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Cite this: DOI: 10.1039/c5ob01651g Received 6th August 2015, Accepted 21st August 2015 DOI: 10.1039/c5ob01651g Dithiafulvenyl-substituted phenylacetylene derivatives: synthesis and structure-propertyreactivity relationships†

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A series of regioisomers for dithiafulvenyl-substituted phenylacetylene derivatives was synthesized and characterized to show structure-dependent electronic properties and different reactivities in their oxidized states.

The application of π -conjugated materials has achieved tremendous advancement in the cutting-edge fields such as molecular electronics, chemo-/bio-sensors, light-emitting devices, and organic photovoltaics.¹⁻³ From the chemistry perspective, continued efforts to explore π -molecular building blocks with novel properties and functions are still imperative and highly needed. For decades, tetrathiafulvalene (TTF)-the first organic conductor-has played an indispensable and central role in the area of molecular electronics.⁴ In modern TTF chemistry, a myriad of π -extended TTF analogues (*i.e.*, exTTF) have been synthesized and many of them were found to show intriguing redox, optoelectronic, and supramolecular properties.⁵⁻⁸ Among numerous types of exTTF, the so-called tetrathiafulvalene vinylogues (TTFVs)9,10 have been actively pursued by us¹¹ and others¹²⁻¹⁴ over the past few years. A particularly appealing aspect of TTFVs, especially aryl-substituted TTFVs, is its redox-controllable conformational switching behaviour.^{11,14,15} Built upon aryl-TTFVs, various functional molecular systems have been prepared and reported recently, including stimuli-responsive conjugated polymers,^{16,17} fluorescent molecular tweezers,^{18,19} redox-active ligands,¹² shapepersistent macrocycles,^{20,21} and molecular rotors.²² Of note is our previous study on certain alkynylated TTFV systems derived from a phenylacetylene-substituted dithiafulvene (DTF) precursor 1a (Fig. 1).^{20,21} Herein the inclusion of the alkynyl group in this DTF compound greatly enhances its synthetic versatility by enabling diverse efficient C-C bond forming reactions to take place on both the DTF unit and the





alkynyl group.^{18–21,23} Our previous study on **1a** naturally led us to ponder on its two other regioisomers, **1b** and **1c** (Fig. 1). As building blocks, acetylenic DTF isomers **1a–c** would give rise to π -conjugated materials with different structures and properties as a result of their different molecular shapes and π -conjugation patterns. In this context, it is of fundamental importance to conduct comparative studies on the properties and reactivities of **1a–c** as well as on the π -extended systems derived from them.

The synthesis of DTFs 1a-c was executed through a phosphite-promoted olefination method,^{24,25} using trimethylsilvlethynyl-substituted benzaldehydes 2a-c and thione 3 as the starting materials (Scheme 1). Experimentally, the three olefination reactions were completed in 3 hours at 105 °C, giving the desired products 1a-c with similar yields (ca. 75-79%). Compounds 1a-c were then subjected to an oxidative coupling²⁶ in CH₂Cl₂ at room temperature using iodine as an oxidant (Scheme 1). The reactions yielded acetylenic diphenyl-TTFVs 3a-c in very good yields. Note that in our previous work, certain ortho-substituents (Br and Me) retarded the deprotonation step in the oxidative dimerization mechanism, leading to the formation of a unique bis-spiro structure other than the typical TTFV product.^{27,28} This scenario however did not happen in the oxidative dimerization of ortho-alkynyl DTF 1c, because the alkynyl group is linear-shaped and much less bulky.

Besides the oxidative dimerization reactions, the acetylenic DTF synthons could also be synthetically elaborated to attain π -extension through alkynyl coupling reactions. Scheme 2 out-

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Scheme 1 Synthesis of acetylenic DTFs 1a-c and related TTFV derivatives 3a-c.



Scheme 2 Synthesis of bis(DTF)-endcapped butadiynes 4a-c via Hay coupling. (a) (i) K_2CO_3 , THF/MeOH, 30 min, (ii) Cul, TMEDA, air, CH₂Cl₂, rt, 5 h.

lines the Hay coupling reactions²⁹ of desilylated **1a–c**. The synthesis began with protiodesilylation with K_2CO_3 to first generate free terminal alkyne intermediates, which after a brief workup were immediately subjected to alkynyl homocoupling catalyzed by CuI/TMEDA, affording bis(DTF)-endcapped conjugated butadiynes **4a–c** in satisfactory yields. Diynes **4a** and **4b** are yellow coloured solids, whereas **4c** shows an intense orange color, suggesting that it has very different electronic absorption properties than the other two isomers (*vide infra*).

The electronic absorption properties of acetylenic DTFs and their related π -derivatives were studied by UV-Vis spectroscopy. As can been from Fig. 2, the three DTF isomers **1a–c** show $\pi \rightarrow \pi^*$ transition bands with similar molar absorptivities. The maximum absorption wavelengths (λ_{max}) shift in the order of **1a** (384 nm) > **1b** (376 nm) > **1c** (357 nm), giving a qualitative comparison of the degree of π -delocalisation among the three isomers. The observation of **1c** showing the lowest λ_{max} value



Fig. 2 UV-Vis absorption spectra of compounds 1a-c, 3a-c, and 4a-c measured in CH₂Cl₂ at room temperature.

in the three isomers can be reasonably correlated with its *ortho*-substitution structure which engenders the most significant steric interactions between the alkynyl group and the DTF moiety to attenuate the π -conjugation.

The $\pi \to \pi^*$ transition bands of TTFV isomers **3a–c** show the same trend of λ_{max} shift (*i.e.*, *para* > *meta* > *ortho*) as that in 1a-c, but the extent of variation is much greater (Fig. 2B). For *para*-isomer **3a** the λ_{max} value is observed at 380 nm, which is nearly identical to that of its DTF precursor 1a and indicates that the central TTFV moiety in 3a does not contribute to increasing π -delocalisation due to its twisted *cis*-like conformation.^{11,15,30} Compared with **3a**, the absorption peak of *meta*isomer 3b shows a significant blueshift to 350 nm, which is suggestive that the π -electron delocalisation in 3b is more disrupted than 3a. The absorption spectrum of ortho-isomer 3c differs pronouncedly from its other two isomers, wherein the $\lambda_{\rm max}$ value at 329 nm is considerably blueshifted. In addition, the spectrum of 3c features a broad absorption slope in the range of ca. 350 to 500 nm. Worth noting is that such a spectral profile bears great resemblance to that of a bis-(1-naphthyl)-TTFV compound³¹ which we had reported

previously. Given that the central TTFV moiety of the bis(1-naphthyl)-TTFV has been confirmed to assume a completely planar *trans* conformation by X-ray analysis,³¹ the molecular shape of **3c** is thus deemed as adopting a similar *trans* motif. Further evidence of this conformational argument can be elicited from cyclic voltammetric analysis (*vide infra*).

The UV-Vis spectral profiles of bis(DTF)-endcapped butadivnes 4a-c varied significantly from one to another (Fig. 2C). The spectrum of divne 4a shows a monotonous absorption peak centred at 423 nm, with absorptivity in the range of 270-350 nm being very low. The spectral envelope of 4b is substantially blueshifted relative to 4a, showing three well-resolved bands at 339, 316, and 295 nm and a shoulder peak at 361 nm, respectively. The spectrum of 4c gives two intense $\pi \rightarrow$ π^* bands at 398 and 332 nm. Overall, the λ_{max} values of the bis(DTF)-butadiyne isomers 4a-c follow an order of para > ortho > meta, which is different from those of DTFs 1a-c and TTFVs 3a-c. An empirical rationalization for such spectroscopic behaviour can be made based on the π -delocalisation degree argument; that is, the DTF-phenylene-diyne skeletons of compounds 4a and 4c are in linear π -conjugation, whereas that of compound 4b is cross-conjugated due to its meta-substitution pattern.

The redox properties of compounds **1a–c**, **3a–c**, and **4a–c** were characterized by cyclic voltammetry (CV), and the detailed voltammograms are shown in Fig. 3. When scanned in the positive direction, DTFs **1a–c** all show a similar anodic peak at *ca.* +0.8 V which can be assigned to the formation of a [DTF⁺¹] radical cation *via* a single-electron oxidation step.^{15,32} In the reverse scan, a cathodic peak at +0.48 V is seen in the voltammograms of *para-* and *meta-*isomers **1a** and **1b**. Compared with the voltammograms of TTFVs **3a** and **3b**, the origin of this current peak can be assigned to the two-electron reduction of the [TTFV²⁺] dication formed on the working electrode surface

through electrochemical oxidative dimerization. For *ortho*isomer **1c**, the reverse CV scan reveals two cathodic peaks at +0.59 and +0.33 V respectively in the positive potential window. These two peaks are ascribed to the stepwise singleelectron transfers occurring on the [TTFV²⁺] product, which is evidenced by the nearly identical reduction features observed in the voltammogram of *ortho*-alkynyl TTFV **3c**. In the negative potential window, all the voltammograms of **1a–c** show a noticeable cathodic peak around -0.47 V. The intensity of this peak shows dependence on scan rate; that is, at relatively slow rate this peak just diminishes. It is therefore reasoned that this particular peak is associated with some intermediary species formed electrochemically which would readily diffuse away from the working electrode surface.

The voltammograms of TTFVs **3a** and **3b** both show a similar reversible redox wave pair, which can be unambiguously assigned to the simultaneous two-electron redox processes on the diphenyl-substituted TTFV moiety which normally would take a twisted *cis*-like conformation.^{11,15,30} Significantly different are the patterns in the voltammogram of *ortho*-isomer **3c**, in which the redox processes of TTFV undergo two distinct single-electron steps. Such CV behaviour, according to the previous studies by Yamashita³⁰ and us,³¹ alludes to a scenario that the TTFV moiety in **3c** prefers a planar *trans* conformation. The *ortho*-alkynyl groups in **3c** are believed to play a key role in directing the central TTFV conformation.

Likewise, the CV analysis of bis(DTF)-endcapped butadiynes **4a–c** also discloses notable substitution effects on their redox properties. The voltammograms of *para-* and *meta-*isomers **4a** and **4b** show the typical DTF oxidation peak at *ca.* +0.8 V along with a steadily growing redox wave pair due to the formation of TTFV during multicycle CV scans. The CV results confirm that **4a** and **4b** can readily undergo electrochemical polymerization



Fig. 3 Cyclic voltammograms of compounds 1a-c, 3a-c, and 4a-c measured in CH_2Cl_2 at room temperature. Experimental conditions: supporting electrolyte: Bu_4NBF_4 (0.1 M), working electrode: glassy carbon, counter electrode: Pt wire, reference electrode: Ag/AgCl (3 M NaCl), scan rate: 200 mV s⁻¹.



Scheme 3 Formation of compound 5 via an intramolecular cycloaddition pathway.

through the oxidative DTF coupling.^{15,20,33,34} Interestingly, such reactivity appears to be disfavoured in the case of *ortho*isomer **4c**, for there is no observation of the characteristic TTFV redox wave pair in its voltammogram. Worth noting is that the CV profile of **4c** exhibits an irreversible pattern with a few weak cathodic current peaks discernible in the reverse scans (see the inset in Fig. 3), suggesting that the oxidation of **4c** is followed by some swift chemical transformations to form redox-inactive species.

To gain a deeper insight into the redox reactivity, bis(DTF)divnes 4a-c were subjected to chemical oxidation using iodine as an oxidant. In line with the electrochemical analysis, diynes 4a and 4b under oxidative conditions were efficiently converted into corresponding TTFV-diyne oligomers, which were validated by MALDI-TOF MS analysis (see ESI†). The same oxidative treatment of ortho-isomer 4c, to our great surprise, ended up with the formation of a major product 5 (Scheme 3) along with a number of intractable byproducts. MS analysis confirmed that the byproducts were not TTFV-diyne oligomeric products. Compound 5 is a red coloured solid and electron-accepting in nature. The cyclic voltammogram of 5 clearly shows a reversible redox wave pair in the negative potential region ($E_{pa} = -1.08$ V, $E_{pc} = -1.25$ V), without any obvious redox peaks in the positive window (see ESI[†]). The mechanism for the formation of 5 is a rather complex one and awaits further investigations to be fully disclosed. At this juncture, a two-fold intramolecular cycloaddition^{35,36} pathway is tentatively proposed (Scheme 3) to account for the formation of the fused aromatic moieties in 5. Density functional theory (DFT) calculations suggest that the two fused aromatic units in 5 are nearly perpendicular to one another (see Fig. 4) in orientation. The biaryl-type structure and the presence of two thioester groups in the scaffold of 5 point to a potential in ligand design and stereoselective catalysis.³⁷

In summary, a series of structurally isomeric acetylenic DTFs and their π -extended derivatives have been systematically synthesized and characterized. Our studies show that the different substitution patterns in these compounds exert important effects on their electronic and redox properties. Worth highlighting is that *ortho*-alkynyl substituted TTFV



Fig. 4 Optimized structure of compound **5** at the B3LYP/6-31G(d) level. (A) Front view, (B) side view.

derivatives and conjugated butadiyne systems behave in a dramatically different way in terms of electronic absorption and redox reactivity compared with their *para-* and *meta-*isomers. The structure–property–reactivity relationships established herein offer a useful guidance to the ongoing research of DTF and alkyne based conjugated materials.

References

- 1 Functional Organic Materials: Syntheses, Strategies and Applications, ed. T. J. J. Müller and U. H. F. Bunz, Wiley-VCH, Weinheim, Germany, 2006.
- 2 G. Inzelt, *Conducting Polymers: A New Era in Electrochemistry*, Springer, Berlin, 2nd edn, 2012.
- 3 A. Facchetti, Chem. Mater., 2011, 23, 733-758.
- 4 TTF Chemistry: Fundamentals and Applications of Tetrathiafulvalene, ed. J. Yamada and T. Sugimoto, Springer, Berlin, 2004.
- 5 D. Canevet, M. Salle, G. Zhang, D. Zhang and D. Zhu, *Chem. Commun.*, 2009, 2245–2269.
- 6 M. Bendikov, F. Wudl and D. F. Perepichka, *Chem. Rev.*, 2004, **104**, 4891–4946.
- 7 J. L. Segura and N. Martín, *Angew. Chem., Int. Ed.*, 2001, 40, 1372–1409.
- 8 M. R. Bryce, J. Mater. Chem., 1995, 5, 1481-1496.
- 9 A. J. Moore and M. R. Bryce, *Tetrahedron Lett.*, 1992, 33, 1373–1376.
- 10 L. Yu and D. Zhu, Chem. Commun., 1997, 787-788.
- 11 Y. Zhao, G. Chen, K. Mulla, I. Mahmud, S. Liang, P. Dongare, D. W. Thompson, L. N. Dawe and S. Bouzan, *Pure Appl. Chem.*, 2012, 84, 1005–1025.
- 12 E. Gontier, N. Bellec, P. Brignou, A. Gohier, M. Guerro, T. Roisnel and D. Lorcy, *Org. Lett.*, 2010, **12**, 2386–2389.
- 13 J. Massue, N. Bellec, M. Guerro, J. F. Bergamini, P. Hapiot and D. Lorcy, *J. Org. Chem.*, 2007, **72**, 4655–4662.
- 14 M. Guerro, R. Carlier, K. Boubekeur, D. Lorcy and P. Hapiot, J. Am. Chem. Soc., 2003, 125, 3159–3167.
- 15 N. Bellec, K. Boubekeur, R. Carlier, P. Hapiot, D. Lorcy and A. Tallec, *J. Phys. Chem. A*, 2000, **104**, 10994–10994.
- 16 S. Liang, Y. Zhao and A. Adronov, J. Am. Chem. Soc., 2014, 136, 970–977.
- 17 S. Liang, G. Chen, J. Peddle and Y. Zhao, *Chem. Commun.*, 2012, **48**, 3100–3102.

- 18 K. Mulla, H. Shaik, D. W. Thompson and Y. Zhao, Org. Lett., 2013, 15, 4532–4535.
- 19 K. Mulla, P. Dongare, D. W. Thompson and Y. M. Zhao, *Org. Biomol. Chem.*, 2012, **10**, 2542–2544.
- 20 G. Chen, I. Mahmud, L. N. Dawe, L. M. Daniels and Y. Zhao, *J. Org. Chem.*, 2011, **76**, 2701–2715.
- 21 G. Chen, I. Mahmud, L. N. Dawe and Y. Zhao, *Org. Lett.*, 2012, **12**, 704–707.
- 22 G. Chen and Y. Zhao, Org. Lett., 2014, 16, 668-671.
- 23 K. Mulla and Y. Zhao, *Tetrahedron Lett.*, 2014, 55, 382–386.
- 24 S. S. Schou, C. R. Parker, K. Lincke, K. Jennum, J. Vibenholt, A. Kadziola and M. B. Nielsen, *Synlett*, 2013, 231–235.
- 25 C. A. Christensen, A. S. Batsanov and M. R. Bryce, *J. Org. Chem.*, 2007, **27**, 1301–1308.
- 26 P. Hapiot, D. Lorcy, A. Tallec, R. Carlier and A. Robert, J. Phys. Chem., 1996, 100, 14823–14827.
- 27 S. Bouzan, L. N. Dawe and Y. Zhao, *Tetrahedron Lett.*, 2013, 54, 4666–4669.

- 28 K. Woolridge, L. C. Goncalves, S. Bouzan, G. Chen and Y. Zhao, *Tetrahedron Lett.*, 2014, 55, 6362–6366.
- 29 A. S. Hay, J. Org. Chem., 1962, 27, 3320-3321.
- 30 Y. Yamashita, M. Tomura, M. B. Zaman and K. Imaeda, *Chem. Commun.*, 1998, 1657–1658.
- 31 S. Bouzan, G. Chen, K. Mulla, L. N. Dawe and Y. Zhao, Org. Biomol. Chem., 2012, 10, 7673–7676.
- 32 J. Massue, J. Ghilane, N. Bellec, D. Lorcy and P. Hapiot, *Electrochem. Commun.*, 2007, 9, 677–682.
- 33 S. Inagi, K. Naka, D. Iida and Y. Chujo, *Polym. J.*, 2006, **38**, 1146–1151.
- 34 K. Naka, S. Inagi and Y. Chujo, J. Polym. Sci., Part A: Polym. Chem., 2005, 43, 4600–4608.
- 35 G. Meazza, G. Zanardi, G. Guglielmetti and P. Piccardi, *J. Fluorine Chem.*, 1997, **82**, 175–180.
- 36 R. Mancuso and B. Gabriele, *Molecules*, 2014, **19**, 15687–15719.
- 37 A. Shockravi, A. Javadi and E. Abouzari-Lotf, *RSC Adv.*, 2013, 3, 6717–6746.