Ion-induced folding of a kink turn that departs from the conventional sequence

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ABSTRACT
Kink turns (k-turns) are important structural motifs that create a sharp axial bend in RNA. Most conform to a consensus in which a three-nucleotide bulge is followed by consecutive G:A and A:G base pairs, and when these G:A pairs are modified in vitro this generally leads to a failure to adopt the k-turn conformation. Kt-23 in the 30S ribosomal subunit of Thermus thermophilus is a rare exception in which the bulge-distal A:G pair is replaced by a non-Watson–Crick A:U pair. In the context of the ribosome, Kt-23 adopts a completely conventional k-turn geometry. We show here that this sequence is induced to fold into a k-turn structure in an isolated RNA duplex by Mg$^{2+}$ or Na$^+$ ions. Therefore, the Kt-23 is intrinsically stable despite lacking the key A:G pair; its formation requires neither tertiary interactions nor protein binding. Moreover, the Kt-23 k-turn is stabilized by the same critical hydrogen-bonding interactions within the core of the structure that are found in more conventional sequences such as the near-consensus Kt-7. T. thermophilus Kt-23 has two further non-Watson–Crick base pairs within the non-canonical helix, three and four nucleotides from the bulge, and we find that the nature of these pairs influences the ability of the RNA to adopt k-turn conformation, although the base pair adjacent to the A:G pair is more important than the other.

INTRODUCTION
Kink turns (k-turns) are commonly found structural elements in the architecture of RNA–protein complexes and large RNA species. The k-turn motif introduces a sharp kink into the axis of the helix, thereby exerting an important influence on the long-range structure of RNA. k-turns were first identified in both subunits of the ribosome (1). They have also been found in the nucleolar RNA species that guide RNA modification (2–5), U4 snRNA (6,7) and in untranslated regions of mRNA (8,9), including within riboswitches (10–12). Therefore, k-turns seem to be involved in almost every aspect of RNA function, including the translation and modification of RNA, spliceosome assembly and the control of gene expression.

The great majority of k-turn motifs conform to a consensus secondary structure comprising a three-nucleotide bulge flanked by regular base pairing on its 5' side (the C helix) and two consecutive A:G pairs on its 3' side (the NC helix) (Figure 1). The k-turn Kt-7 in the 23S rRNA of the Haloarcula marismortui ribosome is close to this consensus (1); its structure is very similar to those of a number of other k-turns (2,6,10). The kinking juxtaposes the two minor grooves, with an included angle between the axes of ~60°, and the conformation appears to be stabilized by interactions between the stacked adenosines of the A:G pairs and the C stem, and by stacking of the 5' and central bases of the bulge on the ends of the C and NC helices, respectively.

k-turn motifs do not form a stably kinked structure in the absence of metal ions (13,14), or the binding of proteins of the L7Ae family (15). The kinked conformation can be stabilized by the addition of either divalent or monovalent metal ions (13) and folding can be treated as a two-state process. Thus, it is likely that electrostatic interactions must be screened to allow the folded structure to achieve stability. This suggests a dynamic character for k-turn structures, where they sample both the kinked and a more extended geometry. Computer modelling studies have suggested that k-turns undergo hinge-like motions on a fast timescale (16–18). We have made a detailed study of the hydrogen-bonding interactions that are important in stabilizing the kinked conformation of Kt-7 in the presence of Mg$^{2+}$ ions (19,20). These include critical hydrogen bonds between the 2'O of the L1 ribose and N1 of the adenine nucleobase in the In position, and between the 2'O of the L3 ribose and the proS non-bridging O of the phosphate group linking L1 and L2 (19).

We have also studied the importance of the two A:G pairs adjacent to the bulge of the archetypal k-turn (20).
Both are trans sugar edge (G)/Hoogsteen edge (A) pairs, linked by potential hydrogen bonds G:N2 to A:N7 and A:N6 to G:N3. The 1b\_1n pair is strongly buckled, while the bases of the 2n\_2b pair are much closer to coplanarity. These are conserved in the great majority of k-turns (21), and sequence substitution of any of the four nucleotides within Kt-7 (13), or disruption of hydrogen bonding (20), completely prevents ion-induced folding. On this basis, it would appear that the two A\_G pairs are an indispensable feature of the k-turn motif. It is therefore surprising that within the 16S RNA of the \textit{T. thermophilus} ribosome k-turn Kt-23 lacks the A\_G pair at the 2b\_2n position (1) (Figure 1), being replaced by a reverse Hoogsteen A\_U pair (22). This is also true in Kt-23 of \textit{E. coli} (23); the two Kt-23 structures are closely similar. We have previously shown that the introduction of the equivalent G2nU substitution in Kt-7 completely prevented detectable ion-induced folding by that sequence (13). Yet Kt-23 adopts a normal k-turn geometry in the ribosome (Figure 2A), with an inter-axial angle of 60° between the C and NC helices. Despite the absence of the A\_G pair at the 2b\_2n position, the 1n and 2b adenine nucleobases of Kt-23 become located in virtually identical positions if \textit{H. marismortui} Kt-7 and \textit{T. thermophilus} Kt-23 are superimposed by their backbones (Figure 2B), and are thus able to participate fully in the A-minor interaction (24) with the C helix. The L1 and L2 nucleobases cap the C and NC helices, respectively, and the key hydrogen bonds involving 2’-hydroxyl groups (19) are present in the Kt-23 structures (22,23). The corresponding Kt-23 sequences of human, mouse and \textit{D. melanogaster} are shown in Figure 2C. Many, though not all, k-turn sequences include a non-Watson–Crick pair at the 3b\_3n position; in \textit{T. thermophilus} Kt-23, this is a U\_G pair with a single hydrogen bond from G3n:N2 to U3b:O4. Unusually, the 4b\_4n position is also non-Watson–Crick, a cis A\_A pair hydrogen bonded from A4b:N6 to A4n:N1. The C1’–C1’ distance of the A\_U pair in Kt-23 at 2b\_2n is longer than that of an A\_G pair (Table 1), and the width of the NC helix could be a factor in stabilizing the k-turn geometry. We have therefore examined the effect of functional group substitutions designed to probe the effect of the stability of the 3b\_3n and 4b\_4n positions on the ability of \textit{T. thermophilus} Kt-23 to undergo ion-induced folding.

\section*{MATERIALS AND METHODS}

\subsection*{RNA synthesis and construction of k-turn species}

The majority of oligonucleotides were synthesized using t-BDMS phosphoramidite chemistry (27), as described in Wilson \textit{et al.} (28), with nucleobase and 2’-deoxyribo substitution as required. Fluorescein and Cy3-conjugated oligonucleotides were attached at 5’-termini as phosphoramidites during synthesis as required. Oligoribonucleotides were deprotected in 25% ethanol/ammonia solution at 55°C for 2 h, and evaporated to dryness. Oligoribonucleotides were redissolved in 300 \mu l 1M tetrabutylammonium fluoride (Aldrich) in tetrahydrofuran to remove t-BDMS protecting groups, and agitated at 20°C for 16 h prior to desalting by G25 Sephadex (NAP columns, Pharmacia) and ethanol precipitation. All oligonucleotides were purified by gel electrophoresis in polyacrylamide, and recovered from gel fragments by electrophoretion or diffusion in buffer followed by ethanol precipitation. Fluorescently labelled RNA was subjected to further purification by reversed phase HPLC on a C18 column (ACE10, Advanced Chromatography Technologies) using an acetonitrile gradient with an aqueous phase of 100 mM triethylammonium acetate pH...
Figure 2. The molecular structure of Kt-23. (A) Parallel-eye stereoscopic view of the structure of Kt-23 taken from the crystal structure of the *T. thermophilus* 30S ribosomal subunit. The view is from the side of the strand not containing the three-nucleotide loop. The NC helix is coloured green, the C helix grey and the nucleotides of the loop purple. Hydrogen bonds forming the base pairs of the NC helix are shown as broken lines. All images of the Kt-23 structure were made using PDB file 1J5E (22). (B) Superposition of the crystallographic structures of Kt-7 and *T. thermophilus* Kt-23. The two structures were superimposed by their backbone atoms (O5', C5', C4', C3', O3', P and the non-bridging phosphate O). The structures are in parallel-eye stereoscopic view, with ribbons to indicate the position of the backbones. Kt-23 is drawn in red and Kt-7 in blue. The structure of Kt-7 was taken from the *H. marismortui* 50S ribosomal subunit, PDB file 1FFK (30). (C) Base pairing in the NC helix of *T. thermophilus* Kt-23. Parallel-eye stereoscopic view of the NC helix of Kt-23, with hydrogen bonds forming the base pairs of the NC helix are shown as broken lines. The structures of the four non-Watson–Crick base pairs are drawn out.
7.0. Oligonucleotides required for electrophoretic analysis were radioactively \[^{32}P\]-labelled using T4 polynucleotide kinase and \[^{32}P\] ATP (Amersham).

Duplex species were prepared by mixing equimolar quantities of the appropriate oligoribonucleotides and annealing them in 50 mM Tris–HCl (pH 7.5), 25 mM NaCl by slow cooling from 90 to 4 °C. They were purified by electrophoresis in 12% polyacrylamide under non-denaturing conditions, and recovered by either electroelution or diffusion into buffer, followed by ethanol precipitation.

Analysis of k-turn folding by comparative gel electrophoresis

Radioactively \[^{32}P\]-labelled RNA species were electrophoresed in 13% polyacrylamide (29:1, acrylamide:bis) gels in 90 mM Tris–borate (pH 8.3), 2 mM Mg\(^{2+}\). Electrophoresis was performed at 120 V at room temperature for up to 68 h, with recirculation of the buffer at >1 h\(^{-1}\). Gels were dried onto Whatman 3 MM paper, exposed to storage phosphor plates and imaged using a Fuji BAS-1500 phosphorimager.

The sequences used for the electrophoretic experiments were (written 5’ to 3’) as follows.

(i) Kt-7 upper strand: CGCAAGCGACAGGAACCT CGCCAGUCAGUCCGGCAAGAAACAGUGACAG GGGACTGTCAAGTTGACAGG
(ii) Kt-7 lower strand: CCTGTCAACTTGAACGT CCCCUGACAUCCGGCAAGAGAGGAGCAGACUGGCG AGGTTCTGTGCTTGCG
(iii) Kt-23 upper strand: CGCAAGCGACAGGAACCT CGCCAGUCAGUCCGGCAAGAAACAGUGACAG GGGACTGTCAAGTTGACAGG
(iv) Kt-23 lower strand: CCTGTCAACTTGAACGT CCCCUGACAUCCGGCAAGAGAGGAGCAGACUGGCG AGGTTCTGTGCTTGCG

The DNA sections of these oligonucleotides are shown underlined. Modified nucleotides were introduced into the RNA sections as required.

Study of Mg\(^{2+}\)-induced folding by fluorescence resonance energy transfer

Absorption spectra were measured in 90 mM Tris–borate (pH 8.3) in 120 µl volumes using a Cary 1E spectrophotometer. Spectra were deconvoluted using a corresponding RNA species labelled only with Cy3, and fluorophore absorption ratios calculated using a MATLAB program. Fluorescence spectra were recorded in 90 mM Tris–borate (pH 8.3) at 4 °C using an SLM-Aminco 8100 fluorimeter. Spectra were corrected for lamp fluctuations and instrumental variations, and polarization artefacts were avoided by setting excitation and emission polarizers crossed at 54.7°. Values of fluorescence resonance energy transfer (FRET) efficiency (\(E_{\text{FRET}}\)) were measured using the acceptor normalization method (29) implemented in MATLAB. \(E_{\text{FRET}}\) as a function of metal ion concentration was analyzed on the basis of a model in which the fraction of folded molecules corresponds to a simple two-state model for ion-induced folding, i.e.

\[
E_{\text{FRET}} = E_0 + \Delta E_{\text{FRET}} \cdot [M]/(1 + K_A [M])
\]

where \(E_0\) is the FRET efficiency of the RNA in the absence of added metal ions, \(\Delta E_{\text{FRET}}\) is the increase in FRET efficiency at saturating metal ion concentration, [M] is the prevailing Mg\(^{2+}\) or Na\(^+\) ion concentration, \(K_A\) is the apparent association constant for metal ion binding and \(n\) is a Hill coefficient. Data were fitted to this equation by non-linear regression. The metal ion concentration at which the transition is half complete is given by [M]\(_{1/2}\) = (1/\(K_A\))^\(1/n\).

The sequences used in the FRET analyses were (written 5’ to 3’) as follows.

(i) Kt-23 upper strand: Fluorescein-CCAGUCAGAUG CGCGAUACCCGGAGAGG
(ii) Kt-23 lower strand: Cy3-CCUCUCCCGGAGUAGCAUCUGACUGG

Single nucleotide substitutions were introduced as required.

RESULTS

Analysis of the ion-induced folding of Kt-23 into k-turn conformation

We can readily detect the formation of the k-turn in RNA by virtue of the markedly increased kinking of the helical axis with folding. In this study, we have explored the ion-induced folding of Kt-23 and variants using two approaches, as in our previous studies of k-turns (13,19,20). In comparative gel electrophoresis, we place the Kt-23-containing RNA into the centre of a 65 bp duplex, of which the outer 20 bp on each side are DNA for synthetic convenience. When the RNA adopts the k-turn geometry, the axial kinking results is a marked retardation of electrophoretic mobility in a polyacrylamide gel. The mobilities can be compared with duplexes of the same length containing oligoadenine bulges of different length, that exhibit progressively greater kinking and hence retardation (25,26). The second method is FRET. This employs a 26 bp RNA duplex that includes a central Kt-23 sequence, with fluorescein and Cy3 fluorophores attached to the 5’-termini. Formation of the k-turn geometry significantly
reduces the separation between the fluorophores, and thus leads to an increase in the efficiency of FRET. All the experimental work described here has been directed at *T. thermophilus* Kt-23 and its derivatives. Kt-23 is induced to fold into the kinked geometry by the presence of metal ions. Using comparative gel electrophoresis, we have previously shown that Kt-7, whose sequence is close to the k-turn consensus, exhibits a mobility similar to a corresponding duplex with a 7-nt bulge. Electrophoresis of Kt-7 and Kt-23 side by side in the presence of 2 mM Mg$^{2+}$ ions shows them to have closely similar mobility, suggesting that they fold to similar extents under these conditions (Figure 3). Ion-induced kinking of Kt-23 can also be observed by FRET (Figure 4), with a change in FRET efficiency ($E_{\text{FRET}}$) of 0.25 in both Mg$^{2+}$ and Na$^+$ ions. The data are well fitted by a two-state folding model. Mg$^{2+}$-induced folding is consistent with non-cooperative ion binding (Hill coefficient $n = 1.2$), while that for Na$^+$-induced folding indicates more cooperative binding of the monovalent ions ($n = 1.7$) although this is not well defined by the data. The half magnesium concentration ([Mg$^{2+}$]$_{1/2}$) obtained by fitting to the two-state folding model (Table 2) is 35 μM, approximately half the value for Kt-7 (19), indicating that the k-turn geometry of Kt-23 is a little more stable than that of Kt-7 (i.e. it requires a lower concentration of Mg$^{2+}$ ions to allow folding into the kinked geometry). However, the [Na$^+$]$_{1/2}$ values for Kt-23 and Kt-7 are closely similar. Both the

Figure 3. Electrophoretic analysis of Mg$^{2+}$-induced kinking of Kt-23 and variants. The scheme (top) shows the 65 bp DNA/RNA/DNA duplex species with a central k-turn sequence used in this analysis. Radioactively [$^{5-32}$P]-labelled k-turn species (with or without modifications in the k-turn) were electrophoresed in 13% polyacrylamide gels in 90 mM Tris-borate (pH 8.3) containing 2 mM Mg$^{2+}$. A$_n$-bulge-containing duplexes were made using the same lower strand, hybridized to complement the non-Watson-Crick pairings, and with the three-nucleotide bulge replaced by an A$_1$, A$_3$ or A$_5$ bulge. A mixture of the three species was electrophoresed in track 1, alongside Kt-7 in track 2. The remaining tracks contain: track 3, unmodified Kt-23; track 4, Kt-23 U2nG; track 5, Kt-23 G3nA; track 6, Kt-23 A4nU; track 7, Kt-23 U2nC; track 8, Kt-23 G1bU; track 9, Kt-23 GL12H; track 10, Kt-23 AL32H. The samples were electrophoresed in two equivalent gels, with the samples Kt-23 U2nG and G3nA common to both.

**Figure 4.** Analysis of ion-induced folding of Kt-23 and variants studied by FRET. The scheme (top) shows the 26 bp RNA duplex with a central Kt-23 sequence, with fluorescein (F) and Cy3 (C) fluorophores attached to the 5' termini. Base substitutions were introduced into this sequence as required. Plots of FRET efficiency ($E_{\text{FRET}}$) as a function of ionic concentration are presented, fitted to two-state transitions (lines). Symbols: closed circles, unmodified Kt-23; open circles, Kt-23 U2nG; closed squares, Kt-23 G3nA; open squares, Kt-23 A4nU; closed triangles, Kt-23 U2nC; open triangles, Kt-23 A4nU. (A) Plot of $E_{\text{FRET}}$ as a function of Mg$^{2+}$ ion concentration. (B) Plot of $E_{\text{FRET}}$ as a function of Na$^+$ ion concentration.
Table 2. Summary of FRET data for folding of Kt-23 and variant k-turns induced by Mg2⁺ and Na⁺ ions

|            | [Mg²⁺]₁/₂ (µM) | ΔE₁₁ | [Na⁺]₁/₂ (mM) | ΔE₂₁ |
|------------|----------------|------|---------------|------|
| Kt-23      | 35             | 0.25 | 32            | 0.25 |
| Kt-7        | 78             | 0.33 | 30            | 0.33 |
| Glb U2nG   | 97             | 0.27 | 26            | 0.22 |
| U2nC       | 97             | 0.27 | 26            | 0.22 |
| G₃₄ A      | 770            | 0.034|               |      |
| G₃₄ inosine| 560            | 0.095|               |      |
| A₄n U      | 290            | 0.10 | 240           | 0.12 |
| A₄n G      | 140            | 0.23 | 68            | 0.16 |
| GL1 2'H    | 120            | 0.12 |               |      |
| AL3 2'H    | 120            | 0.12 |               |      |
| GL1 inosine| 120            | 0.12 |               |      |
| C-2n 2'H   | 120            | 0.12 |               |      |
| U3b 2'H    | 120            | 0.12 |               |      |
| G-1n 2'H   | 120            | 0.12 |               |      |

ΔE₁₁ was measured as a function of ionic concentration and the data fitted to a two-state model in which folding occurs as a result of ion binding (Equation 1). NF, no folding detected.

aData for Kt-7 are taken from Liu & Lilley (19). The absolute value of ΔE₁₁ is not comparable between Kt-7 and Kt-23 since differences in the exact structure adopted in the helical arms could alter the relative placement of the fluorophores.

The ability of the T. thermophilus Kt-23 sequence to adopt the k-turn geometry in the presence of Mg²⁺ ions, unaided by protein binding, despite the presence of the A₄nU pair at the 2b·2n position raises an interesting question. Since the introduction of an A₄nU pair at the corresponding position of Kt-7 completely prevents ion-induced folding (13), what is different about the sequence of Kt-23 that tolerates this change? This is likely to be the sequence context. In both the cases, the 1b·1p pair is G·A, and this cannot be changed in either Kt-7 (13,20) or Kt-23 (Figure 3; Table 2). On the kink-distal side of the 2b·2n pair in H. marismortui Kt-7, there is a further A₄nG pair at the 3b·3n position, followed by Watson–Crick pairing. By contrast, in T. thermophilus Kt-23 there is a U₄nG pair at 3b·3n and an A₄A pair at 4b·4n. However, in the E. coli K-23 these base pairs are Hoogsteen/sugar edge G·G and Watson–Crick A–U, respectively.

The U₄nG pair at the 3b·3n position of T. thermophilus Kt-23 is a Hoogsteen/sugar edge pair that forms a single hydrogen bond donated from G₃₄:n2 to U₃b:o4. It significantly widens the helix, with a C₁′−C₁′ distance of 11.1 Å, compared to 10.0 Å in Kt-7 (Table 1). G₃₄:n substitution removes the 2-NH₃ thus preventing formation of the hydrogen bond. This substitution almost completely prevented folding into the k-turn geometry as judged by both comparative gel electrophoresis (Figure 3) and FRET (Figure 4; Table 2). This could result from formation of a Watson–Crick U–A base pair, but selective removal of the 2-NH₃ from the guanine (inosine substitution, i.e. G₃₄:n) also led to extremely impaired in k-turn folding (Table 2).

The nature of the base pair at the 4b·4n position of Kt-23 is less important

The 4b·4n base pair is usually Watson–Crick in the majority of k-turns. The cis Watson–Crick/Watson–Crick A₄A pair in T. thermophilus Kt-23 again widens the helix, with a C₁′−C₁′ distance of 12.3 Å compared to 10.4 Å in Kt-7 (Table 1). The adenine bases are connected by a single hydrogen bond donated from A₄b:n6 to A₄n:n1. Restoration of a potential Watson–Crick U–A base pair by an A₄nU substitution significantly impaired folding, with a major loss of electrophoretic retardation (Figure 3), and ΔE₁₁ = 0.10 with a [Mg²⁺]₁/₂ = 290 µM (Table 2). Disruption of the A₄A pair by A₄nG substitution (creating a potential A₄G pair) was less disruptive, with ΔE₁₁ = 0.23 with a [Mg²⁺]₁/₂ = 140 µM. In general, it appears that changing the 4b·4n base pair impairs the folding of Kt-23 less than substitutions.
at the 3b3n pair. In the *E. coli* Kt-23, this position has a regular A–U base pair.

**Critical contacts arising from the bulge loop are very important in Kt-23**

The two most conserved hydrogen bonds in crystallographic structures of k-turn RNA involve 2'-hydroxyl groups of the bulge (19). These are formed between the L1 ribose 2'OH and the N1 of the adenine at the invariant 1n position, and between the L3 ribose O2' and the proS O of the L1/L2 phosphate. Both are present in the Kt-23 crystal structures (22,23) (Figure 5A), with distances of 2.7 (2.6) Å and 3.0 (2.7) Å, respectively (*E. coli* values in parenthesis), and with near-tetrahedral geometries in both the cases. Removal of the O2' (by deoxyribose substitution) at either L1 or L3 led to almost complete loss of electrophoretic retardation (Figure 3), and no k-turn folding detectable by FRET (Table 2). We conclude that both interactions are critical to the stability of the k-turn geometry of Kt-23 in the presence of metal ions.

**A possible L1–1n base–base interaction**

In *T. thermophilus* Kt-23, the nucleobase of A1n makes another hydrogen bond. A1n:N3 accepts a proton from GL1:N2H, with a length of 3.3 Å and a near-tetrahedral angle of 114° (Figure 5A) (22). This plainly cannot be a universal interaction, because the L1 nucleotide is not constrained to be guanine in other Kt-23 sequences, and even when L1 is guanine the distance is often long. For example, this interaction is also present in the *E. coli* Kt-23, but the putative bond length is 3.5 Å (23). We examined the importance of this in Kt-23 by replacing

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**Figure 5.** Hydrogen bonding in the Kt-23 k-turn. Parallel-eye stereoscopic views of the structure of Kt-23 taken from the crystal structure of the *T. thermophilus* 30S ribosomal subunit, with key hydrogen bonds indicated by broken lines. **(A)** Hydrogen bonding in the core of the k-turn. The A1n nucleotide is highlighted in yellow. This is held by four hydrogen bonds; in addition to the two bonds with G1b that form the *trans* sugar edge/Hoogsteen edge base pair, there is the critical, conserved A1n:N1–GL1:O2'H interaction, and the A1n:N3–GL1:N2H hydrogen bond. The hydrogen bond between AL3:O2'H and the L1/L2 phosphate proS O is also visible in this image. **(B)** Hydrogen bonding between the C and NC helices. Two hydrogen bonds are shown. One is from G-1n:O2'H to A2b:N1, and between U3b:O2' and C-2n:O2'H.
the guanine by inosine (GL11), selectively removing the exocyclic amine that is the putative donor. The modified RNA exhibited no ion-induced folding detectable by FRET (Table 2), consistent with an important role for this interaction. This hydrogen bond plus the L1:2’O to A1n:N1 bond together hold the nucleobase of adenine 1n by both ring nitrogen atoms. Perhaps its tight positioning becomes important in Kt-23 in order to hold A2b (on which it is stacked) in place in the absence of the A=G pair at 2b•2n.

A base–backbone interaction in Kt-23 involving A2b

Inspection of the structures of both T. thermophilus and E. coli Kt-23 in the crystal suggested a further hydrogen bonding interaction from the 2’-hydroxyl of G-1n and A2b:N1 (Figure 5B). This has a length of 2.6 Å and almost perfect tetrahedral geometry (110°) in both structures. Removal of the 2’-hydroxyl group of G-1n led to impaired folding, with $\Delta E_{\text{FRET}} = 0.12$ (Table 2). Formation of this hydrogen bond would help to fix the position of the adenine nucleobase at 2b; this again could assume greater importance in the absence of the usual A=G pair at 2b•2n.

Possible inter-helical hydrogen bonding in Kt-23

In studies of other k-turns, we have noted that there is hydrogen bonding between the C and NC stems observed in crystal structures, mainly involving 2’-hydroxyl groups (19). However these are variable from one structure to another. In one class (exemplified by Kt-7), there is bonding between O2’ of –2n and 3b, and in the other class (a bigger group, that includes the box C/D k-turn) there is a hydrogen bond formed between O2’ of –1n and 2b. However, removal of selected 2’-hydroxyl groups has suggested that none of these interactions is important to the stability of the k-turn (19). In the Kt-23 crystal structures, there is a hydrogen bond between U3b:O2’ and C-2n:O2’ with a distance of 2.7 Å (T. thermophilus) or 3.1 Å (E. coli) and excellent tetrahedral geometry (Figure 5B). We investigated the importance of this interaction to the stability of the k-turn by removing the 2’-hydroxyl from U3b. The resulting RNA folded well, with a $\Delta E_{\text{FRET}} = 0.3$, and a slightly elevated requirement for Mg$^{2+}$ ions ([Mg$^{2+}$]$_{1/2}$ = 130 μM) (Table 2). Removal of the 2’-hydroxyl at C-2n led to relatively unperturbed folding, with $\Delta E_{\text{FRET}} = 0.29$ and [Mg$^{2+}$]$_{1/2}$ = 240 μM. As with other k-turns (19), it appears that the interhelical hydrogen bonding outside the core region is not very important for the stability of the kinked geometry of Kt-23. In general, these hydrogen bonds form adventitiously, but contribute little stabilization to k-turn conformation.

DISCUSSION

Since Kt-23 breaks the seemingly critical requirement for an A=G pair at the 2b•2n position in both T. thermophilus and E. coli, it might have been thought that it would be unable to adopt the k-turn conformation unaided. While both Kt-23 sequences clearly adopt normal k-turn structure in the context of the 30S ribosomal subunits, we suspected that either long-range tertiary interactions, protein-RNA interactions or both might provide sufficient stabilization to overcome an inherent instability. Our results show that that is not the case; T. thermophilus Kt-23 can adopt the k-turn structure in a simple RNA duplex in the absence of any proteins and with no possibility for tertiary structure. Stable k-turn formation only requires charge neutralization by either divalent or monovalent metal ions. Indeed, folding of Kt-23 occurs at a lower ion concentration compared to Kt-7 with its near canonical k-turn sequence, indicating than the Kt-23 is if anything more stable as a k-turn than is Kt-7. This is supported by a consistently greater electrophoretic retardation of Kt-23 compared to Kt-7, although the effect is small.

This surprising result shows that the sequence requirements for k-turn stability are more complex than we previously supposed. It leaves a question outstanding. Why can Kt-23 adopt the k-turn conformation despite possessing an A•U pair at the 2b•2n position, when conversion of the 2b•2n position in Kt-7 to A•U prevents any detectable ion-induced folding? Clearly, some aspect of the sequence environment of Kt-23 permits the RNA to form a stable k-turn despite the A•U pair at the 2b•2n position, although this does not prevent folding from occurring when an A=G pair is restored at this site. Kt-23 makes the previously identified key hydrogen-bonding interactions within the core of the k-turn involving the 2’-hydroxyl groups of the loop nucleotides L1 and L2 (19). A2b also makes an important interaction with the ribose of G-1n, with a hydrogen bond from G-1n:2’OH to A2b:N1. The A•U pair does not make any additional direct contacts with its neighbours in the NC helix, so perhaps the overall geometry of the NC helix is better able to accommodate this pair. We have shown that the nature of both the 3b•3n and 4b•4n pairs can influence the stability of the k-turn conformation of T. thermophilus Kt-23. The trans Hoogsteen/Watson–Crick edge A=G pair is significantly wider than the usual A=G pair (Table 1), and it is possible that the additional width of the Hoogsteen/sugar edge U=G pair at the 3b•3n position is an important factor. The Hoogsteen/sugar edge G=G pair at the corresponding position in E. coli Kt-23 is also significantly wider, at 11.4 Å. Our results suggest that the nature of the base pair at the 3b•3n position is more important than that at the 4b•4n position, and that changes that should destabilize this base pair markedly impair folding into the k-turn geometry. Normal k-turn sequences with an A=G pair at the 2b•2n position appear less sensitive to the sequence on the bulge-distal side; in the U4 snRNA k-turn there are conventional Watson–Crick G–C base pairs at both 3b•3n and 4b•4n positions (6).

Kt-23 reveals a surprising tolerance to departure from the consensus sequence. It folds into a near perfect k-turn geometry that places the critical 1n and 2b adenine bases into the required positions. The key hydrogen-bonding interactions in the core of the k-turn are preserved. The nucleobase at position 2 is exposed on the splayed-out major groove face of the k-turn, and most generally it is
this face that interacts with L7Ae (2) and related (6) proteins. L7Ae can induce k-turn folding upon binding to Kt-7 and other k-turns (15), and so it is possible that the folding of k-turns (such as Kt-23) that differ in this region might be induced by different proteins. We are in the process of investigating this possibility.

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REFERENCES

1. Klein,D.J., Schmeing,T.M., Moore,P.B. and Steitz,T.A. (2001) The kink-turn: a new RNA secondary structure motif. EMBO J., 20, 4214–4221.
2. Moore,T., Zhang,Y., Fenley,M.O. and Li,H. (2004) Molecular basis of box C/D RNA-protein Interactions; Cocystal structure of archaeal L7Ae and a box C/D RNA. Structure, 12, 807–818.
3. Hamma,T. and Ferre ´-D’Amare´,A.R. (2004) Structure of protein L30 bound to a K-turn derived from an archaeal box H/ACA sRNA at 1.8 A˚ resolution. Structure, 12, 893–903.
4. Wozniak,A.K., Nottrott,S., Kuhn-Holsken,E., Schroder,G.F., Grubmuller,H., Luhmann,R., Seidel,C.A. and Oesterhelt,F. (2005) Detecting protein-induced folding of the U4 snRNA kink-turn by single-molecule multiparameter FRET measurements. RNA, 11, 1545–1554.
5. Mao,H., White,S.A. and Williamson,J.R. (1999) A novel loop-loop recognition motif in the yeast ribosomal protein L30 autoregulatory RNA complex. Nature Struct. Biol., 6, 1139–1147.
6. White,S.A., Hoeger,M., Schwegel,J., Shillingford,A., Shiplov,V. and Zarutskie,J. (2004) Internal loop mutations in the ribosomal RNA guide RNP. RNA, 10, 369–377.
7. Montange,R.K. and Batey,R.T. (2006) Structure of the S-adenosylmethionine riboswitch regulatory mRNA element. Nature, 441, 1172–1175.
8. Blouin,S. and Lafontaine,D.A. (2007) A loop loop interaction and a K-turn motif located in the lysine aptamer domain are important for the riboswitch gene regulation control. RNA, 13, 1256–12567.
9. Reblock,J., Schvetsov,B. and Lafontaine,D.A. (2008) Folding of the SAM aptamer is determined by the formation of a K-turn-dependent pseudoknot. Biochemistry, 47, 1490–1499.
10. Goody,T.A., Melcher,S.E., Norman,D.G. and Lilley,D.M.J. (2003) The kink-turn motif in RNA is dimorphic, and metal ion dependent. RNA, 10, 254–264.
11. Matsumura,S., Ikawa,Y. and Inoue,T. (2003) Biochemical characterization of the kink-turn RNA motif. Nucleic Acids Res., 31, 5544–5551.
12. Turner,B., Melcher,S.E., Wilson,T.J., Norman,D.G. and Lilley,D.M.J. (2005) Induced fit of RNA on binding the L7Ae protein to the kink-turn motif. RNA, 11, 1192–1200.
13. Razga,F., Koca,J., Sponer,J. and Leontis,N.B. (2005) Hinge-like movements in RNA Kink-turns: The role of the second A-minor motif and nominally unpaired bases. Biophys. J., 5, 3466–3485.
14. Cojocaru,V., Klement,R. and Jovin,T.M. (2005) Loss of G-A base pairs is insufficient for achieving a large opening of U4 snRNA K-turn motif. Nucleic Acids Res., 33, 3435–3446.
15. Liu,J. and Lilley,D.M.J. (2007) The role of specific 2’-hydroxyl groups in the stabilization of the folded conformation of kink-turn RNA. RNA, 13, 200–210.
16. Razga,F., Zacharias,M., Reblova,K., Koca,J. and Sponer,J. (2006) RNA kink-turns as molecular elbows: hydration, cation binding, and large-scale dynamics. Structure, 14, 825–835.
17. Wimberly,B.T., Brodersen,D.E., Clemens,W.M. Jr, Morgan-Warren,R.J., Carter,A.P., Vounrhein,C., Hartsch,T. and Ramakrishnan,V. (2000) Structure of the 3OS ribosomal subunit. Nature, 407, 327–339.
18. Razga,F., Koca,J., Sponer,J. and Leontis,N.B. (2005) Hinge-like motions in RNA Kink-turns: The role of the second A-minor motif and nominally unpaired bases. Biophys. J., 5, 3466–3485.
19. Hamma,T. and Ferre ´-D’Amare´,A.R. (2004) Structure of protein L30 bound to a K-turn derived from an archaeal box H/ACA sRNA at 1.8 A˚ resolution. Structure, 12, 893–903.
20. Wozniak,A.K., Nottrott,S., Kuhn-Holsken,E., Schroder,G.F., Grubmuller,H., Luhmann,R., Seidel,C.A. and Oesterhelt,F. (2005) Detecting protein-induced folding of the U4 snRNA kink-turn by single-molecule multiparameter FRET measurements. RNA, 11, 1545–1554.
21. Nissen,P., Ippolito,J.A., Ban,N., Moore,P.B. and Steitz,T.A. (2001) RNA tertiary interactions in the large ribosomal subunit: The A-minor motif. Proc. Natl Acad. Sci. USA, 98, 4899–4903.
22. Nissen,P., Ippolito,J.A., Ban,N., Moore,P.B. and Steitz,T.A. (2001) RNA tertiary interactions in the large ribosomal subunit: The A-minor motif. Proc. Natl Acad. Sci. USA, 98, 4899–4903.
23. Bhattacharyya,A., Murchie,A.I.H. and Lilley,D.M.J. (1999) RNA bulges and the helical periodicity of double-stranded RNA. Nature, 343, 484–487.
24. Nissen,P., Ippolito,J.A., Ban,N., Moore,P.B. and Steitz,T.A. (2001) RNA tertiary interactions in the large ribosomal subunit: The A-minor motif. Proc. Natl Acad. Sci. USA, 98, 4899–4903.
25. Razga,F., Koca,J., Sponer,J. and Leontis,N.B. (2005) Hinge-like motions in RNA Kink-turns: The role of the second A-minor motif and nominally unpaired bases. Biophys. J., 5, 3466–3485.
26. Razga,F., Koca,J., Sponer,J. and Leontis,N.B. (2005) Hinge-like motions in RNA Kink-turns: The role of the second A-minor motif and nominally unpaired bases. Biophys. J., 5, 3466–3485.