Dalton **Transactions**

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Cite this: Dalton Trans., 2016, 45 17095

Received 12th September 2016, Accepted 1st October 2016

DOI: 10.1039/c6dt03545k

<www.rsc.org/dalton>

cis-Conformation of indigo in the coordination complex (indigo-O,O)(Cp*Cr^{II}Cl)⁺

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The interaction of decamethylchromocene ($Cp*_{2}Cr$) with indigo in the presence of a Cl[−] source yields the coordination complex (indigo-O,O)(Cp*Cr^{II}Cl) (1) in which one Cp* ligand at chromium is substituted by indigo. Indigo adopts an unusual cis-conformation in 1, allowing the coordination of both indigo carbonyl groups to one Cr^{II} center. Complex 1 contains Cr^{II} with an $S = 1$ spin state and indigo⁰. At the same time, calculations show that an excited ionic state is positioned close to the neutral ground state, providing the appearance of intense low-energy NIR bands in the spectrum of 1 at 820 and 1002 nm attributed to metal-to-ligand charge transfer. **COMMUNICATION**

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Indigo and its derivatives are well-known dyes now produced on a multi-ton scale.¹ Recently, these dyes have found new applications as electronic materials² and materials for rechargeable batteries.³ Indigo exists as a *trans*-isomer stabilized by intramolecular hydrogen C=O…H–N bonds.⁴ The existence of the cis-isomer of indigo was predicted a long time ago but until now this isomer of indigo has neither been isolated from natural sources nor been synthesized in a pure form most probably because of destabilization resulting from the repulsion of carbonyl dipoles. 5 The *cis*-isomer is observed only for some indigo derivatives and leuco indigo under photoisomerization.⁶ The properties of indigo can be affected not only by chemical modification^{2d} but also by the coordination

of transition metals.⁶ Such complexes are generally obtained for deprotonated indigo in monoanionic (indigoH−) and dianionic states (indigo^{2−}). In all these complexes, indigo also exhibits a *trans*-conformation.⁷ Only some "N-indigo" derivatives formed metal complexes in the cis -conformation $⁸$ but</sup> no information about the coordination complexes of unsubstituted cis-indigo has been found.

This work involved the study of the interaction of indigo with decamethylchromocene $(Cp^*{}_{2}Cr)$ and showed that indigo acts as a strong ligand able to substitute one of the pentamethylcyclopentadienyl ligands at chromium, forming the coordination complex (indigo-O,O)(Cp*CrCl) (1). Moreover, indigo adopts an unusual *cis*-conformation in this complex, thereby allowing the coordination of both carbonyl groups to the chromium center. Transformation of trans- to cis-indigo can be realized in the coordination sphere of a chromium atom.

Indigo shows weak acceptor properties and can be reduced to the radical anion at a redox potential of −0.75 V vs. Ag/AgCl (or -0.795 V vs. SCE).⁹ The reduction ability of Cp^{*}₂Cr, which has the first oxidation potential of -1.04 V vs. SCE,¹⁰ is enough to produce indigo⁻⁻. Indeed, indigo completely dissolves in o-dichlorobenzene following the interaction with $Cp_{2}^{*}Cr$ under strictly anaerobic conditions, and a red solution characteristic of indigo^{-−} (ref. 9) was formed. Slow mixing of the resultant solution with n -hexane did not yield any crystalline product. The addition of a stoichiometric amount of the $chloro(1,5-cyclooctadiene)$ rhodium (i) dimer, $\{Rh^{I}(cod)Cl\}_{2}$ (which was used to affect the crystallization), resulted in a color change of the solution from red to brown over several hours. Slow diffusion of n-hexane into this solution over the course of one month produced crystals as black parallelepipeds. The composition of these crystals determined from single crystal X-ray diffraction was (indigo-O,O)(Cp*CrCl) (1), i.e. (indigo-O,O) chloro-(η⁵-pentamethylcyclopentadienyl)chromium(II).

The crystal structure of 1 was studied at 150 K, \ddagger and the molecular structure of (indigo-O,O)(Cp*CrCl) is shown in Fig. 1a. The indigo molecule had a cis-conformation that allowed the coordination of both carbonyl groups to the chro-

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[†]Electronic supplementary information (ESI) available: Materials, synthesis, details of theoretical calculations and X-ray diffraction experiment, IR spectra and data of magnetic measurements. CCDC 1493779. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c6dt03545k

Fig. 1 (a) Molecular structure of the coordination complex (oxygen atoms are red, nitrogen atoms are blue, chloride anions are green, and chromium atoms are violet). Hydrogen atoms are not shown. (b) van der Waals contacts between the components are shown by green dashed lines.

mium center. The length of the O(indigo)–Cr bonds was 1.911(2)–1.923(2) Å. The length of the C=O bonds in 1 was 1.296(3) \AA . These bonds are elongated in comparison with those in trans-indigo (1.239(2) Å).^{4a} Previously, the elongation of the $C=O$ bonds of indigo has been observed for the coordinated carbonyl in *trans*-indigo-Pd(Bu₃P)Cl $(1.276(3)$ Å), which is significantly longer than the $C=O$ bond with a non-coordinated oxygen atom (1.225(3) Å).^{7a} The central C=C bond and the average C–N bond lengths in 1 were 1.400(4) and 1.392(3) Å, respectively, which were also elongated compared with those in *trans*-indigo (1.343(2) and 1.381(2) Å, respectively^{4a}). These effects can be attributed to the coordination of indigo to chromium. The length of the Cr–Cl bond was 2.3948 (7) Å and the average length of the Cr–C(Cp^{*}) bonds was 2.247 (2) Å. The chromium atom was positioned nearly in the plane of the indigo molecule (the displacement was only 0.020 Å). The *cis* conformation of the indigo molecule resulted in a nonparallel arrangement of the $C=O$ bonds with a dihedral angle of $14.62(2)$ ^o between them. The *cis*-isomer has a slightly twisted shape with a dihedral C(O)–C–C–C(O) angle of 7.0° and the displacement of the oxygen atoms from the 18-atom indigo plane by 0.158 and 0.302 Å in opposite directions. The distance between the hydrogen atoms attached to nitrogen was

only 2.146 Å in the cis-isomer, which is also rather short. These data indicate repulsion between the molecular permanent dipoles resulting from the orientation of the $C=O$ groups and intramolecular repulsion between the N–H hydrogen atoms in the cis-isomer.

The coordination units form $\{ (indigo-O,O)(Cp*CrCl) \}$ π-dimers in 1 (Fig. 1b) with a short interplanar distance of 3.44 Å and a parallel arrangement. As a result, multiple C, N⋯C,N van der Waals contacts are formed between indigo moieties (shown by green dashed lines in Fig. 1b). The Cp*CrCl groups are positioned on the opposite sides of the dimer and the shortest Cr⋯Cr distance was 8.01 Å. The π -dimers arranged in the layers were packed in such a way that the Cp^{*} group formed weak π -stacks with the indigo molecule of the neighbouring dimer (Fig. 1b).

The magnetic properties of 1 were analysed using EPR spectroscopy and the SQUID technique. Charge transfer (CT) produced Cr^{III} and indigo^{-−} species are generally well resolved in the EPR spectrum. The absence of any signal between room temperature and 4 K indicated the presence of an integer-spin system Cr^H that is EPR silent in X-band EPR.¹¹ Indeed, the complex had an effective magnetic moment of 2.91 μ_B at 300 K (Fig. 2), close to the magnetic moment calculated for a system containing one non-interacting $S = 1$ spin from Cr^{II} (2.83 μ_B). The reciprocal molar magnetic susceptibility of 1 was linear in the 10–300 K range, showing the Weiss temperature of −2 K (Fig. 2, inset). This result indicates weak antiferromagnetic coupling of the spins which can also explain the decrease in the magnetic moment of 1 below 12 K (Fig. 2). The distance between the Cr^{II} atoms in the π-dimers is rather long and, accordingly, diamagnetic indigo molecules cannot effectively mediate intermolecular magnetic coupling.

The trans-cis-isomerization of neutral indigo in the individual state requires 59.5 kcal mol⁻¹. The barrier for the isomerization can be reduced to 38.4 kcal mol⁻¹ under electron transfer. According to the calculations (see ESI, Fig. S1 and S2†), the central $C=C$ bond of *trans*-indigo is elongated from

Fig. 2 Temperature dependence of the effective magnetic moment of 1 in the 1.9–300 K range. The inset shows the temperature dependence of the reciprocal molar magnetic susceptibility of 1 in the same range.

1.368 Å in the neutral state to 1.388 Å in the radical anion state to provide easier isomerization of indigo^{•−}. The isomerization of trans-indigo through the enol form has an even lower activation barrier of 25 kcal mol⁻¹ (Fig. S3†).

The possible way for the formation of 1 is shown in Fig. 3. According to the calculations, the haptotropic shift of one Cp* ring in the outer sphere (trans-indigo'⁻⁻)(Cp^{*}₂Cr⁺) complex allows the formation of an inner sphere complex of indigo with one coordinated carbonyl group (Fig. 4a). This complex is unstable since it has slightly higher energy by 6.1 kcal mol⁻¹ than the energy of non-interacting *trans*-indigo and $\text{Cp*}_2\text{Cr}$. The geometry of the indigo ligand in this complex corresponds well to the geometry of indigo⁻⁻, and the spin density on the Cr atom is equal to 2.74. The trans–cis isomerization of the indigo ligand in the coordination sphere of the chromium atom leads to the formation of an additional Cr–O bond (Fig. 4b) under the η^1 -coordination of one Cp* ring and the 3.7 kcal mol⁻¹ energy release. The calculated activation energy of such isomerization is equal to 25.1 kcal mol⁻¹ (see the structure of the transition state in Fig. S4†). In contrast to the previous case, the $(cis\text{-indigo-O,O})(Cp*_{2}Cr)$ complex is stable with respect to the non-interacting *cis*-indigo and $Cp^*{}_{2}Cr$, the energy gain being 14.2 kcal mol⁻¹. We suppose that since only *cis*-indigo can form a stable complex with $Cp^*_{2}Cr$, namely this isomer is stabilized in the complex with chromium. A possible scenario of the formation of 1 is the substitution of the η^1 -Cp^{*} ligand by chloride anions in the intermediate (cis-indigo-O,O) $(Cp^*{}_2$ Cr) complex. The source of chloride anions is probably $\{Rh^{I}(cod)Cl\}_{2}$ also used in the synthesis (Fig. 3). The η^{5} -coordination Cp^* ligand on the Rh^I center is energy favorable and may be the driving force of the substitution reaction. According to the calculations and magnetic data, 1 has a neutral indigo ligand. Thus, initially the ionic (trans-indigo^{-−}) $(Cp^*{}_2Cr^+)$ pair transforms to neutral complex 1 under the isomerization of *trans*-indigo and the exchange of the donor Cp^{*−} ligand on the chloride anion. **Columbial Consequent Commons Access Article and Consequent Commons Article Commons Article Commons Article is an access Article is article. The isometric common Distribution of 1 as however (commons are also the common D**

The IR spectrum of 1 is shown in Fig. S5† and the main peaks are listed in Table S1.† Coordination resulted in a decrease in the intensity of the bands attributed to the $C=O$ vibrations and their shift from 1613 and 1627 to 1607 and 1615 cm−¹ , respectively. The bands associated with the N–H

 $Cp_{2}^{*}Cr$

 ${({\rm COD})}$ RhCl}₂

 $(\eta^5$ -Cp*)(COD)Rh

 \overrightarrow{H} o \overrightarrow{G}
trans-indigo radical anion

lonic pair (-6.7 kcal/mol)

activation

energy
25.1

kcal/mol

 $Cp^*_{2}Cr$

ő 6.1 kcal/mol

-Cp*)CrCl \overrightarrow{u}

Complex 1 of cis-indigo

Fig. 3 Possible route for the formation of 1, calculated relative energies are given with respect to the non-interacting $Cp[*]₂Cr$ and trans- or cisindigo (in parenthesis).

Fig. 4 (a) The calculated structure of $[(trans-indigo-O)(Cp*₂Cr)]$; (a) and [(cis-indigo-O,O)(Cp*₂Cr)]; (b) the interatomic distances are shown in Å. H atoms of C–H bonds are omitted for clarity.

vibrations in trans-indigo were observed at 3250 and 3270 cm−¹ (Fig. S6a†). These bands were found in the spectrum of 1 at 3236 and 3295 cm⁻¹ (Fig. S6b†), indicating the preservation of hydrogen at the nitrogen atoms. At the same time the essential shift of these bands can be attributed to the disappearance of the hydrogen $C=O \cdots H-N$ bonds in the *cis*conformation of indigo. The solid state spectra of indigo and 1 in the UV–visible–NIR range are shown in Fig. 5. Indigo exhibited bands at 334 and 291 nm and the lowest energy band is observed at 665 nm (the S_0-S_1 transition). The spectrum of 1 contained many bands with absorption up to 1250 nm. The bands of trans-indigo at 291 and 334 nm were reproduced in

Fig. 5 Spectrum of indigo and 1 in the UV–visible–NIR range recorded in KBr pellets. The pellet for 1 was prepared under anaerobic conditions. The inset shows the theoretical spectrum calculated for the optimized structure of 1.

 $Cn^*_{2}C$

Ä Ĥ

Coordinated cis-indigo

2.4 kcal/mol (-14.2 kcal/mol)

 $(\eta^1 - Cp^*)(\eta^5 - Cp^*)Cp$

trans-indigo

Isomerization

and coordina-

tion of two C=O groups

the spectrum of 1 at 285 and 342 nm, respectively, and a new band is manifested at 472 nm (Fig. 5). An intense band at 578 nm and weaker bands at 667 and 708 nm are also manifested in the spectrum of 1 (Fig. 5). The most interesting feature of the spectrum of 1 was the appearance of two relatively intense low-energy absorption bands with maxima at 820 and 1002 nm (Fig. 5). To understand the origin of these bands, the theoretical spectrum of 1 (Fig. 5, inset) was calculated using the TD-DFT approach based on the optimized structure (see the ESI†). The calculated spectrum showed a general resemblance to the experimental spectrum. According to the calculations, complex 1 with a triplet ground state (Cr^{II}) , indigo 0) has a low-lying excited ionic state located 1480 cm^{-1} above the ground state. The spin density on the Cr atom in this state is 2.75, corresponding to three unpaired electrons on chromium and an additional unpaired electron that was located on the indigo ligand, *i.e.* an excited quintet magnetic state is formed with the parallel orientation of the Cr^{III} $(S = 3/2)$ and indigo⁻⁻ $(S = 1/2)$ spins. The corresponding triplet CT state with the antiparallel orientation of the spins should have a higher energy because of the effects of interactions with the ground triplet state. Thus, a spin-allowed triplet–triplet transition is expected to be observable in the NIR range of the spectrum of 1 as metal-to-ligand CT. The upper d-levels of the Cr complex have low splitting energies of 0.11–0.35 eV. This leads to the appearance of several CT transitions, namely NIR bands at 1066 and 1170 nm, located in the 800–1250 nm range of the experimental spectrum. Communication

The spectrum of 1 at 285 and 342 nm, respectively, and a now in the spectrum of 1. The distantivelation in the neural on an excell on the neural on 2016. The commons are common and the neural on the neural

Most probably, the intense band at 578 nm corresponds to the S_0-S_1 transition in *cis*-indigo. It is noticeably blue shifted in comparison with the spectrum of trans-indigo (665 nm). The reasons for this can be *trans–cis* isomerization and/or deviation of the molecule from planarity in 1. It is known that the position of the lowest energy band in the indigo dyes is very sensitive to the conformation of the molecule and can be shifted to the blue side by 50 nm or more on trans–cis isomerization and/or deviation of the molecule from planarity.¹² Indeed, the calculated S_0-S_1 transition in *trans*-indigo at 621 nm is shifted to 526 nm for cis-indigo or to 591 nm for the 'frozen' cis-indigo ligand of 1.

In conclusion, the preparation of coordination complex through the radical anions of indigo allows, for the first time, the observation of the cis-isomer of indigo, whose molecular structure is unknown yet. Up to now the cis-conformation has been observed only for some indigo derivatives.⁶ cis-Indigo coordinates to chromium through two carbonyl groups in 1 and this coordination motif stabilizes the cis-isomer in spite of the fact that this isomer is less stable than trans-indigo. It should be noted that all previously studied transition metal complexes of indigo contain deprotonated trans-indigo anions.7 Some "N-indigo" derivatives can form metal complexes in both *trans*- and *cis*-conformations.⁸ Complex 1 is the first structurally characterized coordination complex of neutral cis-indigo. Though complex 1 has a neutral ground state, an excited charge transition state is positioned unusually close to provide the appearance of new intense NIR absorption bands

in the spectrum of 1. The absorption of light in the NIR range can transfer this complex from the neutral to an excited ionic state which is characterized by a high-spin Cr^{III} magnetic state. We believe that such complexes are promising materials for detectors and switchers operating in the NIR range. The range of transition metal complexes of cis-indigo can be extended by using indigo derivatives and by applying other metallocenes instead of $Cp_{2}^{*}Cr$. This work is now in progress.

This work was supported by JSPS KAKENHI grant numbers JP23225005 and JP26288035 and was partially supported by the JST(ACCEL)27(100150500010) project.

Notes and references

 \ddagger Crystal data for 1: C₂₆H₂₅ClCrN₂O₂, F.W. 484.93, black parallelepiped, 0.38 \times 0.12×0.01 mm³, 150.0(2) K, monoclinic, space group $P2_1/n$, $a = 11.7879(7)$, $b =$ 15.2593(16), $c = 12.5982(8)$ Å, $\beta = 94.897(7)$ °, $V = 2257.8(3)$ Å³, $Z = 4$, $d_{\text{caled}} =$ 1.427 g cm⁻³, μ = 0.651 mm⁻¹, $F(000)$ = 1008, $2\theta_{\text{max}}$ = 65.390°; 26 768 reflections collected, 7588 independent; $R_1 = 0.0548$ for 4721 observed data $[>2\sigma(F)]$ with 294 parameters; $wR_2 = 0.1442$ (all data); final G.o.F. = 1.023. CCDC 1493779.

- 1 (a) C. J. Cooksey, Biotech. Histochem., 2007, 82, 105; (b) N. Stasiak, W. Kukuła-Koch and K. Głowniak, Acta Pol. Pharm., 2014, 71, 215.
- 2 (a) M. Irimia-Vlada, E. D. Głowacki, P. A. Troshin, G. Schwabegger, L. Leonat, D. K. Susarova, O. Krystal, M. Ullah, Y. Kanbur, M. A. Bodea, V. F. Razumov, H. Sitter, S. Bauer and N. S. Sariciftci, Adv. Mater., 2012, 24, 375; (b) E. D. Głowacki, G. Voss and N. S. Sariciftci, Adv. Mater., 2013, 25, 6783; (c) O. Pitayatanakul, T. Higashino, T. Kadoya, M. Tanaka, H. Kojima, M. Ashizawa, T. Kawamoto, H. Matsumoto, K. Ishikawa and T. Mori, J. Mater. Chem. C, 2014, 2, 9311; (d) V. Klimovich, L. I. Leshanskaya, S. I. Troyanov, D. V. Anokhin, D. V. Novikov, A. A. Piryazev, D. A. Ivanov, N. N. Dremova and P. A. Troshin, J. Mater. Chem. C, 2014, 2, 7621.
- 3 (a) M. Yao, K. Kuratani, T. Kojima, N. Takeichi, H. Senoh and T. Kiyobayashi, Sci. Rep., 2014, 4, 3650; (b) M. Yao, H. Sano, H. Ando and T. Kiyobayashi, Sci. Rep., 2015, 5, 10962.
- 4 (a) P. Süsse, M. Steins and V. Kupcik, Z. Kristallogr., 1988, 184, 269; (b) F. Kettner, L. J. Schäfer, K. Röder, U. Purgahn and H. Krautscheid, Acta Crystallogr., Sect. E: Struct. Rep. Online, 2011, 67, o2867.
- 5 B. T. Golding and C. Pierpoint, Educ. Chem., 1986, 23, 71.
- 6 (a) E. C. Nicholls-Allison, G. Nawn, B. O. Patrick and R. G. Hicks, Chem. Commun., 2015, 51, 12482; (b) D. Farka, M. Scharber, E. D. Głowacki and N. S. Sariciftci, J. Phys. Chem. A, 2015, 119, 3563; (c) R. Rondão, J. Seixas de Melo, M. J. Melo and A. J. Parola, J. Phys. Chem. A, 2012, 116, 2826.
- 7 (a) W. Beck, C. Schmidt, R. Wienold, M. Steinmann and B. Wagner, Angew. Chem., Int. Ed. Engl., 1989, 28, 1529; (b) A. Lenz, C. Schmidt, A. Lehmann, B. Wagner and W. Beck, Z. Naturforsch., B: Chem. Sci., 1997, 52, 474; (c) J. Y. Wu, C. H. Chang, P. Thanasekaran, C. C. Tsai, T. W. Tseng, G. H. Lee, S. M. Peng and K. L. Lu, Dalton

Trans., 2008, 6110; (d) P. Mondal, M. Chatterjee, A. Paretzki, K. Beyer, W. Kaim and G. K. Lahiri, Inorg. Chem., 2016, 55, 3105.

- 8 (a) G. Nawn, K. M. Waldie, S. R. Oakley, B. D. Peters, D. Mandel, B. O. Patrick, R. McDonald and R. G. Hicks, Inorg. Chem., 2011, 50, 9826; (b) P. Mondal, S. Plebst, R. Ray, S. M. Mobin, W. Kaim and G. K. Lahiri, Inorg. Chem., 2014, 53, 9348. Oatton Transactions

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	- 9 A. Roessler, D. Crettenand, O. Dossenbach, W. Marte and P. Rys, Electrochim. Acta, 2002, 47, 1989.
	- 10 J. L. Robbins, N. Edelstein, B. Spencer and J. C. Smart, J. Am. Chem. Soc., 1982, 104, 1882.
	- 11 J. Telser, L. A. Pardi, J. Krzystek and L. C. Brunel, Inorg. Chem., 1998, 37, 5769.
	- 12 W. Lüttke, H. Hermann and M. Klessinger, Angew. Chem., Int. Ed. Engl., 1966, 5, 598.