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Syntheses and Optical Properties of Azo-Functionalized Ruthenium Alkynyl Complexes**

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Abstract: The syntheses of *trans*-[Ru(C=C-1-C₆H₄-4-N=N-1-C₆H₄-4-C=C-1-C₆H₄-4-NO₂)Cl(L₂)₂] (L₂ = dppm (**Ru1**), dppe) (**Ru2**)), *trans*-[Ru(C=C-1-C₆H₄-4-N=N-1-C₆H₄-4-(*E*)-CH=CH-1-C₆H₄-4-

NO₂)Cl(dppe)₂] (Ru3), and *trans*-[Ru(C=C-1-C₆H₄-4-(*E*)-CH=CH-1- $C_6H_2-2, 6-Et_2-4-N=N-1-C_6H_4-4-NO_2)CI(dppe)_2$ (Ru4) are reported, together with those of precursor alkynes. Their electrochemical properties were assessed by cyclic voltammetry (CV), linear optical and quadratic nonlinear optical (NLO) properties assayed by UV-Vis-NIR spectroscopy and hyper-Rayleigh scattering studies at 1064 nm, respectively, and their linear optical properties in the formally Ru^{III} state examined by UV-Vis-NIR spectroelectrochemistry. These data were compared to those of analogues with E-ene and yne linkages in place of the azo groups. The Ruilli oxidation potentials shift to higher potential on replacing yne-linkage with E-ene linkage (proceeding from Ru2 to Ru3), changing the coordinated diphosphine ligand from dppm (Ru1) to dppe (Ru2), and locating the azo group proximal to the metal centre (proceeding from Ru4 to Ru1, Ru2, and Ru3). The wavelength of the UV-Vis λ_{max} band red-shifts on proceeding from the complex with a distal azo group (Ru4: 503 nm) to complexes with a proximal azo group (Ru1-Ru3: 540-550 nm). The quadratic nonlinearity β_{1064} increases on replacement of co-ligand dppm by dppe (proceeding from 1 to 2), while location of the azo group distal rather than proximal to the metal (proceeding from Ru2 to Ru4 or from Ru1 to trans-[Ru(C=C-1-C₆H₄-4-C=C-1- C_6H_4 -4-N=N-1- C_6H_4 -4-NO₂)Cl(dppm)₂] (Ru5)) leads to a decrease in quadratic NLO parameters, and replacement of yne-linkage by Eene-linkage (proceeding from 2 to 3) affords data that are within the $\pm 10\%$ error margins. The frequency-dependent β values for the azocontaining complexes from the present study are comparatively large (ca. 3000 x 10-30 esu for Ru2 and Ru3) but resonance enhanced. The formally Ru^{III} complexes exhibit low-energy bands at ca. 11 000 cm⁻¹ for Ru1-Ru4, with a progression to lower frequency on replacement of co-ligand dppm by dppe (proceeding from Ru1 to Ru2), replacement of yne-linkage with E-ene-linkage (proceeding

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from **Ru2** to **Ru3**), or location of the azo group further from the metal centre (proceeding from **Ru2** to **Ru4**). Computational studies using time-dependent density functional theory were undertaken on model compounds (**Ru2'-Ru4'**) to rationalize the optical behaviour of the experimental complexes. The HOMO for all model compounds is mainly located on the CIRuC=C-1-C₆H₄-4- unit, while the LUMOs for **Ru2'** and **Ru3'** are delocalized on the N=N-1-C₆H₄-4-C=C-1-C₆H₄-4-NO₂ group, and in **Ru4'** primarily located at the N=N-1-C₆H₄-4-NO₂ group. The lowest-energy LUMO (L) \leftarrow HOMO (H) transitions involve a mixture of MLCT and ILCT character with a LLCT contribution. The second- and third-lowest energy transitions originate from L+1 \leftarrow H and L \leftarrow H-2 electronic transitions; H-2 is delocalized across the N=N-1-C₆H₄-4-C=C unit, while L+1 primarily involves N=N (\pi*) and 1-C₆H₄-4-NO₂ C-N (\pi).

Introduction

Materials that can modify the propagation characteristics of light (e.g. frequency, phase, path, etc) are urgently needed for applications in current and prospective photonics technologies. Organic molecules possessing such "nonlinear optical" (NLO) properties that can fulfill these desired functions generally contain π -systems with highly delocalizable electron density. Very early in the search for NLO-efficient organic molecules, it was found that dipolar species such as *p*-nitroaniline (PNA) and 4-dimethylamino-4'-nitrostilbene (DANS) were particularly active, although subsequent research with molecules possessing other multipolar charge distributions (quadrupolar, octupolar) has also afforded NLO-efficient materials.^[1] Improvements to the classical donor-*π*-bridge-acceptor composition of molecules such as PNA and DANS have focussed on modifying all three components. Thus, the donor group strength increases on proceeding from amino and dimethylamino to julolidinyl and diphenylamino, and the acceptor group strength increases on proceeding from nitro tricyanovinyl and 2-dicyanomethylen-3-cyano-4,5,5to trimethyl-2,5-dihydrofuran, while delocalization through the π bridge is improved on proceeding from phenyl to heterocyclic moieties with a lower aromatic stabilization energy; all of these modifications favourably impact on NLO parameters. In other structure-NLO property studies, it was noted that the key quadratic NLO coefficient β is increased on certain phenyl-based π -bridge modification (proceeding from biphenyl to Z-stilbene, tolane, and E-stilbene, because biphenyl exhibits unfavourable ortho-H steric pressure, reducing bridge co-planarity, while compounds with a Z-stilbene bridge have decreased charge separation in the key excited state compared to E-stilbene analogues.^[2]



Scheme 1. Preparation of diazene-containing alkynes 3, 6 and 11.

In contrast to the aforementioned structural modifications that improve NLO performance, the incorporation of more electronegative heteroatoms such as nitrogen into the π -bridge in organic NLO materials (proceeding from *E*-ene- to imino- or azo-linked compounds) frequently leads to a reduction in β value due to charge localization at the electronegative nitrogen atom(s).^[3] Nevertheless, the azo-linked examples, in particular, are of interest as archetypical examples of photochemical switches; irradiation of azobenzenes at specific wavelengths can provide a facile means to reversibly convert between the *E*- and *Z*-configured forms, although thermal conversion of the Z-form to the *E*-form imposes constraints on the utility of this specific molecular switch.^[4]

Replacing the organic donor group in the classical donor- π bridge-acceptor molecular composition with a ligated metal unit can confer certain advantages: the flexibility of metal, valence electron count, co-ligands, and coordination geometry can all impact favourably on the molecular NLO coefficients,^[5] while the facile reversible oxidation/reduction undergone by many metal complexes can introduce important functionality and the potential to function as molecular NLO switches.^[6] We have previously reported that azo-containing arylalkynyl complexes can possess significant NLO performance, with the relevant parameters amongst the largest for organometallic complexes.^[7] However, the limited examples thus far generally have short π bridges, and thus the NLO behaviour may not be optimized. We report herein the syntheses of several new azo-containing arylalkyne compounds and derivative alkynyl complexes, spectroscopic and in some cases X-ray structural characterization of the new compounds, spectroscopic, electrochemical, linear and nonlinear optical, and in most cases X-ray structural characterization of the new complexes, comparison to precedents with shorter π -bridges or a related ene/yne-linked composition, and theoretical studies to rationalize the experimental observations.

Results and Discussion

Synthesis and characterization. A specific complex composition was targeted in the present studies. Previous reports had revealed an increase in the key NLO coefficients upon increasing the formal metal valence electron (VE) count,^[8] so 18 VE complexes were targeted in the present work. The earlier studies had also shown the efficiency series iron < ruthenium < osmium for group 8 metal alkynyl complexes,^[9] with the facile synthesis of the ruthenium examples more than compensating for the slightly decreased NLO efficiency compared to the kinetically more inert osmium analogues, so the present studies focused on ruthenium complexes. The precedent research had also revealed that pseudo-octahedral complexes *trans*-[Ru(C=CR)Cl(L₂)₂] [L₂ = dppm, dppe; dppm = bis(diphenylphosphino)methane, dppe 1.2bis(diphenylphosphino)ethane] can be highly NLO-efficient and can possess fully-reversible formally Ru^{II/III} redox processes accompanied by strong linear and nonlinear optical changes.^[10] Complexes with this specific composition were therefore targeted in the present study.

The alkynyl complexes were accessible in a straightforward way employing existing methodology, namely reaction of the

corresponding terminal alkyne with a metal chloride complex.^[11] The syntheses of the alkynes proceeded smoothly (Scheme 1), the products being characterized where possible using IR and multinuclear NMR spectroscopies, high resolution mass spectrometry, and elemental analyses (Electronic Supporting Information). Thus, reaction of 1-(4-bromophenyl)-2-(4-((triisopropylsilyl)ethynyl)phenyl)diazene,[12] with trimethylsilylacetylene under Sonogashira^[13] coupling conditions afforded the difunctionalized trimethylsilylethynyl triisopropylsilylethynyl compound 1 (95%), which was selectively desilylated at the trimethylsilyl group using potassium carbonate to give 2 in excellent yield (93%). A second coupling reaction using 1-iodo-4-nitrobenzene^[14] afforded 3 (71%), with azo and yne linkages separating the silyl-protecting group from the nitro acceptor.

Synthesis of the analogue in which the yne linkage is replaced by an *E*-ene group was also successful, albeit in lower overall yield. Oxidation of 4-aminobenzonitrile using potassium peroxymonosulfate^[15] afforded the corresponding labile nitrosoarene which was employed without further purification, reaction with 4-((triisopropylsilyl)ethynyl)aniline^[16] giving the nitrile (E)-4-((4-((triisopropylsilyl)ethynyl)phenyl)diazenyl)benzonitrile (4, 75%). Reduction of **4** using (diisobutyl)aluminium hydride (DIBAL-H) afforded the aldehyde 5 (47%) which was reacted with diethyl (4-nitrobenzyl)phosphonate^[17] under basic conditions to afford silvl-protected terminal acetvlene the (F)-1-(4-((F)-4nitrostyryl)phenyl)-2-(4-((triisopropylsilyl)ethynyl)phenyl)diazene (6, 19%) in low yield.

The preparation of (E)-1-(3,5-diethyl-4-((E)-4ethynylstyryl)phenyl)-2-(4-nitrophenyl)diazene (**11**), in which the bridging azo and *E*-ene groups of **6** are interchanged, required a sequence including modified Sandmeyer reactions; in contrast to **6**, compound **11** necessitated incorporation of alkyl groups to ensure sufficient solubility. Diazotization of *p*-nitroaniline using



Scheme 2. Syntheses of ruthenium complexes Ru1 - Ru4.

isoamyl nitrite as the source of the nitronium ion,^[18] followed by reaction with 2,6-diethylaniline afforded the diazene (*E*)-2,6diethyl-4-((4-nitrophenyl)diazenyl)aniline (**7**, 71%) in good yield. The arylamino group underwent reaction with isoamyl nitrite and then potassium iodide to give the iodo analogue, (*E*)-1-(3,5diethyl-4-iodophenyl)-2-(4-nitrophenyl)diazene (**8**, 65%), which was in turn reacted with copper(**1**) cyanide/ferric chloride to give the nitrile (*E*)-2,6-diethyl-4-((4-nitrophenyl)diazenyl)benzonitrile (**9**) in 47% yield. Reduction of **9** with DIBAL-H afforded the corresponding aldehyde (**10**, 47%) which was reacted with diethyl (4-((trimethylsilyl)ethynyl)benzyl)phosphonate to give the desired terminal acetylene (**11**, 28%).

The preparation of the new ruthenium complexes proceeded by established methodologies (Scheme 2). Reaction between cis-[RuCl₂(dppm)₂]^[19] and the triisopropylsilyl-protected acetylene 3 in the presence of fluoride afforded Ru1 (59%); the fluoride desilvlated the acetylene in situ, eliminating the need for a further step to isolate the terminal alkyne. The 16-electron, five-coordinate species [RuCl(dppe)₂]PF₆^[20] was the precursor for the three dppe-containing complexes. It reacted under similar in situ desilvlation conditions with the triisopropylsilyl-protected acetylenes 3 and 6 to give Ru2 (44%) and Ru3 (55%), respectively, and reacted with the terminal acetylene 11 in the absence of fluoride to give Ru4 (88%). The vibrational spectra of the complexes possess bands corresponding to the stretching mode of the metal-bound C=C at 2037 - 2055 cm-1 for Ru1 -Ru4: this band moves to higher energy on replacing dope by dppm (proceeding from Ru2 to Ru1), replacing E-ene by ynelinkage (proceeding from Ru3 to Ru2) and locating the azo group closer to the nitro (proceeding from Ru3 to Ru4). The v(NO₂) is relatively invariant at around 1580 cm⁻¹. The ³¹P NMR spectra confirm trans-stereochemistry at the metal center, with a singlet at ca. -7 ppm for the dppm-containing complex Ru1, and at ca. 49 ppm for the dppe-containing Ru2 - Ru4.

The identities of **3**, **4**, **6**, **8**, **Ru1**, **Ru2**, and **Ru3** were confirmed by single-crystal X-ray diffraction studies (Figure 1 and Figures S1-S7, Tables S1 - S3). The bond lengths and angles are unremarkable for the organic compounds and the alkynyl complexes (distortions from co-linearity at the Cl1-Ru1-C1-C2-C3 vector are often seen with alkynyl complexes and commonly attributed to packing forces), while the dihedral angles are consistent with good co-planarity through the π -bridges in all complexes.



Figure 1. Molecular structure of $Ru2 \cdot CH_2 Cl_2$, with thermal ellipsoids set at the 40% probability level. Hydrogen atoms and the lattice dichloromethane

molecule have been omitted for clarity. Selected bond lengths (Å) and angles (°) are provided in Table S2.

Optical and electrochemical studies. The UV-Vis spectra of the silyl-protected and terminal acetylenes **3**, **6** and **11** show absorption maxima at ca. 380 - 400 nm, with the longest wavelength band due to **6** (Table S5, Figure S8). It is possible that the low-energy bands for **3** and **6** correspond to the superposition of two low-energy bands: the lowest-energy band in the UV-Vis spectrum of **11** is only about half as intense as the low-energy bands of **3** and **6**, and the spectrum of **11** contains a further similar-intensity band at ca. 289 nm, suggesting that location of the azo group remote from the nitro group (**3**, **6**) red-shifts this band resulting in its superposition with the existing lowest-energy band.

Coordination of 3 to the ruthenium centre results in a significant bathochromic shift of λ_{max} from 380 nm to 540 nm for the dppm-coordinated complex Ru1, and to ca. 550 nm for the dppe-containing complex Ru2. A similar red shift (from 375 nm to 550 nm) was seen on coupling 11 to the bis(dppe)ruthenium centre in Ru3 (Figure S9). Both of these complexes contain the diazene group proximal to the metal atom. A slightly smaller shift to 503 nm is seen on coordination of 11 to give Ru4, where the diazene group is distal to the ruthenium. Extinction coefficients are similar for Ru1 - Ru3, and slightly lower for Ru4 (Table 1, Figure S9); this absorption band intensity difference is consistent with observations with the precursor alkynes but, in contrast to the alkynes, no clear additional band is apparent at higher energy (although the broad absorption profile makes such analysis difficult). Comparison of the λ_{max} values to that of trans-

 NO_2)Cl(dppm)₂]^[7a] (**Ru5**, Table 2) confirms the red-shift in proceeding from complexes with a distal diazene to complexes with a proximal electron-withdrawing diazene group.

Cyclic voltammetry (CV) studies of solutions of the metalcontaining complexes in dichloromethane show a fully reversible



Figure 2. Progressive changes to the UV-Vis-NIR spectra on application of an oxidizing potential of 0.7 V to a solution of **Ru4**. Measured in CH_2CI_2 at -25 °C. Pt gauze working-, Ag wire reference and Pt wire auxillary electrodes.

Ru^{IVIII} oxidation process at 0.62 V (cf. FcH/FcH⁺ couple at 0.56 V) (Table 1, Figure S10) for **Ru2**, with a significant shift to a higher potential on replacing yne-linkage with *E*-ene linkage

Table 1. Cyclic voltammetric $^{\rm [a]}$ and UV-vis $^{\rm [b]}$ data for resting state and oxidized complexes.

	E ^{1/2} V [∆E V, i _{ac} /i _{pa}	Ru ^{ll} ν _{max} [ɛ̂] ^[c]]	Ru ^{III} ν _{max} [ε] ^[c]
Ru1	0.57 [0.09,1]	18 300 [2.6], 27 300 [2.5], 37 400 [3.4]	11 500 [0.43], 20 300 [sh, 1.5], 25 200 [sh, 2.2], 27 000 [2.3]
Ru2	0.62 [0.09,1]	18 200 [4.0], 27 100 [4.2], 40 100 sh [5.8]	11 100 [2.6], 21 100 [3.6], 38 200 [6.1]
Ru3	0.66 [0.05,1]	18 200 [3.8], 26 000 [3.8], 40 300 [5.2]	11 000 [0.68], 20 900 [4.2], 38 700 [5.7]
Ru4	0.53 [0.09,1]	19 800 [1.1], 26 000 [4.7], 40 000 [5.5]	10 900 [1.8], 20 500 [2.1], 23 500 sh [2.5], 26 000 [3.0], 36 900 [5.7]
Ru6 ^[c]	0.49 [-, 1]	21 500 [1.5]	9 700 [0.18], 10 800 [0.16]

Measured in CH₂Cl₂ [a] Pt disc working-, Pt-wire auxilliary- and Ag/AgCl reference- electrodes, with ferrocene internal standard ($E^{1/2} = 0.56$ V [0.09, 1]; [b] cm⁻¹ [10⁴ cm⁻¹ M⁻¹]; [c] ref. [11b].

proceeding to Ru3 (we recognize that the relevant HOMOs in such ruthenium alkynyl complexes have an alkynyl ligand component, but they certainly have significant metal-centered content, and the redox processes have been labelled as such for simplicity). Changing the coordinated diphosphine ligand from dppe (Ru2) to dppm (Ru1) results in a slight shift to lower potentials. Oxidation of the metal centre is most easily achieved for Ru4 (0.53 V), for which the electron-depleting effect of the azo group is reduced due to its increasing distance from the metal. The reduction processes are nitro-centred, occurring at ca. -0.9 V for Ru1 - Ru3, and -0.8 V for Ru4, again consistent with the varying azo location (the electron-withdrawing azo proximal to the nitrophenyl group in 4 results in more facile reduction).

Spectroelectrochemistry allows the measurement of spectral data, in this case UV-Vis-NIR data, whilst applying a redox potential to a solution of the complex in an optically transparent thin-layer electrochemical (OTTLE) cell. Using the redox data obtained from the CV experiments, oxidation potentials slightly higher than indicated on the CV scans were applied to dichloromethane solutions of Ru1 - Ru4, affording spectral progressions indicating the clean formation of the Ru^{III} species (Figures 2 and S11 - S14); subsequent application of a reducing potential to the oxidized species afforded a return to the starting spectra, with isosbestic points in each case. Frequency maxima and extinction coefficients are collected in Table 1, and show the emergence of a new low-energy band at ca. 11 000 cm⁻¹ for Ru1 - Ru4, with a progression to lower frequency on replacement of co-ligand dppm by dppe (proceeding from Ru1 to Ru2), replacement of yne-linkage with E-ene-linkage (proceeding from Ru2 to Ru3), or location of the azo group further from the metal centre (proceeding from Ru2 to Ru4).

$\begin{array}{c} Ru1 [Ru] = \textit{trans-}Ru2(dppm)\\ O_2N-\underbrace{\frown}_{N}-\underbrace{\frown}_{N}N & Ru2 [Ru] = \textit{trans-}RuC(dppe) \end{array}$								
$N - [Ru] = trans-RuCl(dppe)_2$ $O_2N - P - P - P$								
$N [Ru] = trans-RuCl(dppe)_2$ O ₂ N $ N$								
O ₂ N-{N-{}{								
O ₂ N-()-((dppm)) ₂ Ru7 [Ru] = <i>trans</i> -RuCl(dppm)) ₂ Ru7 [Ru] = <i>trans</i> -RuCl(dppe)) ₂								
Cmpd	λ _{max} ^[b]	β1064 ^[c]	$\beta_0^{[c]}$ $\Delta = 0$	$\beta_0^{[c]}$ $\Delta = 30 \text{ nm}$	$\beta_0^{[c]}$ $\Delta = 50 \text{ nm}$			
Ru1	551	2150	-115	145	185			
Ru2	552	2700	-155	190	240			
Ru3	548	3200	-140	195	260			
Ru4	503	1300	110	125	145			
Ru5 ^[d]	510	1215 ± 146	51±6					
Ru6 ^[e]	466	2090 ± 66	395 ± 12					
Ru7 ^[e]	468	2525 ± 175	460 ± 32					

Table 2. Summary of HRS results,^[a] and including simple and modified two-level calculations.

[a] Measured at 1064 nm; [b] nm, Ru1 - Ru4, Ru6 in CH₂Cl₂, Ru5, Ru7 in THF; [c] 10^{-30} esu in THF; [d] ref. [7a]; [e] ref. [11b].

Hyper-Rayleigh scattering studies. The quadratic nonlinearities of Ru1, Ru2, Ru3, and Ru4 were determined at 1064 nm using the hyper-Rayleigh scattering (HRS) technique and 20 ns pulses; the results are given in Table 2, together with the two-level corrected values, and data for related complexes collected under similar conditions. Problems with the two-level model have been discussed previously.[10a] While it is not generally considered adequate for donor-bridge-acceptor organometallics such as those herein, it may be useful in comparing closely related complexes and, with this caveat in mind, one can compare the data. Both experimental and twolevel corrected nonlinearities are consistent with replacement of co-ligand dppm by dppe leading to an increase in quadratic nonlinearity (proceeding from 1 to 2). Replacement of ynelinkage by E-ene-linkage (proceeding from 2 to 3) affords data that are within the $\pm 10\%$ error margins. Location of the azo group distal rather than proximal to the metal (proceeding from Ru2 to Ru4 or from Ru1 to Ru5) leads to a decrease in quadratic NLO parameters. Further comparison to our previously reported data for Ru6 and Ru7 reveals that the frequencydependent β values for the azo-containing complexes from the present study are comparatively large, but the proximity of the linear optical absorption bands to the second-harmonic wavelength of the Nd:YAG laser employed in the present studies (532 nm) inevitably results in azo-containing compound frequency-independent data that are significantly smaller than those of the *E*-ene-containing analogues.

Various modified two-level models have been proposed to account for the proximity of the second-harmonic wavelength to the optical absorption maximum, and specifically the effects of absorption line broadening and vibrational modes in both the ground and excited states.^[21] We have therefore examined our experimental data utilizing the modified two-level model of Wang,^[21b] in which:

$<\beta>_{HRS} = <\beta_0>F(\upsilon,\delta)$

where $F(\upsilon, \delta) = (1/3) \{[2/(1+2\upsilon)-1/(1-\upsilon^2)+2(1-2\upsilon)/((1-2\upsilon)^2+\delta)]^2 + [2\delta/((1-2\upsilon)^2+\delta^2)]^2\}^{1/2}, \upsilon = \lambda_{max}/\lambda, \delta = \Delta/\lambda_{max}, \Delta \text{ is} a damping factor associated with the broadening of the absorption peak, and <math display="inline">\lambda$ is the fundamental light wavelength (1064 nm in this case). Note that, if $\Delta = 0$, the expression above simplifies to that of the usual two-level model. Using this equation and assuming that Δ is 30-50 nm (estimated from the UV-Vis spectra: Δ likely differs from sample to sample), the recalculated $<\beta_0>$ values are listed in Table 2. For **Ru4**, the recalculated $<\beta_0>$ is close to that calculated with the simple two-level model, while for **Ru1-Ru3**, for which λ_{max} are closer to 550 nm, the modified two-level model yields a greater discrepancy; nevertheless, the trend in experimental maximal values (**Ru3** > **Ru2** > **Ru1** > **Ru4**) is maintained for the various two-level approximations.

Computational studies. Time-dependent density functional theory (TD-DFT) calculations were undertaken on a set of model complexes (Figure 3) to rationalize the experimental UV-Vis spectra. Calculated low-lying (< 30 000 cm⁻¹) singlet excited-state transitions and oscillator strengths are listed in Table S7, alongside the experimental data. Frontier molecular orbitals (FMOs) involved in the main transitions for the model compounds **Ru2'** and **Ru4'** are given in Figures S15 and S16, respectively; the FMOs obtained for **Ru2'** and **Ru3'** are very similar in their composition.

Calculated transitions for all model complexes in the lowerenergy region (< $30\ 000\ \text{cm}^{-1}$) can be divided into two different groups; each model complex exhibits an intense transition in the region $15\ 000\ \text{cm}^{-1}$ to $17\ 000\ \text{cm}^{-1}$ and one or two transitions with



Figure 3. Model compounds used in computational studies. $P-P = Me_2PCH_2CH_2PMe_2$. large oscillator strengths in the region 21 500 cm⁻¹ – 25 000 cm⁻¹. The calculated data are consistent with the experimental data, particularly for model compounds Ru2' and Ru3'. The second spectral band for Ru2 and Ru3 is broader than the lowestenergy band; this can be attributed to the presence of several transitions close in energy in this region, according to the calculated data (Table S7). The experimental lowest-energy band is slightly more intense than the second lowest-energy band, and this behaviour is nicely reproduced by the computational outcomes of the corresponding model compounds (Ru2'/Ru3'). In the case of Ru4', these calculated transitions are substantially red-shifted comapred to the UV-Vis data. However, the incorporation of ethyl substituents in the laboratory complex Ru4 to overcome the solubility problems may induce steric crowding around the E-ene linkage in the coplanar conformation, and thus distort the complex from the ideal coplanar arrangement of the phenylene groups which is assumed in these calculations. Further calculations reveal that distortion from a coplanar structure blue-shifts the low-lying transitions, in particular the lowest-energy band, and decreases their intensity.

Although the highest occupied molecular orbital (HOMO) for all model compounds is broadly similar (i.e. the HOMO is mainly located on the CIRuC=C-1-C₆H₄-4- moiety: Figures S15 and S16), the lowest unoccupied molecular orbitals (LUMOs) in **Ru2'** (and **Ru3'**) are delocalized on the N=N-1-C₆H₄-4-C=C-1-C₆H₄-4-NO₂ group, whereas in **Ru4'** electron density is primarily located at the N=N-1-C₆H₄-4-NO₂ group, particularly at the azophenylene unit. Overall, the lowest-energy LUMO (L) \leftarrow HOMO (H) transition involves a mixture of metal-to-ligand charge transfer (MLCT) and intra-ligand CT (ILCT) character, together with some contribution from ligand-to-ligand CT (LLCT).

The second- and third-lowest energy transitions, which lead to a somewhat broad band in the experimental UV-Vis spectrum of **Ru2'** (and **Ru3'**), originate from L+1 \leftarrow H and L \leftarrow H-2 electronic transitions. H-2 differs from H in that density leaks into the central part of the bridge [i.e. the N=N-1-C₆H₄-4-C=C unit]. π -bonding within the -azo and -yne groups (or the *E*-ene group in **Ru3'**) also contributes to H-2. L+1 is primarily N=N (π^*) and 1-C₆H₄-4-NO₂ C-N (π) based. The second-lowest-energy transition for the model complex **Ru4'** is characterized as promotion from H-2 (dominated by electron density around the CIRuC=C-1-C₆H₄-4-C=C-1-C₆H₄-4 unit) to L.

Static first hyperpolarizability values, β_{tot} , were calculated for all model complexes (Table S8). Replacing an yne linkage with an *E*-ene unit, on proceeding from **Ru2'** to **Ru3'**, has little effect on β_{tot} , an outcome that is consistent with laboratory observations (Table 2 and S8). The position of the azo linkage exerts a significant influence on the β value; the calculated findings clearly indicate that the complex in which the *E*-azo linkage is distal from the metal exhibits a much larger nonlinear response than the proximal isomer (proceeding from **Ru3'** to **Ru4'** leads to an approximately two-fold increase in calculated β), although this is contrary to experimental observations. Given the discrepancy with experimental data for the latter, the performance of calculations in reproducing these structureproperty outcomes was examined as a function of the method used (PBEPBE (the principal method used in these studies), CAM-B3LYP, and LC-PBEPBE: see ESI); all methods suggest that β increases significantly when the *E*-azo linkage changes its position from A to B (Figure 3), whereas replacing an yne group with an *E*-ene linkage has a minor effect.

The simple two-level model (TLM) developed by Oudar and Chemla^[22] is widely-used for extrapolating the experimental hyper-Rayleigh scattering (HRS) data (frequency dependent β value) to the static first hyperpolarizability, but this simple model is not applicable when the scattered photon frequency approaches the resonance frequency of the CT transition^[21b, 23] (see above), and thus the simple TLM is not suitable for evaluating the static hyperpolarizability of laboratory analogues of model complexes Ru2' and Ru3' (in particular) as they exhibit transitions very close to the second harmonic at 532 nm. Hence, equation (3) (ESI), which extends the simple two-level model into the resonant regime.^[21b] was used to calculate the static values. However, equation (3) assumes that only two electronic states contribute to the first hyperpolarizability. B^{CT}. It should be used with great caution when two spectral bands relatively close in energy exhibit MLCT character; this is particularly the case with Ru2 and Ru3, for which the high-energy bands may also contribute to the overall β . It should also be noted that although we considered only the lowest-energy band when deriving the experimental static value for Ru4, contribution from the more intense second-lowest-energy band (which is predominantly MLCT in character) to the $\beta_{0,exp}$ should also be taken into account.

Ru2' and **Ru3'** show similar linear optical behaviour in the low-energy region, possessing almost identical lowest-energy transitions. Moreover, the degree of MLCT character of these bands, on the basis of calculations, is almost the same, and thus it is not surprising that replacing an yne linkage with an *E*-ene group on proceeding from **Ru2'** to **Ru3'** has only a minor effect on the β value. In contrast, the calculated UV-Vis spectrum of **Ru4'** contains a significantly red-shifted lowest-energy transition compared to the lowest-energy transitions for **Ru2'** and **Ru3'**, which may account for the large β_{tot} calculated for model compound **Ru4'**.

Conclusions

The new complexes **Ru1-Ru4** and the previously reported **Ru5-Ru7** afford a suite of complexes that permit comment on the effect of azo group introduction and location on electronic and optical properties of metal alkynyl complexes. The **Ru**^{II/III} oxidation potentials shift to higher potential on replacing ynelinkage with *E*-ene linkage (proceeding from **Ru2** to **Ru3**), changing the coordinated diphosphine ligand from dppm (**Ru1**) to dppe (**Ru2**), and locating the azo group proximal to the metal centre (proceeding from **Ru4** to **Ru1**, **Ru2**, and **Ru3**); the lastmentioned observation is consistent with previous studies of ruthenium alkynyl complexes for which the electron-withdrawing influence of groups is particularly pronounced when they are appended to the proximal (to ruthenium) ring.^[10a] The wavelength of the UV-Vis λ_{max} band can be tuned by bridge variation, red-shifting on proceeding from complexes bearing Eene linkages (e.g. trans-[Ru(C=C-1-C₆H₄-4-(E)-CH=CH-1-C₆H₄-4-(E)-CH=CH-1-C₆H₄-4-NO₂)Cl(dppe)₂] (Ru7): 468 nm) to the complex with a distal azo group (Ru4: 503 nm) to complexes with a proximal azo group (Ru1-Ru3: 540-550 nm). The frequency-dependent β values for the azo-containing complexes from the present study are comparatively large (ca. 3000 x 10⁻³⁰ esu for Ru2 and Ru3) but strongly resonance enhanced; experimental values for Ru2 and Ru3 are similar to that of Ru7, but the two-level-corrected data for the azo-linked compounds are much smaller than that of the ene-linked Ru7 and, while being mindful of the problems associated with this approximation, we conclude that the ene-linked alkynyl complexes are more efficient NLO materials. The formally Ru^{III} complexes exhibit lowenergy bands at ca. 11 000 cm⁻¹ for Ru1-Ru4, and the location of this LMCT band can be tuned by molecular modification; it moves to lower frequency on replacement of co-ligand dppm by dppe (proceeding from Ru1 to Ru2), replacement of yne-linkage with E-ene-linkage (proceeding from Ru2 to Ru3), or location of the azo group further from the metal centre (proceeding from Ru2 to Ru4). Thus, Ru1-Ru4 are in principle electrochemically-(Ru^{II/III}) as well as photochemically- ($E \leftrightarrow Z$ azo) switchable optical materials.

Computational studies were undertaken on models Ru2'-Ru4' to rationalize this optical behavior. Calculations agree well with the experimental linear optical data, although we found a mismatch between the calculated and experimental lowestenergy bands for Ru4 as a consequence of employing a fully planar structure for Ru4' to represent the laboratory compound Ru4. In Ru4', the ethyl substituents used as an aid to mitigate solubility problems in Ru4 are removed because they may cause distortions in the bridge, and thus adversely affect both empirical peak position and intensity of the lowest-energy band. On the basis of calculations, the HOMO for all three model systems is broadly similar and mainly located on the CIRuC=C-1-C₆H₄-4- unit, while the LUMO for Ru4' is more localized compared to that for Ru2' and Ru3'. The LUMOs for Ru2' and Ru3' are delocalized on the N=N-1-C₆H₄-4-C=C-1-C₆H₄-4-NO₂ group, and in Ru4' primarily located at the distal N=N-1-C₆H₄-4- NO_2 group. The lowest-energy bands associated with L \leftarrow H excitation can be ascribed to a mixture of MLCT and ILCT character with a LLCT contribution. Although the decrease in β value on proceeding from Ru2-Ru3 to Ru4 is not reproduced computationally, due to a distorted non-coplanar structure for laboratory compound Ru4, as discussed earlier, both the experimental and calculated data reveal that replacing an yne linkage with an *E*-ene group on proceeding from Ru2 to Ru3 (or **Ru2'** to **Ru3'**) has only a minor effect on the β value due to the similar linear optical properties they possess in the low-energy region.

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Keywords

Organometallic chemistry • Molecular electrochemistry • Nonlinear optics • Transition metals

Supporting Information

Complete details of the syntheses and characterization of compounds 1-11 and complexes Ru1-Ru4, crystallographic studies of 3, 4, 6, 8, Ru1, Ru2.CH₂Cl₂, and Ru3.CH₂Cl₂, cyclic voltammetric, UV-Vis-NIR spectroelectrochemical and hyper-Rayleigh scattering studies (the last-mentioned at 1064 nm) of all new complexes, and theoretical studies of models of Ru2-Ru4 are reported in the Supporting Information.

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Entry for the Table of Contents

FULL PAPER



Dipolar organometallic complexes with a ligated ruthenium donor, nitro acceptor, and a π -delocalizable bridge incorporating an azo group exhibit large quadratic optical nonlinearities. The potentials of their reversible oxidation processes and the wavelengths of their absorption maxima in both the resting and oxidized forms are tuned by systematic structural modification.

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Syntheses and Optical Properties of Azo-Functionalized Ruthenium Alkynyl Complexes