



Cite this: *CrystEngComm*, 2018, **20**, 5990

## Understanding solid-state photoswitching in $[\text{Re}(\text{OMe}_2\text{-bpy})(\text{CO})_3(\eta^1\text{-NO}_2)]$ crystals *via in situ* photococrystallography<sup>†</sup>

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Single-crystal-to-single-crystal nitro  $\rightarrow$  nitrito isomerism is reported for the novel rhenium(I)-bipyridine complex  $[\text{Re}(\text{OMe}_2\text{-bpy})(\text{CO})_3(\eta^1\text{-NO}_2)]$ , achieving a maximum conversion of 66% to a photoinduced nitrito-( $\eta^1\text{-ONO}$ ) isomer under continuous illumination. The 3D X-ray structure of the photoinduced isomer is determined by steady-state and pseudo-steady-state photococrystallographic methods, providing insight into the structural changes required to accommodate photoswitching. Photococrystallographic kinetic studies follow the progress of photoswitching with time, determining a reaction rate constant of  $k = 0.38(2) \text{ min}^{-1}$  at 150 K. Linkage isomerism is fully-reversible on warming, and pseudo-steady-state experiments confirm that the photoexcited state is retained, at measurable occupancy, up to a temperature of 240 K. These results confirm the validity of combining photoactive linkage isomer and Re(I) photocatalyst chemistries, and the detailed determination of the photoexcited state structure will facilitate the future design of new photoactive Re(I) crystals for a range of applications.

Received 11th May 2018,  
Accepted 19th June 2018

DOI: 10.1039/c8ce00774h

rsc.li/crystengcomm

### Introduction

Molecular photoswitches that display a fast, useful change in their physical properties in response to irradiation, are an increasingly important area of photochemistry research. They are broadly applicable across a range of disciplines, from the design of smart materials and optoelectronics,<sup>1</sup> to biomedical applications such as photodynamic therapy<sup>2,3</sup> and fluorescent tracers for *in vivo* imaging,<sup>4</sup> and to environmental solutions such as waste water treatment<sup>5,6</sup> and solar energy.<sup>7</sup>

Sunlight is our most abundant and, as yet, under-exploited energy source, delivering more energy to the earth's surface in one hour than is used by humanity in an entire year,<sup>7</sup> and can produce electricity *via* far cleaner routes than fossil fuels. Though advances continue in this area, solar energy alternatives remain expensive.<sup>7</sup> As such, there is a continual need to design new, more efficient materials for solar harvesting, to achieve the ultimate goal of providing cheap and clean energy for all.

Taking inspiration from nature, much research is devoted to the design of biomimetic materials for artificial photosynthesis. Molecular switches find application here as photosensitizers and photocatalysts, accelerating key chemical reac-

tions that transform light into other useful forms of energy.<sup>8</sup> Photoactive organometallics are a key focus area, particularly in the design of homogeneous photocatalysts in solution. Among the most well studied materials are Re(I) systems of the general formula  $[\text{Re}(\text{R}_2\text{-bpy})(\text{CO})_3(\text{X})]$ ,<sup>9–12</sup> and this popularity is due in part to the fact that they are one of the few systems that combine both photosensitizer and photocatalytic centre within the same molecule,<sup>13</sup> promoting efficient charge transfer. Despite this interest, there remain some key issues for Re(I) photocatalysts. They suffer from problems common to other homogeneous catalysts, including issues with separation and recycling, as well as concerns over their long-term stability in solution. As such, there is significant interest in immobilising Re(I) systems into solid state scaffolds, enabling heterogeneous functionality.<sup>14–16</sup> Current Re(I) complexes also do not absorb across a broad enough range of the solar spectrum,<sup>17</sup> and key aspects of their photocatalytic mechanism are also not well understood. This is because of difficulties distinguishing between many structurally similar potential intermediates *via* solution spectroscopy.<sup>9,11</sup> It is here that dynamic structural chemistry methods, such as *Photococrystallography*,<sup>18</sup> can now make important contributions.

Since the late 1990s, photococrystallographic methods have been used to follow dynamic photoswitching in a variety of crystalline materials *in situ* and in real time.<sup>19</sup> Among the earliest systems to be studied are solid state linkage isomers, in which a long lived, metastable photoexcited isomer is produced at low temperature.<sup>20</sup> Building on seminal work by

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† Electronic supplementary information (ESI) available. CCDC 1842772–1842793.

For ESI and crystallographic data in CIF or other electronic format see DOI: [10.1039/c8ce00774h](https://doi.org/10.1039/c8ce00774h)



Coppens on metal-nitrosyl compounds,<sup>21</sup> a variety of linkage isomer systems have been studied including nitrosyl,<sup>22–25</sup> sulfur dioxide,<sup>26–29</sup> dinitrogen<sup>30</sup> and nitrite species.<sup>20,31–38</sup> In particular, nitro → nitrito isomerism is well studied by photocrystallographic methods, with the first 100% conversion in a single crystal reported in 2009 for  $[\text{Ni}(\text{dppe})(\text{Cl})(\eta^1\text{-NO}_2)]$ .<sup>35</sup> Several other crystals capable of 100% conversion have since been designed, and there is also interest in systems capable of room temperature photoswitching for real world applications.<sup>36</sup> Our recent work has demonstrated that the photoinduced nitrito-( $\eta^1\text{-ONO}$ ) isomer is strongly temperature dependent, enabling tuning of the excited state lifetime across a range of timescales.<sup>39</sup> This high degree of facile control makes these materials ideal test systems for the design of new photocrystallographic methods and equipment.

A handful of previous reports highlight an interest in combining photocatalytic Re(i) species with linkage isomer chemistry. Work by Dominguez *et al.* in 2014 reports the preparation of pure crystals of both  $[\text{Re}(\text{bpy})(\text{CO}_3)(\eta^1\text{-NO}_2)]$  and  $[\text{Re}(\text{Me}_2\text{-bpy})(\text{CO}_3)(\eta^1\text{-ONO})]$  from the reaction mixture. While the authors were able to demonstrate changes in solution state UV/vis and IR spectra on irradiation that may be indicative of photoinduced linkage isomer switching, no single-crystal-to-single-crystal transformation is reported.<sup>40</sup> In 2016, Kia and Safari also reported the synthesis of  $[\text{Re}(\text{Me}_2\text{-phen})(\text{CO}_3)(\eta^1\text{-ONO})]$ , but again no evidence of solid state switching is found.<sup>41</sup> There remains scope for detailed, *in situ* photocrystallographic investigations with  $[\text{Re}(\text{R}_2\text{-bpy})(\text{CO}_3)(\text{X})]$  systems, delivering key structural insight into how these systems respond to irradiation.

This work reports the synthesis, crystallisation and *in situ* photocrystallographic study of the novel complex  $[\text{Re}(\text{OMe}_2\text{-bpy})(\text{CO}_3)(\eta^1\text{-NO}_2)]$ . The 3D single crystal X-ray structure of a metastable state (MS) photoexcited nitrito-( $\eta^1\text{-ONO}$ ) isomer is determined, and the progress of the nitro → nitrito process is followed in real time by photocrystallographic kinetic methods. Further manipulation of the maximum excited state conversion level is also achieved under continuous illumination, *via* pseudo-steady-state photocrystallographic techniques.

## Experimental

### Synthetic procedures

Rhenium pentacarbonyl chloride  $[\text{Re}(\text{CO})_5\text{Cl}]$ , 4,4'-dimethoxy-2,2'-bipyridine ( $\text{OMe}_2\text{-bpy}$ ), silver triflate ( $\text{AgOTf}$ ) and potassium nitrite ( $\text{KNO}_2$ ) were all purchased from Sigma Aldrich and used as received. All solvents were reagent grade and used as purchased without further purification.

**Synthesis of  $[\text{Re}(\text{OMe}_2\text{-bpy})(\text{CO})_3(\text{OTf})]$ .**  $[\text{Re}(\text{OMe}_2\text{-bpy})(\text{CO})_3\text{Cl}]$  was synthesised following a literature procedure.<sup>10</sup>  $[\text{Re}(\text{OMe}_2\text{-bpy})(\text{CO})_3\text{Cl}]$  (0.06 g, 0.1 mmol) was converted to  $[\text{Re}(\text{OMe}_2\text{-bpy})(\text{CO})_3(\text{OTf})]$  by adding an excess of  $\text{AgOTf}$  (0.05 g, 0.2 mmol) in dichloromethane, and heating under reflux in the dark for *c.a.* 1 h. This known triflate species<sup>42</sup> was then isolated by filtration, purified by recrystallisation and its

identity confirmed by NMR, IR and a comparison of its unit cell parameters by XRD.

**$^1\text{H}$  NMR (500 MHz,  $\text{CDCl}_3$ ).**  $\delta_{\text{H}}$  8.87 [2H, d ( $J_{\text{HH}} = 6.4$  Hz), CH],  $\delta_{\text{H}}$  7.54 [2H, d ( $J_{\text{HH}} = 2.7$  Hz), CH],  $\delta_{\text{H}}$  7.04 [2H, dd ( $J_{\text{HH}} = 2.7, 6.5$  Hz), CH],  $\delta_{\text{H}}$  4.05 [6H, s,  $\text{CH}_3$ ].

**IR.**  $\nu(\text{CO})$  2029, 1934, 1893  $\text{cm}^{-1}$

### XRD

*CSD entry = YIYMAW.  $P2_1/c$ ,  $a = 8.725 \text{ \AA}$ ,  $b = 15.160 \text{ \AA}$ ,  $c = 14.750 \text{ \AA}$ ,  $\beta = 91.82^\circ$ ,  $V = 1948.732 \text{ \AA}^3$  (at 100 K).<sup>42</sup>*

*Experimental.*  $P2_1/c$ ,  $a = 8.701 \text{ \AA}$ ,  $b = 15.184 \text{ \AA}$ ,  $c = 14.710 \text{ \AA}$ ,  $\beta = 91.70^\circ$ ,  $V = 1942.5 \text{ \AA}^3$  (at 150 K).

**Synthesis of  $[\text{Re}(\text{OMe}_2\text{-bpy})(\text{CO})_3(\text{NO}_2)]$  – complex 1.** A portion of  $[\text{Re}(\text{OMe}_2\text{-bpy})(\text{CO})_3(\text{OTf})]$  (0.06 g, 0.1 mmol) was dissolved in 2 mL methanol and treated with an excess of  $\text{KNO}_2$  (0.08 g, 1 mmol). The reaction mixture was stirred at room temperature in the dark for *c.a.* 24 hours, before the product solution was collected by filtration and dried under vacuum. The resulting yellow solid was recrystallised from dichloromethane by the slow diffusion of diethyl ether vapour at room temperature. The crystallisation vial was kept in the dark throughout. The identity of 1 was confirmed by NMR, IR and XRD (see below).

**$^1\text{H}$  NMR (500 MHz,  $\text{CDCl}_3$ ).**  $\delta_{\text{H}}$  8.91 [2H, d ( $J_{\text{HH}} = 6.5$  Hz), CH],  $\delta_{\text{H}}$  7.51 [2H, d ( $J_{\text{HH}} = 2.4$  Hz), CH],  $\delta_{\text{H}}$  7.02 [2H, dd ( $J_{\text{HH}} = 2.5, 6.4$  Hz), CH],  $\delta_{\text{H}}$  4.02 [6H, s,  $\text{CH}_3$ ].

**IR.**  $\nu(\text{CO})$  2030, 1931, 1891  $\text{cm}^{-1}$ ,  $\nu(\text{NO})$  1525, 1306  $\text{cm}^{-1}$ .

### Single crystal X-ray crystallography

Single crystal X-ray data were recorded on a dual source Rigaku-Oxford Diffraction Gemini A Ultra diffractometer, equipped with an Atlas CCD detector and an Oxford Cryosystems Cryojet-XL liquid nitrogen flow device for temperature control. Data collection, indexing and integration were performed using the Rigaku-Oxford Diffraction software CrysAlisPro.<sup>43</sup> Structures were solved by dual-space methods in SHELXT,<sup>44</sup> or by direct methods in SHELXS,<sup>45</sup> and refined by full matrix least-squares on  $F^2$  in SHELXL.<sup>46</sup> Hydrogen atoms were positioned geometrically and refined using a riding model. The hydrogen atom isotropic displacement parameters were fixed to  $U_{\text{iso}}(\text{H}) = 1.5 \times$  (for  $\text{CH}_3$ ) or  $U_{\text{iso}}(\text{H}) = 1.2 \times$  (for CH) the  $U_{\text{eq}}$  of the parent atom.

### Photocrystallography

For photocrystallographic studies, the standard single crystal X-ray diffraction set-up was modified to incorporate a purpose-built LED ring array,<sup>47</sup> which positions five LEDs (405 nm: Bivar UV5TZ-405-15,  $\lambda_p = 405 \pm 2.5$  nm, 3.4 V, 15 mA, 40 mW) at a distance of approximately 1 cm from the crystal in a uniform arc. This set-up helps to ensure even irradiation on all sides of the crystal, and enables the collection of complete single-crystal X-ray data while the array remains in place. To further ensure even illumination, for steady-state experiments the crystal was also continuously rotated about the phi axis during irradiation.

## Results and discussion

### Ground state single crystal X-ray structure

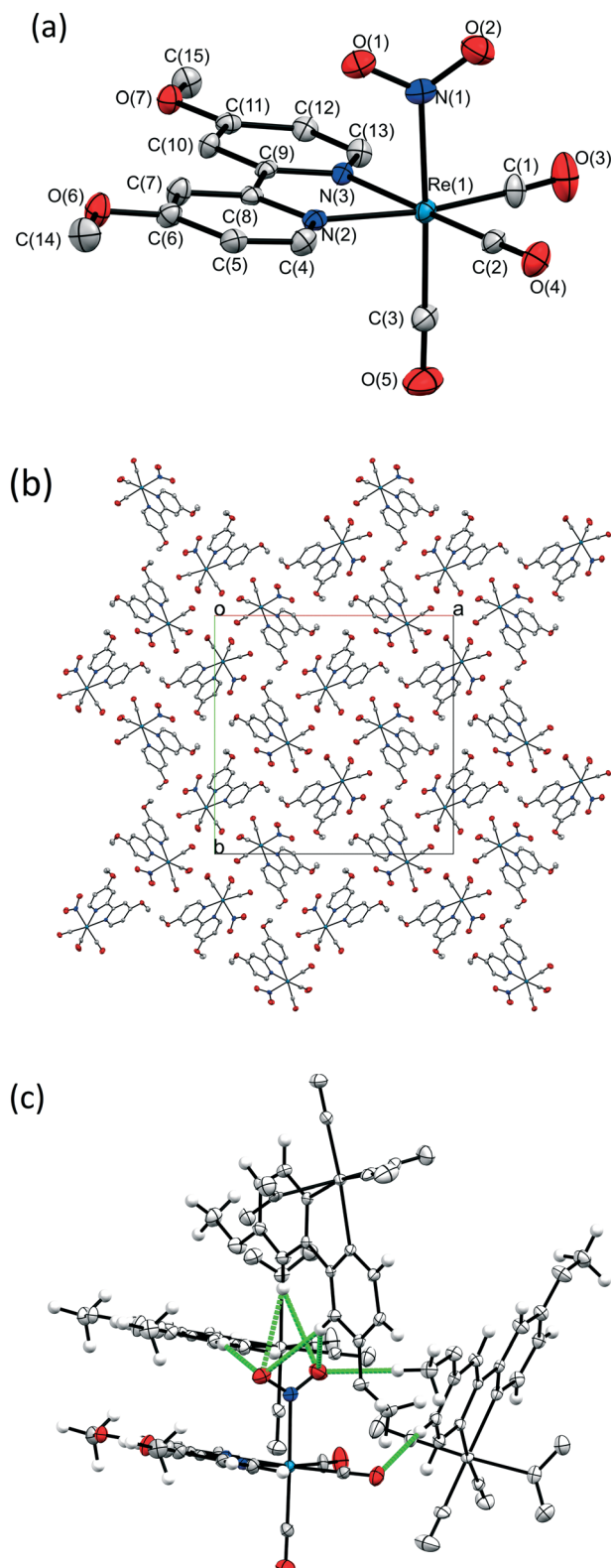
A crystal of **1** was flash frozen at 150 K and the GS single crystal X-ray structure was determined in the dark. **1** crystallises in the tetragonal space group  $I\bar{4}$ , with one Re complex molecule in the asymmetric unit. The Re-centre adopts a distorted octahedral geometry and is bound to three carbonyl ligands, one bidentate  $\text{OMe}_2\text{-bpy}$  ligand and one monodentate nitrite group, which is coordinated solely as the nitro- $(\eta^1\text{-NO}_2)$  isomer. Fig. 1a shows the full atomic connectivity in the asymmetric unit, and selected X-ray data are provided in Table 1.

Fig. 1b shows the crystal packing arrangement in **1**. The  $[\text{Re}(\text{OMe}_2\text{-bpy})(\text{CO})_3(\text{NO}_2)]$  molecules appear closely packed, however a contact surface void space calculation in the CCDC software Mercury<sup>48</sup> (probe radius 1.2 Å, grid spacing 0.1 Å) reveals small pockets of space at points along the 4-fold rotation axes, comprising a total of 65.65 Å<sup>3</sup>, or 2% of the unit cell volume. These voids suggest inefficient packing between the carbonyl ligands oriented towards each other in this region (see Fig. S1, ESI<sup>†</sup>). Any void space in the crystal lattice is significant as it could potentially assist the atomic movements necessary for linkage isomerism to occur. Due to the lack of traditional hydrogen bond donors in **1** there are no strong hydrogen bonds between molecules. However, there are a number of intermolecular  $\text{CH}\cdots\text{O}$  contacts, including several to the potentially photoactive nitro- $(\eta^1\text{-NO}_2)$  group (see Fig. 1c, Table S1 ESI<sup>†</sup>). These intermolecular interactions are significant because, for linkage isomerism to occur, these contacts must be broken and potentially re-formed. As such, they present a likely barrier to the progress of nitro  $\rightarrow$  nitrito photoswitching in **1**.

### Steady-state photococrystallography with **1** at 150 K

The same crystal was held at 150 K and irradiated, *in situ* on the diffractometer, using the LED ring array set-up outlined above. A wavelength of 405 nm, towards the tail of the absorption maxima, was selected by reference to the solid state absorption spectrum of **1** (Fig. S2/S3, ESI<sup>†</sup>). This choice reflects the results of previous investigations, in which wavelengths in the tail of the main absorption band are found to promote solid state photoswitching, helping to maximise the penetration of the excitation light through the crystal.<sup>49</sup>

In light of previous photococrystallographic experiments with metal-nitrite crystals,<sup>20,31,34,36,50</sup> a long-lived metastable isomer is expected on irradiation of **1** at 150 K. Thus, using the steady-state photococrystallographic method,<sup>20</sup> after an illumination period of 1 min the LEDs were switched off and a second X-ray dataset was collected. This revealed that photoactivation had been achieved, with 29% of the crystal now converted to an *endo*-nitrito- $(\eta^1\text{-ONO})$  isomer (Fig. 2a). The steady-state experiment was then continued, with irradiation paused at regular intervals to allow the collection of a complete X-ray diffraction dataset. These results were then used to plot the photoexcitation profile for **1** with respect to irradiation time at 150 K (Fig. 2b). The photoreaction initially pro-



**Fig. 1** Single-crystal X-ray structure of the GS of **1** at 150 K. (a) Atomic connectivity in the asymmetric unit, ellipsoids at 50% probability, hydrogen atoms removed for clarity; (b) crystal packing arrangement; (c)  $\text{C}-\text{H}\cdots\text{O}$  interactions involving the nitro- $\text{NO}_2$  ligand.



**Table 1** Single crystal X-ray data for **1**. Selected crystal data for the ground state (GS) and the metastable state (MS) at 150 K

	Ground state (GS)	Metastable state (MS) <sup>a</sup>
Photoconversion	0%	58%
Irradiation time (total)	0 min	180 min
Temperature	150(1) K	150(1) K
Wavelength	0.71073 Å	0.71073 Å
Empirical formula	C <sub>15</sub> H <sub>12</sub> N <sub>3</sub> O <sub>7</sub> Re <sub>1</sub>	C <sub>15</sub> H <sub>12</sub> N <sub>3</sub> O <sub>7</sub> Re <sub>1</sub>
Formula weight	532.48 g mol <sup>-1</sup>	532.48 g mol <sup>-1</sup>
Crystal system	Tetragonal	Tetragonal
Space group	<i>I</i> 4	<i>I</i> 4
Unit cell parameters (constrained)	<i>a</i> = 22.2989(5) Å <i>c</i> = 6.6471(2) Å <sup>3</sup> 3305.2(2) Å <sup>3</sup>	<i>a</i> = 22.6482(5) Å <i>c</i> = 6.5591(2) Å <sup>3</sup> 3364.4(2) Å <sup>3</sup>
Volume	8	8
<i>Z</i>	2.140 g cm <sup>-1</sup>	2.102 g cm <sup>-1</sup>
Density (calculated)	7.398 mm <sup>-1</sup>	7.268 mm <sup>-1</sup>
Absorption coefficient $\mu$	2032	2032
<i>F</i> (000)	0.0362	0.0748
<i>R</i> (int)	0.987	0.997
Completeness (to $\theta = 25.00^\circ$ )	0.0275	0.0453
<i>R</i> <sub>1</sub> (observed data $I > 2\sigma(I)$ )	0.0554	0.1121
w <i>R</i> <sub>2</sub> (all data)	6711 (3356)	11 493 (3157)
Reflections (independent)	-0.031(11)	-0.048(11)
Flack parameter		

<sup>a</sup> The MS structure at 150 K is recorded in the 180K\_405nm\_150K.cif crystallographic information file, see ESI.

gresses quickly, but the conversion rate rapidly slows with irradiation time, producing a sigmoidal-shaped profile. A maximum conversion of 58% to the *endo*-nitrito isomer was determined after a total of 3 h illumination. Further irradiation caused no appreciable change, confirming that a photo-stationary state had been achieved at this temperature. This excitation level was retained when the crystal was held at 150 K in the dark for a further 1 h, proving that the excited state isomer is metastable at this temperature. A significant reduction in the quality of the diffraction data was seen on irradiation, which is reflected in the increased *R*-indices for the MS structure (Table 1, S3†).

Analysis of the photoconversion profile in Fig. 2b can provide kinetic information on the progress of nitro → nitrito photo-switching in **1**. As shown previously in the literature, the JMAK theory for solid state kinetics provides a good fit for single-crystal-to-single-crystal photoreactions,<sup>36,39,50–53</sup> and can be adapted for both incomplete excitation and decay.<sup>39,51</sup> The general equation for JMAK analysis is given as:

$$\alpha(t) = 1 - e^{-kt^n} \quad (1)$$

where  $\alpha(t)$  = the photoconversion fraction, or MS level, present at a time  $t$ ,  $k$  = the reaction rate constant and  $n$  = the Avrami exponent.<sup>54–56</sup>  $n$  provides a measure of the dimensionality ( $D$ ) of the growth of a new MS phase throughout the crystal, where:

$$D = n - 1 \quad (2)$$

Here, as the photoreaction in **1** remains incomplete at 150 K, the data were fitted with a modified version of eqn (1),<sup>51</sup> which allows the initial and final conversion fractions to be parameterised:

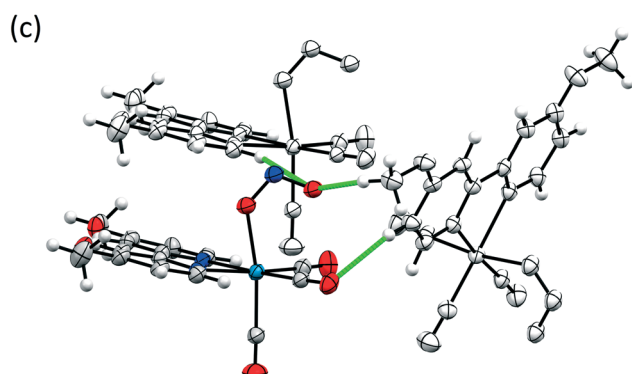
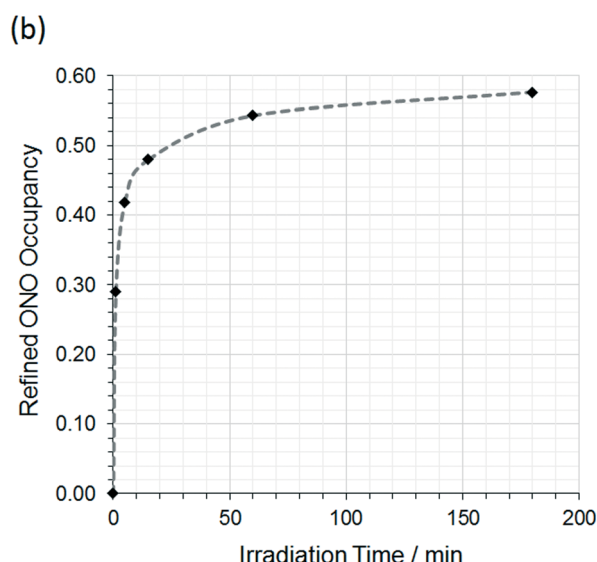
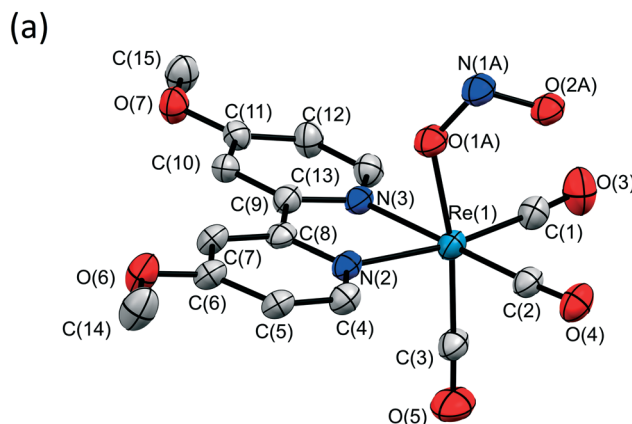
$$\alpha(t) = \alpha(f) - [\alpha(f) - \alpha(0)]e^{-kt^n} \quad (3)$$

where  $\alpha(0)$  and  $\alpha(f)$  are the initial and final conversion fraction values, respectively.

A least squares regression analysis was performed on the data points in Fig. 3 using eqn (3), producing a good fit to the data with a value of 0.996 for  $R^2$ . The initial and final conversion fractions were fixed to their experimental values of  $\alpha(0) = 0$  and  $\alpha(f) = 0.576$ , while the parameters  $k$  and  $n$  were allowed to refine freely. A reaction rate constant of  $k = 0.38(2)$  min<sup>-1</sup> was determined, which is comparable to those reported previously for nitro → nitrito conversion processes in the single crystal at low temperature.<sup>36,39,50</sup> For the Avrami exponent, a refined value of  $n = 0.35(2)$  was determined. While a value of  $n \approx 1$  ( $D = 0$ ) is indicative of zero-dimensional, or homogeneous, growth of the MS phase through the crystal, a value of  $n < 1$  suggests a negative dimensionality for the production of the MS phase, which is not physically realistic. This result indicates that eqn (3) is insufficient to accurately model photoconversion in **1**, and there is likely to be some other physical processes limiting the progress of the nitro → nitrito conversion through the crystal that are not accounted for by simple JMAK theory. Possible limiting factors could include incomplete penetration of excitation light through the crystal bulk, or steric influences resulting from strain in the crystal lattice on photoswitching (see the comparison of GS/MS discussion below).

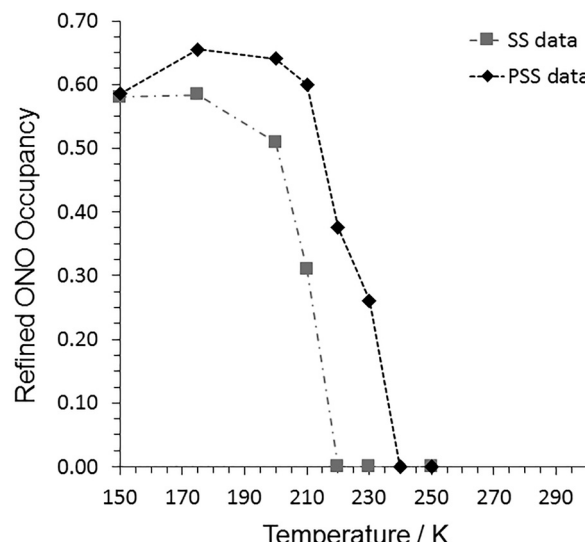
The final step of the steady-state photocrystallographic experiment is to determine the temperature range over which the system is metastable. The crystal in its photoinduced MS was next subject to a variable temperature study between 150 and 300 K (Fig. 3, Tables S2/S4†). The MS occupancy remained constant on heating to 175 K, above which





**Fig. 2** Photo-induced metastable state (MS) structure and refined occupancy levels for **1** at  $\lambda = 405$  nm. (a) Single-crystal X-ray structure showing the atomic connectivity for the MS *endo*-nitrito-(<math>\eta^1\text{-ONO}</math>) isomer at 150 K. (b) Graph of photoconversion level vs. irradiation time in a single-crystal of **1** at 150 K. (c) C-H...O interactions involving the photoinduced nitrito-ONO ligand.

temperature the crystal then began to decay back towards the GS, indicating an approximate temperature for the apparent “metastable limit” for this crystal system.<sup>39</sup> **1** then finally reverts back to a 100% GS nitro-(<math>\eta^1\text{-NO}\_2</math>) isomer by 220 K.



**Fig. 3** Refined nitro:nitrito isomer occupancies from variable temperature parametric studies with **1**, recorded under both steady-state (SS) and pseudo-steady-state (PSS) conditions.

#### Comparison of GS and MS structures of **1** at 150 K

The crystal packing arrangement in **1** is largely unchanged on irradiation, excepting the change in nitrite coordination mode (see Fig. S4/S5, ESI†). However, a comparison of Fig. 1c and 2c shows that there is a significant rearrangement in the intermolecular C-H...O contacts involving the isomerising ligand. For nitro  $\rightarrow$  nitrito isomerism to occur, six C-H...O contacts to the nitro-(<math>\eta^1\text{-NO}\_2</math>) ligand need to be broken, with an average C...O distance of 3.52(2) Å. In the MS, there are just two C-H...O contacts formed by the nitrito-(<math>\eta^1\text{-ONO}</math>) ligand, with a longer average C...O distance of 3.57(4) Å. Full details of the C-H...O distances in the MS are given in Table S5, ESI†. Overall, the intermolecular interactions involving the photoinduced isomer are longer, and therefore likely to be weaker, than those influencing the GS isomer, suggesting that conversion to the MS nitrito-ONO arrangement will not be energetically favourable.

A comparison of the unit cell dimensions in Table 1 for the GS and MS structures reveals that there is a significant increase of 59.2(3) Å<sup>3</sup> (or 1.79%) in the unit cell volume on photoactivation. This is a reasonably large change in unit cell volume in comparison to those reported previously, and may point to an explanation for the incomplete level of photoactivation in **1**. Typically, metal-nitrite crystals displaying high levels of photoactivation, up to 100%, tend to produce a volume change of <1%.<sup>34–36</sup> This larger unit cell change may suggest a significant amount of strain is induced in the crystal as a result of the photoisomerisation, which in turn limits the overall conversion fraction that can be accommodated while maintaining single crystal integrity.

In addition to an assessment of overall changes in the unit cell, the “reaction cavity” concept can provide insight into the local structure changes in the region of the isomerising nitrite group, helping to rationalise how or why the



photoreaction proceeds.<sup>57,58</sup> The reaction cavity is defined as the volume encapsulating the photoactive group and, in particular, the change in the reaction cavity volume  $\Delta V_c$  is a useful measure of how much the local structure must change in order to accommodate  $\text{nitro} \rightarrow \text{nitrito}$  rearrangement. Table 2 provides reaction cavity analyses performed on the GS and MS structures of **1** using the CCDC software Mercury.<sup>48</sup> The reaction cavity volume more than doubles on irradiation of the crystal, with a value for  $\Delta V_c$  of  $10.2 \text{ \AA}^3$  per molecule, equating to a 101.5% increase. In addition, a qualitative visual assessment of the reaction cavities reveals that their shape is also significantly altered on excitation (see Fig. S6, ESI†). Both of these factors indicate that there is a significant change required in the local crystal structure to accommodate the isomerisation, which is expected to induce considerable strain on the surrounding crystalline array. This provides a likely explanation not only for the incomplete photoconversion level in **1**, but also for the significant reduction in data quality on irradiation, and for the low value of the Avrami exponent  $n$  in the JMAK kinetic analysis above.

### Steady-state photocrystallography at 170 K

In our recent work with Pd and Pt-nitrite linkage isomer crystals that also show incomplete conversion at low temperature, we found that irradiation at higher temperatures, just below the “metastable limit” for the system, can help to increase the overall conversion level.<sup>31</sup> As such, a second steady-state photocrystallographic experiment was also conducted with **1** at 170 K. Following a GS X-ray structure determination at 170 K, a crystal was then irradiated for a period of 3 h, before the LEDs were switched off and a second X-ray dataset collected (see Table S6, ESI†). A maximum nitrito-ONO occupancy of 60% was determined from these data, which is comparable, within error, to the photostationary state achieved at 150 K. As such, in the case of the  $\text{Re}(\text{i})$ -complex **1**, it is apparent that raising the temperature does not have a marked effect on the overall level of photoactivation that can be achieved under steady-state photocrystallographic conditions.

### Pseudo-steady-state photocrystallography with **1**

A crystal of **1** was next subjected to a pseudo-steady-state photocrystallographic investigation to determine if the nitro:nitrito ratio can be further manipulated under continuous illumination. Previous work has shown that a photostationary equilibrium is induced between the forwards photoexcitation and the reverse thermal decay processes with continuous irradiation,<sup>32,36,39</sup> and these conditions are easily achieved using

the LED ring array set-up. The crystal was first mounted at 150 K and irradiated for a period of 3 h to ensure that an equilibrium nitro:nitrito occupancy had been reached at this temperature. A complete X-ray dataset was then collected, with the LEDs remaining on and in place, to reveal the pseudo-steady-state photoconversion level. Irradiation was then continued while the temperature was raised, pausing at regular intervals to enable further pseudo-steady-state experiments to be collected.

The refined nitro:nitrito occupancy levels from these data are shown in Fig. 3 and Table S7.† At 150 K, the pseudo-steady-state conversion fraction is the same, within error, as that obtained after 3 h illumination in the steady-state experiments. However, on warming the crystal to 175 K, continuous illumination induces an increase in the maximum conversion level in **1**, achieving a maximum nitrito-( $\eta^1$ -ONO) occupancy of 66%. Selected X-ray crystal data and structure information for the pseudo-steady state data at 150 and 175 K are provided in Table 3. At higher temperatures the equilibrium excited state isomer occupancy gradually decreases, finally returning to the 100% GS nitro-( $\eta^1$ -NO<sub>2</sub>) isomer by 240 K.

This result is interesting as it suggests that, under continuous illumination conditions, raising the temperature helps overcome some of the barriers limiting the overall photoconversion level. To assess this more quantitatively, a direct comparison may be made between the pseudo-steady-state structures at 150 and 175 K. Table 3 shows that there is a negligible  $0.5(10) \text{ \AA}^3$  (or 0.014%) increase in unit cell volume on warming under continuous illumination, and there is similarly no significant increase in the void space in the crystal, using a contact surface calculation in Mercury (probe radius  $1.2 \text{ \AA}$ , grid spacing  $0.1 \text{ \AA}$ ). Interestingly, this observation is at odds with the results of the steady-state studies and tends not to support the theory of steric strain as a barrier to photoconversion, as it would be expected that an increase in excitation level should correlate with an observed increase in free space in the overall crystal structure. Similarly, an assessment of the reaction cavity volumes at 150 and 175 K under continuous irradiation shows no significant change (see Table S8†). However, as irradiation continues throughout the pseudo-steady-state experiment, the data quality is observed to decrease considerably with a significant increase in  $R$ -indices, reduction of  $I/\sigma$  and loss of high angle diffraction data (see Table S9†). As such, it is possible that damage to the crystal may also influence the overall photoconversion fraction observed in these studies (e.g. due to loss of crystallinity at the crystal surface and/or crystal fracturing during the experiment), and so care should be taken when drawing any more detailed conclusions from these data.

**Table 2** Reaction cavity ( $V_c$ ) analysis for the GS and MS of **1**.  $V_c$  determined by removing the nitrite group and performing a contact surface void space calculation in Mercury<sup>48</sup> (probe radius  $1.2 \text{ \AA}$ , grid spacing  $0.1 \text{ \AA}$ ) and subtracting the initial void volume

	$V_c$ per unit cell/ $\text{\AA}^3$	$V_c$ per molecule/ $\text{\AA}^3$	$\Delta V_c$ (MS-GS)/ $\text{\AA}^3$	$\Delta V_c$ (MS-GS)/%
GS	80.26	10.03	10.19	101.5
MS	161.76	20.22		



**Table 3** Selected crystal data for the pseudo-steady-state structures of **1** at 150 and 175 K

	PSS 150 K	PSS 175 K
Photoconversion	59%	66%
Temperature	150(2) K	175(2) K
Wavelength	0.71073 Å	0.71073 Å
Empirical formula	C <sub>15</sub> H <sub>12</sub> N <sub>3</sub> O-Re <sub>1</sub>	C <sub>15</sub> H <sub>12</sub> N <sub>3</sub> O-Re <sub>1</sub>
Formula weight	532.48 g mol <sup>-1</sup>	532.48 g mol <sup>-1</sup>
Crystal system	Tetragonal	Tetragonal
Space group	<i>I</i> 4	<i>I</i> 4
Unit cell parameters (constrained)	<i>a</i> = 22.719(2) Å <i>c</i> = 6.5611(8) Å	<i>a</i> = 22.739(2) Å <i>c</i> = 6.5506(8) Å
Volume	3386.7(7) Å <sup>3</sup>	3387.2(7) Å <sup>3</sup>
<i>Z</i>	8	8
Density (calculated)	2.089 g cm <sup>-1</sup>	2.088 g cm <sup>-1</sup>
Absorption coefficient $\mu$	7.220 mm <sup>-1</sup>	7.219 mm <sup>-1</sup>
<i>F</i> (000)	2032	2032
<i>R</i> (int)	0.0618	0.0652
Completeness (to $\theta = 25.00^\circ$ )	0.996	0.996
<i>R</i> <sub>1</sub> (observed data $I > 2\sigma(I)$ )	0.0430	0.0443
<i>wR</i> <sub>2</sub> (all data)	0.0975	0.0966
Reflections (independent)	6030 (2903)	6047 (2900)
Flack parameter	-0.016(18)	-0.01(2)

## Conclusions

This study delivers the first 3D structure confirmation of photoinduced nitro → nitrito conversion in a Re(i)-bpy single crystal, providing a detailed determination of the photoinduced [Re(OMe<sub>2</sub>-bpy)(CO)<sub>3</sub>(*n*<sup>1</sup>-QNO)] excited state structure. The combination of photoactive linkage isomer chemistry with that of Re(i)-bpy complexes broadens the scope of dynamic structural studies on linkage isomers and also relates the photoactive properties of this and other Re(i) molecules to photocatalytic studies on related systems. The detailed, 3D understanding of the structural changes that occur on photoactivation at the atomic-scale in **1** will benefit the design of new and more efficient Re(i) compounds, in both the solid and solution state, which can be further optimised towards specific photoswitching applications.

The successful use of 405 nm light to induce a considerable (66%) photoactivation level in the system is of particular importance, as many Re(i)-photocatalyst systems are limited to absorption at shorter wavelengths in the UV region.<sup>17</sup> By broadening the range of the UV/vis absorption spectrum in the solid state, these Re(i)-nitrite crystal systems can absorb a wider range of the solar spectrum and therefore have the potential to increase solar conversion efficiency.

The incomplete photoconversion level achieved in **1**, even under continuous illumination, and the considerable reduction in X-ray data quality on irradiation are both considerable limitations for the development of single crystals of **1** for real photoswitchable device solutions. Given the indication of significant steric strain in the crystal, and its potential impact on the photoactivation levels observed, future studies will focus on the development of new Re(i) crystal systems containing bulkier auxiliary fragments (ligands and/or counter-ion moieties) – a crystal engineering route that we

have previously shown can successfully facilitate higher levels of nitro → nitrito conversion by controlling the reaction cavity volume and minimising  $\Delta V_c$ .<sup>34,36</sup>

## Conflicts of interest

There are no conflicts of interest to declare.

## Acknowledgements

LEH would like to thank the UK Engineering and Physical Sciences Research Council (EPSRC) for financial support. This work is supported by EPSRC Programme Grant EP/K004956/1 and EP/M010481/1.

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