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Crystal structure of 3WJ core revealing divalent ion-promoted thermostability and assembly of the Phi29 hexameric motor pRNA

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ABSTRACT

The bacteriophage phi29 DNA packaging motor, one of the strongest biological motors characterized to date, is geared by a packaging RNA (pRNA) ring. When assembled from three RNA fragments, its three-way junction (3WJ) motif is highly thermostable, is resistant to 8 M urea, and remains associated at extremely low concentrations in vitro and in vivo. To elucidate the structural basis for its unusual stability, we solved the crystal structure of this pRNA 3WJ motif at 3.05 Å. The structure revealed two divalent metal ions that coordinate 4 nt of the RNA fragments. Single-molecule fluorescence resonance energy transfer (smFRET) analysis confirmed a structural change of 3WJ upon addition of Mg2+. The reported pRNA 3WJ conformation is different from a previously published construct that lacks the metal coordination sites. The phi29 DNA packaging motor contains a dodecameric connector at the vertex of the procapsid, with a central pore for DNA translocation. This portal connector serves as the foothold for pRNA binding to procapsid. Subsequent modeling of a connector/pRNA complex suggests that the pRNA of the phi29 DNA packaging motor exists as a hexameric complex serving as a sheath over the connector. The model of hexameric pRNA on the connector agrees with AFM images of the phi29 pRNA hexamer acquired in air and matches all distance parameters obtained from cross-linking, complementary modification, and chemical modification interference.

Keywords: RNA crystal; metal ions; three-way junction; RNA nanotechnology; RNA therapeutics

INTRODUCTION

The bacteriophage phi29 DNA packaging motor, one of the strongest biological motors assembled in vitro using recombinant and synthetic materials, is geared by a packaging RNA (pRNA) ring (Guo et al. 1987, 1998; Chen et al. 2000; Shu et al. 2007; Xiao et al. 2008). Since its discovery in 1987 (Guo et al. 1987), the 120-nucleotide (nt) pRNA subunit has been extensively investigated (Reid et al. 1994b,c; Zhang et al. 1994, 1995a,b,c; 1997; Chen and Guo 1997; Chen et al. 2000). The unique features uncovered from the studies of phi29 pRNA include (1) independent folding of the ATPase gpl6 interaction domain (Lee and Guo 2006) and the motor binding domain (Reid et al. 1994a,b,c; Zhang et al. 1994, 1995b,c) and (2) the capacity to harbor and escort therapeutic functional modules without causing misfolding or loss of function of the incorporated functionalities. These special properties of phi29 pRNA have led to the development of this RNA as a novel vehicle for applications in nanotechnology and medicine (Hoeprich et al. 2003; Guo et al. 2005a, 2006, 2012a,b; Khaled et al. 2005; Guo 2010; Abdelmawla et al. 2011; Shu et al. 2011a,b,c; Shukla et al. 2011; Zhou et al. 2011). Recently, it was found that the core structure of phi29 pRNA, a 54-nt three-way junction (3WJ) (Fig. 1A), exhibits extreme thermodynamic stability (Shu et al. 2011a). This trifurcate domain of pRNA can be assembled from three short RNA fragments (<20 mer) with high affinity and low free energy, as demonstrated by its melting curve with a slope of ~90°. Nevertheless, the fundamental mechanism governing the high thermodynamic stability of the 3WJ has been elusive.

The phi29 3WJ has been utilized as a useful platform for the construction of stable RNA nanoparticles. Cell receptor-binding ligands, aptamers, siRNAs, or ribozymes fused to individual arms of this 3WJ motif fold into their authentic
structures and retain their functions such as specific cancer cell binding, cell entry, gene silencing, catalytic function, and in vivo targeting of cancer cells without accumulating in normal organs and tissues (Abdelmawla et al. 2011; Shu et al. 2012; Haque et al. 2012), indicating that the 3WJ core with low ΔG guides the folding of the entire RNA particle to produce its global structure. The important application of the 3WJ in nanotechnology and medicine inspired the interest in further studies on pRNA 3WJ crystal structure. Recently a crystal structure of a part (66 nt) of the phi29 pRNA sequence (120 nt) (Fig. 1B; Ding et al. 2011) has been reported. In their studies, one of the key helices of the 3WJ core (5′-U21U22G23U24-3′/5′-A99A98C97G96-3′) was truncated, several bases of the 3WJ motif were mutated, and five other nucleotides at three different locations were also removed (Fig. 1B). As a result, the right-hand interlocking loop motif was changed from 9 nt to 6 nt, and the left interlocking loop motif was changed from 5 nt to 4 nt. More importantly, two of the 4 nt responsible for the coordination of Mg$^{2+}$ reported here were excluded (Fig. 1B). The truncation, mutation, and alteration of the 3WJ core structure in pRNA molecule may lead to the production of a crystal structure that deviates significantly from the authentic pRNA fold, and therefore, further independent studies on the intact 3WJ crystal structure are necessary.

In this study, we report the crystallization, structure solution, and Mg$^{2+}$-dependent fluorescence resonance energy transfer (FRET) analysis of the pRNA 3WJ, which likely represents the authentic structure of the metal-bound 3WJ core. Crystals were obtained in the presence of either Mg$^{2+}$ or Mn$^{2+}$. The crystal structure revealed two well-coordinated metal ions and a tightly packed core junction structure. To ensure that this result is not a crystal artifact, the effect of the divalent metal ions on the thermodynamic stability and solution structure (via FRET analysis) of pRNA 3WJ was also investigated. Additionally, the crystal structure of metal-coordinated 3WJ was used as a template to build a 3D model of intact, 120-nt monomeric pRNA. Model building indicated that the 120-nt pRNA monomers were capable of assembling into a hexameric pRNA ring, which was constructed and compared with that of AFM images and the previously published hexameric assembly (Hoeprich and Guo 2002), the latter of which was modeled based on secondary structural analysis and distant parameters from numerous biochemical studies. The intact pRNA hexamer model was docked with the 3D structure of the dodecameric ring formed.
by the phi29 connector protein. The resulting models were further evaluated by comparison with distance parameters obtained from single-molecule FRET (smFRET) and additional measurements by photoaffinity crosslinking (Garver and Guo 1997), chemical modification interference (Trottier et al. 2000; Mat-Arip et al. 2001), complementary modification (Zhang et al. 1997), and atomic force microscopy (Chen et al. 2000; Trottier et al. 2000; Mat-Arip et al. 2001).

RESULTS AND DISCUSSION

3WJ overall structure

The 3WJ crystal structure was solved at a resolution of 3.05 Å by piecwise molecular replacement (Table 1; Supplemental Fig. S1B; Robertson and Scott 2008; Robertson et al. 2010). The structure of the 3WJ core is facilitated by “coaxial stacking” of three individual RNA helices. This type of stacking interaction is commonly found in RNA, ribosome, and tRNA junction structures (Batey et al. 1999; Laing et al. 2012). The overall structure presented here is similar to the 3WJ region previously reported for the pRNA_{25-95} crystal structure (Fig. 1C; Ding et al. 2011); however, the H1/H3 interhelical angle is different in the two structures, at least partly due to the presence of metal binding sites that were not included in the previous pRNA_{25-95} construct (see the next section) (Fig. 1C). Helices joined at the 3WJ largely possessed the characteristics of canonical A-form helix but have distortions from A-form geometry mediated by chelating metal ions. Confirmation that metal binding alters the conformation of the 54-nt 3WJ structure was provided through FRET analysis of 3WJ in the presence and absence of Mg^{2+} (see the section of smFRET studies). At the core of the 3WJ domain is the junction formed by the intersection of three RNA helices, whose conformation, including that of the UUU bulge (Fig. 1D,E), dictated the relative orientation of three domains connected by RNA helices H1, H2, and H3. The metal-bound 3WJ presented here displayed an overall packing at the core junction with smaller void volumes and additional putative bonding interactions than that of pRNA_{25-95} (Fig. 1E). In the 3WJ structure, atoms O1P and O2P of nucleotide G28 lie 3.5 Å and 3.6 Å from atom O2 of unpaired base U74, while coaxial packing between U74 and U29 and the U-U (U29-U72) mismatch exhibit near ideal geometries (Fig. 1E).

Both crystal structures (3WJ and pRNA_{25-95}) possessed a mismatched base pair between nucleotides U29 and U72 (Fig. 1D,E), and a wobble pair between residues G75 and U91. The conformation of the 3WJ is influenced by remote Mg^{2+} (or Mn^{2+}) cations, which could induce a conformational shift of the pRNA to comply with the structure requirement for the function of the motor in DNA packaging (Chen et al. 2000).

Divalent metal ion (Mg^{2+} or Mn^{2+}) coordination of 3WJ

One of the striking features of the 3WJ domain structure was the presence of divalent metal binding sites between helices H1 and H3 (Fig. 1C,E,G). The pRNA 3WJ was coordinated by two divalent metal ions bridging four phosphates (two from helix H1 and two from helix H3) such that the contact distance between phosphates donated by adjacent helices was reduced to 4.5 Å, with a relatively short 2.25 Å average metal-phosphate distance that was consistent with Mn^{2+}/Mg^{2+} binding (Fig. 1F). Bound metal atoms were initially identified via 10–12 σ peaks in the [F_o]-[F_c] difference density map. Although both Mn^{2+} and Ca^{2+} were present in the crystallization conditions, the putative metal sites that bridge adjacent phosphate groups were confirmed as Mn^{2+} by anomalous difference Fourier analysis at a wavelength where Ca^{2+} has no measurable signal (Fig. 1F, inset). Therefore, the 3WJ structure deposited in the Protein Data Bank (PDB) contains Mn^{2+} in the phosphate bridging metal sites. However, low-resolution studies using isomorphous crystals grown with MgCl_{2} in place of MnCl_{2} suggest magnesium binding to the same sites in the 3WJ construct (data not shown).

In the final model, one Mn^{2+}/Mg^{2+} ion was chelated by the phosphates of nucleotides G23 and A90, while the other

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**TABLE 1. Data collection and refinement statistics**

| Data collection |  |
|-----------------|---|
| Space group     | 14 |
| Cell dimensions |  |
| a, c (Å)        | 125.01, 26.99 |
| Wavelength (Å)  | 0.98789 |
| Resolution (Å)  | 30.00–3.05 (3.15–3.05)* |
| No. of measured reflections | 42,020 |
| No. of unique reflections | 4217 |
| Rmerge (%)      | 8.8 (53)* |
| I/σ(I)          | 30.3 (3.2)* |
| Completeness (%)| 99.5 (99.4)* |
| Multiplicity    | 10.0 (9.1)* |

**Refinement**

| Resolution (Å)  | 30.00–3.05 |
| No. reflections | 4201 |
| Rwork/Rfree (%) | 18.2/19.7 |
| No. of RNA and non-RNA atoms | 1134/5 |
| RNA and non-RNA ADPs (Å) | 108/56.4 |
| ML-based error estimates |  |
| Coordinate error (Å) | 0.28 |
| Phase error (°) | 22.43 |
| Average B-factors (Å²) | 108.0 |
| RNA ions/waters | 88.0/57.0 |
| RMS deviations |  |
| Bond lengths (Å) | 0.002 |
| Bond angles (°) | 0.43 |

ADPs indicates atomic displacement parameters; ML, maximum likelihood.

*Highest-resolution shell is shown in parenthesis.

bThe cross-validation (R_free) was calculated with 5% of the data.
Mn$^{2+}$/Mg$^{2+}$ ion bridges the phosphates of C24 and A89 (Fig. 1G). Although the pRNA 3WJ was similar to the core region of the published pRNA35-95 crystal structure (Fig. 1C), the cation-mediated close contact between helices H1 and H3 facilitates distortion from canonical A-form RNA (Supplemental Fig. S1A). The resulting distortion from A-form RNA, in the context of intact pRNA, resulted in readjustment of left- and right-hand interlocking loops such that a model for an intact pRNA multimer could be assembled while conserving base-pairing between the interlocking loops (see the section of construction of hexameric pRNA model). Two of four metal-chelating nucleotides (G23 and C24) were deleted in the construct of the previously determined crystal structure of pRNA35-95 (Fig. 1B; Ding et al. 2011); thus, the divalent ion coordination was not observed. In the 3WJ structure, a metal-induced conformational change in the angles between helices H1 and H3 reposition the left-hand and right-hand loops such that a continuous hexameric pRNA can be assembled with conserved base-pairing between the loops.

**Effect of metal ion on 3WJ structure confirmed by smFRET**

Previous results from various studies, including base deletion and mutation (Bailey et al. 1990; Reid et al. 1994c; Zhang et al. 1994, 1995b, 1997), ribonuclease probing (Reid et al. 1994a; Chen and Guo 1997), oligo targeting (Zhang et al. 1995a), competition assays to inhibit phage assembly (Trottier et al. 1996), UV crosslinking to portal protein (Garver and Guo 1997), psoralen crosslinking, and primer extension (Chen and Guo 1997), indicated two functional domains that could fold independently in pRNA. The two interlocking loops, along with the double-helical packaging domain for the binding of the motor ATPase (Lee et al. 2006), are connected through the 3WJ region in pRNA and a "three-arms around a hinge" model was proposed for pRNA function (Fang et al. 2008). Flexibility at the U72U73U74 bulge region was found to be essential for pRNA activity in gearing the DNA packaging motor (Reid et al. 1994c; Zhang et al. 1997), indicating that a conformational change may be required during DNA packaging.

Conformational transitions are crucial to RNA functions (Al-Hashimi and Walter 2008). smFRET has been widely used to study the ion-induced conformational changes of functional RNAs (Zhuang et al. 2000; Kobitski et al. 2007; Qu et al. 2008; Steiner et al. 2008; Xiao et al. 2008; Brenner et al. 2010; McDowell et al. 2010; Haller et al. 2011). In this study, we investigated the effect of Mg$^{2+}$ on the folding and conformational change of pRNA 3WJ using smFRET. To measure the distance transition, the ends of helices H2 and H3 of the pRNA 3WJ were labeled with a FRET pair (Cy3 and Cy5) and H1 was labeled with biotin for immobilization to the slide (Fig. 2A).

The FRET efficiency between donor Cy3 and acceptor Cy5 at H2 and H3 were compared in the absence and presence of Mg$^{2+}$ (Fig. 2B,C). A reduction in FRET efficiency of 0.14

![Figure 2](image1.png)

**FIGURE 2.** Comparison of distances between the 3WJ H2/H3 distal ends derived from FRET measurements (with and without 10 mM MgCl$_2$) and crystal data. (A) Location of labeling in 3WJ for single-molecule FRET study. (a$_{35}$ indicated by gray sticks; b$_{35}$, black backbone; c$_{35}$, gray backbone.) (B,C) Comparison of the FRET efficiency (E) and distance (C) at 0 mM and 10 mM MgCl$_2$, respectively. The Gaussian curve fittings used to locate the peak of the distributions are shown as black lines. (D) Comparison of the distances from FRET and the crystal structure. The distance was corrected by subtracting the arm sizes of the fluorophore pair (Shiu et al. 2010).
Effect of metal ion on 3WJ structure confirmed by DNA packaging and viral assembly assay

Divalent metal ions play a critical role in stabilizing the tertiary structure of RNA and in facilitating RNA functions (Carter and Holbrook 2002; Draper 2004; Scott 2007; Johnson-Buck et al. 2011). The positively charged metal ions shield the negatively charged phosphate backbone of RNA, rendering the RNA tertiary structure more compact. This interaction between metal ions and the RNA could lead either to the formation of a positively charged cloud around the RNA molecule or to site-specific binding (Draper 2004; Johnson-Buck et al. 2011). The latter effect is often observed in various catalytic RNA structures (Cate et al. 1996, 1997; Cate and Doudna 1996; Murray et al. 1998; Kazantsev et al. 2005, 2009; Vicens and Cech 2006).

Metal ions are required to stimulate an appropriate pRNA conformation for dimer formation, procapsid binding, and DNA packaging (Chen and Guo 1997; Chen et al. 2000). It was found that the optimal Mg$^{2+}$ concentration for phi29 DNA packaging was $>3$ mM (Supplemental Fig. S2). Although the pRNA 3WJ can be assembled from three individual RNA fragments in the absence of metal ions (Shu et al. 2011a), the divalent metal ion sites observed in the 3WJ crystal structure implied that these phosphate-bridged metal coordination may be involved in pRNA function. The specific coordination sites for the Mg$^{2+}$ ions on RNA can be confirmed through the rescue of the activity of phosphothioate-modified RNA by thiophilic metal ions such as Mn$^{2+}$ or Cd$^{2+}$ (Dahm and Uhlenbeck 1991; Peracchi et al. 1997; Basu and Strobel 1999; Kovacheva et al. 2004). However, evaluation by “manganese rescue” of the pRNA activity in DNA packaging and phi29 virion assembly is not feasible since Mg$^{2+}$ is involved in multiple steps of DNA packaging and virion assembly other than facilitating the proper pRNA conformation, as Mn$^{2+}$ can replace Mg$^{2+}$ for the motor to package DNA, albeit at lower efficacy (Chen et al. 2000). In general, the increased stability observed for Mg$^{2+}$-mediated stabilization of RNA over other cations (such as Na$^+$ or Ca$^{2+}$) is facilitated by its small size and high charge density. In addition to favorable electrostatic interactions, the relatively short interatomic distances between the phosphate groups and Mn$^{2+}$/Mg$^{2+}$ in the 3WJ structure (2.17–2.35 Å) can stabilize RNA oxyanion/Mg$^{2+}$ interactions via charge transfer, polarization, and exchange, thus providing a special case of bidentate RNA–Mg$^{2+}$ clamps (Petrov et al. 2011). These findings are consistent with a model in which the pRNA 3WJ (and by extension, intact pRNA hexamers) are stabilized by Mg$^{2+}$ binding at specific sites between the phosphates of helices H1 and H3, optimally orienting the left- and right-hand loops for multimer assembly.

Effect of metal ion on 3WJ structure confirmed by gel shift assay

Urea is a common reagent used to denature DNA and RNA (Carlson et al. 1975; Pagratis 1996). Previous report has compared the assembly of 11 3WJ cores from different biological RNAs and has shown that in the presence of 5 mM MgCl$_2$, the pRNA-3WJ assembly stayed associated in the presence of 8 M urea, while majority of others dissociated under the same condition (Shu et al. 2011a). The stability of the 3WJ complex was further assessed using 15% PAGE gels containing 0 M, 4 M, and 8 M urea both in the presence and in the absence of Mg$^{2+}$ (Fig. 3). Without urea, the 3WJ is stable in both the presence and the absence of 5 mM MgCl$_2$. However, in the presence of 4 M urea, the complex remained intact in the presence of 5 mM MgCl$_2$ while it dissociated in the absence of MgCl$_2$. Moreover, in the presence of 5 mM Mg$^{2+}$, the pRNA 3WJ remained intact up to 8 M urea. The result demonstrated that Mg$^{2+}$ is critical for the resistance to urea denaturation of this pRNA-3WJ core, while it failed to promote the urea resistance in several other 3WJ cores in the previous report (Shu et al. 2011a).

**FIGURE 3.** Effect of magnesium ions on the resistance of pRNA 3WJ to urea denaturation, assayed by 15% PAGE gels with (A) 2 mM EDTA and (B) 5 mM MgCl$_2$. (Lane 1) Single RNA oligo b$_{3WJ}$ (lane 2) sample annealed from two oligos a$_{3WJ}$ and b$_{3WJ}$ (lane 3) 3WJ assembled from three oligos a$_{3WJ}$, b$_{3WJ}$, and c$_{3WJ}$ and (lane 4) DNA ladder.
Construction of a hexamer pRNA model using the 3WJ crystal structure and comparison with AFM images

Extensive investigations on the function and folding of pRNA have been carried out (Reid et al. 1994b,c; Zhang et al. 1994, 1995a,b,c, 1997; Chen and Guo 1997; Chen et al. 2000). Based on previous reports, the pRNA can be divided into the following domains (Fig. 1A), including domain 1, a 5'3'-end double-helix for the binding of the motor ATPase gp16 (Zhang et al. 1994; Lee et al. 2006); domain 2, a stably folded trifurcate domain (3WJ) (Shu et al. 2011a); domain 3, the right-hand interlocking loop for inter-RNA interaction in RNA ring formation via hand-in-hand interactions (Guo et al. 1998; Zhang et al. 1998; Chen et al. 2000); domain 4, the head loop (Reid et al. 1994a,b,c; Guo et al. 1998; Chen et al. 2000; Hoeprich and Guo 2002; Kitamura et al. 2008); and domain 5, the left-hand interlocking loop for inter-RNA interaction in RNA ring formation via hand-in-hand interactions (Guo et al. 1998; Zhang et al. 1998; Chen et al. 2000). The availability of the structure of individual domains makes the assembly of the entire pRNA structure and the ring feasible.

For model structure construction, the crystal structure of the pRNA 3WJ and the published structural data (Hoeprich and Guo 2002; Harris and Schroeder 2010; Ding et al. 2011) were used to build the intact 120-nt pRNA monomer as well as a hexameric pRNA ring (see Materials and Methods) (Figs. 1A, 4A). Briefly, the entire 3WJ structure, including the two metal ions, was used as a central building block for grafting the structures of the other four domains from three PDB files (PDB codes 1L4O, 2KVN, 3R4F) (Hoeprich and Guo 2002; Harris et al. 2010; Ding et al. 2011) to generate an intact 120-nt pRNA (Fig. 1A). The 3WJ served as domain 2, and all other domains were connected to the 3WJ. Domain 1 was grafted from the previous hexameric pRNA model (PDB 1L4O) (Hoeprich and Guo 2002). Domains 3, 4, and 5 were grafted from the crystal structure of pRNA25-95 (PDB 3R4F) (Ding et al. 2011), with missing residues inserted to complete the entire 120-nt pRNA monomer.

The pRNA hexamer was assembled using pRNA with 3-nt extensions at the interlocking loops (Shu et al. 2013b) (see Materials and Methods), which enhances the stability of RNA hexamer, while the wild-type pRNA hexamer is relatively unstable. This re-engineered pRNA binds to procapsid specifically with the same efficiency as the wild-type pRNA. The assembled pRNA hexamer was confirmed by AFM imaging (Fig. 4B), which showed a strong resemblance to the model structure (Fig. 4A). The pRNA ring showed an outer contour diameter of ~30 nm formed through the loop-loop interactions, with a center ring of ~10 nm in diameter to sheath on the connector that has an outer contour diameter of 9.4 nm for its narrower end and 13.8 nm for its wider

![FIGURE 4. Structure of a pRNA hexamer ring and docking of the hexameric pRNA model with the dodecameric connector ring of ph29. (A) A 3D model of hexameric pRNA based on the crystal structure of 3WJ with views from three mutually perpendicular angles. (B) AFM images of hexameric re-engineered pRNA rings show strong correlation in size and shape with the 3D computer models for the intact, full-length pRNA hexamers. (C) Histogram of RNA particles in AFM images with discernible stoichiometry. (D) Top view of the published 3D computer model of hexameric pRNA constructed based on biochemical data (PDB 1L4O) (Hoeprich and Guo 2002). (E) Model of a pRNA hexamer complexed with a dodecameric connector ring based on the 3WJ and connector (PDB 1HSW) (Guasch et al. 2002) crystal structures. (For an animation movie of the ph29 pRNA-connector model, see Supporting Video S1.) (F) A close-up view of proper anchoring of the connector N-terminal helices for optimal interactions with pRNA. (G) Three mutually perpendicular views of the hexameric pRNA-connector assembly. (H) A side-view of the 3D model of the hexameric pRNA-connector assembly based on biochemical data (PDB 1L4P) (Hoeprich and Guo 2002).](https://www.rnajournal.org)
end (Guo et al. 2005b). A comparison of the AFM images with the pRNA hexamer model showed a strong correlation in shape and size, including sixfold rotational axes and protrusions every 60° (these correspond to helix H1, essential for contacting the ATPase). The inner diameter of the hexamer (~9.7 nm) was in good correspondence with the diameter of the pRNA hexamer determined by AFM, supporting the model of the pRNA hexameric ring. This model structure was very similar to the previously published computer model of the pRNA hexamer based on biochemical distance constraints (Fig. 4D).

Docking of the hexamer pRNA model to the motor connector

Connector protein, the foothold for pRNA binding on the phi29 motor, contains a wide end and a narrow end. The wide end is embedded inside the capsid, and the narrow end is exposed for interaction with the pRNA (Jimenez et al. 1986; Simpson et al. 2000; Guo et al. 2005b). The interaction between pRNA and the connector was demonstrated by crosslinking of the two (Garver and Guo 1997). It was found that the N-terminal 14 amino acids of gp10 are essential for pRNA binding since deletion of these 14 amino acids rendered the connector incompetent for RNA binding (Xiao et al. 2005) and mutation in R3K4R5 led to a similar result (Atz et al. 2007). Particularly, these three positively charged amino acids (R3K4R5) are believed to be responsible for the binding to pRNA with the negatively charged phosphate backbone.

Modeling the connector/pRNA complex was initiated by alignment of the sixfold and 12-fold rotational axes of the molecules. The inner diameter of the pRNA hexamer was matched with the variable outer diameter of the connector to establish the depth of sheathing of the pRNA onto the connector. A limited rotational search (by definition, the largest error can be no more than 15°) identified one angle that allowed persistence of the N-terminal helices and their putative recognition of the pRNA (Fig. 4E–H).

Ten residues missing from the N terminus of the connector crystal structure (PDB 1HSW) (Gausch et al. 2002) were incorporated as a 10 residue extension of the N-terminal α helix (Fig. 4F). Of the newly modeled N-terminal 10 amino acids of the connector, all amino acids previously determined to interact with the pRNA lie near phosphates or bases of the pRNA hexamer. The close proximity of nucleotides 28–32, 69–72, 46–50, 62, and 81–84 of the pRNA to connector residue S170 agreed with results from Fe-BABE probing of the motor structure (Supplemental Fig. S3; Atz et al. 2007). In addition, the procapsid binding domain of the pRNA (bases 26–83) was in close contact with the connector protein, in agreement with the results that this region is protected from nuclease degradation and chemical modification upon motor assembly (Reid et al. 1994a; Zhang et al. 2001). (For an animation movie of the phi29 pRNA/connector model, see Supporting Video S1.)

Verification of distances between nucleotides in the hexameric RNA ring with distance parameters obtained by biochemical methods

Previously, various biochemical methods were used to study the tertiary structure of pRNA. The presence of helical regions was predicted by phylogenetics and later confirmed by complementary modification experiments (Reid et al. 1994c; Zhang et al. 1995b, 1997). Disruptions of the helical regions 1–2/117–116, 7–9/112–110, 14–16/103–101, and 76–78/90–88 were found to inactivate pRNA activity in DNA packaging, while activity can be restored by compensation mutations. These base-pairing interactions are reflected in the constructed hexameric pRNA model (bases 1–2 are paired with bases 117–116; bases 7–9 are paired with bases 112–110; bases 14–16 are paired with bases 103–101; and bases 76–78 are paired with bases 90–88) (Fig. 5A).

Photoaffinity crosslinking has been applied to probe bases that are in close proximity in the pRNA structure (Chen and Guo 1997). Crosslinking using psoralen, a small photoactive probe that can covalently link uridines upon UV irradiation (Cimino et al. 1985; Tyc and Steitz 1992), showed that U69 is in close proximity to U31, U33, and U36 in the presence of Mg²⁺ (Chen and Guo 1997). Additional photoaffinity crosslinking using phenphi to crosslink guanosine bases in close proximity showed that base G75 is close to G28 and G30 (Mohammad et al. 1999). Similar data using azidophenacyl (APA) crosslinking also indicates that base G75 is in close proximity to bases 26–30, while G78 is close to U31, and G108 is proximal to bases 10–11 (Mat-Arip et al. 2001). These are all in good agreement with the pRNA model reported here, as shown in Figure 5, B through F.

Chemical modification revealed that nucleotides 18–20, 42–48, 55–57, and 82–86 are solvent accessible in monomer, dimer as well as procapsid-bound pRNA forms (Trottier et al. 2000; Zhang et al. 2001), indicating that these regions are present as single-stranded loops (Zhang et al. 2001).
three bases C18C19A20 are deleted (Reid et al. 1994b; Zhang et al. 2001) or paired with 3′GGUS′ inserted between A99 and A100 (Zhang et al. 1997), the resulting pRNA loses its packaging activity while retaining the ability to form dimers and bind procapsid, suggesting CCA is not involved in procapsid binding. Comparison of these regions in the newly modeled hexameric pRNA reported here agrees with these data.

CONCLUSIONS

The 3WJ structure serves as a core of the phi29 hexameric motor pRNA. Crystal structure analysis, as well as biophysical and biochemical studies, indicates that the binding of divalent metal ions promotes a structural change of the 3WJ of pRNA. Analysis of the 3WJ core structure provides a better understanding in the overall pRNA structure and multiparameterization, which are crucial for pRNA function. In recent years, RNA nanotechnology has emerged to play an important role in cancer therapy and disease treatment, serving as a vehicle to deliver therapeutics (Guo 2010; Guo et al. 2012a). Studies on 3D structures of RNA lead to identifications of useful RNA motifs (Leontis et al. 2006), which can be used to carry therapeutics for delivery (Shu et al. 2011a, 2013a,b; Haque et al. 2012). Several self-assembling, stable, and polyvalent RNA nanoparticles have been produced based on computational designs (Afonin et al. 2010; Dibrov et al. 2011; Grabow et al. 2011), in which information of the individual RNA motifs as building blocks is essential. The pRNA 3WJ core has been demonstrated to be a useful platform to generate therapeutic RNA nanoparticles for targeted delivery (Shu et al. 2011a, 2013a,b; Haque et al. 2012). Our analysis of this 3WJ’s crystal structure of core will give insights in the designs of more effective therapeutics RNA nanoparticles.

MATERIALS AND METHODS

RNA synthesis and crystal screening

Each of the pRNA 3WJ strands (Fig. 1A), a3wj, b3wj, and c3wj (Shu et al. 2011a, 2013a,b; Haque et al. 2012), was purchased from IDT. Individual RNA strands were designed to produce blunt ends when they are annealed to facilitate crystal packing through base stacking interactions (Reyes et al. 2009). The nucleotide U24 was replaced with C24 in strand a3wj, and this change did not affect the pRNA’s biological activity in DNA packaging. The three strands were self-assembled to form the 3WJ in DEPC water simply by mixing them at equimolar concentrations. Initial crystallization trials were carried out at 22°C in 96-well plates with the hanging drop vapor diffusion method using a “Mosquito” nano-crystallization robot. Drops consisting of 0.1 μL RNA solution were mixed with an equal volume of reservoir solution and equilibrated against 100 μL reservoir solution. Lead conditions were optimized to 100 mM acetate (pH 4.6), 24%–26% 2-methyl-2,4-pentanediol (MPD), 20 mM CaCl2, and 10 mM MnCl2.

3WJ crystallization and structure determination

The initial crystallization trials were carried out at 295 K in 24-well plates using the hanging-drop vapor-diffusion method with commercially available screening kits for nucleic acids or nucleic acid/protein complexes, including Natrix, Nucleix (Qiagen), and Nuc-Pro (Jena Biosciences). Drops consisting of 0.5 μL RNA solution in DEPC-treated water (7 mg/mL) were mixed with an equal volume of reservoir solution and equilibrated against 500 μL reservoir solution. Initial hits were identified, and conditions yielding small crystals were further optimized by variation of crystallization parameters and additives. The final conditions that produced the best diffraction quality crystals (Supplemental Fig. S1C) contained 24%–26% (v/v) MPD, 20 mM CaCl2, 10 mM MnCl2, and 100 mM sodium acetate (pH 4.6). Crystals appeared within a week and continued to grow until they reached average dimensions of 0.03 × 0.03 × 0.3 mm (Supplemental Fig. S1C). The crystals were mounted on cryoloops and directly plunged into liquid nitrogen without additional cryoprotectant prior to data collection.

Native data sets were collected from single crystals cryoooled to 100 K on a MAR225 CCD detector at the SER-CAT 22BM beamline at APS. The data sets were processed with HKL2000 and scaled and merged with Scalepack (Otwinski and Minor 1997). The best crystals, space group I41, diffract to 3.05 Å (Supplemental Fig. S1D) and have unit-cell parameters a = b = 125.01 Å and c = 26.99 Å. Final data collection statistics are summarized in Table 1.

The structure was determined by piecewise molecular replacement using three copies of a seven base-paired polyadenine standard A-form double helix (stem) as a model. Solvent content analysis indicated one 3WJ molecule in the asymmetric unit (Vm=3.07 Å³/Da, 68.2% solvent content assuming RNA density of 1.7 g/cm³). The molecular replacement search for three stems (which potentially constitute a monomeric 3WJ structure) was carried out with Phaser (McCoy et al. 2007). The best solution contained three stems and had a Z score of 7.7 for the translation function, 12% higher than the next highest potential solution. Subsequent rigid body refinement using CNS (Brüger et al. 1998) resulted in R and Rfree values of 46.1% and 49.6%, respectively. The phases were improved through rounds of manual rebuilding with Coot (Emsley and Cowtan 2004) and composite omit map calculation implemented in CNS, which enabled building of >80% of the structure. Subsequent rounds of model building and refinement using Phenix (Adams et al. 2002) identified the proper asymmetric unit and sequence register. The metal and water sites were identified and validated by normal and anomalous difference Fourier maps as well as bonding geometry.

The structure was refined with Phenix (Adams et al. 2010) to final Rwork and Rfree values of 18.2% and 19.7%, respectively. Four TLS groups matching the three helices, i.e., (1) residues A1-8/C9-16, (2) residues C1-8/B13-20, (3) residues B1-5/A14-18, and (4) the central region of the 3WJ with residues B6-12/A9-13 were defined for the refinement of atomic displacement parameters. The final model displays a good stereochemistry according to MolProbity (Chen et al. 2010) and consists of 54 nucleotides, three Mn2+ ions, and two solvent molecules. Refinement statistics are given in Table 1, and the final 2Fo–Fc electron density map is shown in Supplemental Figure S1B. The coordinates and the structure factors have been deposited in the PDB (4KZ2).

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Hexamer model building

The 3WJ crystal structure was used as a template to build a 120-nt pRNA monomer. Three regions of duplex RNA, designated 1, 2, and 5 (the nomenclature follows a previous publication) (Hoeprich and Guo 2002) are connected at the 3WJ, while regions 3 and 4 comprise the right-hand loop and head loop, respectively. Region 1 was grafted from the hexamer pRNA model and includes the ATPase interaction surface. Region 2 displays the left-hand loop, and region 5 comprises the RNA helix from the 3WJ and includes region 3, designated the right-hand loop.

The head loop and right-hand loop (regions 3 and 4) were positioned by superposing our 3.0Å 3WJ crystal structure with the crystal structure of pRNA25,95 (PDB 3RF4) (Ding et al. 2011) on region 5 using the sugar-phosphate backbone for the 7 bp prior to the right-hand loop. Residues from these regions were grafted onto the 3WJ crystal structure. The left-hand loop was aligned by superposing 3WJ and PDB 3RF4 for 6 bp of region 2 leading up to the left-hand loop, followed by grafting the loop from 3RF4 onto the 3WJ core. The RNA helix containing region 1, including the ATPase interaction domain, came from PDB 1LAO, previously published 3D model for the pRNA hexamer (Hoeprich and Guo 2002). Residues 1–20 and 100–117 were grafted from 1LAO onto the 3WJ structure after superposition of the sugar-phosphate backbone for 4 bp leading up to the CCA bulge. Finally, two single-residue insertions missing from any current model in the database were built into the pRNA monomer, including a "flipped out" conformation for U35 and an in-line conformation for residue U81 of the left-hand loop. To complete the monomer model, three unpaired residues (118–120) were added to the terminus of region 1. For all molecular "grafts," bonds and angles were energy minimized using the "regularize" feature in Coot. All duplex regions from our 3WJ structure were included in the initial model.

To facilitate building a symmetric hexameric pRNA, the monomer model was put into a P6 pseudo-cell, and the orientation was optimized to obtain the best alignment of the 4 bp of the left- and right-hand loops within a hexamer. The resulting hexameric assembly was energy minimized using AMBER force fields (with no symmetry restraints) implemented with Chimera (UCSF). When the six individual monomers output from energy minimization were analyzed as symmetric hexameric assemblies, one model displayed superior packing of left- and right-hand loops relative to the other five. This model was subjected to another round of model building and energy regularization with Coot before expansion into the final sixfold symmetric pRNA hexamer model (Fig. 4).

AFM imaging of re-engineered pRNA ring

Loop-extended pRNAs were constructed by extending the 4-nt complementary loops to 7-nt loops and synthesized through in vitro transcription (Shu et al. 2013b). Six different re-engineered pRNA molecules containing intermolecular complementary loops were mixed at equal concentrations to form stable pRNA hexamers in solution, and the hexameric pRNA was purified from a native PAGE gel. The sample was fixed to the specially modified mica surface (APS) (Lyubchenko and Shlyakhtenko 2009) for AFM imaging. The AFM images were acquired in air using Super Sharp DIL probes (K-Tek Nanotechnology) and a MultiMode AFM NanoScope IV system (Bruker Nano/Veeco) operating in Tapping mode.

smFRET of pRNA 3WJ

Individual biotin-labeled or fluorophore (Cy3 or Cy5)-labeled RNA strands were purchased from IDT, and the labeled 3WJ was produced by mixing the three strands at equimolar concentrations in DEPC water. The 3WJ was then diluted in buffers containing 50 mM Tris (pH 8) and 100 mM NaCl, with 0 mM or 10 mM MgCl₂ for comparison in the absence or presence of Mg²⁺. The sample was immobilized at the biotin-BSA-coated surface of the sample chamber via biotin/streptavidin interaction according to methods previously described (Shu et al. 2010). Concentration of the samples was carefully picked to ensure the observation of individual fluorescent spots while keeping sufficient number of these spots in each field of view. Prism-type total internal reflection fluorescence (TIRF) imaging was performed according to methods previously described (Shu et al. 2010). A laser beam of wavelength 532 nm was used to excite the FRET pair, and dual-color fluorescence images were taken. The images were analyzed to obtain FRET efficiencies and to calculate distances between the FRET pairs.

Effect of Mg²⁺ ion on the resistance of 3WJ to urea denaturation

The 3WJ samples were assembled from the individual RNA strands in a buffer containing 25 mM Tris (pH 8), 50 mM NaCl, and 5 mM MgCl₂ and were assayed by 15% PAGE gels. The gels were made in 1× TB buffer (89 mM Tris-borate at pH 8) containing 5 mM MgCl₂ (TBMB) or 2 mM EDTA (TBE) to compare the stability of 3WJ with or without Mg²⁺. Different concentrations of urea (0 M, 4 M, and 8 M) were mixed into the gel to test the stability of 3WJ. The gels were run in their corresponding TBMB or TBE buffers.

DATA DEPOSITION

The coordinates and the structure factors of the pRNA 3WJ crystal structure have been deposited in the PDB (accession code 4KZ2).

SUPPLEMENTAL MATERIAL

Supplemental material is available for this article.

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Crystal structure of 3WJ core revealing divalent ion-promoted thermostability and assembly of the Phi29 hexameric motor pRNA

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