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## Studying the cellular distribution of highly phototoxic platinated metalloporphyrins using isotope labelling†

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Novel tetraplatinated metalloporphyrin-based photosensitizers (PSs) are reported, which show excellent phototoxic indexes (PIs) up to 5800 against HeLa cells, which is, to the best of our knowledge, the highest value reported for any porphyrin so far. Furthermore, <sup>67</sup>Zn isotope labelling allowed the determination of the ratio of zinc to platinum inside the cells using ICP-MS.

PDT is a clinically approved and minimally invasive therapeutic treatment for dermatological, ophthalmic, and infectious diseases, as well as cancer, which involves the application of a photosensitizer (PS) that ideally localizes disease-specifically. The application is followed by low energy irradiation to activate the PS leading to the generation of reactive oxygen species (ROS) which in turn induce cell death. A number of publications have reported about PSs conjugated with a chemotoxic unit, e.g., a cisplatin analogue.<sup>2</sup> While this approach normally renders the conjugate more toxic,3 it might diminish one of the main advantages of PDT, namely the special selectivity by the applied light source.

Previously, we have reported a series of easily accessible tetrapyridyl-porphyrins (tPt-H<sub>2</sub>4PyP, cPt-H<sub>2</sub>4PyP, dPt-H<sub>2</sub>4PyP) that were coordinated by four platinum(II) complexes to yield highly phototoxic agents (Fig. 1). These compounds displayed a

light toxicity down to 19 nM and at the same time low dark

Based on our previous studies, 1c,4 we investigated a set of important follow-up questions:

- (1) How does the insertion of either zinc or copper<sup>1c</sup> modulate the phototoxicity of tPt-H<sub>2</sub>4PyP, cPt-H<sub>2</sub>4PyP and dPt-H<sub>2</sub>4PyP? The singlet oxygen quantum yield of porphyrins was reported to increase or not to change at all after insertion of zinc into the macrocycle. In another case, the metal-free porphyrin generated mainly free hydroxy radicals, which were in vitro more cytotoxic than the singlet oxygen produced by the zinc porphyrin.8
  - (2) What is the fate of the complex tPt-H<sub>2</sub>4PvP in the cell?
- (3) What is the reason for the low dark toxicities of our platinum-porphyrin conjugates?

cPt-H<sub>2</sub>4PyP: M = H<sub>2</sub>

cPt-Zn4PyP: M = Zn

tPt-H<sub>2</sub>4PvP: M = H<sub>2</sub> H<sub>3</sub>N

Fig. 1 Studied tetraplatinated porphyrins.

toxicities of around 45 µM in human cancer cell lines. 4 The ratio of dark to light toxicity is referred to as the phototoxic index (PI), and it should be as high as possible. Our systems displayed PI values higher than 1000 in HeLa cells after being excited with light of a wavelength of 420 nm. These PI values are better than currently clinically used second generation PSs that display a phototoxicity of > 260.5

tPt-Zn4PyP: M = Zn tPt-<sup>67</sup>Zn4PyP: M = <sup>67</sup>Zn cPt-Cu4PvP: M = Cu tPt-Cu4PyP: M = Cu

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Fig. 2 Ellipsoidal plot of the crystal structure of [trans-PtCl(NH<sub>3</sub>)<sub>2</sub>]<sub>4</sub>-5,10,15,20-tetra(4'-pyridyl)-zinc(||)porphyrin tetraphenylborate-(DMF)7. Hydrogen atoms attached to a carbon atom, non-coordinated anions and DMF molecules not coordinated to the zinc metal center were omitted for clarity

# (4) What is the origin of the slow reaction of tPt-H<sub>2</sub>4PyP with

To study the effect of a metal inside the center of the porphyrins tPt-H<sub>2</sub>4PyP, cPt-H<sub>2</sub>4PyP and dPt-H<sub>2</sub>4PyP, the novel complexes tPt-Zn4PyP, tPt-Cu4PyP, cPt-Zn4PyP, cPt-Cu4PyP, dPt-Zn4PyP and dPt-Cu4PyP were synthesized following our previously published procedures. 4 Subsequently, these complexes were characterized using 1H- and 195Pt-NMR, IR, UV-Vis, MS and elemental analysis (ESI†). Additionally, the cationic unit of tPt-Zn4PyP could be crystallized as the tetraphenylborate salt using the under-oil technique (ESI†) and was studied using X-ray analysis (Fig. 2). We were particularly pleased to obtain this crystal structure, as it is the first example of expanding our robotic, aqueous crystallization trials9 towards organic solvents. The zinc metal center is in a square pyramidal geometry with one DMF molecule as the fifth ligand. This is only the third crystallographic report about a porphyrin with an exocyclic, nonorganometallically bound platinum within the porphyrin plane.<sup>10</sup>

The compounds were then tested in the cancerous HeLa cell line and the non-cancerous MRC-5 cell line composed of fibroblasts derived from lung tissue to assess their antiproliferative properties. The results of the cytotoxicity studies of the compounds are presented in Table 1. The insertion of zinc into the complex tPt-H24PyP to yield the zinc complex tPt-Zn4PyP improves the phototoxicity and lowers the dark toxicity at the same time. Together these two effects improve the PI by almost a factor of five. These results can be attributed to the higher singlet oxygen quantum yield (determined as described in ref. 11) of tPt-Zn4PyP compared to tPt-H<sub>2</sub>4PyP (Table S3, ESI†) and are in line with previous reports about other (metallo)porphyrin containing photosensitizers. 6a,b,d

However, the zinc complexes cPt-Zn4PyP and dPt-Zn4PyP are less phototoxic than their corresponding metal-free porphyrins cPt-H<sub>2</sub>4PyP and dPt-H<sub>2</sub>4PyP. This difference in behaviour can be explained for cPt-Zn4PyP. After the insertion of zinc, the complex becomes rather photosensitive, which leads to photobleaching under irradiation and therefore to a lower singlet oxygen quantum yield (Table S3, ESI†). The higher phototoxic indexes of complexes tPt-H24PyP, tPt-Zn4PyP and tPt-Cu4PyP compared with cPt-H<sub>2</sub>4PyP, cPt-Zn4PyP and cPt-Cu4PyP, respectively, are further explained below. Furthermore, tPt-Cu-4PyP is, to the

Table 1 Anti-proliferative effects of the various compounds on noncancerous MRC5 and cancerous HeLa cells in the dark and upon irradiation

Compound	IC <sub>50</sub> MRC5 dark (μM)	50	IC <sub>50</sub> HeLa 420 nm (μM)	PI (HeLa)	Ref.
Photofrin Transplatin (tPt) tPt-H <sub>2</sub> 4PyP tPt-Zn4PyP tPt-Cu4PyP Cisplatin (cPt) cPt + H <sub>2</sub> 4PyP (1:1) cPt-H <sub>2</sub> 4PyP cPt-Zn4PyP cPt-Zn4PyP cPt-Cu4PyP cisPtDMSO (dPt) dPt-H <sub>2</sub> 4PyP	$\begin{array}{c} \text{n. d.} \\ 86^{\pm 8} \\ 93.4^{\pm 6.5} \\ > 100 \\ 82.5^{\pm 17.5} \\ 7.9^{\pm 1.2} \\ > 100 \\ 50.2^{\pm 0.6} \\ 65.0^{\pm 0.6} \\ > 100 \\ > 100 \\ > 100 \\ > 100 \end{array}$	$\begin{array}{c} >41 \\ \sim 130 \\ 44.9^{\pm 8.0} \\ >100 \\ 56.4^{\pm 6.5} \\ 11.5^{\pm 2.9} \\ >100 \\ 35.4^{\pm 4.4} \\ 17.1^{\pm 5.1} \\ 13.9^{\pm 1.6} \\ >100 \\ >100 \end{array}$	$\begin{array}{c} 4.3^{\pm 0.2} \\ \text{n. d.} \\ 0.037^{\pm 0.02} \\ 0.017^{\pm 0.004} \\ 14.1^{\pm 6.3} \\ 22.3^{\pm 5.7} \\ 5.78^{\pm 1.9} \\ 0.054^{\pm 0.01} \\ 0.25^{\pm 0.02} \\ > 100 \\ > 100 \\ 0.15^{\pm 0.02} \end{array}$	>9.5 n. d. 1210 >5882 4.0 0.52 >17 655 68.4 <0.14 n. a. >680	t. w. 12/13 4 t. w. t. w. 4 4 t. w. t. w. t. w.
dPt-Zn4PyP dPt-Cu4PyP	>100 >100	$> 100 \ 61.4^{\pm 1.5}$	$1.28^{\pm0.4}$ > 100	>78 <0.61	t. w. t. w.

PI = phototoxic index, n. d. = not determined, t. w. = this work, n. a. = not applicable.

best of our knowledge, only the second ever reported copper containing PS, 1c which is phototoxic, as normally copper porphyrinoids are non-phototoxic. The DMSO containing complexes dPt-H<sub>2</sub>4PyP and dPt-Zn4PyP have comparable PIs to the cisplatin derived complexes cPt-H24PyP and cPt-Zn4PyP. For a careful discussion on the influence of DMSO upon any platinum containing drug, the reader is directed to the excellent study by Gottesman.14 The singlet oxygen quantum yields  $(\Phi_{\Delta})$  of **tPt-H<sub>2</sub>4PyP**, **cPt-H<sub>2</sub>4PyP**, tPt-Zn4PyP and cPt-Zn4PyP are shown in Table S3 (ESI†).

As previously reported by us, tPt-H24PyP reacts extremely slowly with guanosine, as the platination of N7 progressed to less than 50% completion within 10 days. We speculate that the high charge (4+) of the cation of tPt-H24PyP might be responsible for this slow reaction, since the platinum chloride bond must be hydrolysed before the platinum can react with N7 of guanosine. The hydrolysis yields an intermediate that possesses an even higher charge of +5 (Fig. S1, ESI†). In order to test this hypothesis, we synthesized trans-[Pt(NH<sub>3</sub>)<sub>2</sub>(pyridine) Cl]NO<sub>3</sub> (tPt-Py)<sup>15</sup> and repeated the earlier mentioned reaction with guanosine (Fig. S2, ESI†). The reaction of tPt-Py and one equivalent of guanosine was monitored using <sup>1</sup>H-NMR spectroscopy in a 1:1 DMF-d<sub>7</sub>:D<sub>2</sub>O mixture (see ESI†), showing that 50% of the binding occurred after nearly three days. Compared to previous results of tPt-H24PvP, the reaction of tPt-Pv is at least three times faster. This supports the assumption that the rather slow reaction rate of tPt-H24PyP with guanosine is caused by unfavourable electrostatic interactions. Since guanosine and tPt-Py react with a 1:1 stoichiometry, the assumption was made that the reaction can be treated as a second order reaction (see ESI†). The determined slope S is equal to a second order reaction constant k of 0.0034  $\pm$  0.0001 M<sup>-1</sup> s<sup>-1</sup>. This value compares well with the pseudo-first order reaction rate  $(k_1 \text{ of } 0.00036 \pm 0.00002 \text{ s}^{-1})$  for the reaction of the mono aquated phenanthriplatin with 9-methylguanine. 16 If one takes the concentration of the excess 9-methylguanine into account, one can convert the pseudo-first order reaction rate into a second order rate ( $k_2$  of 0.012  $\pm$  0.007 M<sup>-1</sup> s<sup>-1</sup>). This rate is

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about 3.5 times faster than the determined rate of tPt-Py with guanosine, which is reasonable since tPt-Py must first hydrolyse before it can react with guanosine.

Next, we investigated the uptake and distribution of the Zn-Pt complex in HeLa cells using ICP-MS. The expected ratio of zinc to platinum is 1:4 in tPt-Zn4PyP, provided the compound does not disintegrate. To study the Zn and Pt uptake, we first synthesized a highly enriched isotope <sup>67</sup>Zn complex tPt-67Zn4PyP. 17 Using the highly enriched zinc isotope makes it possible to differentiate between naturally occurring zinc and 67Zn18 that was introduced into the cells with compound tPt-67Zn4PyP. As 67Zn is expensive, initially, a careful and lengthy series of optimizations of the synthesis of zinc-5,10,15,20-tetra-(4'-pyridyl)-porphyrin (Zn4PyP) starting from elemental zinc with a natural distribution of isotopes was performed (Table S2, ESI†). Using these optimized conditions, the synthesis of tPt-Zn4PyP was then repeated starting from elemental <sup>67</sup>Zn, which delivered complex tPt-<sup>67</sup>Zn4PvP (Fig. S5-S10, ESI†). The uptake study shows that only one quarter of all applied platinum atoms entered the HeLa cells, whereas three quarters of <sup>67</sup>Zn atoms entered the cells (Fig. S13 and S14, ESI†). Subsequently, we studied the distribution of platinum and 67Zn in the different cell compartments (Table S4, ESI†). In the nucleus, the ratio of <sup>67</sup>Zn to Pt was found to be 3:1 instead of 1:4. These results clearly demonstrate that the complex tPt-67Zn4PvP is disintegrated into the <sup>67</sup>Zn containing porphyrin and the relatively non-toxic monoactivated transplatin. 19 The analogous complex tPt-Zn4PyP with the natural Zn isotope distribution is expected to behave in the same way as tPt-67Zn4PyP. The transplatin moiety helps to improve solubility and transport the PS into the cell while not being toxic after the cleavage of the conjugate. Transplatin is known to be effluxed faster from the cell than cisplatin.<sup>20</sup> This explains why there is less platinum found inside of the cells than <sup>67</sup>Zn.

We also investigated the differences between the dark cytotoxcities of the transplatinated series of compounds, tPt-H<sub>2</sub>4PyP, tPt-Zn4PyP and tPt-Cu4PyP and the cisplatinated series, cPt-H24PvP, cPt-Zn4PvP and cPt-Cu4PvP. The transplatinated series generates, as described before, the transdiamminemonochloroplatinum(II) moiety<sup>21</sup> after breakage of the N(pyridine)-platinum bond. This fragment has a much lower cytotoxcity than cis-diamminemonochloroplatinum(II), which is the first intermediate of cisplatin that must be generated in order to achieve any biological impact of cisplatin. Both, fragments of *cis*- and *trans*-diamminechloridoplatinum(II) in the conjugates c/tPt-H<sub>2</sub>4PyP, c/tPt-Zn4PyP and c/tPt-Cu4PyP serve as a cancer-cell penetrating vector. However, the trans moiety has a much lower dark toxicity after hydrolysis. The cis-diamminemonochloroplatinum(II) fragment has been previously used as a chemotoxic part of a dual acting reagent.<sup>2b</sup> Our results imply that the trans-diamminemonochloroplatinum(II) conjugate would be even more interesting to study as well, maybe yielding a photosensitizer with an improved PI. Next, we examined whether the compounds induce damage to the DNA. It is established, that platinum drugs form drug-DNA adducts,

leading to DNA damage, which triggers cell cycle arrest and DNA repair.<sup>22</sup> The drug-induced DNA damage is followed by the phosphorylation of the histone H2AX, which is involved in the recruiting and localizing of the DNA damage repair proteins.

Therefore, YH2AX is considered an important marker to observe DNA damage, in particular double strand breaks.<sup>23</sup> To determine the effectiveness of light-triggered DNA damage by the metal porphyrin conjugates, we treated HeLa cells with 500 nM tPt-H<sub>2</sub>4PyP and tPt-Zn4PyP for 14 h and subsequent light irradiation. Cells were then stained with \( \gamma H2AX-specific \) Alexafluor 594 antibody and visualized by confocal microscopy.

As shown in the immunofluorescent images (Fig. 3 and Fig. S15, ESI†), we could observe that tPt-H<sub>2</sub>4PvP and tPt-Zn4PvP both induced severe DNA damage upon light irradiation, as observed by the increased accumulation of γH2AX. This means that tPt-H<sub>2</sub>4PyP and tPt-Zn4PyP generate significantly more DNA damage after light irradiation compared with the experiments in the dark.

To assess the intracellular localization of the photosensitizers, the compounds were co-stained with a marker for the Golgi organelle. tPt-Zn4PvP co-localized in the Golgi apparatus (Fig. 4). This experimental finding is pretty inspiring, as some reported drugs directed against the Golgi complex have been shown to be effective in both androgen-dependent and androgen-independent prostate cancer, through targeting abnormal glycosylation.24

In summary, the insertion of a zinc cation into the phototoxic photosensitizer tPt-H24PvP vielded an even superior PS tPt-Zn4PyP with a phototoxic index of higher than 5880. This is likely due to the higher singlet oxygen quantum yield  $(\Phi_{\Lambda})$  of the zinc porphyrin platinum complex tPt-Zn4PyP compared to the free-base porphyrin platinum complex tPt-H24PyP. The <sup>1</sup>H-NMR experiments to follow the kinetics of the reaction of trans-[Pt(NH<sub>3</sub>)<sub>2</sub>(pyridine)Cl]NO<sub>3</sub> (tPt-Py) with guanosine revealed that the monocation trans-[Pt(NH<sub>3</sub>)<sub>2</sub>(pyridine)Cl]<sup>+</sup> reacts faster than the tetracation tPt-H24PyP. The tetracation has to overcome the barrier of generating an even five-fold charged state during the reaction with N7 of the guanine base. ICP-MS

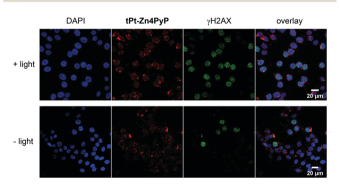


Fig. 3 DNA damage induced by light activated photosensitizer tPt-Zn4PyP. Immunofluorescent images of HeLa cells treated with 500 nM tPt-Zn4PyP for 14 h, then treated with light compared to the non-irradiated control. Cells were stained with yH2AX-Alexafluor 594 antibody (green) and DAPI. **tPt-Zn4PyP**: 458 nm ex., 630–750 nm em.; γH2AX-Alexafluor 594:594 nm

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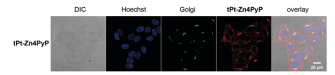


Fig. 4 Co-localization of tPt-Zn4PyP with the Golgi apparatus. Representative images of live-cell imaging of HeLa cells treated with 50  $\mu$ M **tPt-**Zn4PyP and stained with GOLGI-ID® Green dye (ENZO ENZ-51028).

studies of the <sup>67</sup>Zn labelled complex tPt-<sup>67</sup>Zn4PyP showed that the trans-diamminemonochloroplatinum(II) units of the complex tPt-Zn4PyP are dissociated from the conjugate inside the HeLa cells. For the first time, such a <sup>67</sup>Zn labelled conjugate was employed to monitor the fate of the zinc conjugate within cells. The trans-diamminemonochloroplatinum(II) unit is thought to act as a cell-penetrating vector for the photosensitizer, which permits its activation inside the cancer cell by light. We could demonstrate that within the cell, the PS co-localizes with the Golgi apparatus and additionally, induces DNA damage in the nucleus. Unlike conjugates with a cisplatin, transplatinphotosensitizer conjugates have no increased dark toxicity. Therefore, we propose the further investigation of transplatin-drug conjugates for PDT and other oncological treatments.

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#### Conflicts of interest

There are no conflicts to declare.

#### References

1 (a) T. J. Dougherty, C. J. Gomer, B. W. Henderson, G. Jori, D. Kessel, M. Korbelik, J. Moan and Q. Peng, J. Natl. Cancer Inst., 1998, 90, 889–905; (b) P. Agostinis, K. Berg, K. A. Cengel, T. H. Foster, A. W. Girotti, S. O. Gollnick, S. M. Hahn, M. R. Hamblin, A. Juzeniene, D. Kessel, M. Korbelik, J. Moan, P. Mroz, D. Nowis, J. Piette, B. C. Wilson and J. Golab, Ca-Cancer J. Clin., 2011, 61, 250-281; (c) P. M. Antoni, A. Naik, I. Albert, R. Rubbiani, S. Gupta, P. Ruiz-Sanchez, P. Munikorn, J. M. Mateos, V. Luginbuehl, P. Thamyongkit, U. Ziegler, G. Gasser, G. Jeschke and B. Spingler, Chem. - Eur. J., 2015, 21, 1179-1183; (d) S. Mallidi, S. Anbil, A.-L. Bulin, G. Obaid, M. Ichikawa and T. Hasan, Theranostics, 2016, 6, 2458-2487; (e) D. van Straten, V. Mashayekhi, H. S. de Bruijn, S. Oliveira and D. J. Robinson, Cancers, 2017, 9, 19.

- 2 (a) C. Lottner, K.-C. Bart, G. Bernhardt and H. Brunner, J. Med. Chem., 2002, 45, 2064-2078; (b) J. Mao, Y. Zhang, J. Zhu, C. Zhang and Z. Guo, Chem. Commun., 2009, 908-910; (c) G. Klein Couto, J. C. Rodrigues, B. S. Pacheco, L. Damé Simões, J. D. Paschoal, F. K. Seixas, T. V. Acunha, B. A. Iglesias and T. Collares, *Photodiagn*. Photodyn. Ther., 2020, 31, 101942; (d) G. K. Couto, B. S. Pacheco, V. M. Borba, J. C. Rodrigues, T. L. Oliveira, N. V. Segatto, F. K. Seixas, T. V. Acunha, B. A. Iglesias and T. Collares, J. Photochem. Photobiol., B, 2020, 202, 111725.
- 3 V. Ramu, S. Gautam, P. Kondaiah and A. R. Chakravarty, Inorg. Chem., 2019, 58, 9067-9075.
- 4 A. Naik, R. Rubbiani, G. Gasser and B. Spingler, Angew. Chem., Int. Ed., 2014, 53, 6938-6941.
- 5 J. Berlanda, T. Kiesslich, V. Engelhardt, B. Krammer and K. Plaetzer, J. Photochem. Photobiol., B, 2010, 100, 173-180.
- 6 (a) D. Praseuth, A. Gaudemer, J.-B. Verlhac, I. Kraljic, I. Sissoëff and E. Guillé, Photochem. Photobiol., 1986, 44, 717-724; (b) K. Wang, C. T. Poon, C. Y. Choi, W.-K. Wong, D. W. J. Kwong, F. Q. Yu, H. Zhang and Z. Y. Li, J. Porphyrins Phthalocyanines, 2012, 16, 85-92; (c) X. Hu, K. Ogawa, S. Li, T. Kiwada and A. Odani, Chem. Lett., 2017, 46, 764-766; (d) P. J. Gonçalves, F. C. Bezzerra, A. V. Teles, L. B. Menezes, K. M. Alves, L. Alonso, A. Alonso, M. A. Andrade, I. E. Borissevitch, G. R. L. Souza and B. A. Iglesias, J. Photochem. Photobiol., A, 2020, 391, 112375.
- Pavani, A. F. Uchoa, C. S. Oliveira, Y. Iamamoto and M. S. Baptista, Photochem. Photobiol. Sci., 2009, 8, 233-240.
- 8 J. M. Dabrowski, B. Pucelik, M. M. Pereira, L. G. Arnaut and G. Stochel, J. Coord. Chem., 2015, 68, 3116-3134.
- 9 (a) P. P. Nievergelt, M. Babor, J. Čejka and B. Spingler, Chem. Sci., 2018, 9, 3716-3722; (b) M. Babor, P. P. Nievergelt, J. Čejka, V. Zvoníček and B. Spingler, *IUCrJ*, 2019, 6, 145–151; (c) R. Alvarez, P. P. Nievergelt, E. Slyshkina, P. Müller, R. Alberto and B. Spingler, Dalton Trans., 2020, 49, 9632-9640.
- 10 (a) A. K. Bar, R. Chakrabarty, G. Mostafa and P. S. Mukherjee, Angew. Chem., Int. Ed., 2008, 47, 8455-8459; (b) P. Lang, J. Habermehl, S. I. Troyanov, S. Rau and M. Schwalbe, Chem. - Eur. J., 2018, 24, 3225-3233.
- 11 L. Schneider, M. Larocca, W. Wu, V. Babu, R. Padrutt, E. Slyshkina, C. König, S. Ferrari and B. Spingler, Photochem. Photobiol. Sci., 2019, 18, 2792-2803.
- 12 M. Leng, D. Locker, M.-J. Giraud-Panis, A. Schwartz, F. P. Intini, G. Natile, C. Pisano, A. Boccarelli, D. Giordano and M. Coluccia, Mol. Pharmacol., 2000, 58, 1525-1535.
- 13 J. M. Pascoe and J. J. Roberts, Biochem. Pharmacol., 1974, 23, 1345-1357
- 14 M. D. Hall, K. A. Telma, K.-E. Chang, T. D. Lee, J. P. Madigan, J. R. Lloyd, I. S. Goldlust, J. D. Hoeschele and M. M. Gottesman, Cancer Res., 2014, 74, 3913-3922.
- 15 M. Huxley, C. Sanchez-Cano, M. J. Browning, C. Navarro-Ranninger, A. G. Quiroga, A. Rodger and M. J. Hannon, Dalton Trans., 2010, 39, 11353-11364.
- 16 I. A. Riddell, T. C. Johnstone, G. Y. Park and S. J. Lippard, Chem. -Eur. J., 2016, 22, 7574-7581.
- The maybe obvious alternative to <sup>67</sup>Zn-4PyP would have been Pd-4PyP. We have tried many times to synthesize this compound, however the reactions products were always insoluble precipitates. Most likely, the metal coordinates to the exocyclic pyridyl nitrogen atoms, forming an infinite 2D-network.
- 18 S. R. Hennigar and J. P. McClung, J. Nutr., 2016, 146, 2167-2173.
- 19 (a) G. Singh and J. Koropatnick, J. Biochem. Toxicol., 1988, 3, 223-233; (b) K. S. Blisard, D. A. Harrington, D. A. Long and J. E. Jackson, J. Comp. Pathol., 1991, 105, 367-375.
- 20 X. Wu, S. Yuan, E. Wang, Y. Tong, G. Ma, K. Wei and Y. Liu, Metallomics, 2017, 9, 546-555.
- 21 M. Coluccia and G. Natile, Anti-Cancer Agents Med. Chem., 2007, 7, 111-123.
- 22 T. C. Johnstone, K. Suntharalingam and S. J. Lippard, Chem. Rev., 2016, 116, 3436-3486.
- 23 L. J. Kuo and L.-X. Yang, In Vivo, 2008, 22, 305–309.
- 24 T. Migita and S. Inoue, Int. J. Biochem. Cell Biol., 2012, 44, 1872-1876.