Visualizing the ai5γ group IIB intron

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ABSTRACT

It has become apparent that much of cellular metabolism is controlled by large well-folded noncoding RNA molecules. In addition to crystallographic approaches, computational methods are needed for visualizing the 3D structure of large RNAs. Here, we modeled the molecular structure of the ai5γ group IIB intron from yeast using the crystal structure of a bacterial group IIC homolog. This was accomplished by adapting strategies for homology and de novo modeling, and creating a new computational tool for RNA refinement. The resulting model was validated experimentally using a combination of structure-guided mutagenesis and RNA structure probing. The model provides major insights into the mechanism and regulation of splicing, such as the position of the branch-site before and after the second step of splicing, and the location of subdomains that control target specificity, underscoring the feasibility of modeling large functional RNA molecules.

INTRODUCTION

Group II introns are self-splicing ribozymes and retroelements that are found in bacteria, archaea and eukaryotes (1,2). Understanding the structure and behavior of group II introns is of great biological significance owing to their impact on gene expression and genomic organization in modern organisms and their evolutionary relationship with the eukaryotic spliceosome (3). Group II introns are also significant because they show great potential as gene targeting agents that may ultimately be applicable for gene therapy (4–6). From a biophysical perspective, they serve as excellent model systems for exploring basic principles of RNA folding, structure and catalysis (7,8).

Group II introns are structurally classified into three distinct groupings (Group IIA, IIB and IIC) (9,10). The group IIB introns are of particular interest because they are the most structurally complex, they are exceptionally sequence-specific and they are believed to represent a lineage most closely related to the spliceosome (11). While there is no available structure of a group IIB intron, crystal structures of a group IIC intron have become available (12–14). Group IIC introns lack many of the structural domains and enzymatic capabilities of the IIB introns (9,11,15). However, they represent a useful starting point for visualizing the basic architectural plan of all group II introns (12). Much has been learned from previous attempts to model group IIA (16) and IIB (17,18) introns, but the availability of group IIC intron crystal structures now makes it possible to use homology modeling methods. In the field of protein structural biology, a growing trend is to solve one crystal structure of a protein family, and then obtain structures of the relatives by homology modeling (19). This trend is supported by increasingly powerful tools, which yield accurate models for novel proteins that would be impossible to express or crystallize (20). Inspired by this growing trend in protein homology modeling, we have used a similar approach to build an atomic structural model of the ai5γ group IIB intron (ai5γIIB) from yeast mitochondria (21). To make this possible on such a large multidomain RNA (~800 nt), we adapted existing computational approaches and combined them with a new tool we developed for structure building and refinement of exceptionally large RNAs.

A diversity of excellent computational tools has made it increasingly possible to predict RNA structures from sequence and prior knowledge of architectural organization [see recent reviews (22,23)]. The ability to model large RNAs is of critical importance given the explosion of information on large, highly structured noncoding RNAs that direct everything from chromatin remodeling to translational regulation (24,25). Homology modeling tools such as ModeRNA (26) use sequence alignment and template structures to predict RNA structures. Fragment assembly based methods, such as FARNAs (27) and MC-Sym (28), build RNA by assembling nucleotide fragments extracted from crystal structures. NAST...
(29) and iFoldRNA (30) predict structures by performing coarse-grained molecular dynamics simulations. These methods use different input criteria, and there are various advantages and disadvantages associated with each method. For example, homology modeling tools are relatively fast and accurate, but they require template structures with good homology. On the other hand, de novo methods do not require template structures but are computationally expensive and therefore limited to small RNAs. Methods for backbone optimization and refinement are nearly nonexistent. Also, current tools are not fully automated and require manual manipulation at various levels, especially to model large RNAs.

To facilitate structural modeling of large RNAs, we expanded the capabilities of RCrane (31,32), which is a program that we originally developed for automated modeling of RNA into electron density, and which is now used for building RNA crystal structures (32). Here, we adapted RCrane for correcting backbone conformations within large RNA 3D models in the absence of electron density. Using this new tool, together with existing homology and de novo building methods, we built an all-atom 3D model of a group IIB intron. The resulting structure is consistent with known genetic and biochemical data on IIB architecture, and it reveals fundamentally important new insights into the mechanism of self-splicing.

**MATERIALS AND METHODS**

**Computational methods**

A homology model of the core structure was constructed with ModeRNA (26) using the OiIC crystal structure (PDBID: 3IGI) (12) as a template. Sequence alignment was performed manually with BioEdit (33). Additional subdomains specific to the IIB intron were modeled using MC-Sym (28) (see Supplementary Materials for an example of MC-Sym script) and manually docked onto the core structure with PyMol (Schrodinger, LLC) and Discovery Studio (Accelrys Software Inc) All helices were rebuilt using MC-Sym, replaced using PyMol (Schrödinger, LLC) and Discovery Studio (Accelrys Software Inc) and finally, the backbone was refined with RCrane. The new application of RCrane is included in the current version 1.1, which is available at http://www.pylelab.org/software/index.html, and also with Coot 0.7, which is available from http://www.biop.ox.ac.uk/coot/. The final model was subjected to energy minimization using the AMBER10 force field and generalized Born implicit solvent model available in the AMBER package (34). Minimization was started with 500 steps of steepest descent followed by conjugate gradient minimization for at least 500 steps. The quality of resultant structure was assessed using Molprobity (35) and the number of minimization cycles was increased in cases of high clash score. See supplementary Table S1 for complete list of software and additional details.

**RNA constructs and purification**

Plasmid pSS01 encoding the β-β' variant (stem loop #252-264 replaced with UUCG tetraloop) was constructed from pQL71 using the QuickChange site-directed mutagenesis kit. Both plasmids pQL71 (wild type) and pSS01 (β-β' variant) were linearized with HindIII before transcription and purified as described before (18).

**Hydroxyl radical footprinting**

RNA samples (4 pmoles) were incubated in a buffer containing 25 mM potassium cacodylate, pH 7.0, 500 mM KCl and 0.2 mM EDTA at 90°C for 2 min and then cooled to room temperature. Folding was initiated by incubating RNA (100 μl of final volume) at 42°C for 30 min in a buffer containing 25 mM potassium cacodylate, pH 7.0, 500 mM KCl and 100 mM MgCl₂ (or water instead of MgCl₂ for cleavage in the absence of Mg²⁺ ions). Footprinting reactions were initiated by applying 2 μl of each 2.5 mM (NH₄)Fe(SO₄)₂₆, 2.75 mM EDTA (pH 8.0), 1.5% H₂O₂ and 50 mM sodium ascorbate to the inside wall of the tube followed by vortexing the sample. After 15 s, the reactions were quenched with 20 μl of stop mix (containing 100 mM thiourea and 200 mM EDTA). RNA was precipitated with 2.5 volumes ethanol (100%), 10% 3 M sodium acetate (pH 4.8) and 0.5 μl of glycogen (Roche).

**SHAPE**

For selective 2'-hydroxyl acylation analyzed by primer extension (SHAPE) experiments, 4 pmoles of RNA (in 40 μl of buffer containing 500 mM KCl, 50 mM HEPES pH 7.4, 0.1 mM EDTA) were denatured at 90°C for 2 min and then cooled to room temperature. Folding was initiated by incubating RNA (150 μl of final volume) at 42°C for 30 min in a buffer containing 500 mM KCl, 50 mM HEPES pH 7.4, 0.1 mM EDTA and 100 mM MgCl₂ (or water instead of MgCl₂ for probing in absence of Mg²⁺ ions). RNA was divided equally (72 μl) between two tubes, and 8 μl of Dimethyl sulfoxide (DMSO) (for control) or 10 mM 1M7 (1-methyl-7-nitroisatoic anhydride) (36) were added. The modification reaction was allowed to proceed for 5 min at 37°C, and samples were precipitated with 2.5 volumes ethanol (100%), 10% 3 M sodium acetate (pH 4.8) and 0.5 μl of glycogen (Roche).

**Fragment analysis**

Cleavage products from both hydroxyl radical footprinting and SHAPE were analyzed by primer extension using fluorescently labeled primers. The fluorescent dye coding and instrument precalibration was applied as described (37). Briefly, primers with dyes (Anaspec) 6-JOE (reagent present) and 5-FAM (reagent absent) were used for reverse transcription, and those with 6-TAMRA (ddT) and 5-ROX (ddC) were used for cycle sequencing (Affymetrix cycle sequencing kit). Each primer sequence was corrected for mobility shift in ShapeFinder. Primers for reverse transcription were annealed by incubating 2 pmol RNA with 1 μl of 2 μM primer (12 μl of total volume) at 95°C for 3 min, 4°C for 5 min and 42°C for
2 min. Reverse transcription was initiated by adding 8 µl of mixture containing 4 µl of 5× First strand buffer, 1 µl of 100 mM DTT, 1 µl of 10 mM dNTPs, 0.5 µl of Superscript III (U/µl) and water at 48°C for 45 min. The (+) and (−) reagent reactions were quenched and combined followed by ethanol precipitation. Recovered cDNAs were resuspended in deionized formamide and mixed with sequencing ladders and were analyzed on ABI 3730xl DNA analyzer. Data were analyzed with ShapeFinder (38) as described before (39).

RESULTS

Three-dimensional modeling of a group IIB intron

Homology modeling

The core region of the ai5γIIB (Figure 1) was modeled with ModeRNA (26), using a published crystal structure of the Oceanobacillus iheyensis IIC intron (PDBID: 3IGI) (12) as the template (OiIIC). Sequence alignment was performed manually using BioEDIT (33). Before performing the alignment, regions of the IIB intron that are not found in the OiIIC template were removed, including D1c2, D1d2a, D1d2b, D3a, D3b, D3c and D6. The lengths of several helices and loops vary between the two introns, and these regions were left as gaps within the sequence alignment. All the helices were later rebuilt as idealized A-form helices using MC-Sym (28). Among the various helices generated by MC-Sym, the helix that fits best with the existing helix in the core homology model was selected and replaced using PyMol and Discovery Studio (DS) Visualizer. Long-range interactions with different structural forms were adjusted manually: For example, the ζ-ζ' interaction (GAAA tetraloop-receptor) (40,41) was modeled from a similar motif taken from the PDB (PDBID: 2JYF). The α-α' kissing-loop interaction was modeled as a short helix and replaced using PyMol and DS Visualizer. The resulting model consists of a full-length D5, the basal stem of D3, a truncated D2 and D4 and all elements of D1 except D1c2a, D1d2a and D1d2b. The model coordinates are available in the Supplementary Material.

De novo modeling

D6 and various subdomains that are only found within the ai5γIIB intron, such as D3a, D3b, D3c, D1c2a, D1d2a

Figure 1. Schematic secondary structure of ai5γIIB intron. Domains 1–6 are labeled in capital letters, subdomains are labeled in small letters. Exons are colored in magenta. Exon and intron binding sites are labeled as EBS and IBS, respectively. Regions highlighted in light blue are common in ai5γIIB and OiIIC introns (core region); they were built with homology modeling using OiIIC crystal structure as the template. Additional insertions specific to ai5γIIB (highlighted in orange) were built de novo and docked on to the homology model. D6 (highlighted in green), which is missing in all available crystal structures of OiIIC, was also modeled de novo. Long-range tertiary interactions are labeled in red.
and D1d2b, were modeled de novo using MC-Sym (28) and then docked manually onto the core model using PyMol and DS visualizer (Figure 2). For a given secondary structure, MC-Sym generates an ensemble of decoy structures using fragment-based libraries. As the input for MC-Sym, the domain to be modeled was appended to a known region that was already part of the core model, and which served as an anchor for later docking steps. The ensemble of structures predicted by MC-Sym were clustered, docked onto the core model and finally filtered based on known biochemical data.

To model D6 in the context of the intron core, our strategy involved sampling several possible conformations of D6 that are sterically allowed, but which do not significantly alter the structure of the intron core. D6 is covalently linked to D5 via a 3-nt linker, and both domains are part of a higher-order six-way junction. Five helices of this six-way junction are already present in the homology model. In the absence of the other helices, D5 and D6 can obviously sample many conformations; however, relatively few of these are sterically allowed when they are part of a six-way junction. We predicted conformations of D5 and D6 as a two-way junction using MC-Sym. The resulting conformations were clustered and docked by aligning D5 with the corresponding region of the core model, and then filtered to remove conformations with steric clashes. From the remaining conformations, we selected one conformation where the bulged adenosine of D6 is proximal to the active site. We then manually adjusted D6 such that the 2'-OH group of bulged adenosine occupies the position of a nucleophilic water that had been visualized crystallographically in structures of the OiIIC intron (42).

Additional subdomains D3a, D3b, D3c, D1c2a, D1d2a and D1d2b were modeled in a similar fashion (see Supplementary Results).

Model refinement

Backbone refinement: a novel application of RCrane

The RNA backbone is highly flexible, and given that seven torsion angles describe each individual nucleotide, computational modeling can be a daunting task (43). Previously, we developed an alternative conformational description that reduces the dimensionality of RNA structure. Using this approach, each nucleotide is represented by two pseudotorsions η and θ, where η represents torsion between C4₀⁻₋₁, Pᵣ, C4₀ⁱ and Pᵣ₊₁, and θ represents torsion between Pᵣ, C4₀ⁱ, Pᵣ₊₁ and C4₀ᵢ₊₁ (44). Further, using both the RNA pseudotorsions and the RNA backbone conformers (45) we developed the program RCrane, which is a computational tool for semiautomated model building of RNA into electron-density maps (31,32). Here, we developed a novel application of RCrane, using it to automatically correct the backbone configurations in the models even before an electron-density map is available. In the absence of an electron-density map, RCrane first predicts the appropriate backbone conformer based on the positions of phosphate and base, and then subsequently improves the placement of both the phosphate and the backbone sugar. This diversification of RCrane’s capabilities demonstrates that the approach has additional

![Figure 2. Overview of model building. Modeling was performed in three steps, in the first step the core structure of the intron was generated with homology modeling using crystal structure from IIC as the template. Next, all additional regions specific to the IIB intron were modeled using MC-Sym and docked onto the core structure. Finally, the model was refined using RCrane (see text) and AMBER.](https://academic.oup.com/nar/article-abstract/42/3/1947/1056107)
applications in building, modeling and analyzing RNA structures. This new application is included in the current version of RCrane 1.1 available at http://www.pylelab.org/software/index.html and also with Coot 0.7 available from http://www.biop.ox.ac.uk/coot/.

We used RCrane to refine the backbone of the ai5γIIB model at various stages of the modeling process (see Figure 2). The resulting model was subjected to energy minimization with AMBER (34) to remove steric clashes, resulting in an overall clash score that is well within the acceptable range (0.2, 99th percentile) (35).

Model validation

The resulting computational model was validated experimentally using hydroxyl radical footprinting. Hydroxyl radical footprinting probes the solvent-accessible regions of a molecular structure (46). Recent work has shown that there can be good correlation between hydroxyl radical reactivity and number of neighboring nucleotides (47) in the crystal structures of various RNAs. In this work, we calculated the number of contacts of each nucleotide in the model and correlated with hydroxyl radical reactivity.

First, we performed hydroxyl radical footprinting followed by primer extension and fragment analysis with capillary electrophoresis on D135, which is an ai5γIIB construct that contains full length D1, D3 and D5, a truncated form of D2 and D4, while it lacks D6 and both exons. To be consistent with the experiment, the exons and D6 were removed from the model before calculating the number of contacts. Numbers of contacts for each nucleotide in the model were calculated as described previously (47), using 20 Å distance cutoff. There is good correlation between numbers of contacts in the model and hydroxyl radical reactivities (Figure 3A). The observed correlation coefficient ($r = -0.61$) is as good as that observed for crystal structures of other RNAs (47) and it is also statistically significant with $P < 10^{-24}$ (Figure 3A).

The locations of D1c2 and D1d2a (stems containing the loops that form the β-β' interaction) were also validated with mutation and hydroxyl radical footprinting. Previous results have shown that both D1c2 and D1d2a are exposed to solvent, and they are believed to lie at the periphery of D1 (18). However, disrupting the kissing-loop interaction between these stems should disengage them and expose regions that lie protected beneath these stems. To test this hypothesis, and thereby test the model, we disrupted the β-β' kissing-loop by replacing the D1d2a stem loop (nucleotides 252–264) with a UUCG tetraloop (the β-β' mutant).

Hydroxyl radical footprinting was then performed on both the WT D135 and the β-β' mutant. In the case of the β-β' mutant, both the D1c2 and the D1d2a stems showed significant decreases in hydroxyl radical protection (Figure 3B). Disruption of the β-β' kissing-loop therefore causes the two stems to become more dynamic in solution and more solvent exposed relative to their position in WT. Significant decreases in hydroxyl radical protection were also observed in regions of D1b (57–59), D1d'' (232–236), D1d3' (326–328), D3b (632–634, 638–644), D3c (655–657) and D4 (808–813). Among these regions, D1b (57–59), D1d'' (232–236) and D1d3' (326–328) lie beneath the interacting D1c2 and D1d2a stems in the model, indicating that mutation has exposed them and that the model is in good agreement with the experimental data (Figure 3C).

Interestingly, there are a few regions that are more protected in the mutant construct: D1d' (216–218), D1d2b (302–306), D1d3 (357–365), D1d (386–391, 394–398) and D1 (i) (404–408). Most of these regions are in the vicinity of D1c2 or D1d2a, suggesting that when the β-β' kissing-loop is disrupted, D1c2 and D1d2a do not simply move around randomly. Rather, they may form new types of discrete stabilizing interactions.

To ensure that the observed increases in reactivity for the β-β' mutant were not simply a result of localized RNA unfolding, we performed SHAPE experiments to evaluate whether secondary structural elements are still intact in this mutant. As expected, nucleotides (145–149) involved in the β-β' kissing-loop interaction showed a significant increase in SHAPE reactivities, as the other half of this kissing-loop pair was replaced with a tetraloop (Supplementary Figure S1). This local reactivity increase was consistent and comparable with the SHAPE reactivity differences in WT when probed in the presence of 100 mM Mg$^{2+}$ (folded, β-β' interactions formed) and without Mg$^{2+}$ (not folded, β-β' interaction not formed). For the remaining regions of the mutant, there were no significant changes in SHAPE reactivity when compared with WT, suggesting that the secondary structure has not been affected by the mutations (Supplementary Figure S1). Taken together, these results support the location of D1c2 and D1d2a in our model.

Finally, to validate the location of D3, we compared previously published experimental footprinting data on the ai5γIIB D135 construct with data on a D15 construct that lacks D3 (48). On deletion of D3, regions in contact with D3 should become solvent exposed and are expected to be less protected in D15 construct compared with D135 construct. Again, there is good agreement between the model and experimental data, as most of the regions that are less protected in the D15 construct are located beneath D3 in the model (Supplementary Figure S2).

Structural architecture of ai5γIIB

As expected, the active-site region and the overall architecture of the ai5γIIB model is similar to the known structure of OIIIC. However, IIB introns possess additional subdomains and tertiary interactions (such as β-β', μ-μ' and EBS2-IBS2) that are important for both folding and catalysis (11,49). Unlike IIC introns, IIB introns react with high sequence-specificity and undergo branching efficiently. The ai5γIIB model allowed us to visualize the additional subdomains and thereby provided valuable hints on functional roles of structural features specific to IIB introns. For example, the β-β' and μ-μ' tertiary interactions form on opposite sides of D5 (Figure 4 and Supplementary Movie S1). D1c2 and D1d2a (the stems supporting the β-β' interaction) are located on the ‘catalytic face’, while D3 falls on the ‘binding face’ (50) of D5. D1c2 and D1d2a protrude like two extra arms from...
opposite sides of D1, and the β-β kissing-loop interaction between these hairpin loops interlocks like a bridge that is likely to rigidify the scaffold of D1 (Figure 4A). On the other side of D5, D5a and D5b form a wall along the ‘binding face’ that may mediate interactions between D5 and the rest of the intron (Figure 4B and C).

The second exon binding site, or EBS2, is a major distinguishing feature of the highly derived IIB and IIA introns. The model shows that EBS2 is far more solvent exposed than EBS1, which is consistent with footprinting data on ai5γIIB intron (18). The model also shows that the EBS1-IBS1 and EBS2-IBS2 helices, which form on target or exon binding, are not coaxial with each other, consistent with previous spectroscopic and mutational analysis of the ai5γIIB intron (51) and the model of Pylaiella littoralis intron (Pl.LSU/2) (17). We see that the EBS2-IBS2 helix is located immediately adjacent to the β-β interaction (Figure 5), suggesting that function of the two motifs may be linked. Importantly, the modeling process revealed that, when no exon is bound, the EBS2 region and adjacent helices are flexible. This indicates that exon binding, and formation of the EBS2-IBS2 helix, rigidifies EBS2 and the surrounding substructures. Thus, formation of the EBS2-IBS2 helix induces conformational rearrangements that are likely to considerably reduce overall entropy, with important implications for the known role of EBS2-IBS2 in substrate specificity and reaction chemistry (51,52).

Among the subdomains that are specific to IIB introns, D1d2b is the only substructure that appears to be pointed away from the intron core and has no contacts with the rest of the intron. However, D1d2b is coaxial with D1d2a, and it may therefore play an indirect role in the formation of β-β kissing-loop between D1d2a and D1c2.

The active and silent conformers of D6
D6 can exist in two different conformations (Figure 6A). In the conformation capable of branching, (the active form) the 2'-OH group of the branch-site adenosine is positioned precisely to initiate the first step of splicing within the active site (Figure 6B). After the first step, the branch-point nucleotide flips out of the active site, resulting in an alternate conformation (the silent form) (11). The toggling of D6 between active and silent forms is likely to represent an important regulator of splicing in group II introns, and potentially, in the spliceosome as well, for cognate domains (53). To prepare the active site for second step of splicing, and to position the 3'-exon appropriately, D6 will need to exit from the active site. However, cross-linking studies (54,55) have shown that nucleotides in D5 and D6 are located approximately in the same position through the entire cycle of splicing, which is consistent with studies showing that the same active-site elements play a role in both steps of group II intron splicing (56).
Understanding this paradox is of utmost importance, and various studies already attempted to model D6 in active form (16, 57). Here, we modeled both active and silent forms of D6 and sought to visualize its conformational toggling. The location of D6 in active form is consistent with recent studies on the \textit{P. littoralis} intron (Pl.LSU/2) (57). The silent conformation was modeled such that the 3' exon was positioned within the active site, where it forms the EBS3-IBS3 interaction and the \(\gamma-\gamma'\) interaction between the last nucleotide of the intron (U887) and residue A587 (Figure 6C). Surprisingly, the overall spatial location of D6 is similar in the active and silent forms (Figure 6A). Close comparison of the two states reveals that D6 undergoes more of a rotation than a translation along its helical axis on toggling, so that the global position of D6 with respect to the intron remains similar, but individual nucleotides within D6 shift their position. This is consistent with the fact that, after the first step of splicing, D6 cannot move too far from the intron core, as the branched adenosine is now covalently linked (2'-5' lariat bond) with 5'-end of the intron.

Further, we noticed that this conformational change is sufficient to support formation of the \(\eta-\eta'\) interaction between D6 and D2 (Supplementary Figure S3). The crystal structure of OiIIC contains only the basal stem of D2, while the D2 stem that participates in \(\eta-\eta'\) is longer, and part of a four-way junction in ai5\gammaIIB intron. Surprisingly, we were able to model this junction in a way that allows formation of \(\eta-\eta'\) by moving D6 only slightly from the ‘active’ conformation and without disturbing the position of D2 basal stem and the \(\theta-\theta'\) interaction. It is interesting to point out that the tip of D6 in ‘active’ form is not too far from the receptor of \(\eta-\eta'\) interaction in D2, suggesting that D2 may play a role in positioning D6. This is consistent with previous studies showing that mutations of \(\eta-\eta'\) affect exon ligation (58).

\textbf{DISCUSSION}

Here, we report an all-atom model of an intact ai5\gammaIIB, which was built using a combination of homology and \textit{de novo} methods. This approach has enabled us to visualize...
key group II intron structural features, such as the two conformations of intron D6, and the EBS-IBS2 specificity helix. The relative locations and orientations of these subdomains, along with structural innovations such as β-β', explain many of the functional attributes unique to group IIB introns, such as their extreme sequence-specificity and their propensity to splice through branching.

One of the features most readily apparent from the model is that D5 is surrounded by a larger and more protective shell in the ai5γIIB intron relative to OiIIC (Figure 7). This well-packed IIB intron scaffold, along with added interactions such as β-β' and μ-μ', is likely to affect the overall dynamics of the intron, influence the electrostatic potential at the active site and provide additional stability to the active site as it assembles on the surface of D5. Previous studies of ai5γIIB have shown that D5 binds so tightly to the rest of the intron (59,60) that it can be added as a separate domain to truncated variants of the intron (59–61).

Perhaps most importantly, the model provides insights into the branching mechanism of group II introns. Splicing through branching is one of the most notable features of group II introns, but it has remained difficult to visualize because the existing IIC crystal structures lack electron density for D6, and IIC introns do not readily

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**Figure 5.** Location of the EBS2 binding site. The EBS2-IBS2 duplex is shown in orange, the EBS1-IBS1 duplex in shown in magenta. The linker between IBS1 and IBS2 is shown in red. The rest of the intron is shown as a transparent surface representation. The EBS2-IBS2 interaction is located next to the kissing-loop (β-β') interaction that joins D1c2 (green) and D1d2a (blue), suggesting that function of the two motifs may be linked.

**Figure 6.** Active versus silent form of D6: (A) D6 can exist in two different conformations: the active (green) and silent forms (magenta). (B) In the conformation needed for the first step of splicing (the active form) the 2'-OH group of branch-site adenosine (green) is positioned in the active site to initiate the first step of splicing. Active-site elements (catalytic triad, 2-nt bulge and J2,3 junction) are shown in blue, 5'-exon is shown in yellow, first nucleotide of the intron G1 is shown in purple. (C) In the conformation that forms after the first step of splicing, and which is required for the second step (the silent-form), the branch-site adenosine flips out and positions the 3'-exon in the active site and forms two new interactions (shown in red) IBS3-IBS3 and γ-γ'. Based on the model, toggling between the ‘active’ and ‘silent’ conformations of D6 involves a simple rotation of the domain rather than a large translational movement away from the active site.
branch (they splice only through hydrolysis in vitro) (62,63). Previous studies have proposed two candidate receptors of D6 for branching: D1c1 (57) and the coordination loop (55). In the model provided here, D6 (in the active form) is located close to D1c1; however, the coordination loop is also in the vicinity of D6, and it is possible that D6 interacts with the coordination loop before the branch-site adenosine is recruited into the active site. During the modeling process, we observed that the presence of additional subdomains (D1c1 and D1d2a) restrict the possible conformations for D6 in aI5yIIB relative to D6 in OIIC, which is likely to enhance branching relative to hydrolytic attack at the 5’-splice site. In addition, we observe that toggling between the ‘active’ and ‘silent’ conformations of D6 involve a simple rotation of the domain rather than a large translational movement away from the active site as proposed previously (11), thereby explaining the many biochemical studies, showing that intron domains do not radically reorganize between the two steps of splicing (54) (Figure 6).

The EBS2-IBS2 interaction is another distinguishing feature of group IIB and IIA introns, and the model helps to show why this structural innovation has come to play such an important role in intron function. There are introns, such as Clostridium beijerinckii I2 (64) and Pelotomaculum thermopropionicum strain SI I2 (64), which contain an EBS2-IBS2 interaction with no apparent role in self-splicing (64,65). When we compared

Figure 7. Comparison of the ai5yIIB model with the crystal structure of OIIC. (A) Front and side views of the ai5yIIB model. Regions common to both introns are colored in light blue; D5 is colored in red. Long-range interactions that are not present in OIIC are labeled in Greek letters. D1c2 is shown in purple, D1d2a is shown in magenta, D3a and D3b are shown in orange. EBS2-IBS2 helix is shown in green. For comparison with OIIC, D6 is not shown in both views. (B) Front and side views of OIIC crystal structure.
the secondary structures of those introns with ai5γIIB, we noticed that the β-β′ interaction is absent in C. beijerincki I2 (64), P. thermopropionicum strain SI I2 (64) and Sinorhizobium melliloti Rm1021 (65). In the model, the β-β′ interaction lies in a cleft between the EBS1-IBS1 and EBS2-IBS2 duplexes (Figure 5), consistent with a known cross-link between IBS1 and Dbc2 (66) (one of the stems involved in β-β′). Thus, a functional EBS2-IBS2 duplex may require the presence of β-β′ and the two may act synergistically to rigidify the overall intron structure and promote specificity. Another important observation is that EBS2 is more solvent-exposed than EBS1. This suggests that EBS2 may interact with its cognate IBS2 site first, triggering a conformational change that makes EBS1 more accessible, and then leading to EBS1-IBS1 formation. Such a mechanism might be important during specific retro-homing events, whereby the intron protects EBS1 from binding to nonconspecific sites, but allows binding at EBS1 to occur only after a specific EBS2-IBS2 duplex is formed. Finally, the lack of coaxial stacking between the EBS-IBS duplexes explains the extraordinary sequence-specificity of group IIB introns. Because each duplex is fully independent, and the overall exon interaction energy is not supplemented by coaxial stacking, the recognition helices are less tolerant of mismatches (51), leading to the large specificity index for target site recognition that is observed for group IIB introns (51). The specificity index may also be enhanced by the insertion of β-β′ into the cleft between the EBS-IBS duplexes (Figure 5), as the resultant steric clashing may give rise to the energetic penalty that is known to accompany binding of an oligonucleotide to both EBS1 and EBS2 simultaneously. In fact, it is known that binding of long oligonucleotide targets (which bind both EBS1 and EBS2) is ~8 kcal/mol less favorable than binding of short oligos to EBS1 and EBS2 individually (52).

Although the RNA computation field has significantly progressed in the past decade, it was necessary for us to create new tools to complete and refine the model of the IIB intron. Here, we developed a broadly applicable computational tool for automatically correcting the backbone configurations in the models even before an electron-density map is available. This new tool, which is part of the RCrane program (32), will be helpful in modeling large RNA structures in conjunction with existing RNA modeling methods. Several features of RCrane as a 3D modeling tool are notable: First, existing, homology modeling methods (including those used for protein structures) derive models by copying coordinates from a template structure, which is a process that often leads to improper backbone configurations and steric clashes, especially in regions with low sequence similarity. RCrane rebuilds the entire backbone while keeping the position of the bases constant, and thereby provides the unique advantage of preserving secondary structure throughout refinement. In addition, large RNAs like group II introns may use different structural motifs but still share similar global structures, and RCrane preserves this information content. For example, the tetraloop and receptor interaction of D5 in IIC introns has a different structural form compared with other group II introns (40) and various long-range interactions also display slight differences, such as the number of base pairs involved in a kissing-loop. Currently, no methods exist to automatically correct for such differences, and users would have to manually adjust or rebuild these regions. By contrast, RCrane semiautomates this process, allowing the user to adjust or rebuild the nucleotide base, while RCrane rebuilds the backbone accordingly and ensures that it conforms to allowed rotameric states. RCrane can also be used to enhance de novo modeling methods, such as MC-Sym, which build structures by assembling fragments from nucleotide libraries. This process often results in structures with poor backbone connectivity. Again, RCrane can be used to fix backbone inconsistencies in these models without disrupting the secondary structure. In this work, we used RCrane at various stages to successfully build the model of ai5γIIB intron.

The ability to create a detailed homology model of a large multidomain RNA is of particular interest at this time, as it is becoming apparent that large RNAs are central to most aspects of gene expression in biology (25). There is also increasing evidence that large noncoding RNAs are, in many cases, highly structured (24). Since it is not currently feasible to experimentally determine structures for a majority of these RNAs, there will be an increasing demand for computational methods to predict RNA structures. The modeling strategies used here can be applied to other group II introns and to other multidomain RNA molecules for which empirical structural data are available.

In conclusion, we have shown that it is now possible to model large RNAs even from remote homologs. We report and test a 3D model of the ai5γIIB intron that provides structural insights into the mechanistic behavior of group II introns and explains functional differences between group IIC introns and more evolutionarily derived group II introns such as ai5γIIB.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online, including [9,11,26,28,32–35,48,49,67].

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