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Zhu, Congzhi Ji, Xiaozhou You, Di <u>et al.</u>

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Ladder-Type Conjugated Molecules Produced by B ← N Coordination-Promoted Delocalization and Hyperconjugation

⁴ Congzhi Zhu,[†][®] Xiaozhou Ji,[†] Di You,[†] Teresa L. Chen,[§] Anthony U. Mu,[†] Kayla P. Barker,[†] ⁵ Liana M. Klivansky,[§] Yi Liu,[§][®] and Lei Fang^{*,†,‡}[®]

⁶ [†]Department of Chemistry and [‡]Department of Materials Science and Engineering, Texas A&M University, 3255 TAMU, College
 7 Station, Texas 77843, United States

⁸ [§]The Molecular Foundry, Lawrence Berkeley National Laboratory, One Cyclotron Road, Berkeley, California 94720, United States

9 Supporting Information

ABSTRACT: The introduction of $B \leftarrow N$ coordination—isoelectronic to C–C 10 single bond—into π -systems represents a promising strategy to impart exotic 11 redox and electrochromic properties into conjugated organic molecules and 12 macromolecules. To achieve both reductive and oxidative activities using this 13 14 strategy, a cruciform ladder-type molecular constitution was designed to accommodate oxidation-active, reduction-active, and $B \leftarrow N$ coordination units 15 into a compact structure. Two such compounds (BN-F and BN-Ph) were 16 synthesized via highly efficient N-directed borylation. These molecules 17 demonstrated well-separated, two-reductive and two-oxidative electron-transfer 18 processes, corresponding to five distinct yet stable oxidation states, including a 19 20 rarely observed boron-containing radical cation. Spectroelectrochemical measurements revealed unique optical characteristics for each of these 21



reduced/oxidized species, demonstrating multicolor electrochromism with excellent recyclability. Distinct color changes were observed between each redox state with clear isosbestic points on the absorption spectra. The underlying redox mechanism was elucidated by a combination of computational and experimental investigations. Single-crystal X-ray diffraction analysis on the neutral state, the oxidized radical cation, and the reduced dianion of **BN-Ph** revealed structural transformations into two distinct quinonoid constitutions during the oxidation and reduction processes, respectively. B \leftarrow N coordination played an important

27 role in rendering the robust and reversible multistage redox properties, by extending the charge and spin delocalization, by

modulating the π -electron density, and by a newly established hyperconjugation mechanism.

29 INTRODUCTION

30 Ladder-type conjugated molecules, constituted with an 31 uninterrupted sequence of fused adjacent rings that share 32 two or more atoms with one another, have shown great 33 promise for applications on multiple fronts that demand 34 superior optical, electronic, or mechanical properties.^{1–10} 35 Stemming from the strong intramolecular electronic couplings 36 throughout fused conjugated backbones, ladder-type conju-37 gated compounds also demonstrate intriguing electrochemical 38 behaviors, such as the capability of undergoing multiple 39 electron transfers.^{7,11} The extended conjugation and rigid 40 backbones can greatly stabilize the highly reactive radical and 41 ionic intermediates that are generated during the redox 42 processes. For instance, ladder-type oligothiophene exhibited 43 stepwise oxidations.^{7,12} A recently reported series of ladder-44 type conjugated polycyclic hydrocarbons, featuring extensive 45 graphitic constitutions, demonstrated four distinct reversible 46 redox processes.¹³ In this context, ladder-type molecular design 47 represents an important strategy to achieve exotic molecular 48 materials desirable for sophisticated electrochemical applica-49 tions.^{11,14,15}

To date, redox-active ladder-type compounds are mostly 50 fused with entirely covalent bonds.^{1,3,4} Alternatively, intra- 51 molecular noncovalent $B \leftarrow N$ coordination¹⁶⁻¹⁹ has been 52 exploited to serve as a bridge to construct ladder-type 53 conjugated molecules while imparting modification to the 54 aromatic character.^{20,21} Compared to the isoelectronic and 55 isosteric C–C single bond, intramolecular $B \leftarrow N$ coordination 56 in a conjugated molecule not only extends the π -delocalization 57 but also dramatically changes the electronic structures, $_{58}$ electronic and optical properties, $^{17-19,22-32}$ and reactiv- $_{59}$ ities,^{33,34} because of the intrinsically different valence electron 60 configurations and electronegativities of boron and nitrogen 61 compared to carbon. For instance, the incorporation of B \leftarrow N $_{62}$ coordination into conjugated molecules deepens the lowest 63 unoccupied molecular orbital (LUMO) energy levels, which 64 facilitate reductive electron-transfer processes.^{17,19,25–27,35} 65 Additionally, boron atoms can couple with a neighboring 66 unpaired spin, providing additional stabilization effects for 67

Received: October 21, 2018 Published: December 3, 2018 68 radical intermediates during redox processes.^{35–38} Therefore, 69 the employment of $B \leftarrow N$ coordination represents a powerful 70 strategy to develop novel conjugated molecules and macro-71 molecules with new redox properties that were not accessible 72 before.

73 Despite the great potential of this noncovalent approach, 74 however, it is still challenging to achieve robust redox activities 75 in B ← N bridged ladder-type molecules, especially oxidation 76 processes, because of the electron-deficient nature of boron 77 moieties and the resulting low stability of the corresponding 78 oxidized states. Meanwhile, the underlying mechanism of the 79 redox processes of B ← N bridged ladder-type molecules 80 remains unclear, especially on the impacts of tetracoordinated 81 boron centers on the electronic structures and optical 82 characteristics of these molecules. Herein, we report a 83 molecular design strategy that allowed for the access to five 84 redox states, both reduced and oxidized, of B ← N bridged 85 ladder-type molecules, as well as the corresponding mecha-86 nistic investigations.

87 RESULTS AND DISCUSSION

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88 To pursue both robust reductive and oxidative electron-89 transfer processes in $B \leftarrow N$ bridged ladder-type molecules, we 90 envisioned that the molecular design should first integrate 91 multiple redox-active components in an orthogonal and 92 compact manner, so that multiple electron-transfer processes 93 can be enabled in a small molecule without interfering with 94 each other. Second, $B \leftarrow N$ coordination needs to be installed 95 to rigidify the π -system to facilitate the desired delocalization 96 of charges and spins, therefore stabilizing the reduced or 97 oxidized species. On the basis of these principles, we designed 98 a ladder-type molecule composed of an oxidation-active 99 indolo[3,2-b]carbazole (ICBZ) unit and two reduction-active 100 benzo[d]thiazole (BTH) units (Figure 1a). The ICBZ unit is



Figure 1. (a) Designed ladder-type molecule featuring cruciform arrangement of redox-active units (reduction-active in blue and oxidation-active in red) and $B \leftarrow N$ coordination that can rigidify and coplanarize the entire molecule; (b) representative orbital interactions of the boron center in the designed conjugated backbone.

101 covalently linked to the BTH units at the 6 and 12 positions, 102 leading to a cruciform-like geometry, generating two distinctive 103 delocalization regions for oxidation and reduction, respectively. 104 On one hand, ICBZ is known to undergo two-electron 105 oxidation processes into a relatively stable quinonoid 106 structure.^{39,40} The two N-H functionalities on ICBZ provided 107 the point to covalently attach the boron centers.⁴¹⁻⁴³ Indeed, 108 B \leftarrow N bridged ladder-type ICBZ derivatives reported by 109 Curiel et al.⁴³ showed the expected reversible electrochemical 110 oxidation behavior. In our design, tertiary butyl groups are 111 installed at the 2 and 8 positions of ICBZ to ensure the 112 solubility of this rigid compound. On the other hand, BTH was 113 selected as the reduction-active unit.⁴⁴ The Lewis basic 114 nitrogen atoms on the thiazole rings were employed to form 115 the B \leftarrow N coordination.

The $B \leftarrow N$ coordination in this molecular design was 116 formed in a geometrically favored six-membered heterocycle 117 architecture, which fused the ICBZ and BTH units into a 118 ladder-type structure. The N-B \leftarrow N constitution was 119 expected to be stable not only in ambient condition but also 120 during redox processes, similar to the known stability of 121 BODIPY dyes. 45,46 The strong B \leftarrow N coordination 122 significantly withdrew the electron density from BTH units, 123 resulting in an enhanced electron affinity.^{17,19,25} Consequently, 124 the reduction of such a molecule could fall into an accessible 125 potential range. Moreover, this boron-containing six-mem- 126 bered heterocycle architecture reassembles the structure of 127 cyclohexa-1,3-diene, in which a series of hyperconjugative 128 interactions were previously observed to significantly impact 129 the molecular conformation, properties, and stability.^{47,48} In 130 this design, the sp³ hybridized boron atom carried two ligands 131 that were expected to undergo hyperconjugative interactions 132 similar to those existing in cyclohexa-1,3-diene (Figure 1b). 133 We hypothesized that these hyperconjugative interactions 134 could further stabilize the radical intermediates upon 135 reduction/oxidation, by assisting the charge and spin 136 delocalization. Consequently, the low-energy singly occupied 137 molecular orbital (SOMO) to LUMO transition of these 138 stabilized organic radicals could be utilized to achieve 139 electrochromism in the near-infrared (NIR) region,⁴⁹ which 140 is highly desired for various applications including data storage, 141 smart windows, and sensing ^{11,50–52} smart windows, and sensing. 142

On the basis of this design principle, two molecular 143 candidates, **BN-F** and **BN-Ph**, with fluoride and phenyl groups 144 attached to the boron center as ligands, respectively, were 145 synthesized. The syntheses were accomplished (Scheme 1) in 146 s1

Scheme 1. Synthesis of BN-F and BN-Ph



three steps starting from 2,8-di-*tert*-butyl-5,11-dihydroindolo- 147 [3,2-*b*]carbazole (compound 1). First of all, treatment of 1 148 with 2.0 equiv of *N*-bromosuccinimide (NBS) selectively 149 brominated the 6- and 12-positions to give a dibromo 150 derivative, 2. Compound 2 was subsequently subjected to 151 Stille coupling reaction with 2-(tributylstannyl)benzo[*d*]- 152 thiazole. CuI and CsF were used to facilitate the trans- 153 metalation step⁵³ during the Stille reaction to give intermediate 154 3. In this step, temperature control was crucial to achieve a 155 high yield. Significant side reactions were found above 130 °C, 156

157 which was likely a result of undesired copper-catalyzed 158 reactions, such as Ullmann coupling.⁵⁴ Finally, borylation of 159 3 with BF₃ and BPh₂Cl in the presence of non-nucleophilic 160 bases afforded the desired products in excellent isolated yields 161 (96% and 72%, respectively). Notably, the isolation yield of 162 BN-Ph was significantly improved compared to a previously 163 reported reaction using BPh₃ to introduce $B \leftarrow N$ coordinate 164 bonds into a similar ICBZ backbone,⁴³ likely due to the higher 165 electrophilicity and smaller steric hindrance of BPh₂Cl. BN-F 166 and BN-Ph showed good stabilities in ambient conditions. For 167 example, BN-Ph was purified by normal-phase silica gel 168 chromatography. Both final products were fully characterized 169 by NMR spectroscopy and mass spectrometry. In addition, 170 BN-Ph was characterized unambiguously by single-crystal X-171 ray diffraction analysis. Single crystals of BN-Ph suitable for X-172 ray diffraction analysis were obtained by vapor diffusion of pentane into a chloroform solution. Tetrahedral geometry of 173 174 the boron centers was revealed, with a short $B \leftarrow N$ coordinate 175 bond length of 1.643 Å. The anticipated rigid conformation of 176 **BN-Ph** bridged by the $B \leftarrow N$ coordination was validated in 177 the solid-state crystal structure. The dihedral angles between 178 ICBZ units and BTH units were measured to be 20.0° in BN-179 Ph, significantly smaller than that observed in the density 180 functional theory (DFT) computed, energy-minimized geometry of the precursor 3 (Figure S12). 181

DFT calculations on the π -conjugated backbone revealed 182 183 extended π -delocalization on **BN-F** and **BN-Ph**. The visualized 184 highest occupied molecular orbitals (HOMOs) of both 185 molecules were delocalized over the entire conjugated 186 backbones with non-negligible contributions from the boron 187 centers and the ligands (Figure S11). The majority of LUMOs 188 were on the electron-deficient BTH units and the boron 189 centers. Time-dependent DFT calculations afforded multiple 190 optical transitions with significant intensities, including the 191 lowest band gap HOMO \rightarrow LUMO transitions, and the 192 higher-energy HOMO-1 \rightarrow LUMO transitions. These 193 transitions matched well with experimental spectra. Both 194 BN-F and BN-Ph possessed low band gap NIR absorptions 195 and fluorescent emissions. For example, the UV-vis-NIR 196 spectrum of a dilute solution of BN-F in CH₂Cl₂ showed the 197 low-energy HOMO \rightarrow LUMO absorption peak at 695 nm (ε = 198 $1.17 \times 10^4 \text{ M}^{-1} \cdot \text{cm}^{-1}$) (Figure 2a). For BN-Ph with phenyl 199 ligands on the boron center, such HOMO \rightarrow LUMO band 200 red-shifted into the NIR region ($\lambda_{max} = 768 \text{ nm}, \varepsilon = 1.80 \times 10^4$ 201 $M^{-1} \cdot cm^{-1}$) (Figure 2b). The presence of vibrational 202 progression in both spectra indicated the rigid nature of BN-203 F and BN-Ph.^{1,2} Fluorescence emission of BN-F and BN-Ph 204 appeared in the NIR region with λ_{max} at 720 and 812 nm in 205 CH₂Cl₂, respectively (Figure 2a and b). Small Stokes shifts (576 cm⁻¹ for BN-F and 705 cm⁻¹ for BN-Ph) also implied 206 207 their backbone rigidity, due to the lack of energy loss 208 associated with conformational changes during the excita-209 tion-emission processes. The fluorescence quantum yields of 210 BN-F and BN-Ph were measured using reference standards (zinc phthalocyanine for BN-F and indocyanine green for BN-211 212 Ph) to be 2.8% and 1.2% in CH₂Cl₂, respectively.

On the basis of the molecular design, **BN-F** and **BN-Ph** were texpected to undergo multistage electron-transfer processes. Indeed, cyclic voltammograms (CVs) unveiled four wellelectron-transfer processes in both **BN-F** and **BN-Ph** (Figure 3), indicating five accessible redox states for both compounds. The excellent reversibility of these processes indicated the high stability of all the states including



Figure 2. UV-vis-NIR absorption (solid lines) and emission (dashed lines) of (a) BN-F and (b) BN-Ph in CH_2Cl_2 with transition energies calculated by time-dependent DFT (TD-DFT) using B3LYP/6-311g(d,p) with CH_2Cl_2 CPCM solvation.

radical anions (-1), dianions (-2), radical cations (+1), and 220 dications (+2). The measured electrochemical band gaps of 221 **BN-F** and **BN-Ph** (1.81 and 1.62 eV, respectively) matched 222 well with the optical band gap (1.78 and 1.60 eV, respectively). 223 Among these two voltammograms, the redox processes of **BN-** 224 **Ph** appeared cathodically shifted (Figure 3b) compared to that 225 of **BN-F**, likely due to the more electron-rich nature of the 226 phenyl ligand compared to fluoride. Such an effect was more 227 significant on the oxidation processes, leading to a narrower 228 band gap of **BN-Ph** than that of **BN-F**. 229

All the anodic and cathodic peaks of BN-F and BN-Ph 230 during both oxidation and reduction sweeps were well- 231 separated, suggesting the good stabilities of the radical anion 232 and radical cation intermediates toward disproportionation. 233 For example, the potential gaps between the two reduction 234 waves for both BN-F and BN-Ph were ~0.3 V (Figure 3a and 235 b). These values corresponded to large radical comproportio- 236 nation constants of the reduced forms $(K_{\text{com-re}})$ of 5.41 \times 10⁴ 237 for BN-F and 1.74×10^5 for BN-Ph at 25 °C (equilibrium 1 in 238 Figure 3d), indicating a highly stable radical anion. On the 239 oxidation side, the potential gaps between the two oxidative 240 processes were even larger (0.45 V for BN-F and 0.51 V for 241 BN-Ph), corresponding to remarkably high radical compro- 242 portionation constants ($K_{\text{com-ox}}$) of 4.04 × 10⁷ and 4.17 × 10⁸ 243 at 25 °C, respectively (equilibrium 2 in Figure 3d). Such high 244 stabilities of the radicals were attributed to the strong spin- 245 delocalization on the rigid backbones of BN-F and BN-Ph as 246 designed. In contrast, precursor 3 without B ← N 247 coordination, as a control, showed overlapping two-electron 248 oxidation processes (Figure 3c) and irreversible reduction 249 processes. Such a drastic difference on the redox behaviors 250 between BN-F/BN-Ph and 3 demonstrated profound changes 251

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Figure 3. CV curves of (a) **BN-F**, (b) **BN-Ph**, and (c) **3** in CH_2Cl_2 [0.10 M tetrabutylammonium hexafluorophosphate (TBAPF₆) as the electrolyte; Ag/AgCl as the reference electrode; scan rate = 100 mV/s]. The half-wave potentials of each redox process were noted in (a) **BN-F** and (b) **BN-Ph**. (d) Comproportionation equilibriums of radical cations and radical anions.

252 in the electronic structures after the installment of the B \leftarrow N 253 coordination. These electrochemical data demonstrated the 254 highly efficient stabilization effect of the B \leftarrow N coordination 255 on the oxidized/reduced states through extending charge/spin 256 delocalization.

The excellent redox reversibility allowed spectroelectro-257 258 chemical measurements on BN-Ph, i.e., UV-vis-NIR 259 absorption spectra recorded during each of the electrochemical 260 redox processes, which gave highly reversible multicolor 261 electrochromism (Figure 4a). According to time-dependent 262 DFT calculations, these drastic color changes were attributed 263 to the substantial differences on the transition energy diagrams of the different redox states (Tables S4 and S5). When the 2.64 265 applied potential swept from 0.70 to 1.05 V (vs Ag/AgCl), 266 BN-Ph was gradually oxidized into the radical cation form BN-267 Ph^{•+}. During this process, the HOMO-LUMO transition of 268 the neutral BN-Ph at 768 nm diminished, while a new NIR 269 absorption at 1180 nm emerged (Figure 4b). This low-energy 270 absorption band corresponded to the transition of HOMO- β \rightarrow LUMO- β of the delocalized radical cation (Figure 4d), 271 272 according to DFT calculation. On the basis of the cutoff shape 273 of this NIR absorption peak, BN-Ph⁺⁺ can be categorized as a 274 class III mixed valence system,⁵⁵ in which the two nitrogen-275 centered redox moieties in ICBZ were strongly coupled with 276 one another. The absorption spectra exhibited no significant

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change from 1.05 to 1.25 V, indicating the good stability of BN-Ph⁺⁺ over a wide range of potentials. When BN-Ph⁺⁺ was oxidized into the dication form BN-Ph²⁺ by increasing the potential from 1.30 to 1.50 V, the characteristic radical cation absorption peak at 1180 nm gradually diminished, while multiple absorption bands covering a broad visible light region emerged (Figure 4c). To test the reversibility of these oxidation processes, iterative potential sweepings were conducted between +0.20, +1.10, and +1.50 V for >15 cycles (Figure 4h). Excellent reversibility was demonstrated without notable degradation of the absorption intensity (monitored at 470 nm).

Similarly, the radical anion and dianion of BN-Ph were 289 investigated by spectroelectrochemistry study. Upon the 290 application of a negative potential of -0.75 V, BN-Ph started 291 to be reduced as the absorption intensity at 768 nm decreased 292 (Figure 4e) while a highly intensive peak at 677 nm and a weak 293 absorption peak at ~940 nm emerged, corresponding to the 294 optical transitions, HOMO-1- $\alpha \rightarrow$ LUMO- α and HOMO- β 295 \rightarrow LUMO- β of radical anion **BN-Ph**^{•-} (Figure 4g). The 296 Gaussian shape of this low energy peak indicated that BN-Ph^{•-} 297 was a class II mixed valence system, in which the two reduced 298 BTH moieties were coupled in a weaker manner.^{55,56} Further 299 decrease of the potential from -1.05 to -1.35 V resulted in the 300 second step of reduction into the dianion, BN-Ph²⁻, 301 accompanied by the diminishing of all the peaks in the long- 302 wavelength region and the emerging of a higher-energy peak 303 centered at 544 nm as the new HOMO \rightarrow LUMO transition 304 (Figure 4f). Again, excellent reversibility of the reduction 305 processes was confirmed by monitoring the absorption 306 intensity at 680 nm while sweeping the potential iteratively 307 between -0.30, -1.05, and -1.40 V (Figure 4i) for >15 cycles. 308 Spectroelectrochemical measurements were also conducted in 309 a CH₂Cl₂ solution of BN-F, which showed a similar reversible 310 multistage electrochromism over a wide range of electric 311 potentials (Figure S9). 312

To shed light on the mechanism of these remarkable redox 313 and electrochromic activities of BN-Ph and BN-F, DFT 314 computations were performed starting with geometry opti- 315 mization [B3LYP/6-311g(d,p)] for all five redox states of BN- 316 Ph in the gas phase. At the neutral state of BN-Ph, the lengths 317 of bonds c, d, and e in the central ring "A" (1.412-1.427 Å) 318 were close to that in a benzene ring (Figure 5a). Upon 319 f5 oxidation into the radical cation BN-Ph+, DFT calculation 320 showed that the bond length alternation (BLA) between these 321 three bonds increased, indicating a decrease of aromaticity. 322 The BLA increased further in the diradical form BN-Ph²⁺ 323 (Figure 5b), so that bonds c and e were elongated to 1.451 and 324 1.438 Å, respectively, while bond d was shortened to 1.388 Å, 325 close to the bond length of a typical C=C double bond. 326 Additionally, the C=N double bonds b and f were shortened 327from 1.366 Å in BN-Ph to 1.330 Å in BN-Ph²⁺. This 328 computational study revealed a tendency toward quinonoid 329 character of the ICBZ unit after oxidation (Figure 5e). In 330 contrast, optimized geometries of the reduced forms exhibited 331 that a different quinonoid structure was formed in between the 332 two BTH units involving bonds *j*, *k*, *l*, *e*, and *m* (Figure 5c). In 333this case, bonds *i*, *l*, and *m* were shortened, accompanied by 334increased lengths of bonds e and k, during the transformation 335 into BN-Ph^{•-} and BN-Ph²⁻. These computational results 336 suggested that, in the B \leftarrow N bridged molecule, two different 337 yet orthogonal pathways of BLA changes were accessible to 338 two different quinonoid constitutions during the oxidation and 339



Figure 4. (a) Photographs of the BN-Ph solutions at different redox states in the presence of a honeycomb working electrode in a spectroelectrochemical cell (in CH_2Cl_2 with 0.10 M TBAPF₆). UV–vis–NIR absorption changes of the BN-Ph solutions upon stepwise applications of potentials on the honeycomb working electrode (step height = 0.05 V, vs Ag/AgCl) from (b) +0.70 to +1.05 V, (c) +1.25 to +1.50 V, (e) -0.70 to -1.05 V, and (f) -1.05 to -1.35 V. Molecular frontier orbitals and calculated transition energies of (d) BN-Ph⁺⁺ and (g) BN-Ph⁺⁻ [UB3LYP/6-311g(d,p) with CH_2Cl_2 CPCM solvation]. Multiple cycles of absorption changes of the BN-Ph solutions when the potential was switching between (h) 0.20, 1.10, and 1.50 V and (i) -0.30, -1.05, and -1.40 V.



Figure 5. (a) Redox-active core structure of **BN-Ph** labeled with bond indices from *a* to q_j (b) DFT calculated lengths of the bonds a-g before and after oxidation; (c) DFT calculated lengths of the bonds *e* and h-o before and after reduction [B3LYP/6-311g(d₁p)]; (d) calculated HOMA and NICS(1)_{zz} values for the central ring "A" of **BN-Ph** in different redox states. (e) Constitutional structures of the five different redox states of **BN-Ph**.



Figure 6. Single-crystal X-ray structures of (a) **BN-Ph**, (b) **BN-Ph**^{•+}**SbCl**₆⁻, and (c) **BN-Ph**²⁻[**CoCp** $*_{2}^{+}$]₂ and bond lengths of the area around the central ring "A". Hydrogen atoms and solvent molecules were omitted for clarity. Thermal ellipsoids are scaled to the 50% probability level.

340 reduction (Figure 5e), respectively, contributing to the 341 excellent stability in all the oxidized and reduced forms of 342 **BN-Ph.** Considering the noncovalent nature of $B \leftarrow N$ 343 coordinate bonds, the molecular geometries were also 344 optimized at the level of B3LYP/TZVP with D3 version of 345 Grimme's dispersion correction, which revealed similar 346 structural transformations (Figure S15). Similar BLA changes 347 and structural transformations were also observed in the 348 optimized molecular geometries for different redox states of 349 **BN-F.** In addition, the aromaticity of ring "A" was calculated 350 using the harmonic oscillator model of aromaticity (HOMA) 351 and the nucleus-independent chemical shift (NICS) val-352 ues⁵⁷⁻⁵⁹ (Figure 5d). Both calculations confirmed a strong 353 aromaticity of ring "A" in the neutral state and decreased 354 aromaticity upon oxidation or reduction.

To validate these DFT computational results, the bond 355 356 lengths and molecular geometries were examined by using 357 experimental data from single-crystal X-ray diffraction of the 358 neutral BN-Ph, the radical cation salt BN-Ph^{•+}SbCl₆⁻, and the 359 dianion salt $BN-Ph^{2-}[CoCp^*{}_2^+]_2$ (Figure 6). The radical 360 cation was chemically accessed by one-electron oxidation of 361 BN-Ph with 1.2 equiv of tris(4-bromophenyl)ammoniumyl 362 hexachloroantimonate (Magic Blue) in a CH₂Cl₂ solution. The 363 single crystals suitable for X-ray diffraction were grown by slow 364 vapor diffusion of pentane into a CH2Cl2 solution of BN- $_{365}$ Ph^{•+}SbCl₆⁻. The dianion, on the other hand, was chemically 366 prepared after two-electron reduction of BN-Ph by the ₃₆₇ addition of 3.0 equiv of decamethylcobaltocene ($CoCp*_{2}$). 368 Vapor diffusion of diethyl ether into a concentrated solution of 369 BN-Ph²⁻ in acetonitrile yielded single crystals suitable for X-370 ray diffraction analysis. Bond lengths measured from the single-371 crystal structures of BN-Ph, BN-Ph⁺SbCl₆, and BN-372 $Ph^{2-}[CoCp*_{2}^{+}]_{2}$ matched well (Figure 7) with the DFT 373 calculated bond lengths, further confirming the mechanism of 374 the orthogonal quinonoid transformations upon oxidation and 375 reduction, respectively. The average difference between the $_{376}$ bond lengths calculated at the level of B3LYP/6-311g(d,p) and

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Figure 7. Electron paramagnetic resonance (EPR) spectra of (a) BN-Ph^{•+} in CH₂Cl₂ (0.13 mM) and (c) BN-Ph^{•-} in acetonitrile (0.03 mM) at 288 K. DFT computed spin density maps (isovalue = 0.0008) of (b) BN-Ph^{•+} and (d) BN-Ph^{•-} [UB3LYP/6-311g++(d,p)].

experimentally measured bond lengths was only 0.010 Å (see 377 Table S6 for a detailed list). Such an average difference 378 obtained from calculation at the level of B3LYP/TZVP with 379 D3 version of Grimme's dispersion correction was also ~0.010 380 Å (see Table S7 for a detailed list), suggesting highly reliable 381 computation results. The only exception was bond *d*, which 382 was longer in the neutral state (1.444 Å) from the experimental 383 result than that from the calculation, likely because of the steric 384 strain between ICBZ and BTH units. In both **BN-Ph⁺⁺** and 385 **BN-Ph²⁻**, the B \leftarrow N coordinate bonds retained their strength 386 so that the rigid backbone scaffolds were maintained. In fact, 387 the B \leftarrow N coordinate bond was even shorter in **BN-Ph²⁻** 388 (1.619 Å) as a result of the stronger Lewis basicity of the 389 nitrogen center on BTH after reduction. Moreover, the 390



Figure 8. NBO plots of hyperconjugative interactions in different redox states of BN-Ph (a, b, c, d) and BN-F (e, f, g, h).

³⁹¹ backbone of **BN-Ph²⁻** was more coplanar compared to that of ³⁹² **BN-Ph** (dihedral angles between ICBZ and BTH units ³⁹³ reduced to 6.3°), due to an enhanced double-bond character ³⁹⁴ of bonds *j* and *m*.

Additional characterizations were performed on the reduced 395 396 and oxidized states of BN-Ph in order to further understand 397 their electronic structures during the redox processes. Electron paramagnetic resonance (EPR) spectroscopy of the BN-398 399 Ph^{•+}SbCl₆⁻ solution showed a broad resonance signal (Figure 400 7a). The spin concentration $(7.8 \times 10^{17} \text{ spin/mL})$ was close to 401 the theoretical value $(8.9 \times 10^{17} \text{ spin/mL})$ when assuming the 402 formation of an organic monoradical (S = 1/2) on each 403 molecule, indicative of a high conversion from BN-Ph to BN-404 Ph^{•+}. The lack of hyperfine splitting in the EPR spectrum was 405 attributed to the delocalized spin that experienced multiple 406 couplings with a number of nuclei in this molecule,³⁷ in 407 accordance with DFT computed substantial delocalization of the unpaired spin density (Figure 7b and Figure S13). 408

409 Overall, experimental investigations together with computa-410 tional studies clearly revealed the underlying mechanism of the 411 unique redox processes of ladder-type **BN-Ph** molecules, 412 which involved two distinct pathways of constitutional changes 413 from a benzenoid structure into two types of quinonoid 414 structures upon oxidation and reduction, respectively. The 415 oxidation-active moiety and reduction-active moiety were 416 orthogonally compacted into a single molecular scaffold, giving 417 rise to the reversible multistage redox processes. In addition, B 418 ← N coordination promoted the rigid ladder-type structure in 419 which separated redox centers were strongly coupled with each 420 other, and the unpaired electron of radicals was stabilized by 421 extensive delocalization.

In **BN-Ph** and **BN-F**, the boron centers stabilized the redox state through a number of different mechanisms, including spin coupling, extending pi-delocalization, and modulating electron designed the electronic structures of these molecules molecules impacted the electronic structures of these molecules through hyperconjugation similar to that existing in cyclohexaly3-diene (Figure 1b), providing additional stabilization effects. According to the crystal structures of **BN-Ph**, **BN-Ph**^{•+}, and **BN-Ph**²⁻, the two phenyl ligands on the boron center adopted different bonding orientations: one of the B–C σ-bonds was almost perpendicular to the conjugated backbone (denoted as the "axial" bond), while the other one was almost in the plane

of the backbone (denoted as the "equatorial" bond). Such an 434 unsymmetrical geometry of the boron ligand (B—L, L = Ph or $_{435}$ F) bonds was also predicted by DFT optimized structures of 436 both BN-Ph and BN-F. The nonequivalent bonding 437 orientations suggested hyperconjugative interactions between 438 the axial bonds and the conjugated backbone in these ladder- 439 type molecules. To unveil the hyperconjugations of these axial 440 B—L σ/σ^* orbitals, natural bond orbital (NBO) analysis was 441 performed on BN-Ph, BN-F, and their dianionic and dicationic 442 forms.^{48,60} The strengths of these interactions can be 443 correlated with second-order perturbation energies $[E^{(2)}]$ 444 computed from NBO analysis.^{61,62} In the neutral state of 445 **BN-Ph**, a hyperconjugative donor-acceptor interaction $[E^{(2)} = 446]$ 4.18 kcal/mol] was identified from the lone pair [lp(N)] on 447 the ICBZ unit to the antibonding $[\sigma^*(B-C)]$ orbital on the 448 axial position (Figure 8a).⁶¹ After the two-electron oxidation, 449 f8 the quinonoid ICBZ unit of BN-Ph²⁺ lost the lone pair on the 450 nitrogen center and had a lower electron density. Despite a 451 weaker electron-donating ability, the π (C–N) orbital of the 452 quinonoid ICBZ still interacted hyperconjugatively $[E^{(2)} = 2.05 \ 453]$ kcal/mol] with the axial $\sigma^*(B-C)$ orbital (Figure 8b). More 454 interestingly, a unique back-donating interaction $[E^{(2)} = 2.70 455]$ kcal/mol] was observed from the $\sigma(B-C)$ orbital to the 456 $\pi^*(C-N)$ orbital of the electron-deficient, quinonoid ICBZ 457 unit (Figure 8c), reinforcing the hyperconjugative stabilization 458 effect by diluting the positive charges on the dicationic 459 backbone of BN-Ph²⁺. In the reduced form of dianionic BN- 460 **Ph**²⁻, the NBO analysis showed that the $\sigma^*(B-C)$ orbital also 461 served as an acceptor to stabilize the negative charges on the 462 reduced BTH units. In BN-Ph²⁻, bonds p and q (Figure 5a) 463 possessed a partial double-bond character (1.387 Å in the 464 crystal structure) so that their π orbitals participated in a 465 hyperconjugation $[E^{(2)} = 3.27 \text{ kcal/mol}]$ with the $\sigma^*(B-C)$ 466 orbital (Figure 8d). In this case, the hyperconjugation further 467 delocalized the negative charges and, therefore, enhanced the 468 stability of the highly charged dianion.

In **BN-F** with the fluoride ligands, NBO analysis at different 470 redox states demonstrated similar but even stronger hyper- 471 conjugative interactions between the axial $\sigma^*(B-F)$ orbital and 472 the conjugated backbone, as a result of the high electro- 473 negativity of fluorine atoms (Figure 8e-h). In the optimized 474 structure of **BN-F**, the axial $\sigma^*(B-F)$ orbitals withdrew the 475 electron density through hyperconjugation [$E^{(2)} = 7.36$ kcal/ 476

477 mol] from the electron lone pair of the nitrogen atom on the 478 ICBZ unit (Figure 8e). Meanwhile, another donor-acceptor 479 interaction of 2.84 kcal/mol was found in BN-F between the 480 π (C–N) bond in the BTH unit and the axial σ^* (B–F) orbital 481 (Figure 8f). Therefore, these profound hyperconjugative 482 effects in BN-F, together with the inductive effect from 483 fluorine atoms, contributed to its low-lying HOMO and 484 LUMO levels compared to BN-Ph.

According to DFT calculation, these hyperconjugative 485 486 interactions were also present in the paramagnetic radical 487 cations and radical anions. Isotropic Fermi couplings of the 488 axial ligands with the already delocalized spin were much 489 higher than those of the equatorial ligands, indicating strong 490 interactions between the axial bond with the backbone through 491 hyperconjugation. Overall, the hyperconjugative stabilization 492 effect further assisted the desirable charge and spin 493 delocalization during the redox processes. Meanwhile, such 494 orbital interactions provided an additional mechanism to 495 impact the electronic structures and properties of the entire 496 ladder-type molecules.

497 CONCLUSION

498 In conclusion, we demonstrated herein a molecular design 499 strategy to achieve highly reversible, multistage redox activities 500 and multicolor electrochromism in compact ladder-type 501 molecules bridged with $B \leftarrow N$ coordination. The structural 502 transformations from a benzenoid constitution into two 503 distinct types of quinonoid constitutions during both reduction 504 and oxidation processes were elucidated, giving a clear 505 mechanistic picture of these robust multistage electron-transfer 506 processes. Combined theoretical and experimental investiga-507 tions demonstrated that the $B \leftarrow N$ coordination played a 508 pivotal role in rendering the remarkable redox properties of 509 these molecules by extending the charge and spin delocaliza-510 tion and by enforcing the rigid conformation. We systemati-511 cally established a hyperconjugation mechanism that impacted 512 the electronic structures and further stabilized the different 513 redox states. These results advanced fundamental knowledge of s14 sp³ boron-containing π -systems, providing practical design 515 principles for the development of p-block element-derived 516 molecules and macromolecules with exotic optical, electronic, 517 and spin properties, such as fully fused ladder polymers 518 bridged by these $B \leftarrow N$ coordinate bonds.

519 **ASSOCIATED CONTENT**

S20 Supporting Information

521 The Supporting Information is available free of charge on the 522 ACS Publications website at DOI: 10.1021/jacs.8b11337.

- General methods, synthesis, characterization data, and 523
- computational data (PDF) 524
- Crystallographic data (CIF) 525
- Crystallographic data (CIF) 526
- Crystallographic data (CIF) 527
- DFT calculation optimized structures (XYZ) 528
- DFT calculation optimized structures (DOCX) 529

AUTHOR INFORMATION 530

- 531 Corresponding Author
- *fang@chem.tamu.edu 532
- 533 ORCID 0
- 534 Congzhi Zhu: 0000-0002-1302-7187
- 535 Yi Liu: 0000-0002-3954-6102

539

552

- Lei Fang: 0000-0003-4757-5664 536 Notes 537
- The authors declare no competing financial interest. 538

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