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An Alternating Sheared AA Pair and Elements of Stability for a Single Sheared Purine-Purine Pair Flanked by Sheared GA Pairst,,‡

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Abstract

A previous NMR structure of the duplex $\frac{331}{PCCG}$ $\frac{331}{AAG}$ $\frac{331}{CCG5}$, revealed an unusually stable RNA internal loop with three consecutive sheared GA pairs. Here, we report NMR studies of two

duplexes, ${\rm ^{5'}GGU}\over{\rm PCCG}$ ${\rm ^{GGA}\over AAG}$ ${\rm ^{GGCU}}\over{\rm ^{CCG5'}}$ (replacing a UG with a UA closing pair) and

5'GGU GAA GGCU

 \overline{PCCG} \overline{AAG} $\overline{CCG5}$, (replacing the middle GA with an AA pair). An unusually stable loop with

three consecutive sheared GA pairs forms in the duplex $\frac{5'GGU}{PCCG}$ $\frac{GGA}{AAG}$ $\frac{GGCU}{CCG5'}$. The structure contrasts with that reported for this loop in the crystal structure of the large ribosomal subunit of Deinococcus radiodurans [Harms, J., Schluenzen, F., Zarivach, R., Bashan, A., Gat, S., Agmon, I., Bartels, H., Franceschi, F., and Yonath, A. (2001) Cell 107, 679-688]. The middle AA pair in

the duplex ${\rm \,^{5'}GGU} \over \rm \,PCCG$ ${\rm \,GAA} \over \rm \,GCG5},$ rapidly exchanges orientations resulting in alternative base

stacking and pseudo-symmetry with exclusively sheared pairs. The $\frac{U}{G}$ $\frac{GAA}{AAG}$ $\frac{G}{C}$ internal loop is

2.1 kcal/mol less stable than the $_G^U$ $_{AAG}^{GAG}$ $_C^G$ internal loop at 37 °C. Structural, energetic, and dynamic consequences upon functional group substitutions within related 3×3 and 3×6 internal loops are also reported.

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Tables listing chemical shift assignments, tables of NMR distance restraints, and figures of TOCSY, one-dimensional proton spectra (9-14.5 ppm), and 2D NOESY spectrum at 1 M NaCl are available. This material is available free of charge via the Internet at http:// pubs.acs.org.

Non-canonical pairs within the internal loops of RNA are important elements for folding and function. Understanding the sequence dependent folding free energy and dynamics of internal loops can facilitate prediction of structure (1, 2), dynamics, and functional significance from sequence.

AA and GA can form isosteric sheared-type (*trans* Hoogsteen/sugar edge A–A or A–G) non-canonical pairs (Figure 1a) (3–9). Typically, the AA pair is thermodynamically destabilizing but the GA pair is stabilizing (7, 10–13). Depending on sequence context, GA often forms a sheared pair, but AA is more flexible (Figure 1). Two A's can potentially switch base pairing orientation in a sheared AA pair (i.e. *trans* Hoogsteen/sugar edge A1-A2 or A2-A1) without loss of base–base hydrogen bonding. In a sheared GA pair, the equivalent interchange of bases would result in the loss of the two hydrogen bonds between G and A in a sheared GA pair.

The duplex, ${\rm {^{5'}GGU}\over PCCG}$ ${\rm {\overset{GGA}{GGCU}}\over AAG}$ ${\rm {\overset{GGCU}{CCG5'}}}$ (P^1 = purine riboside), contains an unusually stable and

relatively abundant internal loop, $\frac{GGA}{AAG}$ (9). The NMR structure of this duplex reveals three consecutive sheared GA pairs (*trans* Hoogsteen/sugar edge A–G) with separate stacks of three G's (G4/G5/G14 in the major groove) and three A's (A6/A15/A16 in the minor groove), which are closed by wobble UG (*cis* Watson-Crick/Watson-Crick U–G) and Watson-Crick CG pairs (9). (Throughout the paper, each top strand is written from 5' to 3' in going from left to right. Numbering starts at the left most (5') nucleotide of the top strand and ends at the left most (3') nucleotide of the bottom strand.)

Helix 68 of the crystal structure of the large ribosomal subunit of Deinococcus radiodurans

contains a $_G^U$ $\frac{GAA}{AAG}$ $\frac{G}{C}$ loop that has only one sheared GA pair (shown in bold) (14). There is less hydrogen bonding and the base stacking pattern is equivalent to A6/G5/A16 in the minor groove instead of the A6/A15/A16 found in the NMR structure for the equivalent loop with a UG rather than UA closing pair.

Here, we report NMR and thermodynamic studies of $\frac{GGU}{PCCG}$ $\frac{GGA}{AAG}$ $\frac{GGCU}{CCG}$ (A17 duplex)

and $_{\rm PCCG}^{\rm GGU}$ $_{\rm AAG}^{\rm GGCU}$ (A5 duplex) to determine the effects of replacing a UG closing

pair with UA or a middle GA pair with AA, respectively, relative to $\frac{\text{GGU}}{\text{PCCG}} \frac{\text{GGA}}{\text{AAG}} \frac{\text{GGCU}}{\text{CCG}}$ (3GA duplex) (Figure 2). NMR restrained molecular dynamics reveals a conformation of

three consecutive sheared GA pairs for the loop in $\frac{GGU}{PCCA}$ $\frac{GGA}{AAG}$ $\frac{GGCU}{CCG}$. A5 and A15 in

 $\begin{array}{c} {\rm GGU} \ \ \overline{\rm GAA} \ \ \overline{\rm GGCU} \\ {\rm PCCG} \ \ \overline{\rm AAG} \ \ \overline{\rm CCG} \end{array} \ \ {\rm rapidly\ exchange\ positions\ forming\ alternative\ sheared\ AA\ pairs\ (i.e.\ exchanging\ between\ \it trans\ Hoogsteen/sugar\ edge\ A15-A5\ and\ Langle A15-A5$

¹Abbreviations: a, deoxyadenosine; C_T , total concentration of oligonucleotide strands; D, 2,6-diaminopurine riboside; g, deoxyguanosine; I, inosine; M, 2'-O-methyl adenosine; P, purine riboside; R, any purine nucleotide; T_M , melting temperature in kelvin; T_m , melting temperature in degrees Celsius.

A5-A15) flanked by sheared GA pairs. The exchanging AA pair results in alternative base stacking of A6/A15/A16 or A6/A5/A16 in the minor groove. The flexibility of alternative orientations of a middle adenine base edge in the minor groove, i.e. from A15 (N3-C2-N1)

(as observed in $\frac{GGU}{PCCG}$ $\frac{GGA}{AAG}$ $\frac{GGCU}{CCG}$ and $\frac{GGU}{PCCA}$ $\frac{GGA}{AAG}$ $\frac{GGCU}{CCG}$) to A5 (N1-C2-N3), might provide switching between different binding partners for dynamic functions.

Functional group substitutions (atomic mutations) have been extensively used for studying elements of molecular recognition in RNA (10, 15–24). Here, the structural, energetic, and dynamic consequences of functional group substitutions are explored by studying duplexes

GGU GRA GGCU of the form $_{PCCG}$ $_{\overline{AQG}}$ $_{CCG}$, where R and Q are various purine nucleotides (Figures 1a and 2). Single predominant conformations form in the a5, P5, I5, and I15 duplexes. Functional group substitutions also facilitate interpretation of NMR data. The thermodynamic effects of functional group substitutions within 3×6 internal loops are also reported.

MATERIALS AND METHODS

Oligonucleotide Synthesis and Purification

Oligonucleotides were synthesized using the phosphoramidite method (25, 26) and purified as described previously (9, 12). CPG supports and phosphoramidites were acquired from Proligo, Glen Research, or ChemGenes. The mass of all oligonucleotides was verified by ESI MS with a Hewlett Packard 1100 LC/MS Chemstation. Purities were checked by reverse phase HPLC or analytical TLC on a Baker Si500F silica gel plate (250 µm thick) and all were greater than 95% pure.

UV Melting Experiments and Thermodynamics

Concentrations of single-stranded oligonucleotides were calculated from the absorbance at 280 nm at 80 °C and extinction coefficients predicted from those of dinucleotide monophosphates and nucleosides (27, 28) with the RNAcalc program (http://www.meltwin.com) (29). The extinction coefficients were estimated by replacing purine riboside, 2,6-diaminopurine riboside, deoxyadenosine, and 2'-O-methyl adenosine with adenosine; and replacing inosine and deoxyguanosine with guanosine. Although extinction coefficients differ upon functional group substitutions, individual nucleotides contribute only a small portion of the oligomer extinction and thus do not significantly affect thermodynamic measurements. UV melting buffer conditions were 1.0 M NaCl, 20 mM sodium cacodylate, and 0.5 mM disodium EDTA at pH 7 or 80 mM NaCl, 10 mM sodium phosphate, 0.5 mM disodium EDTA, pH 7. Curves of absorbance at 280 nm versus temperature were acquired using a heating rate of 1 °C/min with a Beckman Coulter DU640C spectrophotometer having a Peltier temperature controller.

Melting curves were fit to a two-state model with the MeltWin program (http://www.meltwin.com), assuming linear sloping baselines and temperature-independent H° and S° (29–31). Additionally, the temperature at which half the strands are in duplex, T_{M} ,

at total strand concentration, C_T, was used to calculate thermodynamic parameters for non-self-complementary duplexes according to (32):

$$T_{M}^{-1} = (R/\Delta H^{\circ}) \ln(C_{T}/4) + (\Delta S^{\circ}/\Delta H^{\circ}) \quad (1)$$

Here R is the gas constant, 1.987 cal/mol·K. All of the H° values from T_{M}^{-1} versus $\ln(C_{T}/4)$ plots and from the average of the fits of melting curves to two-state transitions agree within 15%, suggesting that the two-state model is a reasonable approximation for these transitions. The equation $G^{\circ}_{37} = H^{\circ} - (310.15)$ S° was used to calculate the free energy change at 37 °C (310.15 K).

NMR Sample Preparation

With minor modification, sample preparation was similar to that previously reported (7, 9). The sample buffer conditions were 80 mM NaCl, 10 mM sodium phosphate, 0.5 mM

disodium EDTA, pH 5.1 for H_2O , pD 7.3 for D_2O for $\frac{GGU}{PCCG}$ $\frac{GAA}{AAG}$ $\frac{GGCU}{CCG}$ and pH 5.4 for

 $\rm H_2O,\,pD$ 6.8 for $\rm D_2O$ for ${\rm GGU\over PCCG}$ ${\rm \overline{GGA}\over AAG}$ ${\rm GGCU\over CCG}$. Exchangeable proton spectra at pH 6.0 for

GGU GAA GGCU were very similar to those at pH 5.1. Moreover, chemical shifts and critical loop NOEs involving non-exchangeable protons were essentially the same in water at pH 5.1 and pD 7.3. Total volumes were 300 μ L with 90:10 (v:v) H₂O:D₂O for exchangeable proton spectra and 99.996% D₂O (Cambridge Isotope Laboratories) for non-exchangeable spectra. The total duplex concentrations were ~2 mM. The total duplex concentrations of other sequences were 0.5–1.2 mM.

NMR Spectroscopy

Unless otherwise noted, all exchangeable and non-exchangeable proton spectra were acquired on a Varian Inova 500 MHz (¹H) spectrometer (33). One-dimensional imino proton spectra were acquired with an S pulse sequence (33) with a sweep width of 12 kHz and temperatures ranging from 0 to 55 °C. SNOESY spectra were recorded with a 150 ms mixing time at 5 and 30 °C. NOESY spectra of samples in D2O were acquired at 30 °C with 100, 200, and 400 ms mixing times. TOCSY spectra were acquired at 30 °C with 8, 20, and

40 ms mixing times. Natural abundance $^1H^{-13}C$ HMQC specta for $^{\mbox{GGU}}_{\mbox{PCCG}}$ $^{\mbox{GAA}}_{\mbox{AAG}}$ $^{\mbox{GGCU}}_{\mbox{CCG}}$

and $_{PCCG}$ $_{\overline{AAG}}$ $_{CCG}$ were acquired with a 5000 Hz spectral width for proton and 15000 Hz spectral width for carbon. The $^{1}H^{-31}P$ HETCOR and natural abundance $^{1}H^{-13}C$ HSQC spectra were acquired on a Varian Inova 600 MHz (^{1}H) spectrometer. The 1D $^{1}H^{-13}C$ decoupled ^{31}P spectra (referenced to external standard of 85% H $_{3}PO_{4}$ at 0 ppm) were acquired on a Bruker Avance 500 MHz (^{1}H) spectrometer at 30 °C. Proton spectra were referenced to H $_{2}O$ or HDO at a known temperature dependent chemical shift relative to 3-(trimethylsilyl) tetradeutero sodium propionate (TSP). The Felix (2000) software package (Molecular Simulations Inc.) was used to process 2D spectra.

Restraint Generation

Very similar restraints were generated for $\frac{GGU}{PCCG}$ $\frac{GGA}{AAG}$ $\frac{GGCU}{CCG}$ (Supporting Information

Table S1) as for $\frac{GGU}{PCCG}$ $\frac{GGA}{AAG}$ $\frac{GGCU}{CCG}$ (9). For $\frac{GGU}{PCCG}$ $\frac{GAA}{AAG}$ $\frac{GGCU}{CCG}$ and

 $\begin{array}{ccc} \rm GGU & \rm GGA & \rm GGCU \\ \rm PCCG & \overline{\rm AAG} & \rm CCG \end{array}, 15 \ hydrogen \ bond \ restraints \ limiting \ proton \ and \ hydrogen-bond \end{array}$ acceptor distances within 1.8 to 2.5 Å were applied for the five Watson-Crick GC pairs, but no hydrogen bond restraints were used within the loop and UG or UA pair. Dihedral angles of residues in the Watson-Crick stems and UG or UA pair were loosely restrained: α $(0\pm120^{\circ})$, β $(180\pm30^{\circ})$, γ $(60\pm30^{\circ})$, δ $(85\pm30^{\circ})$, ϵ $(-140\pm40^{\circ})$, ζ $(0\pm120^{\circ})$ (ζ was mistakenly given as ξ in ref (9).), and χ (-170±40°). For loop residues, glycosidic bond dihedral angles, χ 's, were loosely restrained (-120±90°) because there was no indication of a syn glycosidic

conformation. For the structural modeling of $_{PCCG}$ $\stackrel{GGU}{\overline{AAG}}$ $\stackrel{GGCU}{CCG}$, the δ dihedral angle for G5 was restrained to be C2'-endo with δ (160±30°), and for A6, G14, U10, and P20, the δ dihedral angles were restrained to cover both C2'-endo and C3'-endo conformations with δ $(122.5\pm67.5^{\circ}).$

Two sets of distance and dihedral angle restraints (set I: A6/A15/A16 and set II: A6/A5/

A16) were run for $\frac{GGU}{PCCG}$ $\frac{GAA}{AAG}$ $\frac{GGCU}{CCG}$ because NOEs were inconsistent with a single

structure. The previous NMR structure of $\frac{GGU}{PCCG}$ $\frac{GGA}{AAG}$ $\frac{GGCU}{CCG}$ (9) facilitated segregation of restraints for structural modeling. For inter-proton distance restraints that differ for A6/A15/A16 and A6/A5/A16 structural modeling, lower and upper bounds were loosened; all other restraints are the same for sets I and II (Supporting Information Table S2).

Two sets of δ dihedral angle restraints were generated for loop residues (G4, A5, A6, G14,

A15, and A16) in $\frac{GGU}{PCCG}$ $\frac{GAA}{AAG}$ $\frac{GGCU}{CCG}$. For set I (A6/A15/A16), the δ dihedral angle for A5 was restrained to be C2'-endo with δ (160±30°), all other loop residues and the two 3'dangling residues, U10 and P20, were restrained to cover both C2'-endo and C3'-endo conformations with δ (122.5±67.5°). For set II (A6/A5/A16), all the loop residues were restrained to cover both C2'-endo and C3'-endo conformations, with the other dihedral angle

restraints the same as those of set I (A6/A15/A16) and ${\rm GGU\over PCCG}$ ${\rm \overline{GGA}\over AAG}$ ${\rm CCG}$.

In summary, a total of 222 distance restraints (110 intra-nucleotide, 112 inter-nucleotide) including hydrogen bond restraints, and 98 dihedral angle restraints were used for the

structural modeling of $\frac{GGU}{PCCG}$ $\frac{GGA}{AAG}$ $\frac{GGCU}{CCG}$ (Supporting Information Table S1). For the

 $\begin{array}{c} {\rm GGU} \ \, {\rm GAA} \ \, {\rm GGCU} \\ {\rm structural\ modeling\ of\ }_{\rm PCCG} \ \, {\rm \overline{AAG}} \ \, {\rm CCG} \end{array} \ \, , a\ \, {\rm total\ of\ } 250\ \, {\rm distance\ restraints} \ \, (128\ intranucleotide,\ 122\ inter-nucleotide)\ \, including\ \, hydrogen\ \, bond\ \, restraints,\ \, and\ \, 98\ \, dihedral\ \, angle \end{array}$ restraints were used for the structural modeling of set I (A6/A15/A16), and a total of 249 distance restraints (128 intra-nucleotide, 121 inter-nucleotide) including hydrogen bond

restraints, and 98 dihedral angle restraints were used for the structural modeling of set II (A6/A5/A16) (Supporting Information Table S2).

Structural Modeling

NMR restrained molecular dynamics and energy minimization was done with the Discover 98 package on a Silicon Graphics computer. An A-form like RNA starting structure was generated with the Biopolymer module of Insight II (2000). The AMBER 95 force field (34) was used with addition of flat-bottom restraint pseudo-potentials, with force constants of 25 kcal/(mol•Ų) for NOE distance restraints and 50 kcal/(mol•rad²) for torsion angle restraints and with a maximum force of 1000 kcal/mol. Group-based summation with an 18 Å cutoff was used for calculating van der Waals interactions. The cell-multipole method (35), with distance dependent dielectric constant ($\varepsilon = 2r$), was used for calculating electrostatic interactions. The progression of the structure simulation was the same as previously reported (7, 9, 17). Several figures were generated with the PyMOL program (36).

RESULTS

Functional Group Substitutions and Thermodynamics of Molecular Recognition

Measured thermodynamic parameters for several duplexes and internal loops with and without functional group substitutions are listed in Tables 1 and 2, respectively. Most were measured at 1 M NaCl to allow comparison to existing databases, but four were also measured in the 80 mM NaCl buffer used for most NMR experiments. The lower salt makes duplex formation less favorable on average by 3.41 ± 0.02 kcal/mol at 37 °C, which is consistent with a sequence independent salt effect. Measured thermodynamic parameters for formation of the internal loops (Table 2) are calculated according to the following equation which relies on the nearest neighbor model for predicting duplex stability (37):

$$\Delta G^{^{\circ}}_{37,loop} = \Delta G^{^{\circ}}_{37(duplex\ with\ loop)} - \Delta G^{^{\circ}}_{37(duplex\ without\ loop)} + \Delta G^{^{\circ}}_{37(interrupted\ base\ stake)} \quad \text{(2a)}$$

For example,

$$\Delta G^{\circ}_{37} \stackrel{U}{G} \stackrel{GGA}{AMG} \stackrel{G}{C} = \Delta G^{\circ}_{37} \stackrel{GGU}{PCCG} \stackrel{GGA}{AMG} \stackrel{GGCU}{CCG} - \Delta G^{\circ}_{37} \stackrel{GGU}{PCCG} \stackrel{GGCU}{CCG} + \Delta G^{\circ}_{37} \stackrel{UG}{GC} \stackrel{(2b)}{CCG}$$

Here, ΔG°_{37} $\frac{GGU}{PCCG}$ $\frac{GGA}{AMG}$ $\frac{GGCU}{CCG}$ is the measured value of the duplex containing the internal loop (Table 1), ΔG°_{37} $\frac{GGU}{PCCG}$ $\frac{GGCU}{CCG}$ is the measured value of the duplex without

the loop (9), and $^{\Delta}G^{^{\circ}}_{37}$ $^{UG}_{GC}$ is the free energy increment for the nearest neighbor base stack interaction interrupted by the internal loop (1, 31). $^{\circ}H^{\circ}_{loop}$ and $^{\circ}S^{\circ}_{loop}$ are calculated similarly. All the thermodynamic parameters used in this calculation are derived from T_{M}^{-1} versus $ln(C_{T}/4)$ plots (eq 1). When eq 2a was applied to 2×2 nucleotide internal loops of non-canonical pairs flanked by different stems, the values for $^{\circ}G^{\circ}_{37,\ loop}$ for a given loop sequence differed by an average of 0.40 kcal/mol (2). The model should be even better for sequences with identical stems.

Functional Group Substitutions, NMR Assignments, and Structural Features

The base – (H1'/H5) "NOESY walk" regions of the 400 ms NOESY spectra at 30 °C are shown in Figure 3. NMR resonances were assigned essentially as described previously (7, 9,

38, 39). Comparison of spectra with those of the duplex $\frac{GGU}{PCCG}$ $\frac{GGA}{AAG}$ $\frac{GGCU}{CCG}$ (9)

 $\begin{array}{c} GGU \\ facilitates \\ NMR \\ assignments \\ of \\ \begin{array}{c} PCCG \\ \hline AAG \\ \\ \end{array} \\ \begin{array}{c} GGU \\ \hline AAG \\ \hline AAG \\ \end{array} \\ \begin{array}{c} GGU \\ \hline AAG \\ \hline AAG \\ \end{array} \\ \begin{array}{c} GGU \\ \hline AAG \\ \hline AAG \\ \end{array} \\ \begin{array}{c} GGU \\ \hline AAG \\ \end{array} \\ \begin{array}{c} GGU \\ \hline AAG \\ \hline A$

structural modeling). U3A17 in PCCG AAG CCG forms a Watson-Crick pair as indicated by a strong NOE between U3H3 and A17H2. The imino proton of U3H3 is relatively broad and shifted upfield (12.53 ppm) (13) relative to the usual range of 13 to 15 ppm for a Watson-Crick UA pair. A similar upfield shift (11.75 ppm) was observed in the 2

 $\times\,2\,loop\, \frac{U}{A}\,\,\frac{GA}{AG}\,\,\frac{A}{U}$ (40). Three consecutive sheared GA pairs form in

medium to strong NOEs, which are similar to NOEs observed for $\frac{GGU}{PCCG}$ $\frac{GGA}{AAG}$ $\frac{GGCU}{CCG}$, define the loop structure, e.g. G14H2'–G5H1, A6H1'-A15H2, A15H1'-A6H2, A16H1'-A15H2, G7H1'–A6H2, and A17H1'–A16H2 (compare to G17H1'–A16H2 in

GGU GGA GGCU (Figure 3a and Table 3) (9). In the loop of A $\frac{U}{AAG}$ GGA and G5 have C3'-endo and C2'-endo sugar puckers, respectively, and G14 is populated in both conformations as evidenced by TOCSY (Supporting Information Figure S1a) and NOESY

(Figure 3a) spectra. The same sugar puckers are found in ${{\rm GGU}\over{\rm PCCG}}$ ${{\rm GGA}\over{\rm AAG}}$ ${{\rm GGCU}\over{\rm CCG}}$ (9).

Similarity of sugar edges (6) of adenosine, purine riboside, and inosine (Figure 1) facilitates H2 assignments of middle purine-purine pairs. Also, the NOEs of G14H1'/H2'-A5H2 in

 $\frac{GGU}{PCCG}$ $\frac{GAA}{AAG}$ $\frac{GGCU}{CCG}$ (Figure 3b) are similar to those of G14H1'/H2'–G5H1 as present for

GGU GGA GGCU GGU GGU GGCU (7) and PCCG AAG CCG (7) and PCCG AAG CCG (8) and PCCG AAG CCG (7) and PCCG AAG CCG (8) and PCCG AAG CCG (8)

 $\begin{array}{c} {\rm GGU} \ \, \frac{{\rm GIA}}{{\rm AAG}} \ \, \frac{{\rm GGCU}}{{\rm CCG}} \ \, \text{(Figure 3e)}. \ \, \text{The chemical shift of I5H1, 12.1 ppm (Supporting Information Figure S2)} \ \, \text{is in agreement with the formation of a sheared IA pair (41)}. \\ {\rm Downfield \ \, chemical \ \, shifts \ \, for inosine \ \, imino \ \, protons \ \, beyond 14 ppm \ \, were \ \, observed \ \, in face-to-face IA pairs (4, 42).} \end{array}$

While comparison of NOESY spectra (Figure 3, Table 3, and Supporting Information Table S2) indicates very similar base pairing and stacking geometries with three sheared-type

 $purine-purine\ pairs\ for\ \frac{\rm GGU}{\rm PCCG}\ \frac{\rm GaA}{\rm AAG}\ \frac{\rm GGCU}{\rm CCG} \ (a5\ duplex), \\ \frac{\rm GGU}{\rm PCCG}\ \frac{\rm GPA}{\rm AAG}\ \frac{\rm GGCU}{\rm CCG} \ (P5)$

duplex), and ${\rm \frac{GGU}{PCCG}}~{\rm \frac{GIA}{AAG}}~{\rm \frac{GGCU}{CCG}}$ (I5 duplex), the NMR spectra provide evidence for two

populations of structures for the middle A5A15 pair in ${}_{PCCG}$ $\overline{}_{AAG}$ $\overline{}_{CCG}$ (A5 duplex) (Figure 6). For example, in addition to NOEs of R5H2-G14H1'/H2' (R is any purine), A6H2-A15H1', A15H2-A6H1', and A15H2-A16H1' (shown in yellow circles) for the a5, P5, and I5 duplexes, one extra set of NOEs A15H2- G4H1'/H2', A16H2-A5H1', A5H2-

A16H1', and A5H2- A6H1' (shown in green circles) are present for GGU GAA GGCU (Figure 3). Moreover, on the basis of TOCSY (Supporting Information Figure S1) and NOESY (Figure 3) spectra, sugar puckers of a5, P5, and I5 are C2'-*endo* as indicated by strong H1'–H2' couplings, which corresponds to the C2'-*endo* conformation of the G5 sugar

pucker in $\frac{GGU}{PCCG}$ $\frac{GGA}{AAG}$ $\frac{GGCU}{CCG}$ (9). Sugar puckers for A15 are C3'-*endo* in each of these

duplexes with the possible exception of the a5 duplex. For $\frac{GGU}{PCCG}$ $\frac{GAA}{AAG}$ $\frac{GGCU}{CCG}$, however, A5 and A15 are populated in both C2'-*endo* and C3'-*endo* conformations with A5 having higher C2'-*endo* population than A15. Evidently, the middle A5A15 pair is more populated in *trans* Hoogsteen/sugar edge A15-A5 than in *trans* Hoogsteen/sugar edge A5-A15 and related conformations. This is in agreement with relatively stronger NOEs (Figure 3b) observed for A5H2-G14H1'/H2', A6H2- A15H1', A15H2- A6H1', and A15H2- A16H1' (shown in yellow circles) than A15H2- G4H1'/H2', A16H2- A5H1', A5H2- A16H1', and

A5H2-A6H1' (shown in green circles), respectively, in $\frac{GGU}{PCCG}$ $\frac{GAA}{AAG}$ $\frac{GGCU}{CCG}$. Presence of a

single set of chemical shifts for $\frac{GGU}{PCCG}$ $\frac{GAA}{AAG}$ $\frac{GGCU}{CCG}$ indicates that the middle A5A15 pair is alternating rapidly (fast exchange on the NMR time scale), with the sugar edge of either A5 or A15 on the base pairing edge of the other, forming *trans* Hoogsteen/sugar edge A15-A5 (shown in yellow) or A5-A15 (shown in green) and related pairs (Figure 5).

NMR spectra acquired at 1 M NaCl indicate that the structural and dynamical properties of the A5 loop are similar at 80 mM and 1 M NaCl (Figure 3b and S3, respectively). Chemical shifts show only modest salt dependence and the same pattern of NOEs for the major and minor conformations are observed, including the A5H2-A15H8 (major) and A15H2-A5H8 (minor) cross peaks (data not shown).

Relatively downfield chemical shifts of H2 protons on the base pairing edge (Figure 1) are observed for P5H2, 9.30 ppm (compared to P20H2, 8.16 ppm (9)), a5H2, 8.44 ppm (compared to A15H2, 7.57 ppm in the a5 duplex), and I5H2, 8.69 ppm, in

GGU GPA GGCU GGU GaA GGCU GGU GGU GGU GGU GGU GGU GGU PCCG \overline{AAG} CCG , \overline{AAG} CCG , \overline{AAG} CCG , \overline{AAG} CCG , respectively (Figure 3 and Table 4). Such downfield chemical shifts of H2 on the base pairing edge of sheared purine-purine pairs are expected due to ring current de-shielding effects (Figure 1a),

as observed previously in 2 \times 2 loops: A4H2, 8.19 ppm in $\frac{GGU}{UCCG}$ $\frac{AA}{AA}$ $\frac{GGCU}{CCG}$ (compared to 7.88 ppm for A5H2, which is not on a base pairing edge) and P4H2, 8.97 ppm in

 $\begin{array}{cc} \operatorname{GGU} & \operatorname{\underline{PA}} & \operatorname{GGCU} \\ \operatorname{UCCG} & \operatorname{\underline{AP}} & \operatorname{CCG} \end{array} (18).$

On the edge that is not base paired, relatively upfield chemical shifts of H2 protons are

observed: A15H2 chemical shifts are 7.69, 7.57, and 7.55 ppm in $\frac{GGU}{PCCG}$ $\frac{GPA}{AAG}$ $\frac{GGCU}{CCG}$,

for A15H2 protons in $\frac{GGU}{PCCG}$ $\frac{GGA}{AAG}$ $\frac{GGCU}{CCG}$ and $\frac{GGU}{AAG}$ $\frac{GGCU}{CCG}$, respectively, which might reflect stronger base pairing and better stacking of the motif with three consecutive sheared GA pairs, resulting in larger ring current shielding effects from A6 and A16. Intermediate chemical shifts of A5H2 (8.28 ppm) and A15H2, (7.83 ppm) are observed

in $\frac{GGU}{PCCG}$ $\frac{GAA}{AAG}$ $\frac{GGCU}{CCG}$, which is consistent with a rapidly alternating sheared A5A15 pair.

The I15 duplex provides further evidence for two structures of the A5 duplex. Several features in the NMR spectra suggest that the loop conformation in the I15 duplex is most similar to the less populated conformation observed in the A5 duplex. NOEs observed in the I15 duplex that were weak in A5 duplex, and not observed at all in a5, P5, or I5 duplexes include 15H2-G4H1', A16H2–5H1', 5H2-A16H1', 5H2-A6H1', and 15H2–5H8 (Figure 3).

NOEs not observed in I15 duplex, but observed in PCCG AAG GCCU and all other duplexes include 5H2-G14H1', A6H2–15H1', 15H2-A6H1', and 15H2-A16H1'. The relatively downfield chemical shift of I15H2 (8.55 ppm) and the relatively upfield shift of A5H2 (7.51 ppm) are consistent with those protons being at the A5-I15 base-pairing edge and out in the minor groove, respectively, as discussed for the other duplexes. A strong scalar coupling, I15H1'-H2', and a large downfield ³¹P shift of A16 (2.75 ppm) indicate a C2'-endo ribose conformation at I15. In contrast, A5H1'-H2' scalar coupling is much smaller and the ³¹P shift of A6 is 1.10 ppm. These contrast with a5, P5, and I5 duplexes which show strong 5H1'-H2' coupling and weak 15H1'-H2' coupling. Moreover, ³¹P shifts of I5 duplex are 0.86 and 2.58 ppm for A16 and A6, respectively. Additionally, the G4H1'-H2' and G14H1'-H2' couplings are moderate and zero, respectively, in the I15 duplex while they are zero and moderate, respectively, in a5, P5, and I5 duplexes. The moderate couplings probably indicate dynamic interconversion of sugar puckers. The ³¹P chemical shifts for the A5 duplex are 1.41 and 2.20 ppm for A16 and A6, which suggest that the A5 duplex is more populated in a conformation similar to a5, P5, and I5 duplexes.

There is a large chemical shift difference between the H2 resonances for the central purine-purine paris, and the linewidths of these resonances are consistent with two rapidly interconverting structures for the A5 duplex (Table 4). The linewidths for A5H2 and A15H2 are both about 14 Hz for the A5 duplex. In contrast, the H2 resonance not at the base pairing edge in a5, P5, I5, and I15 duplexes has an average linewidth of 4.75 ± 1 Hz, which is approximately the same as the average of the stem resonance G2H8 (4.1 ± 0.5 Hz). This is consistent with a model in which duplexes having a single conformation have a relatively narrow linewidth, while the A5 duplex resonances are broadened due to switching between two conformations. The base pairing edge H2 resonance in a5, P5, I5, and I15 duplexes has average linewidth of 10.2 ± 2.3 Hz. This may indicate that this residue is slightly less stable

than the pairing partner on the other strand, and/or that the chemical shift of the proton in this position is more sensitive to slight structural fluctuations.

Structural statistics for $\frac{GGU}{PCCA}$ $\frac{GGA}{AAG}$ $\frac{GGCU}{CCG}$ and $\frac{GGU}{PCCG}$ $\frac{GAA}{AAG}$ $\frac{GGCU}{CCG}$

A total of 24 of 40 modeled structures (PDB: 2DD1) were selected for analysis of

 $\frac{GGU}{PCCA}$ $\frac{GGA}{AAG}$ $\frac{GGCU}{CCG}$. The average root-mean-square deviation of all selected structures to the average structure for all atoms is 0.80 ± 0.15 Å. No distance or dihedral angle restraint violations were greater than 0.2 Å or 2°, respectively. The average of the final energies at 300 K from the force field is -428.0 ± 4.9 kcal/mol.

A total of 27 of 40 modeled structures (PDB: 2DD2) were selected for analysis of the

A6/A15/A16 structure of $\frac{GGU}{PCCG}$ $\frac{GAA}{AAG}$ $\frac{GGCU}{CCG}$. The average root-mean-square deviation of all selected structures to the average structure for all atoms is 0.69 \pm 0.21 Å. Two distance and one dihedral angle restraint violations were greater than 0.2 Å and 2°, respectively. The average of the final energies at 300 K from the force field is -430.6 ± 5.7 kcal/mol.

A total of 18 of 40 modeled structures (PDB: 2DD3) were selected for analysis of the

A6/A5/A16 structure of $\frac{GGU}{PCCG}$ $\frac{GAA}{AAG}$ $\frac{GGCU}{CCG}$. The average root-mean-square deviation of all selected structures to the average structure for all atoms is 0.97 \pm 0.25 Å. One distance and no dihedral angle restraint violations were greater than 0.2 Å and 2°, respectively. This minor conformation is less convergent than the other structures due to loosened restraints in the loop region. The average of the final energies at 300 K from the force field is -423.4 ± 6.3 kcal/mol.

Other known AA geometries are not consistent with the NMR data

As illustrated in Figure 1b, several non-sheared AA pairs have been observed in crystal and NMR structures (6, 43–49). These potential structures can be compared to the NMR spectra

$$\label{eq:formula} \text{for} \frac{\text{GGU}}{\text{PCCG}} \; \frac{\text{GAA}}{\text{AAG}} \; \frac{\text{GGCU}}{\text{CCG}}.$$

A potential A-zipper motif, as seen in a DNA internal loop $\frac{C}{AAG}$ $\frac{GAA}{C}$ and NMR structure

of an RNA tetraloop receptor $_G^C$ $\frac{UAA}{UA}$ $_C^G$ (50, 51), which would place A5 and A15 in

PCCG AAG CCG stacking on each other, is ruled out because no NOE of A5H2 –

A15H1' or A15H2–A5H1' is observed in $\frac{\mathrm{GGU}}{\mathrm{PCCG}}$ $\frac{\mathrm{GAA}}{\mathrm{AAG}}$ $\frac{\mathrm{GGCU}}{\mathrm{CCG}}$.

Both AH2 protons are exposed in the minor groove for a *trans* Watson-Crick/Hoogsteen A–A pair (Figure 1b) (6, 43). This conformation is ruled out for the middle AA pair in

 $\begin{array}{ccc} U & GAA & G\\ G & \overline{AAG} & C \end{array} \ because it would not give the observed cross-strand G4H1'/H2'-A15H2 \ and G14H1'/H2'-A5H2 \ cross-peaks \ (Figure 3b, Table 3, and Supporting Information Table S2). \end{array}$

The cis Watson-Crick/Watson-Crick A-A conformation (6, 45), with both AH2 protons

exposed in the minor groove, is ruled out for the middle AA in $_G^U$ $\stackrel{GAA}{\overline{AAG}}$ $\stackrel{G}{C}$ because it would not give all the observed G14H1'/H2'-5H2, A6H1'-A15H2, A15H1'-A6H2, A16H1'-A15H2 (shown in yellow circles); and G4H1'/H2'-A15H2, A16H1'-A5H2, A5H1'-A16H2, and A6H1'-A5H2 (shown in green circles) cross-peaks (Figure 3b).

A trans Hoogsteen/Hoogsteen A-A pair (Figure 1b) (6, 46-49) is ruled out for the middle

AA in $_G^U$ $\frac{GAA}{AAG}$ $\frac{G}{C}$ because there is no indication of a syn glycosidic conformation as evidenced by A5H1'-A5H8 and A15H1'-A15H8 cross-peaks and because the 14H1'/H2'-

5H2 and 4H1'/H2'–15H2 cross-peaks seen in $_G^U$ \overline{AAG} $_C^G$ are not expected for a trans Hoogsteen/Hoogsteen A–A with two AH2 protons exposed in the minor and major groove, respectively.

DISCUSSION

Understanding relationships between sequence, energetics, structure, dynamics, and function can facilitate rapid extraction of the information encoded in the constantly expanding databases of RNA sequences. The internal loop is a common RNA motif where such relationships are not fully understood (9, 12, 52-54). Detailed understanding of interactions such as hydrogen bonding and base stacking in internal loops will allow prediction of the contributions of internal loops to RNA folding and function.

Three Consecutive Sheared GA Pairs in $_{ m A}^{ m U}$ $_{ m AAG}^{ m GGA}$ $_{ m C}^{ m G}$

The previous NMR structure of ${\rm PCCG} \over {\rm PCCG} \over {\rm AAG} \over {\rm CCG}$ reveals three consecutive sheared GA pairs in the unusually stable internal loop (9). Formation of three consecutive sheared

GA pairs in $_{A}^{U}$ $_{AAG}^{GA}$ $_{C}^{G}$ (-2.27 kcal/mol) (Figure 4 and 5), similar to $_{G}^{U}$ $_{AAG}^{GA}$ $_{C}^{G}$ (-2.62 kcal/mol), is consistent with the thermodynamic stabilities (Table 2) and the occurrences of both loops in helix 41a of small subunit rRNA (52). (Throughout the paper, the values in parenthesis after the duplex are the measured free energy at 37 °C for loop formation in 1 M NaCl unless otherwise noted.)

In contrast to the NMR structures, helix 68 of the crystal structure of D. radiodurans large

subunit rRNA contains a $_{A}^{U}$ $_{AAG}^{GA}$ $_{C}^{G}$ loop that has only one sheared GA pair (shown in bold) (14). The major difference is that the corresponding G5 and A15 bases are shifted, opposite to a sheared GA pair, to the minor and major groove, respectively. This results in the loss of hydrogen bonding, and in a corresponding base stacking pattern equivalent to

> A6/G5/A16 in the minor groove, instead of the A6/A15/A16 stacking pattern found in the NMR structure (Figure 5). Several critical cross-strand NOEs define the stacking pattern of A6/A15/A16 in the NMR structure with three consecutive sheared GA pairs, e.g. A15H2-A6H1', A15H1'-A6H2, and A15H2-A16H1' (Figure 3a, Supporting Information Table S1). The distances between the protons in each pair exceed 5 Å in the crystal structure (PDB: 1NKW) when hydrogens are added (Supporting Information Table S1). Interestingly, the A6/G5/A16 stacking pattern in the crystal structure (Figure 5, top panel) is similar to the A6/A5/A16 stacking pattern determined for the minor NMR structure of

> GGU GAA GGCU $\frac{GH}{AAG}$ $\frac{GH}{CCG}$ (Figure 5, bottom panel), although fewer hydrogen bonds are formed in the crystal structure (14).

> There are several differences between the environments of the ${\rm U \over AAG} {\rm G \over C}$ loop in the crystal and in NMR buffer. The crystals were grown from ribosomal subunits in 10 mM MgCl₂, 60 mM NH₄Cl, 5 mM KCl, and 10 mM HEPES, pH 7.8 (14). The NMR buffer has 80 mM NaCl and 10 mM sodium phosphate, pD 6.8. It would be surprising, however, if Mg²⁺ shifted the local structure. The thermodynamics of the 3GA duplex (Figure 2) were essentially the same in 1 M NaCl and in 10 mM MgCl₂, 150 mM KCl (9). It is quite possible, however, that other interactions in the ribosomal subunit crystal are strong enough to break hydrogen bonds and rearrange stacking. There is no tertiary interaction with or

protein binding to the loop $_{A}^{U}$ $\underset{AAG}{\overset{GGA}{G}}$ $_{C}^{G}$ in the crystal, but the loop, $_{C}^{G}$ $\underset{GAAGU}{\overset{UCAAG}{GAAGU}}$ $_{A}^{U}$, which is

directly 5' to the UA closing pair of ${\rm GGA \over AAG}$ ${\rm G \over C}$, has tertiary interactions with helix 75 by consecutive A-minor interactions. Similar A-minor tertiary interactions are observed in the crystal structures of the large ribosomal subunits of Haloarcula marismortui and E. coli between helix 75 and helix 68 (55, 56). While long range effects may affect local structure, it may also be difficult to determine such fine details in a large crystal refined to 3.1 Å.

Tandem sheared GA pairs closed by UA Watson-Crick pairs have been reported in an NMR

structure of $\frac{GGGCU}{UCCGA}$ $\frac{GA}{AG}$ $\frac{AGCCU}{UCGGG}$ (40). The sugars of G's in $\frac{U}{G}$ $\frac{GA}{AG}$ $\frac{A}{U}$ are in C2'-endo

conformation. In the loop, A AAG C, of the A17 duplex, G4 and G5 have C3'-endo and C2'-endo sugar puckers, respectively, and G14 is populated in both conformations as evidenced by the TOCSY spectrum (Supporting Information Figure S1a). Evidently, a C2'-

endo sugar pucker for G is not required to form a sheared GA pair in a $_{A}^{U}$ $_{\overline{A}}^{G}$ motif.

Two Alternating Structures for $\frac{\mathrm{GGU}}{\mathrm{PCCG}}$ $\frac{\mathrm{GAA}}{\mathrm{AAG}}$ $\frac{\mathrm{GGCU}}{\mathrm{CCG}}$

The sheared AA pair in ${\rm PCCG} \over {\rm AAG} \over {\rm CCG}$ is rapidly exchanging between alternative conformations (Figure 5, bottom panel and Figure 6). This exchange is consistent with an intrinsically flexible AA pair with fewer hydrogen bonds than a GA pair. The pseudosymmetry of the dynamic AA pair allows an estimate of the lower limit for the exchange

rate. The a5 duplex has only one conformation and it is the same as the major conformation of the A5 duplex. The chemical shifts for a5H2 and A15H2 in this conformation are 8.44 and 7.57 ppm, respectively. With the assumption that the minor conformation of the A5 duplex would result in an A5H2 chemical shift of 7.57 ppm, the lower limit for the exchange rate is estimated as $0.87 (500) = 435 \, \mathrm{s}^{-1}$. A calculation based on linewidths (57, 58) of H2 resonances suggests an even faster exchange rate of between 20,000 and 65,000 s^{-1} (Table 4) for estimates of the major conformation ranging from 90% to 60 % of the population. Fast exchange has also been detected between *syn* and *anti* G's in a single GG pair in an RNA duplex (17). A recent theoretical study has provided insight into possible mechanisms for such rapid exchange in the absence of duplex dissociation (59).

In contrast to $\frac{GGU}{PCCG}$ $\frac{GAA}{AAG}$ $\frac{GGCU}{CCG}$, duplexes a5, P5, I5, and I15 all have a single predominant conformation. As indicated by NOE patterns (Figure 3), there is little structural

change for a5, P5, and I5 duplexes relative to $\frac{GGU}{PCCG}$ $\frac{GGA}{AAG}$ $\frac{GGCU}{CCG}$ (9) and to the major conformation of the A5 duplex. NMR data for the I15 duplex suggest that its structure differs from that of a5, P5, and I5 duplexes, but resembles the minor structure of the A5 duplex.

The duplex with a deoxyadenosine (a5), $\frac{GGU}{PCCG}$ $\frac{GaA}{AAG}$ $\frac{GGCU}{CCG}$ (-0.28 kcal/mol), has a single structure with a *trans* Hoogsteen/sugar edge A15-a5 pair (Figure 1). This is consistent with the fact that G5 is predominantly in C2'-*endo* sugar pucker conformation with a *trans*

Hoogsteen/sugar edge A15-G5 pair in $\frac{\text{GGU}}{\text{PCCG}} \frac{\text{GGCU}}{\text{AAG}} \frac{\text{GGCU}}{\text{CCG}}$ (-2.62 kcal/mol) (9). Presumably, the a5 duplex has a single structure because the deoxy sugar favors C2'-endo sugar pucker which thus favors a single conformation similar to that of

 $\begin{array}{c} {\rm GGU} \ \, \frac{{\rm GGA}}{{\rm AAG}} \ \, \frac{{\rm GGCU}}{{\rm CCG}} \ \, (9). \ \, \text{This is also consistent with the observation that the deoxy g5} \\ {\rm substitution \ in} \ \, \frac{{\rm GGU}}{{\rm PCCG}} \ \, \frac{{\rm GgA}}{{\rm AAG}} \ \, \frac{{\rm GGCU}}{{\rm CCG}} \ \, (-3.04 \ kcal/mol) \ \, \text{enhances loop stability by 0.42} \\ {\rm kcal/mol \ \, despite \ \, loss \ \, of \ \, hydrogen \ \, bonds \ \, G5 \ \, (2'-hydroxyl) - G14 \ \, (imino/amino). \ \, The} \\ \end{array}$

opposite change in thermodynamic stability is observed for $\frac{GGU}{PCCG}$ $\frac{GgA}{AAGAAA}$ $\frac{GGCU}{CCG}$

(0.67 kcal/mol) compared with $\frac{GGU}{PCCG}$ $\frac{GGA}{AAGAAA}$ $\frac{GGCU}{CCG}$ (0.17 kcal/mol). Perhaps the greater flexibility of a 3 \times 6 loop negates the necessity for a C2'-endo sugar.

Previous NMR studies showed no orientation exchange for the tandem sheared AA pairs of

 $\frac{C}{G}$ $\frac{AA}{AA}$ $\frac{G}{C}$ (7), presumably because switching the orientation would result in making the backbone too narrow for the adjacent Watson-Crick pair (60, 61). Evidently, switching the side where backbone narrowing occurs is not a problem when an AA pair is flanked by sheared GA pairs.

It has been pointed out that $\frac{GA}{AA}$ might be a potential groove binding and/or intercalation site

(43). The alternating sheared AA pair in $_G^U$ $\stackrel{GAA}{AAG}$ $\stackrel{G}{C}$ could potentially serve as a switch between different binding partners for dynamic functions because the smooth N1-C2-N3 edge of either A5 or A15 is presented differently in the minor groove in alternative orientations (Figure 5, bottom panel).

Energetics of molecular recognition

The consistency of structures for the 3GA, I5, P5, and a5 duplexes (Figures 2 and 3) provides models for studying the interactions determining the energetics of a 3×3 loop with

three sheared pairs. The P5 duplex $\frac{GGU}{PCCG}$ $\frac{GPA}{AAG}$ $\frac{GGCU}{CCG}$ (-0.53 kcal/mol) is

thermodynamically similar to the a5 duplex $_{\mbox{\scriptsize PCCG}}^{\mbox{\scriptsize GGU}}$ $_{\mbox{\scriptsize AAG}}^{\mbox{\scriptsize GGCU}}$ (-0.28 kcal/mol) and to

 $\begin{array}{ccc} GGU & GAA & GGCU \\ PCCG & \overline{AAG} & CCG \end{array} \ (-0.48 \ kcal/mol) \ (Table \ 2). \ This \ is \ in \ agreement \ with \ the \ formation \end{array}$ of a sheared PA pair (trans Hoogsteen/sugar edge A15-P5) without loss of hydrogen bonds compared with a sheared AA pair (Figure 1).

The I5 duplex $\frac{GGU}{PCCG}$ $\frac{GIA}{AAG}$ $\frac{GGCU}{CCG}$ (-1.22 kcal/mol) is 1.40 kcal/mol less stable than

 $\frac{GGU}{PCCG}$ $\frac{GGA}{AAG}$ $\frac{GGCU}{CCG}$ (–2.62 kcal/mol). Similar destabilizations of 1.74 and 2.00 kcal/mol

are observed with the G to I substitutions in ${{
m GGU}\over{
m PCCG}}$ ${{
m GAA}\over{
m AIG}}$ ${{
m GGCU}\over{
m CCG}}$ (-0.88 kcal/mol) and

 $\frac{\mathrm{GGU}}{\mathrm{PCCG}} \frac{\mathrm{GIA}}{\mathrm{AAGAAA}} \frac{\mathrm{GGCU}}{\mathrm{CCG}} (2.17 \text{ kcal/mol}), \text{ respectively (Table 2)}. \text{ This is presumably}$ primarily due to the loss of hydrogen bonds of G5 amino to A15N7 and to the A15 nonbridging oxygen in the IA pair compared to GA (Figure 1). The free energy of about 1.5 kcal/mol attributed to two hydrogen bonds is a lower limit because subtle rearrangement of three-dimensional structure is expected upon inosine substitution and this can strengthen the remaining hydrogen bonds (62).

Interestingly, $\frac{\rm GGU}{\rm PCCG}$ $\frac{\rm GIA}{\rm AAG}$ $\frac{\rm GGCU}{\rm CCG}$ (–1.22 kcal/mol) is more stable than

GGU GaA GGCU

 $rac{GGC}{PCCG} = rac{GGC}{AAG} = rac{GGCC}{CCG}$ (-0.28 kcal/mol) (Table 2 and Figure 2), even though the number of base-base hydrogen bonds is expected to be the same (Figure 1). A water-mediated hydrogen bond between the G imino proton and the non-bridging oxygen of A was predicted (15) and observed in a crystal structure of a GNRA tetraloop (63). A similar water-mediated hydrogen bond might exist between I5H1 and an A15 non-bridging oxygen in the I5 duplex,

 $\frac{GGU}{PCCG}$ $\frac{GIA}{AAG}$ $\frac{GGCU}{CCG}$. This water-mediated hydrogen bond might explain the extra stability of the I5 duplex relative to P5, A5, and a5 duplexes (Figure 2).

The D5 (2,6-diamino purine) duplex, $\frac{GGU}{PCCG}$ $\frac{GDA}{AAG}$ $\frac{GGCU}{CCG}$ (-1.45 kcal/mol), is about 1.2

kcal/mol less stable than $\frac{GGU}{PCCG}$ $\frac{GGA}{AAG}$ $\frac{GGCU}{CCG}$ (-2.62 kcal/mol) (Table 2 and Figure 2), even though no base–base hydrogen bonds are lost (Figure 1). The destabilizing effect upon substituting D5 for G5 may also be due to loss of the proposed water-mediated hydrogen bond between G5H1 and an A15 non-bridging oxygen (15, 63). It is also possible that the 2-amino group of G is a better hydrogen bond donor than that of D (2,6-diamino purine) because of relatively larger positive partial charges on the G amino hydrogens (64). Larger

destabilization of 2.3 kcal/mol is observed for $\frac{\mathrm{GGU}}{\mathrm{PCCG}}$ $\frac{\mathrm{GDA}}{\mathrm{AAGAAA}}$ $\frac{\mathrm{GGCU}}{\mathrm{CCG}}$ (2.47 kcal/mol)

compared with $\frac{GGU}{PCCG}$ $\frac{GGA}{AAGAAA}$ $\frac{GGCU}{CCG}$ (0.17 kcal/mol). Perhaps the greater flexibility of the size asymmetric loop allows binding of a water molecule in a GA pair to be more

favorable. In contrast, D16 substitution for A16 to give $\frac{GGU}{PCCG}$ $\frac{GGA}{DAG}$ $\frac{GGCU}{CCG}$ (-3.32 kcal/

mol) and $\frac{GGU}{PCCG}$ $\frac{GGA}{DAG}$ $\frac{GGCU}{CCG}$ (-0.78 kcal/mol) stabilizes the loop by -0.70 kcal/mol and -0.30 kcal/mol, respectively, even though the 6-amino group on D (2,6-diamino purine) has essentially the same partial charges as those of A (64). Possibly the extra amino group on D allows better stacking and/or extra hydrogen bonding to the backbone.

A 2'-O-methyl substitution favors the C3'-endo sugar conformation (65–67). The 2'-O-

methyl A15 substitution in ${\rm \frac{GGU}{PCCG}}~{\rm \frac{GGA}{AMG}}~{\rm \frac{GGCU}{CCG}}$ (–3 83 kcal/mol) and

 $\frac{GGU}{PCCG}$ $\frac{GGA}{AMG}$ $\frac{GGCU}{CCG}$ (-0.83 kcal/mol) stabilizes the duplexes by -1.21 and -0.35

kcal/mol relative to $\frac{GGU}{PCCG}$ $\frac{GGA}{AAG}$ $\frac{GGCU}{CCG}$ (- 2.62 kcal/mol) and $\frac{GGU}{PCCG}$ $\frac{GAA}{AAG}$ $\frac{GGCU}{CCG}$ (-0.48 kcal/mol), respectively (Table 2). Part of the reason for the smaller effect in the

U GAA G G AAG C loop may be that the 2'-O-methyl substitution limits the loop to a single conformation. If the two conformations of the natural loop have equal concentrations, then the dynamics would favor loop formation by $G^{\circ}_{37} = -T$ $S = -310(1.987)\ln 2 = -0.4 \text{ kcal/mol}$.

The $\frac{GA}{\Delta}$ motif within other internal loops

Other $\frac{GA}{AA}$ motifs have non-sheared pairs. Only face-to-face (\emph{cis} Watson-Crick/Watson-

Crick) purine-purine pairs (Figure 1) form in the loop $_G$ $\frac{U}{AAG}$ $\frac{C}{G}$ in helix 23 of the crystal structure of T. thermophilus 16S rRNA (44), which is consistent with previous NMR studies that imino GA (face-to-face, cis Watson-Crick/Watson-Crick A–G) is favored in the

 G C C O O O motif (68). The crystal structure of a symmetric 4×4 loop C O O O shows

little base overlap for the $\frac{GA}{AA}$ nearest neighbors with sheared GA (*trans* Hoogsteen/sugar edge A–G) pairs and *trans* Watson-Crick/Hoogsteen A–A pairs (43). (In the *trans* Watson-Crick/Hoogsteen A–A pairs, the A paired with Hoogsteen edge is shown in bold (Figure 1).)

Evidently, $\frac{GA}{AA}$ is an intrinsically flexible structure within size-symmetric internal loops.

The free energy increment for $^{C}_{G}$ $\frac{GAAA}{AAAG}$ $^{G}_{C}$ at 37 $^{\circ}C$ in 1 M NaCl is 0.96 kcal/mol as

calculated from the measurement of the duplex ${\rm CGC \over GCG}$ ${\rm GAAA \over AAAG}$ ${\rm GGC \over CCG}$ (13). In contrast,

internal loops with consecutive GA pairs are very stable, e.g. $_G^U \frac{GGAA}{AAGG} \frac{G}{C}$ (-4.27 kcal/mol) (13), which is consistent with extensive stacking and hydrogen bonding as observed in the

crystal structure of the loop $_G^C$ \xrightarrow{GGAA} $_C^G$ with four sheared GA pairs (69). Previous

thermodynamic studies showed that the destabilizing $2 \times 2 \log_{\frac{1}{2}} \frac{AA}{AA} \frac{G}{C}$ (1.2 kcal/mol) with two sheared AA pairs has similar base pairing and stacking geometries but fewer

hydrogen bonds than ${\rm G \over AG} {\rm G \over C}$ (-0.7 kcal/mol) with two sheared GA pairs (2, 5, 7, 8). Evidently, the thermodynamic and structural effects of replacing a GA pair with an AA pair are context dependent.

A recently proposed "reverse kink-turn" motif involves a size asymmetric 2 × 5 internal

 $\begin{array}{c} G & GA \\ loop, & \overline{C} & \overline{AAACA} & G \\ N & AACA & C \end{array}$ with a sheared GA followed by a symmetric AA pair (\$trans\$ Hoogsteen/Hoogsteen A-A) (Figure 1b) (49). Such a conformation is also observed in some

Loop E motifs, $\frac{GAUGA}{AA}$ (47, 48). The glycosidic bond of the A (in bold) 3' to the G of the sheared GA pair is in a *syn* conformation. Thus it might facilitate the packing between two stems via major grooves, with the smooth N1-C2-N3 edge of the bold A flipped to the major

groove. Different detailed structures of an AA pair adjacent to a sheared GA pair in $\frac{GA}{AA}$ nearest neighbors with the bold A paired on its Hoogsteen side are also observed in kinkturn motifs (70) within internal loops, such as kt-11 (*trans* Watson-Crick/Hoogsteen A-A), and multibranch loops such as kt 94/99 (*trans* Hoogsteen/Sugar edge A-A), kt 4/5 (*trans* Hoogsteen/Hoogsteen A-A, with the \underline{A} in a *syn* glycosidic conformation). These kink-turns facilitate local and long-range tertiary interactions (70). In these cases, the A 3' to the G of a

sheared GA pair prefers to base pair with its Hoogsteen edge. Evidently, the $\frac{GA}{AA}$ nearest neighbor is intrinsically flexible compared with the motif of consecutive GA pairs (13) in both size symmetric and asymmetric internal loops.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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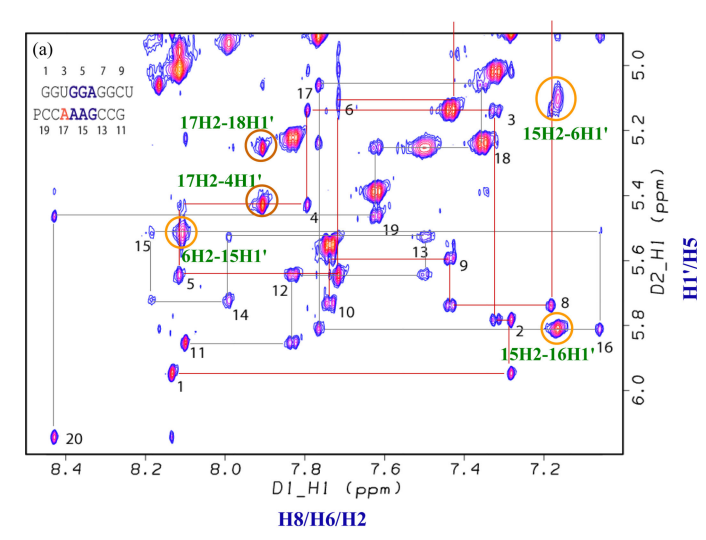
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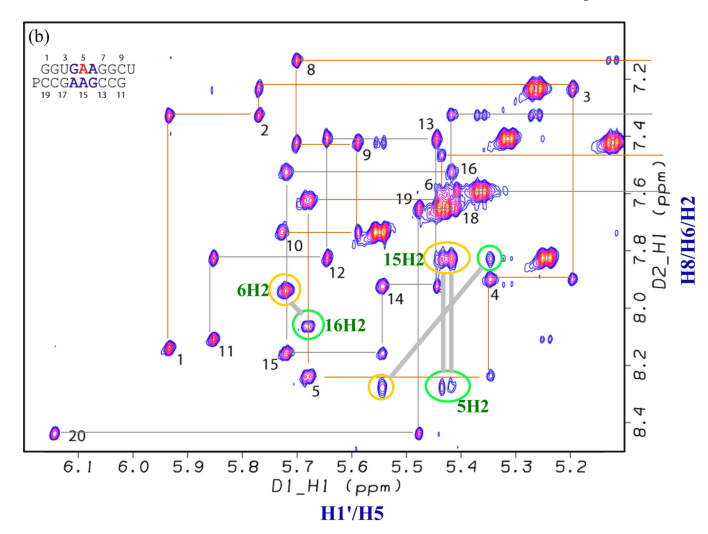
Schematic representation of (a) different sheared pairs, and (b) a GA and various AA pairs mentioned in this paper. The hydrogen bonds between base and backbone are not shown. Note that two conformations with one base-base hydrogen bond are possible for an AA pair because the amino group of either A can form the hydrogen bond. Only one such conformation is possible for the PA and IA pairs because neither P nor I have amino groups.

		$\Delta G^{\circ}_{37, loop}$
		(kcal/mol)
3GA	1 3 5 7 9 GGU GGA GGCU PCCG AAG CCG 19 17 15 13 11	-2.62
A17	1 3 5 7 9 GGU GGA GGCU PCC AAAG CCG 19 17 15 13 11	-2.27
15	1 3 5 7 9 GGU GIA GGCU PCCG AAG CCG 19 17 15 13 11	- 1.22
I15	1 3 5 7 9 GGU GAA GGCU PCCG AIG CCG 19 17 15 13 11	-0.88
P5	1 3 5 7 9 GGU GPA GGCU PCCG AAG CCG 19 17 15 13 11	- 0.53
A5	1 3 5 7 9 GGU GAA GGCU PCCG AAG CCG 19 17 15 13 11	- 0.48
a5	1 3 5 7 9 GGU GaA GGCU PCCG AAG CCG 19 17 15 13 11	-0.28

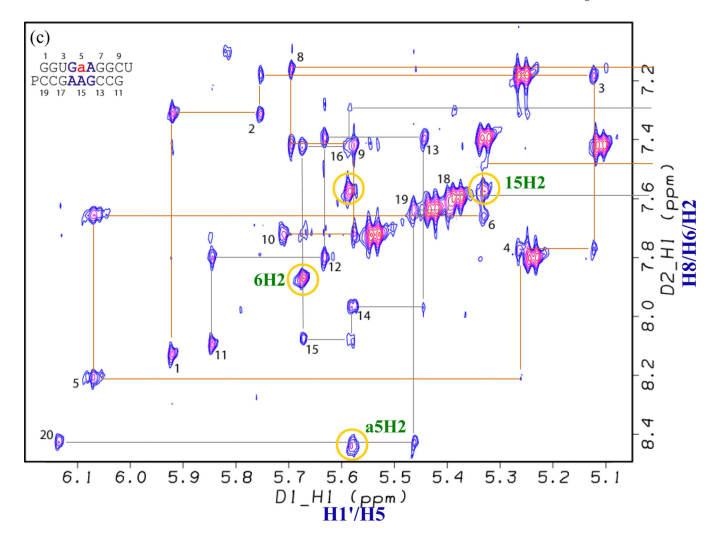
Secondary structure, numbering, and abbreviations for the duplexes studied previously (9, 13) and here. Lower case "a" represents deoxyadenosine. The value to the right of each duplex is the free energy increment in kcal/mol for formation of the internal loop at 37 °C, at pH7 in 1 M NaCl.

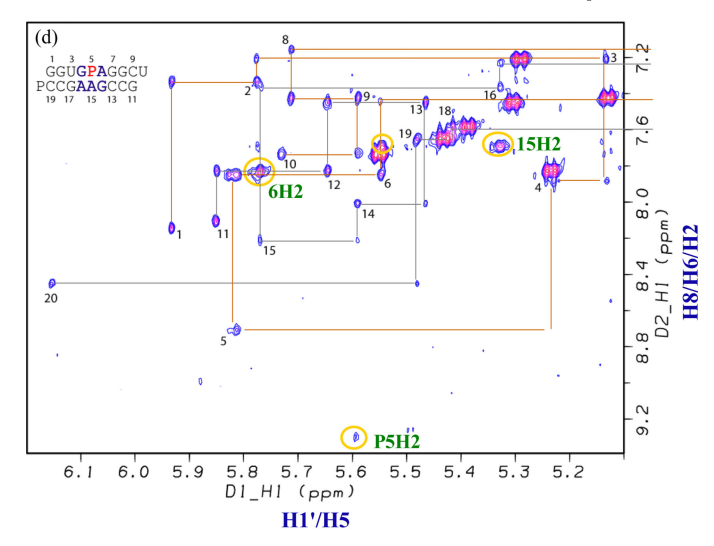


a

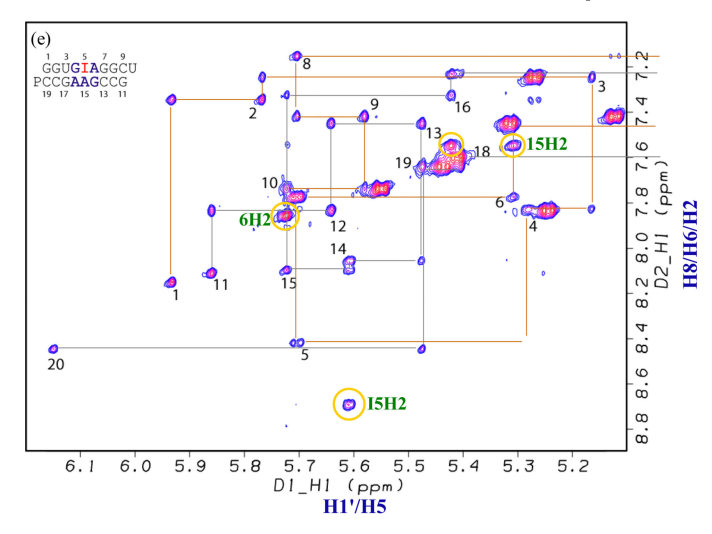


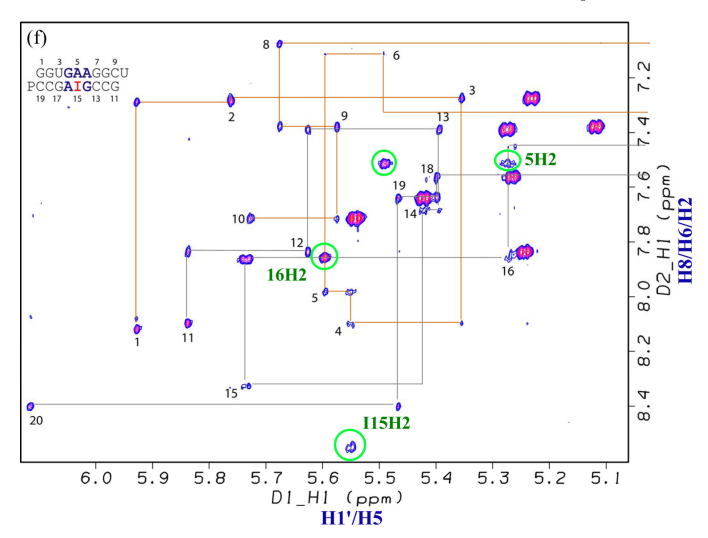
b





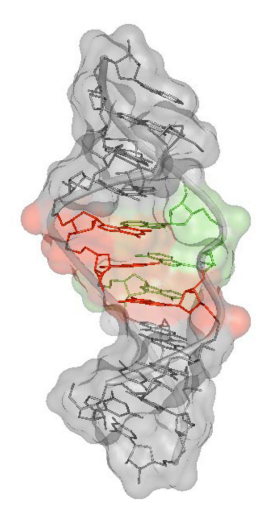
d





f

Figure 3. (H8/H6/H2)-(H1'/H5) region of the 400 ms mixing time NOESY spectra of duplexes (Figure 2) (a) A17, (b) A5, (c) a5, (d) P5, (e) I5, and (f) I15 at 30 °C in 80 mM NaCl, 10 mM sodium phosphate, 0.5 mM disodium EDTA, pD 7 except I5 duplex at pD 6. For A5 sequence in panel (b), yellow and green circles connected by gray lines identify related cross peaks of major and minor conformations, respectively. Yellow and green circles in other spectra identify cross peaks related to those in circles of the same color for A5 duplex.



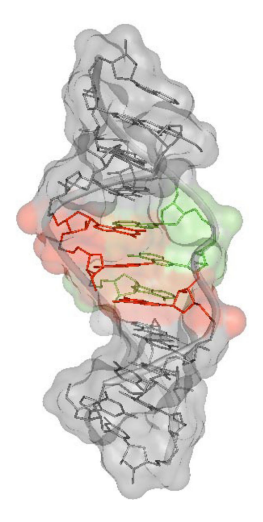


Figure 4.

Major groove stereo view of $\frac{GGU}{PCCA} \frac{GGA}{AAG} \frac{GGCU}{CCG}$, with A6/A15/A16 stack in minor groove shown in green, and G4/G5/G14 stack in major groove shown in red. Hydrogen and non-bridging oxygen atoms are deleted for clarity.

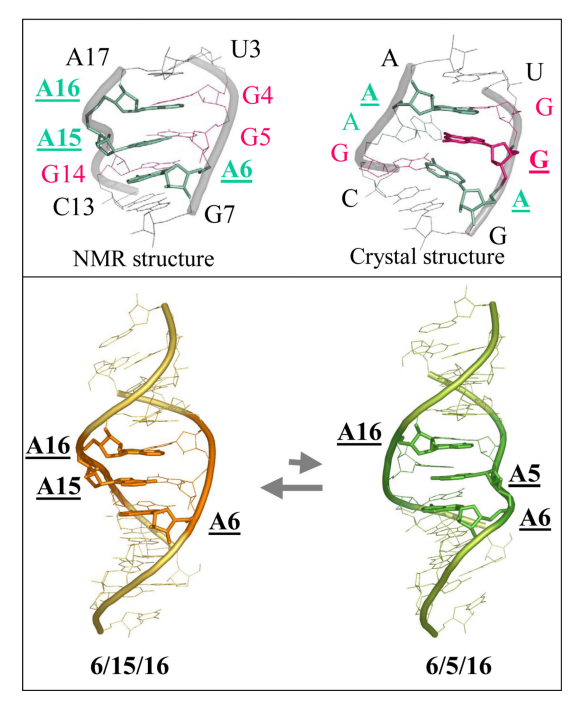


Figure 5.Comparison of minor groove view of the crystal and NMR structures. Residues stacking in minor groove are shown in sticks and labeled in bold. Hydrogen and non-bridging oxygen atoms are deleted for clarity. Top panel: NMR and crystal structures of the internal loop

 $U = \frac{GGA}{A \ AG} = \frac{G}{C}.$ Three G's and three A's in the loop are shown in red and green, respectively.

Bottom panel: NMR structures of the duplex $\frac{GGU}{PCCG}$ $\frac{GAA}{AAG}$ $\frac{GGCU}{CCG}$ with an alternating middle sheared AA pair.

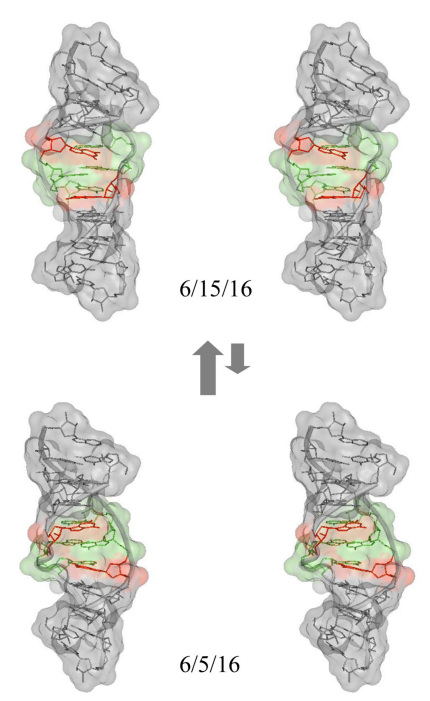


Figure 6.

Major groove stereo views of two alternating structures of $\frac{GGU}{PCCG}$ $\frac{GAA}{AAG}$ $\frac{GGCU}{CCG}$ with A6/A15/A16 or A6/A5/A16 stacks in minor groove. Two G's and four A's in the loop are shown in red and green, respectively. Hydrogen and non-bridging oxygen atoms are deleted for clarity. The hydrogen bonding shown for the AA pair in the minor conformation is similar to that shown in Figure 1a, but a variety of hydrogen bonding patterns are seen in the ensemble of structures generated with the restraints from NMR data.

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Table 1

Measured Thermodynamic Parameters for Duplex Formation in 1 M NaCl, pH7 and in 80 mM NaCl if Listed in Parentheses.

		T_{xx}^{-1} vs $\ln(C_x/4)$ plots (eq. 1)	nlots (ea 1)			Average of melt curve fits	curve fits	
i		((ba) mard			0		
Sequences	- H° (kcal/mol)	(en)	- G°37 (kcal/mol)	T_m^a	- H° (kcal/mol)	(ea)	$-\mathbf{G}^{\circ}_{37}$ (kcal/mol)	T_m^{a}
GGU <u>GGA</u> GGCU PCCGAM G CCG	108.4±3.5	302.9±10.4	14.47±0.24	61.5	105.1±1.2	293.0±3.6	14.24±0.12	61.6
GGU <u>GGA</u> GGCU PCCG DAG CCG	104.2 ± 3.1	290.9±9.4	13.96±0.22	8.09	105.4 ± 3.0	294.4±9.0	14.04±0.23	8.09
GGU <u>GgA</u> GGCU PCCGAAGCCG	103.9 ± 3.1	290.8±9.3	13.68±0.20	59.9	100.1 ± 3.0	279.3±8.9	13.44±0.21	0.09
$ ext{GGU} \overline{ ext{GGA}} ext{GGCU}^b$	94.3±8.2 (82.2±3.4)	261.2±24.5 (233.5±10.7)	13.26 ± 0.57 (9.79 ± 0.10)	60.8 (49.8)	94.5±2.4 (87.8±4.5)	261.9 ± 7.2 (251.1±14.1)	13.27 ± 0.25 9.94 ± 0.15	60.8 (49.5)
GGU <u>GGA</u> GGCU ^b PCCA AAG CCG	$92.7\pm2.2 \\ (84.6\pm4.3)$	258.0 ± 6.6 (243.2±13.6)	12.64 ± 0.13 (9.15 ± 0.13)	58.9 (46.9)	93.2 ± 3.9 (81.9 ±9.3)	259.7 ± 12.0 (234.9 \pm 29.2)	12.67 ± 0.23 (9.09 ± 0.30)	58.9 (47.0)
GGC <u>GGA</u> GGCU ^b PCCGAAGCCG	$\begin{array}{c} 81.2{\pm}7.0 \\ (56.1{\pm}1.9) \end{array}$	$223.4\pm21.2 \\ (152.6\pm6.1)$	11.92 ± 0.47 (8.74 ± 0.05)	59.1 (49.7)	77.8 ± 5.8 (60.7±8.3)	213.1±17.7 (167.2±26.3)	11.76 ± 0.36 (8.88±0.16)	59.3 (49.5)
GGU <u>GDA</u> GGCU PCCG AAG CCG	93.6±1.9	262.9±5.9	12.09±0.11	56.6	89.4±4.4	250.1±13.6	11.88±0.22	56.7
GGU <u>GIA</u> GGCU PCCGAAGCCG	90.7±2.5	254.3±7.7	11.86±0.14	56.4	89.6±5.0	250.8±15.3	11.84±0.27	56.5
GGU <u>GAA</u> GGCU PCCGA IG CCG	90.6±3.4	255.0±10.4	11.52±0.18	55.1	92.0±3.4	259.0±10.2	11.61±0.20	55.1
GGU <u>GAA</u> GGCU PCCGAM G CCG	92.6±3.8	261.7±11.7	11.47±0.19	54.5	90.3±4.6	254.4±14.1	11.38±0.23	54.6
GGU <u>GAA</u> GGCU PCCG DAG CCG	89.2±1.8	250.8±5.6	11.42±0.09	55.0	88.5±4.6	248.6±14.1	11.41±0.23	55.1
GGU <u>GPA</u> GGCU PCCGAA G CCG	88.9±2.1	250.5±6.4	11.17±0.10	54.1	84.3±2.4	236.6±7.5	10.97±0.10	54.2
$ ext{GGU}_{ ext{GAA}} ext{GGCU}^{b}$	84.2 ± 6.1 (75.7±3.3)	235.7 ± 18.6 (219.4±10.4)	11.12 ± 0.32 (7.64 ±0.05)	54.9 (41.6)	86.5 ± 5.4 (74.9±3.6)	242.8 ± 16.9 216.8 ± 11.4	11.23 ± 0.25 (7.64\pm0.11)	54.8 (41.7)
GGU <u>GaA</u> GGCU PCCGAM G CCG	88.7±3.4	250.9±10.6	10.93±0.16	53.2	85.5±2.4	240.9±7.3	10.79 ± 0.14	53.3
GGU <u>GaA</u> GGCU PCCG AAG CCG	86.7±1.5	244.2±4.7	10.92±0.07	53.5	82.9±1.5	232.7±4.8	10.75 ± 0.07	53.6
GGU <u>GGA</u> GGCU ^c PCCGAAGAAACCG	90.8±1.9	259.1±6.0	10.47±0.07	51.1	84.5±4.2	239.4±13.0	10.25 ± 0.19	51.3

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	$\mathbf{T}_{\mathbf{l}}$	$T_{\rm M}^{-1} \ vs \ ln(C_T/4) \ plots \ (eq \ 1)$	plots (eq 1)		₹ ⁷	Average of melt curve fits	curve fits	
Sequences	- H° (kcal/mol)	(ea)	$-\mathbf{G}^{\circ}_{37}$ (kcal/mol)	T_{m}^{a}	- H° (kcal/mol)	(ea)	$-G^{\circ}_{37}$ (kcal/mol)	T_m^a
GGU GgA GGCU PCCGAAGAAACCG	89.4±2.9	256.1±8.9	9.97±0.09	49.4	81.3±5.2	230.8±16.2	9.74±0.20	49.7
GGU <u>GIA</u> GGCU PCCG AIG CCG	80.7±1.4	228.9±4.4	9.70±0.05	49.7	78.5±1.9	222.2±5.9	9.61±0.08	49.7
GGU GIA GGCU PCCG AAGAAA CCG	80.3±4.1	231.7±13.0	8.47±0.09	44.7	71.8±2.5	204.8±7.8	8.31±0.14	44.9
GGU <u>GDA</u> GGCU PCCGAAGAAACCG	70.0±3.4	199.3±11.0	8.17±0.06	4.4	65.2±2.9	184.0±9.3	8.12 ± 0.13	8.44.8
GGU GAA GGCU PCCGAAGAAACCG	73.6±4.3	211.1±13.7	8.07±0.08	43.6	67.8±7.1	192.8±22.5	8.01±0.12	43.9
GGU GaA GGCU PCCG AAGAAA CCG	72.7±4.8	209.0±15.3	7.91±0.09	43.0	66.1±3.3	187.7±10.4	7.84±0.14	43.3
GGU <u>GPA</u> GGCU PCCGAAGAAACCG	66.1±3.9	188.7±12.3	7.57±0.07	42.0	59.2±2.2	166.6±7.3	7.56±0.14	42.5
GGUGGCU ^b PCCGCCG	75.6±3.9	205.0±11.7	12.05±0.28	61.4	80.8±2.3	220.5±7.0	12.39±0.17	61.3

 a At CT = 0.1 mM;

 $b \\ Data$ measured in 1 M NaCl are from Ref (9).

^cRef (13).

Table 2

Measured Thermodynamic Parameters for Internal Loop Formation in 1 M

NaCl, pH7 ^a				
Sequence	$G^{\circ}_{37, loop}$ (kcal/mol)	H° _{loop} (kcal/mol)	S°_{loop} (eu)	G° _{37, loop} (kcal/mol)
GGU <u>GGA</u> GGCU PCCGAMGCCG	-3.83±0.59	-38.4±9.9	-111.4±30.3	-1.21^{b}
GGU <u>GGA</u> GGCU PCCG DAG CCG	-3.32±0.58	-42.0±9.7	-99.4±30.0	-0.70^{b}
GGU <u>GgA</u> GGCU PCCGAAGCCG ^C	-3.04±0.57	-33.9±9.7	-99.3±29.9	-0.42^{b}
GGU <u>GGA</u> GGCU ^c PCCG AAG CCG	-2.62±0.78 [-2.39]	-24.3±12.4	-69.7±37.5	0.00^{b}
GGU <u>GGA</u> GGCU ^C PCCA AAG CCG	-2.27±0.59 [-1.44]	-23.9 ± 9.7	-69.5±29.5	-
GGU <u>GDA</u> GGCU PCCGAAGCCG	-1.45±0.54	-23.6±9.4	-71.4±29.0	$1.17^b, -0.97^d$
GGU <u>GIA</u> GGCU PCCGAAGCCG	-1.22±0.55	-20.7±9.6	-62.8±29.4	$1.40^b, -0.74^d$
GGU <u>GAA</u> GGCU PCCG AIG CCG	-0.88±0.56	-20.6±9.8	-63.5±30.3	$1.74^b, -0.40^d$
GGU <u>GAA</u> GGCU PCCGAMGCCG	-0.83±0.57	-22.6±10.0	-70.2±30.8	-0.35^{d}
GGU <u>GAA</u> GGCU PCCG DAG CCG	-0.78 ± 0.54	-27.0±9.4	-59.3±29.0	-0.30^{d}
GGU <u>GPA</u> GGCU PCCGAAGCCG	-0.53±0.54	-18.9±9.4	-59.0±29.1	$2.09^b, -0.05^d$
GGU <u>GAA</u> GGCU ^C PCCGAAGCCG	-0.48±0.57 [-0.03]	-14.2±11.1	-44.2±34.0	0.00^{d}
GGU <u>GaA</u> GGCU PCCG AMG CCG	-0.29±0.56	-26.5±9.8	-59.4±30.4	0.19^{d}
GGU <u>GaA</u> GGCU PCCGAAGCCG	-0.28 ± 0.54	-16.7±9.3	-52.7±28.8	0.20^{d}
GGU <u>GGA</u> GGCU ^e PCCGAAGAAACCG	0.17±0.54 [0.20]	-20.8±9.4	-67.6±29.0	0.00 ^f
GGU <u>GgA</u> GGCU PCCGAAGAAACCG	0.67±0.54	-19.4±9.6	-64.6±29.8	0.50 ^f
GGU <u>GIA</u> GGCU PCCG AIG CCG	0.94±0.54	-10.7±9.3	-37.4±28.8	-
GGU <u>GIA</u> GGCU PCCGAAGAAACCG	2.17±0.54	-10.3±10.1	-40.2±31.2	2.00^f , -0.40^g
GGU <u>GDA</u> GGCU PCCGAAGAAACCG	2.47±0.54	0.0±9.8	-7.8±30.5	2.30^f , -0.10^g
GGU <u>GAA</u> GGCU PCCGAAGAAACCG	2.57±0.54 [2.56]	-3.6±10.2	-19.6±31.6	0.00^{g}
GGU <u>GaA</u> GGCU PCCGAAGAAACCG	2.73±0.54	-2.7±10.4	-17.5±32.3	2.56 ^f , 0.16 ^g
GGU <u>GPA</u> GGCU PCCGAAGAAACCG	3.07±0.54	3.9±10.0	2.8±31.0	2.90 ^f , 0.50 ^g

^aExperimental error for $G^{\circ}37$, H° , and S° for the canonical stems are estimated as 4%, 12%, and 13.5%, respectively, according to ref (31). There is less error in comparisons between these sequences because the stems are either identical or different by only one or two base pairs. Values in square brackets are predicted according to ref (13);

 $\begin{array}{c} {\rm GGU} \ \ {\rm GGA} \ \ {\rm GGCU} \\ {\rm Compared \ with \ PCCG} \ \ \overline{\rm AAG} \ \ {\rm CCG} \end{array} \ \ {\rm from \ ref \ (9)}; \\$

 c From ref (9);

 $\begin{array}{ccc} & \text{GGU} & \text{GAA} & \text{GGCU} \\ ^{d}\text{Compared with} & \text{PCCG} & \overline{\text{AAG}} & \text{CCG} & \text{from ref (9);} \end{array}$

^eFrom ref (13);

 $f_{\mbox{Compared with }\mbox{PCCG}} \xrightarrow{\mbox{GGU}} \frac{\mbox{GGA}}{\mbox{AAGAAA}} \xrightarrow{\mbox{GGCU}} \frac{\mbox{GGCU}}{\mbox{from ref (13);}}$

GGU GAA GGCU

 g Compared with PCCG $\overline{\mathrm{AAGAAA}}$ CCG \cdot

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Table 3

Distance Restraints Involving A5 and A15 Residues for the Structural Modeling of $\frac{GGU}{PCCG}$ $\frac{GAA}{AAG}$ $\frac{GGCU}{CCG}$ and comparison with that of

 $\begin{array}{ccc} \text{GGU} & \text{GGA} & \text{GGCU} \\ \text{PCCG} & \overline{\text{AAG}} & \text{CCG} \end{array} (9).$

		A6/A	A6/A15/A16					A6/A	A6/A5/A16		
			Dista	Distance (Å)					Dista	Distance (Å)	
Atom1	Atom2	Lower	Upper	NMR	$Model^b$	Atom1	Atom2	Lower	Upper	NMR	$Model^b$
G4H3'	A5H8	1.80	5.00	3.35	3.50/4.73	G14H3'	A15H8	1.80	5.00	2.79	4.47/2.80
G4H8	A5H8	3.40	00.9	4.86	5.41/6.68	G14H8	A15H8	\sim^{c}	×	×	6.68/5.51
A5H1'	A6H8	1.80	4.12	3.17	3.11/4.98	A15H1'	A16H8	1.80	4.34	3.34	5.20/2.75
G14H1'	A5H2	1.80	5.00		4.62/10.32	G4H1'	A15H2	1.80	5.00		10.22/4.23
G14H2'	A5H2	2.00	5.00		2.21/9.41	G4H2'	A15H2	1.80	5.00		9.98/2.27
A5H2	A15H8	2.00	5.00		2.30/10.41	A5H8	A15H2	1.80	5.00		10.47/2.35
A5H4'	A16H2	2.83	5.26	4.05	4.02/5.38	A15H4'	A6H2	3.00	00.9		5.52/4.44
A6H1'	A15H2	1.80	4.86	3.74	2.55/7.66	A16H1'	A5H2	1.80	5.00		7.11/2.88
A15H1'	A6H2	1.80	4.14	3.19	2.84/6.60	A5H1'	A16H2	1.80	5.00		6.08/2.74
G14H1'	A15H8	2.00	5.50	4.23	4.69/5.99	G4H1'	A5H8	1.80	5.52	4.24	5.73/4.82
A16H1'	A15H2	1.80	4.30	3.31	2.80/7.39	A6H1'	A5H2	1.80	5.00		7.75/2.80
A15H8	A16H8	3.08	5.73	4.41	4.08/6.16	A5H8	A6H8	3.27	00.9	4.67	6.58/4.06
A15H2'	A16H8	1.80	3.99	3.07	3.32/4.64	A5H2'	A6H8	1.80	4.66	3.58	5.07/3.13
A15H3'	A16H8	1.80	5.00		2.48/4.80	A5H3'	A6H8	1.80	4.91	3.77	4.86/2.47
		T)	istance re	straints th	Distance restraints that are the same for A6/A15/A16 and A6/A5/A16	e for A6/A	15/A16 and	1 A6/A5/A	.16		
			Dista	Distance (Å)					Dist	Distance (Å)	
Atom1	Atom2	Lower	Upper	NMR	Model b	Atom1	Atom2	Lower	Upper	NMR	qlodel p
G4H2'	A5H8	1.80	5.00	4.86	3.08/3.10	G14H2'	A15H8	1.80	4.76	3.66	2.83/3.44
A5H1'	A5H2'	1.80	3.94	3.03	2.85/2.45	A15H1'	A15H2'	1.80	3.65	2.80	2.61/2.71
A5H1'	A5H3'	1.80	4.81	3.70	3.70/3.56	A15H1'	A15H3'	1.80	5.00	4.19	3.66/3.69
A 51111	V 5117'	1 80	177	3 30	3 26/3 16	A 15111	A 1511A	001	,	,	0

		A6/A	A6/A15/A16					A6/A	A6/A5/A16		
			Dista	Distance (Å)					Dist	Distance (Å)	
Atom1	Atom2	Lower	Upper	NMR	$q_{ m log}$	Atom1	Atom2	Lower	Upper	NMR	$Model^b$
A5H1'	A5H8	1.80	4.97	3.82	3.72/3.41	A15H1'	A15H8	1.80	5.04	3.88	3.48/3.68
A5H2'	A5H8	1.80	3.19^{d}	2.45	2.10/3.97	A15H2'	A15H8	1.80	5.00	2.79	4.11/2.03
A5H3'	A5H8	1.80	5.00	3.35	4.09/3.42	A15H3'	A15H8	1.80	4.27	3.28	3.42/3.89
<u> </u>	stance rest	aints invo	Vino G5 a	nd A15 re	GGU Distance restraints involving G5 and A15 residues for the structure modeling of PCCG	structure n	odeling of	GGU	GGA	CCG	J.
			Dista	Distance (Å)			0				
Atom1	Atom2	Lower	Upper	NMR	Model	Atom1	Atom2	Lower	Upper	NMR	Model
G4H3'	G5H8	2.41	4.47	3.44	3.87	G14H3'	A15H8	×	×	×	4.50
G4H8	G5H8	2.36	00.9	3.37	5.69	G14H8	A15H8	×	×	×	6.35
A5H1'	A6H8	2.30	4.27	3.29	3.55	A15H1'	A16H8	3.28	00.9	4.69	4.81
G14H1'	A5H2	£			1	G4H1'	A15H2	×	×	×	8.77
G14H2'	G5H1	2.50	5.84	4.17	4.05	G4H2'	A15H2	×	×	×	8.74
A5H2	A15H8	,	,	,		G5H8	A15H2	×	×	×	10.38
G5H4'	A16H2	×	×	×	5.21	A15H4'	A6H2	×	×	×	5.49
A6H1'	A15H2	2.15	3.99	3.07	2.45	A16H1'	A5H2	,	,		
A15H1'	A6H2	2.13	3.95	3.04	2.92	G5H1'	A16H2	×	×	×	66.9
G14H1'	A15H8	3.27	00.9	4.67	4.44	G4H1'	A5H8	×	×	×	6.35
A16H1'	A15H2	2.06	3.83	2.94	2.53	A6H1'	A5H2	,			
A15H8	A16H8	×	×	×	4.82	G5H8	A6H8	×	×	×	7.16
A15H2'	A16H8	1.94	3.60	2.77	2.20	G5H2'	A6H8	×	×	×	5.47
A15H3'	A16H8	2.39	4.44	3.42	2.87	G5H3'	A6H8	×	×	×	5.14
G4H2'	G5H8	2.73	5.07	3.90	3.66	G14H2'	A15H8	2.02	3.74	2.88	2.50
G5H1'	G5H2'	2.23	4.14	3.19	2.99	A15H1'	A15H2'	2.30	4.26	3.28	2.73
G5H1'	G5H8	3.29	00.9	4.70	3.91	A15H1'	A15H8	3.15	5.85	4.50	3.74
				1							

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	Dis	tance rest	raints tha	t differ f	Distance restraints that differ for the structural modeling of A6/A15/A16 and A6/A5/A16	ural modeli	ng of A6/≜	x15/A16 aı	nd A6/A5/	A16	
		A6/A]	A6/A15/A16					A6/A	A6/A5/A16		
			Dista	Distance (Å)					Dista	Distance (Å)	
Atom1	Atom2	Lower	Upper	NMR	Atom1 Atom2 Lower Upper NMR Model ^b	Atom1 Atom2 Lower Upper NMR Model ^b	Atom2	Lower	Upper	NMR	$Model^b$
G5H3'	G5H8	G5H8 2.46 4.57 3.52 4.05	4.57	3.52	4.05	A15H3'	A15H3' A15H8 2.09		3.88	2.99 3.14	3.14
G5H1	G4H1	2.34	5.46	3.90	3.52	A15H2	A6H8	3.84	00.9	5.48	4.70
G5H1	G14H1	G14H1 2.50 7.00	7.00		5.41	A15H2	A15H2 A16H2 3.49		00.9	4.99	4.54

 $[^]a$ All other distance restraints for both structural modelings are identical and are provided in Supporting Information.

 $^{^{}b}$ Distances measured for the averaged structure of A6/A15/A16 followed by A6/A5/A16;

 $^{^{}c}$ Cross-peaks not observed and labeled with \times ;

 $[\]frac{d}{L}$ Loosened upper bound to 4.50 Å for the structure modeling of A6/A5/A16;

GGU GGA GGCU The distances in the columns of models are measured from a representative structure of PCCG \overline{AAG} CCG (9);

 $[^]f$ not applied and labeled with -.

Table 4

Chemical shifts (ppm) and full widths (Hz) at half height of H2 peaks of the central loop residues, 5H2 and 15H2, in A5, a5, I5, P5, and I15 duplexes at 30 °C in 80 mM NaCl. G2H8 from the stem is included for reference. Error limits are listed in parentheses.

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	Lir	Linewidths (Hz))	Chem	Chemical shifts (ppm)	(mdd) s
Duplex	5H2	15H2	G2H8	5H2	15H2	G2H8
Y2	15.0 (1.5) ^a	13.1 (1)	4.1 (1)	8.28	7.83	7.32
a5	12.0 (2)	6.2 (1.5)	3.8 (1)	8.44	7.57	7.31
5d	10.0(1)	4.3 (1)	3.6 (0.5) 9.30	9.30	69.7	7.33
SI	7.1 (1)	4.1 (1)	4.4 (0.5)	8.69	7.55	7.35
511	4.4 (1)	12.0 (1.5)	12.0 (1.5) 4.8 (0.5) 7.51 8.55	7.51	8.55	67.Z

^aIf the chemical shift of the A5H2 resonance in the A5 duplex differs by 0.87 ppm between major and minor conformations and the fraction of A5 duplexes in the major conformation ranges between 0.6 and 0.9, then a rough calculation (67, 68) assuming an inherent linewidth of 4 Hz suggests the rate of exchange between the two conformations of the A5 duplex is between 20,000 and 65,000 s⁻¹. Page 40