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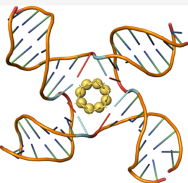


Organometallic Pillarplexes That Bind DNA 4-Way Holliday Junctions and Forks

James S. Craig, Larry Melidis, Hugo D. Williams, Samuel J. Dettmer, Alexandra A. Heidecker, Philipp J. Altmann, Shengyang Guan, Callum Campbell, Douglas F. Browning, Roland K. O. Sigel, Silke Johannsen, Ross T. Egan, Brech Aikman, Angela Casini, Alexander Pöthig,* and Michael J. Hannon*



ABSTRACT: Holliday 4-way junctions are key to important biological DNA processes (insertion, recombination, and repair) and are dynamic structures that adopt either open or closed conformations, the open conformation being the biologically active form. Tetracationic metallo-supramolecular pillarplexes display aryl faces about a cylindrical core, an ideal structure to interact with open DNA junction cavities. Combining experimental studies and MD simulations, we show that an Au pillarplex can bind DNA 4-way (Holliday) junctions in their open form, a binding mode not accessed by synthetic agents before. Pillarplexes can bind 3-way junctions too, but their large size leads them to open up and expand that junction, disrupting the base pairing, which manifests in an increased hydrodynamic size and lower junction thermal stability. At high loading, they rearrange both 4-way and 3-way junctions into Y-shaped forks to increase the available junction-like binding sites. Isostructural Ag pillarplexes show similar DNA junction binding behavior but lower solution stability. This pillarplex binding contrasts with (but



complements) that of metallo-supramolecular cylinders, which prefer 3-way junctions and can rearrange 4-way junctions into 3way junction structures. The pillarplexes' ability to bind open 4-way junctions creates exciting possibilities to modulate and switch such structures in biology, as well as in synthetic nucleic acid nanostructures. In human cells, the pillarplexes do reach the nucleus, with antiproliferative activity at levels similar to those of cisplatin. The findings provide a new roadmap for targeting higher-order junction structures using a metallo-supramolecular approach, as well as expanding the toolbox available to design bioactive junction binders into organometallic chemistry.

■ INTRODUCTION

As the repository of our genetic code, DNA is an important and fascinating target. Agents that can bind to it and potentially regulate its processing have enormous potential. Indeed, synthetic agents that target the DNA duplex such as cisplatin and doxorubicin are key agents in the clinic in the fight against many cancers.¹⁻⁴ The duplex is the most common form of DNA in the body, but when the information is being accessed and processed, a range of other structures are formed, notably the fork structures associated with DNA transactions.^{5,6} The importance of non-duplex structures in genomic DNA is increasingly recognized. Junctions^{7,8} such as the Holliday junction are involved in DNA repair and viral insertion,⁹ and G4-quadruplexes are implicated in the regulation of some gene promoters as well as in telomere stability,^{10–16} while the roles of other known cellular DNA structures such as i-motifs,^{17–19} B-Z junctions,²⁰ and three-way junctions as found at triplet repeat expansions,^{21,22} are still being elucidated. These are all attractive as targets that afford a level of selectivity in DNA binding-a structural selectivity

that complements and expands traditional attempts at sequence selective recognition.

Among these noncanonical structures, the most explored has been the G4-quadruplex, with many elegant agents reported that offer large flat planar surfaces to interact with the ends of the quadruplex stack.^{10–16,23–26} Junctions are less well studied,^{7,27,28} though we have shown that dinuclear metallosupramolecular cylinders bind inside the heart of three-way junctions (3WJ) and characterized this binding by X-ray crystallography and NMR.^{29–34} The striking features of this binding are the cavity fit and the face–face pi-stacking the surface of the cylinder makes with the DNA bases at the junction point. The key to this is that the aryl rings in the

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center of the cylinder present their aryl faces to the outside of the structure to create pi-surfaces^{27–31} and contrast with typical polypyridyl metal complexes (such as the archetypal ruthenium tris bipyridine or tris phenanthroline complexes), which present their CH lined aryl edges instead. Monchaud has screened libraries of agents for 3WJ binding and identified as his lead agent an organic 3-arm cryptand-type system with aryl rings that could potentially rotate to present a very similar aryl surface conformation to the cylinders.^{35–38} Although the binding is not yet structurally characterized, very recent MD simulations suggest this cryptand may also be able to thread into a 3WJ and bind as the cylinder does.³⁹ Other 3WJ binders^{35,37,38,40–49} that lack these outward-facing pi-surfaces are also reported—some of them may bind at the outside of (rather than within) the junction cavity.

Four-way or Holliday junctions (4WJ) are different in nature from 3WJ. While an open cruciform-style structure is possible and is seen in complex with proteins, in their absence, the junction cavity closes up, allowing the arms to come together in coaxial stacks forming a stacked-X configuration. Searcey has taken advantage of this and designed binders that contain two linked intercalators to bind two adjacent arms and with Cardin has structurally characterized this binding mode.^{50,51} Bonnet⁵² has characterized a very interesting 4WJ-like discontinuous structure assembled from a hexanucleotide and containing a small metallo-intercalator inserted between inter-duplex pairs. Binding of organic intercalators in duplex arms has also been shown adjacent to the junction point in 4WJs.^{53–55} The open cruciform-style 4WJ junction cavity has not been addressed with synthetic binders.

Based on our analysis of key binding features, we sought other designs that also offer external pi-face surfaces and thus might be suited to junction binding, and identified another new class of cationic supramolecular organometallic complexes⁵⁶ called pillarplexes.^{36,57} These structures are composed of two organic macrocyclic ligands, each with 6 aryl (imidazole/pyrazole or triazole) rings, that linearly coordinate 8 gold or silver centers (Figure 1). The pillarplexes are a similar width to the cylinder but of a more circular girth (the junction binding unit of the cylinder is in fact a twisted triangular prism that could be circumscribed by the pillarplex—see SI Figure S1). The pillarplexes are significantly shorter than the cylinders (1.2 cf. 1.9 nm), but importantly, they have the same tetracationic charge that will contribute to the binding to anionic DNA. We show herein that these pillarplex structures can bind to open DNA 4-way junctions as well as 3-way junctions and Y-shaped forks. Since they incorporate N-heterocyclic carbene ligands, they also now extend design of nucleic acid junction binders into organometallic chemistry with the different design opportunities this offers for the future.

RESULTS AND DISCUSSION

Our initial experiment was to explore whether this pillarplex structure would bind with DNA three-way junctions (3WJ). We carried out a PAGE gel with three 14-mer strands whose sequences (Figure 2) are designed to assemble a perfect 3WJ.³² In this assay, the 3WJ is not formed at room temperature in the absence of drugs, and only single-stranded DNA is observed (Figure 3 lane 1). If a drug binds in and stabilizes the 3WJ, then a new 3WJ band will be observed in the gel (as seen for the cylinder used as a positive control^{28–34} in lanes 11-14).

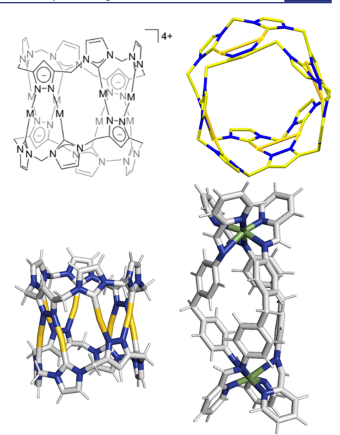


Figure 1. Top: structure of the pillarplex (M = Ag(I)/Au(I)) showing its chemical composition and a view through its cylindrical axis emphasizing its 4-fold symmetry. Bottom: a comparison of the dimensions of the Au pillarplex and the Ni(II)/Fe(II) cylinders that are compared in this work. See also Figure S1 for further comparative images.

On addition of 0.5, 1, and 2 equiv of the Au pillarplex (Figure 3, lanes 3–5), a 3WJ is formed, confirming that the Au pillarplex can bind the 3WJ. However, at 4 equiv of pillarplex (lane 6), we see the 3WJ structure is replaced by a large, smeared band corresponding to 2-stranded Y-shaped fork junctions. This suggests that while the pillarplex prefers 3WJ over Y-shaped (or partially double stranded) forks, this preference is not strong, and the pillarplex would rather bind forks than be unbound (the strands can form 50% more forks (2 strands) than perfect 3WJs (3 strands)). The broad nature of the Y-shaped band likely reflects the three different Y-structures that can be formed by the 3 strands and the flexibility of the single-stranded branches. Controls with metal-free pillarplex pro-ligands show no interactions with the DNA (Figure S2).

An interesting feature of the bound 3WJ band is that it has a slightly lower mobility in the gel compared to the 3WJ bound to the cylinder. This indicates that the 3WJ structure formed with the Au pillarplex has a larger hydrodynamic radius.

While the cylinder crystallizes with a DNA hexamer,^{29,30} attempts to crystallize the pillarplex with the same DNA hexamer were unsuccessful, and the pillarplex precipitated the DNA at the concentrations needed for NMR experiments, preventing detailed structure characterization. We, therefore, attempted to gain some atomistic insight using computational methods, using approaches previously used to explore cylinder and azacryptand 3WJ binding.^{34,39,58,59} As our starting point,

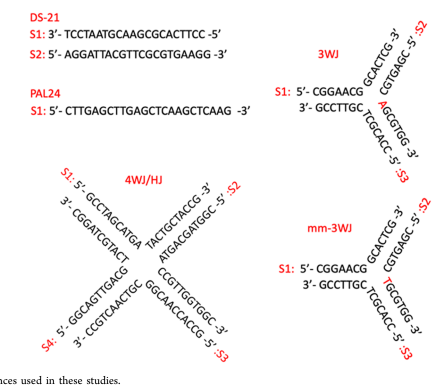


Figure 2. DNA sequences used in these studies.

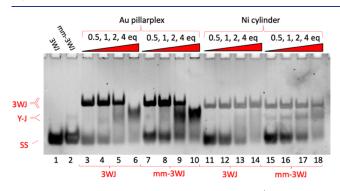


Figure 3. PAGE gel of two different DNA structures (3WJ and mm-3WJ) incubated with different complexes (Au pillarplex and Ni cylinder as the positive control) at varying ratios (all gel lanes read from left to right). The 3WJ and mm-3WJ strand sequences used are shown in Figure 2. Controls (3 strands, no complexes) are shown in lanes 1-2, then 3WJ is mixed with a complex at 0.5, 1, 2, and 4 equiv (lanes 3-6 and 11-14), and mm-3WJ is mixed with a complex at 0.5, 1, 2, and 4 equiv (lanes 7-10 and 15-18). Samples were prepared using 1 μ M of each strand and the appropriate amount of metal complex in TBN buffer (89 mM Tris, 89 mM boric acid, 10 mM NaCl, pH 7.05). Gel stained with SYBR gold. Differential staining of bands and fluorescence quenching effects are detailed in Figure S21.

we used the crystal structure of the cylinder in complex with a 3WJ formed from a palindromic DNA hexamer³⁰ (this structure-pdb 3I1D-is extremely similar to that of a 3WJ in complex with a protein⁶⁰ and thus more broadly representative). On removing the bound cylinder, the unsupported 3WJ was not stable under molecular dynamics simulations, collapsing within 1 microsecond. For this reason, we first used docking experiments to bring the pillarplex close to the 3WJ, where it sat outside the cavity, and then used this as the starting point for molecular dynamics simulations. The pillarplex reproducibly entered the cavity across 3 independent simulations (two with this 3I1D 3WJ, and one where we

mutated two of the central AT bases surrounding the cavity for GC bases-termed AGG-3WJ), adopting a position with its pisurfaces facing the walls of the cavity (Figure 4) where it remained. Simulations where the unstable unbound 3WJ DNA collapsed before entry were halted because the timescale for the reformation of the DNA structure is too long to be captured in an MD simulation of this type. A further 5 simulations were carried out with the AGG-3WJ, using a starting point of the pillarplex inside the open cavity (taken from the initial AGG-3WJ simulation). Each of the 8 simulations was of time length between 1 and 5 μ s (>18 μ s cumulative time). In each case, the binding was associated with the breaking of a base pair at the junction point to enlarge the cavity and accommodate the pillarplex. The larger width dimensions/shape of the pillarplex (compared to the cylinders) is forcing the central cavity of the 3WJ to expand, which is consistent with our experimental observation of an increased hydrodynamic radius in the gel studies.

The base-pair breaking is a dynamic process, and the AGG-3WJ was designed to allow us to probe if there was a base preference; in all simulations, transient breaking of an AT base pair was observed, while one simulation also captured a GC breaking event. After breaking, two of the simulations showed an instability in the arm with the base-pair break and collapse of the junction structure. Reasoning that this was likely an inherent instability of the short arms in the 3I1D 3WJ, three additional simulations were run using a 3WJ with longer arms (three 14-mer strands, adapted from PDB 1F44, with three AT base pairs at the branch point) for 1 μ s each. In each simulation, base-pair breaking at the junction cavity was again observed, but the arms of the 3WJ structure remained intact and stable on the timescale of the simulations, confirming that the breaking itself does not cause the 3WJ to become unstable.

By contrast, in our simulations of the cylinder with the 3WJ DNA, the cylinder resided in the cavity with no disruption to

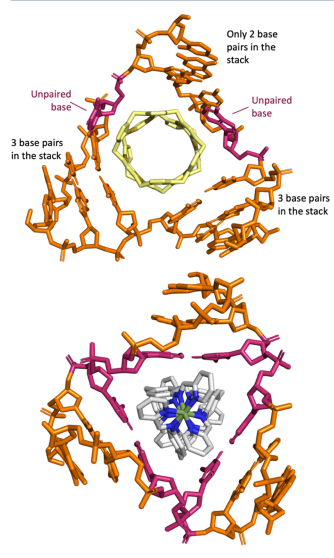


Figure 4. Top: MD simulations of Au pillarplex with palindromic DNA hexamer oligo (from pdb 311D). The MD shows Au Pillarplex inserting into the central cavity of the 3WJ, where it opens up a base pair (highlighted in pink) down one arm, thereby expanding the 3WJ cavity. Bottom: MD simulation of the Fe cylinder in the 3WJ where the central bases (pink) remain paired throughout the simulations, as observed experimentally in the X-ray crystal structures.^{28–30} Further details available in the SI and Figure S22.

the base pairing even over long simulations (4 simulations; one >10 μ s and three of 1 μ s each).

To investigate the potential junction opening further, the binding to an analogous "mismatch" 3WJ structure where the central cavity has a mismatched T-T base pair (Figure 3—mm-3WJ) was studied. This structure should promote 3WJ junction cavity expansion and allow the Au pillarplex to have a more comfortable fit, but it will also reduce the underlying stability of the 3WJ structure.^{61–64} This is reflected in the binding of the cylinder (Figure 3, lanes 15–18) where binding to both mm-3WJ and 2-stranded Y-shaped fork structures are now observed even at low loading, indicating that the mismatch has made the junction less attractive to the cylinder. By contrast, the binding of the Au pillarplex to the mismatched-3WJ is very similar to its 3WJ binding with initial mm-3WJ binding observed (and increased hydrodynamic

radius) followed by a transition to fork binding at higher loading (Figure 3, lanes 7-10).

The data confirm that the Au pillarplex can bind to and stabilize both 3WJ and Y-fork structures. The binding to 3WJ is stronger than to Y-fork (as expected purely on electrostatic grounds), but both Y-junctions (3WJ or fork) are preferred over binding to single-stranded DNA. The similarity between the Au pillarplex's observed binding to both 3WJ and mm-3WJ is consistent with the supramolecule expanding the 3WJ cavity (breaking some base-pairing at the junction).

DNA-melting experiments (monitored by the change in UVabsorption) with the 3WJ formed from 14-base sequences as used for the gels showed no melting curve in the absence of the metallo-supramolecular cations because the 3WJ is unstable at room temperature and single-stranded DNA is present throughout. By contrast, a DNA-melting curve is observed in the presence of both the pillarplex and the cylinder (Figure S3a). The melting temperature (T_m) is 56.2 (±0.1) °C with the Ni cylinder and 48.8 (± 0.5) °C with the Au pillarplex, indicating a greater stabilization of the 3WJ structure when the cylinder is bound, which is consistent with the pillarplex binding breaking base pairs. Using longer 18-base sequences, melting of the free 3WJ is observed (38.5 (± 0.7) °C), and a similar greater stabilization is observed for Ni cylinder ($T_{\rm m}$ = 59.8 (±0.6) °C; $\Delta T_{\rm m}$ = 21.3 (±0.9) °) than Au pillarplex $(T_{\rm m}$ = 56.5 (± 0.6) °C; $\Delta T_{\rm m}$ = 18.0 (± 0.9) °) (Figure S4). When the (14-base) 3WJ is replaced by (14-base) mm-3WJ (Figure S3b), the melting temperatures are now very similar for both Ni cylinder $(T_m = 46.4 (\pm 0.9) \circ C)$ and Au pillarplex $(T_m =$ 44.9 (± 0.3) °C), and these values are very close to that observed for the Au pillarplex with 3WJ. These observations are again consistent with the pillarplex breaking one base pair when it binds 3WJ.

The pillarplex has D_{2d} symmetry with a total of 12 aryl rings arranged in four sets of three, each comprising two imidazoles from one ligand and a pyrazole from the other bridging a pair of coinage metal ions (Figures 1 and S5). Given this tetragonal arrangement of the pi-surfaces and the indication that the pillarplex is a little too wide for the perfect 3WJ, we were intrigued to see whether it might also bind a four-way junction structure (4WJ). The open 4WJ form will have a larger central cavity than the 3WJ.

Analogously to the 3WJ gel studies, a 4WJ was assembled from 4 complementary strands (Figure 2) and exposed to increasing concentrations of Au pillarplex (Figure 5). The 4WJ strands were selected as they have been used by Searcey *et al.* previously to study 4WJ-binding of organic molecules using gel electrophoresis.⁵⁰ At room temperature, the strands form a mixture of 4WJ and individual single strands, with the 4WJ further stabilized by high concentrations of MgCl₂ and NaCl. A very small proportion (<1%) of a higher-order structure can just be discerned with these cations (Figure 5, lane 2).

The Au pillarplex interacts with the 4WJ, causing the appearance of a bound species running slightly slower than the unbound 4WJ. At a 1:1 ratio, both are present, and the 4WJ band is split into two bands (Figure 5, lane 4). In the absence of pillarplex, the closed stacked-X junction structure is expected, and the band shift is consistent with the bound structure being an open junction and cruciform-shaped. As the ratio of Au pillarplex to 4WJ increases, the 4WJ population shifts to favor the bound species band. At high loading, the discrete bound band is further retarded, perhaps indicating a second binding can occur but also reduces in intensity. This

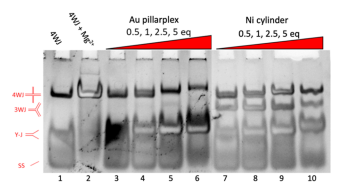


Figure 5. PAGE gels of 4WJ (S1, S2, S3, and S4) incubated with Au pillarplex and Ni cylinder at varying ratios. Controls in lanes 1-2 (lane 1: strands alone. Lane 2: in the presence of cations—2 mM Mg²⁺ and 450 mM Na⁺). Samples were prepared using 1 μ M of each strand and the appropriate amount of metal complex in TBN buffer (89 mM Tris, 89 mM boric acid, 10 mM NaCl, pH 7.05). Gel stained with SYBR gold.

decrease in intensity coincides with an increase in the intensity of a Y-fork double-stranded band, just as seen in the 3WJ gel, indicating again the pillarplex's preference for fork structures. However, at high loading, the 4WJ is still present (whereas at the same loading, the 3WJ band had disappeared—Figure 3, lane 6), and this suggests that the 4WJ may be a better binding target than 3WJ, likely due to the increased size of the central cavity. A very small proportion of discrete three-strand structure is also observed. Once again, controls with the proligand do not show binding (Figure S6).

We also investigated the binding of the Au pillarplex to the 4WJ strands individually and in all combinations (Figure 6). The pillarplex causes band smearing of the individual strands (S1-3), but for strand S4, we see induction of higher-order structures. S4 has the possibility to form a partial duplex containing two 2-base bulges (SI Figures S7 and S8) and it appears that the pillarplex binds and stabilizes this and/or other higher-order structures. The other strands do not have the capability to form such a dimer.

Where combinations of 2 strands are complementary and can form a Y-fork (i.e., S1S2; S2S3; S3S4; S1S4), the pillarplex stabilizes this structure, which is consistent with the observed binding to these structures when all 4 strands are present. All four combinations of 3 strands can form a fork-like 3WJ (or bulged 3WJ with partial interactions in one arm—SI Figure S9), but this seems only stabilized for the S2S3S4 combination and, even then, in only small amounts. Instead, when S4 is present, other *tetra*-stranded structures are stabilized (likely 4WJ containing two S4 strands—SI Figure S10). This again demonstrates the pillarplex's preference for 4WJ and duplex forks.

By contrast, when the cylinder $[Ni_2L_3]^{4+}$ interacts with the 4WJ structure, the most striking feature is the immediate formation of 3WJs and duplex Y-forks (Figure 5). There is some splitting of the 4WJ band, indicating that the cylinder can also bind to the 4WJ (and that bound and unbound species 4WJ are present), but the cylinder preference for Y-shaped structures (especially 3WJs—SI Figure S9) serves to highlight the greater comparative 4WJ preference of the pillarplex. Studies with the individual strands and their combinations (Figure 6) reinforce this observation, with the cylinder also promoting the formation of 3WJ structures in those experiments, including in two strand experiments when S4 is present.

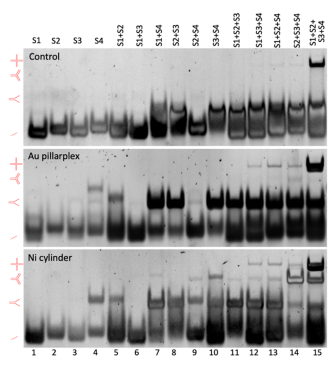


Figure 6. PAGE gels of the individual 4WJ strands (S1, S2, S3, and S4) alone and in combination, alone (top) and incubated with 1 equiv of Au pillarplex (middle) and Ni cylinder (bottom). Samples were prepared using 1 μ M of each strand and the appropriate amount of metal complex in TBN buffer (89 mM Tris, 89 mM boric acid, 10 mM NaCl, pH 7.05). Gel stained with SYBR gold.

MD simulations of Au pillarplex with a 4WJ DNA model (based on pdb 1XNS⁶⁵) were conducted. The DNA structure is a crystallographically characterized example in complex with Cre proteins and small peptides (which we removed before introducing pillarplex) and was selected as a starting point for the simulations because its junction cavity is partially open. The starting junction cavity is rectangular in shape, and at the junction, only two of four potential base pairings are present at the outset (SI Figure S11a).

From starting positions with the pillarplex outside, but close to $(\sim 1 \text{ nm})$, the junction cavity, the pillarplex was observed to rapidly (within 10 ns) enter the cavity (5 simulations from two different starting positions). With the pillarplex inside, the junction rapidly rearranged to a more open and square forman open 4WJ structure with the four DNA arms distinct and not stacked together (Figure 7). This is consistent with the small gel band shift observed. As the cavity rearranged about the pillarplex, some transient breaking of pairs and stacking of individual bases around the pillarplex was observed, as illustrated in Figure 7 (top). However, as the simulations proceeded, full base pairing at the junction point was recovered and retained (Figure 7 bottom). Importantly, during all of the simulations (three at 2 μ s each), the pillarplex remained within the central cavity throughout. In simulations in the absence of pillarplex, the free DNA rapidly closed up into the closed Holliday junction form with stacked coaxial arms, as would be expected (SI Figure S11b). In simulations starting from this closed form, we did not see pillarplex insertion over three 1 μ s simulations, though this is unsurprising, given the timescales of the dynamic opening and closing of a DNA 4WJ. This illustrates the power of sampling different known DNA starting conformations to explore the complex conformational land-

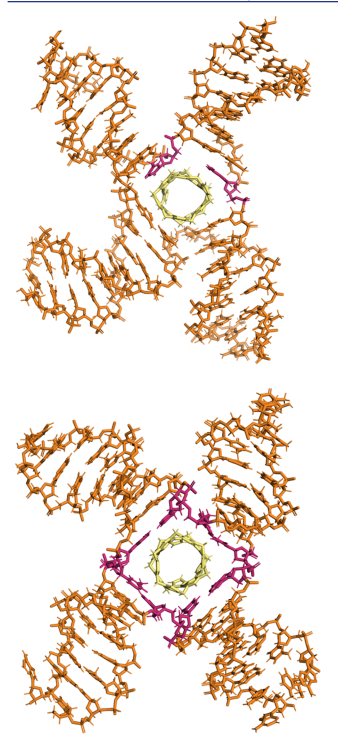


Figure 7. MD simulations of Au pillarplex with a 4WJ DNA. The MD shows Au pillarplex entering and residing in the central cavity of the 4WJ with the cavity closing up around the pillarplex but an open 4WJ structure resulting with the four arms being distinct and not stacked together. Following the entry of the pillarplex, much of the base pairing at the junction point is maintained, though initially, there is transient breaking of pairs and stacking of individual bases around the pillarplex as the cavity flexes to accommodate the pillarplex. This is seen in the upper image, where two bases (highlighted in pink) have become unpaired. As the simulations proceed, the pairings re-form and an open 4-way junction with fully paired bases contains the pillarplex. This is seen in the lower image, where the 8 bases at the junction are highlighted (pink). Further details available in the SI and Figure S22.

scapes of nucleic acids even within relatively short timescales afforded by MD simulations and is similar to simulation approaches we explored previously for RNA binding of cylinders.⁵⁸

While junctions are an exciting target, the majority of genomic DNA is found in duplex form, so we also explored the effects of pillarplex on duplex DNA, starting with a doublestranded DNA (DS21) formed from two complementary strands (Figure 2). In a gel electrophoresis experiment (Figure S13), the Au pillarplex caused a retardation of the duplex band, indicating an interaction, along with some smearing. The cylinder shows a similar (though less marked) effect, while the metal-free pillarplex pro-ligand showed no binding. Similar effects were seen with a palindromic 24-mer (Pal24). Likewise, with a longer, biological DNA (pBR-322 plasmid DNA-4361 base pairs), the linear form showed the same response of a sequential retardation as more Au pillarplex is introduced (Figure S14). DNA precipitation in the well was also observed at high loading, as for the cylinder which is known to coil and condense duplex DNA.⁶⁶⁻⁶⁸ The supercoiled, closed circular form of the plasmid DNA showed unwinding on Au pillarplex binding, with the extent again similar to that of the cylinder.^{69,70}

To probe this duplex interaction in more depth, we used flow linear dichroism (LD) with calf-thymus (CT) DNA as a representative genomic DNA. Flow LD allows the orientation of the DNA and the complex to be probed spectroscopically in a Couette cell by comparing the absorption of linearly polarized light parallel and perpendicular to the direction of flow.^{67,68,71} DNA alone shows a strong negative signal at 260 nm due to the DNA bases that become oriented as the DNA helix orients in the flow. The cylinder coils the DNA, and this leads to a loss of orientation and a dramatic decrease in the magnitude of the 260 nm signal.^{67,68} The same effect is observed with the Au pillarplex at low loading (Figure 8); however, at higher loadings (from 1 pillarplex per 20 base pairs), a second effect is superimposed as a strong positive signal appears. The signal might arise from pillarplex spectroscopy^{57c,72} (Figures S17–S19) or from the DNA bases (or both), but the chromophores are no longer orientated perpendicular to the flow direction but more parallel to the flow orientation, perhaps indicating coiling into some sort of tube conformation. The corresponding circular dichroism spectroscopy shows the B-DNA structure retained at low loading but the spectra becoming more complex at higher loading, which is likely to represent pillarplex-induced CD signals overlain on the DNA spectroscopy. The gel and spectroscopic studies show that the pillarplex is condensing duplex DNA, as other polycations do, but may induce an ordered condensed structure.

Molecular dynamics simulations of multiple pillarplexes and a 25-base pair double-stranded B-DNA (Figure 9) showed positioning of the pillarplexes along the DNA minor groove, with orientation of the pillarplexes inducing changes to the length of the groove, but did not directly inform on any DNA condensation in the timescales that can be accessed through simulations. However, a second frequent binding mode was observed, which involved an opening up of the terminal base pairings at the fraying ends of the double-stranded DNA and insertion of the pillarplex inside the DNA helix. Such a binding mode is reminiscent of both the 3WJ simulations where a base pair is opened, and the 4WJ simulations where a base pair initially breaks to better pack the bases around the pillarplex. It

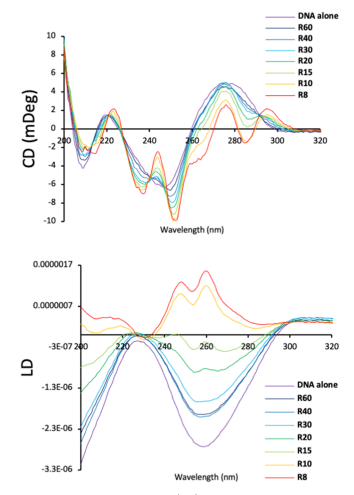


Figure 8. Circular dichroism (top) and flow linear dichroism (bottom) spectroscopic studies of Au pillarplex with CT-DNA. CT-DNA (100 μ M in base pairs), sodium chloride (10 mM), and sodium cacodylate buffer (1 mM, pH 7.3). The *R*-value is the ratio of DNA base pairs to complex.

is also a likely model of the binding to Y-fork junction structures that are observed in both the 3WJ and 4WJ gel electrophoresis experiments. It is clear from the experimental data that the pillarplex prefers such junction structures over a conventional duplex and so a potential role in the DNA condensation is an interesting possibility.

Melting curves of the 4WJ in complex with pillarplex and cylinder (Figure S12) are complicated by biphasic melting involving multiple solution species (biphasic 4WJ melting has also been observed in other studies $^{73-75}$), so we used a gel competition assay to compare the binding preference of pillarplex for the different DNA structures. 3WJ was fluorescently labeled with FAM (carboxyfluorescein) on S1. Since the 3WJ is not formed in the absence of pillarplex, the intensity of its electrophoresis band reflects the extent to which the competitors have scavenged the pillarplex. The data (Figure 10) show that the pillarplex preferentially binds 4WJ and Y-forks, with a preference for 4WJ but a similar affinity for both. This preference and affinity is consistent with the 4WJ gel studies (Figure 5), where binding to Y-forks is observed at increased loading. While some weaker dsDNA binding is also observed, the pillarplex prefers 4WJ and Y-fork over 3WJ, and 3WJ over dsDNA.

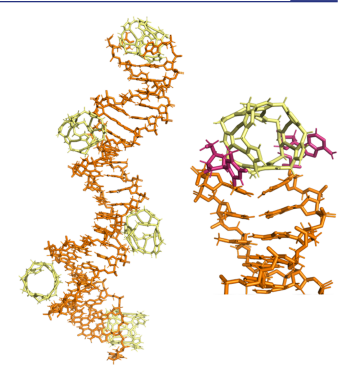


Figure 9. MD simulations of Au pillarplexes with a double-stranded 25-base pair DNA oligonucleotide. Pillarplexes can be seen binding in the minor groove (left) as well as at the fraying ends of the DNA (right—with unpaired bases highlighted in pink).

The corresponding assay with Ni cylinder shows a preference for junctions (3WJ and 4WJ) over Y-forks, with dsDNA not a competitor at these ratios. Figures 5 and 6 have already shown that the cylinder prefers and can create 3-stranded "junction" structures from the 4WJ strands. In the Figure 10 competition experiment, a labeled perfect 3WJ from 14-mer oligos is thus also competing with an imperfect 3-strand junction (Figure S20), which (being formed from 22-mers) is more polyanionic.

We have also studied the DNA binding of the corresponding Ag pillarplex structure (SI Figures S2, S6, and S13–15). This silver organometallic complex has the same shape and charge as the Au pillarplex, but lower stability.⁷² The Ag and Au pillarplexes are stable in aqueous solution, but for the Ag pillarplex, potential complex degradation (e.g., in the presence of high chloride concentrations)⁷² and lower water solubility are complicating factors in the experiments. Nevertheless, the DNA binding is similar to that of the Au pillarplex, confirming the importance of the structural motif in DNA binding.

To explore whether the Au pillarplex can enter the cell and access cellular DNA, we explored the uptake and accumulation in A549 lung cells by treating with the compound (6 μ M) for 24 h, quantifying the intracellular metal content by inductively coupled plasma mass spectrometry (ICP-MS), and also fractionating to isolate the nucleus and assess its Au content. The initial results confirm that the Au pillarplex can enter the cell with the amount of Au accumulated in the cell extract (ca. 0.08 ng metal/ μ g protein) very similar to the amount of platinum from cisplatin (used as a control). Approximately 10% of that Au is found in the nuclei after 24 h, which is again comparable with cisplatin.

Antiproliferative activity studies conducted on both Au and Ag pillarplexes against human A549 lung and SKOV-3 ovarian

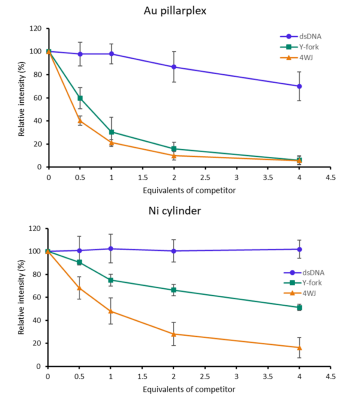


Figure 10. Gel electrophoresis competition of dsDNA (ds-21; blue circles), Y-fork (S2 and S3 of the 4WJ sequence; green squares), and 4WJ (orange triangles) against FAM-labeled 3WJ bound to pillarplex (top) and cylinder (bottom). The 3WJ is unstable in the absence of pillarplex or cylinder, so the relative intensity reflects the extent to which the competitors have scavenged the stabilizing drug. Further details of the experiment are in Figure S20.

cancer cell lines are reported in SI Table ST1. At 24 h, the Ag pillarplex shows some cytotoxic activity (IC₅₀ 6–10 μ M) with a similar activity observed at 72 h. By contrast, the Au pillarplex is not cytotoxic at 24 h (despite having already entered the nucleus) but is moderately cytotoxic at 72 h (IC₅₀ 11–12 μ M) at levels comparable to those of cisplatin.

CONCLUSIONS

We have shown herein that the pillarplex binds to open cruciform-style 4WJs which is exciting, both because these are believed to be the biologically active form of the Holliday junction⁹ and because this now provides a roadmap for targeting higher-order junction structures using this metallo-supramolecular approach.

The increased girth of the pillarplex compared to the previously studied cylinders brings new properties to its binding. Although the pillarplex does bind 3WJ, in doing so, it disrupts the hydrogen bonding between bases and opens up that junction structure, and at higher loading, it breaks apart the 3WJs into Y-shaped forks. Indeed, the contrast with the cylinder is striking, with the cylinder rearranging the 4WJ into its preferred 3WJ targets (or Y-forks), whereas the pillarplex does not transform 4WJs into 3WJ structures.

That the Au pillarplex is an effective junction and fork binder indicates the value of developing cationic agents which display polyaryl surfaces as a junction-binding design strategy. Featuring an organometallic scaffold and a pore (that can be used for rotaxane formation),⁵⁷ it further expands the chemical toolbox available to construct nucleic acid junction binders.

The pillarplex is a little small to completely fill an open 4WJ cavity, and there is scope for further elaboration to create slightly larger agents to target and reside in this cavity. This minor size mismatch helps to explain the competition between 4WJ-binding and Y-shaped fork binding observed at high pillarplex loading.

The 24 h timepoint where the Au pillarplex has entered the nucleus but not yet caused cytotoxicity offers the opportunity for future detailed study of the in cellulo effects of Holliday junction binding. Alongside their biological roles, DNA junctions—and especially 4WJs—are also widely used in nucleic acid nanostructure construction. The ability of the pillarplex to not only form and stabilize DNA junctions but also modulate and switch them is particularly intriguing in this context. We are actively exploring these aspects.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.3c00118.

- Experimental details, Supporting Table ST1, and Supporting Figures S1–S20 (PDF)
- Movies of representative MD simulations illustrated in Figures 4, 7, 9, and S11 (MPG)
- Additional Supporting Information (relating to MD simulation data) has been placed at Zenodo: https://doi: 10.5281/zenodo.7799316 (MPG)
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Notes

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REFERENCES

(1) Thorn, C. F.; Oshiro, C.; Marsh, S.; Hernandez-Boussard, T.; McLeod, H.; Klein, T. E.; Altman, R. B. Doxorubicin Pathways: Pharmacodynamics and Adverse Effects. *Pharmacogenet. Genomics* **2011**, *21*, 440–446.

(2) Dasari, S.; Tchounwou, P. B. Cisplatin in Cancer Therapy: Molecular Mechanisms of Action. *Eur. J. Pharmacol.* **2014**, 740, 364– 378.

(3) Riddell, I. A.; Lippard, S. J. Cisplatin and Oxaliplatin: Our Current Understanding of Their Actions. In *Metals in Life Sciences;* Sigel, R.; Sigel, H.; Sigel, A., Eds.; De Gruyter: Berlin, 2018; Vol. 18, 1–42 DOI: 10.1515/9783110470734-001/MACHINEREADABLE-CITATION/RIS.

(4) Martins-Teixeira, M. B.; Carvalho, I. Antitumour Anthracyclines: Progress and Perspectives. *ChemMedChem* **2020**, *15*, 933–948.

(5) Boer, D. R.; Canals, A.; Coll, M. DNA-Binding Drugs Caught in Action: The Latest 3D Pictures of Drug-DNA Complexes. *Dalton Trans.* **2009**, 399–414.

(6) Hannon, M. J. Supramolecular DNA Recognition. *Chem. Soc. Rev.* **2007**, *36*, 280–295.

(7) Ivens, E.; Cominetti, M. M. D.; Searcey, M. Junctions in DNA: Underexplored Targets for Therapeutic Intervention. *Bioorg. Med. Chem.* **2022**, *69*, No. 116897.

(8) (a) McQuaid, K. Y.; Pipier, A.; Cardin, C. J.; Monchaud, D. Interactions of small molecules with DNA junctions. *Nucleic Acids Res.* **2022**, *50*, 12636–12656. (b) McQuaid, K. Y.; Pipier, A.; Cardin, C. J.; Monchaud, D. Correction to 'Interactions of small molecules with DNA junctions'. *Nucleic Acids Res.* **2022**, *50*, 13198.

(9) Liu, Y.; West, S. C. Happy Hollidays: 40th Anniversary of the Holliday Junction. *Nature Rev. Mol. Cell Biol.* **2004**, *5*, 937–944.

(10) Neidle, S. Quadruplex Nucleic Acids as Novel Therapeutic Targets. J. Med. Chem. 2016, 59, 5987–6011.

(11) Kosiol, N.; Juranek, S.; Brossart, P.; Heine, A.; Paeschke, K. Gquadruplexes: a promising target for cancer therapy. *Mol. Cancer* **2021**, *20*, 40.

(12) Mendes, E.; Aljnadi, I. M.; Bahls, B.; Victor, B. L.; Paulo, A. Major Achievements in the Design of Quadruplex-Interactive Small Molecules. *Pharmaceuticals* **2022**, *15*, 300.

(13) Kwok, C. K.; Balasubramanian, S. Targeted Detection of G-Quadruplexes in Cellular RNAs. *Angew. Chem., Int. Ed* **2015**, *54*, 6751–6754.

(14) Zhang, X.; Spiegel, J.; Martínez Cuesta, S.; Adhikari, S.; Balasubramanian, S. Chemical Profiling of DNA G-Quadruplex-Interacting Proteins in Live Cells. *Nat. Chem.* **2021**, *13*, 626–633.

(15) Spiegel, J.; Cuesta, S. M.; Adhikari, S.; Hänsel-Hertsch, R.; Tannahill, D.; Balasubramanian, S. G-Quadruplexes Are Transcription Factor Binding Hubs in Human Chromatin. *Genome Biol.* **2021**, *22*, 117.

(16) Hui, W. W. I.; Simeone, A.; Zyner, K. G.; Tannahill, D.; Balasubramanian, S. Single-Cell Mapping of DNA G-Quadruplex Structures in Human Cancer Cells. *Sci. Rep.* **2021**, *11*, 23641.

(17) Martella, M.; Pichiorri, F.; Chikhale, Rv.; Abdelhamid, M. A. S.; Waller, Z. A. E.; Smith, S. S. I-Motif Formation and Spontaneous Deletions in Human Cells. *Nucleic Acids Res.* **2022**, *50*, 3445–3455.

(18) Zeraati, M.; Langley, D. B.; Schofield, P.; Moye, A. L.; Rouet, R.; Hughes, W. E.; Bryan, T. M.; Dinger, M. E.; Christ, D. I-Motif DNA Structures Are Formed in the Nuclei of Human Cells. *Nature Chem* **2018**, *10*, 631–637.

(19) Tang, W.; Niu, K.; Yu, G.; Jin, Y.; Zhang, X.; Peng, Y.; Chen, S.; Deng, H.; Li, S.; Wang, J.; Song, Q.; Feng, Q. In Vivo Visualization of the I-Motif DNA Secondary Structure in the *Bombyx mori* Testis. *Epigenet. Chromatin* **2020**, *13*, 12.

(20) Ravichandran, S.; Subramani, V. K.; Kim, K. K. Z-DNA in the Genome: From Structure to Disease. *Biophys. Rev.* 2019, *11*, 383–387.

(21) Zell, J.; Sperti, F. R.; Britton, S.; Monchaud, D. DNA Folds Threaten Genetic Stability and Can Be Leveraged for Chemotherapy. *RSC Chem. Biol.* **2021**, *2*, 47–76.

(22) Duskova, K.; Lejault, P.; Benchimol, É.; Guillot, R.; Britton, S.; Granzhan, A.; Monchaud, D. DNA Junction Ligands Trigger DNA Damage and Are Synthetic Lethal with DNA Repair Inhibitors in Cancer Cells. J. Am. Chem. Soc. **2020**, 142, 424–435.

(23) Garci, A.; Castor, K. J.; Fakhoury, J.; Do, J. L.; di Trani, J.; Chidchob, P.; Stein, R. S.; Mittermaier, A. K.; Friščić, T.; Sleiman, H. Efficient and Rapid Mechanochemical Assembly of Platinum(II) Squares for Guanine Quadruplex Targeting. J. Am. Chem. Soc. 2017, 139, 16913–16922.

(24) Sperti, F. R.; Dupouy, B.; Mitteaux, J.; Pipier, A.; Pirrotta, M.; Chéron, N.; Valverde, I. E.; Monchaud, D. Click-Chemistry-Based Biomimetic Ligands Efficiently Capture G-Quadruplexes In Vitro and Help Localize Them at DNA Damage Sites in Human Cells. *JACS Au* **2022**, *2*, 1588–1595.

(25) Georgiades, S. N.; Abd Karim, N. H.; Suntharalingam, K.; Vilar, R. Interaction of Metal Complexes with G-Quadruplex DNA. *Angew. Chem., Int. Ed* **2010**, *49*, 4020–4034.

(26) Monchaud, D.; Teulade-Fichou, M. P. A Hitchhiker's Guide to G-Quadruplex Ligands. Org. Biomol. Chem. 2008, 6, 627–636.

(27) Cardo, L.; Hannon, M. J. Non-Covalent Metallo-Drugs: Using Shape to Target DNA and RNA Junctions and Other Nucleic Acid Structures. In *Metals in Life Sciences;* Sigel, R.; Sigel, H.; Sigel, A., Eds.; De Gruyter: Berlin, 2018; Vol. 18, 303–324 DOI: 10.1515/9783110470734-017.

(28) Phongtongpasuk, S.; Paulus, S.; Schnabl, J.; Sigel, R. K. O.; Spingler, B.; Hannon, M. J.; Freisinger, E. Binding of a Designed Anti-Cancer Drug to the Central Cavity of an RNA Three-Way Junction. *Angew. Chem., Int. Ed.* **2013**, *52*, 11513–11516.

(29) Boer, D.; Kerckhoffs, J. M. C. A.; Parajo, Y.; Pascu, M.; Lincoln, P.; Usón, I.; Hannon, M. J.; Coll, M. Self-Assembly of Functionalizable Two-Component 3D DNA Arrays through the Induced Formation of DNA Three-Way-Junction Branch Points by Supramolecular Cylinders. *Angew. Chem., Int. Ed.* **2010**, *49*, 2336–2339.

(30) Oleksi, A.; Blanco, A. G.; Boer, R.; Usón, I.; Aymamí, J.; Rodger, A.; Hannon, M. J.; Coll, M. Molecular Recognition of a Three-Way DNA Junction by a Metallosupramolecular Helicate. *Angew. Chem., Int. Ed*, **2006**, 45, 1227–1231.

(31) Cerasino, L.; Hannon, M. J.; Sletten, E. DNA Three-Way Junction with a Dinuclear Iron(II) Supramolecular Helicate at the Center: A NMR Structural Study. *Inorg. Chem.* 2007, *46*, 6245–6251. (32) Malina, J.; Hannon, M. J.; Brabec, V. Recognition of DNA Three-Way Junctions by Metallosupramolecular Cylinders: Gel Electrophoresis Studies. *Chem. - Eur. J.* 2007, *13*, 3871–3877.

(33) Cardo, L.; Sadovnikova, V.; Phongtongpasuk, S.; Hodges, N. J.; Hannon, M. J. Arginine Conjugates of Metallo-Supramolecular Cylinders Prescribe Helicity and Enhance DNA Junction Binding and Cellular Activity. *Chem. Commun.* **2011**, *47*, 6575–6577.

(34) Hooper, C. A. J.; Cardo, L.; Craig, J. S.; Melidis, L.; Garai, A.; Egan, R. T.; Sadovnikova, V.; Burkert, F.; Male, L.; Hodges, N. J.; Browning, D. F.; Rosas, R.; Liu, F.; Rocha, Fv.; Lima, M. A.; Liu, S.; Bardelang, D.; Hannon, M. J. Rotaxanating Metallo-Supramolecular Nano-Cylinder Helicates to Switch DNA Junction Binding. *J. Am. Chem. Soc.* **2020**, *142*, 20651–20660.

(35) Vuong, S.; Stefan, L.; Lejault, P.; Rousselin, Y.; Denat, F.; Monchaud, D. Identifying Three-Way DNA Junction-Specific Small-Molecules. *Biochimie* **2012**, *94*, 442–450.

(36) Novotna, J.; Laguerre, A.; Granzhan, A.; Pirrotta, M.; Teulade-Fichou, M. P.; Monchaud, D. Cationic Azacryptands as Selective Three-Way DNA Junction Binding Agents. *Org. Biomol. Chem.* **2015**, *13*, 215–222.

(37) Stefan, L.; Bertrand, B.; Richard, P.; le Gendre, P.; Denat, F.; Picquet, M.; Monchaud, D. Assessing the Differential Affinity of Small Molecules for Noncanonical DNA Structures. *ChemBioChem* **2012**, *13*, 1905–1912.

(38) Duskova, K.; Lamarche, J.; Amor, S.; Caron, C.; Queyriaux, N.; Gaschard, M.; Penouilh, M. J.; de Robillard, G.; Delmas, D.; Devillers, C. H.; Granzhan, A.; Teulade-Fichou, M. P.; Chavarot-Kerlidou, M.; Therrien, B.; Britton, S.; Monchaud, D. Identification of Three-Way DNA Junction Ligands through Screening of Chemical Libraries and Validation by Complementary in Vitro Assays. *J. Med. Chem.* **2019**, *62*, 4456–4466.

(39) Zell, J.; Duskova, K.; Chouh, L.; Bossaert, M.; Chéron, N.; Granzhan, A.; Britton, S.; Monchaud, D. Dual Targeting of Higher-Order DNA Structures by Azacryptands Induces DNA Junction-Mediated DNA Damage in Cancer Cells. *Nucleic Acids Res.* **2021**, *49*, 10275–10288.

(40) Zhu, J.; Bošković, F.; Nguyen, B. N. T.; Nitschke, J. R.; Keyser, U. F. Kinetics of Toehold-Mediated DNA Strand Displacement Depend on $Fe_4^{II}L_4$ Tetrahedron Concentration. *Nano Lett.* **2021**, *21*, 1368–1374.

(41) Zhu, J.; Haynes, C. J. E.; Kieffer, M.; Greenfield, J. L.; Greenhalgh, R. D.; Nitschke, J. R.; Keyser, U. F. Fe₄^{II}L₄ Tetrahedron Binds to Nonpaired DNA Bases. *J. Am. Chem. Soc.* **2019**, *141*, 11358–11362.

(42) Gómez-González, J.; Bouzada, D.; Pérez-Márquez, L. A.; Sciortino, G.; Maréchal, J. D.; Vázquez López, M.; Vázquez, M. E. Stereoselective Self-Assembly of DNA Binding Helicates Directed by the Viral β -Annulus Trimeric Peptide Motif. *Bioconjugate Chem.* **2021**, 32, 1564–1569.

(43) Gómez-González, J.; Pérez, Y.; Sciortino, G.; Roldan-Martín, L.; Martínez-Costas, J.; Maréchal, J. D.; Alfonso, I.; Vázquez López, M.; Vázquez, M. E. Dynamic Stereoselection of Peptide Helicates and Their Selective Labeling of DNA Replication Foci in Cells. *Angew. Chem., Int. Ed.* **2021**, *60*, 8859–8866.

(44) Gómez-González, J.; Martínez-Castro, L.; Tolosa-Barrilero, J.; Alcalde-Ordóñez, A.; Learte-Aymamí, S.; Mascareñas, J. L.; García-Martínez, J. C.; Martínez-Costas, J.; Maréchal, J. D.; Vázquez López, M.; Vázquez, M. E. Selective Recognition of A/T-Rich DNA 3-Way Junctions with a Three-Fold Symmetric Tripeptide. *Chem. Commun.* **2022**, *58*, 7769–7772.

(45) Gamba, I.; Rama, G.; Ortega-Carrasco, E.; Maréchal, J. D.; Martínez-Costas, J.; Eugenio Vázquez, M.; López, M. V. Programmed Stereoselective Assembly of DNA-Binding Helical Metallopeptides. *Chem. Commun.* **2014**, *50*, 11097–11100.

(46) Barros, S. A.; Chenoweth, D. M. Recognition of Nucleic Acid Junctions Using Triptycene-Based Molecules. *Angew. Chem., Int. Ed.* **2014**, *53*, 13746–13750.

(47) Yang, Z.; Chen, Y.; Li, G.; Tian, Z.; Zhao, L.; Wu, X.; Ma, Q.; Liu, M.; Yang, P. Supramolecular Recognition of Three Way Junction DNA by a Cationic Calix[3]Carbazole. *Chem. - Eur. J.* **2018**, *24*, 6087–6093.

(48) Brabec, V.; Howson, S. E.; Kaner, R. A.; Lord, R. M.; Malina, J.; Phillips, R. M.; Abdallah, Q. M. A.; McGowan, P. C.; Rodger, A.; Scott, P. Metallohelices with Activity against Cisplatin-Resistant Cancer Cells; Does the Mechanism Involve DNA Binding? *Chem. Sci.* **2013**, *4*, 4407–4416.

(49) Qu, C. Y.; Zhou, L. S.; Shu, L. H.; Huang, Q. $Fe_4^{II}L_4$ Tetrahedron-Assisted Three-Way Junction Probe for Multiple MiRNA Detection. *ACS Omega* **2021**, *6*, 3330–3335.

(50) Howell, L. A.; Waller, Z. A. E.; Bowater, R.; O'Connell, M.; Searcey, M. A Small Molecule That Induces Assembly of a Four Way DNA Junction at Low Temperature. *Chem. Commun.* **2011**, 47, 8262–8264.

(51) Brogden, A. L.; Hopcroft, N. H.; Searcey, M.; Cardin, C. J. Ligand Bridging of the DNA Holliday Junction: Molecular Recognition of a Stacked-X Four-Way Junction by a Small Molecule. *Angew. Chem., Int. Ed.* **2007**, *46*, 3850–3854.

(52) van Rixel, V. H. S.; Busemann, A.; Wissingh, M. F.; Hopkins, S. L.; Siewert, B.; van de Griend, C.; Siegler, M. A.; Marzo, T.; Papi, F.; Ferraroni, M.; Gratteri, P.; Bazzicalupi, C.; Messori, L.; Bonnet, S. Induction of a Four-Way Junction Structure in the DNA Palindromic Hexanucleotide 5'-d(CGTACG)-3' by a Mononuclear Platinum Complex. *Angew. Chem., Int. Ed.* **2019**, *58*, 9378–9382.

(53) Satange, R.; Kao, S.-H.; Chien, C.-M.; Chou, S.-H.; Lin, C.-C.; Neidle, S.; Hou, M.-H. Staggered Intercalation of DNA Duplexes with Base-Pair Modulation by Two Distinct Drug Molecules Induces Asymmetric Backbone Twisting and Structure Polymorphism. *Nucleic Acids Res.* **2022**, *50*, 8867–8881.

(54) Canals, A.; Arribas-Bosacoma, R.; Albericio, F.; Álvarez, M.; Aymamí, J.; Coll, M. Intercalative DNA Binding of the Marine Anticancer Drug Variolin B. *Sci. Rep.* **2017**, *7*, 39680.

(55) Hays, F. A.; Watson, J.; Ho, P. S. Caution! DNA Crossing: Crystal Structures of Holliday Junctions. J. Biol. Chem. 2003, 278, 49663-49666.

(56) Pöthig, A.; Casini, A. Recent Developments of Supramolecular Metal-based Structures for Applications in Cancer Therapy and Imaging. *Theranostics* **2019**, *9*, 3150–3169.

(57) (a) Altmann, P. J.; Pöthig, A. Pillarplexes: A Metal–Organic Class of Supramolecular Hosts. J. Am. Chem. Soc. 2016, 138, 13171– 13174. (b) Altmann, P. J.; Jandl, C.; Pöthig, A. Introducing a Pyrazole/Imidazole Based Hybrid Cyclophane: A Hydrogen Bond Sensor and Binucleating Ligand Precursor. Dalton Trans. 2015, 44, 11278–11281. (c) Guan, S.; Pickl, T.; Jandl, C.; Schuchmann, L.; Zhou, X.-Y.; Altmann, P.; J Pöthig, A. Triazolate-based pillarplexes: shape-adaptive metallocavitands via rim modification of macrocyclic ligands. Org. Chem. Front. 2021, 8, 4061–4070. (d) Altmann, P. J.; Pöthig, A. A pH-Dependent, Mechanically Interlocked Switch: Organometallic [2]Rotaxane vs. Organic [3]Rotaxane. Angew. Chem., Int. Ed 2017, 129, 15939–15942.

(58) Melidis, L.; Styles, I. B.; Hannon, M. J. Targeting Structural Features of Viral Genomes with a Nano-Sized Supramolecular Drug. *Chem. Sci.* **2021**, *12*, 7174–7184.

(59) Melidis, L.; Hill, H. J.; Coltman, N. J.; Davies, S. P.; Winczura, K.; Chauhan, T.; Craig, J. S.; Garai, A.; Hooper, C. A. J.; Egan, R. T.; McKeating, J. A.; Hodges, N. J.; Stamataki, Z.; Grzechnik, P.; Hannon, M. J. Supramolecular Cylinders Target Bulge Structures in the 5' UTR of the RNA Genome of SARS-CoV-2 and Inhibit Viral Replication. *Angew. Chem., Int. Ed.* **2021**, *60*, 18144–18151.

(60) Woods, K. C.; Martin, S. S.; Chu, V. C.; Baldwin, E. P. Quasi-Equivalence in Site-Specific Recombinase Structure and Function: Crystal Structure and Activity of Trimeric Cre Recombinase Bound to a Three-Way Lox DNA Junction. *J. Mol. Biol.* **2001**, *313*, 49–69.

(61) Zhong, M.; Rashes, M. S.; Kallenbach, N. R. Effect of T-T Base Mismatches on Three-Arm DNA Junctions. *Biochemistry* **1993**, *32*, 6898–6907.

(62) Zhong, M.; Marky, L. A.; Kallenbach, N. R.; Kupke, D. W. Thermodynamics of DT-dT Base Pair Mismatching in Linear DNA Duplexes and Three-Arm DNA Junctions. *Biochemistry* **1997**, *36*, 2485–2491.

(63) Leontis, N. B.; Kwok, W.; Newman, J. S. Stability and Structure of Three-Way DNA Junctions Containing Unpaired Nucleotides. *Nucleic Acids Res.* **1991**, *19*, 759.

(64) Leontis, N. B.; Hills, M. T.; Piotto, M.; Ouporov, Iv.; Malhotra, A.; Gorenstein, D. G. Helical Stacking in DNA Three-Way Junctions Containing Two Unpaired Pyrimidines: Proton NMR Studies. *Biophys. J.* **1995**, *68*, 251–265.

(65) Ghosh, K.; Chi, K. L.; Guo, F.; Segall, A. M.; van Duyne, G. D. Peptide Trapping of the Holliday Junction Intermediate in Cre-LoxP Site-Specific Recombination. J. Biol. Chem. 2005, 280, 8290–8299.

(66) Malina, J.; Hannon, M. J.; Brabec, V. Iron(II) Supramolecular Helicates Condense Plasmid DNA and Inhibit Vital DNA-Related Enzymatic Activities. *Chem. - Eur. J.* **2015**, *21*, 11189–11195.

(67) Meistermann, I.; Moreno, V.; Prieto, M. J.; Moldrheim, E.; Sletten, E.; Khalid, S.; Mark Rodger, P.; Peberdy, J. C.; Isaac, C. J.; Rodger, A.; Hannon, M. J. Intramolecular DNA Coiling Mediated by Metallo-Supramolecular Cylinders: Differential Binding of P and M Helical Enantiomers. *Proc. Natl. Acad. Sci. U.S.A.* **2002**, *99*, 5069– 5074.

(68) Hannon, M. J.; Moreno, V.; Prieto, M. J.; Moldrheim, E.; Sletten, E.; Meistermann, I.; Isaac, C. J.; Sanders, K. J.; Rodger, A. Intramolecular DNA Coiling Mediated by a Metallo-Supramolecular Cylinder. *Angew. Chem., Int. Ed.* **2001**, *40*, 879–884.

(69) Malina, J.; Hannon, M. J.; Brabec, V. Interaction of Dinuclear Ruthenium(II) Supramolecular Cylinders with DNA: SequenceSpecific Binding, Unwinding, and Photocleavage. Chem.-Eur. J. 2008, 14, 10408-10414.

(70) Malina, J.; Hannon, M. J.; Brabec, V. DNA Binding of Dinuclear Iron(II) Metallosupramolecular Cylinders. DNA Unwinding and Sequence Preference. *Nucleic Acids Res.* 2008, *36*, 3630–3638.
(71) Rodger, A.; Nordén, B. Circular Dichroism and Linear

Dichroism; Oxford University Press: Oxford, UK, 1997. (72) Pöthig, A.; Ahmed, S.; Winther-Larsen, H. C.; Guan, S.; Altmann, P. J.; Kudermann, J.; Andresen, A. M. S.; Gjøen, T.; Åstrand,

O. A. H. Antimicrobial Activity and Cytotoxicity of Ag(I) and Au(I) Pillarplexes. *Front. Chem.* **2018**, *6*, 584.

(73) Carr, C. E.; Marky, L. A. Melting Behavior of a DNA Four-Way Junction Using Spectroscopic and Calorimetric Techniques. *J. Am. Chem. Soc.* **201**7, *139*, 14443–14455.

(74) Carr, C. E.; Marky, L. A. Effect of GCAA stabilizing loops on three- and four-way intramolecular junctions. *Phys. Chem. Chem. Phys.* **2018**, *20*, 5046–5056.

(75) Wang, W.; Nocka, L. M.; Wiemann, B. Z.; Hinckley, D. M.; Mukerji, I.; Starr, F. W. Holliday Junction Thermodynamics and Structure: Coarse-Grained Simulations and Experiments. *Sci. Rep.* **2016**, *6*, 22863.

(76) Thompson, S. J.; Thompson, S. E. M.; Cazier, J.-B. CaStLeS (Compute and Storage for the Life Sciences): A Collection of Compute and Storage Resources for Supporting Research at the University of Birmingham. 2019 DOI: 10.5281/ZENODO.3250616.