Secondary structure confirmation and localization of Mg\(^{2+}\) ions in the mammalian CPEB3 ribozyme

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ABSTRACT
Most of today’s knowledge of the CPEB3 ribozyme, one of the few small self-cleaving ribozymes known to occur in humans, is based on comparative studies with the hepatitis delta virus (HDV) ribozyme, which is highly similar in cleavage mechanism and probably also in structure. Here we present detailed NMR studies of the CPEB3 ribozyme in order to verify the formation of the predicted nested double pseudoknot in solution. In particular, the influence of Mg\(^{2+}\), the ribozyme’s crucial cofactor, on the CPEB3 structure is investigated. NMR titrations, Tb\(^{3+}\)-induced cleavage, as well as stoichiometry determination by hydroxyquinoline sulfonic acid fluorescence and equilibrium dialysis, are used to evaluate the number, location, and binding mode of Mg\(^{2+}\) ions. Up to eight Mg\(^{2+}\) ions interact site-specifically with the ribozyme, four of which are bound with high affinity. The global fold of the CPEB3 ribozyme, encompassing 80%–90% of the predicted base pairs, is formed in the presence of monovalent ions alone. Low millimolar concentrations of Mg\(^{2+}\) promote a more compact fold and lead to the formation of additional structures in the core of the ribozyme, which contains the inner small pseudoknot and the active site. Several Mg\(^{2+}\) binding sites, which are important for the functional fold, appear to be located in corresponding locations in the HDV and CPEB3 ribozyme, demonstrating the particular relevance of Mg\(^{2+}\) for the nested double pseudoknot structure.

Keywords: CPEB3; ribozyme; pseudoknot; Mg\(^{2+}\); metal ion; NMR

INTRODUCTION
The discovery of small self-cleaving ribozymes in mammalian genomes is rather recent. Only in the last decade, a discontinuous hammerhead ribozyme in the 3′-UTR of C-type lectin type 2 genes (Martick et al. 2008), a hammerhead ribozyme in an intron of a tumor suppressor gene (de la Pena and García-Robles 2010), the CoTC motif in the 3′-untranslated region of the β-globin gene (Teixeira et al. 2004) (although in a later work [Dye et al. 2006], autocatalytic RNA cleavage of the CoTC element in the context of intron 2 could not be confirmed), and the CPEB3 ribozyme in the second intron of the cpeb3 gene (which encodes a cytoplasmic polyadenylation element binding protein) (Richter 2007) were described. In this study, we focus on the CPEB3 ribozyme, which was discovered via an in vitro selection approach, starting from a human genomic library (Salehi-Ashtiani et al. 2006).

The function of the CPEB3 ribozyme in the cell is yet unknown but it is likely to regulate the levels of the CPEB3 protein in the cell by its self-cleavage. The CPEB family of proteins is involved in diverse processes such as germ cell development and synaptic plasticity (Christerson and McKearin 1994; Lantz et al. 1994; Luitjens et al. 2000; Darnell and Richter 2012). A single nucleotide polymorphism (SNP) in the human CPEB3 ribozyme has been shown to alter the rate of self-cleavage and to influence episodic memory (Vogler et al. 2009). This supports the suggested correlation between ribozyme activity, CPEB3 protein concentration and learning and memory.

When the CPEB3 ribozyme’s cleavage reaction was initially characterized, it became clear that its catalytic mechanism was very similar to that of the HDV ribozyme. The HDV ribozyme is involved in the separation of genome copies of the HDV satellite generated during rolling-circle replication (Macnaughton et al. 1993; Jeng et al. 1996). Both ribozymes cleave upstream of their first nucleotide, have nearly constant activity over a pH range of ∼6 to 8.5, show a solvent kinetic isotope effect, require Mg\(^{2+}\) for catalysis and are inhibited by hexamminecobalt(III) (Nakano et al. 2001; Salehi-Ashtiani et al. 2006). This strongly suggests that both ribozymes use a very similar combination of acid–base catalysis mediated...
by their nucleobases and Mg\(^{2+}\)-assisted catalysis. Support for the mechanistic resemblance of both ribozymes comes from the fact that all CPEB3 ribozyme sequences can be folded (in silico) into the nested double pseudoknot structure, which is known to occur only in the HDV ribozyme (Fig. 1A). Moreover, the mutation of nucleotides in positions corresponding to catalytically relevant nucleotides of the HDV ribozyme, interferes with CPEB3 self-cleavage (Salehi-Ashtiani et al. 2006).

The nested double pseudoknot structure of the HDV ribozyme comprises two helices (P1 and P2), two hairpin structures (the internal P3 and the P4 hairpin) and two single-stranded linker regions J1/2 and J4/2 (Fig. 1A; Perrotta and Been 1991; Ferre-D’Amare et al. 1998) and is required for self-cleavage activity of both the genomic and antigenic forms (Wadkins et al. 1999). A small pseudoknot consisting of P3 and P1.1 is embedded in a larger one consisting of P1 and P2. Based on the most recent crystal structure and molecular dynamics studies of the HDV ribozyme, a reverse G·U wobble base pair has been suggested to be formed at the base of P3 (G25·U20 wobble, circled residues on Fig. 1; Krasovska et al. 2006; Chen et al. 2010; Veeraraghavan et al. 2011; Kapral et al. 2014), which was not observed in the first two crystal structures (Ferre-D’Amare et al. 1998; Ke et al. 2004). This G25·U20 wobble is assumed to position the catalytic Mg\(^{2+}\) ion in the active site, which comprises also C75, the catalytic cytosine (C57 in the CPEB3 ribozyme) (Chen et al. 2013). N3-protonated C75 has been proposed to act as the general acid in the cleavage mechanism that protonates the 5’-O leaving group with an estimated pK\(_a\) of 6.4 (Gong et al. 2007). The catalytic Mg\(^{2+}\) ion has recently been proposed to serve as Lewis acid, Bronsted base, or both to activate the 2’-OH nucleophile, as well as to coordinate to the pro-R\(_P\) oxygen of the scissile phosphodiester (Chen et al. 2013; Thaplyal et al. 2013).

The strong resemblance of the CPEB3 and HDV ribozymes with regard to their catalytic reactions and (suggested) secondary structures has inspired the idea that the ribozymes are evolutionarily related, despite their dissimilar sequences. Given that the CPEB3 sequence is conserved in mammals, and that HDV is infecting humans, it seems most probable that the CPEB3 ribozyme is evolutionarily young and has been passed from human to HDV (Webb et al. 2009; Webb and Luptak 2011).

Although the complicated nested double pseudoknot motif is unlikely to have arisen independently several times (Nix et al. 1999; Nehdi and Perreault 2006), extensive bioinformatic searches have discovered the motif in the genomes of many organisms such as insects, nematodes, an insect virus, and bacteria. These HDV-like ribozymes are not conserved in their sequences and vary greatly in the length and structure of L1/2 and P4 (Webb et al. 2009; Webb and Luptak 2011; Riccitelli et al. 2014). This argues for the nested double pseudoknot motif being frequently used in self-cleaving sequences.

The HDV and CPEB3 ribozymes are different from all other small ribozymes in their obligatory requirement for divalent metal ions in their catalytic mechanism (Murray et al. 1998). This study, which is one of two NMR studies of such a full-length nested double pseudoknot construct (Tanaka et al. 2002), validates the proposed structure of the CPEB3 ribozyme in solution and elucidates the role of Mg\(^{2+}\) for structure formation. The constructs used were the human and chimpanzee (Fig. 1B) sequence of the ribozyme, which differ in only one nucleotide in P1 (G30 → A). A partial assignment

**FIGURE 1.** Secondary structures of the genomic HDV ribozyme (A), the chimpanzee CPEB3 ribozyme (B), and the small model constructs P1, P2, and P4 (C) used for resonance assignment. P1 is shown in red, P1.1 in orange, P2 in green, P3 and L3 in dark blue, and P4 in light blue; the J1/2 and J4/2 linkers are shown in black, the catalytic cytosine is in boldface; the two nucleotides G25·U20 that form a wobble pair only in the presence of Mg\(^{2+}\) (for the CPEB3 ribozyme, see far infra) are circled in A and B; in C, the difference between the chimpanzee and human sequence is shown in gray (G30A substitution); (C) the nucleotides added to the natural sequences of P1, P2, and P4 are shown in gray.
of the CPEB3 proton and nitrogen resonances was obtained using three model hairpin constructs for P1, P2, and P4 (Fig. 1C). We have used NMR, Tb³⁺-induced cleavage and ion counting techniques for a detailed investigation of Mg²⁺ binding to the CPEB3 ribozyme. Our results are compared to the crystal structures of the HDV ribozyme, confirming strong structural parallels of both ribozymes and shedding light on the critical role of site-specific Mg²⁺ binding for the native fold.

RESULTS

The wild-type human and chimpanzee CPEB3 ribozyme constructs are catalytically active

To confirm that the ribozyme constructs were self-cleaving, we performed cotranscriptional cleavage studies of the human and the chimpanzee sequences, which were elongated by 18 nt of the native genomic sequence on the 5'-end. In vitro transcription reactions were stopped after different time intervals and the cleavage products were separated and analyzed on denaturing PAGE gels (Fig. 2). Both constructs self-cleave, as evidenced by the band of the post-cleavage ribozyme control (67mer) and a band corresponding to the cleaved 18mer that intensifies with time. Also, the two DNA template strands, the 24mer (TS) and the 107mer (OT) are visible. The band of the uncleaved 85mer ribozyme is also detected but it is much less intense than the post-cleavage ribozyme, which indicates that the majority of molecules are active. Consequently, the 67mer constructs used in the structural and metal ion binding studies can indeed be derived from ribozyme self-cleavage and therefore are likely to represent the native fold of the post-cleavage ribozyme.

The self-cleavage rates of the human (Fig. 2A) and chimpanzee (Fig. 2B) sequences are distinctly different, although their sequences differ only in nucleotide 30, which is a G in the chimpanzee and an A in the human sequence, the chimpanzee one seems to cleave at least four times faster. This is in agreement with previous studies, which explain the lower cleavage rate of the human sequence by a higher propensity of mispairing between P1/P1.1 and the upstream sequence (Chadalavada et al. 2010).

NMR confirms the HDV-like secondary structure

NMR proton resonance assignment of the CPEB3 ribozyme proved to be challenging because of the severe spectral overlap in the [¹H,¹H]-NOESY spectra, which is mainly due to the large size of the ribozyme. The chimpanzee construct was chosen for NMR studies, as its spectra looked slightly better resolved compared to the human one (probably due to the canonical base-pairing which additionally stabilizes the P1 helix in the chimpanzee construct [C7–G30, see Fig. 1]). All other experiments (Tb³⁺-induced cleavage, DOSY NMR and ion-counting techniques) were performed on the human CPEB3 ribozyme.

Assignments were made on partially deuterated CPEB3 RNA in the absence of Mg²⁺, in which the signal overlap is drastically alleviated. The assignment was supported by the following strategies: (i) G,C, or A,U-¹³C, ¹⁵N-labeled RNA or uniformly ¹⁵N-labeled RNA helped to identify [¹H,¹H]-NOESY or [¹H,¹⁵N]-HSQC correlations belonging to certain nucleotides; (ii) four samples containing only one natural abundance nucleotide (with the remaining three being fully deuterated); and (iii) three small model constructs for the P1 and P2 helices and for the P4 hairpin (Fig. 1C) were produced and analyzed to confirm the assignment of the CPEB3 ribozyme spectra. Combining these data, the majority of the nonexchangeable H1', H2', H6/8, H2 proton resonances, encompassing the 5' strand of P1, P2, P3, P4, and the J4/2 linker, were assigned (Supplemental Table S1). Assignments for J1/2, L3 and P1.1, regions from the center of the ribozyme, could not be obtained. Comparison of the proton chemical shifts of the three model constructs and the full-length CPEB3 ribozyme reveal that P4 chemical shifts are in excellent agreement (Skilandat et al. 2014). The chemical shift values of the P1 and P2 model constructs only approximate those of P1 and P2 within the ribozyme, however, the relative arrangement of chemical shifts within the sequences is the same (Supplemental Figs. S1, S2; Supplemental Tables S2, S3).

For the three long helical domains, P1, P2, and P4, imino protons and nitrogens were fully assigned (Figs. 3, 4; Supplemental Table S1) and all [¹H,¹H]-NOE cross peaks, which are expected from the model constructs could be attributed in the CPEB3 ribozyme (Fig. 3). Please note that in the
discussion below, the nucleotide numbering corresponds to the full-length construct (not to auxiliary helices). Typical NOEs for A-helical arrangements including also several stacking interactions were observed in the NOESY spectra of nonexchangeable protons of P1, P2, and P4 regions. Moreover, characteristic features such as the upfield shifted and very similar H8 chemical shift values of G2 to G5 and their similar and downfield shifted H1′ resonances as well as the far upfield shifted A13H2 and G14H8 are observed both within the CPEB3 ribozyme and in the P1 and P2 model constructs (Supplemental Figs. S1, S2; Supplemental Tables S1–S3). Such upfield shifts are indicative of stacking interactions between the nucleobases and are thus typically found in stable helices. Taken together, the data clearly demonstrate that the longer P1, P2, and P4 helices are formed and base paired according to the proposed secondary structure.

The formation of those helices is additionally confirmed by a J_{NN} HNN-COSY experiment (Fig. 4), which makes use of the N,N coupling across the hydrogen bond as a direct reporter for base pair formation. Except for the signals belonging to G–C base pairs in P1, P2, and P4, three additional peaks that correspond to G–C base pairs are present (Gω, Gβ, and Gγ, Fig. 4; please note that exactly three G–C base pairs are expected in the small P3 helix). Two of the respective imino protons also show NOE cross peaks, which could, however, not be assigned to specific Gs or Cs, due to the lack of conclusive NOE cross peaks (Fig. 3). The presence of the additional imino signals, together with the high upfield shift of G17H8 and a G17H8–C18H6 cross peak, which suggest stacking interactions and the formation of a stable helix, are an indication that the P3 helix might be formed. Assuming that is the case, then the most likely reason for the lack of sequential NOEs is conformational dynamics in P3 and L3/P1.1, which would also explain why we could not assign any NOE correlations for L3 and P1.1. Given that L3 is mostly unpaired and P1.1 also contains one G–C and a less stable U–U base pair, a flexibility of this region is quite probable.

Internucleotide H2′(n)–H6/8(n + 1) and H1′(n)–H6/8(n + 1) NOE cross peaks are observed throughout the 4-nt long J4/2 linker, which contains the active site cytosine. While the lack of further NOEs prohibits a more detailed view on the structure of this linker, the presence of these sequential-walk resonances suggests a rather stable assembly of the J4/2 linker nucleotides stacked on top of each other.

Mg^{2+} is required for a compact fold of the CPEB3 ribozyme

As Mg^{2+} is known to be required by many RNAs to adopt their native structure, and is obligatory for CPEB3 activity, we first assessed the influence of Mg^{2+} on the global structure of the CPEB3 ribozyme. Using Diffusion Ordered Spectroscopy (DOSY) NMR experiments, K⁺ and Mg^{2+} titrations of the ribozyme were performed to determine changes in the hydrodynamic radius (r_{H}) upon addition of different amounts of metal ions (Table 1). The theoretical r_{H} of the CPEB3 ribozyme can only be estimated based on the related HDV ribozyme architecture: We assume that the folded ribozyme has a length a, which corresponds to 19 base pairs, i.e., the sum of base pairs in P2, P3, P4, and P1.1, and a width b of about twice the diameter of an A-form helix. With 0.26 nm and 2.4 nm being the distance between two base pairs and the diameter of a standard A-form RNA, respectively,
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a = 18.026 nm = 4.68 nm, and b = 2.24 nm = 4.8 nm. Because b/a < 2, a spherical model can be applied, in which \( r_H \) is defined as half of the length (Eimer et al. 1990; Lapham et al. 1997). Consequently, \( r_H \) should be between 2.34 nm and 2.4 nm as a first estimate.

CPEB3 does not achieve the native fold in the absence of significant amounts of counter ions (Table 1): The large hydrodynamic radius of 3.91 nm in the absence of salt is probably an averaged value of different extended states. Also, 10 mM K+ are not sufficient for the ribozyme to obtain a compact fold, which is not surprising, considering the intricate pseudoknot topology of CPEB3. A significant compaction is only achieved in the presence of 100 mM K+ (2.83 nm), but the addition of small concentrations of Mg2+ further reduces the \( r_H \) (2.56 nm in 2 mM Mg2+). Mg2+ titrations were performed with the partially deuterated CPEB3 RNA (Fig. 5; Supplemental Fig. S3; Supplemental Table S1), as well as with the four samples containing one natural abundance and three fully deuterated nucleotides, and with the P1, P2, and P4 constructs (data not shown). All titrations showed consistent results. Nonexchangeable \([1^H,1^H]-NOESY\) spectra show that six regions are strongly affected by the addition of Mg2+: (i) the central part of the P2 helix (G11–C12 and A13H2 broaden to baseline, A13–G14 H8 and H1’ and G65–G66 H1’ and H2’ [from the opposite side of the strand] broaden and shift, Fig. 5A; Supplemental Table S1); the (ii) J4/2 linker with the catalytic C57, of which the respective cross peaks (C57–A59) are broadened to baseline at 5 mM Mg2+, indicating a strong interaction (Fig. 5A); (iii) A60H8 in the upper part of the J4/2 linker is not broadened, but strongly shifted (0.273 ppm, Fig. 5A; Supplemental Table S1); Proton resonances of (iv) the base pairs adjacent to the P4 tetraloop (G50, A51), (v) in the P4 stem (A40, G41) and (vi) in the P1 stem (G3–G5) also display pronounced chemical shift changes (Fig. 5A; Supplemental Table S1).

To probe the above described Mg2+ binding sites for possible outer-sphere coordination, \([1^H,1^H]-NOESY\) spectra of the nonexchangeable (data not shown) and exchangeable protons were recorded in the presence of 1.5 mM [Co(NH3)6]3+ (Fig. 5B, purple spectrum). The [Co(NH3)6]3+ protons show cross peaks to G2H1, G4H1, G5H1 (Fig. 5B), and G5H8 (data not shown) of P1, suggesting that the P1 stem prefers outer-sphere binding. More NOE cross peaks are observed between the protons of [Co(NH3)6]3+ and G41H8, C54H6, and U55H3 in P4, demonstrating outer-sphere coordination to the P4 stem. [Co(NH3)6]3+ does not show cross peaks to the Mg2+ binding sites in P2 and in the L4 tetraloop. Consequently, Mg2+ binding to these sites requires a partial dehydration of the Mg2+ ion. The position and coordination preference of both binding sites in the P4 domain are in line with what has been observed in the P4 model construct (Sklandat et al. 2014). In the J4/2 linker, cross peaks to the [Co(NH3)6]3+ protons are only observed for A60H1’ and A60H8, suggesting this metal ion and assessed, whether inner- or outer-sphere binding occurs at each site. The term “inner-sphere” describes a direct coordination of Mg2+ to the RNA, e.g., to N7 of guanine or to the nonbridging phosphate oxygens (Freisinger and Sigel 2007; Sigel and Pyle 2007). Outer-sphere coordination, in contrast, is mediated by water ligands in the first coordination shell of the Mg2+ ion. The latter can be probed using hexammincobalt(III), [Co(NH3)6]3+, as an exchange-inert substitute of hexaaquamagnesium(II) (Rowinska-Zyrek et al. 2013).

Mg2+ binding is detected through chemical shift changes and by line broadening of the proton resonances in \([1^H,1^H]-NOESY\) spectra. In the case of [Co(NH3)6]3+, NOE cross peaks between the protons of the amine ligands and the coordinating nucleotide can be observed (Robinson and Wang 1996; Rowinska-Zyrek et al. 2013). Mg2+ titrations were performed with the partially deuterated CPEB3 RNA (Fig. 5; Supplemental Fig. S3; Supplemental Table S1), as well as with the four samples containing one natural abundance and three fully deuterated nucleotides, and with the P1, P2, and P4 constructs (data not shown). All titrations showed consistent results. Nonexchangeable \([1^H,1^H]-NOESY\) spectra show that six regions are strongly affected by the addition of Mg2+: (i) the central part of the P2 helix (G11–C12 and A13H2 broaden to baseline, A13–G14 H8 and H1’ and G65–G66 H1’ and H2’ [from the opposite side of the strand] broaden and shift, Fig. 5A; Supplemental Table S1); the (ii) J4/2 linker with the catalytic C57, of which the respective cross peaks (C57–A59) are broadened to baseline at 5 mM Mg2+, indicating a strong interaction (Fig. 5A); (iii) A60H8 in the upper part of the J4/2 linker is not broadened, but strongly shifted (0.273 ppm, Fig. 5A; Supplemental Table S1); Proton resonances of (iv) the base pairs adjacent to the P4 tetraloop (G50, A51), (v) in the P4 stem (A40, G41) and (vi) in the P1 stem (G3–G5) also display pronounced chemical shift changes (Fig. 5A; Supplemental Table S1).

Locating Mg2+ binding sites by NMR spectroscopy

Having established that Mg2+ ions are necessary to reach the native fold of the ribozyme, we located the binding sites of this metal ion and assessed, whether inner- or outer-sphere binding occurs at each site. The term “inner-sphere” describes a direct coordination of Mg2+ to the RNA, e.g., to N7 of guanine or to the nonbridging phosphate oxygens (Freisinger and Sigel 2007; Sigel and Pyle 2007). Outer-sphere coordination, in contrast, is mediated by water ligands in the first coordination shell of the Mg2+ ion. The latter can be probed using hexammincobalt(III), [Co(NH3)6]3+, as an exchange-inert substitute of hexaaquamagnesium(II) (Rowinska-Zyrek et al. 2013).

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that the 3' part of the linker is involved in an outer-sphere binding site, whereas the 5' part preferentially binds Mg2+ in an inner-sphere manner, which is expected for the active site cytosine (Nakano et al. 2001; Gong et al. 2007, 2009).

The [1H,1H]-NOESY spectra of the exchangeable protons in the presence of Mg2+ or [Co(NH3)6]3+ reveal another binding site at the G1·U36 wobble base pair. This site is identified by broadening of the cross peaks involving U36H3 and G1H1 and G2H1 and by an NOE between U36H3 and the protons of [Co(NH3)6]3+ (Fig. 5B).

In total, up to eight sites in the CPEB3 ribozyme are affected by Mg2+: (i) the tetraloop of P4, (ii) the upper half of the P2 helix, and (iii) the nucleotides surrounding C57 prefer inner-sphere coordination of Mg2+. The (iv) G1-U36 wobble, (v) the upper part of the P4 helix, and (vi) A60 located in the upper part of the J4/2 linker can also accommodate outer-sphere coordinated Mg2+, whereas (vii) the middle part of the P1 stem seems to be the only site that prefers [Co(NH3)6]3+ over Mg2+. The eighth and probably one of the most important Mg2+ binding sites, the G25-U20 wobble, is discussed in the paragraph below.

**Mg2+-induced structural changes**

In the [1H,1H]-NOESY spectrum in 90% H2O as well as in the [1H,15N]-TROSY spectrum, new peaks appear upon addition of Mg2+ (bold labels in Figs. 5B, 6A). Judging by their 15N chemical shifts, these new peaks correspond to four guanines (Gd, Ge, G6, G8) and five uracils (Ua, Ub, Uc, Ud, Ue).

There are six guanines (G17, G19, G25, G28, G37, and G58), and four uracils (U20, U21, U26, U38) in the sequence, whose imino protons and nitrogens were not yet conclusively assigned. Given that another three guanine resonances, Gp, Gq, and Gs (suggested to be the three Gs from the P3 helix), were already observed in the absence of Mg2+ in the JNN HNN-COSY (Fig. 4) or NOESY (Fig. 3), there is at least one surplus correlation of both a uracil and a guanine, possibly as a result of conformational exchange. Four of the six unassigned guanine H1 are located in G–C base pairs (Ga, Gb, Gc, and Ge) (Fig. 6B). This is an important finding, given that exactly four G–C base pairs are expected to be formed for P3 and P1.1, according to the proposed secondary structure (Fig. 1B). We hence assigned the newly appearing guanines and uracils based on the [1H,1H]-NOESY correlations as far as possible.

The most intense cross peak that appears in the presence of Mg2+ (and also in the presence of [Co(NH3)6]3+) is the one between a GH1 and a UH3 (Uc–Gg resonance, Fig. 5B). The Uc–Gg NOE cross peak is in a position that is typical for a G·U wobble pair. As the formation of an extra G·U wobble at the base of the P3 helix has been suggested for the HDV ribozyme (Krasovska et al. 2006; Chen et al. 2010; Veeraraghavan et al. 2011; Kapral et al. 2014), we speculated that the corresponding G25-U20 wobble forms as well in the CPEB3 ribozyme upon addition of Mg2+, giving rise to the observed NOE correlation. The hypothesis was both tempting and probable, however it longed for more substantial evidence, since theoretically, also other scenarios might
have been occurring, such as, e.g., the formation of a G19·U26 wobble. To test our theory, we studied the imino proton behavior of a U20C mutant, both in the presence and absence of Mg$^{2+}$ ions. In monovalent metal ions only, the mutant seems to form the same base-pairing interactions as the wild-type CPEB3 construct (also, no canonical C20–G25 base pair is formed, or the signal is overlapped with other cross peaks). In the presence of 5 mM Mg$^{2+}$, no additional peaks appear (Supplemental Fig. S4) and a further increase of Mg$^{2+}$ concentration does not cause any changes, apart from a general broadening of observed signals. This clearly confirms that in the case of the wild-type construct, the Uc·Gg resonance which appears in the presence of magnesium(II) is indeed the U20·G25 wobble. This is an important finding which again shows a structural and functional similarity between the CPEB3 and HDV ribozymes—in the HDV ribozyme, this G-U wobble is thought to present the catalytically active Mg$^{2+}$ ion to the active site cytosine (Chen et al. 2010). This correlates well with our finding that this signal is only visible after the addition of Mg$^{2+}$. UcH3, now assigned to U20H3, has a strong cross peak to the [Co(NH$_3$)$_6$]$^{3+}$ protons. Since the G-U wobble in the active site of the HDV ribozymes makes outer-sphere contacts to Mg$^{2+}$ with its minor groove, the observed cross peak is in agreement with our suggested assignment.

Although the presence of the Uc, Gc, Gg, Gp, and Ud cross peaks suggests the formation of the small P3–P1.1 pseudoknot (according to the proposed secondary structure) and strong evidence suggests that Uc–Gg is the substantial U20·G25 wobble (see above), there are too few cross peaks to relate the imino protons within P3 to other helices. We suspect that the combined effects of Mg$^{2+}$-induced line broadening and of flexibility in the P3–P1.1 pseudoknot cause the low intensity or lack of NOE correlations. One argument for the flexibility is the fact that at least two surplus NH correlations are observed in the [1H,15N]-TROSY spectrum (Fig. 6A). Another feature that might indicate conformational exchange is the prominent X–Gf cross peak appearing upon Mg$^{2+}$ addition (Fig. 5B). X is either a uracil (Uf) or a guanine overlapped with G41, according to the TROSY correlations (Fig. 6A). Both X and Gf are correlated to another unpaired guanine Gd. It is not clear which nucleotides could give rise to this cross peak, as the sequence does not contain unpaired guanines close to each other. It is thus possible that X–Gf is not actually an NOE but an exchange cross peak between two different conformations of the same proton.

Beyond any doubt, Mg$^{2+}$, apart from binding to the previously described sites in P1, P2, and P4, has a very strong influence on the nucleotides in the remaining ribozyme domains, forming the inner pseudoknot and active site. Mg$^{2+}$ induces a gain of structure in this region by enabling additional internucleotide contacts or by stabilizing them. This is in perfect agreement with the overall compaction of the ribozyme, which was observed in the presence of Mg$^{2+}$ and it again underlines the importance of this metal ion for the structure and function of the CPEB3 ribozyme.

**Tb$^{3+}$-induced cleavage reveals metal ion binding in the P3–P1.1 pseudoknot**

Tb$^{3+}$-induced RNA cleavage experiments were carried out in order to confirm the Mg$^{2+}$ binding sites determined from the NMR titration data and to obtain information on metal ion binding sites in the inner pseudoknot (P1.1, P3, L3) of the...
CPEB3 ribozyme, where the lack of resonance assignment prevents an NMR study of the metal ion binding sites. Tb³⁺ can be used to map Mg²⁺ binding sites in RNA through hydrolytic cleavage experiments (Giesiolka et al. 1989; Sigel and Pyle 2003; Erat and Sigel 2011; Choudhary et al. 2014). Lanthanide ions are assumed to be good mimics of Mg²⁺, since they occupy the same binding sites in RNA molecules, but bind with at least three orders of magnitude higher affinity (Kayne and Cohn 1974; Sigel and Pyle 2003; Erat and Sigel 2011). In a Tb³⁺-induced cleavage experiment, Tb³⁺ is added to a folded ribozyme, where it binds to RNA, displacing the previously bound Mg²⁺ ions. The partially deprotonated Tb³⁺ aqua-complex deprotonates the 2′-OH of a close-by nucleotide, which enables a nucleophilic attack of the 2′-oxyanion on the adjacent phosphodiester bond. This triggers backbone scission and the resulting fragments can be separated by PAGE and visualized to identify the metal ion binding sites (Fig. 7; Choudhary et al. 2014).

Tb³⁺-induced cleavage of the human CPEB3 ribozyme takes place in three main regions: the P2 helix (C12), the P4 tetraloop (G48), and several positions in the core of the ribozyme encompassing P1.1, P3, L3, J4/2, and the upper part of P4. The most intense cleavage bands correspond to those of the junction of P4 to J4/2 (G56, C57, G58) and to P1.1 (C22) and L3 (U20, A23). While the P2 stem and L4 clearly represent two distinct binding sites, the number of binding sites in the P1.1–P3 pseudoknot cannot be determined, as the distance and relative orientation of the affected nucleotides in the folded structure are not known (Fig. 7).

However, judging by the number of affected nucleotides and the secondary structure, we estimate two to four ions to bind to the ribozyme core comprising the active site and the P1.1–P3 pseudoknot.

The binding sites obtained from Tb³⁺-induced cleavage confirm Mg²⁺ binding to L4, J4/2, and the P2 stem, which was detected by NMR spectroscopy. The binding sites of [Co(NH₃)₆]³⁺ in the P4 and P1 stem are not occupied by Tb³⁺. This is probably due to different local structures being required for [Co(NH₃)₆]³⁺ binding and for permitting a Tb³⁺-induced cleavage reaction (Soukup and Breaker 1999).

**Counting bound Mg²⁺ ions**

We used two different ion counting methods to independently estimate the number of Mg²⁺ ions bound to the CPEB3 ribozyme. The first method uses the dye 8-hydroxyquinoline-5-sulfonic acid (HQS), whose 1:1 Mg²⁺–HQS complex is fluorescent (Romer and Hach 1975; Grilley et al. 2009). In the second method, equilibrium dialysis of the RNA sample with a known concentration of free Mg²⁺, followed by an atomic absorption spectroscopy (AAS) measurement of total Mg²⁺ concentration of both solutions is used (Bina-Stein and Stein 1976; Das et al. 2005; Leipply et al. 2009).

For our measurements, both methods gave corresponding values within error limits, showing that the four Mg²⁺ ions are associated with the CPEB3 ribozyme. This number seems to be independent of the concentration of monovalent ions in the buffer. At 100 mM KCl, fluorescence titrations yielded 4.13(±0.32) Mg²⁺ ions bound to the ribozyme, while dialysis results suggested that 3.95(±0.25) Mg²⁺ ions bound. At 50 mM KCl, 4.36(±0.24) and 4.41(±0.05) Mg²⁺ ions were detected by the fluorescence and dialysis methods, respectively. A drastically increased concentration of monovalent ions did not significantly alter the result. At 2 M KCl the number of bound ions detected by fluorescence is 4.33 (±0.08), and 3.96 as determined by equilibrium dialysis and AAS (Table 2).

The fact that four bound Mg²⁺ ions are counted in the presence of different concentrations of K⁺ indicates that these four ions are site-bound with higher affinity and occupancy factors and less diffusion into the surrounding ion cloud (Das et al. 2005). Probably, they are bound to specific sites in noncanonical regions in the center of the ribozyme, which are at least partially protected from the bulk solvent, i.e., in the catalytic core of the enzyme (around C57) and the inner P1.1–P3 pseudoknot.

**DISCUSSION**

In this study, we demonstrate that the post-cleavage CPEB3 ribozyme adopts its global fold in monovalent ions alone but that Mg²⁺ binding to several sites is required to fully
form the inner pseudoknot, active site, and the compact native structure. The P1–P2 pseudoknot and the P4 domain of the ribozyme are base paired according to the HDV-like secondary structure proposed earlier (Salehi-Ashtiani et al. 2006), independent of the presence of Mg²⁺. These structures (P1, J1/2, P2, P4, and J4/2) account for the largest part of the ribozyme sequence (52 of 67 nt, 21 of 26 predicted base pairs) and are constrained in their orientation due to the short linkers in between the helices. Accordingly, the overall fold of the CPEB3 ribozyme is largely determined by base-pairing and thus corresponds to the one of the HDV ribozyme, which is subject to very similar structural constraints.

So far, the extent to which the inner pseudoknot is formed is not entirely clear. Judging by the number of G–C correlations in the JNN HNN-COSY, probably P3 is at least partially formed in the absence of Mg²⁺. Nevertheless, the addition of Mg²⁺ has a very pronounced effect on the CPEB3 structure and is known to be strictly required for catalysis (Salehi-Ashtiani et al. 2006). Unfortunately, the severe line broadening upon addition of Mg²⁺ prevents a detailed pH titration study to investigate a possible shifted pKₐ for C57, as has been suggested for the HDV ribozyme (Gong et al. 2007). However, Mg²⁺ significantly reduces the ribozyme’s hydrodynamic radius at near-physiological concentrations and binds to several sites in the ribozyme, thereby promoting the formation of additional structure in the catalytic core. Our data provide strong evidence that the full predicted secondary structure (Salehi-Ashtiani et al. 2006) is formed only in the presence of Mg²⁺. This includes the G25-U20 wobble at the base of P3 and the smaller P1.1–P3 pseudoknot, but that the latter retains some flexibility. This is in line with the data on the HDV ribozyme, whose L3 region has been shown to be mobile, a feature that is probably meaningful for self-cleavage activity (Krasovska et al. 2006).

To clarify the structural role of Mg²⁺ in the CPEB3 (and HDV) ribozyme, we now discuss the metal ion binding sites we have observed in more detail and compare them to those observed in the HDV ribozyme crystal structures (Fig. 8). Our NMR data show evidence for the formation of an outer-sphere Mg²⁺ binding to the G-U wobble base pair (G25-U20). A corresponding Mg²⁺ is supposed to take part in active site formation in the HDV ribozyme (Mg²⁺ [1] in Fig. 8A,B, red nucleotides in Fig. 8C). As this site presumably presents the catalytic, outer-sphere bound Mg²⁺ ion to the catalytic cytosine and the scissile bond between –1 nucleotide and G1, it is possible that the strong line broadening effect that we observe for C57 and G58 as well as the Tb³⁺-induced cleavage band at C57 are due to the same metal ion. Also, the Tb³⁺-induced cleavage bands at C22 and A23, which according to the HDV crystal structure are in direct vicinity (Fig. 8A), may result from Tb³⁺ binding to the active site. This binding site illustrates very well the double role of Mg²⁺ in structure and function for this ribozyme. Apart from its role in the catalytic mechanism, it also stabilizes the interaction of three different regions of the secondary structure being the J4/2 linker, the P1 terminus, and the base of the P3 helix.

The second Mg²⁺ binding site of possible structural relevance for the CPEB3 and the HDV ribozymes lies between the sugar–phosphate backbones of the two adenines in J4/2.

### TABLE 2. Number of Mg²⁺ ions bound to CPEB3

| [K⁺]/mM | Equilibrium dialysis | HQS fluorescence |
|---------|----------------------|-----------------|
| 50      | 4.41 ± 0.05          | 4.36 ± 0.24     |
| 100     | 3.95 ± 0.25          | 4.13 ± 0.32     |
| 2000    | 3.96*                | 4.33 ± 0.08     |

Values represent arithmetic means and errors of a series of measurements from two individual experiments.

*Only one measurement was performed.

**FIGURE 8.** Location of metal ion binding sites in the HDV and human CPEB3 ribozymes. (A,B) Crystal structure of the inhibited HDV ribozyme, pdb entry 3NKB (Chen et al. 2010): Full structure (A) and a close up (B) of the active site. Mg²⁺ ions are shown as dark blue spheres, J4/2 is shown in red, and the G25-U20 wobble formed only in the presence of Mg²⁺ is shown in yellow. (C) Mg²⁺ binding sites in the CPEB3 ribozyme. Each color represents a different binding site. The difference between the chimpanzee and human sequence (G30A substitution) is shown in gray.
and the P1 stem in the HDV crystal structure (Mg2+ [2] in Fig. 8A, blue nucleotides in Fig. 8C). We have demonstrated [Co(NH3)6]3+ (and to a lesser extent Mg2+) binding to the corresponding sites of the CPEB3 ribozyme. [Co(NH3)6]3+ inhibits ribozyme activity indicating that at least one partially dehydrated Mg2+ is needed. The coordination of such a partially dehydrated Mg2+ in this location might help to overcome backbone repulsion between J4/2 and P1, which is another aspect of how Mg2+ can promote the formation of a compact native structure of both CPEB3 and HDV. Also, the outer-sphere binding site at the G1-U36 wobble (khaki nucleotides in Fig. 8C) (G1–U37 in HDV) occurs in both ribozymes and has been suggested to stabilize the P1.1 mini-helix (Veeraraghavan et al. 2011), which maintains stacking interactions with P1. Probably, there are further binding sites for Mg2+ in the core region of the CPEB3 ribozyme (orange and green-blue nucleotides in Fig. 8C), which requires stabilization due to the higher content of unpaired bases and the proximity of the different backbone segments. The binding sites in P2 and P4 (yellow, green, and violet nucleotides in Fig. 8C) are more likely to stabilize local structure such as the UGGU tetraloop (Skilandat et al. 2014) because these regions should not be in direct contact with other domains. In conclusion, a direct comparison of Mg2+ binding sites in CPEB3 and HDV ribozymes is not as straightforward as it might seem because sites that stabilize the local structures of separate helices are not conserved, as are the sequences of those helices not conserved. On the contrary, sites that are involved in the stabilization of the pseudoknot or that are necessary for catalysis, bind Mg2+ ions both in the case of the HDV and the CPEB3 ribozyme.

The question remains, which are the four binding sites that we counted in the CPEB3:Mg2+ stoichiometry experiments? As these ions remain bound in up to 2 M KCl, they are tightly bound to sites, which are at least partly shielded from the solvent or offer a favorable geometry for inner-sphere contacts between the RNA and the Mg2+ ion (Das et al. 2005). Accordingly, the three binding sites in the stem of P1, P4, and at the G1-U36 wobble (blue, green, and khaki nucleotides in Fig. 8C) are unlikely to be counted, because they can accommodate outer-sphere coordination and, judged by the HDV ribozyme crystal (Fig. 8) and the P4 NMR structure (Skilandat et al. 2014) are accessible for the solvent. It is not possible to univocally determine which metal ion binding sites are detected by the ion counting. Based on our combined data, we hypothesize that one of them is between the active site cytosine C57 and the G25·U20 wobble, and the second one is also in the core of the structure, possibly between the sugar–phosphate backbones of the two adenines in J4/2 and the P1 stem. The two remaining binding sites could be located in the P2 stem and P4 tetraloop (yellow and violet nucleotides in Fig. 8C). Both sites offer a suitable geometry for inner-sphere coordination of Mg2+ and this type of binding could be thermodynamically and kinetically more stable. It is remarkable that a comparably small structure such as the CPEB3 RNA contains four such high-affinity sites, and underscores the extraordinary significance of Mg2+ for the particular structure and mechanism of this ribozyme.

The CPEB3 and HDV ribozymes are different from all other small ribozymes investigated so far in that they crucially depend on the presence of divalent metal ions for catalysis, both in low and in high concentrations of monovalent ions. Also, their nested double pseudoknot fold is far more complicated than the secondary structures of the other small ribozymes. The data presented here (i) confirm that the CPEB3 ribozyme folds the same way as the related HDV and (ii) illustrate why the Mg2+ dependence does not only stem from the catalytic mechanism but is inherent to the particular structure of the CPEB3 and HDV ribozymes, by completely structuring their inner catalytic core.

MATERIALS AND METHODS

DNA oligonucleotide templates of the human and chimpanzee CPEB3 ribozymes and their corresponding elongated, precleavage constructs were purchased from Microsynth. Natural abundance, partially deuterated and fully deuterated nucleoside 5′-triphosphates were purchased from GE Healthcare and Cambridge Isotope Laboratories and uniformly 15N-labeled and 13C, 15N-labeled nucleotides from Silantes. The T7 RNA polymerase used for in vitro transcription was produced in house according to standard procedures (Gallo et al. 2005). [Co(NH3)6]3+Cl3 and 8-hydroxyquinoline-5-sulfonic acid were purchased from Sigma Aldrich and 100% D2O was purchased from Armar Chemicals. The electrophoresis apparatus Elutrap was from Whatman. For desalting, Vivaspin Concentrators (5000 MWCO for the full-length constructs and 2000 MWCO for the auxiliary short ones) from Sartorius-Stedim biotech were used. Thermosensitive shrimp alkaline phosphatase and T4 polynucleotide kinase as well as the appropriate buffers were purchased from Promega.

Cotranscriptional self-cleavage assays of CPEB3

In vitro transcription reactions with an elongated 107mer DNA template (designed to transcribe into the CPEB3 ribozyme with additional 18 nucleotides at the 5′ end) were performed in aliquots of 40 µL in the presence of 5 mM Mg2+. For human and chimpanzee CPEB3, single aliquots were removed after 1, 2, 3, 4, 5, and 7 h or 15, 30, 45, 60, 90, and 120 min, respectively, and immediately frozen in liquid nitrogen to stop both transcription and cleavage. All samples were then centrifuged for 15 s and a mixture of 82% formamide and 10 mM EDTA was added to the supernatant (1:1). The transcription/cleavage products were subsequently separated by 12% denaturing PAGE.

NMR sample preparation

All constructs were synthesized by in vitro transcription with T7 polymerase from a double-stranded DNA template (Gallo et al. 2005). Reaction mixtures typically contained 4.5–6 mM of each NTP, 0.9–1.2 µM of each strand of the synthetic, PAGE-purified, double-stranded DNA template, 30–50 mM MgCl2, 40 mM Tris–HCl
(pH 7.5), 40 mM DTT, 2 mM spermidine, and 0.01% Triton X. The amount of T7 RNA polymerase was adapted according to activity of each enzyme batch. Transcription was allowed to proceed at 37°C for 6 h. The transcribed RNA was purified by denaturing 12% PAGE, UV-shadowed, excised from the gel, and recovered by electroelution. During ultrafiltration in Vivaspin devices, the RNA was washed repeatedly with 2 M KCl, pH 8 to remove Tris and afterward several times with water. After lyophilization, the sample was dissolved in 250 µL D2O or 90% H2O, 10% D2O containing an appropriate amount of KCl (100 mM for CPEB3 and P1, 20–30 mM for P2 and P4) and 10 µM EDTA. The pH was adjusted to 6.8 in H2O or 6.4 in D2O, corresponding to a pD of 6.8 (Glasoe and Long 1960). The RNA concentration of the samples varied between 0.4 and 0.9 mM and was determined on a Varian Cary 100 Scan UV-Vis spectrophotometer by using extinction coefficients of ε260 of 203.4 mM⁻¹ cm⁻¹ (P1), 212 mM⁻¹ cm⁻¹ (P2), and 725 mM⁻¹ cm⁻¹ (CPEB3). Before acquisition of NMR data, all samples were annealed by 2-min incubation at 90°C, followed by rapid cooling in ice water.

NMR spectroscopy

All spectra were recorded on a Bruker Avance 600 MHz spectrometer with a 5-mm CRYO TCI inverse triple-resonance probehead with z-gradient coil or on a Bruker Avance 700 MHz spectrometer with a 5-mm CRYO TXI inverse triple-resonance probehead with z-gradient coil. All samples contained 0.4–0.9 mM of sample being either natural abundance RNA, partially deuterated [5, 3, 4, 5, 6, 7, 8]H₂RNA, a combination of fully deuterated and wild-type RNA, G,C or A,U-C, 13N-labeled RNA or uniformly 15N-labeled RNA. Nonexchangeable proton resonances were assigned from [1H,1H]-NOESY spectra, typically recorded with a mixing time of 250 ms at 298 K, in 100% D2O. Suppression of the residual water signal was achieved by presaturation pulses. Exchangeable proton resonances were assigned using [1H,1H]-NOESY spectra with a WATERGATE pulse sequence for water suppression recorded in 90% H2O/10% D2O. To aid assignment, F1,F2-filtered [1H,1H]-NOESY spectra of G,C, 15N-labeled CPEB3 samples were recorded. 13N resonances of CPEB3 were assigned from [1H,15N]-TROSY spectra recorded at 278 K in 90% H2O/10% D2O. Base pair formation was verified by recording 32P-labeled CPEB3 in aliquots of 60 pmol/50 µL by adding 5 µL of 32P-labeled RNA had to be removed prior to 5′-end labeling with 32P. CPEB3 RNA was dephosphorylated using thermosensitive shrimp alkaline phosphatase (TSAP) in aliquots of 60 pmol/50 µL. Additional 3 µL of TSAP was added after 15 min. Two rounds of standard phenol–chloroform extraction and subsequent ethanol precipitation were performed to remove the TSAP. The RNA pellet was dissolved in a buffer of 10 mM MOPS, 1 mM EDTA, pH 6.5.

Mg²⁺ and [Co(NH₃)₆]³⁺ titrations

For metal-ion titrations, a 0.4–0.9 mM of partially deuterated CPEB3 and four samples with only one natural abundance nucleotide (with the remaining three being fully deuterated) were titrated at 298 K with MgCl₂ in steps of 0, 0.5, 1, 1.5, 2, 2.5, 3.5, 5, 9, and 18 mM and a [1H,1H]-NOESY spectrum of the exchangeable (recorded at 278 K, sample in 90% H2O and 10% D2O) and of nonexchangable (recorded at 298 K, sample in 100% D2O) proton regions were recorded at each step. In addition, a [1H,1H]-NOESY spectrum with an excitation sculpting pulse sequence for water suppression was recorded in the presence of 1.5 mM and 2.5 mM [Co(NH₃)₆]³⁺ at 298 K. Cross peaks between RNA protons and [Co(NH₃)₆]³⁺ protons were assigned in each spectrum. Mg²⁺-induced chemical shift perturbations of the nonexchangable protons of the P1 and P2 model constructs were determined as described for the full-length CPEB3, titrating samples with increasing amounts of MgCl₂ (0, 0.5, 1, 1.5, 3, and 5 mM for P1 and 0, 1, 2, 3, 5, and 7 mM for P2). Mg²⁺-induced chemical shift perturbations of the exchangeable protons of the P2 model construct were obtained by titrating a sample with increasing amounts of MgCl₂ (0, 1, 2, 3, 4, 5, 6.5, 8, 9 mM), recording a 1D-1H spectrum in each step.

K⁺ and Mg²⁺-induced changes of the hydrodynamic radius of CPEB3 were determined by titrating a sample with increasing amount of KCl and MgCl₂ (0, 10, 100 mM of KCl plus 2, 5, 10, 20 mM of MgCl₂) recording a DOSY experiment at each step and calculating D and rH for each concentration. Prior to data acquisition, the pH was readjusted after having reached 100 mM KCl, and 10 and 20 mM MgCl₂, respectively. In addition, the RNA was reannealed after both additions of KCl. DOSY experiments with excitation sculpting water suppression were recorded with a diffusion delay Δ of 400 msec and a gradient duration d of 2 msec at 298 K in D2O. The gradient strength was raised incrementally from 1.7 to 32.4 G/cm in 56 steps. The peak areas at each gradient strength were determined in TopSpin (Bruker BioSpin AG). By plotting A₀ against the square of the gradient strength (G²) and fitting to a straight line, the diffusion coefficient can be obtained from

\[
\ln \frac{A}{A_0} = -D\gamma^2 G^2 d (\Delta - \frac{d}{3})
\]

where \( \gamma = 2.67522205 \times 10^{-8} \text{ rad}^{-1} \text{T}^{-1} \) is the proton gyromagnetic ratio. The hydrodynamic radius is then calculated according to the Stokes-Einstein equation

\[
r_H = \frac{k_B T}{6\pi \eta_D}
\]

where \( k_B = 1.3806488 \times 10^{-23} \text{ J/K} \) is the Boltzmann constant and \( \eta_D = 1.0998 \) mPa is the viscosity of D2O.

Tb³⁺-induced cleavage reactions

The 5′-triposphates resulting from in vitro transcription of CPEB3 RNA had to be removed prior to 5′-end labeling with Tb³⁺. CPEB3 RNA was dephosphorylated using thermostable shrimp alkaline phosphatase (TSAP) in aliquots of 60 pmol/50 µL. Additional 3 µL of TSAP was added after 15 min. Two rounds of standard phenol–chloroform extraction and subsequent ethanol precipitation were performed to remove the TSAP. The RNA pellet was dissolved in a buffer of 10 mM MOPS, 1 mM EDTA, pH 6.5.

For Tb³⁺ radiolabeling, 480 pmol of dephosphorylated CPEB3 in 31 µL ME-buffer was mixed with 4.5 µL T4 polynucleotide kinase and 4.5 µL of the commercial kinase buffer (5×), and 5 µL of Tb³⁺-γ-ATP (6000 Ci/mmol, 150 µCi/µL) to a total volume of 45 µL. Phosphorylation was left to proceed for 30 min at 37°C and 300 rpm and then quenched by adding 45 µL of formamide loading buffer (82% formamide, 10 mM EDTA, pH 8, 0.16% xylene cyanol, 0.16% bromophenol blue). Tb³⁺-labeled CPEB3 was purified by 8% PAGE. The CPEB3 RNA band was visualized by phosphorimaging, excised, crushed, and mixed with crush and soak buffer (10 mM

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MOPS, 1 mM EDTA, 250 mM NaCl, pH 6.5). The RNA was eluted into the buffer by shaking for 90 min at 4°C. The supernatant was precipitated with ethanol to recover the RNA. The amount of radiolabeled CPEB was determined by scintillation counting.

For Tb\(^{3+}\)-induced cleavage reactions, 2.4–2.5 nM of the 32P-labeled CPEB RNA were refolded in 25 mM MOPS (pH 7), 100 mM KCl. One micromole of cold CPEB RNA was added to reduce the extent of nonspecific RNA degradation by Tb\(^{3+}\) and to facilitate RNA precipitation (Harris and Walter 2003). For annealing, the RNA was heated to 90°C for 90 s, then transferred to ice and 5 or 20 mM MgCl\(_2\) was added immediately. Refolding was left to proceed for at least 15 min on ice. Tb\(^{3+}\)-induced cleavage was carried out in aliquots containing typically 22.5 fmol of refolded 32p-labeled CPEB3 RNA. An increasing amount of TbCl\(_3\) [0–100 µM, in 5 mM sodium(1) cacodylate, pH 5.5], was added to each aliquot and the mixtures were incubated for 30 min at RT. Cleavage reactions were quenched by adding a mixture of 82% formamide and 10 mM EDTA at a 1:1 ratio to each aliquot and by subsequent ethanol precipitation of the cleaved CPEB3 RNA. The RNA fragments were separated by 16% denaturing PAGE and visualized on a phosphor-imager. On each gel, an alkaline hydrolysis ladder (OH\(^{–}\)) and a T1-ribonuclease cleavage ladder of CPEB3 were loaded (Harris and Walter 2003). The OH\(^{–}\) ladder was prepared by incubating the same amount of radiolabeled CPEB3 as used in the Tb\(^{3+}\)-induced cleavage reactions in 50 mM sodium carbonate buffer (pH 9) and 1 mM EDTA for 4–5 min at 90°C and by quenching with formamide loading buffer (1:1). The T1 ladder was generated by incubation of the same amount of CPEB3 with 0.1 µM T1 in 25 mM sodium citrate buffer (pH 5), 55%–60% formamide, ~7 mM EDTA at 55°C for 4–5 min. T1 digestion was quenched by adding a 7:3 mixture of water/formamide loading buffer (1:1).

Mg\(^{2+}\) counting with 8-hydroxyquinoline-5-sulfonic acid (HQS)

The Mg\(^{2+}\)-dependent indicator dye 8-hydroxyquinoline-5-sulfonic acid (HQS) served as a Mg\(^{2+}\) chelator and a fluorescent reporter at the same time (Mg\(^{2+}\) binds to HQS at a 1:1 ratio, strongly enhancing emission at 550 nm). Measurements with HQS were carried out at 298 K on a Varian Carry Eclipse Fluorescence Spectrometer at pH 7, with excitation wavelength 400 nm and with emission wavelength 550 nm (as previously described by Romer and Hach 1975; Grilley et al. 2009). Two samples (0.1 mM HQS, 20 mM MOPS buffer, 50 mM, 100 mM, or 2 M KCl), one of which contained 24 µM of CPEB3 RNA, were titrated with Mg\(^{2+}\) in exactly the same way (2–260 µM Mg\(^{2+}\)) in 29–42 steps. In the second sample, CPEB3 ribozyme (annealed by a 2-min heating at 90°C and rapidly cooled on ice to ensure the proper folding) was present. After each MgCl\(_2\) addition, the sample was mixed thoroughly and the fluorescence intensity at 550 nm was recorded.

The effect of the concentration of K\(^{+}\) ions on Mg\(^{2+}\) association to the RNA was found to be negligible in the measured range of ionic strength; experiments carried out at monovalent ion concentration similar to physiological (50 and 100 mM KCl) were repeated three times and the control experiment at 2 M KCl was performed twice. By plotting the difference in fluorescence intensities between the blank and CPEB3-containing samples against the ratio between [Mg\(^{2+}\)] and [CPEB3], a plateau is reached when the equivalent of Mg\(^{2+}\) binding to the RNA has been added. Fluorescence spectra did not change with increased exposure to photoexcitation or longer sample mixing, indicating that photobleaching of HQS can be neglected. The equilibration of the sample was fast and longer waiting times between Mg\(^{2+}\) additions did not result in a change of fluorescence.

Mg\(^{2+}\) counting with atomic emission spectroscopy

Measurements were performed as previously described in the literature (Bina-Stein and Stein 1976; Leipply et al. 2009). Tube-O-DIALYZER Micro filters with a 1-kDa molecular weight cut-off were used to dialyze the ribozyme samples overloaded with Mg\(^{2+}\)-containing buffer (20 mM MOPS buffer, pH 7, 50 mM, 100 mM, or 2 M KCl and 0.5–100 mM Mg\(^{2+}\), 24 µM CPEB3 ribozyme, annealed by a 2-min heating at 90°C and rapid cooling on ice; the total volume of the dialyzed sample was 100 µL) against 5 mL of the same buffer without Mg\(^{2+}\) ions for 4–10 h at 298 K. The concentration of Mg\(^{2+}\) in the dialysate was measured with AAS on a Varian SpectraAA 110 spectrometer, and the amount of Mg\(^{2+}\) ions bound to the ribozyme was calculated as the ratio between the difference of the amount of Mg\(^{2+}\) added and present in the dialysate and the amount of CPEB3.

SUPPLEMENTAL MATERIAL

Supplemental material is available for this article.

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