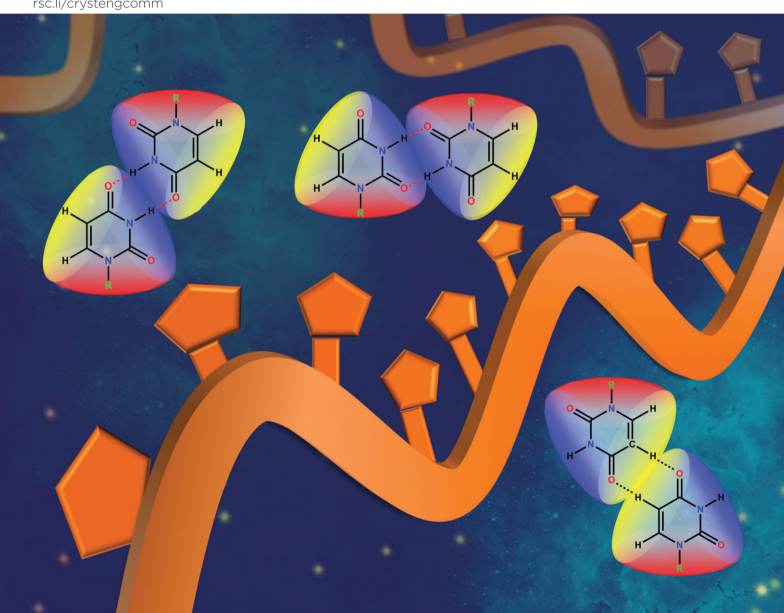
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Mimic uracil-uracil base pairing: self-assembly and single crystal structure†

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The controllable synthesis of non-classical nucleobase pairs that can mimic partial biological behavior is very important for the in-depth study of DNA or RNA. Here, three different interaction modes of uraciluracil base pairs (U-motif) have been explored in the coordination complexes of 2'-deoxyuracil 5'monophosphate (dUMP) and uracil 5'-monophosphate (UMP) with transition metal ions by single crystal structural analysis and DFT calculations for the first time. The results indicate that the structural diversity of non-classical nucleobase pairs may be a bank of supramolecular interactions, which will contribute to the study of supramolecular chemistry and biochemistry.

Introduction

Nucleobase recognition is a crucial biological phenomenon, which underpins many intricate processes within living systems, including gene transcription and translation. It is the cornerstone of life. 1-4 One fundamental question is the recognition mode between base pairs, specifically the type of hydrogen bond between bases that dominates the structure.5-7 In the classical DNA double helix structure, nucleobase recognition is limited to a few choices, namely the complementary adenine:thymine base pair (A·T) and cytosine:guanine base pair (C·G).8,9 Numerous non-classical base pairs, involving at least two hydrogen bonds between any two bases except for the Watson-Crick base pairs, have been discovered based on diverse characterizations, which are of great significance in revealing genetic diseases and DNA rearrangements. 10-13 However, among these characterizations, only single crystal X-ray diffraction (SC-XRD) can offer welldefined structural information, enabling the precise determination of the protonation position of the nucleobase, the hydrogen bond geometry, and the interaction mode

With the growing field of self-assembly chemistry, noncovalent interactions have been employed to construct complicated supramolecular systems. 16-18 The precise design and synthesis of self-assembly systems that can partially mimic biological behaviours is an effective way to address the above issue. For decades, scientists have been dedicated to exploring novel base pairing patterns through hydrogen bond directed assembly of nucleotides. The nucleobase edges have been classified into three categories: Watson-Crick, Hoogsteen, and Sugar edges, according to their different roles in hydrogen bond formation (Fig. S1†). And this has greatly increased the possible orientation and the number of interacting bases, leading to the discovery of innovative basepairing.19

Wobble base pairs in RNA sequences, such as G·U, I·A, I·U and I-C, play a pivotal role in biochemistry. 20,21 Particularly, research focused on G·U wobble base pairs has demonstrated the potential to regulate biological processes. 22-24 Compared to the widely studied guanine base pairs and cytosine base pairs, 25-27 the uracil-uracil base pair (U-motif) has received less attention. Although seven kinds of U-motifs have been reported, most of these are based on planar nucleobase derivatives (Fig. S2†). Additionally, previous research²⁸⁻³⁰ has identified general U-motifs in RNA tertiary structures. Compared to DNA or RNA macromolecules, the crystal structures of smaller native nucleotide complexes have higher resolution.³¹ However, there are few experimental and structural data on native nucleotide molecules. 10 Here, by using aromatic ligands and cobalt or cadmium transition metals, a hydrophobic space is constructed during the

between bases in the solid state. Unfortunately, the lack of a well-defined structure for non-classical base pairs hinders investigations into this issue. 14,15

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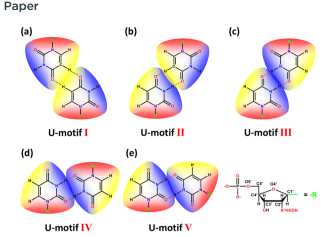


Fig. 1 The structural scheme of five types of U-motifs. R represents the sugar ring and monophosphate group of UMP.

coordination process, which induces the formation of Umotifs. 32-34 For the convenience of our study, a structural overview of the five U-motif types is presented in Fig. 1.

Experimental section

Materials and methods

All reagents and organic solvents were obtained from commercial sources and used without further purification. ¹H NMR spectra were recorded on a 400 MHz Bruker FT-NMR spectrometer using DMSO- d_6 . FT-IR spectra (KBr pellet) were performed on a Thermo IS5 FT-IR spectrometer in the range of 400-4000 cm⁻¹. Powder X-ray diffraction (PXRD) patterns were carried out on a Bruker D8 Advanced X-ray diffractometer with a scanning rate of 10° min⁻¹ (2θ angle measurement range: 5-50°). Elemental analysis (C, H, and N) was obtained by using an EA3000 elemental analyser. UV-vis absorption spectra were measured on a Persee TU-1950 spectrophotometer. Circular dichroism spectra in solution were measured in a concentration of 0.025 mmol L⁻¹. The solid circular dichroism (CD) spectra were measured with a KBr pellet (1:200) at room temperature using a JASCO J-1500 spectropolarimeter under a stable nitrogen stream purge. Thermogravimetric analysis (TGA) was carried out using a DTG-60H thermal analyzer under nitrogen atmosphere from 50 °C to 600 °C with a heating rate of 10 °C min⁻¹.

Synthesis and physical properties

Synthesis of 1,2-bis(pyridin-4-ylmethylene)hydrazine (bpda). 4-Pyridinecarboxaldehyde (3 mL, 22 mmol) and hydrazine hydrate (50 wt%, 1 mL, 11 mmol) were dissolved in ethanol (15 mL) and 2 drops of formic acid were added as a catalyst. The mixture was stirred at room temperature for 24 h. The yellow product was filtered and washed with ethanol/ether mixed solvent (15 mL). The product was recrystallized in ethanol (20 ml) to obtain highly crystalline bpda (yield: 73.4%).

Synthesis of $\{[Co(dUMP)(bipy)(H_2O)_3] \cdot 2H_2O\}_n$ (1). An aqueous solution (3 mL) of Co(Ac)₂·4H₂O (9.9 mg, 0.04 mmol) was added to an aqueous solution (3 mL) of Na₂dUMP (14.1

mg, 0.04 mmol), and the mixture was stirred for 15 minutes. Then the auxiliary ligand 4,4'-bipy (9.4 mg, 0.06 mmol) in ethanol solution (1.5 mL) was added and the ternary system was stirred for 30 min before filtration. The orange-red blockshaped single crystals suitable for single-crystal X-ray diffraction analysis were obtained after one week using the solvent evaporation technique at room temperature (yield: 70.1%, based on Co). Anal. Calcd (%): C, 36.97; H, 4.70; N, 9.31. Found (%): C, 37.33; H, 4.78; N, 9.16. FT-IR (ν_{max}): 3567w, 3345s, 3093w, 3060w, 2981w, 2875w, 2362w, 1670vs, 1605s, 1533m, 1489m, 1473m, 1434m, 1413m, 1385s, 1262m, 1221m, 1100s, 1071vs, 1006w, 981s, 809vs, 631m, 528m, 422m cm⁻¹.

Synthesis [Co(dUMP)(azpy)(H₂O)₃]·[Co(azpy)₂(H₂-O)₄]₂·(dUMP)·7H₂O (2). Complex 2 was obtained similarly to complex 1 except that azopyridine (azpy) was used. Yield: 60.7%, based on Co. Anal. Calcd (%): C, 38.25; H, 5.31; N, 15.57. Found (%): C, 38.12; H, 5.19; N, 15.33. FT-IR (ν_{max}): 3388s, 3113w, 3091w, 3059w, 3039w, 2803w, 2366w, 1698vs, 1603m, 1568w, 1472w, 1423m, 1275m, 1231w, 1195w, 1090s, 1063s, 976s, 849m, 810m, 571m, 556m, 532m cm⁻¹.

Synthesis of $\{[Co(H_2O)_6]\cdot[Co(dUMP)_2(bpda)(H_2O)_2]\cdot 7H_2$ $\mathbf{O}\cdot\mathbf{DMF}_{ln}$ (3). An aqueous solution (3 mL) of $\mathbf{Co}(\mathbf{Ac})_2\cdot\mathbf{4H}_2\mathbf{O}$ (8.7 mg, 0.035 mmol) was added to an aqueous solution (3 mL) of Na₂dUMP (12.3 mg, 0.035 mmol), and the mixture was stirred for 15 minutes. Then the auxiliary ligand bpda (7.4 mg, 0.035 mmol) in DMF solution (3 mL) was added and the ternary system was stirred for 30 min before filtration. The orange-red block-shaped single crystals suitable for single-crystal X-ray diffraction analysis were obtained after one week using the solvent evaporation technique at room temperature (yield: 65.4%, based on Co). Anal. Calcd (%): C, 32.43; H, 3.41; N, 10.37. Found (%): C, 30.68; H, 4.98; N, 9.54. FT-IR (ν_{max}): 3376s, 3167s, 3096s, 3047s, 2306m, 1697vs, 1673vs, 1609s, 1556m, 1470m, 1418m, 1310w, 1276m, 1120s, 1088s, 1063s, 984s, 934m, 874m, 810m, 763w, 689w, 562m, 522m cm⁻¹.

Synthesis of $\{[Co(H_2O)_6]\cdot[Co(UMP)_2(bpda)(H_2O)_2]\cdot H_2O\}_n$ (4). An aqueous solution (3 mL) of Co(NO₃)₂·6H₂O (14.6 mg, 0.05 mmol) was added to an aqueous solution (3 mL) of Na₂UMP (18.4 mg, 0.05 mmol), and the mixture was stirred for 15 minutes. Then the auxiliary ligand bpda (10.5 mg, 0.05 mmol) in ethanol solution (3 mL) was added and the ternary system were stirred for 30 min before filtration. The orange-red block-shaped single crystals suitable for single-crystal X-ray diffraction analysis were obtained after one week using the solvent evaporation technique at room temperature (yield: 63.8%, based on Co). Anal. Calcd (%): C, 29.81; H, 4.28; N, 9.41. Found (%): C, 27.79; H, 5.29; N, 8.64. FT-IR (ν_{max}): 3381s, 3099s, 3047s, 2366m, 1698vs, 1672vs, 1610s, 1557m, 1468m, 1271m, 1234w, 1112s, 1085s, 1062s, 984s, 816m, 689w, 562m, 520m cm⁻¹.

Synthesis of $\{[Cd_2(UMP)_2(bpda)_3(H_2O)_4] \cdot 2H_2O\}_n$ (5). Complex 5 was obtained similarly to complex 4 except that Cd(NO₃)₂·4H₂O was used and all the molar amount changed to 0.03 mmol. The yellow rhombic single crystals suitable for single-crystal X-ray diffraction analysis were obtained after three days using the

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solvent evaporation technique at room temperature (yield: 80.3%, based on Cd). Anal. Calcd (%): C, 36.76; H, 4.57; N, 12.86. Found (%): C, 33.50; H, 5.41; N, 11.57. FT-IR (ν_{max}): 3382s, 1677vs, 1631m, 1606s, 1552m, 1475m, 1420m, 1308m, 1263m, 1436m, 1073vs, 1009m, 978m, 939m, 817s, 762w, 686m, 628w, 565m, 513m cm⁻¹.

Single-crystal X-ray diffraction

Single crystals of 1-5 with suitable dimensions were selected for single-crystal X-ray diffraction analysis, respectively. Data collection was conducted on a Bruker DUO APEX II-CCD area diffractometer using graphite monochromatic Mo K α (λ = 0.71073 Å) radiation at room temperature (296.15 K). The XSCAN program is used to find different peaks and determine the unit cell under the condition of 45 KV and 30 mA. Diffraction data collection adopts the ω -2 θ scanning mode and all scanned data are subjected to empirical absorption correction. Using Olex2,35 the crystal structure was solved with the SHELXT³⁶ structure solution program using the intrinsic phasing method, and refined with the SHELXL³⁷ refinement program using least minimisation. All non-hydrogen atomic positions were obtained from different Fourier maps and anisotropically. All hydrogen atoms were located at their geometrically generated positions by different Fourier maps combined with geometric analysis and refined isotropically. In the five heavy-atom structure, (as it was not possible to see clear election-density peaks in different maps, which would correspond with an acceptable location for part of the water oxygen sites), the refinement was completed with no allowance for these H atoms in the model. Crystallographic data are given in Tables S1 and S2.†

Computational details

Gaussview 6 is used to establish the calculation model, and Gaussian 16W is used for DFT calculation. The calculation parameters of structural optimization³⁸⁻⁴⁰ are as follows: #p opt freq b3lyp/6-311g(d,p) em = gd3bj. The parameters of energy calculation are: #b3lyp/6-311g(d,p) em = gd3bj. The parameters of BSSE (basis set superposition error) corrected energy calculation are: #b3lyp/6-311g(d,p) em = gd3bj counterpoise = 2. The surface electrostatic potential was plotted using Multiwfn⁴¹ and VMD programs.⁴²

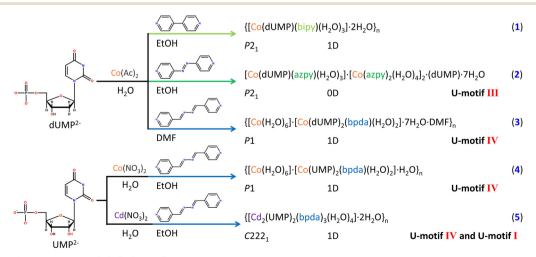
Results and discussion

Crystal structures

In this work, five types of coordination complexes of 2'deoxyuracil 5'-monophosphate (dUMP) and uracil 5'monophosphate (UMP) with transition metal ions were designed and synthesized successfully (Scheme 1 and Fig. S3†), and the crystallographic data are shown in Tables S1 and S2.† Three different types of U-motifs, U-motif I, U-motif III, and U-motif IV, with well-defined structures have been reported for the first time based on SC-XRD. Notably, there are two different U-motifs in 5 (U-motif I and U-motif IV) and they form a chain like structure. It is the first example that realizes the fully compensated base-pairing with all hydrogen donors and acceptors of the uracil base being fully utilized.

1 is a 1D coordination polymer bridged by 4,4'-bipy (Fig. S8 and S10†). The orientation of the uracil base was initially fixed by an intrachain π - π stacking interaction of 4.014(2) Å from the pyridine ring of the auxiliary ligand. And an interchain π - π stacking interaction of 3.837(2) Å was found between adjacent chains, which further stabilized the orientation of the nucleobase and extended the chains into a two-dimensional structure (Fig. S11†). In-depth structural analysis shows that there is no U-motif structure in 1, which may be attributed to the short length of 4,4'-bipy that limited the formation of a U-motif within and between the chains.

Clearly, the length of auxiliary ligand may be an important factor to the construction of U-motifs. Longer auxiliary ligands, azpy and bpda, were used to replace 4,4'-bipy for experiments (Fig. S3†). As expected, U-motifs were successfully constructed in complexes 2-5, and the single



Scheme 1 Synthetic procedures of 1, 2, 3, 4, and 5.

crystal structures of the four complexes have been obtained fortunately. 2 is a supramolecular complex containing azpy, and there are both coordinated nucleotides and guest nucleotides in the asymmetry units (Fig. S8b and S12a†). For the auxiliary ligand azpy, only one end participates in the coordination with metal ions, and it doesn't act as a bridging ligand. There are face to face π - π stacking interactions between azpy ligands, which provided the hydrophobic cavity and promoted the formation of the U-motif III (Fig. 2a and d). The U-motif III has been found in adjacent bases from coordinated dUMP and guest dUMP (Fig. S12b and c†). The orientations of uracil bases were stabilized by edge to face π – π stacking interactions The details of the H-bonding in the U-motif III are: N3-H3···O4: 1.92 Å, 2.778(8) Å, 173.7° and N3-H3···O4: 2.01 Å, 2.868(8) Å, 175.1° (atom names are converted to standard nucleotide atom names), and the dihedral angle involved in the U-motif is 2.2(3)°, which suggest that it is a perfect U-motif structure with high coplanarity.

The length of the auxiliary ligand bpda is longer than azpy and is applied in complexes 3-5 (Fig. S3†). 3 and 4 are 1D coordination polymers bridged by bpda. In 3, strong face to face π - π stacking interactions fixed the orientation of nucleobases within the chain (Fig. 2b). The U-motif IV has been found between adjacent chains (Fig. 2e and S14†). The structural details of the U-motif IV are: N3-H3...O2: 1.98 Å, 2.829(10) Å, 170.8° and N3-H3···O2: 1.97 Å, 2.825(9) Å, 172.5° (atom names are converted to standard nucleotide atom names), and the dihedral angle involved in the U-motif IV is 4.3(4)°, which is an approximately coplanar structure. This indicates that bpda is suitable for the construction of the U-motif. To further verify the role of bpda in inducing the formation of the U-motif, dUMP was replaced by UMP in 4

(Fig. S15 and S16†). The crystal structural analysis has shown that there are similar U-motif IV existing in 4. The structural details of the U-motif IV in 4 are similar with those in 3 (Fig. S17†). Clearly, bpda is suitable for the construction of the U-motif regardless of dUMP and UMP.

In 1-4, the central metal ions are all Co(II) ions in order to verify the influence of the length of auxiliary ligands. The above studies reveal that the length of auxiliary ligands can significantly influence the formation of the U-motif, which can provide a hydrophobic microenvironment for nucleobases, fix the orientation of nucleobases and induce the formation of non-classical base pairs. So, it is a feasible strategy to regulate the U-motif by changing the length of auxiliary ligands.

To further explore the interaction mode between uracil bases, our investigation focused on the central metal ions. The Cd2+ ion which have stronger bonging ability with nucleotide was used in construction of 5.43 Different from 1-4, the asymmetry unit of 5 was a binuclear Cd(II) species bridged by the phosphate group (µ2-O2) of UMP and extended into a 1D chain by the link of the bpda ligand (Fig. S9b and S18†). Notably, there is an edge-to-face π - π stacking between the nucleobases and the auxiliary ligands, which initially fixes the orientation of the nucleotide bases and induces the formation of the U-motif I within the 1D chain, with the details of C5-H5···O4: 2.55 Å, 3.405(6) Å, and 161.2° (Fig. 2c (pink dashed line), f (pink dashed line), and S18†). The U-motif IV was found between adjacent chains 1.94 Å, Å, (N3-H3···O2: 2.785(5)168.1°) (Fig. 2c (blue dashed line), f (dark blue dashed line), and S19†). It is worth noting that two different U-motifs, U-motif I and U-motif IV, appear alternately to form a 1D chain like structure (Fig. S20†), and the dihedral angle between the

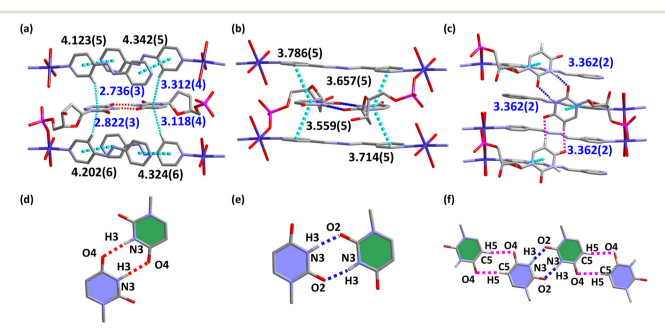


Fig. 2 The hydrophobic chemical environment formed by the auxiliary ligand in (a) 2, (b) 3, and (c) 5, in which nucleobases were fixed by π - π stacking interaction. The formed U-motifs III and IV in (d) 2 and (e) 3 respectively. The U-motifs I (pink dashed line) and IV (dark blue dashed line) formed in (f) 5.

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adjacent uracil bases is 32.8(2)°, whose coplanarity is violently distorted. The edge to face π - π stacking interaction mentioned above stabilizes this structure. It is the first reported nucleobase pairing pattern based on uracil bases, which makes full use of the intrinsic hydrogen bond donors and acceptors of the uracil nucleobase, and this unique structure was names as the full uracil-uracil base pair structure. It provides a basis for discussing the deep interaction between nucleotide bases.

DFT calculations

The electrostatic surface potential (ESP) of the three auxiliary ligands is shown in Fig. S21.†44 The minimum surface electrostatic potentials of 4,4'-bipy, azpy and bpda are concentrated on pyridine nitrogen atoms, and the energy of these are -35.23 kcal mol⁻¹, -33.94 kcal mol⁻¹ and -34.90 kcal mol⁻¹, respectively. So, azpy has a weaker binding ability with metal ions than 4,4'-bipy and bpda. The binding ability of the terminal pyridine nitrogen atom to the metal ion and the size of the auxiliary ligand can regulate the structure of the complexes, which is verified by the coordination mode of azpy in 2. Additionally, the introduction of nitrogen atoms in the middle of azpy and bpda leads to the appearance of new ESP extreme values, which are related to the formation of edge to face π - π stacking interaction between the auxiliary ligand and uracil bases in 2 and 5 and promoted the formation of U-motifs.

DFT calculation based on 1-methyluracil reveals the values of interaction energy of the five different U-motifs and shows their thermodynamic stability without considering the effect of sugar ring substitution (Fig. 3). Combined with structure analyses, we found that the U-motif containing the nonclassical hydrogen bond (C-H···O) have higher interaction

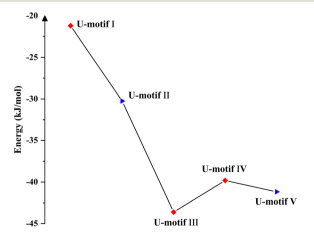


Fig. 3 Interaction energy of optimized U-motif structures based on BSSE correction: U-motif I: -21.19565 kJ mol⁻¹; U-motif II: -30.24775 kJ mol⁻¹; U-motif III: -43.61591 kJ mol⁻¹; U-motif IV: -39.81219 kJ mol⁻¹; U-motif V: -41.18331 kJ mol⁻¹ (red square points represent a symmetric structure; blue triangle points represent an asymmetric structure).

energy, while the U-motif containing the classical hydrogen bond (N-H···O) have lower one. Clearly, it suggests that classic hydrogen bond is more helpful in stabilizing the formation of base pairs.

Additionally, H-bonding parameters of base pairs obtained by theoretical calculation were very similar to those in crystal structures (Fig. S22 and Table S18†), which indicates that the crystal lattice is suitable for the stable binding between uracil bases. To further visualize the interaction between the uracil bases, DFT calculation was done based on the single crystal structure in the solid state (Fig. S23 and S24†). Strong electrostatic surface potential penetration was observed between the positive electrostatic potential (red colour) and the negative electrostatic potential (green colour) in the two uracil bases of the U-motif, which proves the existence of non-covalent base interaction.

CD spectroscopy

Complexes 1-5 crystallized in the chiral space group, and flack factor indicates that they are all homochiral single crystals (Tables S1 and S2†). The molecular and supramolecular chirality of 1-5 were explored via circular dichroism (CD) spectra in solution and the crystallized solidstate. Two characteristic peaks of the Na₂dUMP/Na₂UMP ligand were observed in solution: (1) two negative shoulder peaks, 221 nm(-) and 238 nm(-), centered at 229 nm(-), which correspond to the $n-\pi^*$ in nucleotide; (2) positive peaks at 269 nm(+) (Fig. S25†), which correspond to the n- π * and π - π * transition in nucleotide (Fig. S26a†). ⁴⁵ The solution CD spectra for 1-5 revealed that the molecular chirality of 1-5 depended on the intrinsic chirality of nucleotide molecules.

In the crystallized solid-state CD spectra, two characteristic peaks of Na2dUMP/Na2UMP were observed: (1) negative shoulder peaks centered at 241/245 nm(-), which correspond to the n- π^* and π - π^* in nucleotide molecule; (2) positive peaks at 286/289 nm(+) (Fig. 4), which may be caused by intermolecular electron transitions induced by base accumulation (Fig.

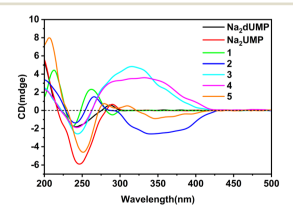


Fig. 4 The CD spectra of Na₂dUMP, Na₂UMP, and 1-5 in solid state (KCl:[sample] = 200:1).

S26b†).46 The peaks in the range of 200-300 nm of complexes 1-5 are attributed to σ - π *, n- π *, and π - π * transitions in the nucleotide ligands, respectively. The signal flip at 289 nm(-) in 1 is caused by the changes in symmetry of electronic transitions in nucleotide. Two CD peaks at 243 nm(-) and 290 nm(+) in 3 were a characteristic chiral signal caused by exciton coupling.⁴⁷ The peaks in the range of 300-400 nm appeared in complexes 2-4 are typical EAC (extended axial chirality) induced by the coordination of the auxiliary ligand and nucleotide with metal ions. 48 Meanwhile in 5, the peaks at 308 nm(+) and 345 nm(-) suggests that a new chiral source has been generated by the induced coordination of metal ions. However, due to the disorder of the auxiliary ligand, there is no strong chiral signal in the range of 300-400 nm. The coordination processes with metal ions along with the participation of auxiliary ligands can facilitate the extension and transfer of chiral sources, while the interactions between bases in chiral nucleotide molecules can effectively affect the chiral signal of nucleotide complexes.

Conclusions

Paper

In summary, the construction of U-motifs and the chirality of uracil nucleotide complexes are systematically investigated. On the basis of the high-quality single crystal structure, three types of U-motifs have been successfully constructed by adjusting the auxiliary ligands and metal ions, which provides a feasible route for the construction of base pairs. In particular, an unusual full uracil-uracil base pairing consisted of U-motifs I and IV has been observed. The interaction mode of nucleobases has been confirmed by implementing DFT calculations. Finally, the chirality of these complexes during self-assembly has been examined using solid-state CD spectroscopy in combination with crystal structure analysis.

Author contributions

All authors are contributed in work and writing of this manuscript. Prof. Hui Li is the project leader and a corresponding author of this manuscript.

Conflicts of interest

There are no conflicts to declare.

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