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Stable dicationic dioxoliums and fate of their dioxolyl radicals†

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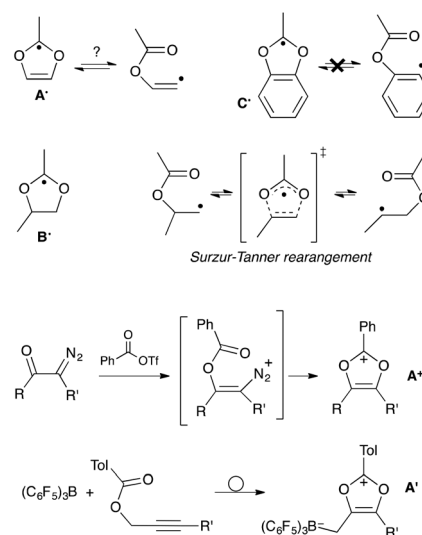
Stable dicationic dioxolium salts featuring an ancillary vinamidinium pattern were synthesized and characterized. Although highly reactive, they were found otherwise easy to handle under inert atmosphere. This offered the opportunity to generate and study the fate of unknown 1,3-dioxolyl radicals. Depending on substituents, reduction led to the formation of dimers of either dioxolyl or cyclohexadienyl radicals, stemming from a process that is related to the Surzur–Tanner rearrangement. The cyclohexadienyl radical could be characterized in the case of a tri(*tert*-butyl)phenyl group, which prevents dimerization processes.

Introduction

The design of stabilized C-centered reactive molecules, such as carbeniums, carbanions or radicals, through the introduction of hetero-substituents is a well-proven concept.^{1–4} Indeed, the diversity of main group elements provides a wide palette of electronic effects, from prototypical π -accepting B- or Al-based functions, up to strong electron-donating amino groups.⁵ Oxygen stands out as the most electronegative element of the periodic table (apart from fluorine),^{5a,b} while at the same time RO-alkoxy groups are almost equivalent to amino substituents in terms of π -donation.⁵ Strikingly, this combination isn't synergistic and the strong, but opposite, effects often counter-balance each other. In other words, the *O*-substituents fail to decrease the basicity of electron-rich carbanions because of their +M donation,³ whereas at the same time the electronegativity of oxygen prevent them for taming the reactivity of electron deficient carbenium centers.^{1,6} Thus, simple aryl- and alkyl-oxoniums are highly reactive and their study have been essentially confined to super acidic media.^{2,7,8} This is in marked contrast with their nitrogen counterparts: countless bench-stable iminiums and amidiniums have been reported.⁹ Similarly, the introduction of alkoxy groups have little beneficial effects on the stability of C-centered radicals. Even their combination with an electron-withdrawing group, so-called capto-dative substitution,⁴ results in modest enhancements, especially when compared to amino groups. As a matter of

fact, to date, all isolated C-radicals with a simple capto-dative substitution pattern feature *N*-substituents as donors.^{10,11}

In this article, we consider the case of 1,3-dioxolyl scaffolds **A** (Scheme 1). Radicals **A**[•] have never been evidenced experimentally,¹² although computational studies already assessed their possible role as reactive intermediates, especially in the rearrangement of β -(acyloxy)vinyl radicals.^{13–15} Note that known¹⁶ parented 1,3-dioxolanyl radicals **B**[•] should be similarly the intermediates in the shift of β -(acyloxy)alkyl radicals, so-called Surzur–Tanner rearrangement.¹⁷ However, extensive experimental and theoretical works have demonstrated that this reaction proceeds in fact through a closely-related transition state, which lies lower in energy (see Scheme 1).^{17c,18}



Scheme 1 Dioxolyl, dioxolanyl and benzodioxolyl radicals **A**[•]–**C**[•]; their relation with “Surzur–Tanner”-type rearrangements; synthesis of dioxoliums salts **A**⁺ and **A'**⁺.

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Even more, the ester-shift doesn't occur in the case of *ortho*-(acyloxy)aryl radicals, neither through 1,3-benzodioxolyl radicals C' nor through another pathway, because of the inability of the aryl radical to achieve suitable orbital overlap.¹⁹

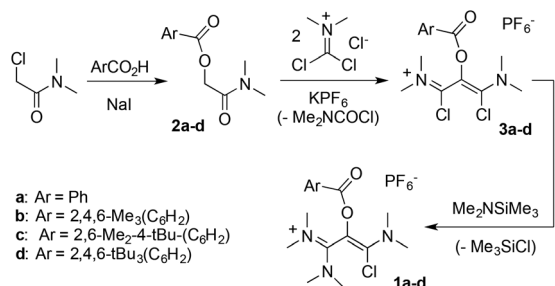
The absence of viable synthetic route towards the generation of A' certainly explains why this class of radicals still constitutes a *terra incognita*. In principle, they could be unambiguously produced through one-electron reduction of the corresponding dioxolium salts A⁺. However, these carbeniums are highly reactive themselves,^{20,21} apart from derivatives benefiting from further stabilization, such as a benzo-fused ring²² or an additional hetero-substituent.²³ Even in this latter case, attempts to assess the fate of the corresponding dioxolyl-like radicals were inconclusive.²⁴ To date, only neutral zwitterionic borate-based derivatives A', which were recently prepared by Stephan *et al.* from prop-2-yn-1-yl benzoates, have been structurally characterized.^{21b}

Herein we report the synthesis of isolable dicationic dioxolium salts, featuring a vinamidinium pattern. Though reactive, they are easily handle-able under dry inert atmosphere. This allowed not only for full spectroscopic and structural characterizations, but also stepwise reactivity studies, especially the generation and fate of the corresponding cationic 1,3-dioxolyl radicals.

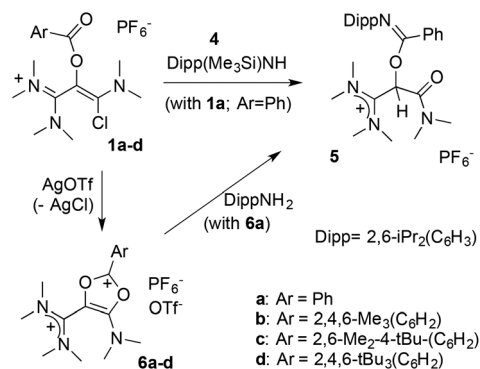
Results and discussion

This study was initially motivated by our interest in the reactivity of salt **1a** (Scheme 2), in the context of our work on unusual vinamidinium scaffolds.²⁵ The three-step synthesis of this novel synthon was straightforward. We first performed an iodide-catalyzed esterification of benzoic acid with *N,N*-(dimethyl)chloroacetamide. The resulting 2-(benzoyloxy) acetamide **2a** was reacted with dichloromethylene-dimethyl iminium chloride²⁶ to yield, after anion metathesis, 1,3-(dichloro)vinamidinium hexafluorophosphate salt **3a**. Finally, the addition of one equivalent of dimethyl(trimethylsilyl) amine allowed for a clean selective mono-substitution, affording **1a** in 78% yield.

Originally, we then wished to introduce a bulky anilino group by direct nucleophilic acyl substitution of the remaining chloride. A clean and complete reaction of **1a** with



Scheme 2 Synthesis of 1-(chloro)vinamidinium salts **1a–d**.



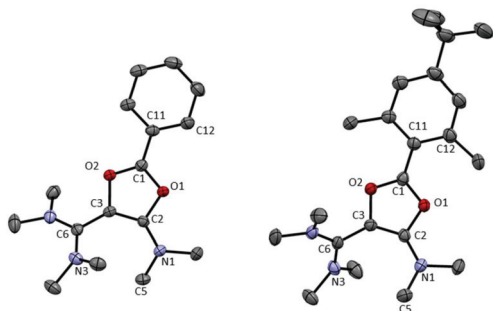
Scheme 3 Reaction of **1a** with trimethylsilylanilines; synthesis of dioxolium salts **6a–d**.

N-(trimethylsilyl)-2,6-di(isopropyl)aniline **4** was observed after 3 days at 60 °C in acetonitrile (Scheme 3). However, whereas HR-MS analysis was consistent with the expected substitution product, the presence of a C–H signal in ¹³C NMR at δ = 72.3 ppm indicated the formation of a different structural isomer. A single crystal X-ray diffraction study finally allowed for the identification of benzimidate **5** (see ESI†).²⁷

The outcome of this reaction suggested the transient formation of dicationic 1,3-dioxolium **6a**, followed by fast aminolysis.^{20,21a} In order to probe this hypothesis, a solution of **1a** was stirred in presence of silver triflate. A bright yellow precipitate appeared, while NMR monitoring of the supernatant showed the gradual disappearance of the starting material with complete conversion after 2 days. The product was purified by crystallization and isolated in 56% yield. An X-ray diffraction analysis confirmed the formation of the dicationic dioxolium **6a** (see further below).

We reacted **6a** with 2,6-di(isopropyl)aniline and observed the formation of **5**, as initially suggested by the reactivity of **1a**. As expected, **6a** is also very sensitive to moisture and solubilization in tetrahydrofuran immediately triggers the oligomerization of the solvent. However, we found it otherwise easy to handle under inert atmosphere, which encouraged us to synthesize a set of parent dicationic dioxoliums **6b–d**, featuring methyl or *tert*-butyl groups in *ortho* and *para* positions of the aryl moieties. Starting from the corresponding benzoic acids, we synthesized 2-(aryloxy)acetamide **2b–d**. Their reaction with dichloromethylene-dimethyliminium chloride first performed very poorly, certainly due to the use of bulkier aryl substituents. Among several modifications to Viehe's original protocol,²⁶ the use of acetonitrile as a solvent was found critical to finally isolate 1,3-di(chloro)vinamidinium salts **3b–d** in 44–62% yields. Next, treatment with dimethyl(trimethylsilyl) amine yielded 1-chlorovinamidinium salts **1b–d**. As frequently observed for vinamidinium cations, **1a–d** consist in mixtures of interconverting *E* and *Z* isomers, which can't be separated. Moreover, the conformers have distinct or fluxional ¹³C and ¹H NMR signals at room temperature, resulting in equivocal spectra. We confirmed further their structure by X-ray diffraction analysis. Note that, in the solid state, **1a–c** adopt a *Z* con-

Of note, although spectroscopic data for **6a-d** are overall analogous, alkyl substituents in *ortho* and *para* positions of the aryl group results in shielding the *ipso* carbon atom (^{13}C NMR chemical shift for **6a**: $\delta = 117.3$ ppm, for **6b-c**: $\delta = 113.1$ ppm). This is especially the case for **6d** ($\delta = 108.3$ ppm). In addition, **6a-c** have almost identical ^{13}C NMR chemical shifts for the carbenium center C1 ($\delta = 171.0\text{--}171.9$ ppm), whereas the signal in **6d** is deshielded by 7 ppm. We inter-



Next, we examined the fate of these dicationic species upon reduction. Cyclovoltammograms of all dioxoliums **6a-d** feature an irreversible reduction wave at about -1.1 V. As a two-electron reduction would lead to an unreasonable “acetal anion” equivalent, we hypothesized an initial one-electron transfer. Note that no reversibility could be evidenced even at high scan rates (up to 10 V s^{-1}), thus indicating that the resulting radicals **7a-d** must undergo a fast chemical transformation.

In order to get further insight, we performed the chemical reductions of dioxolium salts **6a–d**. We first reacted **6a** with half an equivalent of zinc powder. The monitoring of the reaction by ^1H NMR showed the gradual disappearance of **6a** and the appearance of a new set of signals, while the reaction mixture remained EPR silent. At full conversion, the crude mixture consists in closely related isomers. Fractional crystallization allowed for the isolation of two diastereomeric forms of the dimer of dioxolyl radical **7a**, the bis(dioxole)s *d,l*-**8a** and *meso*-**8a**, in 53% and 7% yield respectively (Scheme 3). They were fully characterized and their structure was ascertained by X-ray analysis (Fig. 2). In the solid state, *meso*-**8a** displays a *anti* conformation of the two dioxolyl groups whereas both *anti* and *gauche* conformers are observed in the case of *d,l*-**8a**.

Importantly, we couldn't evidence any reversible dissociation of the dimers. Neither *d,l*-**8a** nor *meso*-**8a** afford equilibrated mixtures of diastereomers when heated in acetonitrile for several hours at 80 °C. This indicates that Gibbs free activation energies for a C–C bond breaking in dimers **8a** are at least 30 kcal mol $^{-1}$. Note that DFT calculations³⁰ at the uB3LYP/6-311 g(d,p) level of theory failed to predict such a strong bond. For instance, the dimerization of **7a** to afford the *anti* conformer of *d,l*-**8a** is predicted to be endergonic by $\Delta G = +12.1$ kcal mol $^{-1}$ and exergonic by only -24 kcal mol $^{-1}$ when introducing the Polarizing Continuum Model (PCM) for acetonitrile. As dispersion forces can play a critical role in the stability of such encumbered dimers,³¹ we considered the long-range corrected functional ω B97XD,³² which implements a version of Grimme's D2 model for dispersion forces.³³ As a matter of fact, the ω B97XD/6-311 g(d,p)/PCM level of theory predicts a more exergonic dimerization ($\Delta G = -47$ kcal mol $^{-1}$; -11 kcal mol $^{-1}$ without PCM), but also affords optimized geometries that better fit experimental solid-state structures, including the length of the C1–C1' bond that is formed upon dimerization of **7a** (for the *anti* conformer of *d,l*-**8a**, B3LYP: 1.569 Å, ω B97XD: 1.558 Å, X-ray: 1.552(4) Å). For consistency throughout this work, all geometry optimizations were carried out at this level of theory. Note that the *anti* conformer of *d,l*-**8a** was found more stable than the *gauche* conformer and than the *meso* diastereomer as well, but by only few kcal mol $^{-1}$, in line with the experimental observation of the three forms.

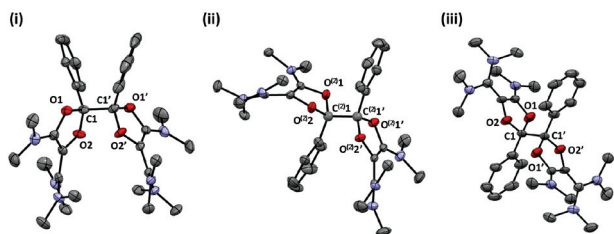


Fig. 2 Representation of the X-ray structure of *d,l*-**8a** (i: *gauche* conformer; ii: *anti* conformer) and *meso*-**8a** (iii) with 50% probability ellipsoids. Counter-anions, hydrogen atoms and solvent are omitted for clarity.

We wondered whether replacing the phenyl group of **7a** with a bulkier 2,4,6-trimethylphenyl could prevent the dimerization of the corresponding dioxolyl radical **7b** into the bis(dioxole) **8b**. Therefore, we performed the reduction of dication **6b** with 0.5 equivalent of zinc. The ^1H NMR analysis of the crude mixture revealed the selective formation of a new compound **9b** that could be isolated as yellow crystals. The ^1H NMR spectrum of **9b** features an olefinic resonance signal at 5.96 ppm suggesting the dearomatization of the aryl group and the formation of a cyclohexadiene moiety. This assumption was confirmed by a single crystal X-ray diffraction analysis. Dication **9b** is a symmetrical dimer of 2-oxaspiro[4.5]deca-3,6,9-triene-1-one units, connected at their C8 position (Fig. 3). Its formation suggests that dioxolyl radical **7b** undergoes a ring opening through C–O bond cleavage to generate vinyl radical **10b** (Scheme 4). This latter undergoes a spiro-cyclization to afford cyclohexadienyl radical **11b**, which is apparently persistent enough to build-up in solution and dimerizes.

In the case of the reduction of **6c**, which features an even more bulky 2,6-dimethyl-4-*tert*butylphenyl group, the crude mixture yielded very complex NMR spectra, suggesting that the corresponding radical **7c** evolved following multiple pathways. Nevertheless, a small amount of pure material could be isolated as a crystalline solid, which was attributed to dimer **8c**. Indeed, although no suitable crystals for X-ray diffraction could be obtained, NMR data are clearly reminiscent of bis(dioxole) **8a**, including signals for untouched aryl substituents and a peak in $^{13}\text{C}\{^1\text{H}\}$ NMR at 115.9 ppm for C1 and C1' carbons, which are linking the two monomeric units (**8a**: 111.2 ppm). This attribution is also supported by HR-MS analysis, which is consistent with a dicationic dimer.

DFT calculations indicate that radicals **7a–c** have almost identical electronic structures. Most of the Mulliken spin density is centered on the C1 carbon atom (51–53%, see Table 2 and Fig. 4a), the rest being spread over the aryl group (30–33%), and the two oxygen atoms (12–13%). Similarly to the corresponding dioxolium carbeniums, the π -systems of the dioxolyl and the vinamidinium moieties poorly interact, the latter bearing less than 5% spin density. Dioxolyl **7a** (aryl = phenyl), vinyl **10a** and cyclohexadienyl radicals **11a** were found very close in energy and it is likely that the three forms co-exist in solution. In the case of bulkier aryl substituents, the balance is more shifted towards the cyclohexadienyl forms **11b–c**. Note that the isolated products **8a,c** and **9b** do not correspond to the dimer of the most stable radical form but to

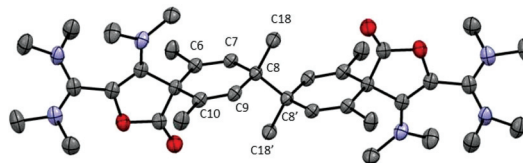
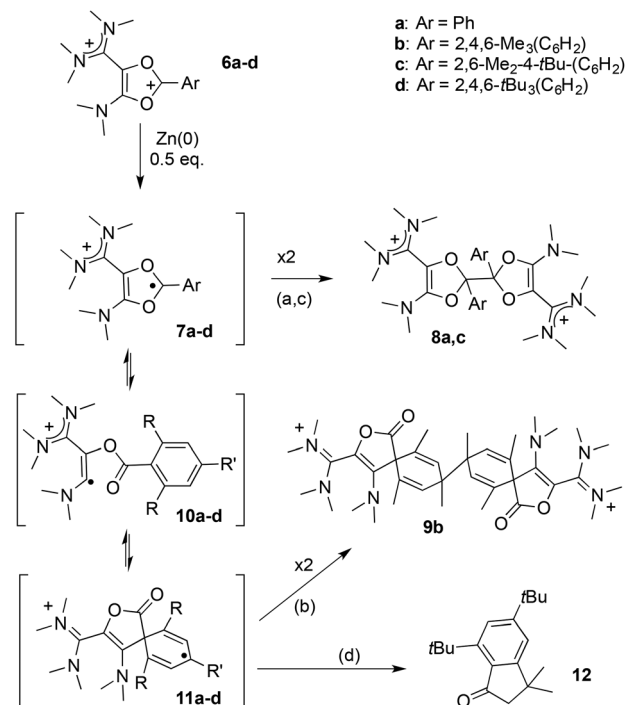


Fig. 3 Representation of the X-ray structure of **9b** with 50% probability ellipsoids. Counter-anions, hydrogen atoms and solvent are omitted for clarity.



Scheme 4 Chemical reduction of 6a–d.

Table 2 Gibbs free enthalpy of radical isomers 10 and 11 (relative to 7), and for the formation of dimers 8 and 9

Dioxolyl radical	7a	7b	7c	7d
R	H	Me	Me	<i>t</i> Bu
R'	H	Me	<i>t</i> Bu	<i>t</i> Bu
Mulliken spin density				
C1	51%	51%	53%	80%
Aryl	32%	33%	30%	1%
O1 and O2	13%	12%	12%	15%
ΔG (relative to 7a–d)^a				
10a–d	+1.0	–1.6	–0.8	–0.8
11a–d	–4.7	–16.7	–15.7	–12.2
$[G(8a-d)-G(9a-d)]^a$				
7a–d (×2) → 8a–d	–11.4	+10.2	–32.0	n.a. ^b
7a–d (×2) → 8a–d	–51.2	–68.5	–69.8	n.a. ^b
11a–d (×2) → 9a–d	–30.4	–45.3	–6.4	+2.7

^a Energies are in kcal mol^{–1}. ^b No minimum found for 8d.

the overall most stable dimer, thus suggesting that dimerization is essentially under thermodynamic control. Interestingly, the introduction of methyl *ortho*-substituents is not detrimental to the formation of dimers 8a–c from radicals 7a–c, the dimerization being even more exergonic for 7b,c (ΔG about –70 kcal mol^{–1}) than for 7a (ΔG = –51 kcal mol^{–1}). Similarly, the formation of dimer 9b from cyclohexadienyl radical 11b is predicted to be more exergonic than the formation of 9a from 11a. However, the *para tert*-butyl group in 11c is clearly prejudicial to the formation of the corresponding dimer 9c (ΔG =

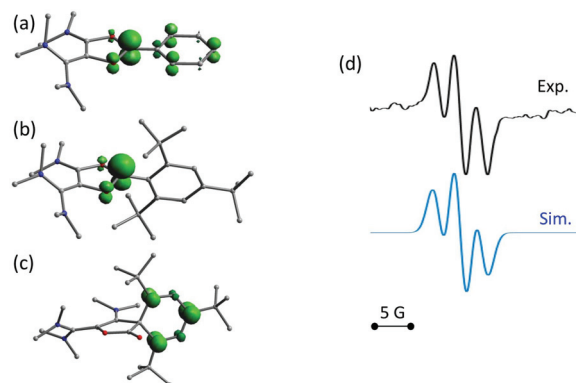


Fig. 4 Representation of computed Mulliken spin densities for, (a) 6a, (b) 6d and (c) 11d; (d) experimental X-band isotropic EPR spectrum at room temperature in acetonitrile after reduction of 6d (top, in black), the simulated spectrum (bottom, in blue) was obtained with a Lorentzian line-broadening parameter of 0.25 and an hyperfine constant $a(^1\text{H})$ of 8.0 MHz (2 nuclei).

–6 kcal mol^{–1} only). In line with this trend, radicals 7d and 11d, which stem from dioxolium 6d with a 2,4,6-tri(*tert*-butyl) phenyl substituent, should not dimerize. Indeed, the formation of dimer 9d from 11d was found slightly endergonic and no minimum on the hypersurface of energy could be found for the putative dimer 8d. Note that 7d is also predicted to be fundamentally different from 7a–c. The aryl group is not conjugated with the rest of the π -system anymore and does not feature significant spin density, 80% of it being now localized on the carbenium center C1 (Table 2 and Fig. 4b). The latter has no planar environment anymore (sum of bond angles around C1 in 7d: 350.6°, in 7a–c: >359.9°), indicating an increased sp³ character. This illustrates the key role of the aryl group in the stabilization of dioxolyl radicals through spin delocalization. In addition, not only is 7d poorly stabilized, but the cyclohexadienyl form is only –12 kcal mol^{–1} lower in energy and is likely to be highly reactive as well.

In order to experimentally assess the fate of these radicals, we examined the reduction of dioxolium 6d with half an equivalent of zinc(0) powder. EPR monitoring of the reaction showed the appearance of a 1 : 2 : 1 triplet, corresponding to an hyperfine coupling constant a = 8 MHz with two equivalent $\frac{1}{2}$ spins³⁴ (Fig. 4d). This value is in perfect agreement with the expected spectrum for 11d. In particular, DFT calculations predict that this organic π -radical should only feature significant isotropic hyperfine coupling constants with the two hydrogen atoms in *meta* position of the aryl group, with a computed value $a(^1\text{H})$ of about 9 MHz. Though persistent at room temperature, the EPR signal evolved after few hours into a more complex unsymmetrical bandshape, indicating a mixture of radical species, and finally faded away (see ESI†). Unsurprisingly, all attempts to isolate 11d failed. Ultimately, known indanone 12³⁵ was isolated from the crude mixture in 30% yield (Scheme 4).

Note that, at a pinch, the formation of dimers 8a,c could have been the result of an ionic mechanism. However, the formation of 9b, the observation of radical 11d, as well as DFT

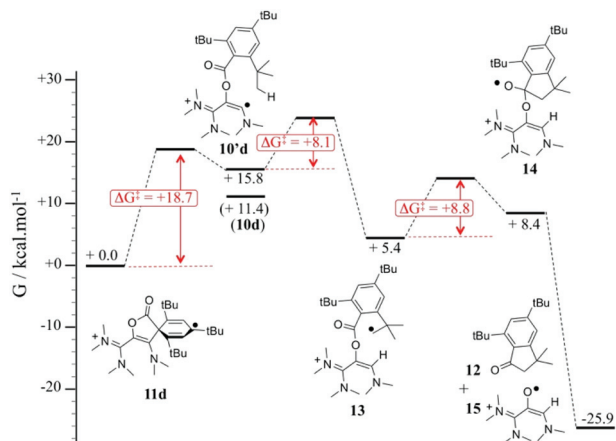


Fig. 5 Computed mechanism for the formation of indenone **12** from **11d**, **b** at the; ω B97XD/6-311 g(d,p)/PCM(acetonitrile) level of theory relative Gibbs energies of intermediates and Gibbs energies of activation.

results as a whole, definitely support radical pathways, stemming from a one-electron reduction of the dioxoliums.

We propose a mechanism for the decay of **11d** into **12**, which is supported by DFT investigations (Fig. 5). The rate-determining step of the process is the ring opening of **11d**, yielding vinyl radical **10d'** ($\Delta G^\ddagger = +18.7$ kcal mol⁻¹). The latter is only +4.4 kcal mol⁻¹ higher in energy than the most stable conformer **10d**. It features the relevant conformation for a hydrogen shift from an *ortho tert*-butyl group ($\Delta G^\ddagger = +8.1$ kcal mol⁻¹). The resulting alkyl radical **13** adds intramolecularly to the carbonyl group ($\Delta G^\ddagger = +8.8$ kcal mol⁻¹), yielding five-membered ring **14**, which undergoes a barrier-less homolytic C–O bond breaking to afford indenone **12** and radical **15**. Note that we previously showed that so-called oxyallyl radical cations, which are parented to **15**, are remarkably stabilized and could even be highly air-persistent.^{10a,c,25a,d,e} Although it appears that **15** evolve further in the reaction conditions, it is likely that it contributes to the transiently observed EPR spectrum upon decay of **11d**. However, we couldn't obtain an unambiguous simulation for the EPR bandshapes by only considering a mixture of the cyclohexadienyl and oxyallyl radicals. This suggested the formation of a more complex mixture of paramagnetic species.

Conclusion

Dicationic dioxoliums **6a–d** were readily synthesized from acetamides **2a–d**.^{36–39} All spectroscopic and structural data indicate a poorly aromatic five-membered ring and a modest interaction between the π -systems of the vinamidinium and carbenium moieties. This situation is reminiscent of that of the neutral borate-based zwitterions from Stephan *et al.*,^{21b} which were the only non-stabilized dioxoliums to have been previously structurally characterized by X-ray analysis.

The fate of **6a–d** upon reduction highly depends on the aryl substituent. All dioxolyl radicals **7a–d** can undergo a ring-

opening, which is reminiscent of the Surzur–Tanner rearrangement of β -(acyloxy)alkyl radicals. They are in equilibrium with the resulting vinyl radical forms **10a–d** and their spiro-cyclization products, the cyclohexadienyl radicals **11a–d**. In the case of a bulky tri(*tert*-butyl)phenyl aryl group (**7d**), dimerization processes are disfavored. The radical essentially exists in the cyclohexadienyl form **11d**, which was observed at room temperature by EPR spectroscopy.

DFT studies show that spin-delocalization on the aryl group plays a major role in the stabilization of dioxolyl radicals, with only second-order effects of the *O*-substituents. As a matter of fact, dioxolyl radicals **7a–c** could be depicted as well as benzyl radicals with ancillary *O*-substituents: more than 30% of spin density is found on the fully conjugated aryl groups, only 12–13% being spread on the two oxygen atoms. In **7d**, the bulky *tert*-butyl *ortho*-substituents twist away the π -systems and prevent this delocalization. The radical is highly localized, with more than 80% of spin density on one carbon atom, which features some sp^3 hybridization.

Finally, although DFT ω B97XD/6-311 g(d,p)/PCM calculations could fairly account for the experimental data and observations of this work, it is too early to fully ascertain their accuracy. We are actually considering experimental gas phase studies with a modified QhQ mass spectrometer to better calibrate DFT level of theory and further explore the uncharted territory of the reactivity of these species.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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