RSC Medicinal Chemistry



RESEARCH ARTICLE

View Article Online
View Journal | View Issue



Cite this: RSC Med. Chem., 2025, 16, 2576

Received 12th December 2024, Accepted 4th March 2025

DOI: 10.1039/d4md00984c

rsc.li/medchem

Towards catalytic fluoroquinolones: from metal-catalyzed to metal-free DNA cleavage†

Moshe N. Goldmeier, ‡^a Alina Khononov, ‡^a Tomasz Pieńko, ‡^a Valery Belakhov, ^a Feng-Chun Yen, ^b Limor Baruch, ^b Marcelle Machluf^b and Timor Baasov [®]*

A library of eight new fluoroquinolone-nuclease conjugates containing a guanidinoethyl or aminoethyl auxiliary pendant on the 1,4,7-triazacyclononane (TACN) moiety was designed and synthesized to investigate their potential as catalytic antibiotics. The Cu(II) complexes of the designer structures showed significant *in vitro* hydrolytic and oxidative DNA cleavage activity and good antibacterial activity against both Gram-negative and Gram-positive bacteria. The observed activity of all the Cu(II)-TACN-ciprofloxacin complexes was strongly inhibited in the presence of Cu(III)-chelating agents, thereby demonstrating "vulnerability" under physiological conditions. However, selected TACN-ciprofloxacin conjugates in their metal-free form efficiently cleaved plasmid DNA under physiological conditions. The lead compound 1 showed good DNase activity which was retained in the presence of strong metal chelators and exhibited excellent antibacterial activity against both Gram-negative and Gram-positive bacteria. Density functional theory calculations combined with quantum mechanics/molecular mechanics simulations suggest a general base-general acid mechanism for the hydrolytic DNA cleavage mechanism by compound 1.

Introduction

The rapid increase in antimicrobial resistance poses a significant threat to public health, with warnings about entering a post-antibiotic era already issued.^{1,2} This has sparked renewed efforts to formulate novel antibacterial strategies that can combat existing bacterial resistance and slow down the development of new resistance in the future. Currently, the most promising approaches include combination therapies, modification of existing antibiotics and efficacy enhancement by metabolic stimulation or employing more efficient delivery systems. In addition to these conventional approaches, alternative strategies based on bacteriophages, anti-biofilm agents, probiotics, vaccines, and antibodies have also been proposed.^{3,4}

In recent years, several new antibiotics, such as delafloxacin or plazomicin, have been approved for use in the clinic to deal with multidrug-resistant bacteria. ⁵ However, it has been well-documented that once a new antibiotic is

requirements, and delaying new resistance development.8

introduced into the clinic, whether it is a novel chemical

entity acting on a distinct bacterial target or a semisynthetic

derivative that counters the resistance to its parent drug, it is

only a matter of time until new resistance will yet again

emerge.6 Therefore, new generations of classical antibiotics

that allow us to survive another round of bacterial combat do

not necessarily affect the roots of resistance development.

Recent studies have highlighted multifunctional antibacterial metallopeptides that modify nucleic acids, proteins, or phospholipids *via* reactive oxygen species (ROS) generated by an amino terminal copper and nickel (ATCUN) binding motif. Research by Cowan^{9,10} and Angeles-Boza and co-workers^{11,12} has shown that Cu(II)–ATCUN conjugates with antimicrobial peptides significantly improve antibacterial activity compared to the parent peptides. These conjugates can also act synergistically with conventional antibiotics to reduce biofilm formation¹³ which is challenging to eradicate.¹⁴ However, the observed improvements in minimal inhibitory

A promising strategy to overcome antibacterial resistance is the development of catalytic antibiotics, which are meant to act as small-molecule-based therapeutic agents that catalytically inactivate specific bacterial targets. This approach, akin to the Michaelis–Menten enzyme model, involves noncovalent binding to the target followed by its chemical modification. Catalytic antibiotics promote multiple turnovers of a catalytic cycle, thus potentially improving antibacterial activity, reducing toxicity due to lower dosage

^a Edith and Joseph Fischer Enzyme Inhibitors Laboratory, Schulich Faculty of Chemistry, Technion – Israel Institute of Technology, Haifa 3200003, Israel. E-mail: chtimor@technion.ac.il

^b Faculty of Biotechnology and Food Engineering, Technion – Israel Institute of Technology, Haifa 3200003, Israel

[†] Electronic supplementary information (ESI) available: Characterization of compounds, additional biological assay results, theoretical methods and additional computational results. See DOI: https://doi.org/10.1039/d4md00984c

[‡] These authors contributed equally to this work.

concentration (MIC) were not as significant as expected from catalytic metallodrugs, indicating a need for more complex design strategies to fully realize their clinical potential.

To address these challenges, we explored repurposing known classical antibiotics as selective binding motifs to facilitate the cleavage of critical chemical bonds in bacterial targets, leading to their immediate deactivation. Initially, we applied this concept to aminoglycoside antibiotics¹⁵ and more recently to the fluoroquinolone scaffold.^{8,16}

Fluoroquinolones are highly effective, broad-spectrum antibiotics that are widely prescribed worldwide. They exhibit concentration-dependent bactericidal activity by inhibiting DNA gyrase and topoisomerase IV, essential enzymes for DNA replication and transcription. However, resistance to fluoroquinolones is limiting their use, often due to specific mutations in DNA gyrase and topoisomerase topoisomerase type Consequently, new bacterial inhibitors have been developed to offer alternative binding modes or mechanisms of action. 17,18

Within the concept of catalytic fluoroquinolones, we synthesized ciprofloxacin-nuclease conjugates (see firstgeneration cyclen-based conjugates, Fig. 1) designed to catalytically cleave a specific phosphodiester bond at the site of the fluoroquinolone-topoisomerase-DNA ternary complex. Under oxidative conditions, Cu(II)-complexes of these conjugates fragmented supercoiled plasmid DNA into linear DNA in the presence of DNA gyrase, demonstrating a proof of concept in vitro. The lead compound also showed enhanced antibacterial activity under oxidative conditions compared to the metal-free ligand.8

However, the hydrolytic nuclease activity of the first-Cu(II)-cyclen-ciprofloxacin complexes inhibited under normal cellular conditions due to Cu(II)-aqua

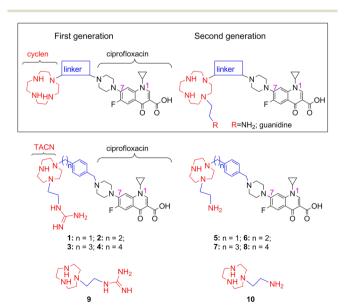


Fig. 1 Designed structures of cyclen-based first-generation⁸ and second-generation 16 conjugates of ciprofloxacin, along with the new TACN-based, third-generation conjugates (1-8) and the reference TACN-warheads (9 and 10).

ligand exchange with endogenous ligands such as glutamate, spermidine or ATP. This "vulnerability" is a significant issue in vivo for metallodrugs working via a hydrolytic mechanism. address we designed second-generation this, ciprofloxacin-nuclease conjugates with dynamic a intramolecular cap to protect the metal center (Fig. 1).16 We have shown that installing the amine pendant at the Co(III)cyclen warhead preserved in vitro DNase activity of the conjugates under physiological conditions. Due to the low pK_a of the amine arm (below 4), the Co(III)-cyclen complex remained capped in solution. At the DNA interface, the complex was uncapped as it formed strong electrostatic interactions with the DNA phosphates. However, the high positive charge of the Co(III) ion and ammonium pendant (+4 in total) could significantly slow down dissociation from the DNA and thereby limit catalytic performance.¹⁶

Consequently, we were motivated to solve the vulnerability issue of the Cu(II) derivatives that bear less positive charge, thus potentially enhancing catalytic turnover. For the secondgeneration cyclen complexes of Cu(II), the pK_a of the aminoethyl or guanidinoethyl pendants was around 9; therefore, the Cu(II) complexes remained uncapped and thus unprotected under physiological conditions. 16 To lower the pK_a of the Cu(II) complexes, we substituted cyclen with 1,4,7triazacyclononane (TACN) as it has been shown that the pK_a of the guanidinoethyl pendant in the Cu(II)-TACN complex is 5.6, ¹⁹ thus favoring the capped state at physiological pH.

With this premise, we assembled a library of thirdgeneration conjugates of ciprofloxacin and Cu(II)-TACN with guanidinoethyl or aminoethyl pendants (compounds 1-8, Fig. 1), along with their corresponding warheads as reference compounds (compounds 9 and 10, Fig. 1). We found that the new designer compounds demonstrated excellent in vitro hydrolytic and oxidative DNA cleavage activity and good antibacterial activity against both Gram-negative and Grampositive bacteria, with improvements regarding the respective cyclen analogues. Surprisingly however, their nuclease activity was inhibited under "normal" cellular conditions indicating that the metal vulnerability issue remained unsolved. Nevertheless, we were pleased to discover a new lead compound based on the TACN-guanidinoethyl warhead that cleaved DNA hydrolytically in a "metal-free" mechanism under physiological conditions and exhibited significant antibacterial activity against both Gram-negative and Grampositive bacteria. By using a variety of in silico tools, we suggest the potential molecular mechanism of DNA cleavage and discuss the benefits of metal-independent nuclease activity for future development of catalytic antibiotics.

Results and discussion

Synthesis of 1-8 and their Cu(II) complexes

Compounds 1-8 were synthesized in four or five chemical steps from terminal bromide derivatives 11a-d (ref. 16) and common intermediate 12 (TFA salt form) incorporating a TACN ring appended to a two-carbon

Research Article

Scheme 1 Synthesis of compounds 1-4 (A) and compounds 5-8 (B). Reagents and conditions: (a) CH₃CN, K₂CO₃, -15 °C; (b) CH₃CN, Cs₂CO₃, 60 °C; (c) dry DCM, Et₃N, Boc₂O; (d) EtOH, CH₃NH₂, K₂CO₃, 0 °C-rt, 21% over 3 steps (13a), 35% over 3 steps (13b), 30% over 3 steps (13c), 40% over 3 steps (13d); (e) CH₃CN, N,N-di-Boc-1H-pyrazole-1-carboxamide, K₂CO₃, rt, 72% (14a), 75% (14b), 53% (14c), 49% (14d); (f) 6 M HCl(aq), 90 °C, 37% (1), 41% (2), 34% (3), 45% (4), 48% (5), 45% (6), 42% (7), 47% (8).

linked protected amine (Scheme 1). It is notable that derivative 12 was prepared from the known bis-Bocprotected derivative 19 of 12 by treatment with TFA, as detailed in the Experimental section.

First, the coupling of primary bromides 11a-d with 12 under basic conditions (Cs2CO3), followed by Boc-protection with Boc₂O, provided the corresponding phthalimide derivatives, which after treatment with methylamine (in the presence of K₂CO₃ (ref. 20)) afforded the corresponding amines 13a-d, which also contained a methyl amide instead of a methyl ester in the ciprofloxacin moiety. The amines were then treated with N,N-di-Boc-1H-pyrazole-1-carboxamide in the presence of a base (K₂CO₃) to afford the corresponding protected guanidine derivatives 14a-d.

One-step deprotection using 6 M HCl at 90 °C, followed by purification of the crude products (using an LH-20 sizeexclusion column), provided the desired compounds 1-4 as hydrochloride salts (Scheme 1A), with guanidinium pendants on the TACN warhead. Alternatively, one-step deprotection of amine derivatives 13a-d using 6 M HCl at 90 °C provided the desired compounds 5-8 as hydrochloride salts (Scheme 1B), with amine pendants. The final 1-8 structures were characterized by combined 1D and 2D nuclear magnetic resonance (NMR) and high-resolution mass spectrometry (HRMS) techniques. Finally, the reference compound 9 was prepared according to a published procedure, 19 and the second reference compound 10 was prepared from the known bis-Boc-protected derivative¹⁹ of 10 by treatment with TFA, as detailed in the Experimental section.

The new set of ligands 1-8 as hydrochloride salts were treated with a stoichiometric amount of CuCl2, followed by pH adjustment (using 1 M NaOH(aq)) to facilitate metal-toligand coordination. The complexes were characterized by UV-vis and HRMS (see the Experimental section for details).

DNA cleavage assays

Biologically relevant nuclease activity tests of metal-free compounds 1-8 and their Cu(II) complexes were performed as previously reported by our group.8,16 Briefly, agarose gel electrophoresis (1% agarose) and ethidium bromide staining were used to monitor the conversion of supercoiled (form I or F-I) pHOT-1 plasmid DNA into its nicked form (form II or F-II) or into multiply nicked DNA; no linear form (form III or F-III) was observed. As reported previously, 8,16 due to the presence of multiple positive charges in the ligands, it is necessary to use an ion-exchange based microscale procedure for removal of the

Chart 1 Compound A from our first-generation cyclen-ciprofloxacin conjugates,8 which was used in this study as the positive control.

ligands prior to electrophoresis to ensure the migration of DNA in the electric field. In all DNA cleavage assays, we used compound A (Chart 1) from our first-generation cyclenciprofloxacin conjugates⁸ as the positive control.

Cleavage experiments of Cu(II) compounds in the absence of adjuvants. For the guanidine series, only 1-Cu(II) exhibits significant DNase activity that is comparable to the parent complex 9-Cu(II)¹⁹ at millimolar concentrations (see Fig. 1 and 2). In contrast, 3-Cu(II) and 4-Cu(II) show no significant activity, whilst 2-Cu(II) exhibits very strong DNA binding rendering it a poor candidate for effective catalysis (see Fig. S1†).

For the amine series, however, three of the Cu(II) complexes, 5-Cu(II), 6-Cu(II) and 8-Cu(II), show significantly enhanced DNase activity compared to the parent complex 10-Cu(II) (see Fig. 1 and 2), whilst 7-Cu(II) shows no significant activity (see Fig. S1†).

Collectively, these data indicate that (i) DNA binding and (ii) DNase activity are highly dependent on the linker length, both for the guanidine and amine series. However, the intercalating cipro scaffold apparently mediates enhanced DNase activity for the amine series, but not for the guanidine series. It is reasonable to suggest that the conformational restrictions imposed upon the bulky guanidine pendant by scaffold-intercalation offset any potential **DNase** enhancement mediated by the "proximity effect" (or its increased effective molarity²¹). Indeed, we have previously reported16 a similar discrepancy between the guanidine and amine pendants for the cyclen-based analogues (i.e., Cu(II) complexes of second-generation derivatives, Fig. 1), for which the guanidine pendant (but not the amine pendant) shows a strong inhibitory effect on DNase activity.¹⁶

Finally, to investigate whether the amine or guanidine pendants provide effective protection (electronic and/or steric) against Cu(II)-aqua ligand exchange,8 we investigated the DNase activity of 1–Cu(π), 5–Cu(π) and 6–Cu(π) in the presence of Cu(II)-chelating Tris buffer. Unfortunately, as depicted in Fig. S2,† all DNase activity was lost under these conditions, which demonstrates that the Cu(II) complexes are vulnerable to ligand exchange. We also performed potentiometric titration (pH 2-12.5) in combination with UV-vis spectroscopy to probe whether the pendants coordinate to the metal. Specifically, we investigated 3-Cu(II) (guanidine series) and 7-Cu(II) (amine series). In both these cases, the UV-vis spectra showed no significant perturbation in the investigated pH range until the pK_a of the pendant was reached, i.e., around 12 for 3-Cu(II) and around 10 for 7-Cu(II). These data indicated that the guanidine and amine pendants do not interact with the metal centre at physiological pH.

Cleavage experiments of Cu(II) compounds in the presence of ascorbic acid. Three of the Cu(II) complexes, 1-Cu(II), 5-Cu(II) and 6-Cu(II), were selected to investigate the DNA cleavage activity in the presence of ascorbic acid under aerobic conditions, which facilitates the production of hydroxy radicals via the Cu(II)/(I) ion couple.8

As depicted in Fig. S3,† 1-Cu(II) shows no significant activity at micromolar concentrations (>40 µM). In stark contrast, the cyclen-based analogues (second-generation derivatives with guanidine pendants, Fig. 1) all show complete conversion of form I to form II at 5 µM.8 This distinction however is not unexpected, as Cowan and coworkers²² have reported that unlike Cu(II)-cyclen, Cu(II)-TACN has no significant DNA cleavage activity in the presence of ascorbic acid under aerobic conditions.

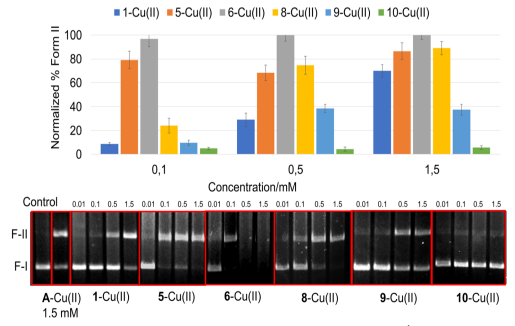


Fig. 2 Comparative concentration-dependent cleavage of (+) supercoiled pHOT-1 plasmid (0.014 μg μL⁻¹) in HEPES buffer (50 mM, pH 7.4) over 5 h. Compounds include A-Cu(II) as a positive control.

Interestingly however, 5–Cu(π) and 6–Cu(π) do exhibit significant DNase activity under these conditions, with complete conversion of form I to multiply nicked DNA (as evidenced by a DNA smear) at 20 μ M and 80 μ M, respectively (Fig. S3†). Although these activities are much less pronounced than their respective cyclen-based analogues (showing complete conversion at 2.5 μ M and 0.5 μ M, respectively¹⁶), these data nevertheless highlight the critical impact of the amine ν s. guanidine pendant on the oxidative DNA cleavage activity of these complexes.

Cleavage experiments of metal-free compounds. For the guanidine series, compounds 1 and 3 exhibit significantly enhanced DNase activity at millimolar concentrations compared to the parent compound 9 (ref. 19) (see Fig. 3). Compounds 2 and 4, however, are poor catalytic candidates since they exhibit very strong DNA binding under the same conditions (see Fig. S4†).

For the amine series, compound 5 exhibits significantly enhanced DNase activity compared to the parent compound 10 (see Fig. 3). In contrast, compounds 7 and 8 both exhibit very strong DNA binding, whilst compound 6 shows no significant DNase activity (see Fig. S4†).

In summary, these data suggest that (i) DNA binding and (ii) DNA cleavage activity are highly dependent on the linker length, for both the guanidine and amine series. Furthermore, compound 1 is shown to be the most effective metal-free nuclease across both series (see Fig. 3).

To investigate whether the mechanism of cleavage for compound 1 is metal-mediated, a scavenging assay was performed in 1 mM EDTA. As illustrated in Fig. 4, whilst 3 loses all DNase activity in the presence of EDTA (and therefore serves as a negative control), compound 1 retains significant activity. Furthermore, in contrast to 1–Cu(II) which loses all DNase activity in the presence of metal-chelating

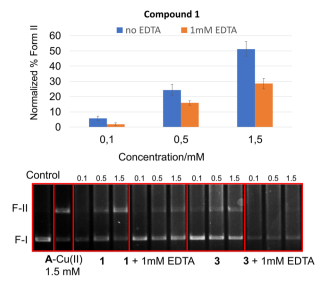


Fig. 4 Comparative concentration-dependent cleavage of (+) supercoiled pHOT-1 plasmid (0.014 μg μL^{-1}) in HEPES buffer (50 mM, pH 7.4) over 5 h, with or without 1 mM EDTA. Compounds include A–Cu(II) as a positive control.

Tris buffer (see Fig. S2†), 1 retains full DNase activity under the same conditions (see Fig. 5).

Collectively, these data strongly suggest that the combination of the DNA-intercalating ciprofloxacin scaffold and the macrocyclic polyamine TACN facilitates metal-free, hydrolytic DNA cleavage. Indeed, several TACN-anthraquinone intercalator conjugates have been reported to exclusively exhibit metal-free DNA cleavage activity. ^{23,24} Those compounds retained their activity in the presence of metal chelators and ROS scavengers, thereby excluding DNA cleavage *via* metals, autooxidation and ROS formation. It is reasonable to speculate that an intercalator moiety (*i.e.*,

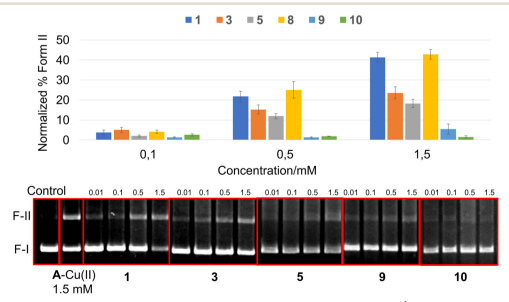


Fig. 3 Comparative concentration-dependent cleavage of (+) supercoiled pHOT-1 plasmid (0.014 μ g μ L⁻¹) in HEPES buffer (50 mM, pH 7.4) over 5 h. Compounds include A-Cu(II) as a positive control.

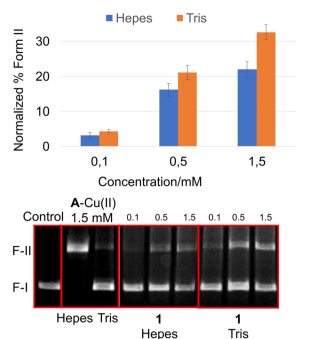


Fig. 5 Comparative concentration-dependent cleavage supercoiled pHOT-1 plasmid (0.014 μ g μ L⁻¹) in HEPES or Tris buffer (50 mM, pH 7.4) over 5 h. Compounds include A-Cu(II) as a positive control

ciprofloxacin or anthraquinone) and an optimal spacer enable TACN to be proximal to the DNA backbone without becoming strongly bound to it.

Antibacterial activity and cytotoxicity

We tested compounds 1-8 and their Cu(II) complexes for their comparative antibacterial activity by determining MIC values against two Gram-negative E. coli strains (R477-100

Table 1 MIC (μg mL⁻¹) of 1–8 and their Cu(II) complexes

Bacteria/compound Ciprofloxacin	а	b	с	d
	Gram-negative		Gram-positive	
	0.05	0.05	0.19	0.02
1	6	3	3	3
1 -Cu(II)	6	3	3	3
2	12	12	12	12
2-Cu(II)	12	6	12	6
3	3	3	3	3
3-Cu(11)	6	6	6	6
4	12	12	12	6
4-Cu(II)	12	12	12	6
5	12	6	6	3
5-Cu(II)	12	12	6	6
6	12	6	12	6
6-Cu(11)	12	12	6	6
7	12	6	3	3
7–Cu(II)	6	6	6	3
8	12	12	3	3
8– Cu(11)	12	12	6	3

^a Escherichia coli R477-100. ^b Escherichia coli 25922. ^c Staphylococcus epidermidis. ^d Bacillus subtilis 6633.

and 25922) and two Gram-positive strains, Staphylococcus epidermidis and Bacillus subtilis (Table 1).

All the new derivatives of ciprofloxacin exhibit significant antibacterial activity against Gram-negative and Grampositive bacteria. The two lead compounds against the two E. coli strains tested (Gram-negative) are 1 and 3 (3-6 µg mL⁻¹), which show 2- to 4-fold improvement in comparison with the other compounds. However, against the two Gram-positive strains tested, 1, 3, 7 and 8 all exhibit similarly good antibacterial activity (3 µg mL⁻¹).

Furthermore, all the Cu(II) complexes of 1-8 exhibit similar antibacterial activity compared to the metal-free ligands. Of particular note in this regard is 5-Cu(II), which exhibits strong DNase activity compared to the metal-free ligand (i.e., full conversion of form I to form II at 0.1 mM vs. no DNase activity whatsoever for 5 - see Fig. 1 vs. 2) but no improvement in antibacterial activity in any of the tested strains. Thus, the MIC data corroborate the notion that these Cu(II) complexes are indeed vulnerable to ligand exchange. However, the new compounds have significantly reduced antibacterial activity relative to the parent ciprofloxacin (Table 1). A very similar reduction in antibacterial potency of ciprofloxacin derivatives also observed was ciprofloxacin was conjugated with other highly positively charged molecules like cyclen8,16 and aminoglycosides to yield hybrid antibiotics.25 It was suggested that the cellular permeability of these ciprofloxacin conjugates is significantly lower than that of the parent drug or the conjugates have increased offsite binding. Both phenomena could be attributed to the high polarity of either cyclen, 8,16 TACNguanidine/TACN-amine (current work), or aminoglycosides, 25 covalently linked to ciprofloxacin.

Given the fact that 1 shows (i) significant, metal-free, hydrolytic DNase activity (see Fig. 4 and 5) and (ii) exhibits strong antibacterial activity against both Gram-negative and Gram-positive bacteria (Table 1), it would be intriguing to investigate the possibility that 1 is a dual-acting antibacterial agent with the potential to slow down resistance development.

To gain a better understanding of the safety of the lead compounds 1, 2, 5 and 8, we determined their comparative cell toxicity to ciprofloxacin by measuring the half-maximal lethal concentrations values (LC₅₀) in baby hamster kidney

Table 2 Comparative cell toxicity LC50 (mM) of ciprofloxacin and lead compounds 1, 2, 5 and 8

Cells/compound ^a	ВНК	HEK293-FT		
Ciprofloxacin	0.20	0.11		
1	0.51	0.48		
2	0.31	0.34		
5	0.42	0.71		
8	0.74	0.46		

^a Cell toxicity was measured in baby hamster kidney (BHK) and human embryonic kidney (HEK293-FT) cells and calculated as a ratio between the number of living cells in cultures grown with and without the presence of the tested compounds.

(BHK) and in human embryonic kidney (HEK293-FT) cells (Table 2). Among the tested compounds, the parent drug ciprofloxacin is the most cytotoxic compound with LC50 values of 0.2 and 0.11 mM in BHK and HEK293-FT, respectively. Compound 2 is the most toxic compound among the synthetic derivatives with an LC₅₀ value of 0.3 mM for both cell lines. Nevertheless, it is still 1.5- and 3-fold less cytotoxic than ciprofloxacin. In BHK cells, the LC50 of compound 8 is 0.74 mM, making it the least cytotoxic compound among all. Finally, the lead compound 1 has a cytotoxicity of 0.5 mM for both cell lines, 2.6- and 4.4- fold less toxic than ciprofloxacin.

Molecular mechanism of metal-free DNA cleavage

Research Article

Intrigued by the metal-free DNase activity of compound 1, we investigated its molecular mechanism by combining several computational methods including quantum chemical calculations based on density functional theory (DFT), molecular docking, classical molecular dynamics (MD), and hybrid quantum mechanics/molecular mechanics (QM/MM) simulations (see Computational methods in the ESI†).

Initially, we focused on studying the structure of the TACN-guanidine warhead in solution. It is of note that the two tertiary amines of the warhead are chiral yielding four possible configurations (RR, RS, SR, SS). We sampled and energy-scored all possible protonation states of the warhead (see Fig. S5 and Table S1†) using GFN2-xTB²⁶ in CREST 2.12 (ref. 27) and CENSO 1.2.0 (ref. 28) followed by DFT calculations in Orca 6.0 (ref. 29) coupled with the COSMO-RS implicit solvation model in OpenCOSMO-RS.30 The freeenergy calculations showed that in all the protonation states

considered, the most energetically stable are stereoisomers with either (R, R) or (S, S) configurations. For the first protonation, we found that the most basic amine within the warhead is the guanidine pendant, for which we estimated a pK_a of 14.48, which agrees with its exceptionally high pK_a in arginine (13.8).31 The second protonation favorably occurs at the secondary amine, with the pK_a calculated to be 12.20. The triprotonation is preferred at the tertiary amine connected with the linker, with the pK_a calculated to be 5.77. The second and the third deprotonation constants correspond to those observed for disubstituted TACN derivatives, such as Hnoapy³² (p K_1 = 12.15 and p K_2 = 5.78) or H_2 noda³³ (p $K_1 = 11.82$ and p $K_2 = 6.70$), supporting the *in* silico predictions. For the quadruple protonated state, we estimated a p K_a of 3.34. The p K_a results indicate that under physiological conditions, the TACN-guanidine warhead is predominantly diprotonated.

To study specific interactions of the diprotonated TACNguanidine warhead with water, we performed classical MD simulations in explicit solvent using the AMBER force field.³⁴ We found that the warhead can recruit a single water molecule from the bulk, bridging the guanidine pendant and TACN amine, stabilizing two positively charged amines in proximity (Fig. 6A). The capability of the TACN-guanidine warhead to bind to a single water molecule has brought us to the hypothesis that this water could be used as a nucleophile for attack on the DNA phosphate in an analogous manner to metal-based catalysis of DNA cleavage.35 To test this hypothesis, we used the DFT-optimized geometries of the complexes of the TACN-guanidine warhead with water observed in the MD simulations, in both (R, R) and (S, S) configurations (Fig. 6A) to perform fixed-anchor docking³⁶ of

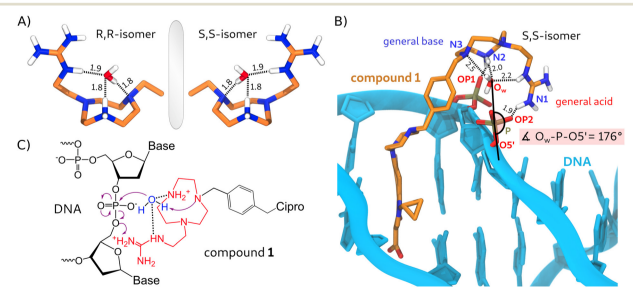


Fig. 6 A: DFT-optimized structures of water complexes of the TACN warhead with the guanidinoethyl pendant in two favorable stereoisomers observed in the MD simulations. For computational efficiency, the ciprofloxacin-linker part is represented by an ethyl group. B: The most probable binding mode of compound 1 during DNA cleavage, derived from the QM/MM simulation and visualized using VMD. 39 The O-P-O angle and key interactions (distances in Å) for catalysis between compound 1 and DNA are depicted. C: General acid-general base mechanism of DNA cleavage by compound 1.

compound 1 to the selected binding sites of DNA (PDB code: 2XKK³⁷) in many conformations which we generated in the Hamiltonian replica-exchange MD (H-REMD)³⁸ simulations (see Fig. S6 and S7†).

Only among the docked structures with the (S, S) configuration of the chiral centers in the warhead, we found poses in which the TACN-guanidine warhead oriented the water molecule in the in-line geometry for nucleophilic attack on the DNA phosphate (with ≥150° angle between the oxygen atom of water Ow, the phosphorus atom P, and the oxygen atom of the DNA phosphate O5' and the distance Ow-P of ≤4.0 Å). The selected structures were further simulated using the classical MD and AMBER force field⁴⁰⁻⁴² (see Fig. S8†). The simulations revealed instability of the complex of the catalytic warhead and DNA on the pico-nanosecond timescale. We observed that using three-point charge representation of water (e.g. OPC3 (ref. 43)) destabilized the complex even more than four-point charge models (e.g. OPC (ref. 40)). The instability issue persisted even after refinement of the AMBER parameters^{44,45} of compound 1 (see Fig. S9†), leading us to the notion that the AMBER force field might be an oversimplified model to study the interactions between compound 1 and DNA prior to the hydrolysis reaction. Therefore, we tested different semi-empirical potentials in the OM/MM approach.46

Only one setup, combining the third-order densityfunctional tight-binding (DFTB3) method corrected with D4 three-body dispersion^{47,48} and using the OPC water model, provided stability of the DNA-warhead complex on the timescale of hundreds of picoseconds (see Fig. S10†). During the QM/MM simulations, the nearly in-line geometry of water approaching the DNA phosphate was stable (with O_w -P \approx 3.78 Å and O_w -P-O5' \approx 163°) (Fig. 6B and S10†). The hydrogen atoms of the water molecule interacted preferably with the TACN tertiary amine connected to the linker (the N3 atom in Fig. 6B) (with H_w -N3 ≈ 2.91 Å) (Fig. 6B and S10†). The in-line orientation of water within the pre-reaction complex is stabilized by two interactions between the DNA and the warhead: i) between the protonated secondary amine of TACN and the adjacent DNA phosphate (N2-OP1 ≈ 2.85 Å), and ii) between the guanidine pendant and the cleaved phosphate (N1-OP2 ≈ 3.02 Å) (Fig. 6B and S10†).

The MD and QM/MM simulations suggested that accurate description of electrostatic and dispersion interactions within the DNA-warhead complex at the DFT level is mandatory to study the next steps of the mechanism of cleavage. Nevertheless, collectively, our calculations suggest that compound 1 cleaves DNA in a general base-general acid mechanism (Fig. 6C). Since the TACN tertiary amine conjugated with the linker (the N3 atom in Fig. 6B) is mostly deprotonated at physiological pH (p $K_a = 5.77$) and favorably interacts with the water molecule in the in-line geometry at the DNA interface, it can serve as a general base, activating the water molecule for attack on the DNA phosphate. Meanwhile, the close interaction of the protonated guanidine arm with the cleaved phosphate group indicates that it may function as a general acid, neutralizing the negative charge of the phosphate and activating the leaving group. Because of the high pK_a of guanidine, its role in shuttling protons on the phosphorane intermediate formed during the hydrolysis has been under debate. 49 In this context, we can rationalize the enhanced DNase activity of compound 1 in Tris buffer (relative to HEPES buffer, Fig. 5), in which a primary amine with a p K_a of 8.6 (ref. 50) could serve as a more potent proton donor for the leaving group than guanidine.

Perspective

To date, metal-free artificial DNases have been commonly built as bis-guanidinium receptors based on the scaffold of phenyl,⁵¹ calix[4]arenes,⁵² Dervan-type polyamides⁵³ and cyclic peptides⁵⁴ decorated with moieties bearing hydroxyl groups, imitating DNA topoisomerase I which uses tyrosine as a nucleophile and catalyzes a transphosphorylation reaction.⁵⁵ An alternative strategy for designing non-metallic nucleases has involved cyclic polyamines like cyclen and TACN. DNase capabilities have been observed in various compounds: N-dimethylated cyclen, 56 bis-cyclen connected with urea,⁵⁷ a cyclen conjugate with peptide nucleic acid⁵⁸ and TACN combined with anthraquinone.²³ Additionally, modifications involving guanidine or amine pendants, and hydroxyethyl were made to TACN perform to transphosphorylation, leading to DNase activity at low micromolar concentrations. 24,59 The transphosphorylation mechanism effectively cuts DNA, yet compounds operating in this manner were thought to become covalently trapped in DNA, preventing subsequent catalytic cycles.

To the best of our knowledge, our study is the first report of an antibiotic molecule bearing a small organic fragment that cleaves DNA with no metal involvement in a general base-general acid mechanism by recruiting a water molecule from bulk. Unlike the transphosphorylation mechanism applied in most artificial metal-free DNases, our proposed DNA hydrolysis mechanism allows the compound to be released after the reaction and perform multiple catalytic turnovers.

Furthermore, metal-free DNA cleavage avoids the problems associated with metals, such as vulnerability under physiological conditions or the associated toxicity due to the ROS formation. With a purely hydrolytic mechanism and strong dependence of the nuclease activity on its structure, it is also feasible to improve site selectivity of cleavage and limit any adverse effects.

It is of note that since ciprofloxacin conjugates of TACN with amine or guanidine pendants were initially designed for metal-catalyzed DNA cleavage, their efficiency in non-metallic catalysis may be limited by the imperfect structure. The QM/ MM simulations revealed that compound 1 can cleave DNA only when ciprofloxacin is arranged in an orientation parallel to the DNA bases, whereas, the H-REMD simulations of the ciprofloxacin-linker conjugate suggested that it preferred perpendicular orientation to the DNA bases (Fig. S7†). Thus,

it can be speculated that, depending on the DNA sequence context, compound 1 can be trapped in catalytically inactive conformations, significantly limiting its global nuclease efficiency. Optimization of the linker connecting the binding and catalytic domains seems crucial for increasing the rate of the hydrolysis reaction. In addition, higher nuclease activity could be provided by modifying the linker attachment site in the catalytic warhead to increase general base pK_a as well as modifying the structure of the guanidine moiety to be a stronger general acid.⁶⁰

Despite those shortcomings, we believe that the presented concept of metal-free DNA cleavage by TACN with a guanidinoethyl pendant can be applied to other antibiotics targeting bacterial nucleic acids, such as aminoglycosides, and paves a new path in the future development of catalytic antibiotics.

Conclusions

In this study, we explored the potential to solve the issue of metal vulnerability for the Cu(II) complexes of the second-generation nuclease-conjugated ciprofloxacin by replacing the cyclen moiety with TACN to effectively enhance the amine or guanidine cap formation at the metal center.

Firstly, the DNA cleavage assays underscore the critical role of the linker length in determining the DNase activity and DNA binding affinity of both guanidine and amine series compounds. For the guanidine series, only 1-Cu(II) exhibited significant DNase activity comparable to the parent complex 9-Cu(π), while 2-Cu(π) showed strong DNA binding but poor catalytic efficiency. In contrast, for the amine series, 5-Cu(II), 6-Cu(II), and 8-Cu(II) demonstrated enhanced DNase activity compared to the parent compound 10-Cu(II), with 7-Cu(II) showing no significant activity. These findings indicate that while the guanidine pendant may exacerbate DNase activity of the Cu(II) complexes due to conformational restrictions, the amine pendant can enhance DNase activity. Furthermore, investigations using Cu(II)-chelating agents like Tris buffer revealed that all tested Cu(II) complexes lost their DNase activity, suggesting vulnerability to ligand exchange.

Surprisingly, the metal-free compounds also displayed significant variability in their DNase activity. Compound 1 emerged as the most effective metal-free nuclease, retaining significant activity even in the presence of EDTA, indicating a metal-free, hydrolytic mechanism of DNA cleavage. This suggests that compound 1 could act as a dual-acting antibacterial agent with both DNA cleavage and antibacterial properties due to inhibiting topoisomerase II enzymes, offering potential to slow down resistance development.

The antibacterial activity assays further reinforced the potential of these compounds. Compounds 1 and 3 were particularly effective against Gram-negative $E.\ coli$ strains, showing 2- to 4-fold improvement over other compounds, while compounds 1, 3, 7, and 8 exhibited strong activity against Gram-positive strains. Notably, the Cu(II) complexes showed similar antibacterial activity to the metal-free ligands,

with no significant enhancement, corroborating the susceptibility of these complexes to ligand exchange.

DFT calculations combined with MD and QM/MM simulations provided deeper insights into the DNA cleavage mechanism of compound 1. The simulations revealed that the DNA sequence context may influence the binding mode of compound 1, which can affect its cleavage capability. Importantly, the structural analysis of the TACN-guanidine warhead in solution indicated that the warhead could recruit a water molecule, which may act as a nucleophile in DNA cleavage, supporting a general base-general acid mechanism of hydrolysis.

In conclusion, this study has identified compound 1 as a promising candidate for further development of antibacterial metal-free nuclease-active agents. The findings suggest a dual mechanism of action, potentially enhancing its therapeutic efficacy and reducing the likelihood of resistance development. Future research should focus on optimizing the DNase activity of these compounds under physiological conditions selectively within the ternary complex of topoisomerase IV or DNA gyrase, exploring their detailed mechanisms of action through advanced simulations and evaluating their efficacy *in vivo*. The insights gained from this study provide a solid foundation for the development of novel antibacterial agents with enhanced metal-free DNA cleavage capabilities.

Experimental

General techniques

¹H and ¹³C NMR, DEPT, COSY, HMQC, and HMBC spectra were recorded on Bruker AvanceTM 600/500 spectrometers. Chemical shifts reported (in ppm) are relative to CHCl₃ (δ = 7.26) with CDCl₃ as the solvent, to HOD ($\delta = 4.79$) or internal TMS with D_2O as the solvent, and to CD_3OH ($\delta = 3.31$) or internal TMS with CD3OD as the solvent. Mass spectral analyses were performed on a Bruker Maxis Impact under electron spray ionization (ESI+) QTOFMS or on a Bruker Autoflex Speed under matrix-assisted laser desorption ionization (MALDI) TOFMS. Reactions were monitored by TLC on silica gel (Gel 60 F254, 0.25 mm, Merck), and spots were visualized using a UV lamp and iodine, or by charring with a yellow solution containing (NH₄)₆Mo₇O₂₄·4H₂O (120 g) and (NH₄)₂Ce(NO₃)₆ (5 g) in 10% H₂SO₄ (800 mL). Flash column chromatography was performed on silica gel 60 (70-230 mesh). The EPR spectra were recorded on a Bruker EMX-10/12 X-band (v = 9.3 GHz) digital EPR spectrometer. The spectra were recorded at a microwave power of 200 mW, 100 kHz, and magnetic field modulation of 3 G amplitude. The digital field resolution was 2048 points per spectrum. Spectral processing was performed with the Bruker WIN-EPR and SimFonia software. The UV-vis spectra were recorded on an Ultrospec 2100 pro spectrometer. All chemicals unless otherwise stated were obtained from commercial sources. The purity of the lead compound 1 was determined by using HPLC analysis, which indicated >95% purity (see the ESI†).

Removal of ligands prior to electrophoresis

The procedure proved to be necessary when more than 10 μ M compound (metal-free, Cu(II)) was employed, that is, (only) for the DNA cleavage assays that were performed in the absence of any adjuvants. An Amberlyst 15 (Sigma-Aldrich) ion-exchange resin was prepared according to the reported literature⁶¹ and then utilized for ligand removal as previously reported by our group.¹⁶

DNA cleavage assays

DNA cleavage activity of 1-8 and their Cu(II) complexes toward supercoiled (+) pHOT-1 plasmid DNA (TopoGEN) was monitored by gel electrophoresis. In a typical experiment, plasmid DNA (400 ng, 0.014 µg mL⁻¹) in HEPES buffer (50 mM, pH 7.4) was mixed with different concentrations of the compounds in the presence or absence of ascorbic acid (0.32) mM). All stock solutions of buffers and of the compounds were prepared using HPLC grade water (ChromAR). Molecular biology reagent grade water (Sigma) was added up to a total reaction volume of 30 µL before incubation for a given time. For the ascorbic acid assays, the reaction was quenched immediately after incubation with EDTA (54 mM). For analysis, 10 µL of loading buffer (40% sucrose, 100 mM Tris-HCl (pH 7.5), 1 mM EDTA, 0.5 mg mL⁻¹ bromophenol blue) was added to the incubated solution, and the mixture was loaded onto an agarose (SeaKemLE) gel [1% in TAE × 1 buffer (40 mM Tris, 20 mM acetic acid, 1 mM EDTA)]. Electrophoresis was carried out at 60 V for 2 h. The gels were then stained with 1 µg mL⁻¹ ethidium bromide for 30 min and de-stained in TAE × 1 buffer. DNA bands were visualized with medium-range ultraviolet light using a Bio-RadGel Doc XR+ imaging system. The quantity of different DNA forms (I or II) was estimated using ImageJ⁶² software.

Antibacterial and cytotoxicity tests

Comparative antibacterial activities were determined by measuring the MIC values using the double-microdilution method according to the Clinical and Laboratory Standards Institute⁶³ (CLSI). A Luria–Bertani growth medium and polypropylene 96-well plates (Thermo) were used. All the experiments were performed in triplicate, and analogous results were obtained in three different experiments.

For the cytotoxicity assays, HEK-293FT cells were seeded overnight in 96-well plates (5000 cells per well) in a DMEM medium containing 10% FBS, 1% penicillin/streptomycin, 1% glutamine, and 1% pyruvate at 37 °C under 5% CO₂. BHK cells were seeded overnight in 96-well plates (5000 cells per well) in a DMEM medium containing 10% FBS and 1% penicillin/streptomycin. Different concentrations of the tested compounds were added to the medium and incubated with the cells for 48 h. A cell proliferation assay (resazurin-based colorimetric assay, Sigma-Aldrich) was performed under 3 h incubation. The fluorescence values were then read by using a plate reader. Cell viability values were normalized to the untreated samples of each tested

compound. The half-maximal lethal concentration (LC_{50}) values were obtained by fitting concentration–response curves to the data of at least four biological repeats within each of 2 independent experiments using an online LC_{50} calculator (AAT Bioquest, https://www.aatbio.com/tools/lc50-calculator).

Compound 13a

Step 1. A suspension of 12 (2.2 gr, 4.1 mmol) and K_2CO_3 (0.6 gr, 4.3 mmol) in dry CH_3CN (80 mL) was set stirring at -15 °C (ice/NaCl bath) for 30 min under an argon atmosphere. Then 11a (1.8 g, 3.4 mmol) was added portion-wise over 18 h (*i.e.*; 0.1 equiv. every 2 hours). The reaction was monitored by TLC using the following system: $CHCl_3$: MeOH: $MeNH_2$ (33% $MeNH_2$ in EtOH) 30: 4:1, which indicated that the reaction stopped progressing after 18 h. The reaction mixture was filtered under gravity and washed extensively with CH_3CN and then the solvent was removed *in vacuo*.

Step 2. The crude was dissolved in dry DCM (80 mL) and set stirring at 0 °C (ice bath) under an argon atmosphere. Et₃N (2.3 mL, 16.5 mmol) and Boc₂O (2.9 mL, 12.5 mmol) were added, and the ice bath was removed after 30 min. The reaction was monitored by TLC using the following systems: CHCl₃: MeOH: MeNH₂ (33% MeNH₂ in EtOH) 30:4:1.3 and DCM:MeOH:Et₃N 30:1:0.2. The reaction mixture was filtered under gravity and washed extensively with DCM and then the solvent was removed in vacuo. The crude product was then loaded onto a DCM-packed silica column as a DCM solution; the desired product was eluted using a DCM: MeOH: Et₃N (30:1:0.1) solvent system. A post-column workup was then performed with DCM and water; the desired product was isolated in the organic phase, dried with MgSO₄ and then evaporated to yield the pure compound (1.39 g, 1.63 mmol, 35% over two steps); ¹H NMR (600 MHz, CDCl₃) $\delta_{\rm H}$ 8.47 (s, 1H, QH-2), 7.95-7.93 (d, 1H, QH-5), 7.77-7.76 (m, 2H, phthalimide Ar), 7.65-7.64 (m, 2H, phthalimide Ar), 7.28-7.18 (m, 5H, QH-8, linker Ar), 3.84 (s, 3H, OCH₃), 3.69-3.67 (m, 2H, CH_2 -phthalimide), 3.52 (m, 2H, $Ph-CH_2$ -piperazine), 3.36-3.33 (m, 1H, cyclopropane CH), 3.22-3.16 (m, 8H, piperazine (4H), TACN (4H)), 2.87-2.74 (m, 6H, TACN (4H), CH_2CH_2 -phthalimide (2H)), 2.61–2.58 (m, 10H, piperazine (4H), TACN (4H), TACN-CH₂-Ph (2H)), 1.41-1.37 (m, 9H, Boc), 1.25-1.19 (m, 2H, cyclopropane CH₂), 1.09-1.06 (m, 2H, cyclopropane CH₂); 13 C NMR (150 MHz, CDCl₃) $\delta_{\rm C}$ 173.11 (cipro C=O), 168.38 (phthalimide C=O), 166.50 (cipro C=O), 155.51 (Boc C=O), 154.27 (cipro), 152.62 (cipro C-H), 148.36 (cipro), 144.63 (linker Ar), 138.02 (cipro), 134.03 (linker Ar), 133.96, 132.11 (C-H phthalimide Ar), 132.05, 131.84 (phthalimide Ar), 129.69 (C-H linker Ar), 129.30, 129.07 (C-H linker Ar), 123.37, 123.22 (C-H phthalimide Ar), 122.96 (cipro), 113.18 (cipro C-H), 110.03 (cipro), 104.79 (cipro C-H), 76.84 (Boc C(CH₃)₃), 62.57 (Ph-CH₂-piperazine), 52.75 (piperazine), 52.08 (OCH₃), 49.89 (piperazine), 34.51 (cyclopropane CH), 29.69, 28.56, 28.43, 28.35 (Boc $C(CH_3)_3$),

8.14 (cyclopropane <u>CH</u>₂). MS (ESI+ QTOFMS) calculated for $C_{47}H_{57}FN_7O_7$ ([M + H]⁺) m/z 850.42; measured m/z 850.42.

Step 3. The third step was carried out according to a previously reported procedure.20 To a stirred solution of the phthalimide derivative (400 mg, 0.47 mmol) in absolute EtOH (15 mL) were added anhydrous K2CO3 (150 mg, 1.09 mmol) and MeNH₂ (15 mL, 33% MeNH₂ in EtOH) at 0 °C (ice bath) under an argon atmosphere. The suspension was then left to stir in the ice bath for 30 min before the system was allowed to return back to r.t. The reaction was monitored by TLC using the same system as in step 1 and indicated complete consumption of the starting material after 18 h. The crude suspension was then filtered and washed extensively with ethanol, and the solvent was removed by evaporation in vacuo. The residue was purified by silica chromatography using a MeOH/MeNH2 elution system to yield the pure compound 13a (203 mg, 0.28 mmol, 60%); ¹H NMR (600 MHz, $CDCl_3$) δ_H 9.83 (m, 1H, CH_3 -NH-CO), 8.76 (s, 1H, QH-2), 7.96-7.94 (d, 1H, QH-5), 7-80-7.78 (d, 1H, QH-8), 7.66, 7.26-7.20 (m, 4H, linker Ar), 3.72-3.69 (m, 2H, Ph-CH₂piperazine), 3.65-3.61 (m, 1H, cyclopropane CH), 3.56-3.50 (m, 4H, CH₂N), 3.47-3.39 (m, 5H, piperazine (4H), CH₂N (1H)), 3.24-3.13 (m, 4H, CH_3 -NH-CO (3H), CH_2 N (1H)), 2.94-2.88 (m, 7H, CH₂N), 2.80-2.78 (m, 3H, piperazine), 2.62, 2.54-2.49 (m, 6H, TACN- CH_2 -Ph (2H), CH_2N (4H)), 1.41-1.39 (m, 9H, Boc), 1.31, 1.12-1.09 (m, 3H, cyclopropane CH₂), 0.82-0.78 (m, 1H, cyclopropane CH₂); ¹³C NMR (150 MHz, CDCl₃) $\delta_{\rm C}$ 1752.54 (cipro C=O), 168.95 (cipro C=O), 156.84 (Boc C=O), 154.14 (cipro), 145.58 (cipro C-H), 145.02 (cipro), 140.76 (linker Ar), 137.52 (cipro), 134.96 (linker Ar), 130.54, 128.71 (C-H linker Ar), 126.57 (C-H linker Ar), 123.75 (cipro), 113.61 (cipro C-H), 110.35 (cipro), 108.98, 105.62 (cipro C-H), 79.06 (Boc $C(CH_3)_3$), 64.75 (Ph- CH_2 -piperazine), 56.05, 52.87, 51.93, 50.41 (CH₂N), 52.76 (piperazine), 51.84 (CH₂N), 50.03 (piperazine), 40.33, 38.05 (CH₂-NH₂), 35.01 (cyclopropane CH), 29.53-27.46 (TACN-CH₂CH₂CH₂-Ph, Boc $C(CH_3)_3$, 22.86 (CH₃-NH-CO), 9.06 (cyclopropane CH₂). MS (ESI+ QTOFMS) calculated for C₃₉H₅₆FN₈O₄ ([M + H]⁺) m/z 719.43; measured m/z 719.43.

General procedure for the synthesis of compounds 13b-d

Step 1. A stirred suspension of **12** (1.25 equiv.) and Cs_2CO_3 (1.25 equiv.) in dry CH_3CN (35 mL g^{-1} **12**) was set stirring at r.t. for 10 min under an argon atmosphere. The reaction mixture was then heated to 50 °C for 30 min. Then the appropriate bromide (**11b–d**, 1 equiv.) was added portionwise over 24 h (*i.e.*; 0.1 equiv. every 3 hours). The reaction was monitored by TLC using the following system: $CHCl_3$: MeOH: MeNH₂ (33% MeNH₂ in EtOH) 30:4:1. The reaction mixture was filtered under gravity and washed extensively with CH_3CN , and then the solvent was removed *in vacuo*.

Step 2. The crude from the previous step was dissolved in dry DCM and set stirring at 0 $^{\circ}$ C (ice bath) under an argon

atmosphere. Et₃N (5 equiv.) and Boc₂O (4 equiv.) were added, and the ice bath was removed after 30 min. The reaction was monitored by TLC using the following systems: CHCl₃: MeOH:MeNH₂ (33% MeNH₂ in EtOH) 30:4:1.3 and DCM: MeOH:Et₃N 30:1:0.2. The reaction mixture was filtered under gravity and washed extensively with DCM, and then the solvent was removed *in vacuo*. The crude product was then loaded onto a DCM-packed silica column as a DCM solution; the desired product was eluted using a DCM: MeOH:Et₃N (30:1:0.1) solvent system. A post-column workup was then performed with DCM and water; the desired product was isolated in the organic phase, dried with MgSO₄ and then evaporated to yield the pure compound.

Step 3. The third step was carried out according to a previously reported procedure.20 To a stirred solution of the phthalimide derivative in absolute EtOH (20 mL g⁻¹ phthalimide derivative) were added anhydrous K2CO3 (2.7 equiv.) and MeNH₂ (20 mL g⁻¹ phthalimide, 33% MeNH₂ in EtOH) at 0 °C (ice bath) under an argon atmosphere. The suspension was then left to stir in the ice bath for 30 min before the system was allowed to return gradually back to r.t. The reaction was monitored by TLC using the same system as in step 1. The crude suspension was then filtered and washed extensively with ethanol, and the solvent was removed by evaporation in vacuo. The residue was purified by silica chromatography using a MeOH/ MeNH₂ elution system to yield pure compounds 13b-d as faintly yellow solids. Chemical yields and analytical data of 13b-d and their precursor intermediates are given in the ESI.†

General procedure for the synthesis of compounds 14a-d

The compounds were prepared from the free-amine derivative using a procedure previously reported by our group. To an anhydrous CH₃CN (50 mL g⁻¹ 13a-d) solution of the appropriate primary amine (13a-d) were added the reagents *N*,*N*-di-Boc-1*H*-pyrazole-1-carboxamide (1 equiv.) and anhydrous K₂CO₃ (2 equiv.) and the mixture was set stirring at r.t. under an argon atmosphere. TLC analysis (CHCl₃/MeOH/Et3N 30:4.5:0.3) indicated complete conversion after overnight. The reaction mixture was filtered and washed extensively with CH₃CN and DCM, and then the solvent was removed *in vacuo*. The dry residue was purified by silica chromatography using a MeOH/DCM/MeNH₂ elution system, to yield pure compounds 14a-d. Chemical yields of 14a-d and their complete analytical data are given in the ESI.†

General procedure for the synthesis of compounds 1-8

The compounds were prepared using a procedure previously reported by our group. ¹⁶ The appropriate derivative (13a–d or 14a–d) was dissolved in an aqueous solution of 6 M HCl (60 mL g $^{-1}$ starting material) and set stirring at 90 $^{\circ}$ C under argon. The reaction was monitored by TLC [MeOH/DCM/MeNH₂ (33% MeNH₂ in EtOH)/H₂O 3.3:3.3:2.3:1.1], mass spectrometry, and 1 H NMR, which indicated complete

conversion after 7 days. The crude solution was evaporated *in vacuo*, washed extensively with diethyl ether to remove any residual HCl, and then dried *in vacuo*. The dry residue was then purified using Sephadex LH20 and then lyophilized to yield pure compounds 1–8.

Compound 1

Following the general procedure, 14a (694 mg, 0.723 mmol) yielded 37% of compound 1 (225 mg, 0.273 mmol); ¹H NMR (600 MHz, D₂O pH 1) $\delta_{\rm H}$ 8.46 (s, 1H, QH-2), 7.76 (d, 2H, linker Ar), 7.67 (d, 2H, linker Ar), 7.35 (d, 1H, QH-5), 7.27 (d, 1H, QH-8), 4.50 (s, 2H, Ph- CH_2 -piperazine), 3.86-3.76 (m, 4H, CH_2N), 3.64-3.58 (m, 6H, cyclopropane CH (1H), CH₂N (5H)), 3.44-3.20 (m, 12H, CH_2 -guanidine (2H), CH_2N (10H)), 3.05 (m, 2H, CH_2CH_2 -guanidine), 3.00-2.98 (m, 1H, CH_2N), 2.81 (m, 2H, TACN-CH2-Ph), 1.34-1.33 (m, 2H, cyclopropane CH_2), 1.07 (m, 2H, cyclopropane CH_2); ¹³C NMR (150 MHz, D_2O pH 1) δ_C 175.71 (cipro C=O), 168.75 (cipro C=O), 156.62 (guanidine C), 154.04 (cipro), 152.38, 148.17 (cipro C-H), 144.04 (cipro), 143.98 (Ar), 138.82 (Ar), 132.45 (C-H linker Ar), 131.33 (linker Ar), 130.31 (C-H linker Ar), 118.64 (cipro), 110.67 (cipro C-H), (cipro C-H), 106.65 (cipro), 105.52, 60.54 (Ph-CH₂-piperazine), 59.71, 52.06, 51.39, 51.10, 49.29, 49.03, 46.64, 46.32, 45.46, 43.30, 40.19 (CH₂N), 36.90 $(CH_2$ -guanidine), 36.16 (cyclopropane CH),8.21 (cyclopropane CH₂), 7.49 (cyclopropane CH₂). HRMS (ESI+ QTOFMS) calculated for $C_{34}H_{47}FN_9O_3$ ([M + H]⁺) m/z648.3786; measured m/z 648.3781.

Compound 2

Following the general procedure, 14b (522 mg, 0536 mmol) yielded 41% of compound 2 (182 mg, 0.218 mmol); ¹H NMR (500 MHz, D_2O pH 1) δ_H 8.48 (s, 1H, QH-2), 7.47-7.44 (d, 2H, linker Ar), 7.38-7.37 (m, 4H, linker Ar (2H), QH-5 (1H), QH-8 (1H)), 4.37 (s, 2H, Ph-CH₂-piperazine), 3.84–3.74 (m, 2H, CH₂N), 3.59-3.45 (m, 7H, cyclopropane CH (1H), CH₂N (6H)), 3.34–3.18 (m, 12H, TACN– CH_2CH_2 –Ph (2H), CH_2 –guanidine (2H), CH₂N (8H)), 2.93-2.90 (m, 4H, CH₂N), 2.85-2.83 (t, CH_2CH_2 -guanidine, 2H), 2.48 (t, 2H, TACN- CH_2CH_2 -Ph), 1.31-1.29 (m, 2H, cyclopropane CH₂), 1.06-1.05 (m, 2H, cyclopropane CH₂); 13 C NMR (125 MHz, D₂O) $\delta_{\rm C}$ 175.92 (cipro C=O), 168.94 (cipro C=O), 156.74 (guanidine C), 154.30, 152.29 (cipro), 148.25 (cipro C-H), 145.02 (cipro), 143.98 (Ar), 138.94 (Ar), 131.82 (C-H linker Ar), 129.70 (C-H linker Ar), 126.64 (linker Ar), 118.89 (cipro), 110.80 (cipro C-H), 106.73 (cipro C-H), 105.67 (cipro), 60.03 (Ph-CH2-piperazine), 57.80, 52.52, 50.89, 50.46, 48.10, 46.77, 46.40, 43.29, 41.94 (CH₂N), 37.61 (CH_2 -guanidine), 36.12 (cyclopropane CH), 34.55, 30.55 (TACN-CH₂CH₂-Ph), 24.48 (TACN-CH₂CH₂-Ph), 8.19, 7.37 (cyclopropane CH₂). HRMS (ESI+ QTOFMS) calculated for $C_{35}H_{49}FN_9O_3$ ([M + H]⁺) m/z 662.3942; measured m/z662.3942.

Compound 3

Following the general procedure, 14c (486 mg, 0.491 mmol) yielded 34% of compound 3 (143 mg, 0.167 mmol); ¹H NMR (600 MHz, D₂O pH 1) $\delta_{\rm H}$ 8.46 (s, 1H, QH-2), 7.44 (d, 2H, linker Ar), 7.34-7.28 (m, 4H, linker Ar (2H), QH-5 (1H), QH-8 (1H)), 4.37 (s, 2H, Ph-CH₂-piperazine), 3.81 (m, 2H, CH₂N), 3.58-3.57 (m, 7H, cyclopropane CH (1H), CH_2N (6H)), 3.32–3.22 (m, 12H, $TACN-CH_2(CH_2)_2-Ph$ (2H), CH_2 -guanidine (2H), CH_2N (8H)), 2.96 (m, 4H, CH₂N), 2.87-2.86 (t, CH₂CH₂-guanidine, 2H), 2.71-2.69 (t, 2H, TACN-(CH₂)₂CH₂-Ph), 2.13-2.04 (m, 2H, TACN-CH₂-CH₂CH₂-Ph), 1.30 (m, 2H, cyclopropane CH₂), 1.05 (m, 2H, cyclopropane CH₂); 13 C NMR (150 MHz, D₂O pH 1) $\delta_{\rm C}$ 175.83 (cipro C=O), 168.86 (cipro C=O), 156.72 (guanidine C), 153.26 (cipro), 148.22 (cipro C-H), 144.07 (cipro), 142.61 (Ar), 138.87 (Ar), 131.69 (C-H linker Ar), 129.36 (C-H linker Ar), 126.14 (linker Ar), 118.74 (cipro), 110.64 (cipro C-H), 106.67 (cipro C-H), 105.58 (cipro), 60.12 (Ph-CH₂-piperazine), 57.72, 52.03, 50.87, 50.70, 50.50, 47.56, 46.64, 46.36, 43.34, 41.12 (CH₂N), 37.15 (CH₂-guanidine), 36.15 (cyclopropane CH), 31.47 (TACN- $(CH_2)_2CH_2-Ph),$ 25.26 $(TACN-CH_2CH_2CH_2-Ph),$ (cyclopropane CH2). HRMS (ESI+ QTOFMS) calculated for $C_{36}H_{51}FN_9O_3$ ([M + H]⁺) m/z 676.4099; measured m/z676.4100.

Compound 4

Following the general procedure, **14d** (403 mg, 0.402 mmol) yielded 45% of compound 4 (156 mg, 0.181 mmol); ¹H NMR (600 MHz, D₂O pH 1) $\delta_{\rm H}$ 8.41 (s, 1H, QH-2), 7.43-7.32 (m, 6H, linker Ar (4H), QH-5 (1H), QH-8 (1H)), 4.36 (s, 2H, Ph- CH_2 -piperazine), 3.83-3.53 (m, 9H, cyclopropane CH (1H), CH_2N (8H)), 3.36–3.25 (m, 12H, $TACN-CH_2(CH_2)_3-Ph$ (2H), CH₂-guanidine (2H), CH₂N (8H)), 2.82 (m, 8H, CH₂N (4H), CH_2CH_2 -guanidine (2H), $TACN-(CH_2)_3CH_2$ -Ph (2H)), 1.73-1.61 (m, 4H, TACN-CH₂(CH_2)₂ CH_2 -Ph), 1.31 (m, 2H, cyclopropane CH₂), 1.04-1.02 (m, 2H, cyclopropane CH₂); ¹³C NMR (150 MHz, D₂O pH 1) $\delta_{\rm C}$ 175.68 (cipro C=O), 168.70 (cipro C=O), 156.78 (guanidine C), 148.07 (cipro), 147.04 (cipro C-H), 144.39 (cipro), 143.94 (Ar), 138.78 (Ar), 131.56 (C-H linker Ar), 129.35 (C-H linker Ar), 125.58 (linker Ar), 121.75 (cipro), 110.68 (cipro C-H), 106.60 (cipro C-H), 105.51 (cipro), 60.13 (Ph-CH₂-piperazine), 58.49, 54.76, 53.65, 53.10, 51.96, 51.85, 50.78, 48.07, 46.54, 46.37 (CH₂N), 38.70 (CH₂-guanidine), 36.14 (cyclopropane CH), 34.14 (TACN- $(CH_2)_3CH_2$ -Ph), 27.41, 24.50-23.02 (TACN- $CH_2(CH_2)_2CH_2$ -Ph), 7.38 (cyclopropane CH_2). HRMS (ESI+ QTOFMS) calculated for $C_{37}H_{53}FN_9O_3$ ([M + H]⁺) m/z690.4255; measured m/z 690.4253.

Compound 5

Following the general procedure, **13a** (268 mg, 0.373 mmol) yielded 48% of product 5 (140 mg, 0.179 mmol); 1 H NMR (600 MHz, D₂O pH 1) $\delta_{\rm H}$ 8.6 (s, 1H, QH-2), 7.63–7.54 (m, 4H, linker Ar), 7.41–7.39 (m, 2H, QH-8 (1H), QH-5 (1H)), 4.37 (s,

2H, Ph- CH_2 -piperazine), 3.80–3.78 (m, 2H, cyclopropane CH (1H), CH₂N (1H)), 3.51–3.44 (m, 7H, CH₂N), 3.33–3.28 (m, 7H, CH₂N) (7H)), 3.18–3.00 (m, 3H, CH₂N), 2.91–2.86 (m, 6H, CH₂N), 2.39 (t, 2H, TACN- CH_2 -Ph), 1.23–1.19 (m, 2H, cyclopropane CH₂), 1.02 (m, 2H, cyclopropane CH₂); ¹³C NMR (150 MHz, D₂O pH 1) δ_C 174.95 (cipro C=O), 167.82 (cipro C=O), 153.02 (cipro), 147.34 (cipro C-H), 142.85 (cipro), 142.44 (Ar), 137.72 (Ar), 131.06 (C-H linker Ar), 127.08 (C-H linker Ar), 125.77 (linker Ar), 117.96 (cipro), 111.75 (cipro C-H), 106.89 (cipro C-H), 104.89 (cipro), 61.23 (Ph- CH_2 -piperazine), 58.95, 51.17, 45.98, 44.78, 43.56, 42.89 (CH₂N), 35.78 (cyclopropane CH_2 H), 35.16 (CH_2 -NH₂), 31.15 (TACN- CH_2 -Ph), 7.22 (cyclopropane CH_2). HRMS (ESI+QTOFMS) calculated for $C_{33}H_{45}$ FN₇O₃ ([M + H]⁺) m/z 606.3568; measured m/z 606.3564.

Compound 6

Following the general procedure, 13b (310 mg, 0.423 mmol) yielded 45% of compound 6 (154 mg, 0.194 mmol); ¹H NMR (600 MHz, D₂O pH 1) $\delta_{\rm H}$ 8.39 (s, 1H, QH-2), 7.40-7.38 (d, 2H, linker Ar), 7.32-7.21 (m, 4H, linker Ar (2H), QH-8 (1H), QH-5 (1H)), 4.31 (s, 2H, Ph-CH₂-piperazine), 3.74-3.69 (m, 2H, cyclopropane CH (1H), CH₂N (1H)), 3.51-3.44 (m, 7H, CH₂N), 3.26-3.17 (m, 9H, TACN-CH₂CH₂-Ph (2H), CH₂N (7H)), 3.11-3.09 (m, 3H, CH₂N), 2.93-2.91 (m, 6H, CH₂N), 2.40 (m, 2H, TACN-CH₂CH₂-Ph), 1.23-1.20 (m, 2H, cyclopropane CH₂), 1.00-0.97 (m, 2H, cyclopropane CH₂); ¹³C NMR (150 MHz, D_2O) δ_C 176.12 (cipro C=O), 169.01 (cipro C=O), 152.38 (cipro), 147.89 (cipro C-H), 144.97 (cipro), 138.72 (Ar), 131.05 (C-H linker Ar), 129.02 (C-H linker Ar), 125.24 (linker Ar), 118.54 (cipro), 110.65 (cipro C-H), 106.84 (cipro C-H), 106.01 (cipro), 61.18 (Ph-CH₂-piperazine), 58.48, 51.74, 50.56, 49.42, 47.48, 44.07, 42.59, 40.97 (CH₂N), 36.08 (cyclopropane CH), 34.50 (CH_2-NH_2) , 28.39 $(TACN-(CH_2)_2CH_2-Ph)$, 24.91 $(TACN-CH_2CH_2CH_2-Ph),$ 22.85 $(TACN-CH_2CH_2CH_2-Ph),$ (cyclopropane HRMS (ESI+ QTOFMS) CH_2). calculated for $C_{34}H_{47}FN_7O_3$ ([M + H]⁺) m/z 620.3724; measured m/z 620.3720.

Compound 7

Following the general procedure, 13c (331 mg, 0.443 mmol) yielded 42% of compound 7 (152 mg, 0.186 mmol);

¹H NMR (600 MHz, D₂O pH 1) $\delta_{\rm H}$ 8.46 (s, 1H, QH-2), 7.44 (d, 2H, linker Ar), 7.35–7.34 (m, 3H, linker Ar (2H), QH-8 (1H)), 7.30–7.28 (d, 1H, QH-5), 4.38 (s, 2H, Ph-C \underline{H}_2 -piperazine), 3.84–3.82 (m, 2H, cyclopropane CH (1H), CH₂N (1H)), 3.60–3.57 (m, 7H, CH₂N), 3.35–3.25 (m, 9H, TACN-C \underline{H}_2 (CH₂)₂-Ph (2H), CH₂N (7H)), 3.16–3.12 (m, 3H, CH₂N), 2.97–2.94 (m, 6H, CH₂N), 2.72–2.70 (t, 2H, TACN-(CH₂)₂C \underline{H}_2 -Ph), 2.14–2.06 (m, 2H, TACN-CH₂C \underline{H}_2 CH₂-Ph), 1.32–1.31 (m, 2H, cyclopropane CH₂), 1.06 (m, 2H, cyclopropane CH₂);

¹³C NMR (150 MHz, D₂O pH 1) $\delta_{\rm C}$ 175.80 (cipro C=O), 168.84 (cipro C=O), 153.28 (cipro), 148.21 (cipro C-H), 144.07 (cipro), 142.56 (Ar), 138.86 (Ar),

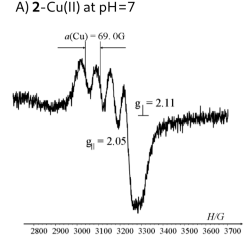
131.66 (C–H linker Ar), 129.36 (C–H linker Ar), 126.18 (linker Ar), 118.76 (cipro), 110.64 (cipro C–H), 106.66 (cipro C–H), 105.56 (cipro), 60.15 (Ph– \underline{C} H₂–piperazine), 57.89, 50.91, 50.68, 50.28, 47.34, 46.38, 43.67, 43.19, 42.06, 41.27 (CH₂N), 36.14 (cyclopropane \underline{C} H), 34.49 (\underline{C} H₂–NH₂), 31.45 (TACN–(CH₂) \underline{C} H₂–Ph), 25.17 (TACN–CH₂ \underline{C} H₂–Ph), 7.47 (cyclopropane \underline{C} H₂). HRMS (ESI+ QTOFMS) calculated for C₃₅H₄₉FN₇O₃ ([M + H]⁺) m/z 634.3881; measured m/z 634.3882.

Compound 8

Following the general procedure, 13d (240 mg, 0.316 mmol0) yielded 47% of product 8 (125 mg, 0.152 mmol); ¹H NMR (500 MHz, D₂O pH 1) $\delta_{\rm H}$ 8.35 (s, 1H, QH-2), 7.43-7.34 (m, 6H, linker Ar (4H), QH-8 (1H), QH-5 (1H)), 4.37 (s, 2H, Ph- CH_2 -piperazine), 3.53-3.48 (m, 9H, CH_2N (8H), cyclopropane CH (1H)), 3.30-3.25 (m, 9H, TACN-CH₂(CH₂)₃-Ph (2H), CH₂N (7H)), 3.18-3.15 (m, 6H, CH₂N), 2.99-2.95 (m, 3H, CH₂N), 2.72-2.66 (m, 2H, TACN-(CH₂)₃CH₂-Ph), 1.75-1.64 (m, 4H, TACN-CH₂(CH₂)₂CH₂-Ph), 1.31 (m, 2H, cyclopropane CH₂), 1.01 (m, 2H, cyclopropane CH₂); 13 C NMR (125 MHz, D₂O) $\delta_{\rm C}$ 175.52 (cipro C=O), 168.58 (cipro C=O), 152.53 (cipro), 147.92 (cipro C-H), 144.56 (cipro), 138.68 (Ar), 131.59 (C-H linker Ar), 129.37 (C-H linker Ar), 125.61 (linker Ar), 118.65 (cipro), 110.63 (cipro C-H), 106.50 (cipro C-H), 105.60 (cipro), 60.16 (Ph-CH₂-piperazine), 57.78, 51.29, 50.83, 50.73, 47.78, 46.79, 43.71, 43.33, 42.09, 41.64 (CH₂N), (cyclopropane CH), 35.02, 34.20 (CH₂-NH₂), 27.56 (TACN- $(CH_2)_3CH_2-Ph)$, 25.68 $(TACN-CH_2CH_2CH_2-Ph)$, 23.62 (TACN-CH₂CH₂CH₂CH₂-Ph), 7.53 (cyclopropane CH₂). HRMS (ESI+ QTOFMS) calculated for $C_{36}H_{51}FN_7O_3$ ([M + H]⁺) m/z648.4037; measured m/z 648.3993.

Compound 12

To a stirred solution of N,N,N-bis-Boc-phthalimide-TACN¹⁹ (2.07 g, 4.12 mmol) in dry DCM (40 mL) was added neat TFA (42 mL) at room temperature under an argon atmosphere. The reaction was monitored by TLC using the following system: CHCl₃:MeOH:MeNH₂ (33% MeNH₂ EtOH) 24:4.5:1.5, which indicated complete conversion after 40 min. The reaction mixture was then set stirring in an ice bath, and diethyl ether was added mL), causing carefully (150)immediate precipitation of the desired product. The reaction mixture was then filtered using a Buchner apparatus and washed thoroughly with diethyl ether to yield a white solid (1.99 g, 3.75 mmol, 91%); ¹H NMR (600 MHz, D_2O) δ_H 7.78-7.63 (m, 4H, phthalimide Ar), 3.75 (t, 2H, CH_2 -phthalimide), 3.54 (s, 4H, TACN CH_2N), 3.26 (t, TACN CH_2N), 3.01 (t, 4H, TACN CH_2N), 2.91 (t, 2H, CH_2CH_2 -phthalimide); ¹³C NMR (150 MHz, D_2O) δ_C 170.50 (phthalimide C=O), 162.89 (TFA C=O), 134.82 (phthalimide Ar), 131.18 (phthalimide Ar), 123.40 (phthalimide Ar), 116.30 (TFA CF₃), 51.80 (CH₂N), 47.33 (CH₂N), 43.68 (CH₂N), 42.06 (CH_2N) , 33.83 $(CH_2$ -phthalimide). MS (ESI+ QTOFMS)



B) **6**-Cu(II) at pH=7

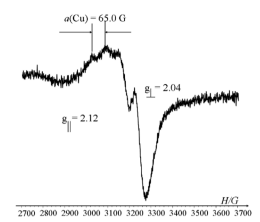


Fig. 7 The spectra shown above are as follows: (A) solid state 2–Cu(II) at 300 K ($g_{perpendicular} = 2.11$; $g_{parallel} = 2.05$, $A_{parallel} (^{63,65}Cu) = 69.0$ G); (B) solid state 6-Cu(II) at 300 K ($g_{perpendicular} = 2.04$; $g_{parallel} = 2.12$, $A_{parallel}(^{63,65}Cu) = 65.0$ G).

calculated for $C_{16}H_{23}N_4O_2$ ([M-H-2TFA]⁺) m/z 303.18; measured m/z 303.22.

Compound 10

The compound 1-(2-aminoethyl)-4,7-bis(tert-butoxycarbonyl)-TACN (390 mg, 1.06 mmol)¹⁹ was dissolved in dry DCM (40 mL) and cooled to 0 °C (ice bath). Then TFA (10 mL) was added and after 30 min, the reaction vessel was allowed to increase to r.t. The reaction was monitored by TLC using the following systems: CHCl₃:MeOH:MeNH₂ 24:4.5:1.5 and CHCl3:MeOH 9:1, which indicated complete conversion after 1 h. Then, the reaction mixture was cooled, Et₂O (450 mL) was added, and precipitation was observed. The precipitate was filtered through a Buchner apparatus and washed with Et₂O. The final crystallized compound was dried in vacuo (152 mg, 0.880 mmol, 83%); ¹H NMR (600 MHz, D_2O) δ_H 3.45 (s, 4H, NHCH₂RNH), 3.17-3.15 (m, 4H, NHCH₂CH₂N), 3.03-3.00 (t, 2H, RCH₂NH₂), 2.86-2.82 (m, 6H, NHCH₂CH₂N). ¹³C NMR (150 MHz, D₂O) $\delta_{\rm C}$ 51.01, 43.59, 42.01, 34.66. MS (ESI+ QTOFMS) calculated for C₈H₂₁N₄ ([M-H-3TFA]⁺) m/z 172.11; measured m/z 172.25.

General procedure for the synthesis and characterization of Cu(II) complexes (1–8)

A stoichiometric amount of CuCl2 was added to an aqueous solution of the hydrochloride salt of the TACN derivative (1-8). The pH was then adjusted to 10 using 1 M NaOH(aq) and then stirring took place overnight at RT. UV-vis spectroscopy ($\lambda_{max} \approx 600$ nm) was used to monitor the progress of the complexation until completion. The reaction solution was pH adjusted to 7.4 and then microfiltered to obtain the stock for analysis and testing. The resulting Cu(II) complexes were characterized by mass spectrometry (see the ESI†). In addition, one representative member each from the guanidine series (compound 2-Cu(II)) and the amine series

(compound 6-Cu(II)) was analyzed by EPR spectroscopy (Fig. 7).

Data availability

Data for this article, including experimental details, purity determination of the lead compound, and NMR and HRMS spectra, are available in the ESI.†

Author contributions

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was supported by research grants from the Israel Science Foundation founded by the Israel Academy of Sciences and Humanities (grant no. 670/20 for T. B.) and by The Israel Innovation Authority (KAMIN Program, grant no. 75044 for T. B.). We acknowledge Dr. Boris Tumansky for his assistance with the acquisition and interpretation of EPR spectra, and Dr. Guy Kamnesky for running HPLC analysis. This work was supported in part by a Technion postdoctoral fellowship for T. P. V. B. acknowledges financial support from the Ministry of Immigration Absorption and the Ministry of Science and Technology, Israel (Kamea Program).

References

- 1 S. Reardon, Nature, 2014, DOI: 10.1038/nature.2014.15135.
- 2 J. H. Kwon and W. G. Powderly, Science, 2021, 373, 471.

- 3 J. Murugaiyan, P. Anand Kumar, G. S. Rao, K. Iskandar, S. Hawser, J. P. Hays, Y. Mohsen, S. Adukkadukkam, W. A. Awuah, R. A. M. Jose, N. Sylvia, E. P. Nansubuga, B. Tilocca, P. Roncada, N. Roson-Calero, J. Moreno-Morales, R. Amin, B. Krishna Kumar, A. Kumar, A. R. Toufik, T. N. Zaw, O. O. Akinwotu, M. P. Satyaseela and M. B. M. van Dongen, Antibiotics, 2022, 11, 200.
- 4 J. Ye and X. Chen, Antibiotics, 2023, 12, 67.

Research Article

- 5 M. Terreni, M. Taccani and M. Pregnolato, Molecules, 2021, 26, 2671.
- 6 H. L. Stennett, C. R. Back and P. R. Race, Antibiotics, 2022, 11, 1237.
- 7 Z. Yu and J. A. Cowan, Chem. - Eur. J., 2017, 23, 14113-14127.
- 8 M. N. Goldmeier, S. Katz, F. Glaser, V. Belakhov, A. Khononov and T. Baasov, ACS Infect. Dis., 2021, 7, 608-623.
- 9 J. C. Joyner, W. F. Hodnick, A. S. Cowan, D. Tamuly, R. Boyd and J. A. Cowan, Chem. Commun., 2013, 49, 2118-2120.
- 10 J. L. Alexander, Z. Thompson, Z. Yu and J. A. Cowan, ACS Chem. Biol., 2019, 14, 449-458.
- 11 M. D. J. Libardo, C. De La Fuente-Nuñez, K. Anand, G. Krishnamoorthy, P. Kaiser, S. C. Pringle, C. Dietz, S. Pierce, M. B. Smith, A. Barczak, S. H. E. Kaufmann, A. Singh and A. M. Angeles-Boza, ACS Infect. Dis., 2018, 4, 1623-1634.
- 12 M. D. J. Libardo, A. A. Bahar, B. Ma, R. Fu, L. E. McCormick, J. Zhao, S. A. McCallum, R. Nussinov, D. Ren, A. M. Angeles-Boza and M. L. Cotten, *FEBS J.*, 2017, **284**, 3662–3683.
- 13 C. M. Agbale, J. K. Sarfo, I. K. Galyuon, S. A. Juliano, G. G. O. Silva, D. F. Buccini, M. H. Cardoso, M. D. T. Torres, A. M. Angeles-Boza, C. De La Fuente-Nunez and O. L. Franco, Biochemistry, 2019, 58, 3802-3812.
- 14 A. Zhao, J. Sun and Y. Liu, Front. Cell. Infect. Microbiol., 2023, 13, 1137947.
- 15 B. Smolkin, A. Khononov, T. Pieńko, M. Shavit, V. Belakhov, J. Trylska and T. Baasov, ChemBioChem, 2019, 20, 247-259.
- 16 M. N. Goldmeier, A. Khononov, V. Belakhov, T. Pieńko, N. Orbach, Y. Gilad Barzilay and T. Baasov, J. Med. Chem., 2022, 65, 14049-14065.
- 17 J. A. Collins and N. Osheroff, ACS Infect. Dis., 2024, 10, 1097-1115.
- 18 A. A. Oviatt, E. G. Gibson, J. Huang, K. Mattern, K. C. Neuman, P. F. Chan and N. Osheroff, ACS Infect. Dis., 2024, 10, 1137–1151.
- 19 L. Tjioe, T. Joshi, C. M. Forsyth, B. Moubaraki, K. S. Murray, J. Brugger, B. Graham and L. Spiccia, Inorg. Chem., 2012, 51, 939-953.
- 20 S. Mahboobi, W. Wagner and T. Burgemeister, Arch. Pharm., 1995, 328, 371-376.
- 21 M. I. Page and W. P. Jencks, Proc. Natl. Acad. Sci. U. S. A., 1971, 68, 1678-1683.
- 22 J. C. Joyner, J. Reichfield and J. A. Cowan, J. Am. Chem. Soc., 2011, 133, 15613-15626.
- 23 W. Xu, X. Yang, L. Yang, Z. L. Jia, L. Wei, F. Liu and G. Y. Lu, New J. Chem., 2010, 34, 2654-2661.

- 24 M. Fang, L. Wei, Z. Lin and G. Y. Lu, Chin. J. Chem., 2014, 32, 142-150.
- 25 V. Pokrovskaya, V. Belakhov, M. Hainrichson, S. Yaron and T. Baasov, J. Med. Chem., 2009, 52, 2243-2254.
- 26 C. Bannwarth, S. Ehlert and S. Grimme, J. Chem. Theory Comput., 2019, 15, 1652-1671.
- 27 P. Pracht, F. Bohle and S. Grimme, Phys. Chem. Chem. Phys., 2020, 22, 7169-7192.
- 28 S. Grimme, F. Bohle, A. Hansen, P. Pracht, S. Spicher and M. Stahn, J. Phys. Chem. A, 2021, 125, 4039-4054.
- 29 F. Neese, Wiley Interdiscip. Rev.: Comput. Mol. Sci., 2022, 12, e1606.
- 30 T. Gerlach, S. Müller, A. G. de Castilla and I. Smirnova, Fluid Phase Equilib., 2022, 560, 113472.
- 31 C. A. Fitch, G. Platzer, M. Okon, B. E. Garcia-Moreno and L. P. McIntosh, Protein Sci., 2015, 24, 752-761.
- 32 A. Marlin, A. Koller, E. Madarasi, M. Cordier, D. Esteban-Gómez, C. Platas-Iglesias, G. Tircsó, E. Boros, V. Patinec and R. Tripier, Inorg. Chem., 2023, 62, 20634-20645.
- 33 L. Pazderová, V. Kubíček, J. Kotek and P. Hermann, Z. Anorg. Allg. Chem., 2021, 647, 1261-1268.
- 34 D. A. Case, H. M. Aktulga, K. Belfon, D. S. Cerutti, G. A. Cisneros, V. W. D. Cruzeiro, N. Forouzesh, T. J. Giese, A. W. Götz, H. Gohlke, S. Izadi, K. Kasavajhala, M. C. Kaymak, E. King, T. Kurtzman, T. S. Lee, P. Li, J. Liu, T. Luchko, R. Luo, M. Manathunga, M. R. Machado, H. M. Nguyen, K. A. O'Hearn, A. V. Onufriev, F. Pan, S. Pantano, R. Qi, A. Rahnamoun, A. Risheh, S. Schott-Verdugo, A. Shajan, J. Swails, J. Wang, H. Wei, X. Wu, Y. Wu, S. Zhang, S. Zhao, Q. Zhu, T. E. Cheatham, D. R. Roe, A. Roitberg, C. Simmerling, D. M. York, M. C. Nagan and K. M. Merz, J. Chem. Inf. Model., 2023, 63, 6183-6191.
- 35 C. M. Dupureur, Curr. Opin. Chem. Biol., 2008, 12, 250-255.
- 36 W. J. Allen, T. E. Balius, S. Mukherjee, S. R. Brozell, D. T. Moustakas, P. T. Lang, D. A. Case, I. D. Kuntz and R. C. Rizzo, J. Comput. Chem., 2015, 36, 1132-1156.
- 37 A. Wohlkonig, P. F. Chan, A. P. Fosberry, P. Homes, J. Huang, M. Kranz, V. R. Leydon, T. J. Miles, N. D. Pearson, R. L. Perera, A. J. Shillings, M. N. Gwynn and B. D. Bax, Nat. Struct. Mol. Biol., 2010, 17, 1152-1153.
- 38 D. R. Roe, C. Bergonzo and T. E. Cheatham, J. Phys. Chem. B, 2014, 118, 3543-3552.
- 39 W. Humphrey, A. Dalke and K. Schulten, J. Mol. Graphics, 1996, 14, 33-38.
- 40 S. Izadi, R. Anandakrishnan and A. V. Onufriev, J. Phys. Chem. Lett., 2014, 5, 3863-3871.
- 41 A. Sengupta, Z. Li, L. F. Song, P. Li and K. M. Merz, J. Chem. Inf. Model., 2021, 61, 869-880.
- 42 M. Zgarbová, J. Šponer and P. Jurečka, J. Chem. Theory Comput., 2021, 17, 6292-6301.
- 43 S. Izadi and A. V. Onufriev, J. Chem. Phys., 2016, 145, 074501.
- 44 J. Wang, R. M. Wolf, J. W. Caldwell, P. A. Kollman and D. A. Case, J. Comput. Chem., 2004, 25, 1157-1174.
- 45 R. M. Betz and R. C. Walker, J. Comput. Chem., 2015, 36, 79-87.

- 46 Y. Tao, T. J. Giese, S. Ekesan, J. Zeng, B. Aradi, B. Hourahine, H. M. Aktulga, A. W. Götz, K. M. Merz Jr. and D. M. York, J. Chem. Phys., 2024, 160, 224104.
- 47 E. Caldeweyher, S. Ehlert, A. Hansen, H. Neugebauer, S. Spicher, C. Bannwarth and S. Grimme, J. Chem. Phys., 2019, 150, 154122.
- 48 B. Hourahine, B. Aradi, V. Blum, F. Bonafé, A. Buccheri, C. Camacho, C. Cevallos, M. Y. Deshaye, T. Dumitric, A. Dominguez, S. Ehlert, M. Elstner, T. Van Der Heide, J. Hermann, S. Irle, J. J. Kranz, C. Köhler, T. Kowalczyk, T. Kubař, I. S. Lee, V. Lutsker, R. J. Maurer, S. K. Min, I. Mitchell, C. Negre, T. A. Niehaus, A. M. N. Niklasson, A. J. Page, A. Pecchia, G. Penazzi, M. P. Persson, J. Řezáč, C. G. Sánchez, M. Sternberg, M. Stöhr, F. Stuckenberg, A. Tkatchenko, V. W. Z. Yu and T. Frauenheim, J. Chem. Phys., 2020, 152, 124101.
- 49 D. M. Perreault, L. A. Cabell and E. V. Anslyn, Bioorg. Med. Chem., 1997, 5, 1209-1220.
- 50 Y. A. Kondratenko, A. A. Nikonorova, A. A. Zolotarev, V. L. Ugolkov and T. A. Kochina, J. Mol. Struct., 2020, 1207, 127813.
- 51 S. Ullrich, Z. Nazir, A. Büsing, U. Scheffer, D. Wirth, J. W. Bats, G. Dürner and M. W. Göbel, ChemBioChem, 2011, 12, 1223-1229.
- 52 R. Salvio, S. Volpi, R. Cacciapaglia, F. Sansone, L. Mandolini and A. Casnati, J. Org. Chem., 2016, 81, 9012-9019.

- 53 D. Wirth-Hamdoune, S. Ullrich, U. Scheffer, T. Radanovic, G. Dürner and M. W. Göbel, ChemBioChem, 2016, 17, 506-514.
- 54 S. Alkhader, A. Ezra, J. Kasparkova, V. Brabec and E. Yavin, Bioconjugate Chem., 2010, 21, 1425-1431.
- 55 M. R. Redinbo, L. Stewart, P. Kuhn, J. J. Champoux and W. G. J. Hol, Science, 1998, 279, 1504-1513.
- 56 S. H. Wan, F. Liang, X. Q. Xiong, L. Yang, X. J. Wu, P. Wang, X. Zhou and C. T. Wu, Bioorg. Med. Chem. Lett., 2006, 16, 2804-2806.
- 57 J. Li, J. Zhang, Q. S. Lu, Y. Yue, Y. Huang, D. W. Zhang, H. H. Lin, S. Y. Chen and X. Q. Yu, Eur. J. Med. Chem., 2009, 44, 5090-5093.
- 58 M. Q. Wang, J. L. Liu, J. Y. Wang, D. W. Zhang, J. Zhang, F. Streckenbach, Z. Tang, H. H. Lin, Y. Liu, Y. F. Zhao and X. O. Yu, Chem. Commun., 2011, 47, 11059-11061.
- 59 X. Sheng, X. M. Lu, J. J. Zhang, Y. T. Chen, G. Y. Lu, Y. Shao, F. Liu and Q. Xu, J. Org. Chem., 2007, 72, 1799–1802.
- 60 M. W. Göbel, C. D. Roussev and U. Scheffer, Helv. Chim. Acta, 2014, 97, 215-227.
- 61 R. Hettich and H.-J. Schneider, J. Am. Chem. Soc., 1997, 119, 5638-5647.
- 62 C. A. Schneider, W. S. Rasband and K. W. Eliceiri, Nat. Methods, 2012, 9, 671-675.
- 63 CLSI, Performance Standards for Antimicrobial Susceptibility Testing. 29th Ed. CLSI supplement M100, Clinical and Laboratory Standards Institute, Wayne PA, 2019.