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Two-way chromic interconversion of the 2,2'biphenol-6,6'-diyl dication with 5H,10H-dioxapyrene or 9H,10H-4,5-dihydroxyphenanthrene†

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Two-proton or two-electron transfer of the title biphenolic dication proceeds nearly simultaneously to induce 2,6'/2',6- or 6,6'-bond formation to give dioxapyrene or dihydrophenanthrene derivatives, respectively, with vivid changes in color (halochromism and electrochromism).

5H,10H-Dioxapyrene (diopy; 5H,10H-[1]benzopyrano[5,4,3-cde][1]benzopyran) is a less-studied heterocyclic skeleton¹ in contrast to its 5,10-dione analogue. We envisaged that the flattened framework of diopy provides a unique opportunity for the development of new bistable molecular response systems, in which an external stimulus induces the cleavage of two C(sp₃)-O bonds to transform into the corresponding biphenyl derivatives with a twisted conformation (Scheme 1). Upon acid treatment of diopy (A) with cation-stabilizing substituents at 5,10-positions, the biphenolic dication (\mathbb{C}^{2+}) would be generated *via* the monocationic intermediate (B⁺). When B⁺ suffers from severe steric repulsion at the bay-region, this amphiprotic species easily undergoes acid-base disproportionation to A and C2+, so that double protonation/ deprotonation between A and C2+ would occur nearly simultaneously. Such a simplified pseudo-two-state switching is favored

Scheme 1 $\,$ Interconversion of diopy A and biphenolic dication C^{2+} upon double protonation/deprotonation via intermediate B+.

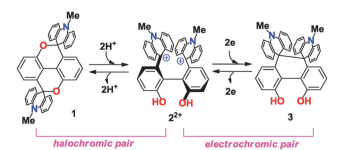
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† Electronic supplementary information (ESI) available: Experimental procedure and characterization data. Supplementary figures of X-ray single-crystal structure analyses, halochromic titration, and cyclic voltammograms. CCDC 1061367, 1061368 and 1061369. For crystallographic data in CIF or other electronic format,

for the construction of promising molecular response systems with a sharp ON/OFF threshold.² When the cationic part in C²⁺ is endowed with a strong absorption in the visible region, interconversion between A and C2+ is accompanied by halochromism,³ since diopy A shows absorptions only in the UV region.

To generate and isolate the dicationic state as a stable entity despite the presence of hydroxy groups within the molecule, the cationic subunit should have a large pK_{R+} value, which prompted us to select the 10-methylacridinium chromophore^{4,5} (Scheme 2). Due to the bulkiness of the chromophore, the biphenol skeleton in 22+ would have a large torsion angle, whereas the diopy skeleton in 1 would be nearly planar, since the spiro(10-methylacridan) units do not induce any steric hindrance. Such a drastic structural change would realize the two-state halochromic interconversion between 1 and 22+. Another interesting point is that, upon reduction, dication 2^{2+} would be transformed into a dihydrophenanthrene (DHP) derivative 3 accompanied by C(sp₃)-C(sp₃) bonding through "dynamic redox (dyrex)" behavior,6 and the interconversion between 22+ and 3 would also exhibit characteristic color and structural changes. Thus, 1, 22+ and 3 can serve as a novel motif for multi-input molecular response systems.⁷

Here we report the preparation and X-ray structures of 1 and 2^{2+} along with their chromic behavior during the interconversion between 1 and 2²⁺ (halochromic³ pair) as well as 2²⁺ and 3 (electrochromic⁸ pair).



Multi-input chromic behavior of diopy 1, biphenolic dication 22+, Scheme 2 and DHP 3.

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Scheme 3 Preparation scheme for 1, $2^{2+}(BF_4^-)_2$ salt, and 3.

6,6'-Dibromo-2,2'-biphenol 4^9 was first reacted with methoxymethyl (MOM) chloride/NaH in DMF to give MOM-protected biphenol 5^{10} in 77% yield. The dilithio derivative derived from 5 and 4 equiv. of 'BuLi in THF was then reacted with 10-methyl-9(10*H*)-acridone to give bis(hydroxy)base 6^{10} in 70% yield. Upon treatment of 6 with HBF $_4$ in MeOH–CHCl $_3$ at reflux afforded the desired $2^{2+}(BF_4^-)_2^{10}$ as yellow-orange crystals in 95% yield. The reaction of $2^{2+}(BF_4^-)_2$ with Et $_3$ N in MeCN gave colorless crystals of diopy 1^{10} in 88% yield (Scheme 3).

Based on the results of an X-ray analysis 11 at 150 K, the diopy core in 1 is nearly planar (largest deviation of an atom from the mean plane: 0.23 Å), although the pyran rings adopt a very shallow twist-chair form (Fig. 1 and Fig. S1, ESI†). The two benzene rings are coplanar (dihedral angle: 0°). To this core are attached the spiro(10-methylacridan) units, which are slightly deformed into a butterfly-shape [dihedral angle between two benzene nuclei of acridan: 21.3(2)°], as found in other structurally-related molecules. 12 In contrast, the two molecular halves in dication 2²⁺ are largely twisted in the crystal of (BF₄⁻)₂ salt¹¹ (Fig. 2 and Fig. S2, ESI†). The dihedral angle of the biphenyl unit is 68.8(1)° (syn-form), and there are no signs to indicate coordination of the hydroxy groups to the acridinium chromophores. If we consider that the two oxygen atoms at the 2,2'-positions are separated by a distance of 3.050(2) Å, intermolecular H-bonding is not effective in 22+ (typical distance for

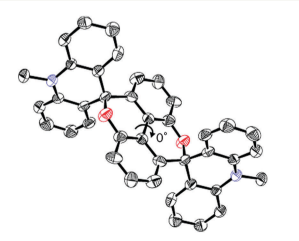


Fig. 1 ORTEP drawing of diopy 1 determined by X-ray analysis at 150 K.

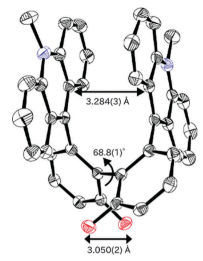


Fig. 2 ORTEP drawing of ${\bf 2^{2+}}$ in ${\bf 2^{2+}}({\rm BF_4}^-)_2$ salt determined by X-ray analysis at 150 K.

the H-bonded O···O: 2.75 ± 0.2 Å). The π - π interaction between two acridinium units must be the major directing force to give the observed *syn*-form (Fig. S2 and S3, ESI†),¹³ and thus the chromophores are stacked nearly in parallel [dihedral angle: $3.92(3)^{\circ}$] with the shortest C···C contact of 3.284(3) Å (sum of van der Waals radii: 3.40 Å).

Diopy 1 is colorless, with absorptions only in the UV region $[\lambda_{\text{max}}/\text{nm}: 339 (4.30) \text{ in } CH_2Cl_2], \text{ whereas } 2^{2+} \text{ exhibits a yellow-}$ orange color [358(3.92) in MeCN] due to the characteristic absorptions of acridinium (Fig. 3a). Although 10-methylacridinium itself is highly fluorescent, 22+ is non-fluorescent due to the chargeshift-type quenching of the excited state by the electron-donating biphenol unit. Upon the aliquot addition of TfOH to a DMSO- d_6 solution of 1, a clean conversion to 22+ was observed (Fig. S4, ESI†). The resulting spectra showed the presence of only two species (1 and 22+), which demonstrated that the steady-state concentration of the intermediary monocationic derivatives is negligible. The halochromic response was examined by the repeated addition of TfOH (100 microL) to a DMSO solution of 1 (1.2 \times 10⁻⁵ M), followed by the addition of Et₃N (200 microL) to the solution of as-generated 22+. By monitoring the color change using UV-vis spectroscopy, we could confirm the reversibility of the present halochromism (Fig. 3b and Fig. S5, ESI†).

According to the results of a voltammetric analysis, 14 3 undergoes irreversible two-electron oxidation at an anodic peak potential $(E_{\rm pa})$ of +0.32 V in CH₂Cl₂/MeCN (4:1) ν s. SCE (Fig. S6a, ESI†). The return peak was observed in the far cathodic region $(E_{\rm pc}=-0.23~{\rm V})$, which corresponds to the reduction process of dication 2^{2+} (Fig. S6b, ESI†). In fact, Zn-reduction of 2^{2+} (BF₄ $^-$)₂ induced C(sp₃)–C(sp₃) bonding at the C6 and C6' positions to give DHP 3. Colorless crystals of 3 $[\lambda_{\rm max}/{\rm nm}: 285~(4.37)$ in CH₂Cl₂] were isolated in 91% yield, and regenerated 2^{2+} (BF₄ $^-$)₂ in 87% yield upon treatment with 2 equiv. of ferrocenium tetrafluoroborate in CH₂Cl₂/MeCN. In this way, reversible redox interconversion between 2^{2+} and 3 accompanied by C–C bond formation/cleavage ("dyrex" behavior) was confirmed. Due to the dynamic geometrical changes, 15 two-electron transfer occurs nearly simultaneously,

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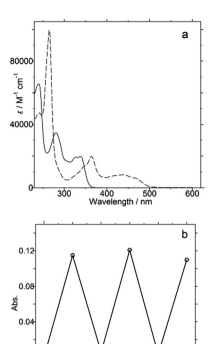


Fig. 3 (a) UV-vis spectra of 1 (solid line) in CH₂Cl₂ and 2²⁺(BF₄⁻)₂ (dashed line) in MeCN. (b) UV-vis spectral changes at 440 nm for halochromic switching from $\mathbf{1}$ [1.2 \times 10⁻⁵ M; 3 mL] to $\mathbf{2^{2+}}$ upon addition of TfOH (100 microL) in DMSO. The reverse conversion was accomplished upon addition of $\mathrm{Et}_3\mathrm{N}$ (200 microL). The halochromic cycles could be repeated without significant loss of response

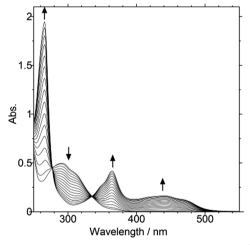


Fig. 4 A continuous change in the UV-vis spectra of **3** [2.1×10^{-5} M; 3 mL] to 22+ in CH₂Cl₂/MeCN (4:1) containing 0.05 M Bu₄NBF₄ upon constantcurrent electrochemical oxidation on a Pt electrode (30 microA, every 1 min).

which was confirmed by the negligible steady-state concentration of the intermediary cation radical upon the electrochemical conversion of 3 to 2^{2+} (Fig. 4).

In this work, we have demonstrated the reversible halochromic and electrochromic interconversion of 2,2'-biphenyl-6,6'-diyl dication with two kinds of neutral molecules (diopy and DHP).

This is the first example of concomitant but independent twoproton or two-electron transfer with a negligible concentration of the intermediates. A drastic structural change is the key to this novel feature, which may represent a new molecular design concept for multi-input response systems with advanced features.

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