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Viral peptide conjugates for metal-warhead delivery to chromatin†

Lucinda K. Batchelor,^{‡a} Louis De Falco,^{‡b} Paul J. Dyson^{ID}*^a and Curtis A. Davey^{ID}*^b

The presence of heavy metal groups can endow compounds with unique structural and chemical attributes beneficial for developing highly potent therapeutic agents and effective molecular labels. However, metallocompound binding site specificity is a major challenge that dictates the level of off-site targeting, which is a limiting factor in finding safer and more effective metal-based drugs. Here we designed and tested a family of metallopeptide conjugates based on two different chromatin-tethering viral proteins and a drug being repurposed for cancer, the Au(I) anti-arthritis auranofin. The viral peptides associate with the acidic patch of the nucleosome while the gold moiety can bind allosterically to the H3 H113 imidazole. To achieve synthesis of the conjugates, we also engineered a sulfur-free, nucleosome-binding Kaposi's sarcoma herpesvirus LANA peptide with a methionine-to-ornithine substitution and coupled the peptide to the metal group in a final step using click chemistry. The four conjugates tested are all selectively cytotoxic towards tumor cell lines, but the choice of viral peptide and mode of linkage to the Au(I) group influences metal binding site preference. Our findings suggest that viral peptide-metalloconjugates have potential for use in chromatin delivery of therapeutic warheads and as nucleosome-specific tags.

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Introduction

Given their ability to form covalent bonds with target biomolecules, metal-based compounds have a proclivity for high potency as cancer drugs. Nonetheless, by virtue of promoting toxicity or other side effects, off-site targeting can be a major disadvantage that limits safety and efficacy. Different methods have been employed to alter or enhance molecular site discrimination that include the attachment of a protein or peptide to the metalloagent.¹ This strategy can be used to improve cell uptake, bioavailability, and/or site selectivity, depending on the nature of the peptide.² Here, we focused on conjugates comprising nucleosome-binding viral peptides, which are based on the chromatin-tethering proteins that certain viruses express in order to maintain their genomes in, or deliver their genomes to, the cell nuclei.³

As the instructional and regulatory foundation of the cell, chromatin presides over an extraordinary variety of DNA and epigenetic therapeutic targets. Many chromatin sites are targets of metal-based drugs, notably anticancer agents like cisplatin

and oxaliplatin.⁴ Nonetheless, beyond a preference for solvent-accessible N7 sites of guanine, these platinum drugs bind otherwise indiscriminately to the DNA.⁵ Much recent work has focused on finding more site selective metallocompounds, and we earlier discovered that RAPTA [(arene)Ru(1,3,5-triaza-7-phosphaadamantane)Cl₂] anti-metastasis/tumor drugs^{6,7} and RAPTA-like agents have a preference for binding to defined histone protein sites, as opposed to the DNA, of chromatin.^{8–10}

RAPTA compounds bind to an epitomal regulatory site of chromatin, the “acidic patch” of the nucleosome^{8–10}—a cleft region on the surface of the H2A–H2B dimer having an abundance of aspartate and glutamate residues.¹¹ We subsequently discovered that RAPTA-T (T = toluene) binding at the acidic patch promotes reaction of auranofin [(3,4,5-triacetyloxy-6-acetyloxymethyl,oxane-2-thio-ate)Au(triethyl-phosphonium)], through an allosteric mechanism, at two 36+ Å-distant symmetry-related imidazole sites, H3 H113 (AU1) and H3' H113 (AU1'), on the H3–H4 tetramer.¹² Auranofin is an antiarthritic drug that is being repurposed for the treatment of cancer and other diseases,¹³ and we found that, in combination with RAPTA-T, it also has a synergistic effect in killing tumor cells.¹² More recently, we demonstrated that conjugating a RAPTA group to an auranofin-like group, ([Au(4-diphenylphosphanylbenzoic acid)X], AuDPPBX; where X = Cl), via a long polyethylene glycol (PEG) linker yields a potent heterobimetallic compound that crosslinks the H2A–H2B dimer with the H3–H4 tetramer in chromatin.¹⁴ In the current study, we aimed to develop a system for highly specific targeting of nucleosomes

^aInstitut des Sciences et Ingénierie Chimiques, École Polytechnique Fédérale de Lausanne (EPFL), 1015 Lausanne, Switzerland. E-mail: paul.dyson@epfl.ch

^bSchool of Biological Sciences & NTU Institute of Structural Biology, Nanyang Technological University (NTU), 637551, Singapore. E-mail: cadavey@protonmail.com

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‡ Equal contributions.



with a non-covalent binding module that could be used to deliver a reactive metal group to chromatin.

Results and discussion

Synthesis of Au(I)-peptide conjugates

Four Au(I)-peptide conjugates were generated by connecting the nucleosome-binding region of either the Kaposi's sarcoma-associated herpesvirus LANA protein¹⁵ or foamy virus GAG protein¹⁶ to a AuDPPBX group *via* a long PEG linker (Fig. 1). Each of two configurations were synthesized, wherein the Au-PEG linker group was attached to the peptide N-terminus (Au-LANA and Au-GAG) *versus* C-terminus (LANA-Au and GAG-Au). In order to avoid potential interactions with the Au(I) cation, the Met residue in the wild type LANA peptide was replaced by ornithine (Orn; Fig. 1 and below). To produce the conjugates, the first synthetic route tested entailed typical amide coupling approaches to connect the Au-PEG group to the peptide, but this resulted in extensive product degradation (Scheme S1†).

In order to circumvent the problematic deprotection step involving the Au(I) center, a synthetic route was devised in

which the complex and the deprotected peptide were conjugated using copper click chemistry.¹⁷ This route involves the introduction of a propargylglycine residue to the peptide and a terminal azide to the complex. The Au(I) complex, containing a long polyethylene glycol linker and a terminal azide, was prepared in five steps (Scheme 1; ESI Experimental details†). The phosphorus ligand, 4-(diphenylphosphino)benzoic acid, was coupled to octaethylene glycol *via* an esterification reaction with *N*-ethyl-*N'*-(3-dimethylaminopropyl)carbodiimide hydrochloride (EDCI) and 4-(dimethylamino)pyridine (DMAP) acting as coupling reagent and base, respectively. The equivalents of reagents were adapted, 1 equiv. 4-(diphenylphosphino)benzoic acid *versus* 1.5 equiv. octaethylene glycol, to promote the formation of the mono-phosphine product, **1**. A carboxylic acid, necessary for the addition of a second octaethylene glycol to the linker, was introduced to the ligand using succinic anhydride under basic, triethylamine conditions to afford **2**. The second octaethylene glycol unit was introduced under the same EDCI/DMAP coupling conditions to yield **3**, substantially increasing the length of the linker. The phosphorus ligand was coordinated to the gold centre *via* a freshly prepared Au(I)

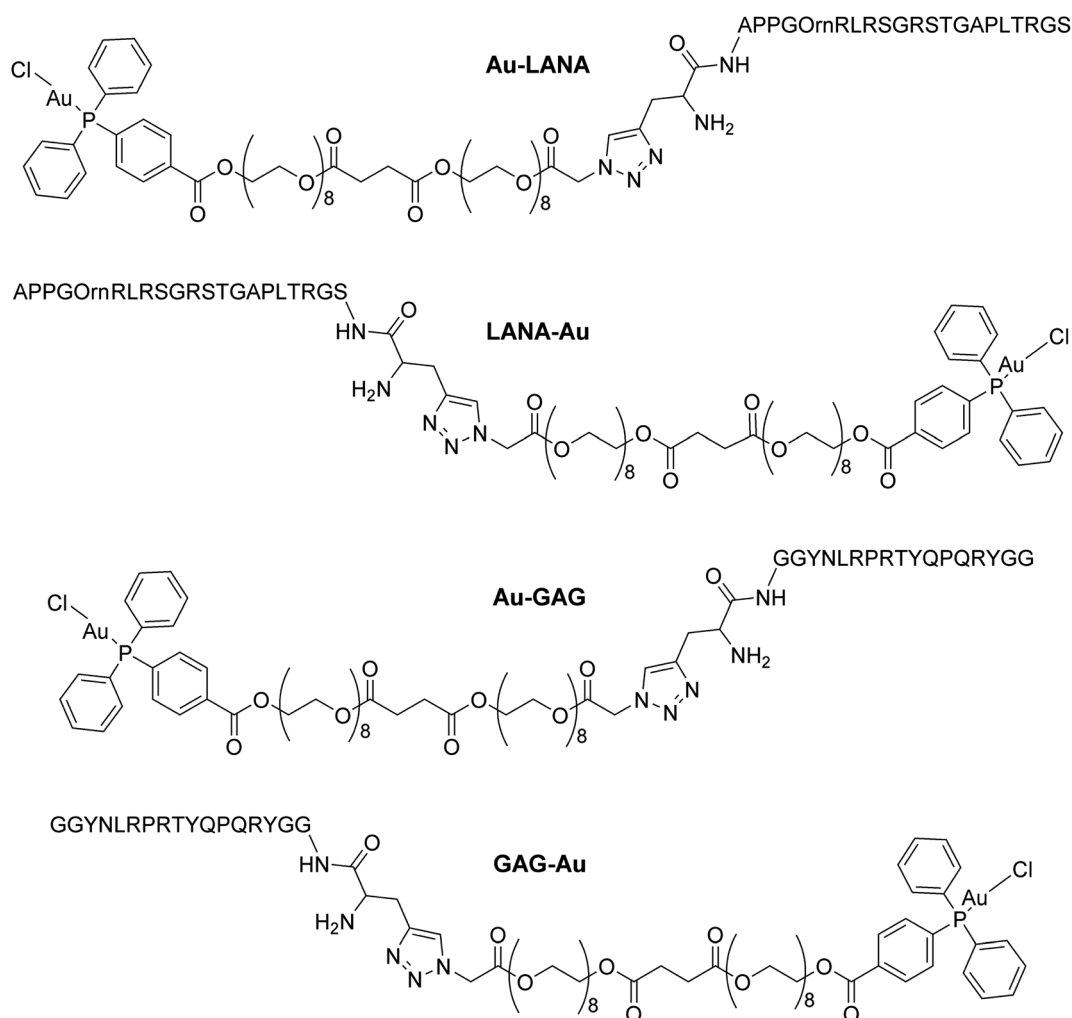
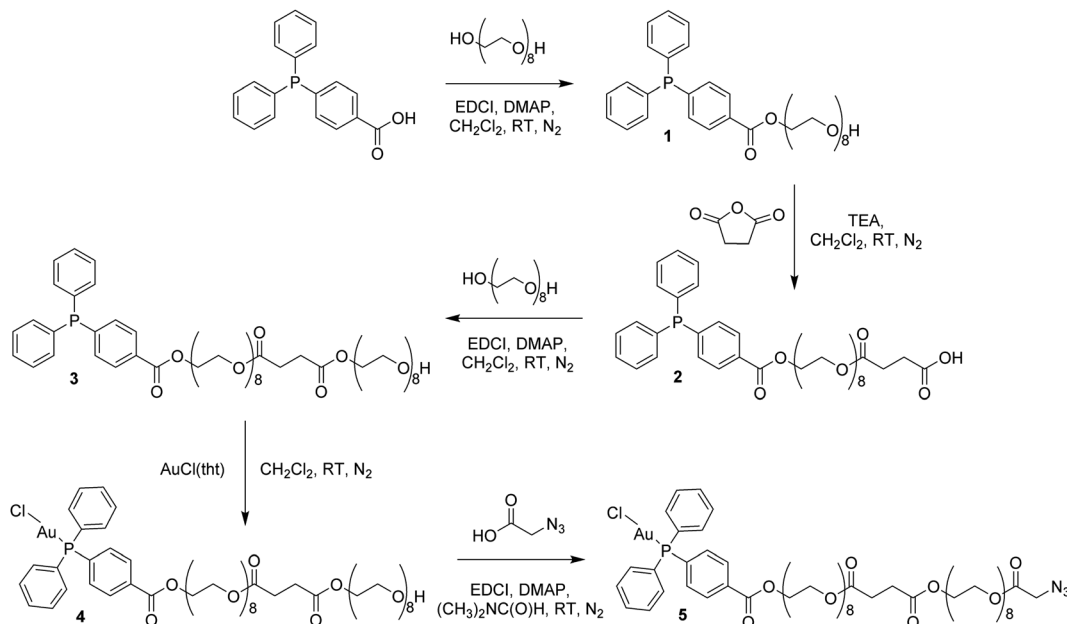


Fig. 1 Structures of the four Au(I)-viral peptide conjugates.



Scheme 1 Synthesis of the Au(I) complex containing a terminal azide, 5.

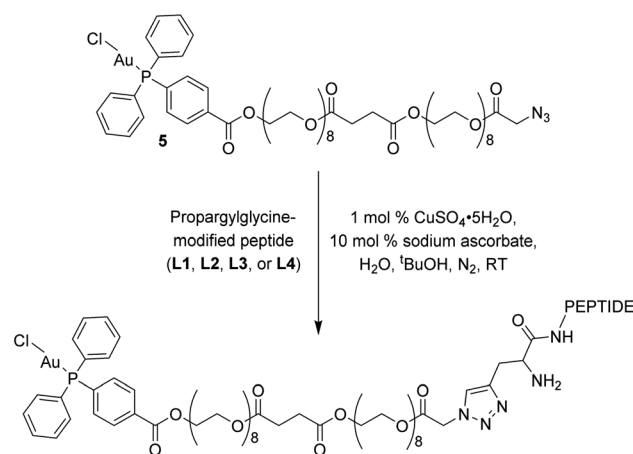
intermediate, AuCl(tht) (tht = tetrahydrothiophene). Finally, the terminal alkyne was introduced *via* an esterification reaction between 4 and 2-azidoacetic acid under EDCI/DMAP conditions in *N,N*-dimethylformamide (DMF) to yield 5. Each product was purified *via* flash column chromatography conducted in the normal phase using CH₂Cl₂/CH₃OH as eluent and isolated as viscous oils.

Compounds 1–5 were fully characterized using ¹H, ³¹P and ¹³C NMR spectroscopy (Fig. S1–S5[†]), mass spectrometry and elemental analysis. The complexation of the phosphorus ligand to the Au(I) centre was confirmed *via* a downfield shift in the ³¹P NMR spectra from –5.07 ppm for 3 to 32.97 ppm for 4. The successful synthesis of 5 was confirmed by NMR spectroscopy, where a characteristic multiplet representing the CH₂–O–(C=O)–CH₂–N₃ protons is observed at 4.31–4.33 ppm in the ¹H NMR spectrum. The phosphorus peak is almost unchanged at 33.00 ppm in the ³¹P NMR spectrum and the [M + Na]⁺ ion was observed at *m/z* 1448.4567 in the electrospray ionization (ESI) mass spectrum.

The LANA (L1 and L2) and GAG (L3 and L4) peptides, modified at either the N-terminus (L1, L3) or the C-terminus (L2, L4) with a propargylglycine (Fig. S6[†]), were synthesized with an automatic peptide synthesizer using standard Fmoc solid phase chemistry on Rink Amide MBHA resin (25 μmol scale, 0.3 mmol per g loading). The amide coupling reactions were performed twice for each amino acid using 1-[bis(dimethylamino)methylene]-1*H*-1,2,3-triazolo[4,5-*b*] pyridinium-3-oxide hexafluorophosphate (HATU) and *N,N*-diisopropylethylamine (DIPEA). The Fmoc protecting groups were removed with piperidine (20% (v/v)) in DMF and the peptides were capped using acetic anhydride (5% (v/v)) and lutidine (6% (v/v)) in DMF. The peptides were deprotected and cleaved from the resin *via* incubation with 90% trifluoroacetic acid (TFA), 2.5% H₂O, 2.5%

thioanisole, 2.5% phenol and 2.5% ethanedithiol (EDT), and the peptides were purified *via* RP-HPLC. The successful synthesis of the peptides, L1–L4, was confirmed *via* high-resolution ESI(+) spectrometry, with the [M + 2H]²⁺ ion observed at *m/z* 1101.6240 (L1), 1101.6228 (L2), 1039.0308 (L3) and 1039.0299 (L4).

The click reactions between 5 and each of the peptides (L1–L4) were conducted in *tert*-butyl alcohol and milliQ water in the presence of 1 mol% Cu(II) sulphate and 10 mol% sodium ascorbate in milliQ water (Scheme 2).¹⁸ The resulting conjugates were purified *via* RP-HPLC and target fractions were lyophilised. The successful synthesis of the conjugates was confirmed *via* high resolution mass spectrometry, where the [M–Cl + 3H]⁴⁺ ion was observed for each conjugate at *m/z*



Scheme 2 Copper click reaction used to form the final Au-peptide conjugates: Au–LANA (derived from L1), LANA–Au (derived from L2), Au–GAG (derived from L3), and GAG–Au (derived from L4).



898.9393 (Au–LANA, derived from L1), 898.9382 (LANA–Au, derived from L2), 867.6437 (Au–GAG, derived from L3) and 867.6442 (GAG–Au, derived from L4). A shift in the ^{31}P NMR signals was observed from 33.00 ppm for the free complex to 42.31–42.67 ppm for the conjugates, indicating that the environment of the phosphorus is influenced by the introduction of the peptide. The purity of the conjugates was determined by analytical HPLC (Fig. S7†). The stability of the conjugates was assessed in 100 mM NaCl in D_2O for 72 h, monitored *via* ^{31}P NMR spectroscopy and ESI mass spectrometry. The phosphorus peak in the ^{31}P NMR spectra remained essentially unchanged (Fig. S8–S11†), and the $[\text{M} - \text{Cl} + 3\text{H}]^{4+}$ peak was observed with no discernible degradation throughout the 72 h incubation. Thus, the Au(I) peptides show good stability with no degradation observed.

Cytotoxicity of Au(I)–peptide conjugates

The cytotoxicity of the Au–peptide conjugates and the free peptide ligands was assessed against cisplatin-sensitive and cisplatin-resistant human ovarian carcinoma (A2780 and A2780cisR) and non-tumoral human embryonic kidney (HEK293) cell lines (Table 1). The Au(I) conjugate group in isolation, AuDPPBX (where $\text{X} = [3,4,5\text{-triacetyloxy-6-acetyloxymethyl,oxane-2-thiolate}]$), was also evaluated as a reference alongside auranofin, cisplatin, and RAPTA-C ($\text{C} = p\text{-cymene}$) as controls. The free ligands L1–L4 were inactive against all cell lines, with IC_{50} values $>200\ \mu\text{M}$. Although several-fold less cytotoxic than AuDPPBX, conjugating the Au(I) complex with the peptides yields moderate cytotoxicity in the 35–53 μM range against both cancer cell lines, A2780 and A2780cisR. Notably, the potency of the Au–peptide conjugates against the sensitive *versus* resistant cell lines is overall nearly equivalent. This is in contrast to the other Au(I) compounds tested, AuDPPBX and auranofin, which both

show significant cross-resistance with cisplatin. On the other hand, while AuDPPBX, auranofin, and cisplatin are (equally or) more cytotoxic to the model healthy (HEK293) cell line compared to the A2780cisR tumor cell line, the Au–peptide conjugates are all significantly less cytotoxic in this regard. Any differences observed between the Au–peptides conjugates are modest, but include a slightly higher potency for Au–GAG relative to GAG–Au against the tumoral cell lines. Also, the two GAG-based conjugates are slightly more cytotoxic towards the A2780cisR cell line compared to A2780 cells, whereas the opposite relationship is seen for the LANA-based conjugates.

Nucleosome binding of Au(I)–peptide conjugates

To confirm compound binding to and reactivity with chromatin, many trials were conducted by incubating nucleosome core particle (NCP) crystals with each of the conjugates at different concentrations and for different durations, followed by analysis with X-ray crystallography (Fig. 2, S12–S14 and Tables S1–S3†). The LANA and GAG viral peptide elements of the four conjugates associate with the nucleosome acidic patch as previously characterized (PDB accession code, 5MLU).¹⁵ Since a Met-to-Orn substitution was introduced into Au–LANA and LANA–Au, we also confirmed the ability of the modified LANA peptide in isolation to bind NCP. The Met side chain of the wide type LANA peptide binds in a cavity formed by hydrophobic elements as well as two glutamate side chains.¹⁵ In its place, the Orn side chain (having nearly the same length as the Met side chain) of bound Au–LANA and LANA–Au is sandwiched between two carboxylate groups, resulting in ion pairing with the ammonium group (Fig. 2c). Alongside association of the peptide, a substantial degree of coordination of the AuDPPB group at the AU1 site is apparent for Au–LANA, which also binds to a lesser extent at the symmetry-related AU1' (Fig. 2a and Table S3†). In addition, there is evidence of non-covalently associated AuDPPBX in the form of residual (apparently depot) binding. For the LANA–Au X-ray data, there is no evidence of AuDPPB group coordination, but this may be at least partly attributable to a more pronounced disordering effect on the crystals for this particular conjugate.

In the case of the two GAG conjugates, pronounced localization of the AuDPPBX group is mostly evident as depot binding in the vicinity of the N- or C-terminal points of peptide attachment for Au–GAG and GAG–Au, respectively (Fig. 2b and Table S3†). This depot association is likely fostered by hydrophobic interactions with GAG residues. Evidence for gold(I)–histone binding is otherwise limited to low levels of adduction at the AU1' site on the opposing nucleosome face and an imidazole site (H2B H46) proximal to the peptide N-terminus, for Au–GAG, as well as a weaker gold atom signal for GAG–Au binding to AU1. The substantial depot binding and lack of pronounced adduct formation at AU1 for the GAG conjugates, contrasting what is seen for Au–LANA, is likely attributable to a steric obstacle posed by the C-terminal Tyr residue of GAG, which binds in close proximity to AU1 (Fig. 2d).

Table 1 Antiproliferative activity of the Au–peptide conjugates, free peptides modified with a propargylglycine group (L1–L4), AuDPPBX, auranofin, cisplatin, and RAPTA-C against human ovarian carcinoma (A2780), human ovarian carcinoma cisplatin-resistant (A2780cisR) and human embryonic kidney 293 (HEK293) cell lines after 72 h exposure^a

Compound	A2780	A2780cisR	HEK293
Au–LANA	35 ± 3	42 ± 3	>100
LANA–Au	36 ± 3	46 ± 3	>100
Au–GAG	40 ± 4	36 ± 2	>100
GAG–Au	53 ± 6	50 ± 5	>100
L1	>200	>200	>200
L2	>200	>200	>200
L3	>200	>200	>200
L4	>200	>200	>200
AuDPPBX	6.9 ± 0.8	12.0 ± 2	11.7 ± 0.4
Auranofin	1.1 ± 0.1	2.6 ± 0.3	1.7 ± 0.8
Cisplatin	1.6 ± 0.9	17 ± 1.8	8 ± 1
RAPTA-C	>200	>200	>200

^a Values are given as the mean ± SD (μM).



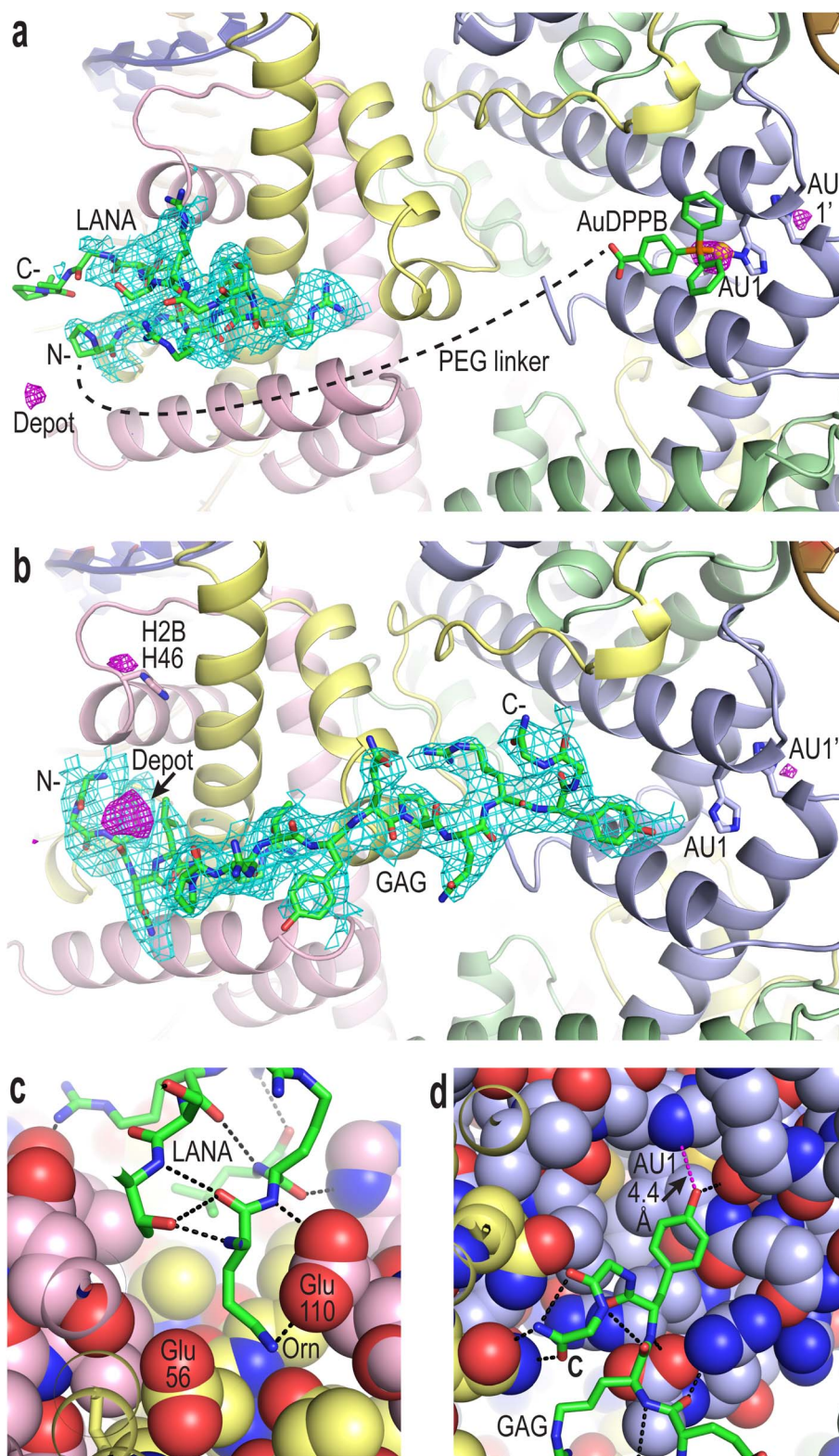


Fig. 2 Viral peptide binding and AuDPPBX coordination/localization in the nucleosome. (a and b) X-ray crystal structure of NCP treated with Au–LANA ((a); 2 N- and 5 C-terminal peptide residues not shown/visible in electron density maps) and Au–GAG (b). A $F_O - F_C$ omit electron density map (cyan; contoured at 2σ ; prior to inclusion of viral peptide) and an anomalous difference electron density map (magenta; contoured at 3.5σ ; indicates locations of gold atoms) are shown superimposed on the model (N- and C-, peptide N- and C-termini). (c and d) Close view of the LANA (c) and GAG (d) conjugate peptides binding in the NCP crystal structures (H-bonds, dashed black lines; close contact, dashed magenta line; histone protein atoms in space filling representation). (a–d) Histone proteins are shown in purple (H3), pale green (H4), yellow (H2A) and pink (H2B), and DNA strands (a and b) are orange and dark blue.



Conclusions

Here we presented a design principle and synthetic scheme for producing biologically active viral peptide metalloconjugates. The four Au(I)-peptide compounds explored are selectively cytotoxic to cancer cells, including cells with acquired resistance to cisplatin, in contrast to auranofin or the isolated AuDPPBX complex. Although the conjugates display relatively modest cytotoxicity against the tested cancer cell types, some distinctions are evident, and these may be linked to the differences in site selectivity/reactivity and impact on the NCP crystals observed between the four agents. In particular, the overall cancer cell cytotoxicity and extent of adduct formation at AU1/AU1' is highest for Au-LANA, but the latter is less than that observed for AuDPPBX or auranofin that is allosterically elicited *via* RAPTA binding in the acidic patch.^{12,14} For the GAG agents, this can be rationalized by limited steric access due to the bound peptide, but it also suggests that binding of the viral peptides does not bring about the same extent of allosteric activation. Nonetheless, we also observe gold adducts at H2B H46, and considering that the chromatin fiber *in vivo* would present a variety of nucleosomal structural contexts,¹⁹ alternative imidazole sites beyond H3 H113 (AU1/1') may also be relevant given the cellular activities observed here.

This metallopeptide conjugate system has potential for further development towards therapeutic and research tool applications. For instance, alternative metal groups with distinct site selectivity attributes could be explored, in addition to viral peptide modifications that modulate cellular uptake, localization, or stability/lability characteristics. Indeed, the distinctions in adduct formation potential and site discrimination on the nucleosome observed here that result from changes to the viral peptide sequence or disposition of coupling to the metal group may assist further conjugate design efforts; in particular, by providing insight into target site accessibility, proximity, and allosteric impact relationships. Moreover, although we did not observe substantial differences in the impact of the conjugates on cancer cells, further studies could illuminate distinctions with respect to other attributes beyond cytotoxicity.

Experimental section

Materials for synthesis

Chemical reagents were purchased from commercial sources (Sigma Aldrich, ABCR, Acros, and TCI Chemicals) and used without further purification. The reactions were performed under an inert atmosphere (N₂) using Schlenk techniques. Dry solvents, dried using a PureSolv solvent purification system (Innovative Technology Inc.), were collected and used under an inert atmosphere (N₂). Thin Layer Chromatography was conducted on Merck TLC silica gel coated aluminium sheets 60 F254 and verified by UV λ_{max} at 254 nm. Purifications were achieved by column flash chromatography using a CombiFlash Rf+ automated column machine operated with prepacked Luknova columns.

Instrumentation for synthetic procedures

¹H (400 MHz), ³¹P (162 MHz) and ¹³C (101 MHz) NMR spectra were recorded on a Bruker Avance II 400 spectrometer at 298 K. Chemical shifts are reported in parts per million (ppm) and referenced to deuterated solvent residual peaks (CDCl₃: ¹H δ 7.26, ¹³C{¹H} δ 77.16 ppm) and coupling constants (*J*) are reported in Hertz (Hz). High resolution electrospray ionization mass spectra (HR ESI-MS) were obtained on a Xevo G2-S QTOF mass spectrometer coupled to the Acquity UPLC Class Binary Solvent manager and BTN sample manager (Waters, Corporation, Milford, MA). Elemental analyses were determined on a Thermo Scientific Flash 2000 Organic Elemental Analyzer.

Compound synthesis

Details for the synthesis and characterization of the Au(I)-peptide conjugates can be found in the ESI.†

Cell culture

Human ovarian carcinoma (A2780 and A2780cisR) cell lines were obtained from the European Collection of Cell Cultures. The human embryonic kidney (HEK293) cell line was obtained from ATCC (Sigma, Buchs, Switzerland). Penicillin streptomycin, RPMI 1640 GlutaMAX (where RPMI = Roswell Park Memorial Institute), and DMEM GlutaMAX media (where DMEM = Dulbecco's modified Eagle medium) were obtained from Life Technologies, and fetal bovine serum (FBS) was obtained from Sigma. The cells were cultured in RPMI 1640 GlutaMAX (A2780 and A2780cisR) and DMEM GlutaMAX (HEK-293) media containing 10% heat-inactivated FBS and 1% penicillin streptomycin at 37 °C and CO₂ (5%). The A2780cisR cell line was routinely treated with cisplatin (2 μ M) in the media to maintain cisplatin resistance.

Cell antiproliferation activity assays

Cytotoxicity was determined using the 3-(4,5-dimethyl 2-thiazolyl)-2,5-diphenyl-2H-tetrazolium bromide (MTT) assay.²⁰ Cells were seeded in flat-bottomed 96-well plates as a suspension in a prepared medium (100 μ L aliquots and approximately 4300 cells per well) and preincubated for 24 h. Stock solutions of compounds were prepared in MilliQ water and sequentially diluted in the appropriate medium to give a final compound concentration range (0–200 μ M). Cisplatin and RAPTA-C were tested as positive (0–100 μ M) and negative (>200 μ M) controls, respectively. The compounds were added to the preincubated 96-well plates in 100 μ L aliquots, and the plates were incubated for a further 72 h. MTT (20 μ L, 5 mg mL⁻¹ in Dulbecco's phosphate buffered saline) was added to the cells, and the plates were incubated for a further 4 h. The culture medium was aspirated, and the purple formazan crystals, formed by the mitochondrial dehydrogenase activity of vital cells, were dissolved in DMSO (100 μ L per well). The absorbance of the resulting solutions, directly proportional to the number of surviving cells, was quantified at 590 nm using a SpectroMax M5e multimode microplate reader (using SoftMax Pro software, version 6.2.2). The percentage of surviving cells was calculated



from the absorbance of wells corresponding to the untreated control cells. The reported IC₅₀ values are based on the means from two independent experiments, each comprising four tests per concentration level.

Crystallographic analysis of treated nucleosome

X-ray crystallographic analysis was conducted using NCP assembled with recombinant *Homo sapiens* histones and a 145 bp DNA fragment.^{10,21} The hanging droplet method was used to grow NCP crystals from buffers containing MnCl₂, KCl and K-cacodylate [pH 6.0].²² Crystals were harvested and transferred into a stabilization buffer (37 mM MnCl₂, 40 mM KCl, 20 mM K-cacodylate [pH 6.0], 24% 2-methyl-2,4-pentanediol and 2% trehalose). MgSO₄ was substituted in place of MnCl₂ by thorough rinsing of crystals with a magnesium buffer (10 mM MgSO₄, 20 mM K-cacodylate [pH 6.0], 24% 2-methyl-2,4-pentanediol and 2% trehalose).⁸

To obtain structural data on adduct formation of the Au-peptide conjugates in the nucleosome core, native NCP crystals were subjected to incubation with magnesium buffer containing 0.5–2 mM of the conjugates. Treated crystals were mounted directly into a cryocooling N₂ gas stream set at –175 °C.⁹ X-ray diffraction data were recorded at beam line X06DA of the Swiss Light Source (Paul Scherrer Institute, Villigen, Switzerland) with a Pilatus 2M-F detector at a wavelength corresponding to the X-ray absorption edge of gold (1.04 Å). Diffraction data were indexed, integrated, merged, scaled and evaluated with a combination of iMosflm,^{23,24} XDS,²⁵ autoPROC,²⁶ SCALA,²⁷ and AIMLESS²⁸ from the CCP4 package^{29,30} and in-house data processing pipelines, *go.com* and *go.py*, developed by the Swiss Light Source macromolecular crystallography beamlines (Paul Scherrer Institute, Villigen, Switzerland).

NCP-conjugate structures were solved by molecular replacement (PHASER³¹ and MOLREP³² from the CCP4 package^{29,30}) using the 1.99 Å resolution crystal structure of native NCP, assembled with the same histones and 145 bp DNA fragment (*pdb* code 6IPU),³³ as the reference model. COOT³⁴ and REFMAC³⁵ from the CCP4 suite^{29,30} were used to conduct model building and structural refinement. Small molecule crystal structures of [Au(2-isopropylimidazole)(Pyclohexyl₃)]³⁶ and [Au(4-(diphenylphosphanyl)-benzoic acid)Cl]³⁷ were used to compose stereochemical restraint parameters for the AuDPPB-histidine adduct.

Data collection and structural refinement statistics are given in Tables S1 and S2,† and gold cation binding evaluation based on anomalous difference electron density maps is shown in Table S3.† Molecular graphics images were produced with PyMOL (DeLano Scientific LLC, San Carlos, CA, USA). Atomic coordinates and structure factors for the LANA, Au–LANA, GAG, and Au–GAG models have been deposited in the Protein Data Bank under accession codes 8Q3M, 8Q3X, 8Q3E, and 8Q36.

Conflicts of interest

There are no conflicts to declare.

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