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# m<sup>1</sup>A and m<sup>1</sup>G Potently Disrupt A-RNA Structure Due to the Intrinsic Instability of Hoogsteen Base Pairs

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# Abstract

The B-DNA double helix can dynamically accommodate G–C and A–T base pairs in either Watson-Crick or Hoogsteen configurations. Here, we show that G–C<sup>+</sup> and A–U Hoogsteen base pairs are strongly disfavored in A-RNA. As a result,  $N^{1}$ -methyl adenosine and  $N^{1}$ -methyl guanosine, which occur in DNA as a form of alkylation damage, and in RNA as a posttranscriptional modification, have dramatically different consequences. They create G–C<sup>+</sup> and A–U Hoogsteen base pairs in duplex DNA that maintain the structural integrity of the double helix, but block base pairing all together and induce local duplex melting in RNA, providing a mechanism for potently disrupting RNA structure through posttranscriptional modifications. The markedly different propensities to form Hoogsteen base pairs in B-DNA and A-RNA may help meet the opposing requirements of maintaining genome stability on one hand, and dynamically modulating the structure of the epitranscriptome on the other.

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# INTRODUCTION

The Watson-Crick (WC) double helix is the most common structural element in RNA and the dominant structure of genomic DNA. It provides the basis for templated replication, transcription, and translation, and also serves as a scaffold that defines the 3D structure of DNA, RNA, and their protein complexes. The canonical double helices formed by RNA (A-form) and DNA (B-form) differ in several important respects (Fig. 1a). In B-form DNA (B-DNA), the five-membered deoxyribose ring is flexible and favors the C2'-endo sugar pucker (Fig. 1a). In contrast, due to the sugar 2'-hydroxyl group (2'-OH), the sugar in A-RNA is more rigid and adopts an alternative C3'-endo conformation<sup>1,2</sup> (Fig. 1a). This in turn brings the oxygen atoms (O5' and O3') adjoining sequential nucleotides into closer proximity effectively compressing and rigidifying the A-form helix, widening its helical diameter, and displacing base pairs (bps) away from the helical axis<sup>1,3</sup>(Fig. 1a). In addition, B-DNA and A-RNA differ considerably with respect to their deformability, with B-DNA being generally more flexible<sup>4</sup>. For example, B-DNA is more bendable than A-RNA, and this property is of fundamental importance for many biochemical processes including the tight compaction of genome within the nucleus in higher order organisms.

Recently, NMR studies have uncovered a new dynamic property in canonical B-DNA; Watson-Crick (WC) dG–dC and dA–dT bps exist in a dynamic equilibrium with alternative Hoogsteen (HG) bps<sup>5,6</sup>. A HG bp is created by rotating a WC purine base  $\approx 180^{\circ}$  around the glycosidic bond to adopt a *syn* rather than *anti* conformation<sup>5</sup> (Fig. 1b). The two bases are also brought into closer proximity by  $\approx 2.0-2.5$  Å to form a unique set of hydrogen bonds (H-bonds) (Fig. 1b). HG bps exist transiently (lifetimes typically 0.1–1 ms) and in low abundance (populations typically <3%) in naked canonical B-DNA<sup>6,7</sup>. However, dA–dT and dG–dC<sup>+</sup> HG bps can become the dominant configuration (for review see ref. 8) in DNAprotein<sup>9</sup> and DNA-small molecule<sup>10</sup> complexes where they contribute to DNA recognition, in damaged nucleotides where they contribute to damage accommodation and repair<sup>11–13</sup>, and in the active sites of translesion synthesis polymerases that use HG pairing to bypass damage during DNA replication<sup>14</sup>. Purine-purine HG bps have also been shown to play important roles in DNA replication errors and in DNA damage accommodation and repair<sup>15,16</sup>.

Here, we set out to study WC-HG dynamics in canonical A-RNA duplexes. We show that unlike the canonical B-DNA double helix, rA–rU and rG–rC<sup>+</sup> HG bps are strongly disfavored in A-RNA duplexes. As a result, while the DNA double helix can absorb damaged nucleotides incapable of forming WC bps such as  $N^{1}$ -methyl deoxyadenosine (m<sup>1</sup>dA) and  $N^{1}$ -methyl deoxyguanosine (m<sup>1</sup>dG) by forming HG bps; the same methyl marks,  $N^{1}$ -methyl adenosine (m<sup>1</sup>rA) and  $N^{1}$ -methyl guanosine (m<sup>1</sup>rG), acting as a posttranscriptional modification in RNA, block base pairing altogether. This provides a direct mechanism for potently modulating the structure of the epitranscriptome. Our results indicate that HG-dependent DNA biochemical transactions may not be as readily supported in RNA duplexes and identify a unique dynamic property in B-DNA that may help enhance its ability to function as the repository of genetic information.

# RESULTS

### Absence of conformational exchange in A-RNA

We used NMR spin relaxation in the rotating frame  $(R_{1\rho})^{17-19}$  to examine whether WC bps in A-RNA duplexes transiently adopt HG bps like in B-DNA. A dynamic equilibrium between a dominant ground state (GS) and short-lived low-abundance 'excited state' (ES) conformation can lead to line-broadening of NMR resonances if the conformational exchange occurs on the  $\mu$ s-ms timescale. The  $R_{1\rho}$  experiment<sup>17</sup> measures this line broadening contribution ( $R_{ex}$ ) to the transverse relaxation rate ( $R_2$ ) during a relaxation period in which a continuous radiofrequency (RF) field is applied with variable power ( $\omega_{SL}$ ) and frequency ( $\omega_{RF}$ ). The resulting dependence of  $R_2+R_{ex}$  on  $\omega_{SL}$  and  $\omega_{RF}$ , referred to as relaxation dispersion (RD), can be fitted to the Bloch-McConnell equations describing *n*-site exchange<sup>20</sup> to extract exchange parameters of interest, including the population of the ES ( $p_{ES}$ ), the rate constant for conformational exchange ( $k_{ex} = k_{forward} + k_{backward}$ ), and the difference between the chemical shifts of the ES and GS ( $\omega = \omega_{ES}-\omega_{GS}$ ).

So far, RD studies have provided evidence for µs-ms conformational exchange in noncoding RNAs involving localized changes in secondary structure in and around noncanonical motifs (reviewed in ref. 18). The RD contributions from such chemical exchange processes can mask the ability to detect WC ≈ HG exchange. To hone in on WC ≈ HG exchange in A-RNA, we carried out <sup>13</sup>C and <sup>15</sup>N  $R_{1\rho}$  RD experiments<sup>21,22</sup> on an RNA duplex (hp-A<sub>6</sub>-RNA) capped by a stabilizing apical loop lacking non-canonical motifs and containing the same sequence (A<sub>6</sub>-DNA) for which we first reported transient HG bps in B-DNA<sup>6</sup> (Fig. 1c and Supplementary Fig. 1a). We targeted purine-C8, C-C6, G-N1, T-N3 and sugar purine-C1' sites (highlighted in orange in Fig. 1b), all of which have previously been (Supplementary Table 1 and Supplementary Note). In stark contrast to B-DNA, all RD profiles measured in in hp-A<sub>6</sub>-RNA were flat with no signs of detectable conformational exchange on the µs-ms timescale (Fig. 1d). No RD was observed across a variety of rG-rC and rA-rU WC bps, under low pH conditions (pH = 5.4) that allow optimal RD detection of WC $\rightleftharpoons$ HG exchange in B-DNA<sup>6,24</sup>, upon increasing the temperature (T = 35°C), and in the presence of 4 mM Mg<sup>2+</sup> (at pH = 6.8 and T = 5 or 25°C) (Figs. 1d and Fig. 2b, Supplementary Fig. 1b).

To broaden the search for WC $\rightleftharpoons$ HG exchange in A-RNA duplexes, we carried out additional RD measurements over a wide range of conditions (pH = 5.4–8.4 and T = 5– 35°C) for another ten rG–rC and seven rA–rU bps embedded in distinct sequence and structural contexts in three additional RNA molecules, including a GC-rich hairpin (hpgc<sup>GU</sup>), elongated duplex (E-gc), the transactivation response element (TAR) and a mutant form of TAR (TAR-UUCG<sup>GU</sup>) that is impaired from undergoing secondary structure chemical exchange<sup>25</sup> (Fig. 2a). In all cases we did not detect any signs of RD (Fig. 2b and Supplementary Fig. 1b). These results, together with our studies<sup>25–27</sup> reporting flat RD profiles for RNA WC bps near non-canonical motifs and mismatches (wtTAR and P5abc, Fig. 2a) and for the reverse wobble rG<sup>*syn*</sup>–rU mispairs in apical loops<sup>28</sup> stand in striking contrast to canonical duplex DNA in which we have robustly observed WC $\rightleftharpoons$ HG exchange

in all 35 dA–dT and dG–dC bps examined to date in a wide variety of positional and sequence contexts in eight different duplexes that have varying lengths and stabilities<sup>6,7</sup>.

The lack of detectable WC $\rightleftharpoons$ HG exchange in A-RNA could in principle result from small differences between the WC and HG NMR chemical shifts ( $\omega < 0.5$  p.p.m. for carbon chemical shifts). However, based on density functional theory calculations (DFT)<sup>6,29</sup> and a survey of *syn* purine base chemical shifts in the Biological Magnetic Resonance Data Bank<sup>30</sup>, it is highly unlikely that such a large transformation in base pairing would result in such small changes in chemical shifts for the different sugar (C1') and base (C8, C6 and N1/N3) sites targeted for RD measurements (Supplementary Note). The absence of RD is unlikely to be due to the exchange rate falling outside the detection limits of the RD experiment given that flat profiles are observed over a wide range of temperatures and pH conditions (Fig. 2b and Supplementary Fig. 1b) known to significantly alter the WC $\rightleftharpoons$ HG exchange rate in B-DNA<sup>6,7</sup>.

A more likely explanation is that HG bps are energetically disfavored in A-RNA duplexes, and have an abundance that falls below the detection threshold of the RD experiment (population < 0.01%). Indeed, a survey of X-ray structures of RNA duplexes in the Protein Data Bank (PDB)<sup>31</sup> failed to identify a single rG–rC<sup>+</sup> or rA–rU HG bp within continuous A-RNA duplexes out of a total of 123,935 rG–rC and rA–rU bps (Methods); while in sharp contrast, a similar survey conducted recently on B-DNA duplexes<sup>32</sup> identified 54 dG–dC<sup>+</sup> or dA–dU HG bps out of a much smaller set of 51,485 bps. The survey identified a single rA–rU HG bp (for example, PDBID: 1GID<sup>33</sup>) within an RNA duplex that fell well outside the A-form structural context being surrounded by a bulge and internal loop. The survey did identify several examples of long-range rG–rC<sup>+</sup> and rA–rU HG bps forming tertiary contacts; rG–rC<sup>+</sup> and rA–rU HG bps in triplexes and reverse rA–rU HG bps within duplexes typically near rG–rA mismatches where purines adopt *anti* rather than *syn* conformation, as well as several examples of HG mispairs in A-RNA duplexes (e.g. rG<sup>*syn*</sup>–rG<sup>*anti*</sup>, and rG<sup>*syn*</sup>–rA<sup>*anti*</sup>) (Supplementary Note).

# m<sup>1</sup>A and m<sup>1</sup>G modified A-RNA

If rG–rC<sup>+</sup> and rA–rU HG bps are indeed thermodynamically disfavored in A-RNA, they should prove more difficult to trap using chemical modifications known to stabilize dG–dC<sup>+</sup> and dA–dT HG bps in B-DNA<sup>6</sup>. We therefore examined whether HG bps could be stably trapped in A-RNA duplexes using m<sup>1</sup>rA and m<sup>1</sup>rG. These modified bases block WC pairing because of steric collisions with the methyl group and because the methylation knocks out one of the WC H-bonds (Fig. 3a). Both m<sup>1</sup>dA and m<sup>1</sup>dG occur in DNA due to alkylation damage<sup>12,13</sup>. In B-DNA, m<sup>1</sup>dA and m<sup>1</sup>dG are accommodated as m<sup>1</sup>dA–dT and m<sup>1</sup>dG–dC<sup>+</sup> HG bps<sup>6,11,34</sup> (Fig. 3a), which can in turn be recognized and repaired by damage repair enzymes<sup>12,13</sup>. m<sup>1</sup>rA and m<sup>1</sup>rG can also occur as a form of alkylation damage in RNA but they are also highly conserved post-transcriptional modifications in transfer and ribosomal RNAs that play critical structural and functional roles often by blocking WC base pairing<sup>35–39</sup>. m<sup>1</sup>rG and m<sup>1</sup>rA have been shown to induce duplex-to-hairpin transitions in palindromic RNA sequences where the modified base favors an unpaired conformation within apical loops<sup>40,41</sup>. Recent genome-wide studies have shown m<sup>1</sup>rA to be a dynamic

In prior studies<sup>6</sup>, we showed that m<sup>1</sup>dA16 and m<sup>1</sup>dG10 form stable m<sup>1</sup>dA16-dT9 and  $m^{1}dG10-dC15^{+}$  HG bps stabilized by unique H-bonds in A<sub>6</sub>-DNA while minimally impacting neighboring WC bps as judged based on observation of HG-specific chemical shifts, Nuclear Overhauser effect spectroscopy (NOESY) cross-peaks, and imino resonances (highlighted in Fig. 3a). In contrast, we did not observe any NMR evidence for HG bps or syn purine bases in the corresponding A<sub>6</sub>-RNA duplex containing m<sup>1</sup>rA16 or m<sup>1</sup>rG10 (Fig. 3b - e and Supplementary Fig. 2). This was the case despite the fact that A<sub>6</sub>-DNA and A<sub>6</sub>-RNA duplexes have very similar thermodynamic stabilities. Rather, the rA-C1' chemical shifts falls in a region that is consistent with A-form helical residues (Fig. 3c and Supplementary Note). We also observe continuous NOE distance-based connectivity between the m<sup>1</sup>rA and its preceding residue that are consistent with an *anti* conformation for the purine base (Supplementary Fig. 2). These data, together with the absence of strong H1'-H8 NOEs expected for syn base (Fig. 3d) and imino and amino resonances indicative of H-bonding (Fig. 3e) suggest that in A<sub>6</sub>-RNA, m<sup>1</sup>rA adopts a predominantly unpaired anti conformation although we cannot rule out transient formation of syn base conformations. The resonances belonging to  $m^{1}rG$  in A<sub>6</sub>-RNA were broadened out of detection suggesting extensive conformational exchange at the µs-ms timescale with no NMR evidence for HG pairing given the absence of downfield-shifted rC-H4 (Supplementary Fig. 1 and Fig. 3e). However, we cannot exclude micro-to-millisecond exchange between syn and anti conformations for the m<sup>1</sup>rG base since the resonances are broadened out of detection<sup>24</sup>.

Compared to  $A_6$ -DNA, m<sup>1</sup>A and m<sup>1</sup>G also induced more significant structural perturbations in  $A_6$ -RNA (Fig. 3b and Supplementary Table 2). We do not observe some of the imino resonances belonging to WC bps neighboring the modified site (Fig. 3f), which suggests loss of H-bonds and the melting of these base pairs. The modifications also induced more extensive chemical shift perturbations (Fig. 3g and Supplementary Fig. 1c; highlighted in orange in Fig. 3b) and line broadening (Supplementary Fig. 1c; highlighted in grey in Fig. 3b) in the sugar and base resonances that extend to the partner strand. The direction of the perturbations is consistent with deviations from a helical conformation (Supplementary Fig. 1c). The perturbations were particularly pronounced for m<sup>1</sup>rG, which broadened all imino resonances out of detection at 35°C, consistent with significant melting of the entire duplex (Supplementary Fig. 1d). Thus, HG bps are so sufficiently disfavored in A-RNA that m<sup>1</sup>rA and m<sup>1</sup>rG prefer to adopt predominantly non-helical conformations that disrupt the duplex structure.

Similar results were obtained in GC-rich ( $gc^{m1A}$ ) and scrambled ( $A_2^{m1A}$ )<sup>6</sup> (B.S., H.Z., Y.X., H.M.A., unpublished) duplexes with m<sup>1</sup>A consistently forming HG bps or adopting a *syn* conformation in B-DNA but not in A-RNA (Fig. 3b,d and Supplementary Fig. 3 and Supplementary Fig. 4) and with the modification more significantly perturbing the structure of A-RNA as compared to B-DNA (Fig. 3b,g). The structural perturbations induced by m<sup>1</sup>rA varied with sequence, and were either distributed across many WC bps ( $A_6$ -RNA and  $A_2$ -RNA) or more severe but localized to the modified and partner base (gc-RNA) (Fig. 3b and Supplementary Fig. 1c and Supplementary Fig. 3). In all cases we did not observe any

evidence for m<sup>1</sup>rA or m<sup>1</sup>rG induced duplex-to-hairpin transition based on spectral overlays with the unmodified counterparts (Supplementary Fig. 1c and Supplementary Fig. 3).

We corroborated the more portent destabilization of A-RNA as compared to B-DNA duplexes by m<sup>1</sup>A and m<sup>1</sup>G using UV melting experiments. m<sup>1</sup>dA destabilized A<sub>6</sub>-DNA, A<sub>2</sub>-DNA, and gc-DNA duplexes by  $G = 1.8-3.4 \text{ kcal mol}^{-1}$  (Fig. 3h) in good agreement with the relative stability of transient HG bps measured by NMR RD (2.1-4.3 kcal mol<sup>-1</sup>) and prior UV-melting studies of m<sup>1</sup>dA containing DNA duplexes (≈2 kcal mol<sup>-1</sup>)<sup>44</sup>. By comparison, m<sup>1</sup>rA and m<sup>1</sup>rG destabilized the corresponding A-RNA duplexes by a larger G = 4.3-6.5 kcal mol<sup>-1</sup>. Interestingly, this greater destabilization is comparable amount to the relative stability of the base-opened state<sup>45</sup>. This suggests that in A-RNA, the modification results in a conformation similar to the base-opened state, consistent with the NMR evidence for local melting. Greater destabilization (by  $\approx 1.1-4.7$  kcal mol<sup>-1</sup>) of A-RNA as compared to B-DNA was robustly observed across different duplex and hairpin contexts in the presence or absence of  $Mg^{2+}$  and with the destabilization being principally enthalpic (Fig. 3h, and Supplementary Table 3). The potent m<sup>1</sup>rA induced destabilization of duplex RNA is highly significant considering recent studies showing it to be a dynamic mRNA modification with roles in post-transcriptional gene regulation<sup>42,43</sup>. For comparison, the other well studied mRNA modification<sup>46</sup> N<sup>6</sup>-methyladenosine (m<sup>6</sup>A) which affects mRNA localization, stability, translation, and splicing destabilizes A-RNA by only 0.5-1.7 kcal mol<sup>-1 47</sup>.

The more potent m<sup>1</sup>A and m<sup>1</sup>G destabilization of A-RNA as compared to B-DNA is unlikely to be due to differences in steric contacts involving the methyl group in a HG bp configuration (Supplementary Note). While the positive charge on m<sup>1</sup>rA may affect stacking and H-bonding interactions, significant destabilization is also observed with the neutral m<sup>1</sup>rG, and the m<sup>1</sup>A destabilization is greater for A-RNA as compared to B-DNA (Fig. 3h). Rather, the greater destabilization observed in A-RNA is likely due to the higher energetic cost of forming HG bps in A-RNA as compared to B-DNA.

## Why are HG bps disfavored in A-RNA?

Why are HG bps so strongly disfavored in RNA as compared to DNA duplexes? The HG bp could in principle be disfavored in RNA due to the sugar 2'-OH at the purine residue. The 2'-OH helps bias the sugar pucker toward the C3'-endo conformation (Fig. 1a) due to unfavorable steric contacts between O2' and O3' and electronic effects involving the 2'-OH group<sup>2,48</sup>. This in turn disfavors the *syn* purine base conformation even in nucleosides<sup>49</sup> and single-stranded polynucleotides<sup>50</sup> due to unfavorable base-sugar steric contacts (N3–H3' and N3–O4'). The *syn* purine conformation may also destabilize water-bridged interactions involving the 2'-OH group on the ribose moiety of the flipping purine base is sufficient to suppress WC $\rightleftharpoons$ HG exchange, we carried out  $R_{1\rho}$  RD experiments on site or strand specifically labeled A<sub>6</sub>-DNA duplexes containing a single ribonucleotide, rA16 or rG10 (Supplementary Fig. 5a,b and Methods). These RD measurements were also of interest given that single ribonucleotides are frequently incorporated in DNA during replication and

can have important biological consequences through mechanisms that are not fully understood<sup>52</sup>.

Both rA16 and rG10 formed the expected rA16–dT9 and rG10–dC15 WC bps<sup>53</sup> and exhibited RD consistent with WC HG exchange (Fig. 4a). The lower  $R_{ex}$  contribution observed for the rA16 and rG10 substituted samples relative to the unmodified DNA duplex can be attributed to  $\approx$ 4-fold faster exchange rate ( $k_{ex} = 2325 \text{ s}^{-1}$  versus 595 s<sup>-1</sup>) in the case of rA16 and a combination of slightly smaller  $\omega$  (1.8 versus 2.1 p.p.m.) and transient HG population (0.8% versus 1.3%) in the case of rG10 (Fig. 4a, Supplementary Table 4 and Supplementary Fig. 5c). Neither rA16 nor rG10 significantly impacted the abundance of the transient HG bps relative to the unmodified A<sub>6</sub>-DNA duplex (Fig. 4a and Supplementary Table 4) indicating that the purine sugar 2'-OH group alone cannot account for the lack of observable WC HG exchange in A-RNA duplexes. We confirmed these findings by analyzing A<sub>6</sub>-DNA duplexes containing  $N^1$ -methylated single ribonucleotide, m<sup>1</sup>rA16 or m<sup>1</sup>rG10. In both cases, we observe stably formed m<sup>1</sup>rA16–dT9 and m<sup>1</sup>rG10–dC15<sup>+</sup> HG bps (Supplementary Fig. 5b). These data suggest that the destabilization of HG bps requires the broader A-form RNA helical context.

Next, we examined whether there were unique steric clashes that could disfavor *syn* purine bases within the compact A-RNA helix context that are absent in the more capacious B-form DNA helix. Indeed, flipping the purine base around the glycosidic  $\chi$ -angle through a range of angles (160°–200°) that span *syn* base conformations found in RNA helices (Supplementary Note) resulted in greater steric clashes in A-RNA as compared to B-DNA. The additional base-sugar (N3–H3' and N3–O4') and base-backbone (N3–O5') clashes observed in A-RNA arise due to both the C3'-endo sugar pucker and unique phosphodiester backbone conformation at the *syn* purine residue (Fig. 4b).

To further examine the energetics of the WC $\rightleftharpoons$ HG transition, we carried out biased MD simulations on the A<sub>6</sub>-DNA duplex and hp-A<sub>6</sub>-RNA hairpin, as well as a  $3' \rightarrow 5'$  inverted sequence of the hp-A<sub>6</sub>-RNA hairpin. A bias was applied on dA16 or rA16 starting in a WC bp configuration to force purine base flipping and a transition to a target HG configuration (Methods and Supplementary Movie 1, 2). The computed mean interaction energy (averaged over an ensemble of biased trajectories) as a function of the  $\gamma$ -angle along the WC $\rightleftharpoons$ HG transition (Methods) reveals a clear two state transition in the case of B-DNA, consistent with previous results<sup>6</sup>, whereas in the cases of A-RNA the resultant HG bp is significantly destabilized relative to its WC bp counterpart; for A-RNA, the energy profile in the syn region has much higher relative energies than in the case of DNA (Fig. 4c). In accord with this energetic destabilization, the simulations reveal that flipping the purine base in A-RNA is accompanied by major structural disruption of the surrounding base pairs (Supplementary Movie 2), consistent with m<sup>1</sup>rA induced NMR chemical shift perturbations, which were more pronounced for residues 3' to the modified nucleotide (Supplementary Fig. 1c). The extent of the disruption for the neighboring bp was far less significant in B-DNA (Supplementary Movie 1).

We corroborated these findings using unbiased MD simulations, which began with a HG bp embedded in various duplex and hairpin contexts (Methods). The HG H-bonding remained

stable during the course of the simulation in the case of B-DNA, B-DNA containing a single rA, and B-DNA containing m<sup>1</sup>dA (Supplementary Table 5 and Supplementary Movie 3–5). In contrast, for A-RNA strong disruption of N7---H3-N3 HG H-bond between A16 and U9 was observed in cases of the hp-A<sub>6</sub>-RNA hairpin (Supplementary Table 5). In  $\approx$ 35% of the trials in the case of 3' $\rightarrow$ 5' sequence hp-A<sub>6</sub>-RNA, the HG bp even transitioned rapidly after equilibration back to a WC bp. Strikingly, in the case of m<sup>1</sup>rA embedded in A-RNA, the HG bp caused melting of the A-form helix (Fig. 4d and Supplementary Movie 6).

Taken together, these results indicate that HG bps are disfavored in the more compact A-RNA helix due to steric contacts that are difficult to alleviate without substantially perturbing the A-form helix structure.

## DISCUSSION

Duplex B-DNA can stably accommodate dA-dT and  $dG-dC^+$  HG bps which can in turn play roles in sequence-specific DNA recognition, damage induction and repair, and in DNA replication. In contrast, our results indicate that rA–rU and rG–rC<sup>+</sup> HG bps are so unstable in the more compressed A-RNA that melting is preferred over the HG bp conformation. It remains to be seen whether the greater instability of HG bps in A-RNA as compared to B-DNA extends to purine-purine HG mispairs (Supplementary Note), which play important roles in replication<sup>14,54</sup> and translation errors<sup>55</sup>, mismatch repair<sup>8</sup>, as well in translational reprogramming<sup>56,57</sup>.

The markedly different stability of the A–T/U and G–C<sup>+</sup> HG bp in RNA and DNA duplexes provides a basis for achieving opposing functions at the genome and transcriptome levels (Fig. 5a). If DNA did not have a capacity to form HG bps, and instead behaved like RNA, lesions such as m<sup>1</sup>dA and m<sup>1</sup>dG that block canonical WC base pairing could greatly destabilize the double helix and potentially cause genomic instability (Fig. 5a). The ability to form HG bps therefore endows DNA with an additional layer of chemical stability over its RNA counterpart that goes beyond resistance to hydrolysis due to the absence of the sugar 2'-OH group. On the other hand, the greater instability of HG bps in A-RNA gives rise to a chemical switch in the form of m<sup>1</sup>rA and m<sup>1</sup>rG that can potently modulate RNA structure (Fig. 5a). While it has long been recognized that m<sup>1</sup>A and m<sup>1</sup>G can modulate the structure and function of tRNA, rRNA, and other non-coding RNAs<sup>37–39,58</sup>, this functionality hinges on the unique instability of HG bps in A-RNA uncovered in this work.

For example, m<sup>1</sup>rA9 has been shown to stabilize the native structures of human mitochondrial tRNAs by blocking helical rA–rU WC bps that would otherwise stabilize alternative secondary structures<sup>58</sup> (Fig. 5b). Likewise, m<sup>1</sup>rG37 next to the anti-codon loop, which is highly conserved in most tRNAs that read the CNN codon, has been shown to prevent +1 frameshifting by blocking base-pairing between G37 and the first rC in the codon sequence<sup>37,39</sup> (Fig. 5c). If RNA behaved like DNA, such posttranscriptional modifications would simply create HG bps and fail to block base pairing and have their intended functional consequence (Fig. 5).

m<sup>1</sup>rA was recently shown to be a reversible mRNA modification in eukaryotic cells, from yeast to mammals, that can respond to changes in physiological conditions<sup>42,43</sup>. It is enriched in the 5' UTR near start codons and was shown to promote translation through mechanisms that are not yet understood<sup>42,43</sup>. The formation of stable mRNA secondary structure around start codons has been shown to reduce translational efficiency<sup>59,60</sup>. Although it is unclear whether these m<sup>1</sup>rA modifications target adenine nucleotides involved in WC base pairing, it is possible that m<sup>1</sup>rA enhances translation in part by destabilizing secondary structure at the 5' UTR near the start codons. Indeed, based on our results, m<sup>1</sup>rA should also be capable of stabilizing alternative RNA secondary structures that feature bulged adenosines even if they are disfavored by as much as ≈5 kcal mol<sup>-1</sup> in the absence of the modification. At the same time, placement of m<sup>1</sup>rA in an unpaired bulged conformation can make it accessible to demethylases for achieving efficient reversible control at the epitranscriptomic level (Fig. 5d). Further studies are needed to test this proposed mechanism for m<sup>1</sup>A-enhanced translation.

# **ONLINE METHODS**

## Sample Preparation

**NMR buffer**—All RNA and DNA samples were buffer exchanged at least three times using a centrifugal concentrator (EMD Millipore) until containing >99.9% of the desired buffer, which unless stated otherwise, consisted of 15mM sodium phosphate, 25 mM NaCl, 0.1 mM EDTA with pH 5.4 or 6.8 and 10% D<sub>2</sub>O.

**Uniformly** <sup>13</sup>**C**/<sup>15</sup>**N labeled RNA and DNA samples**—hp-A<sub>6</sub>-RNA and single-strands of the E-gc and TAR-UUCG<sup>GU</sup> were prepared using *in vitro* transcription with uniformly <sup>13</sup>C/<sup>15</sup>N-labeled ribonucleotide triphosphates (Cambridge Isotope Laboratories), T7 polymerase (Takara Mirus Bio Inc.) and synthetic DNA templates (Integrated DNA Technologies, Inc.), purified by 20% (w/v) denaturing polyacrylamide gel electrophoresis (PAGE) and electro-eluted into 20 mM Tris buffer (pH = 8) followed by ethanol precipitation as described previously<sup>26</sup>. The uniformly labeled T6-strand in A<sub>6</sub>-DNA<sup>rG</sup> and uniformly labeled A<sub>6</sub>-DNA were prepared by the primer-extension approach<sup>61</sup> using uniformly <sup>13</sup>C/<sup>15</sup>N-labeled deoxyribonucleotide triphosphates (Silantes) as previously described<sup>6</sup>.

**m<sup>1</sup>A and m<sup>1</sup>G containing oligonucleotides**—Oligonucleotides were purchased from Keck Oligo Synthesis Resource (W.M. Keck Foundation) with Glen-Pak<sup>TM</sup> DNA/RNA cartridge purification ( $A_6$ -RNA<sup>m1A</sup>,  $A_2$ -RNA<sup>m1A</sup>, gc-RNA<sup>m1A</sup>,  $A_6$ -DNA<sup>m1G</sup>,  $A_6$ -DNA<sup>m1rA</sup>, gc-DNA<sup>m1A</sup>, hp- $A_6$ -RNA<sup>m1A</sup>, hp- $A_6$ -DNA<sup>m1A</sup>, hp- $A_6$ -DNA<sup>m1A</sup>, hp- $A_6$ -DNA<sup>m1A</sup>, hp- $A_6$ -DNA<sup>m1A</sup>) Midland Certified Reagents with reverse-phase (RP) HPLC purification ( $A_6$ -DNA<sup>m1A</sup>), and GE Healthcare Dharmacon Inc. with RP-HPLC purification ( $A_6$ -RNA<sup>m1G</sup>,  $A_6$ -DNA<sup>m1rG</sup> and hp- $A_6$ -RNA<sup>m1G</sup>). To minimize Dimroth rearrangement of m<sup>1</sup>A into m<sup>6</sup>A<sup>62,63</sup>, all DNA and RNA oligonucleotides containing m<sup>1</sup>A were synthesized and deprotected using the UltraMild protocol (http:// www.glenresearch.com/Technical/TB\_UltraMild\_Deprotection.pdf; Glen Research Corporation).

Assessing purity of m<sup>1</sup>A and m<sup>1</sup>G containing oligonucleotides—Samples were assessed using 20% denaturing PAGE, MALDI Mass Spectrometry, Liquid Chromatography-Mass Spectrometry (LC-MS) and NMR spectroscopy. For hairpin constructs hp-gc-RNA<sup>m1A</sup> and hp-A<sub>6</sub>-RNA<sup>m1A</sup>, we obtained evidence for incomplete base deprotection during synthesis based on observation of additional imino proton and acetyl group (the N4 protecting group on the cytosine) resonances and NOE talk between the two. Evidence for the acetyl groups was also obtained by quantitative mass spectrometry (LC-MS). We suspect that incomplete deprotection arises due to formation of stable secondary structure in these hairpin constructs during the UltraMild deprotection step. These impurities could be effectively eliminated by synthesizing individual single strands of duplex versions of the hairpin sequence (gc-RNA<sup>m1A</sup> and A<sub>6</sub>-RNA<sup>m1A</sup>).

In all cases, the NMR chemical shifts of the  $N^1$ -methyl group and base moieties (A-C2, N1C, N1H) were consistent with m<sup>1</sup>A with no evidence for Dimroth rearrangements<sup>62</sup>, which lead to formation of m<sup>6</sup>A (Supplementary Fig. 1c and Supplementary Fig. 3). In particular, we observed  $\approx 4$  p.p.m. upfield shift in m<sup>1</sup>A-C2 (Supplementary Fig. 1c and Supplementary Fig. 3) consistent with base protonation, which is expected for m<sup>1</sup>A but not m<sup>6</sup>A. This, as well as the observation of two amino protons (H61 and H62) with distinct chemical shifts involved in HG H-bonding in DNA (Fig. 3e and Supplementary Fig. 2) indicates a major positively charged amine tautomer rather than a neutral imine tautomeric form<sup>64</sup>. Nevertheless, we cannot rule out the existence of the neutral imine tautomeric form transiently and/or in low-abundance.

**Unmodified oligonucleotides**—Unmodified RNA oligonucleotides were synthesized using an in-house MerMade 6 Oligo Synthesizer employing 2'-TBDMS RNA phosphoramidites (ChemGenes) on 1 µmol standard synthesis columns (1000 Å) from BioAutomation using the option to leave the final 4.4'-dimethoxytrityl (DMT), the 5'protection group on for the cartridge purification. The oligonucleotide was cleaved from each 1 µmol column using  $\approx 1$  mL ammonia methylamine (1:1 ratio of 30% ammonium hydroxide and 30% methylamine) followed by 2-hour incubation at room temperature to allow base deprotection. The solution was then subjected to airflow until complete evaporation, leaving the desired product oligonucleotide as dried crystals. The crystals were then dissolved in 115 µL DMSO, mixed with 60 µL TEA and 75uL TEA"135;3HF, and incubated at 65°C for 2.5 hours for 2'-deprotection. The reaction was quenched using Glen-Pak<sup>TM</sup> RNA quenching buffer and loaded onto Glen-Pak<sup>TM</sup> RNA cartridges (Glen Research Corporation) for purification following the online protocol (http://www.glenresearch.com/ Technical/GlenPak\_UserGuide.pdf). Samples were ethanol precipitated and exchanged into NMR buffer. Unmodified DNA oligonucleotides were purchased from Integrated DNA Technologies with standard desalting.

**Site-specifically** <sup>13</sup>C/<sup>15</sup>N-labeled samples—The A<sub>6</sub> strand of A<sub>6</sub>-DNA<sup>rA16</sup> containing C8-<sup>13</sup>C/<sup>15</sup>N-labeled adenosine was synthesized using an in-house Solid-phase Oligonucleotide Synthesizer (BioAutomation MerMade 6), C8-<sup>13</sup>C/<sup>15</sup>N-labeled adenosine phosphoramidite (see below), and unlabeled DNA phosphoramidites (ChemGenes) using 1 µmol scale 1000 Å CPG DNA columns (BioAutomation). The synthesized oligonucleotides

were cleaved and deprotected as described above for unlabeled RNA oligonucleotides and purified with the Glen-Pak<sup>TM</sup> RNA cartridge (Glen Research Corporation) followed by ethanol precipitation and exchange into the desired NMR buffer.

**Synthesis of 8-<sup>13</sup>C-adenosine phosphoramidite**—The 8-<sup>13</sup>C-adenine nucleobase was synthesized according to a published procedure<sup>65</sup>. The protection of the exocyclic amino group with a benzoyl moiety and the conversion to the 5'-O-DMT-2'-O-TOM-protected 8-<sup>13</sup>C-adenosine 3'-O-phosphoramidite was accomplished according to published procedures<sup>66,67</sup>. A detailed description on the chemical synthesis of 8-<sup>13</sup>C-purine RNA phosphoramidite building blocks will soon be published elsewhere.

#### NMR experiments

**Resonance assignment**—NMR data were collected on an 800 MHz Varian DirectDrive2 spectrometer equipped with a triple resonance HCN cryogenic probe; 700 Bruker Avance III spectrometer equipped with a triple-resonance HCN cryogenic probe; and 600 MHz Varian Inova NMR spectrometer equipped with a Bruker HCPN cryogenic probe. Data were processed and analyzed using NMRpipe<sup>68</sup> and SPARKY (T. D. Goddard and D. G. Kneller, SPARKY 3, University of California, San Francisco), respectively. Resonances were assigned using conventional 2D HSQC, HMQC, NOESY and HCN experiments.

Chemical shift perturbations (CSPs) induced by  $m^1A$  or  $m^1G$  for each residue ( $\omega_{residue}$ ) were calculated using equation (1)<sup>69</sup> from the average Euclidean distance of all measured CSP ( $\omega_C$ ,  $\omega_N$ , and  $\omega_H$ ):

$$\Delta \omega_{\text{residue}} = \sqrt{\frac{1}{N} \sum_{i}^{N} \left(\frac{\gamma_{i}}{\gamma_{1_{H}}} \Delta \omega_{i}\right)^{2}} \tag{1}$$

where  $\gamma_i$  is the gyromagnetic ratio of the i<sup>th</sup> nucleus (C, H or N), *N* is the total number of CSPs measured for each residue, and  $\omega_i$  is the difference in chemical shifts (in p.p.m.) for the i<sup>th</sup> nucleus between the m<sup>1</sup>A or m<sup>1</sup>G modified and unmodified duplex. Residues with  $\omega_{residue}$  0.1 p.p.m. are highlighted on the duplexes in Fig. 3b and Supplementary Fig. 2 and Supplementary Fig. 4. An average CSP ( $\omega_{avg}$ ) was calculated for each duplex by averaging  $\omega_{residue}$  for two base pairs above and below the modified bp.

<sup>13</sup>C and <sup>15</sup>N R<sub>1p</sub> relaxation dispersion—<sup>13</sup>C and <sup>15</sup>N  $R_{1p}$  RD experiments were performed at 600 MHz (14.1 T) and 700 MHz (16.4 T) Bruker spectrometers as previously described<sup>6,21,23</sup> using spinlock powers ( $\omega_{SL} 2\pi^{-1}$  Hz) and offset frequencies ( $\Omega 2\pi^{-1}$  Hz) listed in Supplementary Table 1. Magnetization of the spins of interest was allowed to relax under an applied spinlock for the following durations: [0–120 ms] for N1/N3 in hp-A<sub>6</sub>-RNA and E-gc; [0–60 ms] for C8/C1′ in hp-A<sub>6</sub>-RNA, E-gc, TAR-UUCG<sup>GU</sup>, A<sub>6</sub>-DNA, A<sub>6</sub>-DNA<sup>rA</sup> and A<sub>6</sub>-DNA<sup>rG</sup>.

# Analysis of R<sub>1p</sub> data

Fitting of <sup>13</sup>C and <sup>15</sup>N  $R_{10}$  data—Experimental  $R_{10}$  relaxation rate constants were calculated by fitting peak intensities versus relaxation delay durations to a single exponential decay<sup>21</sup>. Uncertainty in the fitted  $R_{1\rho}$  values (one s.d.) were derived using a Monte-Carlo method<sup>18</sup>.  $R_{1\rho}$  data were fitted to simulated  $R_{1\rho}$  values given by the solution to the Bloch-McConnell (BM) equations<sup>20</sup> at each given  $\Omega$  and  $\omega_{SL}$  combination. Residual sum of squares were minimized using a bounded least-squares algorithm<sup>70</sup> to give best-fit exchange parameters. The uncertainty in the chemical exchange parameters was calculated as the standard error of the fit<sup>18</sup>. A 2-state exchange model was used to fit the  $R_{10}$  RD profiles of A<sub>6</sub>-DNA, A<sub>6</sub>-DNA<sup>rA</sup> and A<sub>6</sub>-DNA<sup>rG</sup> with the initial magnetization aligned either along the effective field of the ground (for slow exchange with  $k_{ex}$   $\omega^{-1} < 1$ ) or average (for fast exchange with  $k_{\text{ex}} = \omega^{-1} = 1$ ) state<sup>71</sup>. For the dA16-C8 RD data measured in A<sub>6</sub>-DNA at low temperatures, both protocols yielded acceptable fits but resulted in different exchange parameters given the slower exchange rate (Supplementary Table 4). The exchange parameters obtained from average alignment protocol were selected based on a van't Hoff analysis<sup>6</sup> (Supplementary Fig. 5d). For the dC15-C6 RD data measured in A6-DNA<sup>rG</sup>, a 3state chemical exchange model without minor exchange with average alignment was statistically favored over 2-state models (Supplementary Fig. 5c). In all cases, Akaike information criterion (AIC)<sup>72</sup> and Bayesian information criterion (BIC)<sup>72</sup> were used to select the models.

#### Analysis of chemical shift and NOESY data

Chemical shifts and NOESY cross-peaks were used to characterize WC versus HG bps. The NOESY cross-peaks unique to HG bps include strong intra-nucleotide H1<sup>'</sup>–H8 NOE for *syn* purine, (i)A-H2–(i-1)H1<sup>'</sup>/H2<sup>'</sup> and (i)A-H2–(i-1)H6/H8 for *syn* adenosine, and H8–H3, A-H6/C<sup>+</sup>-H4–H3, (i)H3–(i+1)/(i-1)H1/H3, (i)A-H6/C<sup>+</sup>-H4–(i+1)/(i-1)H1/H3 NOEs for connectivity involving imino or amino protons in both G-C and A-U/T HG bps<sup>6,73,74</sup>. Absence of the canonical sequential (i-1)H1<sup>'</sup>–(i)H8 NOE is also expected for *syn* purines due to the base flip. As described previously<sup>6</sup>, the HG bps are also characterized by a unique set of chemical shifts relative to WC bps including downfield shifted purine-C8 and purine-C1<sup>'</sup>, protonated cytosine-C6 ( $\approx$ 3 p.p.m.) and upfield shifted protonated cytosine-C5, guanine-N1 and thymine-N3 ( $\approx$ 1–2 p.p.m.). The C1<sup>'</sup> chemical shifts is also sensitive to sugar pucker. In A-RNA, deviations from the A-form C3<sup>'</sup>-endo toward C2<sup>'</sup>-endo sugar pucker leads to an upfield shift ( $\approx$ 4 p.p.m.)<sup>75,76</sup>. In addition, deviations from the A-form conformation due to loss of stacking and bulging out of nucleotides results in a downfield shift in the base C6/C8 and cytosine-C5 and upfield shift on sugar-C1<sup>'18</sup>.

# Density functional theory geometry optimizations and CS calculations

Density functional theory (DFT) calculations<sup>77</sup> using Gaussian 09c (Gaussian, Inc.) were performed as previously described<sup>6</sup> to compute chemical shifts for WC and HG bps in A-RNA and B-DNA. In all cases, protons were added using PyMOL (https://www.pymol.org/) and the phosphate backbone truncated leaving only the nucleoside motifs for each bp<sup>6</sup>. Calculations were performed on rA–rU HG bp obtained from snapshots of rA16–rU9 bp in biased MD simulation of hp-A<sub>6</sub>-RNA, rA–rU HG bp from the X-ray structure of P4-P6

domain of the Group I intron RNA (PDBID:  $1L8V^{78}$ ), tertiary rG–rC<sup>+</sup> HG bp in the structure of 23S ribosomal RNA-protein complex (PDBID:  $3U56^{79}$ ) and rG<sup>syn</sup>–rG<sup>anti</sup> mispair in duplex RNA structure (PDBID:  $3CZW^{80}$ ). Reference rA–rU or rG–rC WC bps were taken from MD snapshot or from the same X-ray structures used to obtain HG bps (Supplementary Fig. 5e). Two runs of geometry optimizations were carried out using the B3LYP functional with 3–21G and 6–311+G(2d,p) basis sets, respectively, with all heavy atoms (C, N and O) frozen. Carbon chemical shifts were computed using the GIAO method within the B3LYP/6–311+G(2d,p) basis set<sup>6</sup> on the converged configuration after the second run of optimization. The isotropic carbon chemical shift ( $\omega_{13C}$ ) was referenced to that of TMS ( $\omega_{TMS} = 182.4656$  p.p.m.), which was optimized and computed at the same level of theory. The carbon chemical shifts computed for reference WC bps were subtracted from those computed for HG bps to yield chemical shift changes upon HG formation ( $\omega = \omega_{HG}-\omega_{WC}$ ).

#### Analysis of UV melting data

The UV absorbance at 260nm (A<sub>260</sub>) as a function of temperature was measured on a Shimadzu UV-3600 UV-Vis-NIR spectrophotometer using the 8-cell sample holder with a Fisher Isotemp Refrigerated Circulator to regulate sample temperature. All DNA and RNA oligonucleotides were diluted directly from NMR samples using the same NMR buffer (15mM phosphate, 25mM NaCl, 0.1mM EDTA at pH = 5.4 or 6.8) unless stated otherwise and triplicate measurements were carried out for each oligonucleotide simultaneously using a sample volume of 125  $\mu$ L in each cell and an additional reference cell containing the same amount of buffer. The temperature was varied between 5°C and 90°C at a ramping rate of 1 °C min<sup>-1</sup>.

The melting temperature  $(T_m)$  and enthalpy ( H) for duplex association and hairpin folding was obtained by fitting the melting curves to equation (2) and equation (3)<sup>81</sup>, respectively,

$$f = \frac{1 + 4e^{(1/T_m - 1/T)\Delta H/R} - \sqrt{1 + 8e^{(1/T_m - 1/T)\Delta H/R}}}{4e^{(1/T_m - 1/T)\Delta H/R}}$$
(2)

$$f = \frac{e^{(1/T_m - 1/T)\Delta H/R}}{1 + e^{(1/T_m - 1/T)\Delta H/R}}$$
(3)

*T* is the measured temperature and *f* is the fraction of the remaining unmelted duplex or folded hairpin over the total concentration of duplex or hairpin ( $C_{\rm T}$ ), which is proportional to the measured A<sub>260</sub>.

The thermodynamic parameters G and S were calculated using equation (4) for duplex association and equation (5) for hairpin folding, respectively.

$$\Delta S = \Delta H / T_m - R \ln(C_T / 2); \Delta G = \Delta H - T \Delta S \quad (4)$$

$$\Delta S = \Delta H / T_m; \Delta G = \Delta H - T \Delta S \quad (5)$$

The fitting was carried out using nonlinear model fitting with Mathematica 10.0 (Wolfram Research). Errors in  $T_m$  and H represent the standard deviation (one s.d.) from the triplicate measurements. The destabilization effects due to  $m^1A$  and  $m^1G$  in DNA or RNA were calculated by taking the difference in free energy for folding i.e.

 $G = G_{\text{mod}} - G_{un\text{mod}}$ ,  $H = H_{\text{mod}} - H_{un\text{mod}}$  and  $S = S_{\text{mod}} - S_{un\text{mod}}$ .

Since the Dimroth rearrangement can occur for m<sup>1</sup>A in both DNA and RNA under basic conditions<sup>62</sup> and high temperatures<sup>63</sup>, melting experiments were repeated for all m<sup>1</sup>A containing duplexes when restricting the temperature to <65°C and when using both neutral (pH = 6.8) and acidic (pH = 5.4) conditions. These control experiments yielded reproducible melting curves and fitted thermodynamic parameters at neutral or acidic pH conditions that are within experimental error (Supplementary Table 3). <sup>1</sup>H 1D NMR spectra recorded for the A6-DNA<sup>m1A</sup> sample following melting showed insignificant changes and no evidence for Dimroth rearrangements (data not shown).

#### Steric analysis and survey of HG bps in RNA

A-T/U and G-C WC bps were obtained from idealized B-DNA and A-RNA helices built using 3DNA<sup>82</sup> (Fig. 4b). 146 and 159 WC bps surrounded by at least one WC bp above and below were obtained from high resolution (< 2 Å) X-ray structures of A-RNA and B-DNA duplexes, respectively in the PDB. Purine bases were flipped around the glycosidic bond and inter-atomic distances measured using an in-house python script. Note that proximity of the exocyclic amino group on guanine to the phosphate group during the base flip was not considered a steric clash given the potential for H-bonding. The survey of HG bps in RNA was carried out following the same protocol reported for B-DNA<sup>32</sup>. Briefly, all RNA X-ray structures with resolution 3 Å were downloaded from the PDB on August 31 2014. The same in-house program was used to identify rA-rU and rG-rC bps in RNA structures using three HG criteria (H-bonding, constricted C1'-C1' distance and syn purine)<sup>32</sup>. Redundancies defined as bps that are surrounded with the same sequence contexts and from same RNA or RNA-protein/ligand complexes were removed by manual inspection as described previously<sup>32</sup>. The survey identified a single rA-rU HG bp that reoccurs in four distinct X-ray structures of the P4-P6 domain of the Tetrahymena thermophila group 1 intron RNA (PDBID: 1GID<sup>33</sup>, 118V<sup>78</sup>, 1HR2<sup>83</sup>, and 2R8S<sup>84</sup>). The RNA HG survey also identified several examples of long-range HG bps forming tertiary contacts, HG bps in triplexes, and reverse rA-rU HG bps within duplexes typically near rG-rA mismatches.

#### **Biased and Unbiased Molecular Dynamics Simulations**

Structure Generation for MD Simulation—hp-A<sub>6</sub>-RNA, hp-A<sub>6</sub>-RNA  $3' \rightarrow 5'$ , and A<sub>6</sub>-DNA helices were built using make-na<sup>85</sup> with all bases in WC conformation. In the case of hp-A<sub>6</sub>-RNA, a duplex structure was generated using make-na and the UUCG loop attached and annealed using the CHARMM simulation package<sup>86</sup>. Structures with HG conformation at A16 were created by rotating along the glycosidic bond angle  $\chi$  by 180°.

**Unbiased MD equilibrium simulations**—All structures were simulated using constant temperature MD with CHARMM36 forcefield<sup>87</sup> and a generalized Born molecular volume (GBMV) implicit solvent<sup>88</sup>; parameters for m<sup>1</sup>A were taken from Xu *et al*<sup>89</sup>. Integration used a velocity-Verlet algorithm with a timestep of 1 fs. The cutoff for non-bonded list generation was 21 Å, the cutoff for non-bonded interactions was 18 Å, and the onset of switching for non-bonded interactions occurred at 16 Å. The SHAKE algorithm was used to constrain the covalent bonds to hydrogen atoms involved. Each structure was heated to 300.0 K with harmonic constraints on all non-hydrogen atoms, heating occurred in1 ps increments of 1.0 K for a total of 300 ps steps, followed by 200 ps equilibration at 300.0 K. Harmonic constraints were then gradually removed during a sequence of 4 reductions for 50 ps each. Unbiased production-run simulations were then run for 3 ns without constraints for each system. Ten independent simulations with hp-A<sub>6</sub>-RNA and A<sub>6</sub>-DNA<sup>rA</sup> with A16 in HG conformation were produced from independent conformations obtained during the heating and equilibration method described above. A<sub>6</sub>-DNA in HG was repeated twice.

Global RMSD was calculated from the single 3 ns trajectories of  $m^1A$  starting in HG for both hp-A<sub>6</sub>-RNA and A<sub>6</sub>-DNA,

$$\text{RMSD} = \sqrt{\frac{\sum\limits_{i=1}^{N} (r_i(t) - r^R)^2}{N}} \quad (6)$$

in which  $r_i(t)$  is the instantaneous coordinate of an atom and  $r^R$  is the position of the reference structure.

H-bond presence was evaluated using CHARMM's COOR HBOND module for each trajectory with cutoff distance and angle of 3.6 Å, and 120° following Goldsmith *et al*<sup> $\rho$ 0</sup>.

**Biased MD Simulations**—The protocols for minimization, heating, and solvation were identical to those used for the unbiased simulations. The biased molecular dynamics method<sup>91</sup> implemented in the CHARMM package was used to force conformational transitions between WC and HG states using a biasing potential W ( $\rho$ (t)) applied according to equation (7),

$$W(\rho(t)) = \begin{cases} \alpha/2(\rho(t) - \rho_a(t))^2, & if \ \rho(t) < \rho_a(t) \\ 0, & if \ \rho(t) \ge \rho_a(t) \end{cases}$$

where

$$\rho(t) = \frac{1}{N(N-1)} \sum_{i=1}^{N} \sum_{j\neq 1}^{N} (r_{ij}(t) - r_{ij}^{R})^{2}$$
(7)

 $\rho(t)$  is a collective distance between the instantaneous  $(r_{ij})$  and the reference structure  $(r_{ij}^R)$ , and  $\alpha$  the strength of the half-harmonic bias. In all cases, biases were placed between pairs of atoms that share a H-bond in the target structure, ensuring that the adenine base would not only perform the roughly 180° flip, but also form the definitive H-bonding structure of the desired WC or HG configuration. After the biased trajectories were generated, they were post-processed in CHARMM, outputting the  $\chi$ -angle dependence of the relative interaction energy value in the absence of the bias. Only successfully flipping trajectories were used, resulting in 40 trajectories for A<sub>6</sub>-DNA, 24 for hp-A<sub>6</sub>-RNA, and 25 for hp-A<sub>6</sub>-RNA  $3' \rightarrow 5'$ . The relative interaction energy was calculated for the base pair that includes the flipping base as well as the base pairs above and below the flipping base. Angle-energy pairs were binned into 50 bins and the mean of the energy was evaluated within each bin. Plots of relative interaction energy as a function of the  $\chi$ -angle were thus generated.

# Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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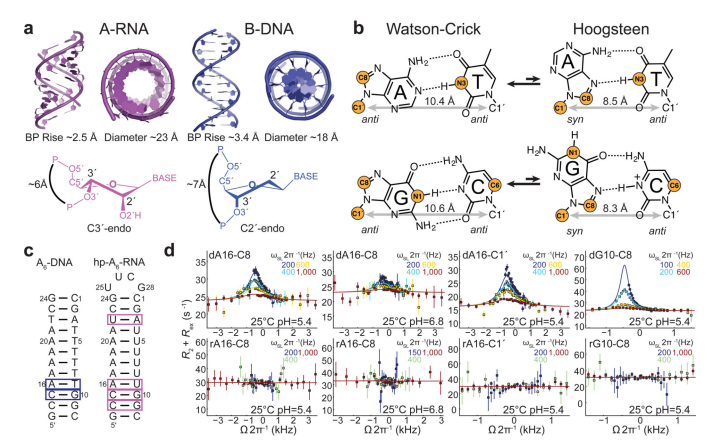
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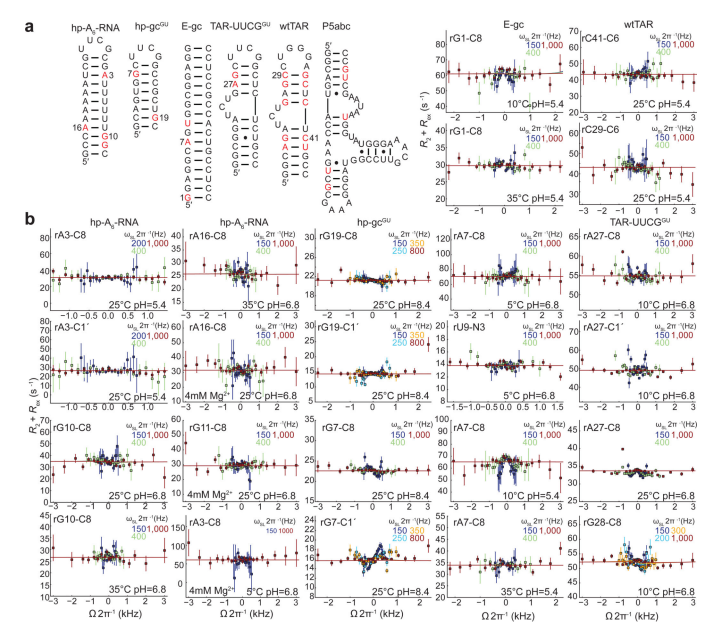
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### Figure 1.

Absence of detectable WC $\rightleftharpoons$ HG exchange in A-RNA by NMR relaxation dispersion. (a) Comparison of A-form RNA (violet) and B-form (blue) DNA double helices. (b) WC and HG bps in dynamic equilibrium in B-DNA. Sites used for RD measurements are highlighted in orange. (c) A<sub>6</sub>-DNA and hp-A<sub>6</sub>-RNA duplexes with bps targeted in RD measurements highlighted. (d) Off-resonance RD profiles showing  $R_2+R_{ex}$  as a function of spin lock offset ( $\Omega \ 2\pi^{-1}$ Hz, where  $\Omega = \Omega_{obs} - \omega_{RF}$ ) and power ( $\omega_{SL} \ 2\pi^{-1}$ Hz, in insets). Error bars represent experimental uncertainty (one s.d.) estimated from mono-exponential fitting using a Monte-Carlo based method (Methods). Solid line represents a fit to two-state exchange<sup>6</sup>.

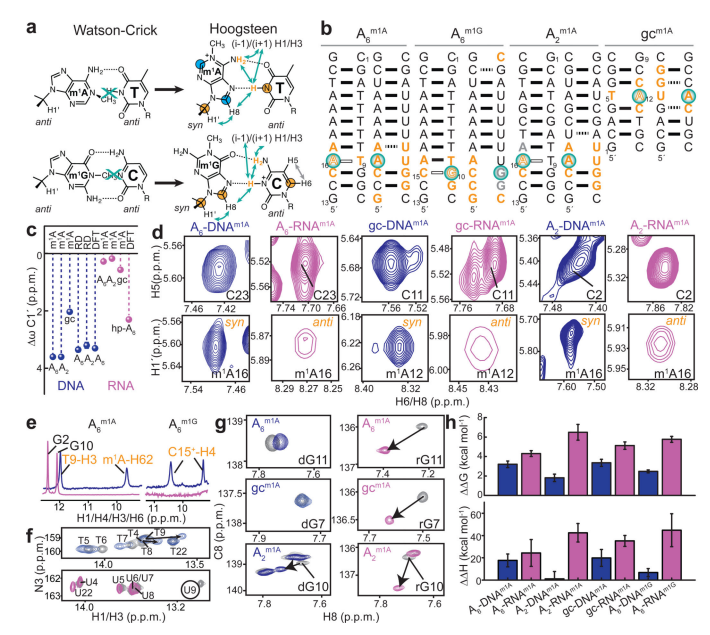
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#### Figure 2.

Lack of detectable exchange across diverse RNA sequence and structural contexts. (a) Secondary structures with bps showing no detectable RD highlighted in red. (b) Off-resonance RD profiles for the highlighted bps with error bars representing experimental uncertainty (one s.d.) estimated from mono-exponential fitting using a Monte-Carlo based method (Methods).

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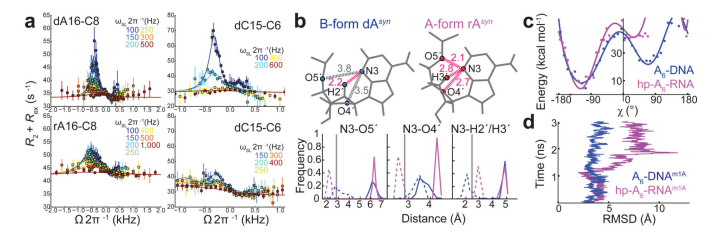


### Figure 3.

m<sup>1</sup>A and m<sup>1</sup>G do not form HG bps and disrupt A-RNA structure. (**a**)  $N^{1}$ -methylated purines trap HG bps in B-DNA. NMR chemical shift probes of HG bps are in orange and of purine methylation state in cyan. Arrows indicate characteristic HG NOE cross-peaks. (**b**) Duplexes containing m<sup>1</sup>A or m<sup>1</sup>G (turquoise circles). *syn* or *anti* purines deduced by NMR are shown as open and filled letters, respectively. HG and partially melted bps as deduced by NMR are indicated using open and dashed lines, respectively. Residues showing significant chemical shift perturbations or line-broadening due to m<sup>1</sup>A or m<sup>1</sup>G are colored orange and grey, respectively. (**c**) m<sup>1</sup>A or m<sup>1</sup>G induced purine-C1' chemical shift perturbations ( $\omega = \omega_{\text{modified}} - \omega_{\text{unmodified}}$ ) in A-RNA (violet) and B-DNA (blue). Shown for comparison are  $\omega = \omega_{\text{HG}} - \omega_{\text{WC}}$  measured for transient dA–dT HG bps by RD ("RD") in unmodified DNA duplexes (error bars showing one s.d.) and computed for adenine residues using DFT

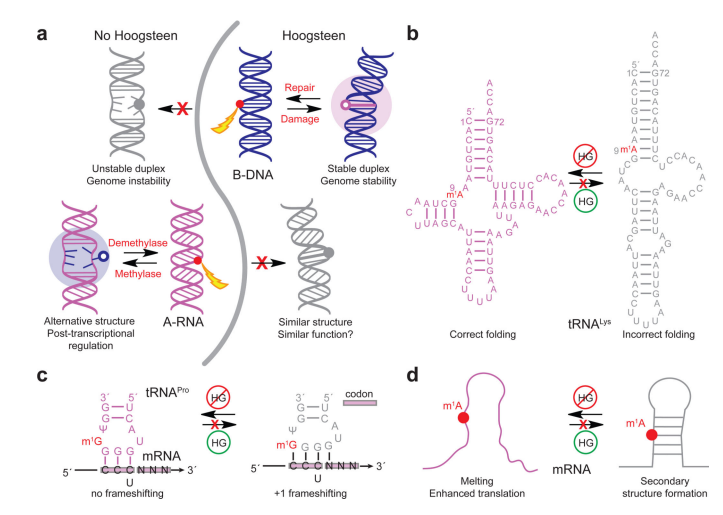
(Methods). (d) NOESY H1<sup>'</sup>-H8 cross-peaks showing *syn* purine bases in B-DNA but not A-RNA. Shown for reference is the cytosine base H5–H6 NOE with inter-atomic distance  $\approx 2.5$  Å. (e) 1D <sup>1</sup>H spectra showing the imino/amino resonances expected for HG type Hbonds in A<sub>6</sub>-DNA<sup>m1A</sup> and A<sub>6</sub>-DNA<sup>m1G</sup> but not in methylated RNA at 5°C and 15°C, respectively. (f) Example showing m<sup>1</sup>G induced loss of a WC imino resonance (highlighted in a circle) in A<sub>6</sub>-RNA but not A<sub>6</sub>-DNA in 2D NMR spectra. (g) Example downfield shifted carbon chemical shifts induced by m<sup>1</sup>A. (h) Free energy (G) and enthalpy (H) destabilization due to m<sup>1</sup>A and m<sup>1</sup>G in DNA (blue) and RNA (violet) duplexes measured by UV melting experiments with error bars, one s.d. (n = 3 independent measurements) (Methods and Supplementary Table 2).

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### Figure 4.

Source of HG instability in A-RNA. (a) Comparison of RD profiles measured in  $A_6$ -DNA<sup>rA</sup>,  $A_6$ -DNA<sup>rG</sup>, and  $A_6$ -DNA. Error bars correspond to one s.d. estimated from monoexponential fitting using a Monte-Carlo based method (Methods). Note that the larger  $R_2$  value in  $A_6$ -DNA<sup>rA</sup> A16-C8 as compared to  $A_6$ -DNA likely reflects decreased flexibility in rA16. Exchange parameters are shown in Supplementary Fig. 4b. (b) Inter-atomic distances (in Å) with unfavorable steric contacts in pink when rotating the purine base 180° around the glycoside bond in WC bps derived from idealized B-DNA and A-RNA duplexes (Methods) to adopt a *syn* conformation. Shown below are corresponding distance distributions in WC bps derived from X-ray structures of A-RNA (total 146) and B-DNA (total 159) duplexes before (solid line) and following (dashed line) 180° rotation of the purine base. The inter-atomic cut-off distance (grey line) was defined based on the van der Waals radii. (c) Relative interaction energy versus  $\chi$ -angle from biased MD trajectories of A<sub>6</sub>-DNA (blue) and hp-A<sub>6</sub>-RNA (violet). (d) Simulation time (ns) versus the global RMSD (Methods) for single A<sub>6</sub>-DNA<sup>m1A</sup> and hp-A<sub>6</sub>-RNA<sup>m1A</sup> trajectories depicting the destabilization of the RNA strand within the time of the simulation.



#### Figure 5.

Different propensities to form HG bps in B-DNA and A-RNA enable contrasting roles at the genome and transcriptome level. (**a**) In DNA, m<sup>1</sup>dA or m<sup>1</sup>dG damage is absorbed as HG bps that can be recognized by repair enzymes (in red). Had B-DNA lacked the ability to form HG bps, damage could result in duplex melting and genomic instability. In RNA, post-transcriptional modifications resulting in m<sup>1</sup>rA and m<sup>1</sup>rG block both WC and HG pairing, melting or modulating RNA secondary structure to favor functional states or effect epigenetic regulation. Had A-RNA had the ability to form HG, the m<sup>1</sup>rA and m<sup>1</sup>rG would form HG bps and potentially fail to more significantly alter RNA structure and function. (**b**) Highly conserved m<sup>1</sup>rA9 in human mitochondrial tRNA<sup>Lys</sup> blocks rA–rU WC base pairing and stabilizes native tRNA structure in which m<sup>1</sup>rA9 is in a single strand<sup>58</sup>. The m<sup>1</sup>rA9 modification would not stabilize native tRNA structure if it were simply absorbed as a HG bp. (**c**) Highly conserved m<sup>1</sup>rG37 next to the anti-codon loop<sup>37</sup> blocks base pairing between m<sup>1</sup>rG37 and the first rC in the codon and prevents +1 frameshifting in tRNA<sup>Pro</sup>, which could occur if m<sup>1</sup>rG37 formed stable HG bp with rC. (**d**) Proposed mechanism for m<sup>1</sup>rA enhanced translation through destabilization of secondary structure in the 5<sup>′</sup> UTR of mRNA.