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Insights into the radical-radical and radical-substrate dimerization processes for substituted phenylmethyleneepyrans

Laurianne Wojcik, a Isidoro Lopez, a Sébastien Gauthier, b Nolwenn Cabon, b Pascal Le Poul, b Frédéric Gloaguen, a* Nicolas Le Poul a*

a Laboratoire de Chimie, Electrochimie Moléculaire et Chimie Analytique (CNRS UMR 6521), Université de Bretagne Occidentale, 6 Avenue Le Gorgeu, CS 93837, 29238 Brest, France

b IUT de Lannion, CNRS UMR 6226, Institut des Sciences Chimiques de Rennes, Université de Rennes 1, Rue Edouard Branly, 22300 Lannion, France

*Corresponding authors: nicolas.lepoul@univ-brest.fr; frederic.gloaguen@univ-brest.fr

Abstract

The electrochemically-induced C-C bond making/breaking for six differently R-substituted phenylmethyleneepyrans has been investigated by voltammetry in organic media. All compounds display an irreversible oxidation peak whose potential is fully dependent on the electrophilic property of the substituent R. The electrochemical oxidation yields bis-pyrylium compounds by σ-σ C-C bond formation. The initial methyleneepyrans are recovered by cleavage of the C-C bond through electrochemical reduction of the bis-pyrylium species. According to the voltammetric analysis, the mechanistic pathway, radical-radical or radical-substrate, for the intermolecular dimerization is fully R-dependent. Electronic structure
calculations show that the spin population in the radical cation and the strength of the σ-σ C-C bond in the dimer strongly depend on the nature of R. In addition, low-temperature electrochemical voltammetry (175 K), and room-temperature high scan rate cyclic voltammetry have been used to unravel the kinetics of the C-C bond formation.

**Keywords**: radical dimerization; methylenepyran; low-temperature voltammetry; C-C bond making; ab initio calculations.

1. Introduction

The reversible dimerization of organic radicals has for a long time fascinated electrochemists, essentially because the analysis of the experimental data, in particular cyclic voltammetry (CV) and chronoamperometry (CA), can give straightforward information on the mechanism of dimerization as well as the kinetics/thermodynamics of the process [1-8]. Different types of organic dimers can be found according to the mode of interaction between monomeric units, upon redox activation. Most of studies have been focused on organic dimers generated by the formation of σ-σ carbon-carbon bond [9-11]. Also, π-π dimers have been more recently developed with supramolecular organic systems [12-15]. While most of reported studies have been carried out for fundamental purposes, recent works have shown some remarkable applications in the domain of redox switches and electrochromic devices for systems displaying optical properties in the UV-Visible spectroscopic domain [9, 11, 16-20]. Among them, methylenepyrans (MPs) are organic molecules which have been fully exploited as dyes for photovoltaic cells [21], but also as component parts of push-pull compounds for non-linear optic (NLO) applications and luminescent devices [22-24]. As their dithiafulvalene (DTF)
analogues [25], MPs are able to form $\sigma-\sigma$ C-C bound dimers upon oxidation at a relatively low redox potential. The generated bis-pyrylium compounds can be reduced chemically and electrochemically to yield back the initial MPs. Our experimental and computational studies on a series of substituted-MPs have emphasized that the redox and spectroscopic properties of these compounds could be modulated by adequate variation of the substituting group [26-30]. Noteworthy, we recently demonstrated by time-resolved spectroelectrochemistry that R-substituted phenyl-MPs 1a-f (Scheme 1) could be switched between their mono-pyran and bis-pyrylium states by redox triggering over several cycles, leading to a concomitant change of their spectroscopic signatures [26]. Moreover, the electrophilic character of the substituting group R on the phenylmethylen moiety (Scheme 1) was shown to affect in a significant manner the oxidation potential of the MP. Contrarily, the reduction potential of the bis-pyrylium species $(2a-f)^{2+}$ remained almost insensitive to the nature of the R group. These experimental results were fully supported by electronic structure and thermochemical calculations [26]. In complement to these first results, we present here our further investigations on the dimerization mechanism and the kinetics associated to the oxidation of compounds 1a-f, by the combination of electrochemical and computational approaches. In particular, the influence of the R-substituting group on the dimerization process has been deeply explored, with the help of low-temperature and high scan rate cyclic voltammetry.

Scheme 1

**Scheme 1.** C-C bond making/breaking by oxidation of R-substituted phenylmethylenepyrans 1a-f and reduction of their bis-pyrylium derivatives $(2a-f)^{2+}$. 
2. Experimental section

2.1 Synthesis of compounds 1a-f

All syntheses and characterization of the phenylmethylenepyran derivatives 1a-f were previously reported [26].

2.2 Electrochemical methods

The electrochemical studies were performed in a glovebox (Jacomex) (O<sub>2</sub> <1 ppm, H<sub>2</sub>O <1 ppm) with a homemade three-electrode cell (WE, Pt; RE, Ag wire; CE, Pt), such that the volume of the solution (hence concentration) was constant all over the duration of measurement (no need for gas bubbling). Ferrocene was added at the end of each experiment to determine the redox potential values. The potential of the cell was controlled by an AUTOLAB PGSTAT 100 (Metrohm) potentiostat monitored by NOVA software (Metrohm). Dichloromethane was freshly distilled from CaH<sub>2</sub> and kept under argon in the glovebox. The supporting salt NBu<sub>4</sub>PF<sub>6</sub> was synthesized from NBu<sub>4</sub>OH (Acros) and HPF<sub>6</sub> (Aldrich). It was then purified, dried under vacuum for 48 h at 100 °C, and kept under N<sub>2</sub> in the glovebox.

Low-temperature voltammetric measurements were performed with a specific home-designed schlenk-type cell (WE: Pt, RE: Pt wire, CE: Pt wire). The cell was introduced at room temperature under nitrogen in a liquid nitrogen cryostat (LN2 Oxford Instruments) [31]. Temperature was adjusted with a temperature controller (ITC Mercury) connected to the cryostat. The temperature inside the electrochemical cell was also checked by a supplementary thermocouple (Pt1000, VWR, resolution 0.1 K).

2.3 Theoretical methods
Calculations were carried using the so-called Hartree-Fock-3c (HF-3c) method as implemented in the Orca 3.03 software [32, 33]. The HF-3c method includes correction terms for London dispersion interactions, basis set superposition error, and basis set deficiency effects. Geometry optimizations and single-point calculations were carried out with a tolerance of $10^{-8}$ hartree on energy change. Electronic structures of singly charged species (i.e. open shell) were calculated using the unrestricted HF method. The preparation of the input files and analysis of the output files was carried out using the Avogadro 1.03 software [34]. The input structures for the geometry optimization in the gas phase of 1a, 1a$^+$, 1f, 1f$^+$, as well as 2$^+$ and 2$^{2+}$ dimers (for R = NMe$_2$, H or NO$_2$) were manually built from the X-ray structures of 1b and [BisFeMP]$^{2+}$, respectively [26, 30]. A vibrational frequency calculation was carried out on each of the structures to confirm that the stationary point is a minimum in energy and to obtain the free energy correction term. Effect of the solvent (i.e. CH$_2$Cl$_2$) was accounted for using a conductor-like screening model (COSMO) as implemented in the Orca 3.03 software [35].

3. Results and Discussion

3.1 Electrochemical studies

Cyclic voltammetric (CV) studies of compounds 1a-f have been first carried out at a Pt working electrode in dichloromethane at room temperature. As previously stated [26], all PhMPs display the same redox behavior, i.e. a single irreversible oxidation peak on the forward scan at $E_{pa}(1)$ and an irreversible reduction peak on the backward scan at $E_{pc}(2)$ (see Table 1 for electrochemical data), the value of $E_{pa}(1)$ and $E_{pc}(2)$ varying with R. For example, Fig. 1 displays the CV obtained for 1a at $v = 0.1$ V.s$^{-1}$ and $C = 1$ mM. The same current density was obtained for all methyleneypyran compounds [26]. Scanning initially in negative direction, by CV or Rotating Disk Electrode Voltammetry (RDEV), does not show any cathodic process except for the nitro derivative, as previously reported [26] (see Fig. S3 for
CV and RDEV). Exhaustive electrolyses at $E_{pa}(1)$ lead to the formation of the bispyrylium derivatives (2a-f)$^{2+}$[26]. Coulometric measurements indicate that the process is monoelectronic (i.e. two electrons for the oxidation of two molecules of phenylmethyleneepyran). For all R-substituted compounds, the resulting species are irreversibly reduced at $E_{pc}(2)$, in agreement with CV studies.

**Fig. 1**

**Fig. 1.** CV ($v = 0.1 \, \text{V.s}^{-1}$) at a Pt electrode (diam. 3mm) of compound 1a ($C = 1\, \text{mM}$) in CH$_2$Cl$_2$/NBu$_4$PF$_6$ 0.1 M under argon at $T = 293 \, \text{K}$.

<table>
<thead>
<tr>
<th>R</th>
<th>$E_{pa}(1)^a$</th>
<th>$E_{pc}(2)^a$</th>
<th>$dE_{pa}(1)/d\log v^b$</th>
<th>$dE_{pa}(1)/d\log C^c$</th>
<th>Dimerization mechanism$^d$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1a</td>
<td>H 0.22</td>
<td>-0.85</td>
<td>33.3</td>
<td>-30.2</td>
<td>RSD</td>
</tr>
<tr>
<td>1b</td>
<td>NMe$_2$ -0.17</td>
<td>-0.83</td>
<td>21.3</td>
<td>-17.2</td>
<td>RRD</td>
</tr>
<tr>
<td>1c</td>
<td>OMe 0.14</td>
<td>-0.81</td>
<td>22.9</td>
<td>-20.6</td>
<td>RRD</td>
</tr>
<tr>
<td>1d</td>
<td>CO$_2$Me 0.34</td>
<td>-0.77</td>
<td>34.1</td>
<td>-37.9</td>
<td>RSD</td>
</tr>
<tr>
<td>1e</td>
<td>Br 0.29</td>
<td>-0.78</td>
<td>32.7</td>
<td>-31.4</td>
<td>RSD</td>
</tr>
<tr>
<td>1f</td>
<td>NO$_2$ 0.41</td>
<td>-0.73</td>
<td>32.1</td>
<td>-38.0</td>
<td>RSD</td>
</tr>
</tbody>
</table>

$^a$ Data obtained at $v = 0.1 \, \text{V.s}^{-1}$ and $C = 0.5 \, \text{mM}$ with ohmic drop correction; $^b$ Determined from data obtained at $C = 0.5 \, \text{mM}$ for $0.02 \, \text{V.s}^{-1} < v < 1 \, \text{V/s}$ with ohmic drop correction; $^c$ Determined from data obtained at $v = 0.1 \, \text{V.s}^{-1}$ with ohmic drop correction for $0.05 < C < 0.5 \, \text{mM}$. $^d$ Abbreviations for: RSD: Radical-Substrate Dimerization; RRD: Radical-Radical Dimerization.
As previously stated [26], monoelectronic oxidation at $E_{pa}(1)$ leads to the dimerization of the phenylmethylenepyran. The generated bis-pyrylium is further reduced at $E_{pc}(2)$, inducing the cleavage of the generated C-C bond, and yielding back the initial species. The value of $E_{pa}(1)$ is strongly dependent on the electrophilic nature of the R substituting group ($\Delta E_{pa}(1) = 580$ mV). Such a result was previously ascribed to the location of the electronic density on both the substituted phenyl and pyran rings for 1a-f. In contrast, the peak potential value at $E_{pc}(2)$ is less affected by the substituent R ($\Delta E_{pc}(2) = 120$ mV), the highest value being obtained for the nitro compound. This suggests that the electron transfer reaction merely occurs on the pyrilium moiety and not on the substituted phenyl groups, in full agreement with previous theoretical studies [26].

Fig. 2

Fig. 2. Plots of $E_{pa}$ vs. log $v$ for compounds A) 1a (R=H); B) 1b (R=NMe$_2$); C) 1c (R=OMe); D) 1d (R=CO$_2$Me); E) 1e (R=Br); F) 1f (R = NO$_2$); (C = 0.5 mM).

In order to further investigate the C–C bond making/breaking mechanism, variation of the anodic peak potential $E_{pa}(1)$ with voltammetric scan rate and concentration has been measured for 1a-f. Fig. 2 displays plots of $E_{pa}(1)$ against log $v$ for $C = 0.5$ mM obtained for the six organic dyes. Clearly, the variation is linear in agreement with the occurrence of coupled chemical and electrochemical reactions. Interestingly, full analysis of the experimental data in the 0.05 to 1.00 mM concentration range shows that the slope value from ($E_{pa}(1)$ vs. log $v$) plots remains constant up to 0.5 mM (Fig. 3).
The values of the slope at a concentration of 0.5 mM for all studied phenylmethyleneypyran 1a-f are given in Table 1. Two different behaviors are clearly observed. When R = NO₂, CO₂Me, Br and H, a slope of ca. 30 mV/decade is obtained. With better donor groups such as R = NMe₂ and OMe, the slope is rather close to 20 mV/decade. The effect of the R group can also be observed when analyzing the variation of the anodic peak \( E_{pa}(1) \) with the concentration in compound 1a-f at a moderate scan rate (\( v = 0.1 \text{ V.s}^{-1} \)). Plots of \( E_{pa}(1) \) vs. log C are linear, as shown in Fig. 4 for all compounds. However, the analysis gives two different slope values according to the nature of R. Hence, it is close to -20 mV/decade for derivatives bearing electrodonating groups (NMe₂, OMe), while the four other compounds display a -30 mV/decade value (Table 1). Noteworthy, according to the classical treatment of dimerization process, these data indicate that the R moiety has a strong influence on the mechanism of dimerization. With withdrawing groups such as NO₂, dimerization occurs merely through a radical-substrate process (RSD) whereas radical-radical reaction (RRD) is preferred as the donor effect of the R group increases as depicted in Scheme 2.
Scheme 2. Schematic pathway for the C-C bond making/breaking of R-substituted phenylmethylenepyran 1a-f compounds upon electron transfer. Inset: RRD vs. RSD mechanisms.

Further investigation of the dimerization process was then carried out by low temperature electrochemistry, in order to find better conditions to obtain a reversible system by CV for the monoelectronic oxidation of the phenyl-MP into its pyrylium cation. As shown above, variation of the scan rate between 0.02 V.s$^{-1}$ and 5 V.s$^{-1}$, or decrease of the concentration in phenyl-MP (0.04 mM < C < 1 mM) did not induce any appearance of reversibility within the defined ranges, meaning that the dimerization process is fast vs. timescale of the CV measurements. We then took advantage of a cryo-electrochemical setup [31, 36], developed by some of us to slow down the kinetics of the chemical process, for the eventual detection of the reduction peak of the electrochemically-generated radical cation $(1a-f)^{••}$ at $E_{pc}(1)$. The cryo-set-up consists in a home-designed air-tight electrochemical cell which can be introduced in a liquid nitrogen cryostat, hence allowing electrochemical measurements above the freezing point of the liquid electrolyte.

**Fig. 5**

**Fig. 5.** CVs ($v = 0.1$ V.s$^{-1}$) at a Pt electrode (diam. 1mm) of compound 1d (0.04 mM) in CH$_2$Cl$_2$/NBu$_4$PF$_6$ 0.1 M under argon for 175 K < T < 183 K.
Fig. 5 displays the voltammetric response at very low temperatures (175 K < T < 183 K) in CH$_2$Cl$_2$/NBu$_4$PF$_6$ of 1d in a relatively low concentration (C = 0.04 mM) and moderate scan rate (ν = 0.1 V.s$^{-1}$). CV shows that the decrease of the temperature induces a significant increase of the $E_{pa}(1)$-$E_{pc}(2)$ peak-to-peak separation. For instance, this value reaches 1.36 V at T=183 K whereas it is equal to 1.11 V at room-temperature. Such effect can be ascribed to a decrease of the electron transfer kinetics for each process (1$^{+•}$/1 and 2$^{2+}$/2) with $T$. More importantly, the anodic peak at $E_{pa}(1)$ remains chemically irreversible, even at 175 K or by increasing the scan rate (up to 1 V.s$^{-1}$), showing that even under these conditions, the dimerization process remains fast (vs. timescale of the experiment).

We then opted for another approach consisting in increasing substantially the scan rate at room temperature in diluted solutions, such that the kinetics of the C-C bond formation could be attained. An example is given with the dimethylamino derivative 1b for which high scan rate CV at $v$ = 100 V.s$^{-1}$ induces the appearance of a cathodic peak corresponding to the reduction of the electrochemically generated radical cation 1b$^{+•}$ at $E_{pc}(1)$ with concomitant decrease of the reduction peak of the bispyrylium 2b$^{2+}$ (Fig. 6, black curve). However, no oxidation peak of the transient bis-pyran species 2b was detected, even at this scan rate, in agreement with a very fast dissociative C-C bond breaking process.

**Fig. 6**

**Fig. 6.** Experimental (black) and simulated (red) CVs at A) $ν$ = 100 V.s$^{-1}$ and B) $ν$ = 1 V.s$^{-1}$ of compound 1b (C = 0.09 mM) at a Pt electrode (diam. 1mm) in CH$_2$Cl$_2$/NBu$_4$PF$_6$ 0.1 M. (for details, see Appendix A).
Voltammetric analysis of the 1b+/1b redox system at this scan rate allows the determination of the standard potential \( E^0(1) = -0.12 \) V vs Fc. Moreover, according CV analysis (see above), 1b dimerizes through a radical-radical coupling upon oxidation. Hence, the dimerization rate constant, \( k_{\text{dim}} \), can be determined by using the equation (1) [37]:

\[
E_p = E^0 = 0.903 \frac{RT}{F} - \frac{RT}{3F} \ln \left( \frac{4RTk_{\text{dim}}C}{3Fv} \right) \tag{1}
\]

The \( k_{\text{dim}} \) term can be extracted from the intercept value from plots of \( E_{pa} \) vs \( \ln v \). Our calculations give an average value for \( k_{\text{dim}} = 4.2 \pm 0.7 \times 10^7 \) M\(^{-1}\).s\(^{-1}\) averaged from three different concentrations in compound 1b \((C = 0.40 \) mM, 0.19 and 0.09 mM\). The calculated \( k_{\text{dim}} \) value found illustrates a fast dimerization process, in good agreement with previous studies on a dithiafulvene analogue which also performs a radical-radical dimerization upon oxidation \( (k_{\text{dim}} = 2 \times 10^8 \) M\(^{-1}\).s\(^{-1}\)) [25]. Alternatively, voltammetric simulation has been carried out by using the DigiElch\(^\circ\) software by assuming an EC\(_{\text{dim}}\) mechanism and \( k_{\text{dim}} = 9 \times 10^6 \) M\(^{-1}\).s\(^{-1}\) (see Appendix A for details). As shown in Fig. 6, a relatively good match was obtained between the experimental and simulated CVs at two different scan rates \((v = 100 \) V.s\(^{-1}\) and \( v = 1 \) V.s\(^{-1}\)) indicating that the value of \( k_{\text{dim}} \) found by using equation (1) was reasonable.

**3.2 Computational studies**

The electrochemical study provided strong evidence that the mechanism of oxidative dimerization of 1a-f derivatives depends on the nature of the R group, i.e. radical-radical (RRD) coupling for R = NMe\(_2\) and OMe, and radical-substrate (RSD) coupling for R = H, Br, CO\(_2\)Me and NO\(_2\) (Table 1). To rationalize this experimental result, the electronic structures of [\( ^{R}\text{PhMP} \)]\(^0/+\) and [\( ^{\text{Bis}}\text{PhMP} \)]\(^+/2+\) have been calculated for R = NMe\(_2\), H and NO\(_2\). The range of
quantum chemical methods applicable to study the dimerization of \( {^R} \text{PhMP} \) derivatives is restricted by the large size of the dimer (ca. 100 atoms). Here, all electronic structure calculations were carried out using the HF-3c method, which is a fast Hartee-Fock method developed for accurate computation of large organic molecules [32].

The calculated structure of 1b (\( R = \text{NMe}_2 \)) is shown in Fig. S1. Selected values of bond length and angle from the X-ray and calculated structures are displayed in Table S1. The average absolute error on the calculated bond lengths is better than 0.02 Å, indicating that the chosen theoretical method appropriately describes the electronic structure of this family of compounds. Similarly, a good agreement was found between the X-ray structure of [Bis-FcMP]\(^{2+}\) [28, 30] and the calculated structure of 2b\(^{2+}\) (Fig. S2). The length of the C–C bond formed in the dimerization process is 1.53 Å and 1.56 Å in [Bis-FcMP]\(^{2+}\) and 2b\(^{2+}\), respectively. Moreover, the dihedral angle between the two Fc groups trans to the C–C bond of the dimer is 180° and that between the two \( \text{NMe}_2 \) groups is 175°. These results further show that electronic structure calculations are consistent with a \( \sigma \)-dimer, in which the \( {^R} \text{Ph} \) moieties have little steric effects.

**Table 2.** Spin population in the gas phase and solvent (i.e. CH\(_2\)Cl\(_2\)) on the carbon C\(_{30}\) involved in the dimerization of 1a\(^{++}\), 1b\(^{++}\) and 1f\(^{++}\) (see atom numbers on Fig. S1).

<table>
<thead>
<tr>
<th></th>
<th>R</th>
<th>Gas</th>
<th>Solvent</th>
</tr>
</thead>
<tbody>
<tr>
<td>1b(^{++})</td>
<td>NMe(_2)</td>
<td>0.716</td>
<td>0.880</td>
</tr>
<tr>
<td>1a(^{++})</td>
<td>H</td>
<td>0.914</td>
<td>0.938</td>
</tr>
<tr>
<td>1f(^{++})</td>
<td>NO(_2)</td>
<td>0.932</td>
<td>0.953</td>
</tr>
</tbody>
</table>
Since dimerization occurs upon oxidation of compound 1, we first analyzed the spin population on the carbon involved in the formation of the σ-dimer for three selected radical cations 1⁺⁺. As shown in Table 2, the spin population on C₃₀ (see atom numbers on Fig. S1) increases with the electrophilic character of the R group in both gas phase and solvent. Noticeably, the spin density on this carbon atom is significantly larger for 1a⁺⁺ (R = H) or 1f⁺⁺ (NO₂) than for 1b⁺⁺ (R = NMe₂), reaching a value of almost +1 for 1f⁺⁺ in CH₂Cl₂ (Table 2). Such a result suggests that the radical-radical dimerization is less favored in the case of 1a⁺⁺ and 1f⁺⁺, probably because of strong electrostatic repulsion, even in presence of solvent.

As shown in Scheme 2, the coupling of two 1⁺⁺ radicals produces a doubly-charged dimer 2⁺⁺, while the radical-substrate dimerization yields a singly-charged dimer 2⁺ which can be further oxidized. The relative stability of these two different dimers (2⁺ and 2⁺⁺) can be estimated from the values of the C₁-C₃₃ bond length (see atom numbers on Fig. S2), as well from its Mayer bond order [38]. Fig. 7 and 8 display the results obtained for the three series of dimers according to the Brown constant of the R group (R=H, NMe₂ and NO₂). The C₁-C₃₃ bond length is smaller for 2b⁺⁺ (Fig. 7, black square) than for 2b⁺ (Fig. 7, red disk), whereas the opposite result is obtained for (2a)⁺⁺⁺ and (2f)⁺⁺⁺ dimers. In addition, the Mayer bond order is higher for 2b⁺⁺ than for 2b⁺, conversely to the two others series (Fig. 8). These results strongly suggest that the doubly-charged dimer 2b⁺⁺ is relatively more stable than the monocharged one 2b⁺, consistent with a RRD process. In contrast, the shorter C₁-C₃₃ bond and the larger Mayer order in the monocationic dimer than in the dicationic one for R=H and R=NO₂ indicates that the RSD process is likely favored in those cases, in agreement with experimental data.

**Fig. 7**
Fig. 7. Plots of calculated C-C bond length vs. Brown contant ($\sigma^+$) of the R group for doubly-charged $2^{2+}$ (black squares) and mono-charged $2^+$ dimers ($R = \text{NMe}_2, H, \text{NO}_2$).

Fig. 8. Plots of Mayer C-C bond order vs. Brown contant ($\sigma^+$) of the R group for doubly-charged $2^{2+}$ (black squares) and mono-charged $2^+$ dimers ($R = \text{NMe}_2, H, \text{NO}_2$).

Table 3. Free energy change (kcal.mol$^{-1}$) in the gas phase and in solvent (i.e. CH$_2$Cl$_2$) for homolytic and heterolytic C–C bond cleavage in $2^{2+}$ and $2^+$ dimers.

<table>
<thead>
<tr>
<th>R</th>
<th>Gas</th>
<th>Solvent</th>
<th>Gas</th>
<th>Solvent</th>
</tr>
</thead>
<tbody>
<tr>
<td>NMe$_2$</td>
<td>-63.2</td>
<td>-36.8</td>
<td>39.9</td>
<td>37.6</td>
</tr>
<tr>
<td>H</td>
<td>-45.6</td>
<td>-18.0</td>
<td>49.8</td>
<td>48.8</td>
</tr>
<tr>
<td>NO$_2$</td>
<td>-120.3</td>
<td>-125.9</td>
<td>93.6</td>
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</tr>
</tbody>
</table>

The free energy changes associated with C–C bond cleavage for $2^{2+}$ and $2^+$ dimers are listed in Table 3. Homolytic cleavage produces two cationic monomers from a doubly charged dimer, i.e. $2^{2+} \rightarrow 1^+ + 1^+$, while heterolytic cleavage produces one cationic and one neutral monomer from a singly charged dimer, i.e. $2^+ \rightarrow 1^+ + 1$. Homolytic bond cleavages are all exergonic whereas heterolytic cleavages are all endergonic, indicating that a symmetric bond dissociation is always thermodynamically favored. On the other hand, the free energy change associated with the heterolytic bond cleavage increases in the order $\text{NMe}_2 < H < \text{NO}_2$, in both
the gas phase and solvent. Although this result does not provide any information on the kinetics of dimerization, it suggests that RS-coupling is less thermodynamically disfavored for R = NO$_2$ than for R = NMe$_2$.

4. Conclusions

The voltammetric studies of the R-substituted phenylmethylenepyrans 1a-f have shown that the irreversible oxidation potential leads to the formation of bis-pyrylium $\sigma$-dimers (2a-f)$^{2+}$. Although the oxidation potential is fully dependent on the electrophilic property of the substituent, the potential of reduction of (2a-f)$^{2+}$ cleaving the C-C bond and yielding back 1a-f, does not vary significantly with R. Our voltammetric analyses also demonstrate that the intermolecular dimerization follows either a radical-radical or radical-substrate pathway according to the nature of the substituting moiety. Hence, dimerization occurs merely through a radical-radical process with electrodonating groups such as NMe$_2$ and OMe, whereas RSD is observed for the other phenylmethylenepyran. Electronic structure calculations suggest that this trend results from the inversion of the relative stabilities of the mono- and doubly-charged dimers with R. The kinetics of the RRD reaction have also been investigated by low-temperature electrochemical voltammetry (175 K), and room-temperature high scan rate CV. The electrochemical analysis, supported by voltammetric simulations, suggest a rather fast process with $k_{\text{dim}} = 10^7$ M$^{-1}$s$^{-1}$. From these results, future works will aim at developing new PhMPs compounds in order to slow down the dimerization coupling, by incorporation of bulky groups on the pyran moiety. Under such conditions, experimental and theoretical approaches will be carried out to further elucidate the mechanistic pathways. These results will be of interest for optimizing further development of redox chromophores with fast switching time, on the basis of our previous studies [26].
Acknowledgments

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Appendix A. Supplementary data

Supplementary data to this article can be found online.

References


Scheme 1. C-C bond making/breaking by oxidation of R-substituted phenylmethylenepyrans 1a-f and reduction of their bis-pyrylium derivatives (2a-f)\(^{2+}\).
Fig. 1. CV ($v = 0.1 \text{ V.s}^{-1}$) at a Pt electrode (diam. 3mm) of compound 1a ($C = 1\text{mM}$) in CH$_2$Cl$_2$/NBu$_4$PF$_6$ 0.1 M under argon at $T = 293 \text{K}$. 
Fig. 2. Plots of $E_{pa}$ vs. log $v$ for compounds A) 1a (R=H); B) 1b (R=NM$_2$); C) 1c (R=OMe); D) 1d (R=CO$_2$Me); E) 1e (R=Br); F) 1f (R = NO$_2$); (C = 0.5 mM).
Fig. 3. Plots of slope of ($E_{pa}$ vs. log $v$) against log $C$ for 1a-f.
Fig. 4. Plots of $E_{pa}$ vs log $C$ for compounds A) 1a (R=H); B) 1b (R=NMe$_2$); C) 1c (R=OMe); D) 1d (R=CO$_2$Me); E) 1e (R=Br); F) 1f (R = NO$_2$); ($v = 0.1$ V.s$^{-1}$).
Scheme 2. Schematic pathway for the C-C bond making/breaking of R-substituted phenylmethyleneypyran 1a-f compounds upon electron transfer.
Fig. 5. CVs ($v = 0.1 \text{ V.s}^{-1}$) at a Pt electrode (diam. 1mm) of compound 1d (0.04 mM) in CH$_2$Cl$_2$/NBu$_4$PF$_6$ 0.1 M under argon for 175 K $< T < 183$ K.
Fig. 6. Experimental (black) and simulated (red) CVs at A) $\nu = 100 \text{V.s}^{-1}$ and B) $\nu = 1 \text{V.s}^{-1}$ of compound 1b ($C = 0.09 \text{mM}$) at a Pt electrode (diam. 1mm) in CH$_2$Cl$_2$/NBu$_4$PF$_6$ 0.1 M. (for details, see Appendix A).
**Fig. 7.** Plots of calculated C-C bond length vs. Brown constant ($\sigma^+$) of the R group for doubly-charged $2^{2+}$ (black squares) and mono-charged $2^+$ dimers (R = NMe$_2$, H, NO$_2$).
Fig. 8. Plots of Mayer C-C bond order vs. Brown content (σ⁺) of the R group for doubly-charged 2⁺ (black squares) and mono-charged 2⁺ dimers (R = NMe₂, H, NO₂).