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Hybrid DNA/RNA nanostructures with 2'-5' linkages†

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Nucleic acid nanostructures with different chemical compositions have shown utility in biological applications as they provide additional assembly parameters and enhanced stability. The naturally occurring 2'-5' linkage in RNA is thought to be a prebiotic analogue and has potential use in antisense therapeutics. Here, we report the first instance of DNA/RNA motifs containing 2'-5' linkages. We synthesized and incorporated RNA strands with 2'-5' linkages into different DNA motifs with varying number of branch points (a duplex, four arm junction, double crossover motif and tensegrity triangle motif). Using experimental characterization and molecular dynamics simulations, we show that hybrid DNA/RNA nanostructures can accommodate interspersed 2'-5' linkages with relatively minor effect on the formation of these structures. Further, the modified nanostructures showed improved resistance to ribonuclease cleavage, indicating their potential use in the construction of robust drug delivery vehicles with prolonged stability in physiological conditions.

In the Marvel superhero universe, Tony Stark builds the Iron Man suit and becomes a superhero. The power of the suit comes not from a single function but from its many tools and functionalities, which Stark continuously upgrades to increase his power. We can draw analogy to this in DNA nanotechnology, where a growing list of molecular tools and biomedical functionalities continue to add power to this emerging field. So far, scientists have created a wide variety of nanostructures using DNA,¹⁻³ with applications in biosensing,⁴ molecular computation,⁵ macromolecular scaffolding,⁶ biomolecular analysis,⁷ and in drug delivery.⁸ While many different assembly strategies have been optimized over the years to construct nucleic acid nanostructures, there is still ongoing work to

functionalize these structures to make them better suited for real life applications. For example, modification of strand terminus with chemical groups (for example, biotin, thiol or click chemistry groups) allow the attachment of nanoparticles,⁹ or proteins.¹⁰ These functionalities are useful in biosensing (for optical readout in the case of nanoparticles)⁹ or in the creation of enzyme cascades¹¹ and synthetic vaccines (for antigens and proteins).¹² In drug delivery, incorporation of chemical groups to attach peptides provide cell targeting¹³ or drug encapsulation¹⁴ while photocleavable linkers are useful in triggered release.¹⁵ Modified DNA or RNA nucleotides or backbones have also been reported,¹⁶ including the use of L-DNA,¹⁷ locked nucleic acids (LNA),¹⁸ peptide nucleic acids (PNA)¹⁹ and incorporation of unnatural base pairs such as 5-Me-isoC, isoG and 2-thioT.²⁰ A prevalent use of such modifications is to enhance the stability of nucleic acid nanostructures²¹ and provide additional nuclease resistance²⁰ so that the structures do not degrade easily in physiological conditions. This increasing library of available modifications is a useful resource for researchers in the field to create and functionalize nucleic acid nanostructures for specific applications.²² Adding to this list, we report here for the first time, the incorporation of 2'-5' linkages in nucleic acid nanostructures. We characterized the tolerance of backbone heterogeneity in the assembled structures and discuss potential use of 2'-5' linkages in enhancing the stability of such structures against ribonucleases.

Although most RNAs have 3'-5' linked phosphate backbone, the 2'-5' phosphodiester bond also exists in nature and plays critical roles in many biological systems such as mRNA processing,²³ neurodegenerative diseases²⁴ and in viral replication²⁵ (Fig. 1). Further, 2'-5' linked oligonucleotides have potential use in antisense therapeutics due to their resistance to enzymatic hydrolysis,²⁶ diminished immune-stimulatory response²⁷ and reduced nonspecific binding to plasma and cellular proteins.²⁸ Other than the therapeutic applications, 2'-5' linked RNAs have also emerged as a plausible prebiotic RNA analogue and been regarded as a well-conserved chemical relic from evolution.²⁹ In addition, RNA aptamers and ribozymes

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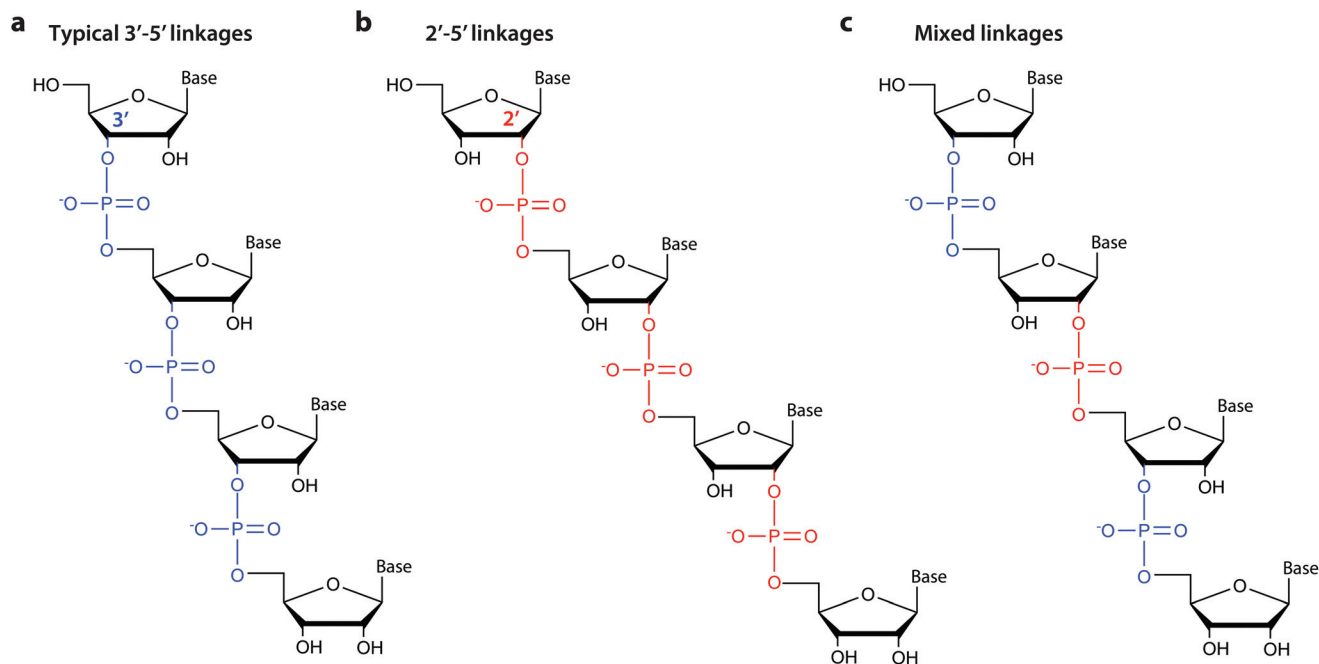


Fig. 1 3'-5'-Linked vs. 2'-5'-linked RNA. Structures of single stranded (a) 3'-5'-linked "regular" RNA, (b) 2'-5'-linked RNA and (c) RNA with mixed linkages (3'-5' linkages shown in blue and 2'-5' linkages shown in red).

with mixtures of 3'-5' and 2'-5' linkages retain their ligand binding ability and catalytic activity,²⁹ indicating that these systems are very flexible to accommodate certain numbers of 2'-5' linkages.

Tools in structural biology have provided insights into the conformation of 2'-5'-linked RNA within the context of RNA duplexes. In previous work, we solved the first two crystal structures for a self-complementary RNA 10-mer duplex containing two and six 2'-5' linkages respectively.³⁰ These data, along with molecular simulation work, provided structural insights into the conformation of 2'-5'-linked nucleotides and how RNA duplexes adjust their global and local structures to accommodate these mixed backbones through flexible sugar puckers and the flanking nucleotides compensation effect.³⁰ Following this work, we also reported two additional high-resolution structures of the same RNA duplex containing four and eight 2'-5' linkages at different positions, providing a dynamic view of RNA structural changes with increased numbers of 2'-5' linkages.³¹ Despite this research progress, incorporation of 2'-5'-linked ribonucleotides in a DNA/RNA hybrid duplex has not been studied so far, and their structural and functional compatibility in the context of nucleic acid nanostructures have not been explored. In this study, we investigated the effect of 2'-5' linkages on hybrid DNA/RNA assemblies using four different structures of increasing complexity: a simple duplex, a four-arm junction (one branch point), a double crossover (DX) motif (two branch points) and a tensegrity triangle motif (three branch points) (Fig. 2a). We analyzed the assembly of these structures through non-denaturing polyacrylamide gel electrophoresis (PAGE), characterized the

assembled motifs using UV melting studies and circular dichroism (CD), and revealed their dynamic conformational changes through molecular dynamics simulations.

We first synthesized RNA strands with different numbers of 2'-5' linkages through solid phase synthesis using commercially available phosphoramidites and purified them using denaturing PAGE (see ESI and Table S1†). We first tested DNA/RNA hybrid duplexes where the RNA strand contains zero to four 2'-5' linkages (Fig. 2b-i, RNA strand shown in red). We validated their formation on native PAGE (Fig. 2b-ii, full gels in Fig. S1†) and characterized conformational changes in the hybrid duplexes using CD (Fig. 2b-iii). The CD spectra showed clear transition from B-form duplex for the DNA/DNA to A-form for the DNA/RNA (evident by a dominant positive band at 260 nm and a negative band at 210 nm) and showed similar spectra for the hybrid duplex containing RNA with one to four 2'-5' linkages. We then analyzed the thermal stability of the hybrid duplexes using UV melting studies (Fig. S2†). Thermal melting profiles showed that the melting temperatures of the hybrid duplexes slightly decreased as the number of 2'-5' linkages in the component RNA strand increased. These results collectively indicate that formation of the hybrid DNA/RNA duplex did not seem to be affected by the heterogeneity of 2'-5' and 3'-5' linkages in the RNA strand.

Moving to motifs related to nanotechnology, we tested DNA structures containing strand crossovers, starting with a 4-arm junction and a double crossover (DX) motif. We characterized formation of the 4-arm junction containing 2'-5'-linked RNA using non-denaturing PAGE and CD, showing that the structure tolerated incorporation of 2'-5' linkages (Fig. 2c and

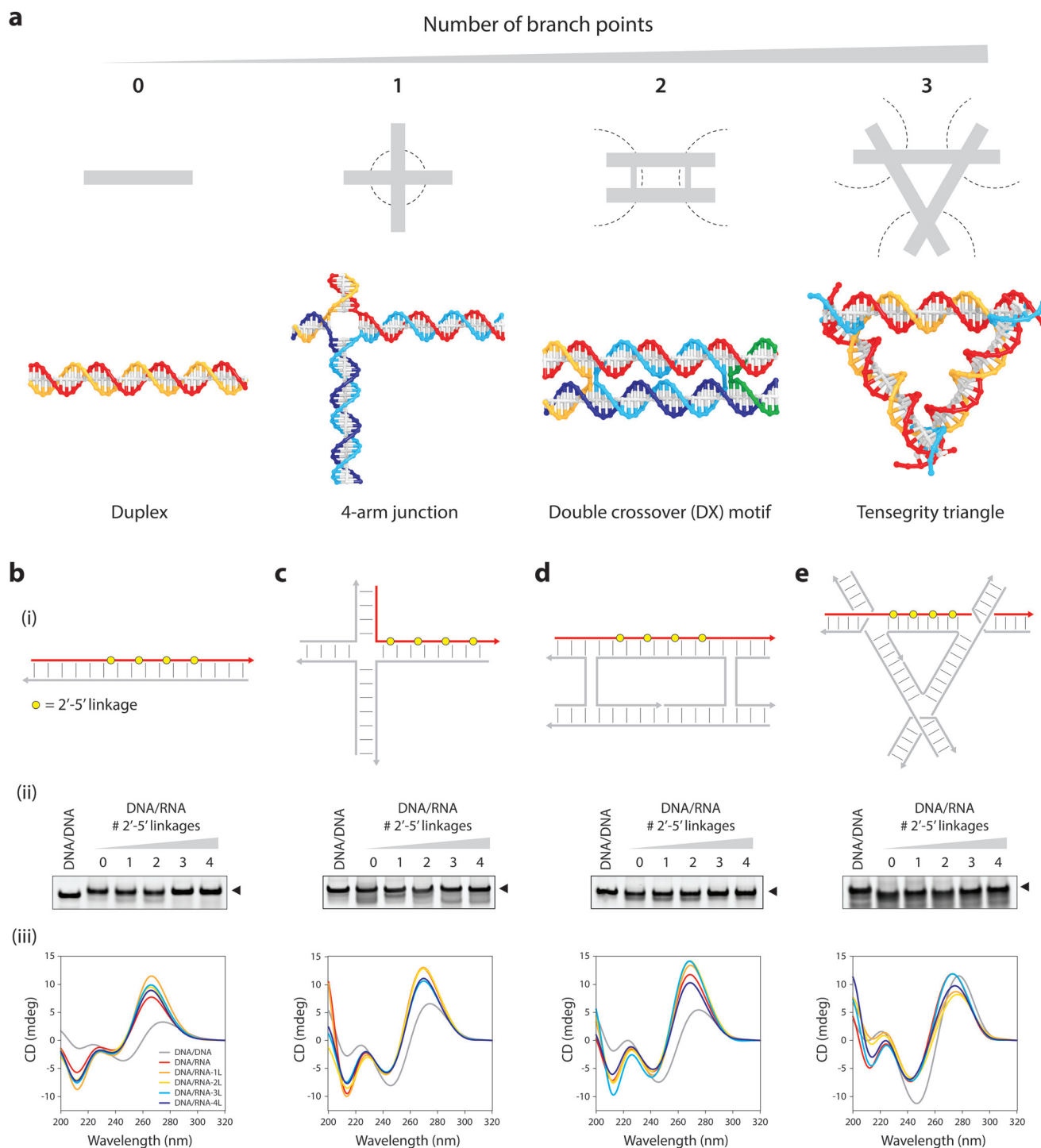


Fig. 2 Analysis of 2'-5' linkages in DNA/RNA hybrid structures. (a) Nucleic acid structures used in this study. A duplex (no branch points), a 4-arm junction (single branch point), a double crossover (DX) motif (two branch points) and a tensegrity triangle motif (each vertex of the triangle is a junction, thus three branch points). Each column shows results from (b) duplex, (c) 4-arm junction, (d) DX motif and (e) tensegrity triangle. In each case, the figure shows (i) scheme of the structures with the RNA strand in red and the four 2'-5' linkage positions indicated as yellow circles, (ii) results from native PAGE, and (iii) CD spectra of the assembled structures. 1 L to 4 L = one to four 2'-5' linkages.

Fig. S1, S3†). Our third system was a DX motif that contains 5 strands folded into two adjacent double helical domains connected by two crossovers (Fig. 2d-i and Fig. S4†).³² DNA/RNA hybrid DX molecules³³ and all-RNA DX molecules³⁴ have pre-

viously been reported, and DX-based two dimensional arrays have been used to study the helicity of duplex domains with modified nucleotides.^{35,36} We modified one of the non-cross-over strands to be RNA with 2'-5' linkages (red strand in

Fig. 2d-i) and tested proper assembly of the motif using native PAGE (Fig. 2d-ii, full gel in Fig. S1†). The formation of many such DNA motifs is based on strict stoichiometry, and in some cases the purity of the component strands. Similar motifs have been characterized using non-denaturing PAGE and other methods in many other studies^{15,37–41} and the presence of higher and lower molecular weight bands in some instances are known to indicate the formation of dimers and improperly folded products, respectively. The presence of the target band (corresponding to the structure) in high yield is indicative of proper assembly of these structures. We then characterized the hybrid assemblies using CD (Fig. 2d-iii) and tested the thermal stability of the motif with increasing numbers of 2'-5' linkages (Fig. S2†). The melting temperature of the DX motif containing an RNA strand (~51 °C) was lower than an all-DNA complex (~57 °C), possibly due to the change in helical parameters around the two crossover points. The melting temperature of the motif decreased slightly with the incorporation of 2'-5' linkages but did not exhibit much variation for the complexes containing one to four 2'-5' linkages (ranging from ~50 °C to ~54 °C).

Our final model structure was the tensegrity triangle motif, previously used in assembly of 2D lattices⁴² and designed 3D crystals.⁴³ The motif consists of three duplex edges connected pairwise by strand crossovers (Fig. 2e-i). The edges of the motif are in an over-and-under fashion, allowing it to be used in 3D construction (model shown in Fig. 2a). In construction of 3D crystals, symmetric versions of these triangle motifs yielded higher resolution;^{43–45} thus, we first tested incorporation of 2'-5'-linked RNA into a symmetric 3-turn tensegrity triangle motif (with each edge containing 31 base pairs). The symmetric version is constructed using 3 unique strands in the ratio 3:3:1 where the three duplex edges all have the same sequence identity (Fig. S5†). We first confirmed formation of the symmetric tensegrity triangle on a non-denaturing PAGE (Fig. S6†). We then modified the non-crossover strand to be RNA with 2'-5' linkages. Since the motif is symmetric, modifications will occur on all three edges when this strand is modified. Non-denaturing PAGE showed that the motifs do not form with an RNA strand containing multiple 2'-5' linkages (Fig. S7†). To test the influence of symmetry, we then tested an asymmetric design of the tensegrity triangle motif. In this version, the motif is constructed using 7 unique strands where all the edges have different sequences (Fig. S5†). We modified the sequences of one of the edges of a triangle reported previously⁴⁶ so that the sequence of the modified RNA strand is the same in both the symmetric and asymmetric designs. We confirmed successful assembly of this altered design on a non-denaturing PAGE (Fig. S6†). We then incorporated an RNA strand on one of the edges (shown in red in Fig. 2e-i, sequences in Fig. S5†). We checked assembly of the motif with the RNA strand as well as RNA containing one to four 2'-5' linkages, and non-denaturing PAGE showed proper assembly of all the motifs (Fig. 2e-ii, full gels in Fig. S1†). The asymmetric motif is modified only on one edge (*i.e.* one out of seven strands is modified) and thus tolerates the 2'-5' linkages better

than the symmetric motif (three out of seven strands modified). CD spectra showed that the positive band at 260 nm is not as dominant as in other structures containing 2'-5' linkages, indicating a predominantly B-form structure, an expected result since only one out of seven strands is changed to RNA (Fig. 2e-iii).

The effects of incorporating 2'-5' linkages in a DNA/RNA hybrid structure has not been studied before. To understand the effect of these modifications on structural and dynamics of DNA/RNA hybrid duplexes, we simulated the DNA/DNA and hybrid DNA/RNA duplexes containing zero to four 2'-5' linkages (Fig. 3). We performed simulations of length >36 ns, each consisting of 420061 atoms on average (418127, 424681, 418017, 418041, 419981, 421521 for DNA/DNA and DNA/RNA with zero to four 2'-5' linkages, respectively) including explicit solvent and K⁺/Cl[−] ions. All the DNA/RNA duplexes were observed to be stable on the simulated timescales and maintained their overall helical structures with well-defined base pairing and base stacking. Root-mean-square deviations (RMSD) of the trajectories with respect to their initial starting structures show that the initial DNA/RNA duplex structures, even containing 2'-5' linkages, remain rigidly folded throughout the simulations (Fig. S8†). RMSD values plateau after 10 ns indicating a rapid equilibration of local bending fluctuations at the modification site. Each trajectory was then structurally clustered excluding the initial 10 ns equilibration period; the centroid of each simulation's highest populated cluster is shown in Fig. 3a. In all cases, a single, highly populated cluster containing >70% of all frames was observed (Fig. 3b and Fig. S9†).

To further understand how these modifications affect the structural dynamics of duplexes, we analyzed the helical bending angle distribution within the main cluster of each duplex over the course of the trajectory (Fig. 3c). This analysis showed a steady increase in bending dynamics when one strand of DNA is replaced with RNA and further increases with the addition of 2'-5' modifications (increase in standard deviation of bending angle values in Fig. 3c). Superposing all six centroid structures illustrates the gradually increasing amounts of helical bending as a function of the 2'-5' linkage numbers (Fig. 3a). We then used the 3DNA structural analysis package⁴⁷ to measure how 2'-5' linkages affect helix structural parameters of these DNA/RNA duplexes. This analysis revealed a transition from predominantly B-form helices in the DNA/DNA duplex to partially A-form in the DNA/RNA duplex irrespective of the presence of 2'-5' linkages (Fig. S10†), in agreement with the experimentally observed CD spectra (Fig. 2b-iii). This data is also in agreement with our experiments where the addition of more 2'-5' linkages in the structure cause a monotonic decrease in melting temperatures (Fig. S2†). Increased availability of structural parameters and crystal studies of DNA motifs might provide additional functions to simulate more complex DNA nanostructures than just the duplexes studied here.

Since RNA strands with 2'-5' linkages are known to resist digestion by ribonucleases (RNases),^{26,28,48} we tested whether

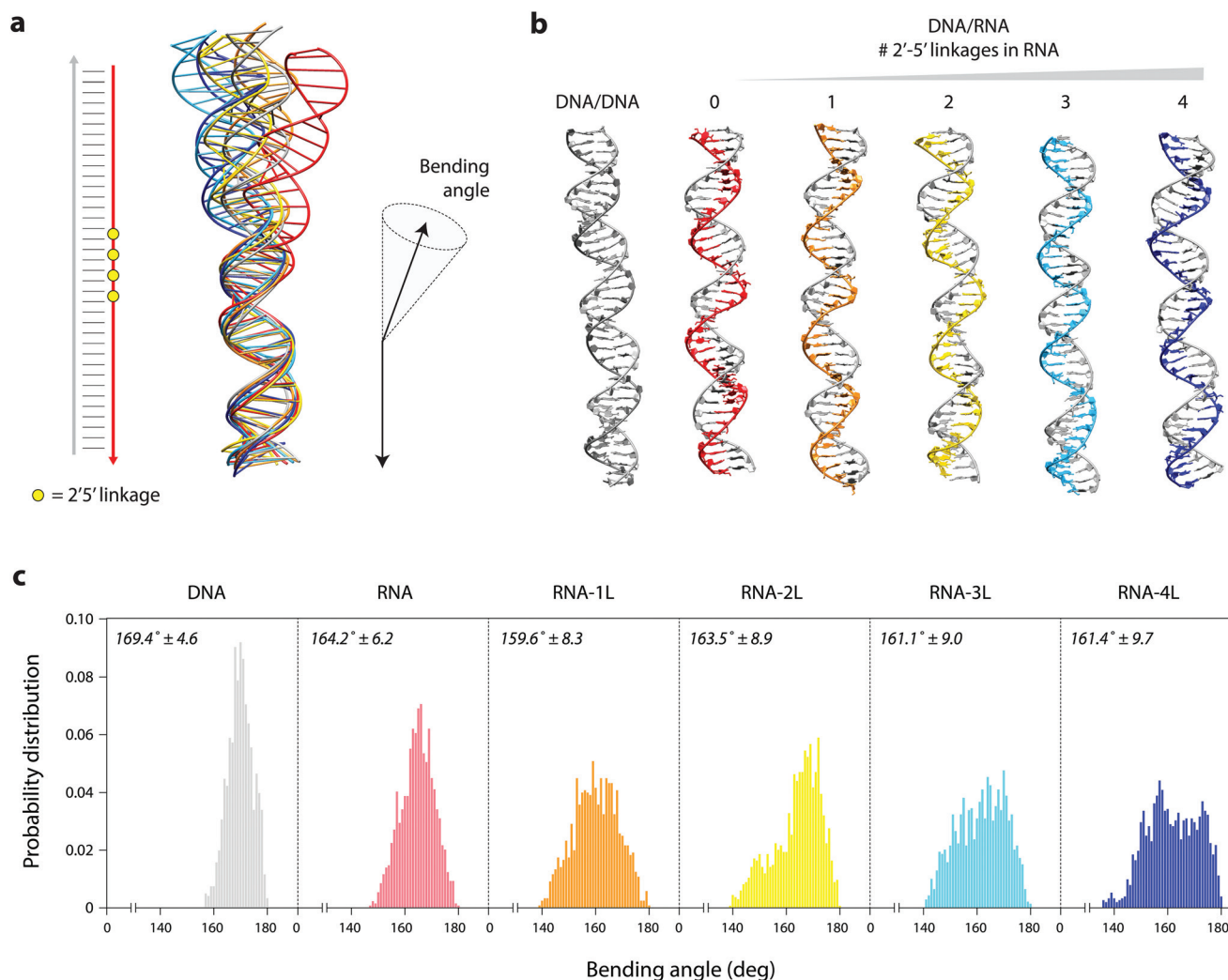


Fig. 3 Molecular dynamics simulations of 2'-5'-linked DNA/RNA hybrid duplexes. (a) Superposition of centroids of all the structures analyzed. (b) Simulated structures of each variation, with the DNA strand shown in grey (constant in all the structures). RNA strands with none to four 2'-5' linkages are shown in different colors for clarity. (c) Interhelical bending angle distribution of the simulated structures, color coded as in (b). 1 L to 4 L = one to four 2'-5' linkages.

these modifications confer RNase protection to DNA/RNA hybrid duplex structures (Fig. 4a). We incubated the native and 2'-5'-linked hybrid duplexes with different amounts of RNase H and analyzed the samples using non-denaturing PAGE.⁴⁹ When incubated with 0.5 units RNase H for 16 minutes, the modified structure showed lower levels of degradation compared to the native structure (Fig. 4b and Fig. S11†). At a lower enzyme concentration (0.1 unit), the duplex with RNA containing four 2'-5'-linkages was almost fully intact while the duplex with unmodified RNA showed higher levels of degradation. We also monitored the digestion of native and 2'-5'-linked structures over a time course, showing that the modified structure is more resistant to RNase digestion than its native counterpart (Fig. 4c).

This study is the first report on incorporating RNA strands with 2'-5' linkages into nucleic acid nanostructures. We have shown that 2'-5' linkages have minimal influence on the

assembly of hybrid duplexes, a 4-arm branched junction and a DX motif. In a larger tensegrity triangle motif, incorporation of one modified strand does not influence the structure. However, incorporation of 2'-5' linkages in a symmetric design where all three edges are modified affects self-assembly of the motif. Future studies on incorporation of such linkages in crossover strands (compared to the non-crossover strands presented here) will also provide useful information on structural effects. Further, the number of 2'-5' linkages incorporated into the structures could also play a role in the assembly process, depending on the sequence contexts.^{27,31,50} Previous studies have shown that RNA/RNA or DNA/DNA duplexes are more stable than hybrid RNA/2'-5' linked RNA or DNA/2'-5' linked RNA.⁴⁸ Further, a fully 2'-5'-linked RNA strand preferably binds with RNA than DNA strands.⁵¹ Thus, we think the use of a fully 2'-5'-linked strand in nucleic acid nanostructures might interfere with the assembly of the motifs to certain extent, but

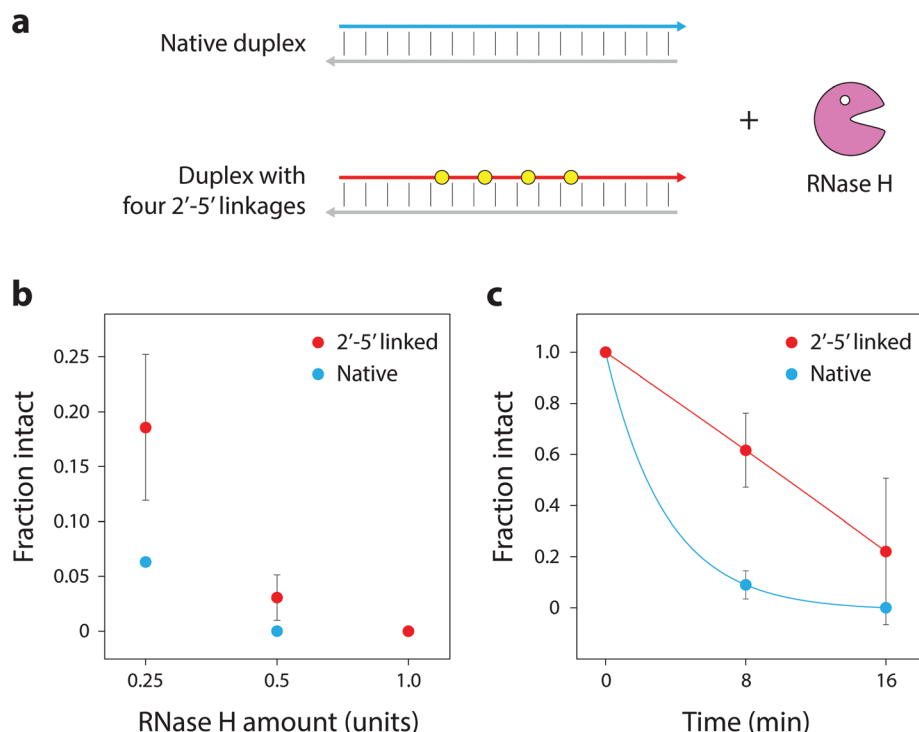


Fig. 4 Testing ribonuclease activity on 2'-5'-linked structures. (a) Native and modified duplexes with four 2'-5' linkages were tested for resistance against RNase H. (b) Percent structure intact after 16 min incubation with different amounts of RNase H. (c) Digestion trend of native and modified duplexes with 0.5 units RNase H.

can be overcome by design optimization. Here, we have shown that hybrid DNA/RNA duplexes tolerate the incorporation of quite a few 2'-5' linked nucleotides, which would be sufficient for ideal functions of nanostructures such as enhanced RNase resistance and binding of small molecule ligands. From synthesis costs and function points of view, our current design of ~10% of 2'-5' links in a strand is representative of the future applications.

Addition of such unusual linkages into nucleic acid nanostructures might provide additional features for certain applications. For example, mixture of 3'-5'/2'-5' linkages have previously been tested in RNA aptamers and ribozymes,²⁹ thus opening up the possibility of using such functionalities within nucleic acid nanostructures. As mentioned above, 2'-5'-linked RNA oligonucleotides have special binding specificity to single stranded RNA over single stranded DNA, and can discriminate between DNA and RNA.⁵¹ This feature could be exploited in future nucleic acid nanomachines where DNA or RNA-preferred strand displacement reactions can drive chemical reactions or trigger release of compounds. In addition, it is known that single stranded RNA with 2'-5' linkages, phosphorothioate or phosphate chimeras resist enzymatic hydrolysis.²⁶ Preliminary ribonuclease assays in this study show that 2'-5'-linked oligonucleotides can potentially be used in DNA/RNA objects that can withstand the effect of RNases better than native structures. Construction of such modified nanostructures could provide new platforms for drug delivery with

better stability and functionality. For the tensegrity triangle motif tested here, one of the main applications (and the original notion of the field of DNA nanotechnology) is in the creation of designed DNA crystals that can act as a framework for the crystallization of other molecules.⁵² These 3D crystals have previously been used to host triplex forming oligonucleotides⁵³ and polyaniline,⁵⁴ and used to study the crystal structure and effect of torsional strain induced by base pair addition or deletion.⁵⁵ In that context, understanding the tolerance of 2'-5' linkages in this motif could lead to designed DNA crystals that can act as scaffolds for structural analysis of such unusual linkages and modifications.

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Author contributions

A.R.C. conceived the project, designed experiments, performed experiments, and wrote the manuscript. J.M. and J.V. per-

formed experiments. P.I. and A.C. performed simulations. K. H. and J.S. supervised the project and edited the manuscript. All authors provided comments on the manuscript.

Conflicts of interest

There are no conflicts of interest to declare.

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