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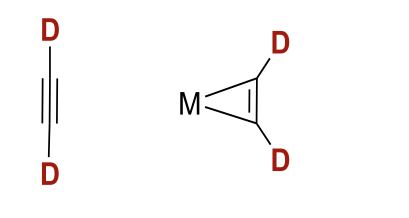
Traditio et Innovatio



New Alkyne Complex Based Framework for Visible Light induced Electron Transfer

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Introduction



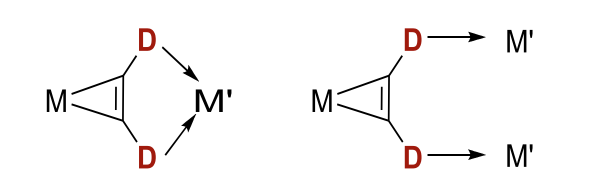
 W(II)-alkyne complexes bearing donor atoms in both α-positions

SPP 2102

 \rightarrow combines redox-active complex moiety with a potentially chelating unit

 \rightarrow valuable building blocks for polynuclear

Synthesis $\begin{array}{c} \stackrel{H}{\longrightarrow} \\ \stackrel{H}{\longrightarrow} \\ \stackrel{N}{\longrightarrow} \\ \stackrel{N}{\longrightarrow} \\ \stackrel{N}{\longrightarrow} \\ \stackrel{N}{\longrightarrow} \\ \stackrel{N}{\longrightarrow} \\ \stackrel{1) Py-CC-H, FcPF_{6}}{DCM} \\ \stackrel{Tp^{*}}{\longrightarrow} \\ \stackrel{N}{\longrightarrow} \\ \stackrel$



compounds with a short metal-metal distance and interesting redox behaviour \rightarrow so far, we were able to form heterobimetallic complexes using P, N, S or O as donor atoms [1]

Tp*

OC

Br

e

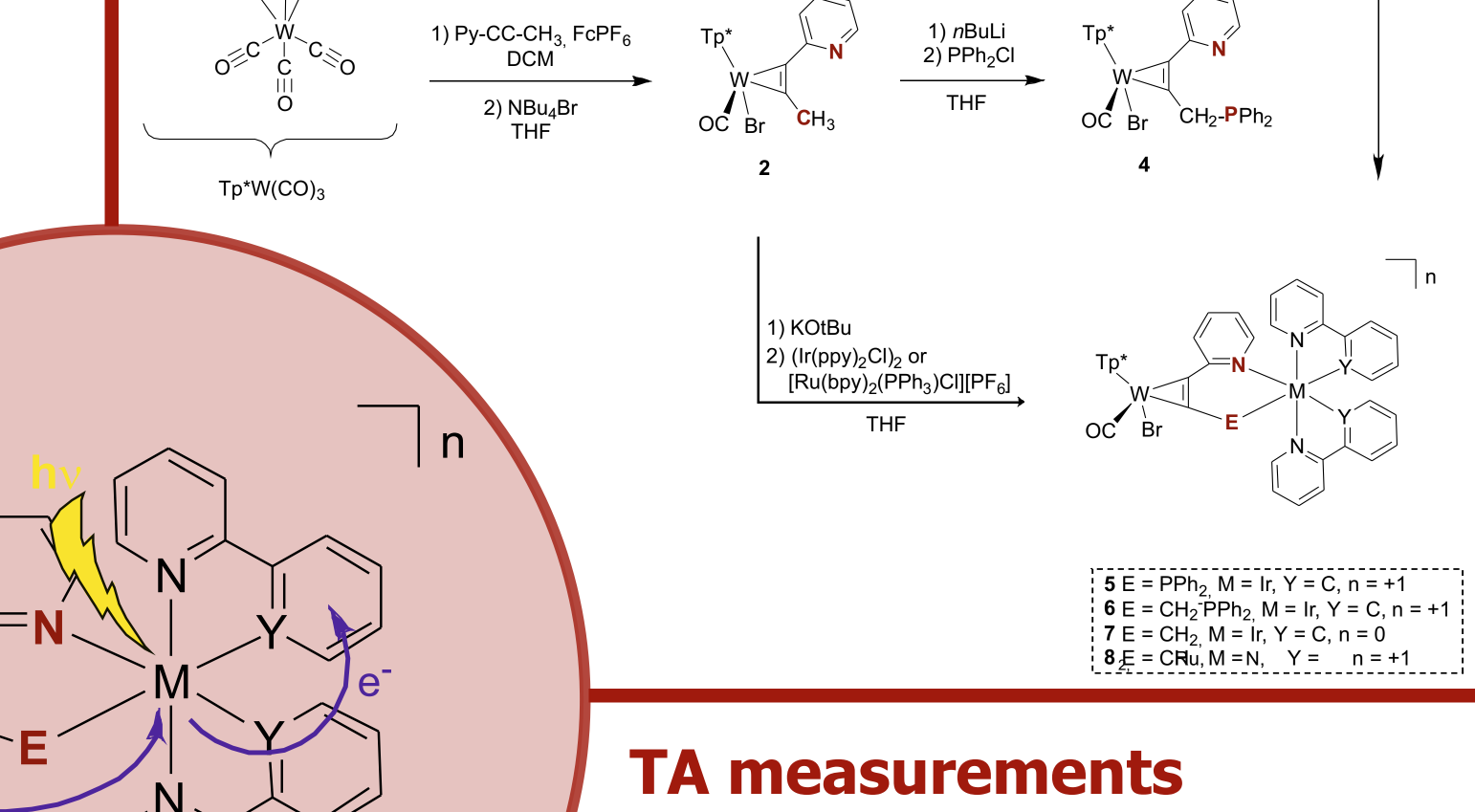
D = P, N, S, O

This project

- coordination of the alkyne complex via a cyclometallating pyridinyl propyne donor to an Ir(III) or Ru(II) centre, to mimic the phenyl pyridine ligand
- for comparison, we also used two different phosphine groups in place of the carbanion, since phosphines are also suitable donor atoms for photoactive polypyridine substituted complexes [2]

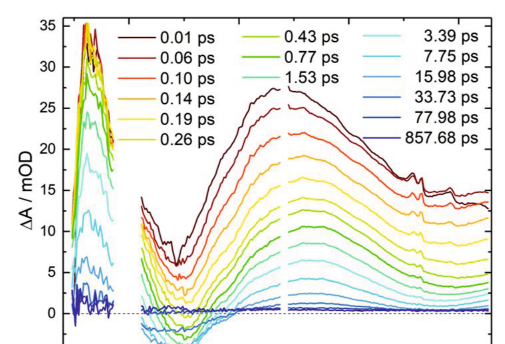
UV/vis and luminescence measurements

- UV/vis of 5-PF₆: shoulder 450 nm is assigned to W(II) to pyridine transitions; weak absorption at 710 nm due to d-d-transitions associated with the W(II)-centre
- UV/vis of 6-PF₆: shoulder at 500 nm is assigned to W(II) to pyridine transitions; absorption features being associated with the W alkyne complex moiety cannot be observed in the visible range
- compared to the mononuclear $Ir(ppy)_3$ the quantum yield of 5-PF₆ and 6-PF₆ is quenched to

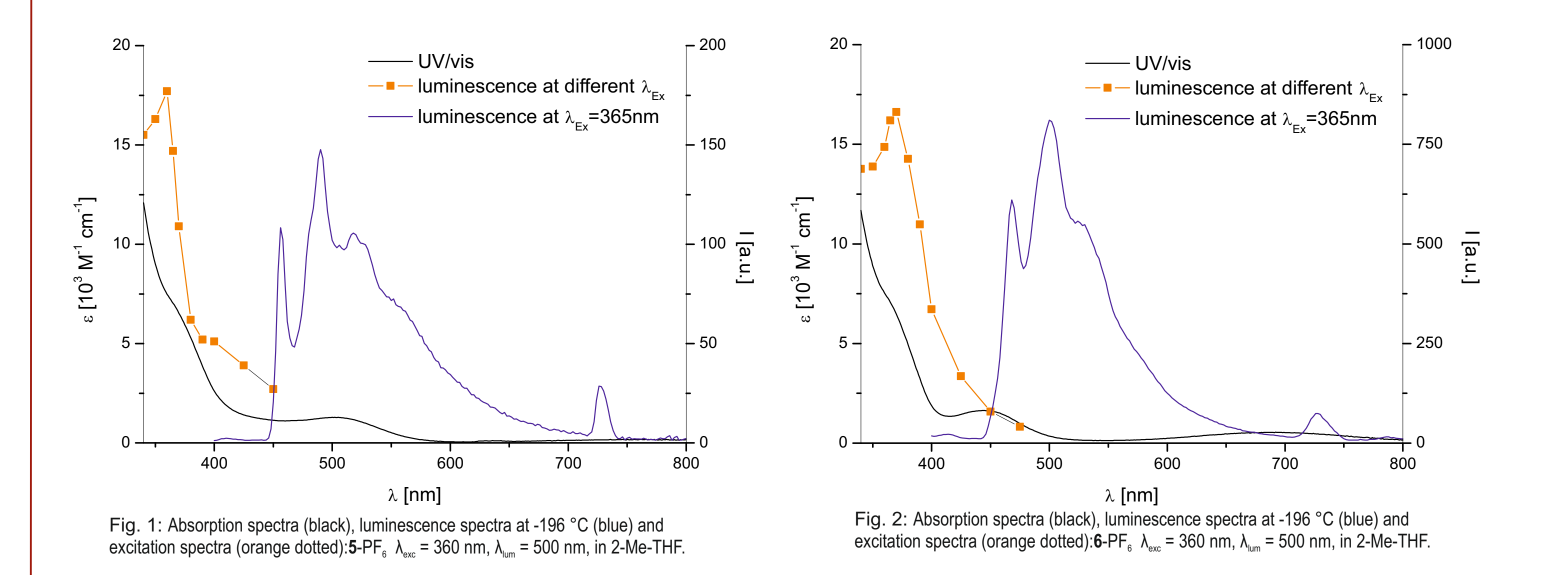


 the predominant broad excited state absorption between 500 and 650 nm, which is typical for a ³MLCT state Ir(IV)(ppy⁻) [4], decreases with a time constant of 5.2 ps

 the resulting state can be assigned to the charge separated W(III)-ppy complex formed by a W(II)



0.3 and 0.15%, respectively, due to the electron transfer from the W(II)-centre



• UV/vis of 7: shoulder 450 nm is assigned to W(II) to pyridine CT transitions; weak absorption at 710 nm due to d-d-transitions associated with the W(II)-centre

- quantum yield is quenched drastically compared to the mononuclear Ir(ppy)₃ (to 0.3 %), due to the electron transfer from the W(II)- centre
- UV/vis of 8-PF₆: transitions between 300 and 420 nm are ligand based π - π *-transitions, while the absorption around 470 nm are assigned to Ru to bipyridine CT transitions
- compared to the mononuclear $[Ru(bpy)_2(ppy)][PF_6]$ the quantum yield is quenched (to 40 %)

to Ir(IV) electron transfer due to the bleach feature at 450 nm

-0.08 ps

----- 0.17 ps

----- 0.31 ps

----- 0.61 ps

---- 1.15 ps

----- 1.98 ps

- 3.42 ps

5.58 ps

700

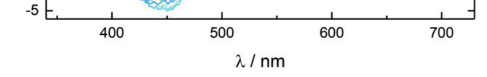


Fig. 5: fs-TA measurements of 7in CH_2Cl_2 ; 70 fs pump pulses at 360 nm.

- the ground state bleach between 520 and 650 nm is due to a ³MLCT state Ru(III)(bpy⁻) [5] and decreases with a time constant of 4.1 ps
- the species with a life time of 18 ps is assigned to the intermediate charge separated W(III)-ppy⁻ state



600

8.82 ps 13.32 ps

23.32 ps

- 59.57 ps - 155.32 ps

400

pulses at 490 nm.

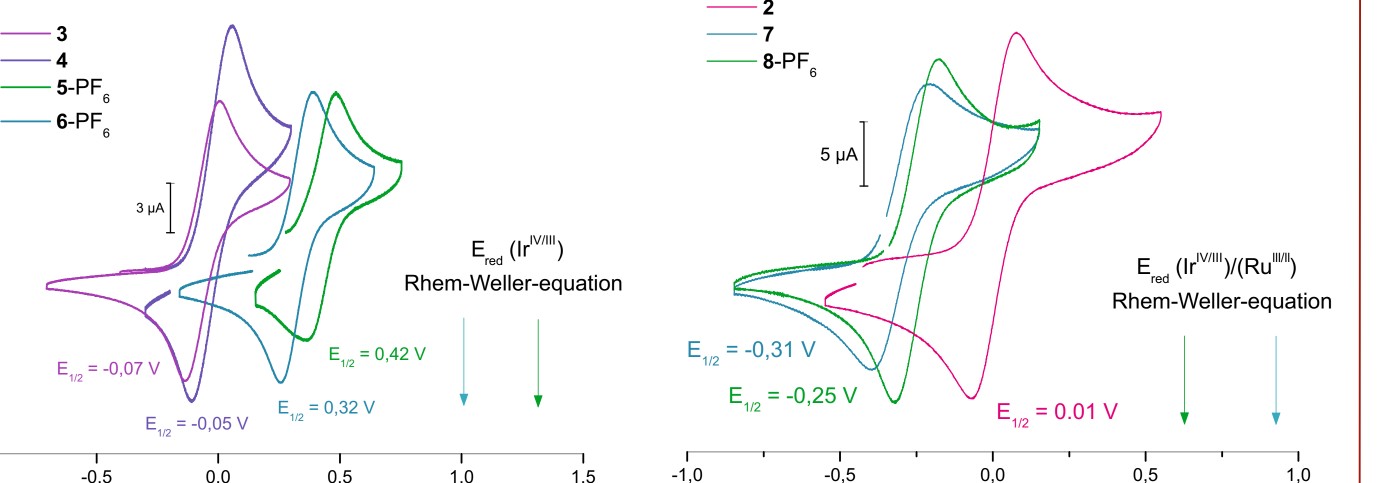
-1.0

- 857.52 ps

500

 λ / nm

Fig.6: fs-TA measurements of **8**-PF₆ in CH₃CN; 70 fs pump



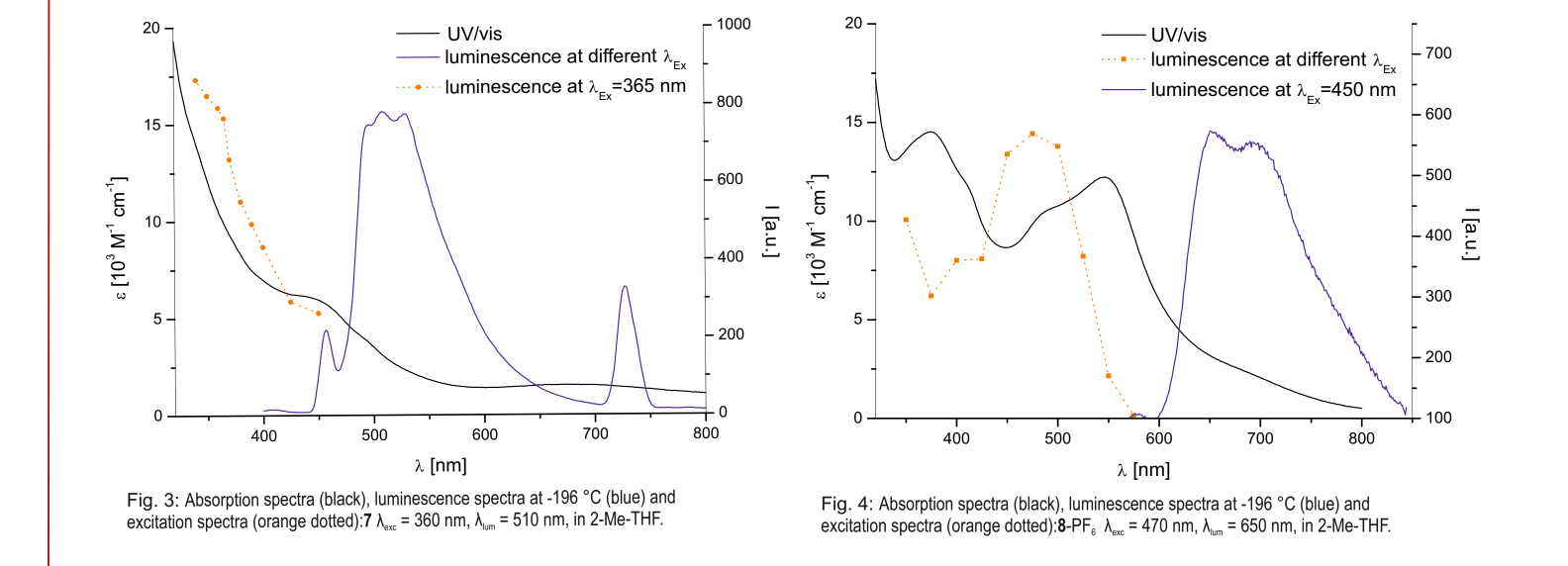




Fig.7: CV measurements of **2** - **8**-PF₆ in CH_2CI_2 vs. Fc/Fc⁺

- the coordination of the Ir(ppy)₂-moiety via the neutral donors leads to the formation of cationic complexes and a lower electron density at the tungsten centre, resulting in higher redox potentials for 5-PF₆ and 6-PF₆ of 0.42 and 0.32 V, respectively
- coordination to either the Ir(ppy)₂- or Ru(bpy)₂-moiety via the neutral pyridine- and the anionic carbon-donor leads to lower redox potentials compared to the mononuclear complex, indicating a higher electron density at the W(II) centre
- the estimated redox potentials for the photoexcited Ir(VI)- or Ru(III)-centre, via the Rehm-Weller equation reveals that such a transfer is certainly thermodynamically possible

[1] (a) J. Rüger, C. Timmermann, A. Villinger, W. W. Seidel, *Inorg. Chem.* 2019, *58*, 9270–9279;
(b) K. Helmdach, A. Villinger, W. W. Seidel, *Organometallics* 2017, *36*, 4844–4853.
[2] T. Suzuki, T. Kuchiyama, S. Kishi, S. Kaizaki, H. D. Takagi, M. Kato, *Inorg. Chem.* 2003, *42*, 785-795.

[3] K. Helmdach, S. Ludwig, A. Villinger, D. Hollmann, J. Kösters and W. W. Seidel, *Chem. Commun.*, **2017**, 53, 5894–5897.
[4] G. J. Hedley, A. Ruseckas, I. D. W. Samuel, *Chem. Phys. Lett.*, **2008**, 450, 292.
[5] Y. Na, J. Pan, M. Wang, L. Sun, *Inorg. Chem.* **2007**, *46*, 3813-3815.

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