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Non-Oxido-Vanadium(IV) Metalloradical Complexes with Bidentate 1,2-Dithienylethene Ligands – Observation of Reversible Cyclization of the Ligand Scaffold in Solution

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Abstract Derivatives of 1,2-dithienylethene (DTE) have superb photochromic properties due to an efficient reversible photocyclization reaction of their hexatriene structure and thus have application potential in materials for optoelectronics and (multi-responsive) molecular switches. Transition metal complexes bearing switchable DTE motifs commonly incorporate their coordination site rather distant from the hexatriene system. Here we describe the redox active ligand 1,2-bis(2,5-dimethylthiophen-3-yl)ethane-1,2-dione, that reacts with [V(TMEDA)₂Cl₂] to give a rare non-oxido vanadium(IV) species 3(M,M/P,P). This blue complex has two bidentate en-diolato ligands which chelate the V(IV) center and give rise to two five-membered metallacycles with the adjacent hexatriene DTE backbone bearing axial chirality. Upon irradiation with UV-A light or prolonged heating in solution, the blue compound 3(M,M/P,P) converts into the purple atropisomer 4(para,M/para,P). Both complexes were isolated and structurally characterized by single-crystal X-ray diffraction analysis (using lab source and synchrotron radiation). The antiparallel configuration (M- or P-helicity) present in both 3(M,M/P,P) and 4(para,M/para,P) is a prerequisite for (reversible) 6π cyclization reactions. A CW EPR spectroscopic study reveals the metallo-radical character for 3(M,M/P,P) and 4(para,M/para,P) and indicates dynamic reversible cyclization of the DTE backbone in complex 3(M,M/P,P) at ambient temperature in solution.

Introduction

Oxido-vanadium compounds, mostly in the oxidation state +V and +VI, are intensively studied for applications in a diverse range of areas. For instance, vanadate and vanadyl derivatives possess significant importance as biologically active species,^{1,2} have been shown to possess insulin-mimetic properties³⁻⁶ and are important catalytic species in various oxidation reactions.⁷⁻⁹ In contrast, non-oxido vanadium(IV) compounds are less frequently reported in the literature. Sproules, Wieghardt and co-workers described V(IV) complexes with non-innocent ligand-sets.¹⁰ A prominent compound is amavadin, which is a naturally occurring example of a non-oxido V(IV) coordination complex¹¹ that was initially isolated from the toadstool Amanita Muscaria. 12 Vanadocene dichloride and derivatives thereof have been investigated with respect to their antitumor activity.13 Vanadium coordination compounds show a broad range of potential applications in medicinal chemistry including the treatment of endemic tropical and various heart muscle diseases. A concise overview was recently given by Rehder.9

Herein we characterize a class of non-oxido vanadium(IV) complexes bearing two chelating en-diolato ligands with a 1,2 dithienylethene (DTE) motif as backbone, Scheme 1 F, G. DTE motifs have received considerable attention due to their superior photochromic properties, that is, DTEs undergo reversible photocyclization reactions with an associated characteristic color change. They have a significant application

Scheme 1. Atropisomers and photocyclization of 1,2-bis(2,5-dimethyl-3-thienyl)perfluorocyclopentane.

potential as a new photo switchable material for optoelectronics and (multi-responsive) molecular switches. The photochromic properties of DTEs are characterized, for instance, by a high fatigue resistance, high quantum yields, the robust thermal stability of all isomers, and a rapid response to the external stimulus. They are also responsive in the solid state.14-17 DTEs have been discussed as functional key components in a range of applications including conductive polymers¹⁸, multi-responsive molecular switches,¹⁹ optical memory devices²⁰, photo-responsive building blocks that can regulate supramolecular architectures,21,22 a selective fluorescent probe for the detection of metal ions,²³ and photoresponsive self-assemblies at liquid/solid interfaces.²⁴ DTE derivatives which show turn-on mode fluorescence promising candidates for applications in super-resolution fluorescence microscopy²⁴ and all-optical transistors.²⁶ A current active field of research addresses the control of motion in covalently bonded molecular systems, which is closely associated with the design of "molecular machines". In this aspect the control of conformational change is crucial. For instance, an established approach is the control of conformational change about a single bond triggered by an external stimulus.²⁷ Against this background, DTEs are reported to possess axial chirality resulting in the formation of atropisomers. Atropisomers (from Greek atropos = not turning) are stereoisomers (rotamers) occurring due to a hindered rotation about a single bond. Their rotational interconversion is hampered by a large-enough barrier to allow the individual 'minus' (M or R_a) and 'plus' (P or S_a) atropisomers to be 'locked' (see Scheme 1, A,B).28 Oki defines atropisomers and distinguishes them from rotamers on the basis that they are fully resolvable at ambient temperature, corresponding to a barrier of interconversion of >23.3 kcal/mol.²⁹ Atropisomerism, especially in biaryl derivatives bearing a chiral axis between two aromatic moieties, has been intensively exploited for the design of chiral ligand systems which find numerous applications in the field of asymmetric catalysis.30-35 Atropisomeric chirality, not only restricted to biarylic systems, is important in medicinal chemistry,³⁶ has pharmaceutical implications as an alternative source for the production of chiral drugs,37 and can be found in a variety of natural products.28,38

1,2-Bis(2,5-dimethyl-3-thienyl)perfluorocyclopentane, a typical example for a photochromic DTE-based derivative with axial chirality, is shown in Scheme 1 A-E. 15,16 The compound forms the open-ring isomer with parallel aligned thienyl groups (C) and the respective enantiomeric antiparallel forms with M- (A) and P-helicity (B).21,39 The photo-induced cyclization reaction can only occur from the antiparallel conformers A and B and gives rise to the closed enantiomeric forms **D** and **E** with (S,S) and (R,R) configurations,21 respectively, based on the asymmetric carbon atoms in 2-position of the thiophene units (see Scheme 1).39 Note, the parallel atropisomer cannot directly undergo photochemical cyclization,⁴⁰ and its coexistence with the antiparallel isomers lowers the efficiency of the switching process. Consequently, effort has been spent to increase the relative amount of the antiparallel atropisomers A and B.41

With this background, we now report on the DTE-based bidentate ligand (**F**) equipped with an en-diolato motif. Chelation of a vanadium(IV) metal center gives rise to the formation of a five-membered metallacycle (**G**). These non-oxido vanadium(IV) atropisomeric complexes with metallo radical character were then isolated and structurally characterized by single crystal X-ray (lab source and synchrotron radiation) diffraction analysis, UV/VIS, and EPR spectroscopy.

Results and Discussion

Scheme 2. Synthesis of the racemic vanadium(IV) compounds **3(***M*,*M*/*P*,*P*) and **4(***para*,*M*/*para*,*P*).

1,2-Bis(2,5-dimethylthiophen-3-yl)ethane-1,2-dione (1) reacts with an equimolar amount of vanadium(II)bis-N,N,N',N'tetramethylethylenediamine-dichloride ([V(TMEDA)2Cl2], 2) to vanadium(IV) enantiomeric $[V(DTE_{(apM)})_2TMEDA]$ 3(M,M) and $[V(DTE_{(apP)})_2TMEDA]$ 3(P,P). Note, the conversion of 1 to the en-diolato ligand is a twoelectron reduction. Therefore half of the vanadium must lead to a side product, presumably a 'VCl₄' chlorido species, which was not further investigated. The racemic mixture 3(M,M/P,P) was isolated as a deep blue micro-crystalline solid in 43 % yield with respect to starting material 2. In 1998, Rehder and coworkers gave structural precedence to a non-oxido bisenolato complex of vanadium(IV). Specifically, the redox reaction of compound 2 and benzil was reported to give the corresponding vanadium(IV) bis benzoin complex [V(Ph(O)C=C(O)Ph)₂(TMEDA)].⁴² We applied a similar synthesis strategy; the V(II) compound 2 serves as both metal precursor complex and reducing agent for the redox-active ligand precursor 1. That is, the 1,2 diketone motif in 1 is reduced to an enediolate, which is stabilized upon coordination to a vanadium(IV) center to form the non-oxido en-diolato complex **3(M,M/P,P)**. Thus a hexatrien motif is formed which is a prerequisite for a (reversible) 6π peri-cyclic reaction. Consistently, the cyclic voltammogram of 1 in THF reveals a quasi-reversible redox-wave at a half potential $E_{1/2}^1 = -1.93 \text{ V}$ with respect to the ferrocene/ferrocenium (Fc/Fc⁺) redox couple. However, whilst the voltammogram suggests a reversible process for the first redox step, the second redox process is observed at very low potential $(E^2_{red} = -2.80 \text{ V} \text{ and } E^2_{ox} = -2.63 \text{ V})$ and is not-fully reversible (see Figure 1).

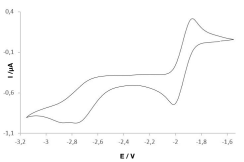


Figure 1. Cyclic voltammogram of **1** (3.9 mm) at 298 K, in 0.1 M [TBA]PF₆ in THF, scan rate 100 mV/s, referenced to Fc/Fc $^+$. Quasireversible redox wave at half potential $E^1_{1/2} = -1.93 \text{ V}$; $E^2_{\text{ox}} = -2.60 \text{ V}$, $E^2_{\text{red}} = -2.80 \text{ V}$.

To the best of our knowledge, no five-membered metallacycle with a dithienylethene (DTE) backbone has been reported to date. The compounds entail two axially chiral en-diolato ligands with both DTE ligands in each enantiomer exhibiting antiparallel configurations with M,M-helicity 3(M,M) and P,Phelicity 3(P,P). Upon irradiation of crystalline 3(M,M/P,P) dissolved in toluene with UVA light (λ_{max} = 370 nm) or prolonged heating of a toluene solution at 80°C, the racemate 3(M,M/P,P)is converted to the closely related [V(DTE_(par))(DTE_(apM))TMEDA] 4(para,M) [V(DTE_(par))(DTE_(apP))TMEDA] **4(para,P)**. Each of the formed enantiomers exhibits one DTE ligand in an open parallel arrangement and one DTE ligand in an antiparallel arrangement with the respective M-helical 4(para,M) or Phelical 4(para,P) configuration. The isomerization reaction is not reversible and is characterized by a significant color change from deep blue to purple (images are shown in Figure 3). The reaction can be followed in situ in consecutively recorded

UV/VIS absorption spectra of crystalline 3(M,M/P,P) in toluene under concomitant irradiation with UVA light as shown in Figure 2.

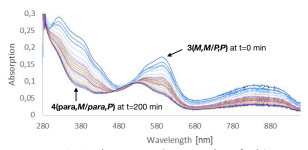


Figure 2. Stacked UV/VIS spectra of complex 3(M,M/P,P) (0.1 mM single crystals dissolved in toluene) under irradiation of UVA light $\lambda_{max} = 370$ nm for t = 0 - 200 min (10 min interval) results in the formation of 4(para,M/para,P).

The characteristic shoulder at 593 nm associated with 3(M,M/P,P) starts to decrease upon irradiation, as well as a decrease in intensity for the strong absorption band at 836 nm and for the signal in the range of 315-430 nm can be observed. After 200 min, the racemic mixture 4(para,M)/4(para,P) is formed as the major component. In this regard it is noteworthy, that photochromic DTE motifs in switchable transition metal complexes commonly feature the coordination site rather remotely located from the DTE moiety.⁴³ For instance, Wing-Wah Yam⁴⁴, Bielawski^{45,46} and co-workers reported the incorporation of the photochromic DTE motif in the backbone of an N-heterocyclic carbene. The coordination of such a NHC ligand to Rh(I) was reported by Bielawski and coworkers to possess switchable catalytic properties in alkene and alkyne hydroborations that are triggered by the photoinduced reversible cyclization of the DTE motif.⁴⁷ Recently, the same group described a photoswitchable olefin metathesis catalyst using a similar NHC ligand.48 Photochromic and luminescence switching properties of a Re(I) triscarbonyl complex with dithienylphenanthroline were published by Wing-Wah Yam, Phillips and co-workers. 49,50 However, nonswitching coordination complexes with DTE entities were reported where no cyclization reaction was observed.51

Single Crystal X-ray Diffraction Analysis

The racemic mixture of compound 3(M,M/P,P) crystallizes in the monoclinic centrosymmetric space group P2₁/n. Therefore, both enantiomers are present in equal amount in the unit cell, that is, the two DTE ligands in each enantiomer are either in an antiparallel M-helical 3(M,M) or antiparallel P-helical 3(P,P) arrangement with respect to the assignment of the highest priority to the methyl groups. Single crystals suitable for X-ray diffraction analysis were obtained by slow diffusion of npentane into a THF solution of 3(M,M/P,P). The determined molecular structure is shown for complex 3(M,M) in Figure 3, top left. The structure reveals the vanadium metal center in an octahedral coordination sphere, which consists of four oxygen donors associated with two bidentate DTE en-diolato ligands and the two nitrogen donors of the TMEDA ligand. The endiolate motif is characterized by short C=C double bonds (C7-C8 = 1.369(7) Å and C21-C22 = 1.365(7) Å) and O-C single (O1–C7 = 1.330(5) Å; O2–C8 = 1.345(5) Å C21 = 1.344(5) Å; O4-C22 = 1.349(6) Å)suggesting dianionic en-diolato ligand motifs coordinated to a formal vanadium(IV) center. Most significantly, due to a restricted rotation about the C-C bond connecting the thiophene units and the ethylene backbone, both DTE ligands in complex **3**(*M*,*M*/*P*,*P*) bear axial chirality and the enantiomeric complexes are atropisomers. Multiple attempts to grow single crystals of **4**(*para*,*M*/*para*,*P*) suitable for an X-ray diffraction study proved to be difficult due to the extraordinarily small size

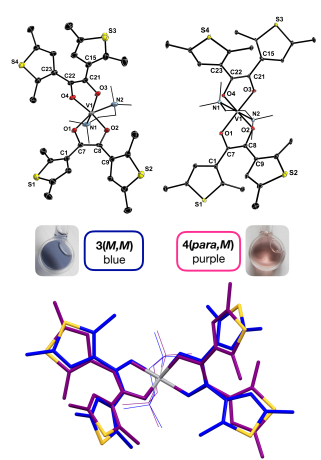


Figure 3. Top: Mercury plot of exemplary complex 3(M,M) (left) and 4(para,M) (right) at 50 % ellipsoid probability. Bottom: Wireframe structural overlap of 3(M,M) (blue) and 4(para,M) (purple). Hydrogen atoms are neglected for clarity. Selected bond lengths [Å]: 3(M,M): C1-C7 = 1.473(6),C7-C8 = 1.369(7),C8-C9 = 1.467(6),C7 = 1.330(5),O2-C8 = 1.345(5),C15-C21 = 1.481(7). C22-C23 = 1.473(7),C22 = 1.365(7), O3-C21 = 1.344(5)C22 = 1.349(6), V-N1 = 2.238(5), V1-N2 = 2.200(5). **4(para,M)**: C1-C7 = 1.475(7), C7 - C8 = 1.366(7), C8 - C9 = 1.471(7), O1 - C7 = 1.349(6),O2-C8 = 1.339(6), C15-C21 = 1.476(7), C21-C22 = 1.368(7), C23 = 1.472(6). O3-C21 = 1.345(6). O4-C22 = 1.351(5). N1 = 2.195(5), V-N2 = 2.215(5).

of the crystals obtained. This prompted us to undertake an Xray diffraction study using a synchrotron radiation source (beamline BL02B1 at SPring-8, Japan). A representative molecular structure is shown in Figure 3 top, right (4(para,M)). The racemic mixture 4(para,M/para,P) crystallizes in the monoclinic centrosymmetric space group P2₁/c with both enantiomers present in the unit cell. The molecular structures of 4(para,M/para,P) exhibit similar interatomic distances around the vanadium center and in the en-diolato backbone with respect to 3(M,M/P,P) [C=C double bond (C7-C8 = 1.366(7) Å and C21-C22 = 1.368(7) Å) and O-C single bonds (O1–C7 = 1.349(6) Å; O2–C8 = 1.339(6) Å and $C21 = 1.345(6) \text{ Å}; \quad O4-C22 = 1.351(5) \text{ Å})$]. However, structure unambiguously indicates a change to the parallel configuration in only one of the DTE ligands due to a rotation about the C-C bond connecting the thiophene rings and endiolato backbone. The other DTE unit retains its corresponding antiparallel M- or P-helicity to give the enantiomers **4(para,M)** and **4(para,P)**, respectively. An overlay plot of the blue compound **3(M,M)** and purple compound **4(para,M)** is shown in Figure 3, bottom. The crystallographic investigation of single crystals of **3(M,M/P,P)** and **4(para,M/para,P)** allows for the unambiguous structural characterization of the four atropisomers.

EPR Spectroscopic Study

With the aim of gaining insight into the solution structure, the blue 3(M,M/P,P) and purple 4(para,M/para,P) compounds were investigated by dissolving crystalline samples in THF. We cannot exclude isomerization upon dissolution, however, both solutions exhibit the characteristic blue and purple color, respectively. Both samples give similar but distinctly different EPR spectra as shown in Figure 4, which indicates the existence of two (or more) atropisomers. All experimental EPR spectra indicate metal centered radicals, that is, V(IV) metalloradicals. The purple solution spectrum at both room temperature and in frozen-solution is accurately simulated with a single EPR component denoted 4. These data were modelled with an electron Zeeman (principal g-values g_1 , g_2 , g_3) and a ^{51}V hyperfine (A_1, A_2, A_3) interaction with the isotropic values g_{iso} and Aiso used for the room temperature spectrum being the average of the three principal values of the corresponding interaction, e.g. $g_{iso} = (g_1+g_2+g_3)/3$ and $A_{iso} = (A_1+A_2+A_3)/3$.

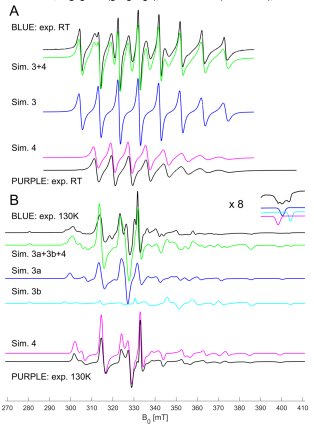


Figure 4. X-band (9.3810 GHz) CW EPR spectra measured at room temperature (A) and at 130 K (B) for the blue and purple solutions in THF. Simulated EPR model components and the sum are shown next to each experimental spectrum.

The blue solution was more complicated. Accurately simulating its isotropic room temperature spectrum requires a two component EPR model comprising of the same component 4 as observed in the purple solution and a new EPR component

denoted 3 which again was modelled by an isotropic g-value and isotropic vanadium hyperfine coupling (Table 1). In the room temperature spectrum, the two resolved EPR components have the relative weighting 0.54×3RT + 0.46×4. However, in frozen solution there are at least three EPR components resolved as can be deduced by examination of the high field feature in the CW EPR spectrum that arises from the largest principal value of the 51V hyperfine interaction, see the blowup of the region around 400 mT in Figure 4B. The appearance of three resolved features indicates the presence of three resolved EPR components. The observation of only two components at room temperature in the blue solution may be because of the presence of several species that interconvert rapidly in solution at room temperature to give an average spectrum $\mathbf{3}^{\text{RT}}$ whereas in frozen-solution the individual components of this average are observed. In the simulation of the frozen-solution data, these components are denoted 3a and 3b. Note that EPR component 4 does not appear to be involved in any room temperature averaging process as it is present in both the room temperature and frozen-solution data. A reasonable model for the blue spectrum was obtained with a simulation employing three EPR components (3ab,4) with approximately equal relative weights. Spin Hamiltonian parameters for all EPR model components are listed in Table 1. Note that the largest hyperfine value of **3a** and **3b** is accurately determined but the other g and A values are estimates as they are difficult to determine given the complexity of the spectrum particularly in terms of EPR component number. For example, component **3b** accurately accounts for the largest ⁵¹V hyperfine coupling observed in the experiment with its spectrum ranging from 280 mT to 403 mT, a range quite distinct from 3a and 4. To help assign the EPR components 3a, 3b and 4 to a chemical structure, various models were tested. For each structural model, density functional theory (DFT) was used to obtain a geometry optimized structure which was subsequently used to compute EPR parameters for comparison to the experimental data as shown in Table 1. The optimized geometries are listed in the Supporting Information. The DFT computed EPR parameters for the structure of the model 4(para,M/para,P) fit the EPR parameters from the purple solution very accurately (Table 1) and thus allow EPR component 4 to be assigned to atropisomer 4(para,M/para,P) with metalloradical character in solution. However, assignment of the EPR components 3a and 3b from the blue solution on the basis of the EPR parameters (g-values and 51V hyperfine couplings) was not as definitive as for 4. Seven structural models were investigated: The model for compound 3(M,M/P,P) and the (partially) closed compounds 5^{DFT}(M,closed-S,S), 6^{DFT}(closed-S,S,closed-S,S) 9^{DFT}(closed-R,R,closed-S,S), and 10^{DFT}(P,closed-S,S), in which the DTE moiety underwent a pericyclic ring-closure. Model 7^{DFT}(M,P) has two DTE ligands with M- and P-helicity, respectively. Structural model 8^{DFT}(para,para) carries both DTE ligands in a parallel alignment. Component 3b exhibits a large and well determined 51V principal hyperfine coupling value of $A_3(51V) = -481$ MHz, which is best matched by the group of (partially) closed compounds 5DFT(M,closed-S,S), 6DFT(closed-S,S,closed-S,S), 9DFT(closed-R,R,closed-S,S), and 10DFT(P,closed-S,S) with $A_3^{DFT}(^{51}V)$ ranging from -413 to -498 MHz. The other models all have significantly lower largest ADFT(51V) values in the 300 MHz range. On this basis, EPR component 3b is tentatively assigned to a compound class bearing cyclized DTE ligands. Component 3a has $A_3(^{51}V) = -385$ MHz and given the limited experimental accuracy of the other A and g principal values, the signal could arise from a compound with both DTE

in their non-cyclized form such as in 3(M,M/P,P), $7^{\text{DFT}}(M,P)$ or $8^{\text{DFT}}(para,para)$ (see Scheme 3). These three DFT models all have similar A and g-values. Given the complexity of the experimental spectrum, we cannot rule out either that a combination of EPR components with slightly different EPR parameters are required to accurately explain the 3b CW EPR signals.

Table 1: EPR parameters for the blue and purple samples derived from experimental and DFT data. Simulation of the CW EPR spectra provide for each component the principal g-values, ^{51}V hyperfine couplings (MHz), linewidth (MHz), and for room temperature data, the rotational correlation time τ_R (ns). For the DFT geometry optimized structures EPR parameters are computed (g-values and ^{51}V hyperfine couplings) and compared to the experimental parameters.

EPR Parameters		Principal Values	Isotropic	
'Purple' component				
Model 4	g	1.9114, 1.9694, 1.9886	1.9564	
	51V (1)	-371, -268, -69	-236	
	LW	58, 42, 23		
	τ_{R}	0.14		
4 ^{DFT} (para,M/para,	g	1.9276, 1.9682, 1.9757	1.9571	
P)	⁵¹ V	−365, −226, −81	-224	
'Blue' component				
Model 3 (RT)	g	NA	1.9735	
	51V (1)		-271.8	
	LW			
	τ_R	1010 1100	0.44	
Model 3a (130K)	g 51V (1)	1.910, ~ 1.97, ~ 1.98	ND	
	LW	-385, ~ -260, ~ -70		
Model 3b (130K)		57, 42, 30 ~ 1.92, ~ 1.94, 1.970 (2)	ND	
Model 36 (130K)	g 51V (1)		ND	
	LW	~ -260, ~ -250, -475		
		57, 42, 30	1	
DFT optimized structures				
$3^{DFT}(M,M/P,P)$	g	1.9276, 1.9670, 1.9730	1.9560	
	51 V	−363, −232, −92	-167	
5 ^{DFT} (M,closed-S,S)	g	1.9083, 1.9377, 1.9752 ⁽²⁾	1.9404	
,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	51V	-131, -306, -413	-283	
6 ^{DFT} (closed-	g	1.9160, 1.9357, 1.9640 ⁽²⁾	1.9385	
S,S,closed-S,S)	51 V	-308, -223, -498	-343	
7 ^{DFT} (<i>M,P</i>)	g	1.9345, 1.9530, 1.9629	1.9501	
	51 V	−390, −198, −270	-286	
8 ^{DFT} (para,para)	g	1.9284, 1.9674, 1.9749	1.9570	
	51 V	−365, −224, −85	-225	
			1	
9 ^{DFT} (closed-	g	1.9149, 1.9328, 2.0386	1.9621	
R,R,closed-S,S)	⁵¹ V	−156, −323, −435	-305	
10DFT(P,closed-S,S)	q	1.8989, 1.9400, 1.9887	1.9425	
10 (1)010364-3,3)	51V	-109, -306, -431	-282	
	'	103, 300, 431	202	
	L	l .	1	

- (1) CW EPR only determines the absolute values of the hyperfine coupling; the sign was set according to the DFT
- (2) Principal g-values and 51 V hyperfine coupling not collinear, the hyperfine interaction in the g-matrix frame is given as A = R*A_{diagonal}*R^T, where R = [0.5121611 0.0082016 0.8588503; -0.0021891 0.9999636 0.0082437; 0.8588866 0.0023421 0.5121604].
- (3) Principal g-values and ⁵¹V hyperfine coupling not collinear.

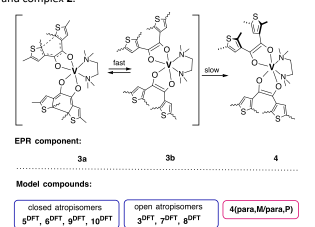
Summary and Conclusion

The redox-active ligand 1,2-bis(2,5-dimethylthiophen-3-yl)ethane-1,2-dione (1) reacts with [V(TMEDA) $_2$ Cl $_2$] (2) in a disproportionation reaction to give rare non-oxido vanadium(IV) species encompassing two bis-enolato ligands

with 1,2-dithienylethene (DTE) backbone. A blue crystalline product was isolated from the reaction mixture. Upon irradiation with a UVA light source or heating of a toluene solution, a color change occurs from blue to purple and a purple crystalline material was obtained from the solution. Our single crystal XRD analysis of both isolated compounds reveals the structure of the racemic complex **3**(*M*,*M*/*P*,*P*) (blue) and **4**(*para*,*M*/*para*,*P*) (purple).The antiparallel configuration of the thiophene rings of the DTE ligands in both complexes display axial chirality corresponding to M- or P-helicity. A parallel configuration is observed in complex **4**(*para*,*M*/*para*,*P*).

The vanadium(IV) center is chelated via the en-diolate of each DTE ligand giving rise to five-membered metallacycles with a hexatriene motif formed in the DTE scaffold. The antiparallel alignment present in both 3(M,M/P,P) and 4(para,M/para,P) is a prerequisite for potential (reversible) 6π cyclization reactions (Scheme 1) and thus for the development of e.g. photo-switchable metallo-radicals.

With the aim of characterizing the solution structures, X-band CW EPR spectra were recorded of solutions of crystalline samples of 3(M,M/P,P) and 4(para,M/para,P) dissolved in THF at room temperature and in frozen-solution. All obtained EPR spectra indicate vanadium centered radicals. The purple solution spectrum of 4(para,M/para,P) at both room temperature and in frozen-solution contained a single EPR active component 4 which is assigned to 4(para,M/para,P). The X-band EPR solution spectrum of the blue 3(M,M/P,P) indicates a complex dynamic behavior; two EPR components are resolved at room temperature comprising of 4 and another denoted 3RT. We assume that 3RT is an average spectrum of rapidly interconverting species as there are three (or more) components in the frozen-solution spectrum (including 4). The frozen-solution spectrum was modelled by 4 and components **3a** and **3b**. While **3a** may be described as a complex with one or two cyclized DTE units represented by model compounds $\mathbf{5}^{\mathrm{DFT}}$, $\mathbf{6}^{\mathrm{DFT}}$, $\mathbf{9}^{\mathrm{DFT}}$, or $\mathbf{10}^{\mathrm{DFT}}$, component $\mathbf{3b}$ is best described as a complex involving only open DTE units represented here by model compounds 3^{DFT}, 7^{DFT}, or 8^{DFT} or a combination thereof (see Scheme 3). Component 4 does not appear to be involved in dynamic averaging processes as 4 is definitively detected at room temperature as well as in frozen-solution. Compound 4 thus appears to be the thermodynamically stable atropisomer as it forms slowly from 3(M,M/P,P) solutions. The formation of the blue component 3 is kinetically controlled in the course of the synthesis of 3(M,M/P,P) starting from ligand precursor 1 and complex 2.



Scheme 3. Dynamic interconversion of EPR components **3ab** and **4** in solution and assignment of DFT models.

In conclusion, switchable transition metal complexes bearing photochromic DTE motifs commonly have the coordination site rather distant from the DTE backbone and its hexatriene system. Here we report the bidentate en-diolato ligand \mathbf{F} , which directly chelates V(IV) centers giving rise to five-membered metallacycles with adjacent DTE hexatriene backbone. Although we do not observe well-defined photochemically triggered cyclization, we found evidence for dynamic reversible cyclization reactions starting from $\mathbf{3}(M,M/P,P)$ at ambient temperature in solution.

Experimental

Experimental details for single crystal X-ray diffraction analysis of complexes 3(M,M/P,P) and 4(para,M/para,P): Diffraction experiments were performed on a Bruker D8 Venture in-house diffractometer using MoKα radiation and at the beamline BL02B1 of SPring-8 in Hyogo, Japan, using a wavelength of 0.3567 Å. The collection of the dataset of compound 4(para,M/para,P) was done under exclusion of light and oxygen by using argon atmosphere while preparing crystals. Further details are shown in Table 2 and in the crystallographic information files (CIFs) deposited with the Cambridge Structural Database (CSD) under deposition numbers 1946375-1946376. The CIFs can be accessed free of charge from https://www.ccdc.cam.ac.uk/structures/.

Table 2. Details on the measurements and refinements of compound **3**(*M*,*M*/*P*,*P*) and **4**(*para*,*M*/*para*,*P*).

	1		
Structure	3(<i>M,M</i>)/(<i>P,P</i>)	4(para,M)/(para,P)	
Sum Formula	C ₃₄ H ₄₄ N ₂ O ₄ S ₄ V		
Mol. mass	723.89		
Space group	P2 ₁ /n	P2 ₁ /c	
a (Å)	10.4187(5)	28.109(6)	
b (Å)	25.2837(11)	8.9569(18)	
c (Å)	13.8148(7)	14.529(3)	
β (°)	108.345(2)	94.47(3)	
V (ų)	3454.2(3)	3646.8(13)	
Z	4	4	
F ₀₀₀	1524	1524	
Radiation	Mo Kα (0.71073 Å)	Synchr. (0.3567 Å)	
Temperature (K)	100(1)	20(1)	
Crystal color	blue	purple	
Diffractometer	Bruker D8 Venture	SPring-8 Rigaku CCD	
Max. res. (Å)	0.82	0.75	
R _{int}	0.089	0.173	
hkl (-min=max)	12/30/16	37/11/19	
# of refln.	35517	63595	
# of indep. refln.	6455	9032	
R	0.064	0.089	
wR(F²)	0.131	0.188	
GoF	1.14	1.11	
Residual (e Å ⁻³)	0.60/-0.38	1.14/-0.73	
Obsv. criteria	F ² >2σ(F ²)		
CCDC entry	1946375	1946376	

Method and materials: If not mentioned otherwise all reactions were carried out in a heavy-walled Schlenk tube with a wide bore Teflon screw stopcock under an atmosphere of dry argon. Solvents and chemicals were used without further purification. UV/VIS Spectroscopy: Experiments were carried out at 298 K on a Varian Cary 50 UV/VIS-spectrophotometer. Cuvettes: 10 mm, synthetic quartz (QS) with Teflon cap. Probes were prepared under an inert atmosphere of argon. Cyclic Voltammetry: Experiments were carried out at 298 K with a Metrohm potentiostat running the NOVA 2.1 software package

using a RHD Instruments electrochemical cell TSC 1600 closed. Solvent: THF; Electrolyte: [TBA]PF6; Electrodes: Pt working electrode, Ag/AgCl pseudo reference electrode, Pt counter electrode. Continuous wave (CW) Electron Paramagnetic Resonance: X-band (9.3810 GHz) CW EPR measurements were carried out on a Bruker ElexSys E540 spectrometer equipped with a Bruker super-high-Q cavity and a Eurotherm LN2 cooling system. Room temperature measurements were made under non-saturating conditions using a microwave power of 20 mW with a modulation frequency of 100 KHz and a modulation amplitude of 0.1 mT. Measurement in frozen solution used a microwave power of 5.0 mW (non-saturating) with a modulation frequency of 100 KHz and modulation amplitude of 0.3 mT. The spectrometer field was calibrated with 2,2-Diphenyl-1-picrylhydrazyl (DPPH: g = 2.0036). Simulations used the XSophe⁵² and Easyspin⁵³ software. Density functional theory: Calculations to compute EPR parameters were performed using ORCA 3.0.3.54,55 Geometry optimization was performed using the UKS B3LYP functional, a T2V basis set and a frozen core for all electrons, and a COSMO model for THF. The g-, hyperfine and nuclear quadrupole tensors were calculated using a spin-unrestricted Kohn-Sham self-consistent field calculation method with a ZORA scalar relativistic Hamiltonian and the B3LYP functional, using the Def2QZVPP basis set for V and the def2TZVPP basis set for all other nuclei,56 and tighter convergence constraints. The spin-orbit coupling was calculated using the SOMF(1X) option in ORCA.

[$V_2(\mu\text{-Cl})_3(\text{THF})_6]_2[\text{Zn}_2\text{Cl}_6]$ was prepared following a slightly modified literature procedure:⁵⁷ (I) $V\text{Cl}_3(\text{THF})_3$: $V\text{Cl}_3$ (0.50 g, 3.18 mmol, 1.00 eq.) was suspended in THF (40 mL) and stirred for 48 h at 75 °C. The formed light purple solution was subsequently filtered and the solvent removed under reduced pressure. Yield: 1.18 g (3.15 mmol, 99%). (II) $V\text{Cl}_3(\text{THF})_3$ (1.00 g, 2.68 mmol, 1.00 eq.), dissolved in THF (50 mL), was treated with zinc powder (0.44 g, 6.69 mmol, 2.50 eq.) and stirred for 48 h at room temperature. After removal of the solvent under reduced pressure, the green product was dissolved in DCM, filtered, dried and washed with small portions of n-pentane. Yield: 0.87 g (0.54 mmol, 80%).

VCl₂(TMEDA)₂ was prepared following a modified literature procedure: 58 [V₂(μ -Cl)₃(THF)₆]₂[Zn₂Cl₆] (0.70 g, 0.43 mmol, 1.00 eq.) was suspended in THF (30 mL), treated with N,N,N',N'-Tetramethylethylenediamine (2.3 mL, 15.07 mmol, 35.00 eq.), stirred at 75 °C for 6 h and then at room temperature for 16 h. After partial removal of the solvent under reduced pressure the red solution separated blue crystals upon standing overnight in a cryostat at 5 °C. The supernatant was decanted and the product washed with small portions of cold THF. Yield: 0.53 g (1.51 mmol, 87%).

1,2-bis(2,5-dimethylthiophen-3-yl)ethane-1,2-dione (DTE, 1): was prepared following a modified literature procedure:59 In a three neck flask, AlCl₃ (9.39 g, 70.42 mmol, 1.00 eq.) was suspended in DCM (40 mL), cooled to -15 °C and treated with pyridine (2.8 mL, $\rho = 0.978 \text{ g/cm}^3$, 35.21 mmol, 0.50 eq.) and 2,5-dimethylthiophene (8.0 mL, ρ = 0.985 g/cm³, 70.42 mmol, 1.00 eq.), each dissolved in DCM (40 mL). Over a time period of minutes, oxalyl chloride (3.6 mL, $\rho = 1.48 \text{ g/cm}^3$, 42.25 mmol, 0.60 eq.), dissolved in DCM (40 mL), was added to the red solution. After stirring for 40 minutes at -5 °C and 1 h at 5-10 °C, the mixture was poured onto ice. The organic layer was separated and the aqueous layer extracted with chloroform. The combined organic phases were then washed with water, sodium carbonate solution and saturated NaCl solution subsequently. Filtering through hot cotton and removal of the solvent under reduced pressure was followed

by flash chromatography (cyclohexane/ethylacetate, 100:3) of the viscous oil and led to pure orange-brown product. Yield: $4.12 \, \mathrm{g}$ ($14.80 \, \mathrm{mmol}$, 42%), $R_f = 0.38$.

[V(DTE_(apM))₂TMEDA] / [V(DTE_(apP))₂TMEDA] (3(M,M/P,P)): Compound 1 (0.20 g, 0.71 mmol, 1.00 eq.) was added to a suspension of [VCl₂(TMEDA)₂] (0.25 g, 0.71 mmol, 1.00 eq.) in THF (15 mL). The resulting dark blue solution was stirred at 75 °C for 24 h. After filtration and partial removal of the solvent under reduced pressure, n-pentane was allowed for seven days to diffuse into the saturated solution allowing for the crystallization of product 3. The supernatant was decanted and the product was washed with small portions of n-pentane and subsequently dried in a stream of argon. Yield: 0.22 g (0.30 mmol, 43%).

 $[V(\mathsf{DTE}_{(\mathsf{par})})(\mathsf{DTE}_{(\mathsf{apM})})\mathsf{TMEDA}]/[V(\mathsf{DTE}_{(\mathsf{par})})(\mathsf{DTE}_{(\mathsf{apP})})\mathsf{TMEDA}]$ (4(para,M/para,P)): Photo-chemical isomerization: In a typical experiment, a 20 mL fused-quartz Schlenk tube with Teflon valve was charged with a toluene solution of compound 3(M,M/P,P) (0.01-0.02 mM). The sample was irradiated with UVA light (λ_{max} = 370 nm), performed in a custom-built photoreactor with two 11W UV-A lamps (Philips UV-A PL-S 11 W/10/2P, 28 mm diameter, 236 mm total length) equipped with a cooling ventilator. Distance to probe: 10 cm. The reaction is characterized by a color change from deep blue to purple. The solvent was removed under reduced pressure after 4-6 h reaction time. The product was isolated as purple powder in quantitative yield. Crystals suitable for single crystal X-ray diffraction analysis using synchrotron radiation at the BL02B1 beamline at SPring-8 in Hyogo, Japan, were obtained from a saturated solution of 4(para,M/para,P) in toluene upon standing for several days.

Thermal isomerization: A toluene solution of **3(***M*,*M*/*P*,*P*) (0.1-0.2 mM) was transferred in a Schlenk tube with a wide bore Teflon screw stopcock and heated to 80 °C for 12 h. The reaction is characterized by a color change from deep blue to purple. The supernatant was removed in *vacuo* leaving compound **4(***para*,*M*/*para*,*P*) as purple solid.

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