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## Raising the (metastable) bar: 100% photo-switching in $[\text{Pd}(\text{Bu}_4\text{dien})(\eta^1\text{-NO}_2)]^+$ approaches ambient temperature†

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100% nitro-nitrito linkage isomerism is reported in single-crystals of the  $\text{Pd}(\text{II})$ -nitrite system  $[\text{Pd}(\text{Bu}_4\text{dien})(\eta^1\text{-NO}_2)]\text{BPh}_4$  ( $\text{Bu}_4\text{dien} = N,N,N',N'$ -tetrabutylmethylenetriamine,  $\text{BPh}_4$  = tetraphenylborate). Complete conversion to a metastable *endo*-nitrito-( $\eta^1\text{-ONO}$ ) isomer is achieved after just 15 min irradiation with 400 nm LED light. The system is entirely metastable below 240 K, while pseudo-steady-state photocrystallographic experiments confirm that the excited state isomer is retained, at substantial conversion levels, under continuous illumination until 260 K. These results show promise for new linkage isomer systems, based on heavier transition metal centres, which display full functionality under near-ambient conditions.

## Introduction

Photo-switchable molecular compounds, based on organic and organometallic chemistries, continue to be important in the search for new functional materials that will address a number of real-world challenges, including renewable energy solutions, sensors and data storage.<sup>1</sup> Light-induced reactions are tunable, site-specific and are often efficient and clean, all of which increases their popularity for a number of applications. In particular, molecules whose photo-switching behaviour is retained in the solid-state can be advantageous as these are often more easily developed into devices. Where possible, single-crystal X-ray diffraction (SC-XRD) continues to be the preferred method to study these solid-state systems, providing highly-accurate, 3-D structural parameters at all stages of the reaction. The detailed study of photochemical switching in crystalline materials was pioneered by Schmidt, Cohen *et al.* in the 1960s,<sup>2–4</sup> whose work with *trans*-cinnamic acid derivatives developed an understanding of the conditions necessary for [2 + 2] photo-dimerisation in single-crystals. While this early work relied primarily on SC-XRD experiments conducted on a crystal in its ground state (GS), dramatic advancements in the SC-XRD method since this time now allow the determination of GS, metastable state (MS) and entirely transient excited state species over a range of time-scales and temperatures.<sup>5,6</sup> Such experiments are conducted *via* so-called

photocrystallographic methods<sup>7</sup> and require *in situ* irradiation of a crystal either before or during the X-ray experiment.

The earliest photocrystallographic studies were conducted by Coppens *et al.* in the mid-1990s and were the first experiments to unambiguously determine the presence of MS linkage isomers at low temperature in the Fe-nitrosyl complex sodium nitroprusside.<sup>8</sup> Metastable linkage isomers have the potential to be exploited as functional materials for applications where a long-lived excited state is desirable. In the last 20 years, single-crystal-to-single-crystal linkage isomerism has been reported for a number of small di- and tri-atomic ligands including nitrosyl,<sup>9–12</sup> di-nitrogen,<sup>13</sup> sulfur dioxide<sup>14–18</sup> and nitrite groups.<sup>19–24</sup> Despite these successes, there remain some key factors that limit the functionality of metastable linkage isomer systems and must be overcome before these can be considered viable candidates for real devices. One such limitation is the maximum level of photo-activation that can be achieved in the crystal. Ideally, 100% activation that is reversible, controllable and reproducible over many repeat cycles should be achieved. The first fully-reversible, 100% photo-activation in a linkage isomer crystal was reported by Warren *et al.* in 2009, in  $[\text{Ni}(\text{dppe})(\text{NO}_2)\text{Cl}]$  (dppe = bis-diphenylphosphinoethane).<sup>19</sup> 100% excitation is rationalised by the inclusion of the bulky di-phosphine ligand as this photo-inert fragment can dominate the packing, providing a “reaction cavity”<sup>25–27</sup> in which the smaller nitrite ligand can rearrange without imparting undue strain to the surrounding lattice. Since that time, this crystal engineering principle has been utilised by Raithby *et al.* in designing several  $\text{Ni}(\text{II})$ -nitrite species with bulky co-ligands, which are capable of very high nitro-nitrito conversion in the single-crystal.<sup>20,22–24</sup> A second factor limiting the efficacy of metastable linkage

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isomers as functional molecular switches is the relatively low critical temperature, or “MS limit”, above which the system regains its GS arrangement. Currently, MS limits for the high-converting Ni(II)-nitrite complexes reported to-date do not exceed 200 K. A few Pd(II) and Pt(II)-nitrite systems boast MS limits between 200 and 240 K,<sup>21,28</sup> while some Ru-SO<sub>2</sub> compounds reach limits as high as 250 K.<sup>14,29</sup> However, these systems do not achieve high levels of GS to MS conversion. MS linkage isomers capable of 100% switching at near-ambient temperature would be most desirable for practical applications. The current study reports here-in a novel Pd(II)-nitrite species capable of 100% photo-activation, with a long-lived nitrito-( $\eta^1$ -ONO) isomer still detectable by SC-XRD methods at 250 K.

## Experimental section

### Synthetic preparations

All manipulations were carried out in air. All reagents were used as purchased, without the need for further purification.

**Potassium tetranitropalladate(II).** Potassium tetranitropalladate(II) was prepared *via* a modified literature procedure.<sup>30</sup> Excess potassium nitrite (2.55 g, 30 mmol) was dissolved in distilled water (2 mL) and added to an aqueous suspension (5 mL) of palladium(II) chloride (0.89 g, 5 mmol). The red suspension initially dissolved to yield a yellow solution, before a yellow solid precipitated over a 2 h period. The product was collected by filtration and dried *in vacuo*.

**Nitro-(*N,N,N',N'*-tetrabutyldiethylenetriamine)palladium tetraphenylborate.** *N,N,N',N'*-Tetrabutyldiethylenetriamine (Bu<sub>4</sub>dien, 0.39 mL, 1 mmol) was dissolved in acetone (5 mL) and added to a solution of potassium tetranitropalladate(II) (5 mL) in water. The resulting yellow solution was stirred overnight. Excess sodium tetraphenylborate (0.69 g, 2 mmol) was dissolved in water/acetone (10 mL), then added to the reaction. The cream precipitate that formed was collected by filtration and dried *in vacuo*.

**Elemental analysis, for C<sub>44</sub>H<sub>65</sub>B<sub>1</sub>N<sub>4</sub>O<sub>2</sub>Pd<sub>1</sub>.** Calc. (%): C = 66.12; H = 8.20; N = 7.01. Found (%): C = 66.25; H = 8.33; N = 7.23.

**<sup>1</sup>H NMR (500 MHz, CDCl<sub>3</sub>).** δ<sub>H</sub> 1.00 (6H, t, CH<sub>3</sub>); δ<sub>H</sub> 1.14 (6H, t, CH<sub>3</sub>); δ<sub>H</sub> 1.29 (8H, m, CH<sub>2</sub>); δ<sub>H</sub> 1.38 (8H, m, CH<sub>2</sub>); δ<sub>H</sub> 2.06 (4H, t, CH<sub>2</sub>); δ<sub>H</sub> 2.16 (4H, t, CH<sub>2</sub>); δ<sub>H</sub> 2.35 (1H, b, NH); δ<sub>H</sub> 2.54 (8H, m, CH<sub>2</sub>); δ<sub>H</sub> 6.99–7.47 (20H, m, C<sub>6</sub>H<sub>5</sub>).

**FT-IR.** ν(N–O) = 1582 cm<sup>-1</sup>, 1333 cm<sup>-1</sup>; ν(N–H) = 3137 cm<sup>-1</sup>.

Single-crystals of the tetrahydrofuran (THF) solvate (1) were obtained by recrystallization from THF/diethyl ether.

### UV/vis diffuse reflectance spectroscopy

UV/vis diffuse reflectance spectra were collected on a Shimadzu UV-2600 spectrophotometer with an integrating sphere. Data were collected on a powdered crystalline sample of (1) pelleted with KBr, between 800 and 200 nm at ambient temperature.

### Single-crystal X-ray crystallography

Standard single-crystal X-ray diffraction (SC-XRD) experiments were conducted at the University of Bath using a Rigaku–Oxford Diffraction Gemini A Ultra diffractometer equipped with an Atlas CCD detector. The crystal temperature was controlled by an Oxford Instruments CryojetXL liquid nitrogen flow device, between 100–300 K. Data collection, indexing and integration procedures were completed with CrysAlis Pro.<sup>31</sup> Structures were solved by dual-space methods using SHELXT<sup>32</sup> and refined by full-matrix least squares on F<sup>2</sup> using SHELXL-2014.<sup>33</sup>

### Photocrystallographic experiments

*In situ* irradiation of the crystal was achieved using a specifically designed LED ring array.<sup>34</sup> Six 400 nm LEDs (Bivar Opto<sup>TM</sup> LED5-UV-400-30 Series, 350 med, 3.8 V, 12 mW) are positioned 1 cm from the sample in a uniform circle. SC-XRD data collection, indexing and reduction were completed as outlined previously. Metastable state (MS) X-ray data were refined against ground state (GS) coordinates and a Fourier electron density difference map (photodifference map) used to identify new features corresponding to the MS. In the final refinement, partially-activated systems were treated as a disorder model with linkage isomer occupancies refined using standard free variables.

**Steady-state photocrystallographic studies.** Steady-state photocrystallographic studies required the crystal to be irradiated for a set time period, whilst the sample was maintained at low temperature (100 K). After the irradiation period, LEDs were removed and a standard SC-XRD experiment conducted in the dark.

**Pseudo-steady-state photocrystallographic studies.** Pseudo-steady-state photocrystallographic studies required continuous illumination of the sample throughout the SC-XRD data collection, which was facilitated by the LED ring set-up. The crystal was irradiated for a period of 1 h prior to the start of any pseudo-steady-state photocrystallographic data collection, to ensure an equilibrium GS/ES occupancy had been reached.

## Results

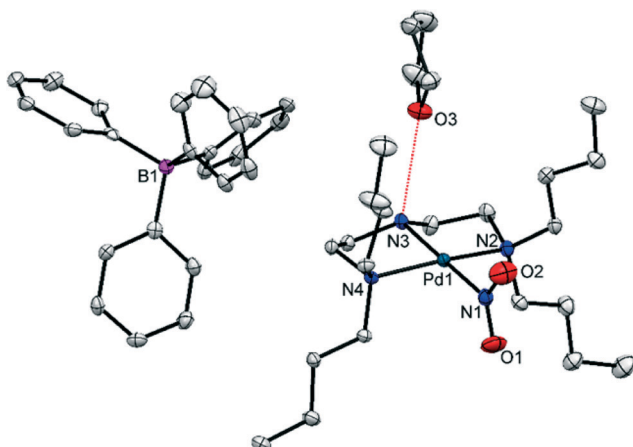
### Ground state (GS) crystal structure

A single-crystal of (1) was mounted in oil at ambient temperature and cooled to 100 K in the dark, before a standard SC-XRD dataset was obtained (Table 1). Fig. 1 shows the structure of the asymmetric unit (the full atomic numbering scheme for the palladium cation is given in Fig. S1†). With this data it is possible to conduct a detailed analysis at the molecular level and, in terms of the crystal packing, establish whether a switch of coordination mode is likely in (1).

(1) crystallises in the monoclinic space group P2<sub>1</sub>/c with one complex cation, one BPh<sub>4</sub> anion and one THF solvent molecule in the asymmetric unit. The Pd(II) cation is square planar, with three coordination sites occupied by the chelating amine. The fourth site is filled by the nitrite ligand,

**Table 1** Single-crystal X-ray data for (1) in the ground state (before any irradiation) and metastable state (after irradiation with 400 nm LED light for 1 h)

|   | Ground state (GS)   | 400 nm metastable state (MS)   |
|---|---|--|
| Photo-conversion                          | 0%  | 100%   |
| Temperature                               | 100(2) K  | 100(2) K   |
| Wavelength                                | 0.71073 Å   | 0.71073 Å  |
| Empirical formula                         | C <sub>48</sub> H <sub>73</sub> B <sub>1</sub> N <sub>4</sub> O <sub>3</sub> Pd <sub>1</sub>  | C <sub>48</sub> H <sub>73</sub> B <sub>1</sub> N <sub>4</sub> O <sub>3</sub> Pd <sub>1</sub>   |
| Formula weight                            | 871.31 g mol <sup>-1</sup>  | 871.31 g mol <sup>-1</sup>   |
| Crystal size                              | 0.17 × 0.05 × 0.03 mm   | 0.17 × 0.05 × 0.03 mm  |
| Crystal system                            | Monoclinic  | Monoclinic   |
| Space group                               | P2 <sub>1</sub> /n  | P2 <sub>1</sub> /n   |
| Unit cell parameters (constrained)        | $a = 11.5457(4)$ Å,<br>$\alpha = 90^\circ$<br>$b = 13.4021(5)$ Å,<br>$\beta = 95.335(4)^\circ$<br>$c = 29.7173(12)$ Å,<br>$\gamma = 90^\circ$ | $a = 11.4775(4)$ Å,<br>$\alpha = 90^\circ$<br>$b = 13.3647(5)$ Å,<br>$\beta = 94.569(3)^\circ$<br>$c = 29.9849(8)$ Å,<br>$\gamma = 90^\circ$ |
| Volume                                    | 4578.4(3) Å <sup>3</sup>  | 4584.9(3) Å <sup>3</sup>   |
| Z   | 4   | 4  |
| Density (calculated)                      | 1.264 M g m <sup>-3</sup>   | 1.262 M g m <sup>-3</sup>  |
| Absorption coefficient $\mu$              | 0.451 mm <sup>-1</sup>  | 0.448 mm <sup>-1</sup>   |
| $F(000)$                                  | 1856  | 1856   |
| $R(\text{int})$                           | 0.0459  | 0.0661   |
| Completeness (to $\theta = 25.00^\circ$ ) | 0.998   | 0.998  |
| $R_1$ (observed data)                     | 0.0428  | 0.0428   |
| $wR_2$ (all data)                         | 0.0863  | 0.0901   |
| Reflections (independent)                 | 19 627 (9335)   | 31 062 (9355)  |



**Fig. 1** Single-crystal X-ray structure for the asymmetric unit of (1) showing the atomic arrangement in the ground state (GS), before any irradiation. Ellipsoids are set at 50% probability and hydrogen atoms are omitted for clarity (red dashed line = hydrogen bonding to THF solvent molecule).

which is monodentate and adopts solely nitro-( $\eta^1$ -NO<sub>2</sub>) coordination in the GS. The N(3)-H(3)…O(3) hydrogen bond, highlighted in Fig. 1, is the only strong intermolecular interaction in the asymmetric unit and does not involve the potentially isomerisable nitro-( $\eta^1$ -NO<sub>2</sub>) group.

Details of the packing arrangement in the GS structure of (1) are outlined in Fig. 2. Pd(II) cations are aligned along the crystallographic *b*-axis, with adjacent cations related by the 2<sub>1</sub>

screw axis symmetry. Although the view in Fig. 2 suggests that adjacent nitro-( $\eta^1$ -NO<sub>2</sub>) ligands point towards one another, they are not close in 3-D space due to the  $\frac{1}{2}$  translation along the *b*-direction. Each NO<sub>2</sub> group is surrounded on the opposite side by butyl chains within the same cation. The BPh<sub>4</sub> anions occupy positions near to the backbone of the triamine ligands, away from the bulky butyl groups. Finally, solvent THF molecules exist in discrete pockets that might reasonably be considered the result of inefficient packing between the other, bulkier components. Void space calculations conducted in Mercury,<sup>35</sup> with the THF solvent molecules removed, provide representations of these cavities (Fig. S2†). Using a contact surface calculation (probe radius 1.2 Å, grid spacing 0.1 Å) a total of 13.2% of the unit cell volume is found to comprise THF solvent. A solvent-accessible void calculation (with the same parameters) reduces this percentage to just 2.2%. Void space calculations conducted on the crystal structure as a whole, including THF, suggest no cavities are present in the GS and indicate a close-packed structure. Further quantification of the packing efficiency is possible by calculating a packing coefficient (see ESI† for method details). Using molecular volumes calculated by Hirschfeld Stockholder Partitioning in CrystalExplorer,<sup>36,37</sup> a packing coefficient of 0.98 is determined. Finally, three intermolecular close-contact interactions exist between nitro-( $\eta^1$ -NO<sub>2</sub>) and the surrounding crystal structure (Fig. S3†). Any intermolecular interactions involving nitro-( $\eta^1$ -NO<sub>2</sub>) in the GS must necessarily be overcome for linkage isomerism to occur in (1) and literature precedent suggests that strong intermolecular interactions can limit photo-activation.<sup>23</sup> However, given that these are all relatively weak intermolecular C-H…O contacts, they are not expected to cause a significant barrier to excitation.

### UV/vis spectroscopy

The UV/vis diffuse reflectance spectrum for (1) contains a single, broad maximum centred at approximately 300 nm, with a tail that extends to approximately 470 nm (Fig. S4†). The spectrum indicates that irradiation in the 200–470 nm range might induce a response. A wavelength in the tail of the UV/vis maxima was considered most suitable for excitation in the solid-state, in line with work by Enkelmann, Abdelmoty *et al.*<sup>38,39</sup> As such, a peak LED wavelength of 400 nm was used for all photocrystallographic experiments with (1).

### Steady-state photocrystallography

**Photoexcitation experiments.** The same crystal was held at 100 K, in the dark, and a series of steady-state photocrystallographic experiments were conducted *via* the methods outlined above. The purpose of this first series of photocrystallographic experiments was to ascertain both the maximum achievable level of photo-conversion in (1) and the rate of activation. The crystal was first irradiated with 400 nm LED light, for a period of 1 min. A subsequent data collection, conducted in the absence of further light, revealed that

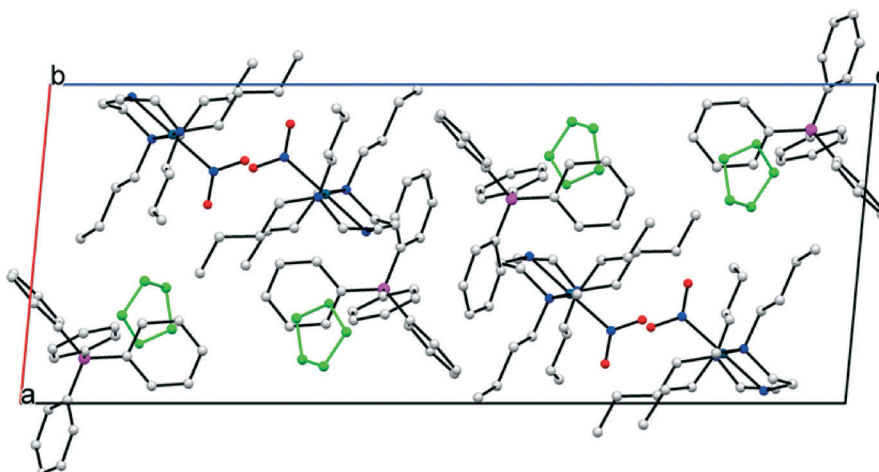


Fig. 2 Crystal packing diagram (one unit cell) for the GS structure of (1), before any irradiation. The packing diagram is viewed along the crystallographic *b*-axis, *c*-axis horizontal (hydrogen atoms omitted and THF solvent molecules highlighted in green for clarity).

a nitro–nitrito linkage isomerisation reaction had been induced, with 25% of the crystal now converted to an *endo*-nitrito-( $\eta^1$ -ONO) isomer. Similar data collections were then conducted for further irradiation times at intervals between 1 and 60 min. The refined GS and MS occupancies are provided in Table S1.<sup>†</sup>

Fig. 3 shows the plot of *endo*-nitrito-( $\eta^1$ -ONO) occupancy *vs.* irradiation time, for 0–15 min illumination. Remarkably, 100% photo-conversion is achieved in (1) after just 15 min. To the author's knowledge, this is the fastest reported example of a fully-complete single-crystal-to-single-crystal nitro–nitrito isomerisation reaction. Kinetic parameters for the photo-reaction can be obtained by application of the JMAK theory for solid-state kinetics<sup>22,40–45</sup> to the data points in Fig. 3:

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

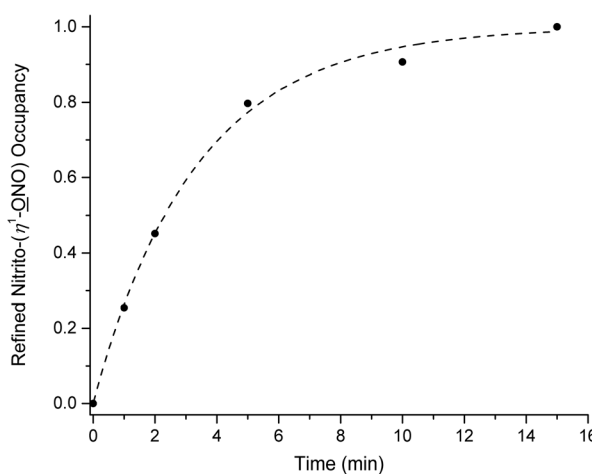


Fig. 3 Plot of crystallographically-refined nitrito-( $\eta^1$ -ONO) occupancy *vs.* irradiation time (excitation at  $\lambda = 400$  nm) for steady-state photo-crystallographic experiments with (1) between 0 and 15 min illumination. JMAK kinetic analysis:  $k = 0.30(2)$  min<sup>-1</sup>, (0.0049(3) s<sup>-1</sup>),  $n = 0.98(7)$ ,  $R^2 = 0.99$ .

where  $\alpha(t)$  is the conversion fraction [here the nitrito-( $\eta^1$ -ONO) occupancy] at a time  $t$ ,  $k$  is the rate constant and  $n$  is the Avrami constant. Least-squares fitting of eqn (1) to the data gives values of  $k = 0.30(2)$  min<sup>-1</sup> (0.0049(3) s<sup>-1</sup>) and  $n = 0.98(7)$ .

A value of  $n \approx 1$  indicates a homogeneous spread of nucleation sites for MS growth occurring throughout the crystal lattice.

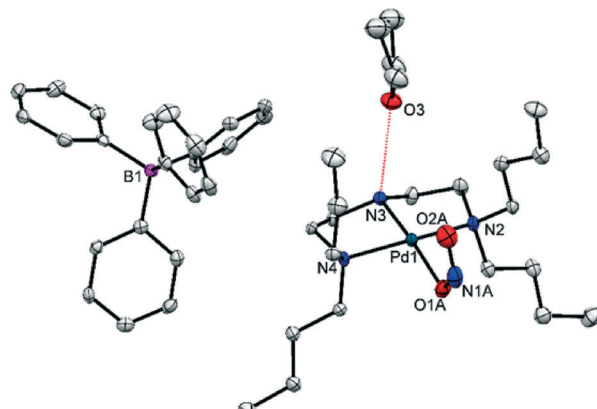
Extended illumination of the crystal for a total of 60 min induced no further change, indicating that a photostationary state had been reached after 15 min. The photo-excited crystal was held in the dark for a further 1 h and a subsequent data collection showed still no change, proving the excited state is metastable at 100 K. SC-XRD data for the MS structure are given in Table 1.

Variable temperature (VT) studies were then conducted on the same crystal, to determine the temperature range over which the system is metastable. The crystal was heated *in situ* and paused at intervals to allow for SC-XRD data collections. Full occupancy data for VT studies are provided in Table S2,<sup>†</sup> with selected data given in Table 2. (1) remains metastable between 100 and 200 K (Table S2<sup>†</sup>). The onset of MS decay occurs on warming to 220 K, with complete recovery of the GS by 240 K.

**Metastable state (MS) crystal structure.** Fig. 4 displays the asymmetric unit in the MS crystal structure (the full atomic numbering scheme for the cation is given in Fig. S5<sup>†</sup>). By comparison with Fig. 1 little change is apparent between the

Table 2 Refined nitro-( $\eta^1$ -NO<sub>2</sub>) and nitrito-( $\eta^1$ -ONO) occupancies for variable temperature studies conducted under steady-state and pseudo-steady-state conditions

| Temp/K | Steady-state                  |                  | Pseudo-steady-state           |                  |
|--------|-------------------------------|------------------|-------------------------------|------------------|
|        | ( $\eta^1$ -NO <sub>2</sub> ) | ( $\eta^1$ -ONO) | ( $\eta^1$ -NO <sub>2</sub> ) | ( $\eta^1$ -ONO) |
| 100    | 0.00                          | 1.00             | —                             | —                |
| 200    | 0.00                          | 1.00             | 0.00                          | 1.00             |
| 220    | 0.71                          | 0.29             | 0.00                          | 1.00             |
| 240    | 1.00                          | 0.00             | 0.00                          | 1.00             |
| 250    | 1.00                          | 0.00             | 0.53                          | 0.47             |
| 260    | 1.00                          | 0.00             | 1.00                          | 0.00             |



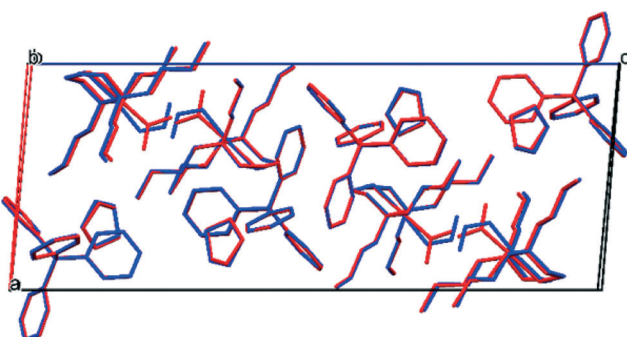
**Fig. 4** Single-crystal X-ray structure for the asymmetric unit of (1) showing the atomic arrangement in the metastable state (MS) after 1 h irradiation at  $\lambda = 400$  nm. Ellipsoids set at 50% probability and hydrogen atoms omitted for clarity (red dashed line = hydrogen bonding to THF solvent molecule).

GS and MS atomic arrangements, excepting the switch in nitrite coordination, and there is little evidence of crystal degradation. The crystal packing arrangement is also largely unchanged as a result of photo-activation (Fig. S6†). An overlay of GS and MS unit cells (Fig. 5) provides a pictorial comparison and the result of a Crystal Packing Similarity calculation in Mercury<sup>35</sup> shows a RMS deviation of 0.098 between structures. This is small and is largely the result of the change in nitrite coordination mode.

There is a small increase in the unit cell volume following 100% excitation, comprising  $6.5(4)$   $\text{\AA}^3$  or 0.14% of the GS volume. The changes are anisotropic: while the *c*-axis is found to lengthen by  $0.268(1)$   $\text{\AA}$ , the *a* and *b* axes contract by a smaller amount ( $\Delta a = -0.068(1)$   $\text{\AA}$  and  $\Delta b = -0.037(1)$   $\text{\AA}$ ). This asymmetry may suggest the changes are a result of the isomerisation, as nitro–nitrito rearrangement occurs approximately along *c*. However, the reduction in  $\beta$ -angle is the most significant, with  $\Delta\beta = -0.766(5)$ °.

### Pseudo-steady-state photococrystallography

Pseudo-steady-state experiments were next conducted to ascertain the response in (1) to continuous illumination at



**Fig. 5** Overlay of unit cell packing diagrams for the GS (red) and MS (blue) of (1), showing subtle changes in the unit cell as a result of excitation at  $\lambda = 400$  nm.

temperatures close to the MS limit (200–260 K). As the crystal is “pumped” by LEDs throughout the experiment, the pseudo-steady-state isomer ratio determined at temperatures above the MS limit describes an equilibrium situation between the forward nitro–nitrito photo-excitation and the reverse nitrito–nitro decay processes. A summary of the refined nitro-( $\eta^1\text{-NO}_2$ ) and nitrito-( $\eta^1\text{-ONO}$ ) occupancies under continuous LED irradiation ( $\lambda = 400$  nm) are given in Table 2. In contrast to the VT experiments conducted in the dark, under continuous illumination 100% excitation is maintained up to 240 K and 47% excitation is still determined at 250 K. By 260 K no evidence of the excited state nitrito-( $\eta^1\text{-ONO}$ ) isomer could be determined and the SC-XRD refinement indicated 100% occupancy of the GS nitro-( $\eta^1\text{-NO}_2$ ) isomer, in the time-frame of the experiment.

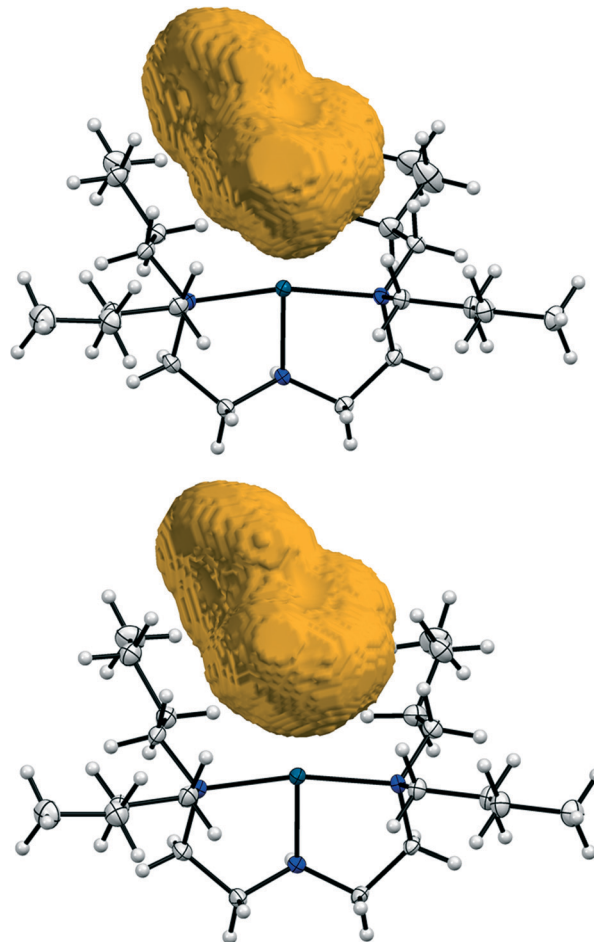
## Discussion

### Photo-activation level

(1) is the first reported example of complete, 0–100% MS photo-activation in the single-crystal for a Pd(II)-nitrite linkage isomer. Previously, nitrito-( $\eta^1\text{-ONO}$ ) occupancies of 46% and 39%, in  $[\text{Pd}(\text{PPh}_3)_2(\text{NO}_2)_2]$  and  $[\text{Pd}(\text{AsPh}_3)_2(\text{NO}_2)_2]$  respectively, were the highest reported MS photo-conversion levels for Pd(II) systems starting from a clean, 100% nitro-( $\eta^1\text{-NO}_2$ ) GS.<sup>21</sup> A photo-induced, 100% nitrito-( $\eta^1\text{-ONO}$ ) isomer is also reported in  $[\text{Pd}(\text{NO}_2)(\text{C}^{\text{N}})(\text{PPh}_3)]$  ( $\text{C}^{\text{N}} = \text{papaverine}$ ) in 2012,<sup>28</sup> however this MS arrangement was produced from a GS structure that already contained 77% of the nitrito-( $\eta^1\text{-ONO}$ ) ligand, meaning the maximum GS-MS conversion in this case was 23%.

In light of previous work, the  $\text{Bu}_4\text{dien}$  ancillary ligand and  $\text{BPh}_4^-$  counterion are chosen specifically for their high steric demands. Analysis of the crystal packing arrangement (Fig. 2, above) confirms that these bulky, photo-inert fragments dominate the structure and outline the reaction cavity within which nitro–nitrito isomerisation can occur. Fig. 6 provides a pictorial representation of the reaction cavity space available for the nitrite ligand in the GS and MS structures. Reaction cavity volumes were determined in Mercury<sup>35</sup> by removing the atoms of the nitrite ligand and conducting a contact surface void space calculation. The results of these reaction cavity calculations are given in Table 3. Visually, there is little change in reaction cavity size between the GS and MS arrangements, while the calculations indicate only a very small change in the reaction cavity volume,  $\Delta V_c$ , of  $+1.46$   $\text{\AA}^3$  (0.05% unit cell volume) per molecule. In line with the Topochemical Postulate outlined by Schmidt and Cohen,<sup>2–4</sup> it is expected that solid-state reactions requiring the least change in the crystalline environment will proceed to highest completeness. As such, the requirement for only small  $\Delta V_c$  on excitation is likely to facilitate the high level of photo-activation achieved in (1). Additionally, the nitrite ligand is not involved in strong hydrogen bonds in the GS or MS and only weaker C–H···O interactions must be broken and re-formed on excitation (Fig. S3†). As stronger intermolecular contacts are expected to raise the





**Fig. 6** Reaction cavity volumes: top = GS cavity, bottom = MS cavity. Reaction cavity volumes are determined by removing the atoms of the nitrite ligand before conducting a contact surface void calculation in Mercury<sup>35</sup> (probe radius 1.2 Å, grid spacing 0.1 Å).

**Table 3** Reaction cavity volumes calculated in Mercury<sup>35</sup>

| Cavity size<br>(per unit cell, u.c) | Cavity size/Z<br>(per molecule) |       | $\Delta V_c$<br>(per molecule) |       |
|-------------------------------------|---------------------------------|-------|--------------------------------|-------|
|                                     | Vol (Å <sup>3</sup> )           | % u.c | Vol (Å <sup>3</sup> )          | % u.c |
| GS                                  | 198.36                          | 4.30  | 49.59                          | 1.08  |
| MS                                  | 204.20                          | 4.50  | 51.05                          | 1.13  |

| Vol (Å <sup>3</sup> ) | % u.c | Vol (Å <sup>3</sup> ) | % u.c | Vol (Å <sup>3</sup> ) | % u.c |
|-----------------------|-------|-----------------------|-------|-----------------------|-------|
| 198.36                | 4.30  | 49.59                 | 1.08  | 1.46                  | 0.05  |
| 204.20                | 4.50  | 51.05                 | 1.13  |                       |       |

energy barrier to linkage isomerism, this also rationalises the high photo-conversion level in (1).

In addition to steric factors, the absorption properties of (1) will likely contribute to the overall activation level. The choice of an excitation wavelength in the tail of the UV/vis profile is expected to help maximise conversion.<sup>38,39</sup> 400 nm light should have sufficient energy to promote a response, but at the same time not be too strongly absorbed in outer layers of the crystal, allowing light to penetrate more evenly throughout the bulk.

#### Reaction rate and speed of response

The photo-induced linkage isomerisation reaction in (1) reaches completion after just 15 min irradiation. This is

remarkably fast, in comparison to the reaction rates reported for similar photo-reactions in single-crystals. Other systems displaying 100% nitro-nitrito conversion, including  $[\text{Ni}(\text{dppe})(\text{NO}_2)\text{Cl}]$ ,<sup>19</sup>  $[\text{Ni}(\text{dppe})(\text{NO}_2)_2]$ ,<sup>24</sup>  $[\text{Ni}(\text{dcpe})(\text{NO}_2)_2]$  (dcpe = bis-dicyclohexyl-phosphinoethane)<sup>24</sup> and  $[\text{Ni}(\text{Et}_4\text{dien})(\text{NO}_2)_2]$  ( $\text{Et}_4\text{dien}$  = *N,N,N',N'*-tetraethylidienetriamine)<sup>22</sup> achieve maximum conversion in 90 min, 60 min, 60 min and 120 min, respectively. It is possible that the butyl substituents in the triamine co-ligand may facilitate faster linkage isomer switching in (1). These butyl chains are expected to be relatively flexible, and so may help to disappate any strain induced in the lattice as a result of linkage isomerism. This should reduce the nitro-nitrito energy barrier and so promote a faster reaction. Such a hypothesis is difficult to quantify by crystallographic data alone, although qualitative support is gained on considering the shape of the GS and MS reaction cavities (Fig. 6). Although there is little overall change in reaction cavity size between the GS and MS, close inspection shows some subtle variation in the shape of each cavity, particularly in the regions surrounded by the butyl substituents, indicating some degree of flexibility.

#### Metastable limit and near ambient-temperature functionality

Under steady-state conditions, the onset of MS decay in (1) occurs around 220 K and this MS limit is comparable to other reported Pd(II)-nitrite linkage isomers,<sup>21</sup> although these do not achieve 100% activation. Other high-converting Ni(II)-nitrite systems generally display lower MS limits. Previous reports highlight the trend for complexes with heavier transition metal (TM) centres to display higher MS limits, suggesting a link between the reduced kinetic lability of the TM(II)-ligand bond with heavier metals and the stability of the MS isomer.<sup>21</sup>

More interestingly, SC-XRD experiments conducted under pseudo-steady-state conditions allow full characterisation of the excited state nitrito-( $\eta^1$ -ONO) isomer with 100% occupancy at 240 K (-33 °C) and 47% occupancy as high as 250 K (-23 °C). These temperatures are approaching ambient conditions and are the highest reported operating temperatures for a Pd(II)-nitrite linkage isomer switch. In contrast to the response under continuous illumination, no evidence of the excited state *endo*-nitrito isomer is found at 240 K and above when the excitation source is removed. As such, complex (1) shows potential as a light-on, functionality-on/light-off, functionality-off molecular switch in the 240–250 K region. Given that the largest difference occurs at 240 K (light on = 100%, light off = 0%) this would likely be the most suitable operating temperature. Before any such device could be considered practicable however, it is important to ascertain the speed of excitation and decay in the crystal at 240 K. Unfortunately, given that the photo-reaction is essentially dynamic at this temperature, it was not possible to conduct full SC-XRD experiments to determine the change in nitro:nitrito ratio with time in this case. Instead, an estimation of the isomerisation reaction progress was made by following the change in unit cell parameters, as this approach required

only a few frames of X-ray data per time-point. These kinetic experiments were completed at 240 K for both the excitation (light on) and decay (light off) processes. The results of these experiments are given in Fig. 7. Only the  $\beta$ -angle showed a clear trend under both conditions, which might be expected considering that this is the parameter which changes the most on photo-activation under steady-state conditions (Table 1). As expected, a clear reduction in the  $\beta$ -angle is observed on photo-excitation and a corresponding increase can be seen with excited state decay. Unfortunately, it was not possible to obtain reliable kinetic parameters from these data by least-squares fitting of eqn (1), given the low level of accuracy in the measurements. However, it does appear that both the excitation and decay processes reach completion in approximately 15 min, a similar time to the excitation process followed at 100 K. These experiments provide a strong starting point for time-resolved synchrotron powder X-ray diffraction (PXRD) experiments with (1) at 240 K, for which very fast data collection times would allow many and more accurate data points to be determined for each reaction profile.

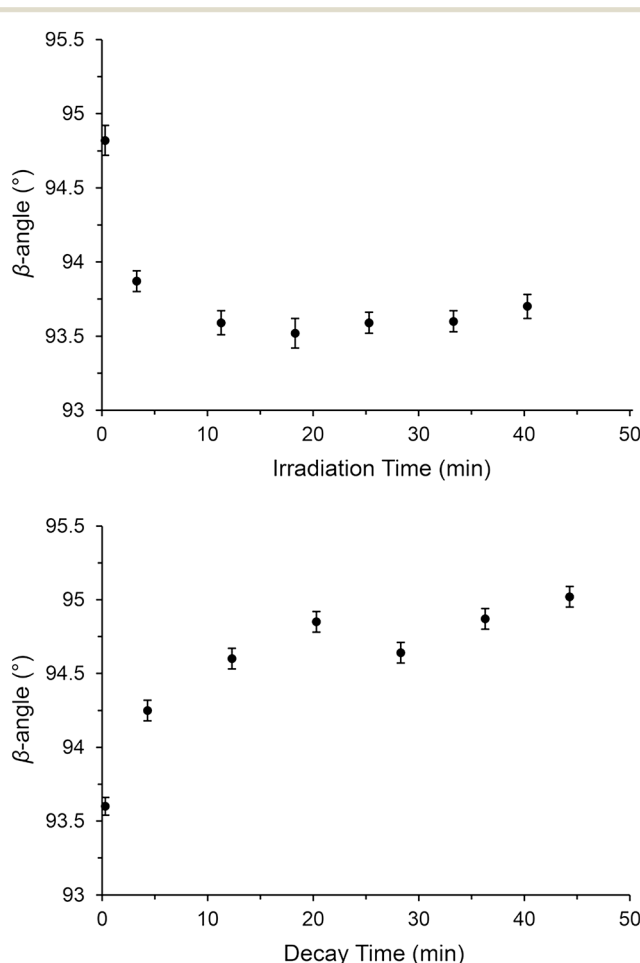


Fig. 7 SC-XRD kinetics experiments at 240 K, following the change in  $\beta$ -angle as a result of nitrite linkage isomerism. Top: Plot of  $\beta$ -angle vs. irradiation time under continuous illumination ( $\lambda = 400$  nm). Bottom: Plot of  $\beta$ -angle vs. decay time in the dark.

## Conclusions

The observation of 100%, fully-reversible linkage isomerism in the Pd( $\eta$ )-nitrite complex (1) is encouraging, as it proves that complete nitro–nitrito conversion in the single-crystal is not limited to complexes containing first-row TM centres. Detailed analysis of the packing effects in both GS and MS structures once again validates the approach of including sterically-demanding ancillary fragments into the crystal system, to facilitate high levels of photo-conversion. In addition, analysis of the size and shape of the reaction cavity provides both a qualitative and quantitative indication of changes in the immediate location of the nitrite ligand as a result of linkage isomer switching, which also supports the hypothesis that bulky photo-inert fragments help the single-crystal-to-single-crystal isomerisation reaction to proceed topotactically.

While steady-state photocrystallographic experiments provide important, accurate and detailed information on the MS structure, pseudo-steady-state SC-XRD experiments are equally important to understand the progress of linkage isomer switching at temperatures close to the MS limit. The determination of a photo-excited *endo*-nitrito-( $\eta^1$ -ONO) isomer by SC-XRD as high as 240–250 K in (1) is promising evidence that linkage isomer switching is approaching viable operating temperatures. The result is suggestive that the design of linkage isomer systems containing either heavier TM centres, or less kinetically labile ambidentate ligands, might offer a new approach to maximising MS limits, and further investigation in this area would be profitable.

There remain a number of key challenges in the development of viable solid-state linkage isomer devices and future work with (1) will focus on obtaining more detailed understanding of both the rate of GS–MS switching at relevant temperatures and the reproducibility of this process over extended operating periods.

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