ChemComm



COMMUNICATION

View Article Online



Cite this: Chem. Commun., 2021, **57** 1320

Received 9th October 2020 Accepted 27th November 2020

DOI: 10.1039/d0cc06755e

rsc.li/chemcomm

Rapid build up of nanooptomechanical transduction in single crystals of a ruthenium-based SO₂ linkage photoisomer†

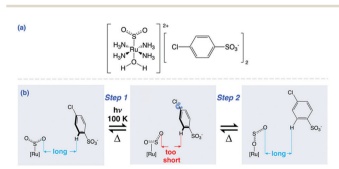
Jacqueline M. Cole, *\oldsymbol{0}** \text{abcd} David J. Gosztola, *\oldsymbol{0}* Jose de J. Velazquez-Garcia, *\oldsymbol{0}** a SuYin Grass Wang oe and Yu-Sheng Chene

Single-crystal nanooptomechanical transduction occurs in [Ru(SO₂) (NH₃)₄(H₂O)]chlorobenzenesulfonate₂, reaching maximal levels within 40 s at 100 K when photostimulated by 505 nm light. Its in situ lightinduced crystal structure reveals the molecular origins of this optical actuation: 26.0(3)% of the η^1 -SO₂ ligand photoconverts into an η^1 -OSO photoisomer which, in turn, induces a 49.6(9)° arene ring rotation in its neighbouring counter ion.

Materials that display optical switching in crystalline media are of interest for a range of solid-state applications, from optical sensing¹ to optical data storage.2 A sub-class of optical switches exhibit single-crystal nanooptomechanical transduction, whereby a lightdriven switching process in one molecule or ion stimulates mechanical motion in a neighbouring molecule or ion. Nanooptomechanical transducers may be employed in light-driven molecular machinery³⁻⁵ or nanotechnology;⁶ when observed in single-crystal form, they have prospective use in futuristic circuitry for quantum computers.⁷ Such functionality in materials is rare, especially in single crystals, which offer a high-quality solid-state medium for single-photon control. Only a few materials have demonstrated such function;⁸⁻¹³ this includes a family of ruthenium-based complexes that undergo SO₂-linkage photoisomerisation in single crystals.8-11 The generic formula for these complexes is trans- $[Ru(SO_2)(NH_3)_4X]^{m+}$ Y_n where X is the ligand that lies trans to the SO₂-linkage isomer and Y is the counter ion, with the integer values

This paper reports the discovery of a new nanooptomechanical transducer in this family of ruthenium-SO₂ based complexes: trans-[Ru(SO₂)(NH₃)₄(H₂O)]chlorobenzenesulfonate₂ (1). Scheme 1 presupposes its two-step operational mechanism based on our previous work.8-11 This is confirmed by its light-induced crystal structure which is revealed by in situ light-induced single-crystal X-ray diffraction, a recently developed technique that has become known as photocrystallography. 14-17 Concerted singlecrystal optical absorption spectroscopy and microscopy is then employed to determine the visible absorption characteristics of a single crystal of 1 as well as reveal its anticipated photochromic effects. Kinetic aspects of this optical metrology will reveal that

[†] Electronic supplementary information (ESI) available. CCDC 2035266 and 2035267. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/d0cc06755e



Scheme 1 (a) Dark-state chemical diagram of 1; (b) η^1 -SO₂ to η^1 -OSO photoisomerisation in 1 which induces nanooptomechanical transduction via arene ring rotation to reduce interionic strain

of m and n being dictated by the charge-balancing requirements of X and Y. Certain combinations of X and Y afford nanooptomechanical transduction.⁸⁻¹¹ A concerted two-step molecular mechanism is responsible for this functionality, whereby an S-bound η^1 -SO₂ ligand photoisomerises to an O-bound \(\eta^1\)-OSO photoisomer. The non-bound oxygen in this photoisomer protrudes from the cation within the crystal lattice, in a direction such that it lies uncomfortably close to one of the counter ions. Pending the anion contains an arene ring, it rotates in order to alleviate the crystal-lattice strain that results from the close proximity of the η^1 -OSO ligand.

^a Cavendish Laboratory, Department of Physics, University of Cambridge,

J. J. Thomson Avenue, Cambridge, CB3 0HE, UK. E-mail: jmc61@cam.ac.uk

^b ISIS Neutron and Muon Facility, STFC Rutherford Appleton Laboratory, Harwell Science and Innovation Campus, Didcot, OX11 0QX, UK

^c Department of Chemical Engineering and Biotechnology, University of Cambridge, West Cambridge Site, Philippa Fawcett Drive, Cambridge, CB3 0AS, UK

^d Argonne National Laboratory, 9700 South Cass Avenue, Lemont, Illinois 60439,

^e NSF's ChemMatCARS Beamline, The University of Chicago, Advanced Photon Source, Lemont, Illinois 60439, USA

the nanooptomechanical transduction in 1 builds up macroscopically very rapidly.

Communication

1 was synthesised from trans-[Ru(SO₂)(NH₃)₄Cl]Cl, which was prepared according to a literature procedure. 18 5 mg (16 mmol) of this precursor was dissolved in a mixture of Na₂CO₃ (1 M, 350 mL) and methanol (350 mL), into which a distilled water/ methanol (1/1, v/v) solution of p-chlorobenzenesulfonic acid (100 mL, 2 M; > 90% purity, Sigma Aldrich) was added dropwise. Red plate-like crystals precipitated from solution which were collected by filtration and washed with methanol.

The light-induced crystal structure of 1 was determined using synchrotron-based X-ray photocrystallography at the ChemMatCARS beamline of the Advanced Photon Source, Argonne National Laboratory, IL, USA. See ESI† for full crystallography details; just the relevant photostimulation procedure is given here. A reference dataset for the dark-state crystal structure was first obtained from a 50 \times 25 \times 25 μm^3 single crystal. A 505 nm light (Thorlabs M505L3 LED) was then shone onto the crystal for 2.5 h while it was held at 100 K on the diffractometer. This light was switched off before acquiring data for the light-induced crystal structure. For control purposes, ambient lighting in the experimental hutch of the beamline was extinguished while all data were being collected.

The 505 nm light-induced crystal structure of 1 is shown in Fig. 1. 26.0(3)% of its dark-state η^{1} -SO₂ ligand has photoconverted into an O-bound η^1 -OSO photoisomer. 22.4(5)% of the arene ring in one of the chlorobenzenesulfonate counter ions has rotated by 49.6(9)° in response to the formation of the photoisomer. This is the nanooptomechanical transduction. The dark-state structure remains the dominant residue, as is witnessed in the majority of cases for this family of complexes. The clean conversion from the η^1 -SO₂ isomer to the η^1 -OSO photoisomer is notable since η^{1} -OSO formation in this series of complexes is generally accompanied by a discernible fraction of the side-bound η^2 -(OS)O photoisomer.^{8–10} To this end, there is only one other reported instance of a clean conversion which

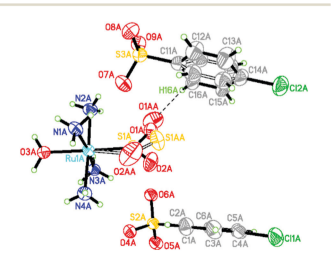


Fig. 1 Light-induced crystal structure of 1. Disordered species have hollow bonds. The close O1AA···H16A contact (2.265(7) Å) between the η^{1} -OSO ligand and arene ring has a dashed line

occurs in trans-[Ru(SO₂)(NH₃)₄(3-phenylpyridine)]Cl₂·H₂O where 52(3)% of the η^1 -OSO photoisomer is shown to form. ¹⁹ Its small counter ions render it to behave as an optical switch. Thus, 1 is the only reported nanooptomechanical transducer whose η¹-OSO photoisomer is not polluted by another type of photoisomer. The practical ramifications of such a clean conversion could ultimately lead 1 to be more favourable than other complexes in prospective applications.

The bond geometry of the light-induced structure of 1 at 100 K is very similar to that of the 90 K structure of trans-[Ru(SO₂)(NH₃)₄(H₂O)]tosylate₂, ²⁰ aside from the SO₂ and an arene ring in one of the counter ions which are affected by nanooptomechanical transduction in 1. The SO₂ ligand in trans- $[Ru(SO_2)(NH_3)_4(H_2O)]$ tosylate₂ photoconverts into an η^2 -(OS)O photoisomer at 100 K, and thus behaves as an optical switch rather than a transducer. The two complexes chemically differ by only one substituent: a Cl rather than methyl substituent in each counter ion. This structural comparison illustrates nicely how a subtle detail of the counter ions can completely alter the optical properties in this series of complexes.

Despite taking all of the usual experimental precautions that are usually successful in our photocrystallography work, the reference dark-state crystal structure of 1 was polluted by some photoinduced species. Thereby, the 'dark-state' structure displayed 12.0(3)% of the η^{1} -OSO photoisomer and 17.6(4)% of a corresponding arene ring disorder which displays 48.7(9)° rotation. One expects a pure η^1 -SO₂ configuration and ordered arene ring in its true dark state, judging from all other materials in this family of complexes. This expectation is at least corroborated by the much lower photoconversion fractions found in this 'dark' structure compared to the deliberately intended light-induced structure of 1.

On the face of it, the light pollution is curious since only ambient light from the experimental hutch of the synchrotron beamline would have impinged upon the sample, once it had been cooled to 100 K where it might be susceptible to photoisomerisation; and even then, the crystal was only exposed to this ambient light for the few minutes that was required to centre the crystal on the diffractometer. The ambient lighting had been switched off for the entire data collection, as per a standard control measure that is employed in our photocrystallography experiments. A pure dark-state structure could not be obtained despite best efforts. This was unusual. So, a series of concerted single-crystal optical absorption spectroscopy and microscopy experiments on this same crystal of 1 were employed in order to understand why.

Single-crystal optical absorption spectroscopy and microscopy were performed on 1, using the custom-built microspectroscopy system described by Cole and co-workers. 19,21 For micro-spectroscopy details, see ESI.† The crystal was photosimulated by the 505 nm LED source at 100 K. The 505 nm light-excitation power measured at the 20x, 0.3 NA objective (Olympus UMPlanFL) was 360 μW, giving an estimated 261 μ W mm⁻² illuminating the field of view.

Single-crystal optical absorption spectra of 1, acquired after t minutes of light exposure at 100 K, are shown in Fig. 2. These ChemComm Communication

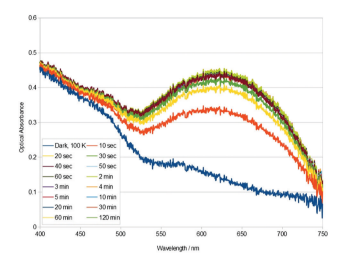


Fig. 2 Single-crystal optical absorption spectra of 1, acquired after t minutes of 505 nm light exposure at 100 K. A reference dark-state spectrum of 1 at 100 K is provided.

results reveal that the photoisomeric fraction in 1 build up very fast: the crystal essentially attains its maximum optical absorbance after only 40 s of 505 nm light exposure. This compares with at least several hours of light exposure that is normally required to maximise photoconversion levels in these types of complexes. 19,21 The build up of nanooptomechanical transduction in 1 will likewise be very fast, judging from the highly correlated percentage growth of the n¹-OSO species and that of the rotated arene ring; and given that the arene ring rotates as a direct consequence of η^1 -OSO formation.⁸ The rapid build up of nanooptomechanical transduction explains our inability to obtain a pure dark-state crystal structure of 1.

The rate constant associated with the photoisomerisation process in 1 was calculated using the light-induced area under each curve in Fig. 2 as a proxy metric for the concentration of photoisomer species. The calculation proceeded in four steps, described in the ESI.† This yielded a rate constant, $k = 7.90 \times 10^{-3} \text{ s}^{-1}$, for the η^1 -SO₂ (dark) to η^1 -OSO (light) photoisomerisation process in 1, from a linear fit whose coefficient of determination, $R^2 = 0.991$. This is the first time that k has been evaluated for the photoisomerisation process in this family of complexes; yet, we have observed qualitatively that it is a faster process than other complexes in the series. $^{8-10}$

Metal-ligand charge-transfer (MLCT) effects tend to dominate the optical absorption of ruthenium-based coordination complexes. $\hat{2}^{22-24}$ Changes in the optical absorption profile of 1 in response to light exposure (Fig. 2) are therefore expected to be indicative of MLCT bands that are influenced by n¹-OSO and dark η^1 -SO₂ isomers, and the arene ring rotation, given the lightinduced crystal structure of 1. The light-induced rise of the broad band in the 525-750 nm spectral region of 1 (Fig. 2) is echoed in the optical absorption spectral changes observed for other complexes in this family of materials, in those that form η^2 -(OS)O or η¹-OSO photoisomers. 19,21 However, this rise in optical absorption is followed by a rise of additional bands in these other complexes of this family of materials. Crystals of such complexes

that display η²-(OS)O photoisomerisation adopt almost 'blackbody' absorption within the visible spectrum, upon external light exposure.²¹ In contrast, the rise in optical absorption of 1 ceases after the formation of the broad band shown in Fig. 2. trans-[Ru(SO₂)(NH₃)₄(3-phenylpyridine)]Cl₂·H₂O represents the only previous report of a η^1 -SO₂ to η^1 -OSO photoisomerisation that is not accompanied by \(\eta^2\)-(OS)O formation. 19 Its changes in optical absorption in response to 505 nm light exposure are broadly similar to those of 1, at least within the part of the crystal that lies under the surface that is shown to peel macroscopically; yet, this peeling effect in that crystal, its twinned nature, and its lack of nanooptomechanical transduction behaviour, render difficult a like-for-like comparison of its optical absorption spectra with those of 1. Fig. 2 would therefore seem to provide a representative optical absorption profile for a complex in this series of materials that exclusively exhibits the \(\eta^1 \)-OSO photoisomer together with nanooptomechanical transduction.

Optical microscopy images that accompany these optical spectroscopy findings show that the crystal of 1 is photochromic, transforming from an orange to bluey-grey colour over the duration of 505 nm light exposure (see Video, ESI†). These photochromic effects corroborate the optical absorption spectral profiles observed.

The light-induced structure of 1 thermally decays back to its dark state; its decay rate was too fast to measure reliably with our instrument set up, although it was observed that full decay occurred within 2 minutes at T = 160 K.

Given the rapid build up of nanooptomechanical transduction in 1 at 100 K, even in the ambient light conditions of the experimental hutch of the synchrotron for just a few minutes, the optical absorption spectral characteristics of 1 were explored using two additional wavelengths of external light stimulation: 405 nm and 455 nm. Single-crystal optical absorption spectra of 1 acquired after t minutes of 405 nm light exposure at 100 K are displayed in Fig. 3. The same rise in

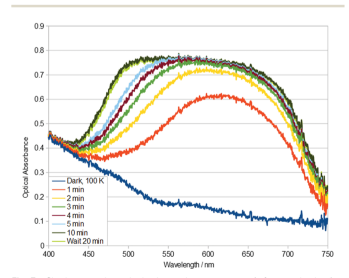


Fig. 3 Single-crystal optical absorption spectra of 1, acquired after t minutes of 405 nm light exposure at 100 K. A reference dark-state spectrum of 1 at 100 K is given.

Communication ChemComm

optical absorption within the 525-750 nm spectral region is observed for 1 upon 405 nm and 505 nm light stimulation. Yet, this rise is slower with 405 nm light activation, and then a new charge-transfer (CT) band opens up in its 450-550 nm region, yielding almost 'black-body' absorption in the visible spectrum. 455 nm light stimulus also exhibits this CT band (see ESI†).

This 'black-body' like absorption is seen in members of this family of complexes that exhibit η^2 -(OS)O photoisomers.²¹ This suggests that the growth of this low-wavelength CT band could be MLCT that arises from η^2 -(OS)O photoisomer formation. Indeed, n¹-OSO photoisomers usually form together with a discernible fraction of η^2 -(OS)O photoisomers. The 405 nm and 455 nm stimulated photoisomers thermally decay to their dark state at 160 K (see ESI†).

Rate constants could not be determined for the 405 nm stimulated photoisomerisation or thermal decay of 1, because the process was not first order over the full data range. However, plots of temperature *versus* ln k were approximately linear for two disparate sets of data, before and after the onset or deterioration of the new CT band, respectively. cf. $k = 3.174 \times 10^{-3} \text{ s}^{-1} (R^2 = 0.936)$ or $k = 6.586 \times 10^{-4} \text{ s}^{-1}$ ($R^2 = 0.984$) for t = 0-3 min or t = 3-5 min of 405 nm light exposure; while $k = 3.398 \times 10^{-3} \text{ s}^{-1} (R^2 = 0.952)$ or $k = 7.838 \times 10^{-3} \text{ s}^{-1} (R^2 = 1.000) \text{ for } t = 0-6 \text{ min or } t = 8-12 \text{ min time}$ lapse after the crystal of 1 had reached 160 K, respectively. These bifurcated results further corroborate the notion that a new CT band opens up in the 405 nm data, while it is absent in the 505 nm data. The 455 nm data behave similarly to the 405 nm data (see ESI†).

The rate constant for 505 nm light exposure of 1 at 100 K is faster that that at 405 nm. Its photoisomerisation process will thus dictate the nature of the 'dark' structure that resulted from modest exposure to ambient light conditions at 100 K.

In summary, nanooptomechanical transduction from 505 nm light activation of a single crystal of 1 has been showcased in this work. Its rapid build up in single crystals of 1 may have practical implications for its ultimate prospects in solid-state optical device technology. Further work is required to confirm the nature of the extra CT band that is seen with 405 nm or 455 nm light stimulation. Pending confirmation that it is the anticipated η^2 -(OS)O photoisomer, it is exciting to consider the prospect that η^{1} -OSO and η^{2} -(OS)O photoisomerisation could be alternated in the same crystal using different light sources.

J. M. C. is grateful for the BASF/Royal Academy of Engineering Research Chair in Data-Driven Molecular Engineering of Functional Materials, which is partly supported by the Science and Technology Facilities Council via the ISIS Neutron and Muon Facility. J. M. C. also thanks the 1851 Royal Commission of the Great Exhibition for the 2014 Fellowship in Design, hosted by Argonne National Laboratory where work done was supported by the U.S. Department of Energy (DOE) Office of Science, Office of Basic Energy Sciences, and used research resources of the Center of Nanoscale Materials and the Advanced Photon Source (APS), Office of Science User Facilities operated for the DOE Office of Science by Argonne National Laboratory, supported by the U.S. DOE, all under contract no. DE-AC02-06CH11357. Within the APS, the NSF's ChemMatCARS Sector 15 is supported by the Divisions of Chemistry (CHE) and Materials Research (DMR), National Science Foundation (NSF), under grant no. NSF/CHE-1346572 and NSF/CHE-1834750. Use of the PILATUS 3X CdTe 1M detector is supported by the NSF under grant no. NSF/DMR-1531283. J. d. J. V.-G. acknowledges the National Council of Science and Technology of Mexico (CONACyT) and the Cambridge Trust for a PhD Scholarship (217553).

Conflicts of interest

There are no conflicts to declare.

Notes and references

- 1 T. Liu, F. Pagliano, R. van Veldhoven, V. Pogoretskiy, Y. Jiao and A. Fiore, Nat. Commun., 2020, 11, 2407.
- 2 M. Imlau, Th. Woike, R. Schieder and R. A. Rupp, Europhys. Lett., 2001, 53, 471-477.
- 3 N. Koumura, R. Zijlstra, R. van Delden, N. Harada and B. L. Feringa, Nature, 1999, 401, 152-155,
- 4 N. Ruangsupapichat, M. M. Pollard, S. R. Harutyunyan and B. L. Feringa, Nat. Chem., 2011, 3, 53-60.
- 5 J. M. Abendroth, O. S. Bushuyev, P. S. Weiss and C. J. Barrett, ACS Nano, 2015, 9, 7746-7768.
- 6 L. Midolo, A. Schliesser and A. Fiore, Nat. Nanotechnol., 2018, 13,
- 7 J. Bochmann, A. Vainsencher, D. D. Awschalom and A. N. Cleland, Nat. Phys., 2013, 9, 712-716.
- 8 S. O. Sylvester and J. M. Cole, Adv. Mater., 2013, 25, 3324-3328.
- 9 S. O. Sylvester and J. M. Cole, J. Phys. Chem. Lett., 2013, 4, 3221-3226.
- 10 S. O. Sylvester, J. M. Cole, P. G. Waddell, H. Nowell and C. Wilson, J. Phys. Chem. C, 2014, 118, 16003-16010.
- 11 K. T. Mukaddem, J. M. Cole, K. A. Beyer and S. O. Sylvester, J. Phys. Chem. C, 2020, 124, 10094-10104.
- 12 F. Tong, W. Xu, T. Guo, B. F. Lui, R. C. Hayward, P. Palffy- Muhoray, R. O. Al-Kaysi and C. J. Bardeen, J. Mater. Chem. C, 2020, 8,
- 13 R. O. Al-Kaysi, F. Tong, M. Al-Haidar, L. Zhu and C. J. Bardeen, Chem. Commun., 2017, 53, 2622-2625.
- 14 P. Coppens, D. V. Fomitchev, M. D. Carducci and K. Culp, J. Chem. Soc., Dalton Trans., 1998, 6, 865-872.
- 15 J. M. Cole, Chem. Soc. Rev., 2004, 33, 501-513.
- 16 J. M. Cole, Acta Crystallogr., Sect. A: Found. Crystallogr., 2008, 64, 259-271.
- 17 J. M. Cole, Analyst, 2011, 136, 448-455.
- 18 L. H. Vogt, J. L. Katz and S. E. Wiberley, Inorg. Chem., 1965, 4, 1157-1163.
- 19 J. M. Cole, J. d. J. Velazquez-Garcia, D. J. Gosztola, S.-Y. G. Wang and Y.-S. Chen, Chem. Mater., 2019, 31, 4927–4935.
- 20 A. Y. Kovalevsky, K. A. Bagley and P. Coppens, J. Am. Chem. Soc., 2002, 124, 9241-9248.
- J. M. Cole, J. d. J. Velazquez-Garcia, D. J. Gosztola, S.-Y. G. Wang and Y.-S. Chen, Inorg. Chem., 2018, 57, 2673-2677.
- 22 E. Tfouni, Coord. Chem. Rev., 2000, 196, 281-305.
- 23 R. S. Lumpkin, E. M. Kober, L. A. Worl, Z. Murtaza and T. J. Meyer, J. Phys. Chem., 1990, 94, 239-243.
- 24 J. J. Rack, Coord. Chem. Rev., 2000, 253, 78-85.