Ribozyme reactions of viroids

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1. Introduction

Viroids replicate through a DNA-independent rolling-circle mechanism involving the synthesis of multimeric strands of both polarities and their subsequent cleavage into monomeric fragments, which are then circularized to produce the progeny. Regarding whether or not the minus multimeric strands are cleaved and ligated to unit-length circular strands, which are then used as templates for the second half of the cycle, the viroid RNA is considered to replicate by either a symmetric or asymmetric mode (see chapter on replication in Section 3). While processing of the multimeric plus RNA intermediates is generally believed to require a host ribonuclease in members of family Pospiviroidae (formerly known as group B viroids), this step is autocatalytic and mediated by hammerhead ribozymes in members of family Avsunviroidae (formerly known as group A viroids) (reviewed in Symons, 1989; Flores et al., 1998; see chapter on classification in Section 3). However, the possibility has been raised that the processing step is RNA-catalyzed in all cases (reviewed in Symons, 1997). The three viroid species of family Avsunviroidae known to date, avocado sunblotch viroid, ASBVd (Symons, 1981; Hutchins et al., 1986), peach latent mosaic viroid, PLMVd (Hernández and Flores, 1992) and chrysanthemum chlorotic mottle viroid, CChMVd (Navarro and Flores, 1997), can adopt hammerhead structures on their plus and minus polarity strands and, as a consequence, they are presumed to replicate according to the symmetric rolling-circle mechanism. The hammerhead structures appear as the only "homologous molecular character" shared by these viroids. Briefly, the hammerhead structure is a small RNA motif consisting of three sequence non-specific helices bordering a catalytic core of 11 conserved residues which form a complex array of non-canonical interactions (Prody et al., 1986; Hutchins et al., 1986; Forster and Symons, 1987; Pley et al., 1994) (Fig. 1). The adoption of this structure in the presence of a divalent cation, usually magnesium, results in the self-cleavage of the RNA chain at a specific phosphodiester bond creating 2',3'-cyclic phosphate and 5'-hydroxyl termini.

Biochemical knowledge in respect to both detailed structural features and molecular mechanism of the hammerhead structures has been reviewed recently (Flores *et al.*, 2000; Stage-Zimmermann and Uhlenbeck, 1998) and, therefore, will not be the main focus of this chapter that has for primarily aim to present the hammerhead structure as an essential molecular feature of the *Avsunviroidae* members and, particularly, of their replication cycle. We will also consider the potential of this self-cleaving motif to act in *trans* targeting cellular RNA and, more specifically, to contribute to viroid pathogenesis.

2. Hammerhead structures of viroids: molecular architecture

Fig. 2 shows the six hammerhead structures described so far in viroids. In ASBVd, the sequences involved in both polarity hammerhead structures are found in the upper and lower strands of the central domain of the quasi-rod-like secondary structure proposed for this viroid, with the remaining nucleotides of the genome, referred here as 'extracatalytic' RNA sequences, flanking the central domain. Therefore, the sequences forming the catalytic core are not contiguous but segregated in two subdomains. In contrast, the sequences involved in the hammerhead structures of PLMVd and CChMVd are contiguous and located in an arm of their proposed branched conformation, with the 'extracatalytic' RNA sequences constituting the rest of the genomes.

There are two classes of viroid hammerhead structures according to their morphology. The monomeric strands of PLMVd and CChMVd can adopt stable hammerhead structures with helices I and II of five-six base pairs closed by short loops 1 and 2 (the CChMVd minus hammerhead structure is an exception in having an unusually long imperfect helix II), and helices

III of six-eight base-pairs (Fig. 2A and B). Conversely, the hammerhead structures that can form the monomeric ASBVd RNAs are thermodynamically unstable, particularly in the plus polarity strand with a stem III of only two base pairs closed by a loop 3 of three residues (Fig. 2C). This very different architecture of the viroid hammerhead structures has deep implications for their *in vitro* and *in vivo* self-cleavage efficiency (see below).

Inspection of natural hammerhead structures shows that they are characterized by a central core with a cluster of strictly conserved nucleotide residues flanked by three double-helix regions (I, II and III) with loose sequence conservation except positions 15.2 and 16.2, which in most cases form a C-G pair, and positions 10.1 and 11.1, which in most cases form a G-C pair (Fig. 2). Some viroid hammerhead structures present unusual features. For example, a transition U to C affecting the conserved U4 in the plus hammerhead structure has been observed in a sequence variant of PLMVd. On the other hand, the common C17 preceding the minus self-cleavage site is A in a sequence variant of ASBVd, and the common pyrimidine residue at position 7 is substituted by an A residue in the minus hammerhead structure of another ASBVd variant. An extra A between A9 and G10.1 of the plus hammerhead structure of CChMVd has been also reported (Fig. 2C). This extra residue, which is compatible with extensive *in vitro* self-cleavage, could either induce a rearrangement of the junction between helix II and three adjacent noncanonical interactions of the central core, or be accommodated as a bulging residue. These and other sequence variations in the hammerhead structures retrieved in the nature from different selfcleaving RNAs have been compilated recently (see Flores et al., 2000). The conservation of the sequences forming the hammerhead structures in the Avsunviroidae members, as well as in most other hammerhead structures known so far, extend beyond the strict requirements for selfcleavage, suggesting that additional selective pressures may act on these sequences. However, the identity of any other selective pressure remains unidentified.

3. Cis-acting hammerhead structures of viroids: in vitro and in vivo function

The similarities found between the plus and minus hammerhead sequences and their genomic organization within each viroid most likely have physical and functional consequences. The PLMVd and CChMVd sequences spanning the two hammerhead domains are almost complementary and can fold into structures with long double-stranded regions; this is the typical arrangement found in the most stable secondary structures predicted for the two viroids (Fig. 2A and B). These stable arrangements, formed by the superposition of the hammerhead sequences of both polarities, have the potential to prevent the adoption of the active hammerhead foldings, which are alternative structures of higher energy (Hernández and Flores, 1992; Beaudry et al., 1995; Navarro and Flores, 1997). More importantly, self-cleavage inhibition permits the accumulation of certain levels of the viroid monomeric circular forms which are the templates of the rolling-circle mechanism of replication. In addition, the compact non-self-cleaving structures may contribute positively to the extra- (e.g. during transmission) and intracellular stability of these RNA species. The peculiar organization of the hammerhead sequences may be also informative of mechanistic requirements. For example, these RNAs may need to have similar hammerheads in order to perform *in vivo* self-cleavage to the essentially the same extent in both strands, as it appears to be the case in PLMVd (Bussière et al., 1999). Interactions with cellular component (e.g. proteins) enhancing self-cleavage may have promote conservation of similar hammerheads. Therefore, a complex synergy between the stability of the viroid RNA as a whole, and the mechanisms of self-cleavage regulation, has probably contributed to the emergence of the superposed hammerhead sequences.

As already indicated, the ability of viroid RNAs that possess autocatalytic sequences to selfcleave depends on their adoption of a conformation different from the most stable structure (Fig. 2). Self-cleavage of viroid strands occurs at either single or double hammerhead structures depending on whether or not the sequences can form stable helices surrounding the catalytic core. Whereas the six hammerhead structures of viroids have stable helices I and II, this is not the case for helix III. Both polarity hammerhead structures of PLMVd have that stable helices III and selfcleave *in vitro*, and likely *in vivo*, through single hammerhead structures (Hernández and Flores, 1992; Beaudry et al., 1995) (Fig. 2A). This is also probably the case with the two hammerhead structures of CChMVd, which also have stable helices III (Navarro and Flores, 1997), although the extended helix II of the minus hammerhead structure might facilitate the adoption of alternative foldings inactive for self-cleavage (Fig. 2B). In contrast, the single hammerhead structures of ASBVd have unstable helices IIII closed by short loops, and their self-cleavage is assumed to occur via double hammerhead structures involving longer-than-unit RNA that allow stabilization of the catalytic core (Forster et al., 1988) (Fig. 2C). ASBVd plus strands self-cleave through a double hammerhead structure during *in vitro* transcription and after gel purification, whereas ASBVd minus strands self-cleave via a double hammerhead structure during in vitro transcription, but mostly via a single hammerheads structure after gel purification (Davies et al., 1991). This is most probably the consequence of the different stability of helix III in both hammerhead structures.

Direct enzymatic sequencing and primer extension experiments have shown that *in vitro* self-cleavage of ASBVd, PLMVd and CChMVd occurs at the positions predicted by the hammerhead structures (Hutchins *et al.*, 1986; Hernández and Flores, 1992; Navarro and Flores, 1997). The efficiency of the *in vitro* self-cleavage reaction can be high; for example around 50-60% of PLMVd strands self-cleave under standard conditions (Hernández and Flores, 1992; Beaudry *et al.*, 1995). However, this efficiency is strongly increased (>95%) when the same RNAs are transcribed under conditions of slow polymerase activity, which favors the adoption of

the active hammerhead structures catalyzing the self-cleavage reactions (Bussière, 1999). The self-cleavage efficiency is also strongly dependent on divalent ions such as Mg^{2+} .

There is also solid evidence supporting the involvement of hammerhead structures in the *in vivo* processing of viroid RNAs with these catalytic domains. For ASBVd (Daròs *et al.*, 1994; Navarro and Flores, 2000), CChMVd (Navarro and Flores, 1997), and PLMVd (Hernández, unpublished data), linear RNAs of one or both polarities with 5' termini identical to those generated in the corresponding *in vitro* self-cleavage reactions have been isolated from infected tissues. Moreover, the frequent occurrence in sequence variants of PLMVd (Hernández and Flores, 1992; Beaudry *et al.*, 1995; Ambrós *et al.*, 1998) and CChMVd (De la Peña *et al.*, 1999) of compensatory mutations or covariations that preserve the stability of the hammerhead structures, further support their *in vivo* role, as also does the correlation existing between the infectivity of different PLMVd and CChMVd variants and the extent of their self-cleavage during *in vitro* transcription (Ambrós *et al.*, 1998; De la Peña *et al.*, 1999).

In vivo, self-cleavage of viroid strands should be under regulation, with two different mechanisms appearing to operate for this purpose. In the case of PLMVd and CChMVd, their most stable secondary structures are transiently lost during transcription with the concurrent adoption of the active single hammerhead structures that promote self-cleavage before synthesis is completed and the most stable secondary structures are reformed. In ASBVd self-cleavage of monomeric strands is restricted because the single hammerhead structures are unstable whereas the multimeric replicative intermediates can adopt stable double hammerhead structures and self-cleave to their unit-length strands. Therefore, in both situations the hammerhead ribozymes are active only during replication. Self-cleavage of PLMVd RNAs *in vivo* appears almost optimal reaching near total processing of the multimeric strands into their corresponding monomeric units (Bussière *et al.*, 1999). For the reasons stated above, this high efficiency may be the result of slow

progession of the host RNA polymerase during replication. The situation seems similar in the case of CChMVd, for which the predominant RNAs accumulating in infected cells are also the monomeric linear strands of both polarities (Navarro and Flores, 1997). In contrast, the most abundant ASBVd RNA in infected tissue is the plus circular monomer, a clear indication of the low efficiency of the corresponding single hammerhead structure, although decreasing levels of multimeric strands up to octamers in size have been also detected (Bruening *et al.*, 1982).

4. Viroid hammerhead structures: potential for *trans*-acting function

In their natural context, the hammerhead structures of viroids operate in *cis* mediating the self-cleavage of the RNAs in which they are contained. However, active hammerhead structures can also be formed by annealing two different RNA fragments in trans, such that one RNA fragment acts as the ribozyme and the other as the substrate (see Fig. 3A). If the complementary regions between the two RNAs are short enough, the cleavage products will dissociate from the ribozyme, thus permitting the binding of new substrate molecules. Via successive rounds of binding and cleavage a single ribozyme molecule can therefore cleave many substrate molecules, thereby establishing a classic enzyme/substrate relationship (Uhlenbeck, 1987). Furthermore, by changing the complementary sequences between the ribozyme and its substrate, it is possible to create a ribozyme with a new substrate specificity. A wide variety of RNAs can be targeted for cleavage by such engineered ribozymes (Fig. 3B). Because of their ability to interact directly with RNA, ribozymes, particularly those of the hammerhead class, are currently being developed as potential therapeutic agents for a wide range of applications based on the specific cleavage of different RNAs of biological relevance including viroids themselves (see the preceding chapter). In the coming years, altered forms of these versatile molecules will surely emerge as a new class of drugs.

Apart from these applications, a detailed description of which fall outside the scope of this chapter, we will consider the possibility that viroid pathogenesis of members of the family Avsunviroidae could result from trans cleavage of host RNAs accidentally recognized by the hammerhead ribozymes (Symons, 1989). No supporting evidence for such a mechanism has been reported yet. The following discussion is based on experiments performed with PLMVd as a model viroid in an attempt to put this intriguing hypothesis to test (XXX and Perrault, unpublished data). Minimal artificial hammerhead ribozymes are prefolded into a quasi catalytically active structure and following the substrate binding, which involves the formation of helixes at both sides of the cleavage site, the break of the scissile bond occurs. Natural hammerhead structures are integral features of viroid RNAs. In the most stable secondary structures of these RNAs, the hammerhead catalytic core is not formed because this is not the most stable structure. For example, active PLMVd hammerhead structures of both polarities are adopted either during the *in vitro* transcription or by a prior heat denaturation coupled to a snapcooling treatment, that favor these active structures over others more stable but lacking catalytic activity. Therefore, the knowledge acquired from studies with minimal hammerhead ribozymes can not be simply extended to situations in which this catalytic motif is included in full-lenght viroid RNAs.

In order to compare the cleavage efficiency of a hammerhead catalytic sequence as a model molecule or as part of a viroid, a series of experiments were performed in which four PLMVd-derived transcripts, acting as the ribozyme (Rz), were tested for their ability to catalyze the cleavage of a short substrate (S). As expected, no cleavage products were detected when the Rz was a PLMVd 250-nt transcript lacking the hammerhead sequences, whereas most of the S (>85%) was cleaved by a Rz with the sequences corresponding to the minimal plus hammerhead structure. This efficient cleavage probably results from the absence of extra sequence interfering

with the adoption of the catalytically active folding. When the Rz was composed by a full-length PLMVd RNA circularized *in vitro* to have either a 3',5'- or a 2',5'-phosphosdiester bond at the self-cleavage site, only a trace amounts of product (<1%) were detected in both cases. Finally, the Rz corresponding to the full-length PLMVd linear RNA that accumulates predominantly in infected peach cells was able to cleave the S, although with an extremely low efficiency (<5%). Further experiments showed that cleavage in this latter case was most likely performed by the hammerhead motifs released during the preliminary denaturation-renaturation treatment, because no cleavage was detected when this treatment was omitted. Altogether, these experiments indicate that the possibility that a viroid may function as a *trans*-acting ribozyme triggering a pathogenic cascade is unlikely. The main impediment comes from the catalytic sequences being embedded in very stable secondary structures, reducing considerably their ability to hybridize in *trans* with other RNAs.

As already mentioned, minimal hammerhed structures can catalyze efficiently *trans* cleavage of small substrates. The possibility that such an active ribozyme could be released by specific hydrolysis during the viroid life cycle seems remote because there is no indication supporting the idea that if the viroid is attacked by host RNases the hammerhead sequences would be protected against degradation. Alternatively, minimal hammerhead structures could be transiently formed during viroid replication. Such a possibility has been evaluated by testing the cleavage of a model substrate during the *in vitro* transcription of a PLMVd dimeric RNA. In the absence of either the DNA template or the T7 RNA polymerase, no cleavage was observed. However, a small fraction of the substrate was cleaved during viroid replication. Additional experiments performed in the presence of protein A1, which has been previously shown to enhance the *trans* cleavage activity of hammerhead ribozymes (***EMBO J. paper from

Herschlag's group?), have allowed to increase slightly the extent of cleavage. Together, these results do not exclude that *in vivo* a host protein could open the compact PLMVd structure and favor the recognition and cleavage of a host RNA possessing a sequence targeted by the viroid hammerhead ribozymes. To this aim, at least two other requisites need to be met. First, the sequences forming the binding region of the ribozyme (i.e. one strand of helices I and II), should be expected to be conserved between sequence variants, but sequencing of numerous PLMVd natural variants (Hernández and Flores, 1992, Ambrós *et al.*, 1998; Pelchat *et al.*, 2000) has shown that this is not the case. And second, a natural substrate(s) for the viroid hammerhead ribozymes should exist. PLMVd replication intermediates have been predominantly detected in chloroplasts but attemps to retrieve with well-established procedures chloroplastic sequence(s) which could be potentially cleaved by one or the other PLMVd hammerhead structures have been unsuccessful. Therefore, the hypothesis that PLMVd, and by extension other hammerhead viroids, may exert their pathogenic effect by hammerhead-mediated cleavage of a cellular RNA appears very unlikely.

The previous discussion points out that in their natural context the hammerhead ribozymes of viroids are not good candidates for targeting cellular RNA of biological relevance. On the other hand, viroid RNAs could be in principle appropriate vehicles for delivery of nucleic acid-based drugs into chloroplasts ,because they are endowed with the information required to be host-and organel-specific and they can be genetically manipulated. It is clear that there is still a long way to go before such a development could be reached. Since the hammerhead motif has to fold into an active conformation to express its catalytic *trans* cleaving potential, one option would be to engineer viroids with pre-folded hammerhead structures. However, whether these modified ribozymes are compatible with normal viroid functions, particularly replication, remain to be known.

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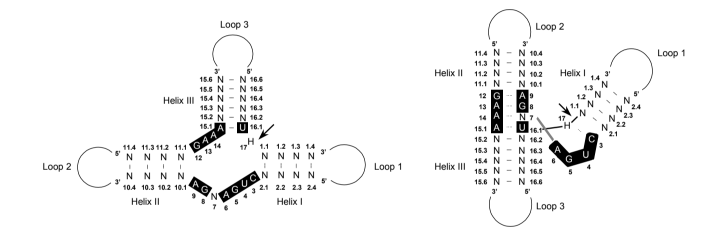
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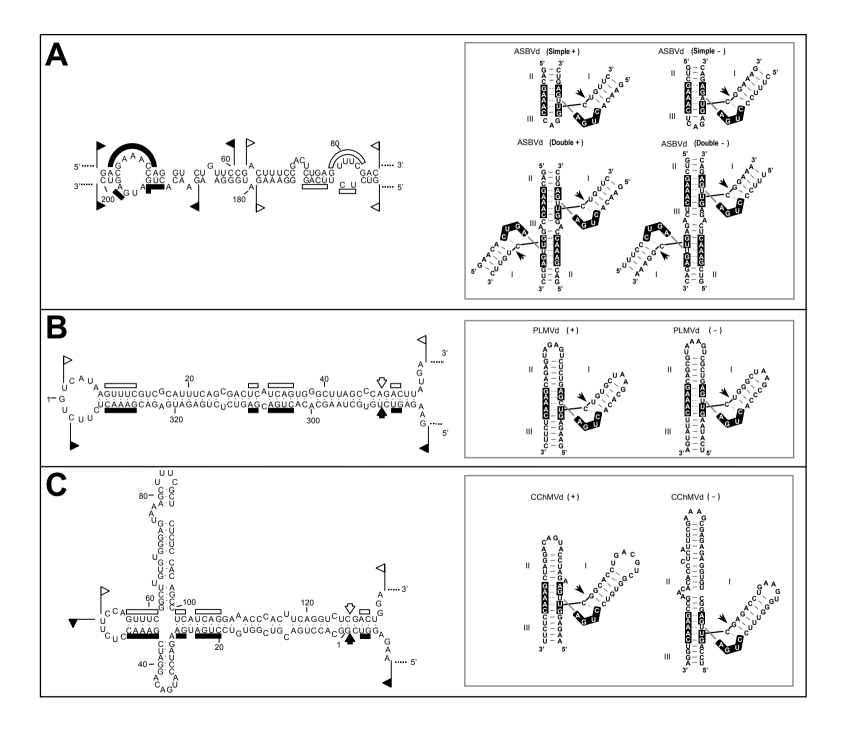
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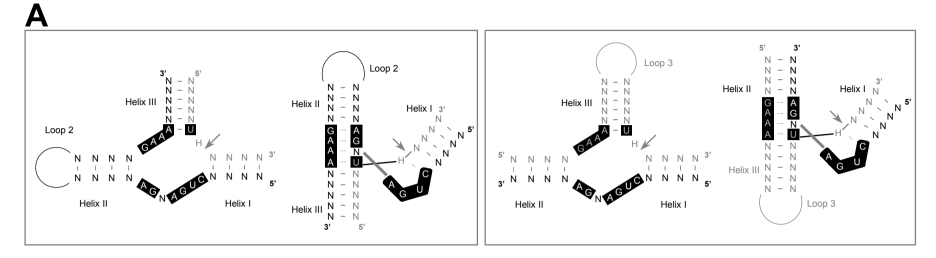
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Fig. 1. Consensus hammerhead structure derived from 23 natural hammerhead structures, schematically represented as originally proposed with its numbering system and names of helices and loops (Hertel *et al.*, 1992) (right), and according to X-ray crystallography data (Pley *et al.*, 1994) (left) Letters on a dark background refer to absolutely or highly conserved residues in all natural hammerhead structures. Arrows indicate self-cleavage sites. Watson-Crick base pairs and non-canonical interactions are denoted with continuous and broken lines, respectively.

Fig. 2. Conservation of the hammerhead structures and sequences. **A** Nucleotide sequence and secondary structure of the hammerhead self-catalytic motif. The three conserved domains are outshadowed. Unconserved positions known to affect the catalytic rate are boxed. N: A, G, C or U; I: inosine; H: A, C or U. The arrow points the cleavage site. **B** Organization and sequences of the hammerhead found on both the plus (+) and minus (-) polarities RNAs. The size of stem-loops and the positions of the loops (underlined nucleotides) are shown. - are gaps while insertion size are between parenthesis. Boxed nucleotides in stems are not involved in Watson-Crick interactions. Arrows represent the position of extracatalytic sequence. When several variants were known a consensus sequence was used (R= A or G; Y= U or C; W= A or U; K= G or U; M= C or A). The (A) is presents only in ASBVd. The written consensus of (+) hammerhead stem-loop II represents the sequence found in at least 11 out of 13 considered RNAs. Sequence where retrieved from the viroid and viroid-like database (http://www.callisto.si.usherb.ca/~jpperra; Pelchat *et al.*, 2000).







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