648  
 <S\*\*2>=0.000  
 105 ->106 0.67983

Excited State 3: Singlet-A 3.6080 eV 343.64 nm f=0.0005  
 <S\*\*2>=0.000  
 103 ->106 0.11873  
 103 ->107 -0.66486

**2d, at T geom**

Excited State 1: Triplet-A 1.5325 eV 809.04 nm f=0.0000  
 <S\*\*2>=2.000  
 103 ->106 0.63822

103 ->107 0.16985  
103 ->112 -0.15780

This state for optimization and/or second-order correction.  
Total Energy, E(TD-HF/TD-DFT) = -1259.47145839  
Copying the excited state density for this state as the 1-particle RhoCI density.

Excited State 2: Triplet-A 2.2994 eV 539.20 nm f=0.0000  
<S\*\*2>=2.000  
99 ->106 -0.32416  
99 ->107 -0.12305  
99 ->109 0.10554  
102 ->106 0.44126  
102 ->107 -0.12646  
105 ->106 0.33278

Excited State 3: Triplet-A 3.0365 eV 408.32 nm f=0.0000  
<S\*\*2>=2.000  
99 ->106 -0.14201  
100 ->110 -0.12128  
102 ->106 -0.35048  
102 ->107 0.19191  
105 ->106 0.46246

## 2e,

Excited State 1: Singlet-A 2.7719 eV 447.29 nm f=0.0025 <S\*\*2>=0.000  
118 ->124 0.63125  
119 ->124 -0.24218

This state for optimization and/or second-order correction.  
Total Energy, E(TD-HF/TD-DFT) = -1487.99999072  
Copying the excited state density for this state as the 1-particle RhoCI density.

Excited State 2: Singlet-A 3.2448 eV 382.10 nm f=0.0028  
<S\*\*2>=0.000  
123 ->124 0.69637

Excited State 3: Singlet-A 3.4351 eV 360.93 nm f=0.0176  
<S\*\*2>=0.000  
122 ->124 0.67723

Excited State 4: Singlet-A 3.9295 eV 315.52 nm f=0.4637 <S\*\*2>=0.000  
121 ->129 -0.10945  
122 ->127 -0.14014  
123 ->126 0.67168

Excited State 5: Singlet-A 3.9731 eV 312.06 nm f=0.1720  
<S\*\*2>=0.000  
114 ->124 -0.10797  
116 ->125 0.11871  
119 ->124 0.20320  
120 ->124 0.63321

## 2e, S1 opt

Excited State 1: Singlet-A 2.1640 eV 572.93 nm f=0.0025 <S\*\*2>=0.000  
120 ->124 -0.10339  
121 ->124 0.66726  
121 ->125 0.11256  
121 ->132 -0.12708

This state for optimization and/or second-order correction.  
Total Energy, E(TD-HF/TD-DFT) = -1488.01029994

Copying the excited state density for this state as the 1-particle RhoCI density.

Excited State 2: Singlet-A 3.2304 eV 383.80 nm f=0.0041  
<S\*\*2>=0.000  
123 ->124 0.69467

Excited State 3: Singlet-A 3.4052 eV 364.10 nm f=0.0402 <S\*\*2>=0.000  
122 ->124 0.67110

### 2e, at T geom

Excited State 1: Triplet-A 1.8986 eV 653.03 nm f=0.0000  
<S\*\*2>=2.000  
103 ->124 0.10788  
116 ->124 0.45497  
116 ->128 -0.10901  
119 ->124 0.12749  
120 ->124 -0.44017  
120 ->125 0.11156  
122 ->124 -0.14884  
116 <-124 0.13057  
120 <-124 -0.10330

This state for optimization and/or second-order correction.

Total Energy, E(TD-HF/TD-DFT) = -1488.01716911

Copying the excited state density for this state as the 1-particle RhoCI density.

Excited State 2: Triplet-A 2.0471 eV 605.66 nm f=0.0000  
<S\*\*2>=2.000  
117 ->124 0.63699  
117 ->128 0.11955  
117 ->132 -0.10194  
123 ->124 -0.12159

Excited State 3: Triplet-A 2.1075 eV 588.31 nm f=0.0000 <S\*\*2>=2.000  
121 ->127 -0.18052  
121 ->129 0.10066  
122 ->127 0.11038  
123 ->126 0.63324  
123 <-126 0.13370

### 2f,

Excited State 1: Singlet-A 2.5531 eV 485.62 nm f=0.0002 <S\*\*2>=0.000  
98 ->102 -0.10787  
101 ->102 0.67932

This state for optimization and/or second-order correction.

Total Energy, E(TD-HF/TD-DFT) = -1256.03595057

Copying the excited state density for this state as the 1-particle RhoCI density.

Excited State 2: Singlet-A 2.7849 eV 445.19 nm f=0.0027  
<S\*\*2>=0.000  
98 ->102 0.66991  
101 ->102 0.11500

Excited State 3: Singlet-A 3.8223 eV 324.37 nm f=0.0003  
<S\*\*2>=0.000  
101 ->103 0.65236  
101 ->104 -0.18500

Excited State 4: Singlet-A 3.9631 eV 312.85 nm f=0.1659  
 <S\*\*2>=0.000  
 93 ->102 -0.13837  
 94 ->103 0.12290  
 99 ->102 0.66856

**2f, S1 opt**

Excited State 1: Singlet-A 1.8261 eV 678.96 nm f=0.0001 <S\*\*2>=0.000  
 101 ->102 0.69516

This state for optimization and/or second-order correction.

Total Energy, E(TD-HF/TD-DFT) = -1256.04986137

Copying the excited state density for this state as the 1-particle RhoCI density.

Excited State 2: Singlet-A 2.7970 eV 443.28 nm f=0.0038 <S\*\*2>=0.000  
 96 ->102 0.68894

Excited State 3: Singlet-A 3.5387 eV 350.36 nm f=0.1106  
 <S\*\*2>=0.000  
 93 ->102 0.11162  
 94 ->102 0.17594  
 99 ->102 -0.66283

**2f, at T geom**

Excited State 1: Triplet-A 1.8133 eV 683.76 nm f=0.0000  
 <S\*\*2>=2.000  
 101 ->102 0.69471

This state for optimization and/or second-order correction.

Total Energy, E(TD-HF/TD-DFT) = -1256.04981757

Copying the excited state density for this state as the 1-particle RhoCI density.

Excited State 2: Triplet-A 1.8936 eV 654.75 nm f=0.0000  
 <S\*\*2>=2.000  
 94 ->102 -0.46104  
 99 ->102 0.48535  
 99 ->103 0.11643  
 94 <-102 -0.13274  
 99 <-102 0.11471

Excited State 3: Triplet-A 2.0616 eV 601.39 nm f=0.0000  
 <S\*\*2>=2.000  
 96 ->102 0.66639  
 96 ->105 0.10623  
 96 ->107 -0.10183

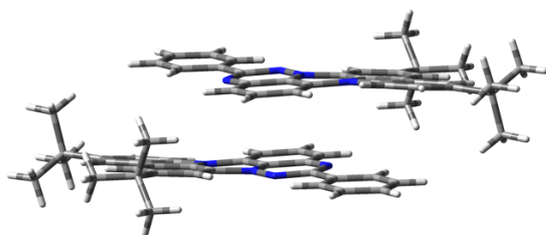
Excited State 4: Triplet-A 2.6284 eV 471.71 nm f=0.0000  
 <S\*\*2>=2.000  
 95 ->108 -0.13009  
 97 ->110 0.18624  
 100 ->106 -0.11561  
 101 ->103 0.12784  
 101 ->104 0.56920  
 101 ->105 0.28399

### f) intermolecular interaction energy calculations

The spin-spin exchange interaction  $J_{DFT}$  for the close pair of molecules in the crystal lattice of **1d** were determined using the Yamaguchi formalism (eq S13):<sup>16</sup>

$$\Delta E_{ST} = 2J_{DFT} = 2 \frac{E_{BS} - E_T}{\langle S^2 \rangle_T - \langle S^2 \rangle_{BS}} \quad \text{eq S13}$$

where the SCF energies of the triplet ( $E_T$ ) and broken symmetry singlet ( $E_{BS}$ ) and total spin angular momenta ( $\langle S^2 \rangle$ ) before spin annihilation were obtained by single point calculations for selected pairs of molecules at crystallographically determined coordinates at the UB3LYP/6-311+G(d) level of theory.<sup>17</sup> The factor “2” in eq S13 makes the experimental and calculated exchange interactions comparable. The input geometries and calculated exchange interaction energies are shown in Figure S44.



crystallographic close pair  $J_{DFT} = +76 \text{ cal mol}^{-1}$ .

**Figure S44.** Pairs of molecules of radical **1d** in their crystallographic coordinates used for DFT calculations and the resulting unprojected spin-spin interaction energies.

## 11. Archive for DFT geometry optimization results.

### 1a

```
1\1\GINC-GAUSSIAN-PRECISION-WORKSTATION-T7500\FOpt\UB3LYP\6-311G(d,p)\
C21H13N4(2)\GAUSSIAN\29-Mar-2024\0\#\p freq=noraman b3lyp/6-311g(d,p)
fopt=tight geom=(nodistance,noangle) #P scf=direct\benzotrazinyl indo
le radical\0,2\N,-0.2511110822,-0.3171637738,0.\N,1.6289855588,1.6830
637674,0.\N,-0.6556188719,0.9759006692,0.\C,0.3133708925,1.8989729284,
0.\C,2.8952711992,-2.3104997987,0.\C,1.5342795901,-2.0280865372,0.\C,1
.1037597319,-0.6808976675,0.\C,2.039928506,0.3796622589,0.\C,3.411458
1958,0.0600145627,0.\C,3.8212060139,-1.2608448152,0.\C,-0.7855726906,-
2.627910587,0.\C,-1.6123626005,-3.7590692972,0.\C,-1.2395237263,-1.312
6504053,0.\C,-3.0050758522,-3.5430705006,0.\C,-2.618374195,-1.11372015
89,0.\C,-3.4729650866,-2.2331854636,0.\H,4.1185478641,0.8791687246,0.\
H,4.8791934103,-1.4953323252,0.\H,3.2344774276,-3.3378814867,0.\C,-0.1
52036837,3.3127405713,0.\C,-1.5166641831,3.6307772263,0.\C,0.788175195
1,4.3508111197,0.\C,-1.9292138074,4.9586776679,0.\H,-2.2443846629,2.83
0695166,0.\C,0.3714731187,5.6777156523,0.\H,1.8393967661,4.0959213915,
0.\C,-0.9875692323,5.9868671512,0.\H,-2.9880740757,5.1923516,0.\H,1.10
91592167,6.4724391838,0.\H,-1.3113687628,7.0218078115,0.\C,-0.71388315
59,-4.8856058731,0.\C,0.5693715808,-4.4016965928,0.\N,0.5348850001,-3.
```

0123690208,0.\H,-3.7000697733,-4.3741111906,0.\H,-4.5423771289,-2.0567  
758609,0.\H,-3.0129302095,-0.1085878432,0.\H,-0.983460087,-5.930015412  
1,0.\H,1.5059378185,-4.9333584324,0.\Version=ES64L-G16RevA.03\State=2  
-A"\HF=-1027.2850617\S2=0.767107\S2-1=0.\S2A=0.750235\RMSD=1.994e-09\R  
MSF=2.623e-07\Dipole=0.1996715,-1.0925411,0.\Quadrupole=4.9272343,7.75  
69219,-12.6841561,-4.547521,0.,0.\PG=CS [SG(C21H13N4)]\@

### 1b

1\1\GINC-GAUSSIAN-PRECISION-WORKSTATION-T7500\FOpt\UB3LYP\6-311G(d,p)\  
C20H12N5(2)\GAUSSIAN\02-Apr-2024\0\#p freq=noraman b3lyp/6-311g(d,p)  
fopt=tight geom=(nodistance,noangle) #P scf=direct\benzotrazinyl benz  
imidazole radical\0,2\N,-0.2580209781,-0.3147744871,0.\N,1.6373961155  
,1.6686033787,0.\N,-0.6521865174,0.9798077744,0.\C,0.3236366748,1.8968  
675409,0.\C,2.8827787882,-2.3322958323,0.\C,1.5254713507,-2.0405745925  
,0.\C,1.095804918,-0.6921869522,0.\C,2.0385455501,0.3627331809,0.\C,3.  
4085727638,0.0348119752,0.\C,3.8142822724,-1.2871027946,0.\C,-0.796329  
4934,-2.613150737,0.\C,-1.5925415942,-3.7551889643,0.\C,-1.2536193879,  
-1.3048380251,0.\C,-2.9866001998,-3.5800465189,0.\C,-2.6365803877,-1.1  
333293133,0.\C,-3.4690347194,-2.2722333856,0.\H,4.1192145892,0.8509497  
246,0.\H,4.8711407976,-1.5257016671,0.\H,3.2147425739,-3.3623693214,0.  
\C,-0.1305217762,3.3133254018,0.\C,-1.4929608501,3.6411268726,0.\C,0.8  
177474965,4.3441758108,0.\C,-1.8954402796,4.9719957201,0.\H,-2.2268969  
111,2.8467486328,0.\C,0.4108399798,5.6739936994,0.\H,1.8671980277,4.08  
21803598,0.\C,-0.9459524166,5.9929542578,0.\H,-2.9524272935,5.21366595  
28,0.\H,1.1542486398,6.4632710029,0.\H,-1.2620256416,7.030227773,0.\C,  
0.4600264696,-4.4078857417,0.\N,0.5139333114,-3.020805595,0.\H,-3.6579  
17379,-4.4290267466,0.\H,-4.540926322,-2.1125177633,0.\H,-3.0555654798  
, -0.1378598007,0.\H,1.3535867479,-5.0124225119,0.\N,-0.7637560792,-4.8  
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0.766609\S2-1=0.\S2A=0.750219\RMSD=9.708e-09\RMSF=1.657e-06\Dipole=0.5  
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4856,0.,0.\PG=CS [SG(C20H12N5)]\@

### 1c

1\1\GINC-GAUSSIAN-PRECISION-WORKSTATION-T7500\FOpt\UB3LYP\6-311G(d,p)\  
C25H15N4(2)\GAUSSIAN\12-Sep-2024\0\#p freq=noraman ub3lyp/6-311g(d,p)  
fopt=tight geom=(nodistance,noangle) #P scf=direct\benzotrazinyl Tcar  
bazole radical\0,2\N,0.5612694871,-0.2876287692,-0.0010660458\N,2.5  
14264207,1.6336923849,-0.0045462006\N,1.863322225,-0.6638914008,0.0032  
350917\C,2.7635729652,0.3252671701,0.0026719566\C,-1.4984907966,2.8091  
046237,-0.0180882543\C,-1.2051393433,1.4480547953,-0.0026228977\C,0.15  
87163934,1.0551794067,-0.0023504062\C,0.8594935027,3.3781145853,-0.017  
7092538\C,-0.4676969783,3.7563520122,-0.0248597292\C,-2.8098449352,-1.  
7880222078,-0.0072403421\C,-0.4044953119,-1.3009994938,-0.0098538808\C  
, -2.5392423384,-3.1642558359,-0.0191438794\C,-0.1416891951,-2.67123958  
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23097\H,-2.5200063576,3.1474509993,-0.030421616\C,4.1882177077,-0.1055  
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64\H,3.7582661984,-2.2091582666,0.0309968677\C,6.5395430978,0.47627179  
15,0.0088184507\H,4.9218105229,1.9045077584,-0.0086430504\C,6.88219688  
85,-0.8746333635,0.023146879\H,6.1373941246,-2.8940625112,0.0424788577  
\H,7.3157950581,1.2333434245,0.0026831895\H,7.9248136455,-1.1727412904  
,0.028244325\C,-3.9985837876,-0.9598911128,0.0085912324\C,-3.578080288  
3,0.4022351873,0.0212722149\N,-2.1709106914,0.423795551,0.008818416\C,  
-5.3583597279,-1.2727104162,0.0172650083\H,-5.6784999421,-2.3083903671  
,0.0072460807\C,-6.2919728064,-0.2451943158,0.0404832995\H,-7.35086384  
16,-0.4746461217,0.0473596954\C,-5.8710315553,1.0896966608,0.057899506  
9\H,-6.6097000613,1.8825937674,0.0798395844\C,-4.5221868419,1.43188724

76,0.0493451717\H,-4.246902966,2.4740121813,0.0703345236\H,-3.34197778  
53,-3.8915703734,-0.02164173\H,-0.985745666,-4.6397932263,-0.035688716  
3\H,0.8832838825,-3.0105186925,-0.0261753693\C,-1.7355277059,-0.895299  
9241,-0.0044415934\C,1.200566014,2.0131711451,-0.0079425801\\Version=E  
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50201\RMSD=4.880e-09\RMSF=2.373e-07\Dipole=-1.092949,-0.0313867,0.0014  
335\Quadrupole=6.4768953,7.3119803,-13.7888756,-3.079821,-0.0684126,0.  
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### 1c-oxo

1\1\GINC-GAUSIANDELL\FOpt\RB3LYP\6-311G(d,p)\C25H14N4O1\PKASZYNSKI\29-  
Mar-2024\0\#P B3LYP/6-311(d,p) FOpt=tight SCRF=(solvent=Benzene) geom  
=(noangle,nodistance)\Flat Blatter Carbazole oxidized, Cs\0,1\N,-0.6  
680907945,-0.1769131444,0.\N,-2.7115694909,1.6609592209,0.\N,-1.944886  
9933,-0.5976317841,0.\C,-2.8992054155,0.3512494707,0.\C,1.2470088512,3  
.0117103858,0.\C,1.0177077872,1.6384390702,0.\C,-0.3334859955,1.184745  
4477,0.\C,-1.4226137302,2.0974693371,0.\C,-1.1396660572,3.4744863234,0  
\C,0.171306267,3.9062906407,0.\C,1.6493483433,-0.6686953042,0.\C,2.75  
06240372,-1.4977259088,0.\C,0.3156566222,-1.1136374437,0.\C,2.57909862  
14,-2.9407811326,0.\C,0.1013305041,-2.5299531797,0.\C,1.1632278567,-3.  
378721855,0.\H,-1.9731481965,4.1639812728,0.\H,0.3846430555,4.96874927  
66,0.\H,2.2506432964,3.3986076475,0.\C,-4.2959747039,-0.1580737769,0.\  
C,-4.5709226188,-1.5325093479,0.\C,-5.362640775,0.7507505348,0.\C,-5.8  
859871773,-1.9855314828,0.\H,-3.7514276566,-2.2381104054,0.\C,-6.67564  
58007,0.2936103579,0.\H,-5.1439726358,1.8100847362,0.\C,-6.9420277935,  
-1.0753124483,0.\H,-6.0870328606,-3.050755078,0.\H,-7.4927557245,1.006  
0050768,0.\H,-7.9663374614,-1.4306433664,0.\C,3.8992753277,-0.63435033  
14,0.\C,3.4309120691,0.7127630891,0.\N,2.019446297,0.6672995413,0.\C,5  
.2730823296,-0.8993458085,0.\H,5.6147064736,-1.9259693082,0.\C,6.15838  
35969,0.1681123164,0.\C,5.6893168076,1.4904404326,0.\H,6.3990378336,2.  
3093086529,0.\C,4.3296732613,1.7826936046,0.\H,4.0160897731,2.81423108  
06,0.\H,-0.9150102418,-2.8964552248,0.\H,7.2260954651,-0.0176885041,0.  
\H,1.0072588523,-4.4511544099,0.\O,3.5099079944,-3.7510778014,0.\Vers  
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.962e-06\Dipole=-0.9673213,2.1416958,0.\Quadrupole=5.2461145,1.958586,  
-7.2047005,20.6217421,0.,0.\PG=CS [SG(C25H14N4O1)]\@

### 1d

1\1\GINC-GAUSSIAN-PRECISION-WORKSTATION-T7500\FOpt\UB3LYP\6-311G(d,p)\  
C33H31N4(2)\GAUSSIAN\26-Mar-2024\0\#p freq=norman b3lyp/6-311g(d,p)  
fopt=tight geom=(nodistance,noangle) #P scf=direct\\benzotrazinyl carb  
azole\0,2\N,1.7099182611,-0.4674908509,0.0578949741\N,3.7113165241,-2  
.3387172277,0.0025449922\N,3.0023513216,-0.0591601479,0.0739524265\C,3  
.9268788147,-1.0251504173,0.0451299235\C,-0.2710914657,-3.6164962878,-  
0.0475520422\C,-0.0117519628,-2.2488757001,-0.0031323873\C,1.341774426  
3,-1.8201349667,0.0137138984\C,2.4073144008,-2.7513866876,-0.013675189  
3\C,2.1009398723,-4.1237477029,-0.0583515721\C,0.7835162719,-4.5361981  
612,-0.0745970468\C,-0.5996704849,0.0752709425,0.0699703926\C,-1.68762  
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38\C,-1.4492052413,2.3267910148,0.1401338076\C,0.9403313523,1.89322716  
69,0.1317364445\C,-0.1370370513,2.8081729696,0.1585658666\H,2.92538586  
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95\H,-1.2850887857,-3.9781847554,-0.0614773978\C,5.3406862374,-0.55920  
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6516,-1.498825565,0.0355818891\C,6.9871030307,1.2168595187,0.124029892  
1\H,4.859288013,1.5321145137,0.1292455085\C,7.7058394802,-1.0825907791  
,0.0519734488\H,6.1234817054,-2.5493841051,0.0013203686\C,8.0152923097  
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8.5004572171,-1.8200989979,0.0301589793\H,9.0502635051,0.5991339812,0.  
1089703985\C,-2.8594719504,0.0892902273,0.0652318419\C,-2.4092177135,-

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-4.2263148154,0.3778283882,0.0713380644\H,-4.5367261336,1.4137729279,0  
.1038037272\C,-5.169717112,-0.6472413509,0.0363459042\C,-4.6920562256,  
-1.9734599119,-0.0050294197\H,-5.4012590967,-2.7921556552,-0.032872499  
1\C,-3.3413131966,-2.297088299,-0.0122965326\H,-3.0551424668,-3.336665  
3441,-0.0449276261\H,-2.2865852349,3.0100152285,0.1600937188\H,1.96383  
93896,2.235471446,0.1450495595\C,0.1849554458,4.3146372714,0.207554907  
6\C,-6.6866971972,-0.3861022063,0.0412299359\C,-1.0852816509,5.1842965  
485,0.2324915689\H,-1.6980663839,5.0363262817,-0.6609222064\H,-1.70213  
74319,4.9795495439,1.1117845401\H,-0.8040334785,6.240032037,0.26695086  
28\C,1.0034940724,4.6263430262,1.4821353251\H,1.9463007716,4.075626872  
,1.5055385717\H,1.239290309,5.69390457,1.5293691885\H,0.4401542703,4.3  
621848761,2.3812203104\C,-7.0193340587,1.1162985156,0.0885796324\H,-6.  
6229515491,1.6484367697,-0.7805194906\H,-8.104084967,1.2515307641,0.09  
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6.8977765401,-0.6538106102,2.203650499\H,-8.4025546157,-0.8804764725,1  
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0.4501179887,4.5011440303,-1.9574190322\C,-7.3169476369,-0.976813005,-  
1.24160545\H,-6.8878278269,-0.5148765508,-2.1348326929\H,-7.1572828331  
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0343,0.003556\Quadrupole=3.906653,8.6423632,-12.5490162,4.9583935,0.19  
66713,0.6910214\PG=C01 [X(C33H31N4)]\@

1e

1\1\GINC-GAUSSIAN-PRECISION-WORKSTATION-T7500\FOpt\UB3LYP\6-311G(d,p)\  
C33H19N4(2)\GAUSSIAN\30-Mar-2024\0\#\#p freq=noraman b3lyp/6-311g(d,p)  
fopt=tight geom=(nodistance,noangle) #P scf=direct\\benzotrazinyl Benz  
carbazole radical\\0,2\N,-0.2926326694,0.5747248284,0.0207453568\N,1.6  
395178259,2.5088824753,-0.1788351282\N,-0.6577027261,1.8799525043,-0.0  
068713212\C,0.3361212993,2.7711330907,-0.0949154815\C,2.7686077681,-1.  
5147612835,-0.0999012425\C,1.4195719112,-1.1976094156,0.0439310847\C,1  
.0422338283,0.165602671,-0.0136980693\C,2.0076164052,1.1937937657,-0.1  
447831891\C,3.3633386069,0.8329792547,-0.244673773\C,3.7231515481,-0.5  
000679106,-0.2365427975\C,-0.9300115505,-1.727283843,0.1376091588\C,-1  
.8102609817,-2.8057584514,0.1875092558\C,-1.3184093278,-0.3771758041,0  
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\H,4.0916349202,1.6271635433,-0.3426217968\H,4.7651211518,-0.778360487  
8,-0.341861171\H,3.086312284,-2.5427668675,-0.1385761886\C,-0.08283913  
93,4.1989505008,-0.1046920204\C,-1.4275249523,4.5648596789,0.040558491  
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9433643978,-0.1358396514\C,-0.9606542439,-3.9878721141,0.2135265532\C,  
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,0.5836007486\H,2.4420576337,-4.0483829907,0.7913252948\H,-2.968856669  
8,0.9378176817,-0.0687035692\C,-4.2151491489,-3.4442061881,0.665157941  
7\C,-4.9965550104,-0.8055821896,0.2393701565\C,-5.544435451,-3.0853112  
024,0.7637042195\C,-5.9463774988,-1.7603272817,0.5187968126\H,-6.99370  
24298,-1.488151821,0.580949482\H,-5.2870856786,0.2298599798,0.09751605  
51\H,-6.2801325371,-3.8305700425,1.0441877415\H,-3.93003336,-4.4571435

571,0.9042605735\C,-2.4051443681,-5.9795021571,-0.3541675365\H,-3.2068  
722215,-5.3368861329,-0.6851269301\C,-0.299521668,-7.6874232257,0.2911  
318364\H,0.5345142195,-8.3422860197,0.5211290815\C,-2.5604938815,-7.34  
52076437,-0.4476201069\H,-3.4961608615,-7.7530060564,-0.8134811636\C,-  
1.5099967219,-8.2136495563,-0.0926793136\H,-1.6475043232,-9.2870349473  
, -0.1534150699\\Version=ES64L-G16RevA.03\State=2-A\HF=-1488.3115342\S2  
=0.768329\S2-1=0.\S2A=0.750272\RMSD=7.510e-09\RMSF=1.656e-07\Dipole=0.  
1438285,-0.8487309,0.0495424\Quadrupole=9.8586221,6.2579398,-16.116561  
8,-5.0854711,0.1243348,-0.8142658\PG=C01 [X(C33H19N4)]\@

## 1f

1\1\GINC-GAUSSIAN-PRECISION-WORKSTATION-T7500\FOpt\UB3LYP\6-311G(d,p)\  
C25H15N4O1(2)\GAUSSIAN\02-Apr-2024\0\#\#p freq=noraman b3lyp/6-311g(d,p  
) fopt=tight geom=(nodistance,noangle) #P scf=direct\\benzotrazinyl ph  
enoxazine radical\\0,2\N,-0.2996144073,0.6893061403,0.2192479929\N,1.7  
02067982,2.560939027,0.368458041\N,-0.5977143443,1.9984079299,0.025482  
6744\C,0.4308874929,2.8523894193,0.0925126788\C,2.6138790104,-1.497302  
0539,0.8358591102\C,1.3067520986,-1.1302904866,0.512220219\C,1.0090700  
469,0.2430229028,0.4091736218\C,2.0076296536,1.2374520787,0.5298429011  
\C,3.3222326404,0.8269450772,0.8117595078\C,3.6027522803,-0.5178723755  
,0.9778967139\C,-1.0637331684,-1.5748359426,0.4522338849\C,-2.09710657  
27,-2.488359879,0.657140439\C,-1.3685648786,-0.2078297291,0.3797131015  
\C,-3.4141450735,-2.0701190336,0.7719220289\C,-2.6920998492,0.22687813  
53,0.5102354215\C,-3.7015058186,-0.7064319198,0.702065613\H,4.08447856  
5,1.5896659957,0.9018316021\H,4.6111474741,-0.8295736983,1.2244627149\  
H,2.8595375257,-2.5383847077,0.985967565\C,0.0902427994,4.2798209433,-  
0.1567694947\C,-1.195815358,4.662226884,-0.5606554453\C,1.0715822399,5  
.2649565752,0.0100952025\C,-1.4920257309,6.0015052907,-0.7895385105\H,  
-1.9526722074,3.9018857598,-0.6992138146\C,0.7706282027,6.6037024331,-  
0.217155939\H,2.0626450049,4.9598038071,0.3180163005\C,-0.5111929366,6  
.9772726461,-0.6171756765\H,-2.4901666956,6.2848807007,-1.1046146169\H  
,1.5385594637,7.3572978537,-0.0822973062\H,-0.7443178204,8.0212376183,  
-0.7953702261\C,-0.6646303496,-4.2652805911,0.1243999545\C,0.402182249  
3,-3.3865147357,-0.1215183881\N,0.2567744401,-2.0461778681,0.316006916  
8\O,-1.8096808953,-3.8326583479,0.7683123012\C,-0.610563232,-5.5922020  
544,-0.2719345753\H,-1.4584639661,-6.2285862198,-0.050888304\C,0.51674  
80169,-6.0724144586,-0.9350126976\H,0.565275646,-7.1127941679,-1.23227  
28304\C,1.5624468851,-5.2031399079,-1.2280928206\H,2.4318107494,-5.556  
5591064,-1.7694421214\C,1.5018416036,-3.8667780143,-0.8380372558\H,2.3  
096937413,-3.1956201908,-1.0924813075\H,-4.1864681402,-2.8124991014,0.  
9258011734\H,-4.7261237735,-0.3704942661,0.8033265457\H,-2.894220464,1  
.2860953179,0.4663397138\\Version=ES64L-G16RevA.03\State=2-A\HF=-1256.  
1843188\S2=0.764376\S2-1=0.\S2A=0.75016\RMSD=8.537e-09\RMSF=3.290e-07\  
Dipole=-0.0770277,-1.1020447,-0.1659398\Quadrupole=5.8453873,5.1448637  
, -10.990251,-4.5257557,-0.0659791,1.7876395\PG=C01 [X(C25H15N4O1)]\@

## 2a, GS

1\1\GINC-LOCALHOST\FOpt\RCAM-B3LYP\6-311G(d,p)\C21H14N4\PIOTR\27-Dec-2  
024\0\#\#P CAM-B3LYP/6-311G(d,p) FOpt=tight SCF=Direct freq(noraman) #P  
Geom=(NoDistance,NoAngle) fcheck SCRF(Solvent=EthylEthanoate)\\benzot  
razine indole\\0,1\N,-0.2145378739,-0.3101522597,-1.2667580259\N,-0.64  
8720431,2.3984149336,-1.4202752345\N,0.6327431304,0.4531548031,-1.8604  
603878\C,0.4096899345,1.8020190268,-1.9124268653\C,-3.4116686408,-0.04  
36890055,0.4453530212\C,-2.2668471162,-0.6078466966,-0.0445106732\C,-1  
.3203223192,0.2299443847,-0.7055650605\C,-1.5585039747,1.6177036723,-0  
.8070203416\C,-2.7508151677,2.1701302864,-0.2851758024\C,-3.6547948259  
,1.3444599236,0.3147151594\C,-1.0045958056,-2.6394604372,0.7111952505\  
C,-1.2586201686,-4.024904774,0.6554115506\C,0.107415189,-2.1194024934,  
1.3702492899\C,-0.3556305863,-4.9096693477,1.2561418466\C,0.9779310347  
, -3.014741865,1.9591299565\C,0.7526665963,-4.3994852904,1.8998061662\H

, -2.9125735565, 3.2354827478, -0.381171527\H, -4.5744087723, 1.7499044978, 0.7175146701\H, -4.1356607179, -0.6690972028, 0.9514350069\C, 1.450237888, 2.610507764, -2.5862384723\C, 2.5811323244, 2.0055019576, -3.1353211285\C, 1.3042169486, 3.9953056278, -2.6757304512\C, 3.5478259863, 2.7757018308, -3.7630418049\H, 2.6923783709, 0.9326697383, -3.0653455574\C, 2.2727008413, 4.7608687017, -3.3032157217\H, 0.4253414474, 4.4589241853, -2.2486765766\C, 3.3971436893, 4.1532745642, -3.8486401357\H, 4.4224767726, 2.2976690619, -4.1869720986\H, 2.150865906, 5.8352750618, -3.3676116621\H, 4.1541385289, 4.7531158519, -4.3394754067\C, -2.4973431035, -4.1989487723, -0.0495303397\C, -2.9439175478, -2.9618643165, -0.3790495585\N, -2.0488671869, -1.997433744, 0.0609962057\H, -0.5309703903, -5.9785777929, 1.2189195857\H, 1.4599801614, -5.072069277, 2.3699574261\H, 1.8537665673, -2.6404718069, 2.4753249016\H, -2.9843074624, -5.1312799638, -0.2870915439\H, -3.8265045309, -2.6612620701, -0.9207814036\H, 0.2907896813, -1.0543586156, 1.4169325525\Version=ES64L-G16RevC.01\State=1-A\HF=-1027.3354817\RMSD=9.766e-09\RMSF=1.311e-06\Dipole=-1.3691419, 1.258514, 0.2137499\Quadrupole=9.9694998, -0.9760925, -8.9934072, 2.1664667, -4.5510987, -0.7268779\PG=C01 [X(C21H14N4)]\@

## 2a, S1

1\1\GINC-LOCALHOST\FOpt\RCAM-B3LYP TD-FC\6-311G(d,p)\C21H14N4\PIOTR\31-Dec-2024\0\#\#P CAM-B3LYP/6-311G(d,p) Fopt TD=(singlets, root=1, NState s=3) SCF=tight #P Geom=(NoDistance, NoAngle) fcheck SCRF(Solvent=EthylE thanoate)\benzotrazine Benzcarbazole TD ex geom\0,1\N, -0.2167145005, -0.2526645528, -1.21203971\N, -0.6563462905, 2.3976746777, -1.3954742582\N, 0.5817607597, 0.5191716905, -1.7706316528\C, 0.4334791824, 1.8481403354, -1.8996007713\C, -3.4754782833, -0.0358830339, 0.4948503803\C, -2.3106575273, -0.6082356017, -0.0083300608\C, -1.3780781302, 0.2071713079, -0.6478176824\C, -1.5883906261, 1.62170029, -0.772950714\C, -2.7684072994, -2.1497640445, -0.2549739385\C, -3.699395649, 1.3256379993, 0.3632173887\C, -1.027793751, -2.6327853184, 0.7221362907\C, -1.2375728683, -4.0235028659, 0.6141051154\C, 0.0687048383, -2.0999661407, 1.3985146151\C, -0.3059075432, -4.8989592178, 1.1838092439\C, 0.969277492, -2.9879522198, 1.9527884818\C, 0.7873966242, -4.376171596, 1.8436806424\H, -2.9354340835, 3.2141670557, -0.3506041876\H, -4.6095781929, 1.7563944429, 0.760225634\H, -4.1944950566, -0.6697017749, 0.9971191441\C, 1.4731029183, 2.6215351995, -2.5852223751\C, 2.6093070995, 1.993326573, -3.0939385816\C, 1.3222719124, 3.9994688353, -2.7291781564\C, 3.5824758913, 2.7371048502, -3.7392374843\H, 2.7239032059, 0.9222494082, -2.9803104989\C, 2.2993042665, 4.7395201698, -3.3758590456\H, 0.4362041934, 4.4753196505, -2.3303270953\C, 3.4299057864, 4.1114816305, -3.8817240016\H, 4.4630643507, 2.2445449004, -4.1326819602\H, 2.1780059169, 5.8101742792, -3.485585912\H, 4.1925403789, 4.6915725707, -4.3869895269\C, -2.4690867868, -4.2083829155, -0.09968603\C, -2.9499617075, -2.9725386418, -0.3883585435\N, -2.0885323667, -2.0002587681, 0.0943406588\H, -0.4469522809, -5.9711243661, 1.1106206898\H, 1.5162545891, -5.0423552121, 2.289480058\H, 1.8331865767, -2.6053903442, 2.4828729003\H, -2.9302080365, -5.1453938435, -0.3684392308\H, -3.8392929014, -2.678348765, -0.9226774097\H, 0.2119687192, -1.0311918429, 1.4878743941\Version=ES64L-G16RevC.01\State=1-A\HF=-1027.3234955\RMSD=6.668e-09\RMSF=6.873e-06\Dipole=-0.3609147, 1.007194, -0.3442113\PG=C01 [X(C21H14N4)]\@

## 2a, T

1\1\GINC-LOCALHOST\FOpt\UCAM-B3LYP\6-311G(d,p)\C21H14N4(3)\PIOTR\31-Dec-2024\0\#\#P UCAM-B3LYP/6-311G(d,p) FOpt=tight SCF=Direct freq(noraman) #P Geom=(NoDistance, NoAngle) fcheck SCRF(Solvent=EthylEthanoate)\benzotrazine Benzcarbazole t\0,3\N, -0.2205321206, -0.2524318672, -1.224630168\N, -0.6522649394, 2.40200827, -1.4077192029\N, 0.6015853956, 0.5112329914, -1.7733130034\C, 0.4254390189, 1.8468985373, -1.901849799\C, -3.4644449082, -0.0330483384, 0.4984737518\C, -2.3074001548, -0.6094655404, -0.0085956574\C, -1.3720975902, 0.204701525, -0.6518520502\C, -1.582712279, 1.6125983595, -0.7740404624\C, -2.7571591148, 2.1486590764, -0.2539360762\C, -3.6889011257, 1.3315865518, 0.3686053554\C, -1.0309019418, -2.6355522538, 0.726

7592458\C,-1.2377234206,-4.0262990586,0.6133298628\C,0.0600079106,-2.1025173535,1.4119592797\C,-0.3084381609,-4.9016387313,1.1872925514\C,0.9582646783,-2.99046175,1.9700650033\C,0.7793146755,-4.3787119701,1.8560907951\H,-2.9201605445,3.2135836414,-0.3531139697\H,-4.5969420786,1.7630152782,0.7693361788\H,-4.1836481949,-0.664877977,1.0036097209\C,1.4673190371,2.6252132501,-2.5885439901\C,2.5991488509,1.9959074118,-3.1033807466\C,1.3201956214,4.0040938683,-2.7261757919\C,3.5737014484,2.7394670947,-3.7485548139\H,2.7104639643,0.9240546299,-2.9954499362\C,2.2970766107,4.7439150143,-3.3723326603\H,0.4367172148,4.480152017,-2.3223462853\C,3.4246486494,4.1141366251,-3.8842303829\H,4.4514481919,2.2458553674,-4.1469360989\H,2.1790212049,5.8153950956,-3.4772074871\H,4.1874509822,4.6943311229,-4.3892287074\C,-2.4630999508,-4.2108409079,-0.1108840531\C,-2.9429956798,-2.9747153168,-0.4007724446\N,-2.0870177367,-2.0027817083,0.0914510622\H,-0.4468410177,-5.9738626446,1.1100449391\H,1.5061831391,-5.0448385904,2.3052000298\H,1.8178252574,-2.6078993796,2.5071502467\H,-2.9209411396,-5.1477328621,-0.3855839608\H,-3.8283906806,-2.6802020808,-0.9414578821\H,0.2006317481,-1.033614507,1.5057644175\\Version=ES64L-G16RevC.01\State=3-A\HF=-1027.2668557\S2=2.028824\S2-1=0.\S2A=2.000404\RMSD=8.662e-09\RMSF=9.653e-07\Dipole=-0.3580375,0.9475862,-0.336578\Quadrupole=8.7809435,-0.711524,-8.0694195,3.5948009,-5.4376822,-1.2416008\PG=C01 [X(C21H14N4)]\@\

## 2b, GS

1\1\GINC-GAUSSIAN-PRECISION-WORKSTATION-T7500\FOpt\RCAM-B3LYP\6-311G(d,p)\C20H13N5\GAUSSIAN\02-Jan-2025\0\#\#P CAM-B3LYP/6-311G(d,p) FOpt=tight SCF=Direct freq(noraman) #P Geom=(NoDistance,NoAngle) SCRF(Solvent=Ethylethanoate)\benzotrazine benzimidazole\0,1\N,-0.6954162497,0.6191849503,1.6329729013\N,-1.2019154127,-1.9829198826,2.3567473074\N,-0.0052793933,0.0542716985,2.5586595149\C,-0.2615178114,-1.2464723083,2.8978743605\C,-3.456000922,-0.173132247,-0.5984315273\C,-2.4413856701,0.5336404691,-0.0195940722\C,-1.6701711107,-0.0802848961,1.0090281857\C,-1.9455987553,-1.4094407811,1.3926812963\C,-3.0041388988,-2.117158209,0.7755638752\C,-3.7415158881,-1.4989269677,-0.189444022\C,-1.0242638729,2.4017112635,-0.9572678323\C,-1.3361312813,3.7427589647,-1.2238542852\C,0.2141473981,1.8503029618,-1.2657502569\C,-0.3831088998,4.578720636,-1.804573478\C,1.1443561588,2.6935155786,-1.8444569195\C,0.8523008188,4.041125974,-2.1087251672\H,-3.199203094,-3.1333179347,1.0914207862\H,-4.5556292674,-2.0258548873,-0.6705828871\H,-4.0405249621,0.2827589914,-1.3870137986\C,0.595997638,-1.8264380005,3.9548891082\C,1.5972725982,-1.0661212878,4.5598810066\C,0.4062339438,-3.1498684915,4.3544691796\C,2.3937708998,-1.6235394557,5.5480192863\H,1.7426701723,-0.0410643702,4.2494608753\C,1.2046284915,-3.7026119944,5.3417039111\H,-0.3715550852,-3.7345022162,3.882291415\C,2.2004846224,-2.9410535653,5.9411507325\H,-3.1685849214,-1.0261871801,6.0127926722\H,1.0503642285,-4.7308644532,5.6453060452\H,2.824464124,-3.3745002866,6.7134882507\C,-3.0972039661,2.8971357878,-0.3794528317\N,-2.1826187903,1.8685429238,-0.407154907\H,-0.6195047261,5.6155089857,-2.0086503652\H,1.6125908961,4.6665333424,-2.5603606264\H,2.1235349239,2.305643689,-2.0974182732\H,-4.0887052522,2.7361950897,0.0187978347\N,-2.641824153,4.0181207832,-0.8431302078\H,0.4474883717,0.8145595658,-1.0585062073\\Version=ES64L-G16RevA.03\State=1-A\HF=-1043.3927245\RMSD=4.224e-09\RMSF=1.420e-06\Dipole=-0.7765075,-2.4035135,-0.0242589\Quadrupole=9.7354998,-9.4365462,-0.2989536,7.7176772,5.6661288,1.8045209\PG=C01 [X(C20H13N5)]\@\

## 2b, S<sub>1</sub> state

1\1\GINC-GAUSSIAN-PRECISION-WORKSTATION-T7500\FOpt\RCAM-B3LYP TD-FC\6-311G(d,p)\C20H13N5\GAUSSIAN\05-Jan-2025\0\#\#P CAM-B3LYP/6-311G(d,p) FOpt TD=(singlets,root=1, NStates=3) SCF=tight #P Geom=(NoDistance,NoAngle) SCRF(Solvent=Ethylethanoate)\benzotrazine Benzcarbazole TD ex\0,1\N,-0.6911429097,0.5571658218,1.6107405235\N,-1.2016388889,-1.9865072385,2.3336546365\N,-0.0435516174,-0.0294916676,2.4960478749\C,-0.23667

7359,-1.2933288146,2.9092398326\C,-3.5062528246,-0.1858046227,-0.64906  
18821\C,-2.4742436207,0.5320024367,-0.0544001075\C,-1.7113386496,-0.06  
82202322,0.9452666582\C,-1.9652033731,-1.4187503955,1.3582396021\C,-3.  
0111932029,-2.1000317463,0.7398593881\C,-3.7719704242,-1.4854166865,-0  
.245466666\C,-1.0431494093,2.3979167558,-0.9658153632\C,-1.3267607695,  
3.7537514993,-1.188550909\C,0.1858996177,1.8303475403,-1.2848916971\C,  
-0.3532268057,4.5875692905,-1.7372997017\C,1.1372071661,2.673422722,-1  
.8291661291\C,0.8734856219,4.0345086072,-2.0507478707\H,-3.2102702192,  
-3.1168729205,1.0502828158\H,-4.5783392572,-2.0341340123,-0.714397959\  
H,-4.0875759559,0.2813456885,-1.433102689\C,0.6189303547,-1.840961691,  
3.9657117244\C,1.6308434334,-1.0681929265,4.534139761\C,0.4170206923,-  
3.1480956468,4.4047782099\C,2.4310740605,-1.599339831,5.5311797936\H,1  
.7853073999,-0.0526751666,4.1904274982\C,1.2207400251,-3.6750042244,5.  
4030817813\H,-0.3714969132,-3.7374898194,3.9558867309\C,2.2279215673,-  
2.9033954125,5.967775587\H,3.2157622889,-0.9955232054,5.9698527789\H,1  
.0606162689,-4.6911875932,5.7417541941\H,2.855142309,-3.3170000004,6.7  
480959316\C,-3.10476812,2.9142025181,-0.3735369167\N,-2.2118305014,1.8  
70295981,-0.4359535226\H,-0.5667741196,5.6348643678,-1.9107887424\H,1.  
6488386989,4.6585516171,-2.4780662908\H,2.1095938555,2.2742930844,-2.0  
907736076\H,-4.1000507875,2.7598942659,0.017938357\N,-2.6279974085,4.0  
413552029,-0.8033826636\H,0.3927465268,0.7819086943,-1.1157200817\\Ver  
sion=ES64L-G16RevA.03\State=1-A\HF=-1043.3807782\RMSD=5.977e-09\RMSF=6  
.201e-06\Dipole=0.1066815,-2.0058041,0.6427243\PG=C01 [X(C20H13N5)]\@

## 2b, T

1\1\GINC-GAUSSIAN-PRECISION-WORKSTATION-T7500\FOpt\UCAM-B3LYP\6-311G(d  
,p)\C20H13N5(3)\GAUSSIAN\04-Jan-2025\0\#\#P UCAM-B3LYP/6-311G(d,p) FOpt  
=tight SCF=Direct freq(noraman) #P Geom=(NoDistance,NoAngle) SCRF(Solv  
ent=EthylEthanoate)\benzotrazine Benzcarbazole t\0,3\N,-0.6947896692  
,0.5591266262,1.621191604\N,-1.198316855,-1.9895529394,2.34493034\N,-0  
.0275383887,-0.0203298877,2.5041659764\C,-0.2441519372,-1.2936006989,2  
.9094241565\C,-3.4948869465,-0.1873968562,-0.6479166009\C,-2.470487792  
4,0.5344543548,-0.0525155684\C,-1.7054977582,-0.0641634815,0.950523001  
8\C,-1.9586984885,-1.4096651032,1.3580311925\C,-2.9991066159,-2.098345  
9769,0.7405674338\C,-3.760477655,-1.4906403352,-0.2463192432\C,-1.0451  
518216,2.4002268179,-0.9715964557\C,-1.3262393642,3.7578830495,-1.1864  
012458\C,0.179332032,1.8297400113,-1.3028169353\C,-0.3544418143,4.5905  
508409,-1.740070183\C,1.1289178644,2.6718327322,-1.8516530181\C,0.8677  
347591,4.0346049488,-2.0658359532\H,-3.1944755535,-3.1152205475,1.0539  
874535\H,-4.5641925378,-2.0404541866,-0.7181653561\H,-4.0762388605,0.2  
778491724,-1.4334612029\C,0.6127076564,-1.8451696527,3.9689717105\C,1.  
6183697565,-1.0696708422,4.5429035394\C,0.4161846352,-3.1548549568,4.4  
02671319\C,2.4195256337,-1.6003811285,5.5404964211\H,1.7682963512,-0.0  
519649418,4.2041328171\C,1.2195292354,-3.6814712352,5.4007780608\H,-0.  
3681691487,-3.7458608337,3.9490728437\C,2.2216902917,-2.9064711313,5.9  
710307989\H,3.199791088,-0.9941347045,5.9836331339\H,1.0638936474,-4.6  
996790609,5.7353779513\H,2.8488432111,-3.3200953959,6.7514591674\C,-3.  
0992070131,2.9208230598,-0.3577358667\N,-2.2104660957,1.8744183952,-0.  
432497825\H,-0.5657032668,5.6392894624,-1.907521016\H,1.6415656956,4.6  
577312868,-2.4972194876\H,2.0977542387,2.2705583187,-2.1229328668\H,-4  
.0913313724,2.7682026212,0.0423669069\N,-2.6228624921,4.0480849252,-0.  
7881570904\H,0.383972101,0.7797195133,-1.1400690335\\Version=ES64L-G16  
RevA.03\State=3-A\HF=-1043.3241454\S2=2.029102\S2-1=0.\S2A=2.000413\RM  
SD=5.383e-09\RMSF=9.238e-07\Dipole=0.111931,-1.9412889,0.6099082\Quadr  
upole=8.3159789,-9.7757245,1.4597456,6.3290663,6.5412559,0.7785389\PG=  
C01 [X(C20H13N5)]\@

## 2c, GS

1\1\GINC-GAUSSIAN-PRECISION-WORKSTATION-T7500\FOpt\RCAM-B3LYP\6-311G(d  
,p)\C25H16N4\GAUSSIAN\01-Jan-2025\0\#\#P CAM-B3LYP/6-311G(d,p) FOpt=tig

ht SCF=Direct freq(noraman) #P Geom=(NoDistance,NoAngle) SCRF(Solvent=EthylEthanoate)\benzotrazine carbazole\0,1\N,0.1957414617,0.0387111446,-1.9634484307\N,0.8019947499,2.7108489815,-2.1838279277\N,1.2055182765,0.4232179163,-2.6599251985\C,1.499731933,1.7565476493,-2.750556002\C,-2.4696090542,1.4972437108,0.0332133416\C,-1.6882807117,0.5440416359,-0.555817059\C,-0.5674243187,0.9605385391,-1.3325236822\C,-0.267269296,2.3334005503,-1.4577838035\C,-1.0961092052,3.2953229314,-0.8340956113\C,-2.1748685587,2.8746615198,-0.1145251128\C,-1.2327377505,-1.7805345847,0.2441597361\C,-1.8889464079,-3.0225939672,0.1742400849\C,-0.0318316468,-1.6263788225,0.9273888151\C,-1.3233008667,-4.1356298095,0.7924383525\C,0.5101358514,-2.7461994875,1.5342526155\C,-0.1247148034,-3.9915041335,1.468277447\H,-0.8490418452,4.3424068783,-0.9486867737\H,-2.8192112128,3.5984373171,0.3684040514\H,-3.3221822757,1.1907666711,0.6254487895\C,2.6899360671,2.1084779049,-3.5566739183\C,3.4516881065,1.1172444159,-4.176095954\C,3.0596077952,3.4460093588,-3.7026402237\C,4.5640472699,1.4615174428,-4.9282805128\H,3.163477925,0.0817807957,-4.0623938162\C,4.1717566392,3.7853560047,-4.4548344781\H,2.4651092785,4.2103301052,-3.2208396276\C,4.927094933,2.794146724,-5.0696660617\H,5.1495807152,0.6854467331,-5.4058989254\H,4.4510195615,4.8263753739,-4.5626110443\H,5.7969120182,3.0607318747,-5.6580081694\C,-3.1062302192,-2.8115760545,-0.5807350181\C,-3.1369056282,-1.4486237479,-0.9296862608\N,-1.9950712756,-0.8281350432,-0.427396544\C,-4.1447327747,-3.64779861,-0.9832737273\H,-4.1345417638,-4.6999497588,-0.7238128865\C,-5.1877704091,-3.1166619018,-1.7218969547\H,-6.0022098049,-3.7548078325,-2.0417147632\C,-5.1997466889,-1.76028389,-2.0636459404\H,-6.0234227317,-1.3655149461,-2.6462748724\C,-4.1789829042,-0.9085132978,-1.6755771853\H,-4.1938770604,0.1389979963,-1.9474592992\H,-1.816707622,-5.0993804154,0.7463590829\H,0.3279728814,-4.8475815231,1.953101711\H,1.4468444911,-2.6544543846,2.0708134802\H,0.4666815019,-0.6674980148,0.9830941567\Version=ES64L-G16RevA.03\State=1-A\HF=-1180.9335696\RMSD=6.270e-09\RMSF=1.034e-06\Dipole=-0.4621819,1.568585,0.4475161\Quadrupole=3.5910365,4.0690773,-7.6601139,-0.820768,-3.8528611,0.0791753\PG=C01 [X(C25H16N4)]\@

## 2c, S<sub>1</sub>

1\1\GINC-GAUSSIAN-PRECISION-WORKSTATION-T7500\FOpt\RCAM-B3LYP TD-FC\6-311G(d,p)\C25H16N4\GAUSSIAN\05-Jan-2025\0\#\#P CAM-B3LYP/6-311G(d,p) Fo pt TD=(singlets,root=1, NStates=3) SCF=tight #P Geom=(NoDistance,NoAngle) SCRF(Solvent=EthylEthanoate)\benzotrazine Carbazole TD ex\0,1\N,0.2117939469,0.0947280247,-1.9362134582\N,0.7961632318,2.7135258819,-2.1683485306\N,1.1822906325,0.5067798269,-2.5952082795\C,1.5376435967,1.7927411792,-2.7557933724\C,-2.5198822361,1.5261578913,0.0786676204\C,-1.7249475109,0.5576914389,-0.5274438634\C,-0.6253698978,0.9605101791,-1.2834214517\C,-0.2909086474,2.3485991984,-1.4310626437\C,-1.1168359329,3.2818690681,-0.8095543299\C,-2.2185238001,2.8711428736,-0.0699666665\C,-1.2490750754,-1.7685812533,0.2528740907\C,-1.8821760824,-3.0218675563,0.1580718662\C,-0.0483337222,-1.6052624874,0.9353616177\C,-1.2943500343,-4.1356594507,0.7534503056\C,0.5159407746,-2.7271345509,1.5182202446\C,-0.0963130168,-3.9822681239,1.4290310778\H,-0.8740996131,4.3301824624,-0.9202580613\H,-2.8451371791,3.6139945734,0.40655152\H,-3.3698168315,1.2097673358,0.6689974735\C,2.7170338937,2.1141624424,-3.5650147465\C,3.4690581296,1.1026250375,-4.161374257\C,3.0888815627,3.4459814436,-3.7386379032\C,4.580542732,1.4221908794,-4.9224218523\H,3.1777874732,0.0682861674,-4.0250468053\C,4.2023597285,3.7605804385,-4.5012518613\H,2.4969033765,4.221713584,-3.2712267179\C,4.9498586175,2.7513265142,-5.0939447632\H,5.1614048893,0.6330695014,-5.3836412541\H,4.487774108,4.7969421497,-4.6336204854\H,5.8199940997,2.9990391523,-5.6897191523\C,-3.1029220456,-2.8172957325,-0.59327595\C,-3.1543261781,-1.4487835072,-0.9187989218\N,-2.0266284326,-0.8177718384,-0.401048759\C,-4.1280240904,-3.662196868,-1.0122328784\H,-4.1027118931,-4.7183545115,-0.770565217\C,-5.1774823042,-3.1340886089,-1.743950618\H,-5.9817971924,-3.7788079788,-2.0760573665\C,-5.2092875033,-1.7722293886,-2.0631038348\

H,-6.0379924363,-1.3805150492,-2.6407104631\C,-4.2025053984,-0.9116758  
092,-1.6585673784\H,-4.231068293,0.140347697,-1.9113954004\H,-1.769844  
6699,-5.1074698207,0.690348572\H,0.3731537902,-4.8391697547,1.89617848  
18\H,1.4518817109,-2.6290198908,2.0550523404\H,0.4290150399,-0.6368628  
099,1.0106579124\\Version=ES64L-G16RevA.03\State=1-A\HF=-1180.9215825\  
RMSD=5.993e-09\RMSF=3.256e-06\Dipole=0.3587305,0.9503436,-0.1934592\PG  
=C01 [X(C25H16N4)]\@

## 2c, T

1\1\GINC-GAUSSIAN-PRECISION-WORKSTATION-T7500\FOpt\UCAM-B3LYP\6-311G(d  
,p)\C25H16N4(3)\GAUSSIAN\13-Jan-2025\0\#\#P UCAM-B3LYP/6-311G(d,p) FOpt  
=tight SCF=Direct freq(noraman) #P Geom=(NoDistance,NoAngle) SCRF(Solv  
ent=EthylEthanoate)\benzotrazine Carbazole t\0,3\N,0.2105053393,0.09  
6708258,-1.9456215338\N,0.8040950116,2.7170559592,-2.1748630524\N,1.19  
27929935,0.4942408469,-2.6066107074\C,1.5296151724,1.7964947664,-2.757  
6885893\C,-2.5005821375,1.5225316398,0.091179729\C,-1.7167889301,0.553  
9465927,-0.5216045541\C,-0.6174446954,0.9556440156,-1.283805994\C,-0.2  
846184631,2.337956823,-1.4259652314\C,-1.0997761735,3.2755051989,-0.79  
86415788\C,-2.1975330427,2.870611591,-0.0532342753\C,-1.2544268905,-1.  
772429706,0.2665406131\C,-1.8888223437,-3.0247923334,0.167777205\C,-0.  
0602072375,-1.6095055122,0.960397523\C,-1.3086016591,-4.1383142467,0.7  
711563701\C,0.4964214083,-2.7311521466,1.5510838812\C,-0.1171276967,-3  
.9854091698,1.4583648957\H,-0.8525544421,4.3228513151,-0.9106887892\H,  
-2.8197258064,3.6126133084,0.4300098686\H,-3.3485374529,1.2073129361,0  
.6855372094\C,2.7092796977,2.1223459144,-3.5729093259\C,3.4562937801,1  
.1116350608,-4.175016972\C,3.0824976887,3.4539257669,-3.7446056396\C,4  
.5658087361,1.4308527537,-4.9404668343\H,3.1634890153,0.077694083,-4.0  
402622733\C,4.1927467015,3.7687636703,-4.5109852125\H,2.4934877027,4.2  
287241401,-3.2723324924\C,4.9360931446,2.7592327241,-5.1096591705\H,5.  
1432065628,0.6419476736,-5.4063306504\H,4.4795861286,4.8048748897,-4.6  
420645815\H,5.8040347287,3.0073359275,-5.7085499685\C,-3.1011029305,-2  
.8194519206,-0.59698721\C,-3.1460687953,-1.4515923783,-0.926402845\N,-  
2.0223644683,-0.821829071,-0.3990254893\C,-4.1240899629,-3.6628911523,  
-1.024176456\H,-4.103825006,-4.7185327825,-0.7797968306\C,-5.165097171  
4,-3.13389262,-1.7671740635\H,-5.9676758501,-3.7774413538,-2.105687944  
4\C,-5.1908873066,-1.7725305538,-2.0892415928\H,-6.0133339841,-1.38013  
06982,-2.675266068\C,-4.1861087888,-0.9134509397,-1.6768286897\H,-4.20  
99684404,0.138286498,-1.9314584078\H,-1.7847590777,-5.109599222,0.7050  
583394\H,0.3462439197,-4.8421250019,1.9318859535\H,1.427134655,-2.6335  
255794,2.0970042407\H,0.4178209867,-0.6415560149,1.0385430747\\Version  
=ES64L-G16RevA.03\State=3-A\HF=-1180.8651428\S2=2.029144\S2-1=0.\S2A=2  
.000413\RMSD=6.768e-09\RMSF=5.429e-07\Dipole=0.3359503,0.8898672,-0.17  
91427\Quadrupole=5.0386644,1.8127011,-6.8513655,0.4208301,-5.6912938,-  
0.745427\PG=C01 [X(C25H16N4)]\@

## 2d, GS

1\1\GINC-LOCALHOST\FOpt\RCAM-B3LYP\6-311G(d,p)\C27H20N4\PIOTR\28-Dec-2  
024\0\#\#P CAM-B3LYP/6-311G(d,p) FOpt=tight SCF=Direct freq(noraman) #P  
Geom=(NoDistance,NoAngle) fcheck SCRF(Solvent=EthylEthanoate)\benzot  
razine carbazole\0,1\N,1.0447221391,0.1312077082,0.2983274198\N,3.177  
6724868,-1.2339346347,-0.7702967612\N,2.2429508376,0.3826300627,0.6908  
69278\C,3.2944038629,-0.2981345517,0.1402006823\C,-0.6702388695,-2.087  
9406109,-2.0155888169\C,-0.4872666913,-1.0956641268,-1.0935638744\C,0.  
8386092041,-0.8093603535,-0.6520200723\C,1.9309653093,-1.5238642727,-1  
.1886295065\C,1.7064452762,-2.5428250884,-2.1432486144\C,0.4286120412,  
-2.815506477,-2.5328352213\C,-1.8484457774,0.9709493709,-0.7272872729\  
C,-3.064303619,1.2815782577,-0.0985147646\C,-1.1228917345,1.9406327063  
,-1.4107948855\C,-3.5518772706,2.5878529131,-0.1421805763\C,-1.6318145  
116,3.2252696713,-1.4389089448\C,-2.8403387953,3.5710945858,-0.8096184  
065\H,2.5593004893,-3.0802247671,-2.5361029185\H,0.2393018341,-3.59314  
23496,-3.2620917968\H,-1.6718903877,-2.3130919725,-2.3583791434\C,4.64

0272924,0.068392487,0.6352171085\C,4.8001693085,1.062870401,1.60052834  
82\C,5.7666225547,-0.5853433547,0.134301695\C,6.0669274175,1.396095011  
,2.054435517\H,3.9264258952,1.5672315225,1.9882659784\C,7.0302100829,-  
0.2494672122,0.590784023\H,5.6367219888,-1.3562563034,-0.6130307201\C,  
7.1839279857,0.7421801331,1.5520398205\H,6.1817934876,2.1695594229,2.8  
03981201\H,7.8987079576,-0.7626960425,0.1962406758\H,8.1729048441,1.00  
41625225,1.9085360213\C,-3.5639648635,0.051780103,0.4789414474\C,-2.63  
11140968,-0.9507666174,0.1679055585\N,-1.5845552333,-0.3877890989,-0.5  
619912188\C,-4.6997573716,-0.2679345914,1.2221896857\H,-5.4229996305,0  
.5022471788,1.4673942126\C,-4.9045709309,-1.5693478797,1.6517827174\C,  
-3.9499870966,-2.5485541789,1.3286361933\H,-4.1089122632,-3.5658229471  
,1.6700084856\C,-2.8139462163,-2.2623007712,0.5939189231\H,-2.09324966  
96,-3.0366705972,0.3644775492\H,-4.4898860948,2.8343659275,0.343434883  
\H,-1.0780209109,3.9954911543,-1.964908969\H,-0.1886313109,1.705466079  
9,-1.9036537335\C,-3.33818694,4.9919366497,-0.8686000035\H,-2.64222268  
52,5.6761517259,-0.3762687451\H,-3.4492339915,5.3313097535,-1.90149182  
25\H,-4.3068326002,5.09120853,-0.3774011352\C,-6.1229840299,-1.9440316  
619,2.4551190986\H,-6.7610155703,-1.0773910666,2.6316942119\H,-6.72074  
83791,-2.7003688186,1.939990961\H,-5.8446863849,-2.358629532,3.4273322  
279\\Version=ES64L-G16RevC.01\State=1-A\HF=-1259.5368259\RMSD=3.274e-0  
9\RMSF=1.054e-06\Dipole=0.2618266,-1.0907159,-1.0118635\Quadrupole=4.5  
437054,2.4280606,-6.971766,0.4819093,-2.3581001,5.6812534\PG=C01 [X(C2  
7H20N4)]\@

## 2d, S<sub>1</sub>

1\1\GINC-LOCALHOST\Fopt\RCAM-B3LYP TD-FC\6-311G(d,p)\C27H20N4\PIOTR\03  
-Jan-2025\0\#\#P CAM-B3LYP/6-311G(d,p) Fopt TD=(singlets,root=1, NState  
s=3) SCF=tight #P Geom=(NoDistance,NoAngle) fcheck SCRF(Solvent=EthylE  
thanoate)\benzotrazine tBuCarbazole TD ex\0,1\N,1.0663939262,0.11236  
66502,0.2321406684\N,3.1634326709,-1.2371524747,-0.7834814367\N,2.2373  
721493,0.3352548431,0.5854401655\C,3.3319587083,-0.2926878526,0.123381  
6676\C,-0.7019791417,-2.1429530973,-2.0830515121\C,-0.5197377599,-1.13  
38083923,-1.1411692234\C,0.772151146,-0.8435680098,-0.7042419099\C,1.9  
104647944,-1.5498307653,-1.2207002404\C,1.6784326712,-2.5528365235,-2.  
1582595691\C,0.3864509732,-2.8452938934,-2.5762418538\C,-1.860354273,0  
.942465303,-0.7599381395\C,-3.0544136772,1.2716756246,-0.097606776\C,-  
1.1276355914,1.9050032278,-1.4471669827\C,-3.5141028668,2.5883629854,-  
0.1160786058\C,-1.6087646594,3.2009201609,-1.4475714506\C,-2.795468539  
5,3.5643523483,-0.7875625769\H,2.5309264838,-3.0923767968,-2.548464396  
5\H,0.2292830016,-3.6270604566,-3.3082140453\H,-1.7047128838,-2.362648  
3411,-2.4253600178\C,4.6491059868,0.0921326731,0.6395466263\C,4.773924  
6275,1.1076431071,1.5868871533\C,5.7875447632,-0.5649917526,0.17667759  
35\C,6.0245998169,1.4601473077,2.0646797973\H,3.88731611,1.6168836748,  
1.9443619588\C,7.0371949658,-0.2085560081,0.6581446549\H,5.6774832436,  
-1.3512880418,-0.5583193721\C,7.1588048173,0.8031723614,1.6016671133\H  
,6.1163808225,2.2496004226,2.8003415186\H,7.9192869087,-0.7221631262,0  
.2959629485\H,8.1364052775,1.0805135224,1.9769386377\C,-3.5634803038,0  
.0456169385,0.4796525132\C,-2.6546017411,-0.9700238451,0.1393661282\N,  
-1.6196588687,-0.4213951727,-0.6151389152\C,-4.6871398367,-0.261810625  
,1.2464907974\H,-5.3925642929,0.5175919488,1.513685576\C,-4.9025074577  
,-1.5631014836,1.671265771\C,-3.9709201112,-2.5550042771,1.3197653323\  
H,-4.1385439611,-3.5719704322,1.6579564425\C,-2.8477013311,-2.28168652  
14,0.5609559929\H,-2.1438655845,-3.0643230999,0.3082337185\H,-4.435921  
7361,2.8495906006,0.3923667012\H,-1.0507015748,3.96574293,-1.976998569  
5\H,-0.2123559141,1.65261077,-1.9667691963\C,-3.2643631068,4.995927315  
6,-0.8195695484\H,-2.5418736649,5.6605236323,-0.3386498255\H,-3.394531  
1973,5.3482468287,-1.8458788717\H,-4.2179430738,5.1103873994,-0.302853  
6999\C,-6.1092181572,-1.9252406921,2.497822029\H,-6.7205240623,-1.0472  
273281,2.70947275\H,-6.7390367657,-2.6547182847,1.9820051943\H,-5.8171  
977299,-2.3684112823,3.4532232847\\Version=ES64L-G16RevC.01\State=1-A\  
HF=-1259.5248815\RMSD=6.820e-09\RMSF=3.497e-06\Dipole=0.6883264,-0.306

6262,-0.2003573\PG=C01 [X(C27H20N4)]\ \@

## 2d, T

1\1\GINC-LOCALHOST\FOpt\UCAM-B3LYP\6-311G(d,p)\C27H20N4(3)\PIOTR\02-Jan-2025\0\ \#P UCAM-B3LYP/6-311G(d,p) FOpt=tight SCF=Direct freq(noraman) #P Geom=(NoDistance,NoAngle) fcheck SCRF(Solvent=EthylEthanoate)\ \benzotrazine tBuCarbazole t\ \0,3\N,1.0703153809,0.1070597988,0.2394071776\N,3.1736108883,-1.2358338159,-0.7809796677\N,2.2415969432,0.346024743,0.6009997058\C,3.3292621425,-0.303804231,0.1249738179\C,-0.6947666895,-2.1162108416,-2.100237093\C,-0.5177586981,-1.1194517495,-1.1492505383\C,0.77547484,-0.8325487943,-0.7053215233\C,1.906845184,-1.5340540759,-1.2246841383\C,1.6827463822,-2.5280906954,-2.1725243568\C,0.3951141219,-2.8177282485,-2.5999149177\C,-1.8699538158,0.9500092455,-0.771053961\C,-3.0628739431,1.2756742282,-0.1048856719\C,-1.1458139959,1.9127689659,-1.4668475236\C,-3.5301771266,2.5896910086,-0.1285190047\C,-1.6345423769,3.2058720138,-1.4722738989\C,-2.8204294126,3.5659621478,-0.8089166881\H,2.5380312147,-3.0644823625,-2.5615573168\H,0.2379495423,-3.5920161497,-3.3395711293\H,-1.6968283034,-2.3326034998,-2.4475811046\C,4.6512756381,0.0734897875,0.6473383626\C,4.7751597325,1.0788163169,1.6044594156\C,5.789661959,-0.581046479,0.1810033354\C,6.0257295582,1.4254825628,2.0890208958\H,3.8886541064,1.5855621035,1.9653159704\C,7.0383975935,-0.2316058935,0.6685796356\H,5.6793618073,-1.359875783,-0.5615955219\C,7.159113203,0.7712711201,1.6223786021\H,6.1168971649,2.2075188517,2.8326033837\H,7.9206854206,-0.7430155792,0.3038428269\H,8.1363969863,1.0432044472,2.0025363698\C,-3.5609341369,0.0499070983,0.482525339\C,-2.6467173988,-0.961752098,0.1443880179\N,-1.6186626684,-0.4110454639,-0.6174118273\C,-4.6791532468,-0.2606917204,1.2561526782\H,-5.388605361,0.5154664487,1.52209879\C,-4.8841553544,-1.5609896402,1.6888733666\C,-3.9479310914,-2.5489642511,1.3382612974\H,-4.107863472,-3.5652268072,1.6822460923\C,-2.8298677189,-2.2725391195,0.5731114802\H,-2.1222959187,-3.0519779176,0.3207809141\H,-4.4508961911,2.8486529773,0.3830882376\H,-1.0834113695,3.9709838217,-2.008490745\H,-0.2313618343,1.6627126018,-1.9892719838\C,-3.2980568599,4.9945324875,-0.8470719158\H,-2.5755634305,5.6667881651,-0.3769649312\H,-3.4389573817,5.3387763693,-1.8747134317\H,-4.247972971,5.1069840047,-0.3232200776\C,-6.0848298463,-1.926446016,2.5227468177\H,-6.7004491758,-1.0511363415,2.7331258138\H,-6.7126301049,-2.662120487,2.0133050794\H,-5.7857259154,-2.3633492547,3.478841545\ \Version=ES64L-G16RevC.01\State=3-A\HF=-1259.4683366\S2=2.029014\S2-1=0.\S2A=2.00041\RMSD=5.912e-09\RMSF=7.092e-07\Dipole=0.6402989,-0.2845218,-0.184359\Quadrupole=7.2453415,0.3810889,-7.6264304,2.2090117,0.0740631,4.9930546\PG=C01 [X(C27H20N4)]\ \@

## 2e, GS

1\1\GINC-LOCALHOST\FOpt\RCAM-B3LYP\6-311G(d,p)\C33H20N4\PIOTR\03-Jan-2025\0\ \#p freq=noraman cam-b3lyp/6-311g(d,p) scrf=(solvent=ethylethanoate) fcheck fopt=tight geom=(nodistance,noangle) scf=direct\ \benzotrazine Benzcarbazole\ \0,1\N,-1.2274902664,-0.4160943673,-1.9979898379\N,-3.4009395457,-2.0523917289,-2.4018113288\N,-2.1173436646,-0.1146441175,-2.8750292133\C,-3.1906638572,-0.9421337146,-3.0672806254\C,-0.5532219249,-3.0351027845,0.4318613386\C,-0.3872772298,-1.8970990937,-0.3013500864\C,-1.3681212494,-1.5501938354,-1.2734790285\C,-2.4848754423,-2.3867649751,-1.4741474809\C,-2.6316881766,-3.5610499783,-0.6971054055\C,-1.6843091563,-3.8665204443,0.2337307856\C,0.8663778133,-0.1230446455,0.9126299393\C,2.0660510808,0.576547839,0.7473324296\C,-0.0364380861,0.1052067934,1.9703917635\C,2.2936480591,1.7163694342,1.5924185928\C,0.2634477745,1.0868932865,2.8606541008\C,1.403321985,1.9249648348,2.685058945\H,-3.4982344297,-4.1864723848,-0.8664665918\H,-1.7835209897,-4.7616848788,0.8343603137\H,0.1913781869,-3.3031075346,1.1703338936\C,-4.1600874918,-0.5228543111,-4.1034949132\C,-3.9654155906,0.65182658,-4.8307054295\C,-5.2842583348,-1.3083875535,-4.3610934531\C,-4.8821690906,1.0316858694,-5.7986279415\H,-3.0940002206,1.2589402922,-4.6302047597\

C,-6.1973264275,-0.9250444579,-5.3290811324\H,-5.4301715817,-2.2179450  
914,-3.7944141894\C,-5.9988871057,0.2460155997,-6.0503880757\H,-4.7233  
982031,1.9452225162,-6.3584744379\H,-7.0670865252,-1.5410981741,-5.522  
1988926\H,-6.7138662552,0.5450882138,-6.8074285767\C,2.7404896603,-0.0  
376626191,-0.385069881\C,1.8484107655,-0.9805695387,-0.9050024681\N,0.  
7326847543,-1.0529859402,-0.0981079005\C,4.0526521919,0.0286481096,-0.  
9677962155\C,4.2839518084,-0.6955331938,-2.1728472577\C,3.2722369949,-  
1.5410825866,-2.7156513626\H,3.4862884927,-2.0765655466,-3.6330902798\  
C,2.0876319257,-1.7255044981,-2.0769030425\H,1.3420606296,-2.412257188  
8,-2.4563708584\H,-0.3940105009,1.2809440108,3.6999668742\C,3.27594792  
3,2.7065747829,1.3575305794\C,1.6143777057,3.0173053322,3.5565708007\C  
,3.4378538864,3.7732266444,2.2037122823\C,2.6197423035,3.9190854176,3.  
3366501894\H,2.7639106031,4.7564590258,4.0083514006\H,0.9409223017,3.1  
421980233,4.3972975613\H,4.1939984579,4.5181602773,1.9858901915\H,3.88  
20850625,2.6511412149,0.4671677838\C,5.1622165441,0.6773984592,-0.3775  
086866\H,5.0566267433,1.1271917914,0.5970920807\C,5.552210136,-0.62941  
06373,-2.7925376089\H,5.6983210768,-1.1658815369,-3.7235795144\C,6.390  
2787094,0.6991199883,-0.9872190306\H,7.2199477375,1.1978497181,-0.5003  
048152\C,6.5859296114,0.06542319,-2.2257559607\H,7.5548858722,0.100552  
9856,-2.7085109832\H,-0.9332008808,-0.4927317936,2.0688311997\\Version  
=ES64L-G16RevC.01\State=1-A\HF=-1488.1018554\RMSD=3.882e-09\RMSF=4.696  
e-07\Dipole=-1.2123521,-1.739597,0.3800671\Quadrupole=0.3558294,1.1041  
121,-1.4599415,8.6294772,-0.8471896,-0.8746341\PG=C01 [X(C33H20N4)]\@

## 2e, S<sub>1</sub>

1\1\GINC-LOCALHOST\Fopt\RCAM-B3LYP TD-FC\6-311G(d,p)\C33H20N4\PIOTR\07  
-Jan-2025\0\#\#P CAM-B3LYP/6-311G(d,p) Fopt TD=(singlets,root=1, NState  
s=3) SCF=tight #P Geom=(NoDistance,NoAngle) fcheck SCRF(Solvent=EthylE  
thanoate)\benzotrazine carbazole TD ex\0,1\N,-1.273806711,-0.4546940  
751,-2.0399355961\N,-3.4095980087,-2.0575851946,-2.4029591899\N,-2.146  
5017399,-0.2247578166,-2.895600641\C,-3.2361666755,-0.9733991156,-3.13  
58990852\C,-0.5842117194,-3.0051763193,0.5267982482\C,-0.4068940582,-1  
.8676916919,-0.2530549629\C,-1.3533355657,-1.5523278264,-1.2248745259\  
C,-2.5014096147,-2.3852924786,-1.4410234488\C,-2.6416836335,-3.5124733  
537,-0.6345390314\C,-1.6953787691,-3.8129946963,0.3367186594\C,0.87810  
1003,-0.1128219772,0.9596290279\C,2.0786814997,0.5830489379,0.78286748  
2\C,-0.0013854304,0.1002265391,2.040217041\C,2.3304203453,1.7054271918  
,1.644508377\C,0.323398699,1.0622679884,2.9430110806\C,1.4645174922,1.  
8972672846,2.7597371956\H,-3.5066132889,-4.143622187,-0.7886899712\H,-  
1.824863675,-4.695602725,0.9495216437\H,0.158002926,-3.2461290182,1.27  
650089\C,-4.1720477826,-0.56082061,-4.185824684\C,-3.9413818345,0.5947  
327104,-4.9313626822\C,-5.3039536599,-1.3332742406,-4.4392635702\C,-4.  
8348989131,0.9715456122,-5.9194462904\H,-3.060982736,1.1934809114,-4.7  
320186648\C,-6.1952771921,-0.9516272124,-5.4292789099\H,-5.4715648272,  
-2.2276345568,-3.8538590316\C,-5.9633967535,0.199639557,-6.1705593033\  
H,-4.6518039188,1.869820416,-6.4959375003\H,-7.0738692893,-1.554540335  
7,-5.6230853465\H,-6.6612590559,0.4965397145,-6.9440189379\C,2.7247030  
564,-0.0123432402,-0.3760199352\C,1.8163089585,-0.9415813045,-0.893242  
9703\N,0.7144605811,-1.0195350259,-0.0673932276\C,4.0239286788,0.05851  
92078,-0.986628333\C,4.2250398704,-0.6452420991,-2.20918765\C,3.198512  
8305,-1.4786672366,-2.7430719098\H,3.391147798,-2.0011654167,-3.672724  
074\C,2.0271373942,-1.6687779785,-2.0818672391\H,1.2706892458,-2.34794  
25577,-2.4531413806\H,-0.3146463834,1.2434123305,3.8000098368\C,3.3131  
79231,2.6949735011,1.408574669\C,1.7004484215,2.9707523287,3.648039529  
2\C,3.4992264219,3.7437626579,2.2720634737\C,2.7062001381,3.8713954318  
,3.4248255569\H,2.8696301001,4.694148476,4.1100548456\H,1.0458709421,3  
.0821522548,4.5054379782\H,4.2552444803,4.4885356793,2.0531924879\H,3.  
900491492,2.6538385657,0.5049190011\C,5.1495519226,0.6919236406,-0.410  
1174978\H,5.0681868345,1.1237520097,0.5748494674\C,5.4790628366,-0.572  
1312041,-2.8565077539\H,5.6018322845,-1.0925212879,-3.7999634084\C,6.3  
63525275,0.7202424641,-1.0471492526\H,7.2061940538,1.206574249,-0.5701

22767\C, 6.5282612288, 0.1086683122, -2.301214296\H, 7.4860949563, 0.1489524404, -2.8053033225\H, -0.8992351916, -0.4950065512, 2.1442156808\\Version=ES64L-G16RevC.01\State=1-A\HF=-1488.0898268\RMSD=3.696e-09\RMSF=3.239e-06\Dipole=-1.3202458, -0.7905591, -0.3827146\PG=C01 [X(C33H20N4)]\@

## 2e, T

1\1\GINC-LOCALHOST\FOpt\UCAM-B3LYP\6-311G(d,p)\C33H20N4(3)\PIOTR\05-Jan-2025\0\#P UCAM-B3LYP/6-311G(d,p) FOpt=tight SCF=Direct freq(noraman) #P Geom=(NoDistance,NoAngle) fcheck SCRF(Solvent=EthylEthanoate)\benzotrazine carbazole t\0,3\N, -1.2418066682, -0.3592406421, -1.8192636544\N, -3.3719266006, -2.0663027087, -2.4460446921\N, -2.2851910444, 0.051899386, -2.5777843141\C, -3.2685796523, -0.7953471375, -2.8466588933\C, -0.1996319275, -3.4315583822, -0.098552173\C, -0.2115669858, -2.1277924111, -0.5757770879\C, -1.265417952, -1.6317552311, -1.3738315568\C, -2.3398569614, -2.5183717937, -1.6789008662\C, -2.310767548, -3.8278225147, -1.1916797523\C, -1.2520940305, -4.2822004444, -0.4107664417\C, 0.8258207273, -0.1618742144, 0.5464495169\C, 2.066842185, 0.5630526069, 0.4770125453\C, -0.2228006769, 0.1867955945, 1.3951391257\C, 2.1419151849, 1.8367117072, 1.1677666413\C, -0.0644132657, 1.3049718771, 2.1646335886\C, 1.0804805876, 2.1606382716, 2.0533548141\H, -3.1407567101, -4.478072422, -1.4413910093\H, -1.24882448, -5.2997737364, -0.0405774521\H, 0.6274630061, -3.7687838733, 0.5144536332\C, -4.3787557511, -0.2621688406, -3.689632626\C, -4.3420267002, 1.0406232263, -4.1884911156\C, -5.4773217083, -1.0650700892, -3.9934019373\C, -5.3782939068, 1.5262473988, -4.9706944914\H, -3.4881575258, 1.6598120215, -3.9511257165\C, -6.5146558366, -0.5783585079, -4.7760504551\H, -5.5000094875, -2.0737129639, -3.6041946059\C, -6.4700128358, 0.7191431266, -5.2682816756\H, -5.3342461529, 2.5402868694, -5.3511972269\H, -7.3618306832, -1.2155807877, -5.0023994016\H, -7.2797472661, 1.0996743342, -5.8799534451\C, 2.9130669847, -0.1931966005, -0.3435753686\C, 2.1053211809, -1.259758449, -0.8726968058\N, 0.8773757529, -1.2458539482, -0.2799439609\C, 4.3171412297, -0.1688729656, -0.7081894327\C, 4.733228249, -1.033471007, -1.7550379456\C, 3.8136708145, -1.963356616, -2.3412567673\H, 4.1733242184, -2.5923119908, -3.1467953102\C, 2.5347105002, -2.1189061576, -1.8834429445\H, 1.8644771977, -2.8611700717, -2.2936869926\H, -0.8483446576, 1.5961014014, 2.8536972289\C, 3.1274474997, 2.8031650541, 0.9448622427\C, 1.1072822678, 3.3708589265, 2.7597189958\C, 3.1209179847, 4.0033070995, 1.6336521108\C, 2.1254746817, 4.2809788456, 2.566540673\H, 2.1331092947, 5.2157872259, 3.1121156369\H, 0.2977967863, 3.593414249, 3.4445505907\H, 3.8945751208, 4.734152434, 1.4339791113\H, 3.8858579223, 2.6383134195, 0.1956939258\C, 5.2983877817, 0.5688938183, -0.0384233803\H, 5.0363281045, 1.1521544238, 0.8304853043\C, 6.0747486883, -1.0388259671, -2.1628057271\H, 6.3707533453, -1.6895611243, -2.977075486\C, 6.6218007412, 0.5278949503, -0.4394991469\H, 7.3590387252, 1.1066690866, 0.1027834979\C, 7.0117693643, -0.2551728867, -1.5231559117\H, 8.0460196733, -0.2706911887, -1.8419354643\H, -1.1172242149, -0.4183906007, 1.4380878318\Version=ES64L-G16RevC.01\State=3-A\HF=-1488.0240448\S2=2.07007\S2-1=0.\S2A=2.003009\RMSD=5.600e-09\RMSF=1.726e-06\Dipole=6.7342027, 1.3352857, 3.3264235\Quadrupole=3.6174354, 1.669811, -5.2872464, -2.0661057, -12.9106749, 14.1680263\PG=C01 [X(C33H20N4)]\@

## 2f, GS

1\1\GINC-LOCALHOST\FOpt\RCAM-B3LYP\6-311G(d,p)\C25H16N4O1\PIOTR\28-Dec-2024\0\#P CAM-B3LYP/6-311G(d,p) FOpt=tight SCF=Direct freq(noraman) #P Geom=(NoDistance,NoAngle) fcheck SCRF(Solvent=EthylEthanoate)\benzotrazine Phenoxazine\0,1\N, 0.5844556146, -0.0000041328, -0.3543197815\N, 2.7185110428, 0.0000274424, 1.3809242259\N, 1.789857197, -0.0000099492, -0.8017256415\C, 2.8404296952, 0.0000062224, 0.0749422143\C, -1.1527499654, 0.0000485245, 2.8355980299\C, -0.9602489758, 0.000024287, 1.4860944566\C, 0.3730107691, 0.000016936, 0.9820711201\C, 1.4656505761, 0.0000337565, 1.8726048934\C, 1.2316598957, 0.000058384, 3.2694609067\C, -0.0510371686, 0.0000653669, 3.7293510518\C, -2.4974178878, 1.2097747526, 0.0207751835\C, -3.5274164342, 1.1793548782, -0.9237443602\C, -1.9606879896, 2.4438297118, 0.36704

89125\C,-3.9971694018,2.3385245954,-1.5043888544\C,-2.4388322421,3.615  
3746827,-0.2106997058\C,-3.4559013208,3.5687213216,-1.1469976693\H,2.0  
845516127,0.0000713784,3.9354083414\H,-0.244276812,0.00008388,4.794640  
2118\H,-2.1636357122,0.000054012,3.2225275825\C,4.1924714552,-0.000000  
7771,-0.5268813054\C,4.3586865382,-0.0000222885,-1.9121782008\C,5.3188  
014981,0.0000142987,0.2967829786\C,5.6317401902,-0.0000288071,-2.46088  
60657\H,3.4855112395,-0.0000332091,-2.5492294616\C,6.5886290157,0.0000  
076658,-0.2557620689\H,5.1839111391,0.0000312507,1.369852552\C,6.74865  
38243,-0.0000139897,-1.6361082741\H,5.751610751,-0.0000453848,-3.53738  
74865\H,7.4570426407,0.0000195372,0.3916748927\H,7.7424734243,-0.00001  
92337,-2.0674730663\C,-3.5274155465,-1.1794031599,-0.9236979288\C,-2.4  
974166163,-1.2097848126,0.0208223092\N,-2.063428269,0.0000063024,0.591  
0332878\O,-4.1327039679,-0.0000316278,-1.2927753283\C,-3.9971684396,-2  
.3385963959,-1.504295578\H,-4.7947912174,-2.2584155004,-2.2320852507\C  
, -3.4559004264, -3.5687786568, -1.1468544392\H, -3.8315245043, -4.47620740  
61, -1.6018147992\C, -2.4388311588, -3.6153940904, -0.210554778\H, -2.00437  
47685, -4.5635596176, 0.0803513924\C, -1.9606864803, -2.443825617, 0.367145  
7834\H, -1.1662073613, -2.4929958939, 1.0993078917\H, -4.794791684, 2.25831  
41508, -2.232175809\H, -3.8315249101, 4.4761315616, -1.6019953448\H, -2.004  
3754619, 4.5635519746, 0.0801675154\H, -1.166208397, 2.4930296762, 1.099208  
4644\\Version=ES64L-G16RevC.01\State=1-A\HF=-1256.1297753\RMSD=4.256e-  
09\RMSF=5.784e-07\Dipole=0.7668215,0.0000319,1.7613802\Quadrupole=-1.5  
072404,-3.5902561,5.0974966,0.0000298,-0.6201796,0.0001115\PG=C01 [X(C  
25H16N4O1)]\@

## 2f, S<sub>1</sub>

1\1\GINC-LOCALHOST\Fopt\RCAM-B3LYP TD-FC\6-311G(d,p)\C25H16N4O1\PIOTR\  
02-Jan-2025\0\#\#P CAM-B3LYP/6-311G(d,p) Fopt TD=(singlets,root=1,NSta  
tes=3) SCF=tight #P Geom=(NoDistance,NoAngle) fcheck SCRF(Solvent=Ethy  
lEthanoate)\benzotrazine phenoxazine TD ex\0,1\N,0.5559872625,-0.000  
0034152,-0.4102516658\N,2.7308375917,0.0000275792,1.3389074143\N,1.828  
9314091,-0.000009931,-0.8676066903\C,2.8259006702,0.0000043355,0.00683  
27035\C,-1.1784415234,0.000047607,2.8261747447\C,-0.9278970009,0.00002  
46679,1.4567132746\C,0.3765987673,0.0000177414,0.9197335638\C,1.470115  
8199,0.0000350315,1.8368282043\C,1.2093807578,0.0000580841,3.211816512  
2\C,-0.0968252251,0.0000643461,3.6950701311\C,-2.4724817732,1.19332821  
56,0.0016978848\C,-3.4601085264,1.1692087765,-0.9987297493\C,-1.999042  
1214,2.4348301215,0.4520827081\C,-3.9695749645,2.3412225615,-1.5330050  
768\C,-2.5055526707,3.5948321798,-0.0821655587\C,-3.4917063624,3.55083  
50214,-1.0745169306\H,2.0545902747,0.0000708637,3.8883688837\H,-0.2736  
101789,0.0000825729,4.7633509833\H,-2.196302284,0.0000520941,3.1933632  
495\C,4.2024851431,-0.0000024384,-0.570966927\C,4.4026274672,-0.000026  
8891,-1.9516395905\C,5.3147152727,0.0000154235,0.2689466071\C,5.685523  
0154,-0.0000337119,-2.4773597445\H,3.5378919773,-0.0000405399,-2.60071  
88692\C,6.5983713931,0.0000087158,-0.2578466747\H,5.1526628993,0.00003  
43639,1.3380950918\C,6.7892333526,-0.0000159646,-1.6330082907\H,5.8253  
768913,-0.0000529922,-3.5521429202\H,7.4530425767,0.0000227423,0.40869  
32678\H,7.7915157356,-0.0000213039,-2.0451894713\C,-3.4601069001,-1.16  
92592309,-0.9986837938\C,-2.4724799621,-1.1933380219,0.0017446069\N,-1  
.9994278113,0.00000541,0.5085446316\O,-3.9457404285,-0.0000349259,-1.4  
754352132\C,-3.9695717586,-2.3412947025,-1.5329130403\H,-4.7271942628,  
-2.2752605686,-2.3016888428\C,-3.4917014289,-3.5508884982,-1.074377452  
5\H,-3.8818589922,-4.470859605,-1.4887558826\C,-2.5055474938,-3.594845  
315,-0.0820245508\H,-2.1344892303,-4.5488750776,0.2668453995\C,-1.9990  
384366,-2.4348215656,0.4521780645\H,-1.2362959703,-2.4578904415,1.2154  
705811\H,-4.7271972562,2.275157182,-2.3017784039\H,-3.8818650758,4.470  
7893199,-1.4889315942\H,-2.1344958049,4.5488761736,0.2666669591\H,-1.2  
362998342,2.4579299982,1.2153744662\\Version=ES64L-G16RevC.01\State=1-A  
\HF=-1256.1169692\RMSD=8.953e-09\RMSF=6.084e-06\Dipole=-5.7481995,-0.  
0000177,-0.2310034\PG=C01 [X(C25H16N4O1)]\@

**2f, T**

```
1\1\GINC-LOCALHOST\FOpt\UCAM-B3LYP\6-311G(d,p)\C25H16N4O1(3)\PIOTR\04-
Jan-2025\0\#\#P UCAM-B3LYP/6-311G(d,p) FOpt=tight SCF=Direct freq(noram
an) #P Geom=(NoDistance,NoAngle) fcheck SCRF(Solvent=EthylEthanoate)\
benzotrazine phenoxazine triplet\0,3\N,0.6162304516,0.0000503265,-0.4
604317588\N,2.7668335063,0.0000277619,1.3341922815\N,1.9006693419,0.00
00695109,-0.887924557\C,2.8833937194,0.0000568806,0.0016592165\C,-1.16
23458564,-0.0000467396,2.7535745075\C,-0.8959506925,-0.000020947,1.391
8334131\C,0.4128825876,0.000010018,0.8714765022\C,1.4902541893,0.00000
39194,1.8051713671\C,1.2122170917,-0.0000253337,3.1753768346\C,-0.0979
486009,-0.0000487222,3.6454766617\C,-2.4868080869,1.191472465,-0.03192
5765\C,-3.5472864312,1.1687270614,-0.9547975861\C,-1.9777649204,2.4348
546374,0.3777670873\C,-4.0957216839,2.3421261716,-1.4523167261\C,-2.52
10916161,3.5923050298,-0.1203019391\C,-3.5829961026,3.5502074818,-1.03
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000674193,4.7105808268\H,-2.1870870532,-0.0000617827,3.1039973861\C,4.
2685191074,0.0000671052,-0.5526225679\C,4.4929391485,0.000052631,-1.93
00226016\C,5.3668413701,0.0000856141,0.3061389141\C,5.7844406482,0.000
0548348,-2.4336173802\H,3.6389189438,0.0000428923,-2.5931129711\C,6.65
92574894,0.0000893922,-0.1987735238\H,5.1862533564,0.0000961191,1.3723
447931\C,6.8738227409,0.0000724423,-1.5705034369\H,5.9426586593,0.0000
431867,-3.5059187796\H,7.5024658771,0.0001043633,0.4823156593\H,7.8830
367605,0.0000740057,-1.9655196159\C,-3.5473647416,-1.1687900386,-0.954
6998839\C,-2.486892616,-1.191529631,-0.0318200397\N,-1.9859288851,-0.0
000249747,0.4442414911\O,-4.0727101413,-0.0000320688,-1.389024109\C,-4
.0958723377,-2.342193053,-1.4521295162\H,-4.9111446901,-2.2743691121,-
2.1596317643\C,-3.5832279525,-3.5502748184,-1.0347757839\H,-4.00148425
68,-4.4712934571,-1.4188993102\C,-2.5213348713,-3.5923688556,-0.120003
2865\H,-2.1217729055,-4.5467836941,0.1959126095\C,-1.9779369496,-2.434
913957,0.3779784123\H,-1.1579849732,-2.4582599276,1.0804223413\H,-4.91
10044193,2.2742994845,-2.1598068101\H,-4.0011974278,4.4712233751,-1.41
92515405\H,-2.1214664987,4.5467192147,0.1955360453\H,-1.1578078082,2.4
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ipole=-6.026714,-0.0001148,-0.1240784\Quadrupole=1.780444,9.2791116,-1
1.0595557,0.0011521,6.7721136,-0.001041\PG=C01 [X(C25H16N4O1)]\@
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**nitroxide**

```
1\1\GINC-LOCALHOST\FOpt\UB3LYP\6-311G(d,p)\C16H17N2O1(2)\PIOTR\08-Apr-
2025\0\#\#P UB3LYP/6-311G(d,p) Fopt=tight geom=(nodistance,noangle) SCF
=direct\carbazole-NO-tBu\0,2\C,-1.2316547433,1.5697056842,-0.0160268
252\C,-1.0240127755,0.1734201382,-0.1467433467\C,-0.1690304832,2.47275
52204,-0.0733127284\C,0.2702907539,-0.3182223142,-0.3294894011\C,1.108
2003984,1.9643798118,-0.246948999\C,1.3366643559,0.5776655915,-0.35789
32441\C,-2.3296092854,-0.4511437567,-0.0700120244\C,-3.2707008954,0.59
51952475,0.1094672951\N,-2.5865919379,1.8001729564,0.1420684708\C,-2.7
813479946,-1.7727011659,-0.132797536\H,-2.0798832066,-2.5883568085,-0.
2693456012\C,-4.1421106183,-2.0292463568,-0.0191389755\H,-4.5038550555
,-3.0496003928,-0.0673862124\C,-5.0579876623,-0.980119899,0.1574700323
\H,-6.1150273255,-1.2049193258,0.2432104814\C,-4.6375016215,0.34332298
36,0.2247595538\H,-5.3499773676,1.1491949392,0.3613365917\H,0.43902760
44,-1.3762592146,-0.4756205066\H,1.9595420826,2.627073376,-0.323766908
1\H,-0.3297077532,3.5415773948,0.0120454834\H,-3.0126715854,2.70607395
45,0.2393705013\N,2.6859884373,0.1385242652,-0.5922092264\O,3.38791403
32,0.8781986438,-1.3713887286\C,3.4893487419,-0.7006699595,0.385108296
8\C,2.6144937747,-1.6672904805,1.188715609\H,1.8478445238,-1.150755287
7,1.7677078165\H,2.1328974122,-2.4093572523,0.5485260463\H,3.256126472
,-2.2067647194,1.8899137208\C,4.1978799419,0.2811525758,1.337094962\H,
3.4701696875,0.8426684089,1.9293552546\H,4.8506117516,-0.2626654182,2.
0249842156\H,4.80028287,0.9861232607,0.7628596448\C,4.5138123288,-1.48
7373243,-0.4437427282\H,5.1708585616,-2.0537680748,0.2210120793\H,4.01
```

01593557,-2.1898007058,-1.1129868832\H,5.1151652237,-0.809371077,-1.04  
71671802\\Version=ES64L-G16RevC.01\State=2-A\HF=-804.8230124\S2=0.7554  
37\S2-1=0.\S2A=0.750024\RMSD=9.280e-09\RMSF=5.024e-07\Dipole=-0.957072  
1,0.1092644,0.941295\Quadrupole=1.3953651,7.4916347,-8.8869998,-8.4612  
26,5.0248205,2.2573305\PG=C01 [X(C16H17N2O1)]\@

**carb-N-CPH<sub>2</sub>**

1\1\GINC-LOCALHOST\FOpt\UB3LYP\6-311G(d,p)\C25H18N1(2)\PIOTR\09-Apr-20  
25\0\\#P UB3LYP/6-311G(d,p) Fopt=tight geom=(nodistance,noangle) SCF=direct\\  
carbazole-CPH<sub>2</sub>\0,2\C,-1.1505803895,1.1204874438,0.9020381819\C  
, -1.0073018303, -0.0316961999, 0.0938416895\C, -0.0865083659, 1.626377758,  
1.6483674396\C, 0.2307778886, -0.6762894661, 0.0249263461\C, 1.1352892887,  
0.9674956116, 1.5615534567\C, 1.2976404161, -0.1702641636, 0.757218464\C, -  
2.3014060987, -0.2924635303, -0.502515807\C, -3.182809262, 0.7071631246, -0  
.028059921\N, -2.4797441153, 1.5712267791, 0.8288501331\C, -2.7718064864, -  
1.2599861265, -1.3945450878\H, -2.1075743324, -2.0308521499, -1.7693273466  
\C, -4.1002316652, -1.2167411152, -1.7994216683\H, -4.4785517831, -1.961453  
4831, -2.4900370694\C, -4.9579618206, -0.2123515713, -1.3268623119\H, -5.98  
97414548, -0.192554697, -1.6590968274\C, -4.5141994687, 0.7626303743, -0.43  
9767839\H, -5.1822076642, 1.5342807343, -0.0806750702\H, 0.3554973321, -1.5  
621986491, -0.5878630685\H, 1.9785932162, 1.3434894058, 2.1297156002\H, -0.  
2033729006, 2.5029469532, 2.2717417213\H, 2.263631898, -0.6596598047, 0.712  
2132493\C, -3.0148198802, 2.6948695042, 1.4986341729\C, -2.3914943138, 3.98  
14522728, 1.2540367925\C, -1.7167981512, 4.231982922, 0.0357671897\C, -2.40  
4584066, 5.0173640479, 2.2182158002\C, -1.1146858582, 5.4565345693, -0.2104  
862531\H, -1.6876597702, 3.4572905729, -0.7201122298\C, -1.7953728023, 6.23  
81450219, 1.9648761876\H, -2.8704135227, 4.8428419143, 3.1797042647\C, -1.1  
505289898, 6.4702416118, 0.7488311099\H, -0.6157106691, 5.6248766154, -1.15  
8341643\H, -1.8115429921, 7.0107302484, 2.7257314453\H, -0.6760753809, 7.42  
51226874, 0.5543940725\C, -4.1462009244, 2.4751810785, 2.3791900799\C, -5.1  
064565305, 3.4832563366, 2.6333979214\C, -4.3411351206, 1.2171838558, 2.996  
9279523\C, -6.1850167181, 3.2480905103, 3.4742447058\H, -5.0141180168, 4.44  
21038514, 2.1394344285\C, -5.4194479679, 0.9926262539, 3.8392644077\H, -3.6  
243225469, 0.4254153353, 2.8190715292\C, -6.3483620257, 2.0048650885, 4.087  
5761743\H, -6.9115989713, 4.0352416418, 3.6431272345\H, -5.5365912081, 0.02  
24658003, 4.3091700768\H, -7.1918669552, 1.8249760913, 4.7439674856\\Versi  
on=ES64L-G16RevC.01\State=2-A\HF=-1018.4866173\S2=0.774948\S2-1=0.\S2A  
=0.750553\RMSD=8.613e-09\RMSF=5.165e-07\Dipole=-0.1860129,0.390618,0.2  
328475\Quadrupole=0.9785687,2.3764254,-3.3549941,0.2100591,-2.8620039,  
3.0927713\PG=C01 [X(C25H18N1)]\@

**N-oxide**

1\1\GINC-LOCALHOST\FOpt\UB3LYP\6-311G(d,p)\C12H8N1O1(2)\PIOTR\09-Apr-2  
025\0\\#p ub3lyp/6-311g(d,p) fopt=tight geom=(nodistance,noangle) scf=direct\\  
carbazole-N-oxide\0,2\C,0.,-1.144639235,0.6066100306\C,0.,-0.  
7311101185,-0.7415799322\C,0.,-2.4761930236,0.9948882647\C,0.,-1.70068  
76283,-1.740783295\C,0.,-3.4287586143,-0.0217959879\C,0.,-3.0463331357  
, -1.3700326662\C, 0., 0.7311101185, -0.7415799322\C, 0., 1.144639235, 0.6066  
100306\N, 0., 0., 1.4317737058\C, 0., 1.7006876283, -1.740783295\H, 0., 1.4209  
847893, -2.78800789\C, 0., 3.0463331357, -1.3700326662\H, 0., 3.8106686561, -  
2.1383776299\C, 0., 3.4287586143, -0.0217959879\H, 0., 4.4817777759, 0.23410  
40245\C, 0., 2.4761930236, 0.9948882647\H, 0., 2.7423241926, 2.0439191984\H,  
0., -1.4209847893, -2.78800789\H, 0., -4.4817777759, 0.2341040245\H, 0., -2.7  
423241926, 2.0439191984\H, 0., -3.8106686561, -2.1383776299\O, 0., 0., 2.6978  
987203\\Version=ES64L-G16RevC.01\State=2-B1\HF=-592.1593573\S2=0.76106  
7\S2-1=0.\S2A=0.750102\RMSD=5.480e-09\RMSF=1.767e-06\Dipole=0.,0.,-1.0  
849064\Quadrupole=-5.916299,8.1246499,-2.2083509,0.,0.,0.\PG=C02V [C2(  
N1O1),SGV(C12H8)]\@

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