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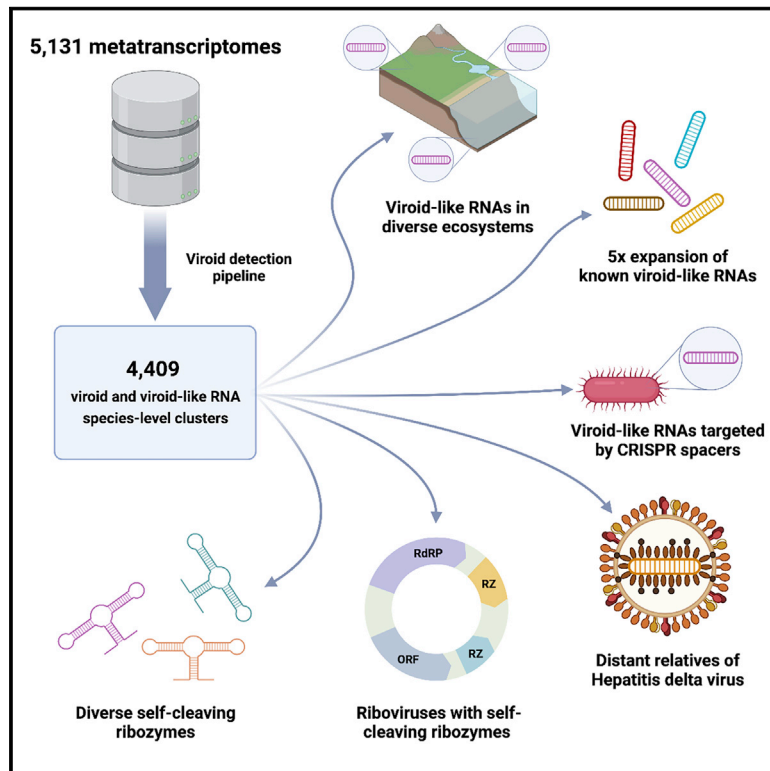
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Peer reviewed

Mining metatranscriptomes reveals a vast world of viroid-like circular RNAs

Graphical abstract



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In brief

A large-scale survey of covalently closed circular RNA across ecosystems reveals that viroids infect a wide range of host species, extending beyond plants, and identifies additional types of ribozyme activity as well as functional features in these molecules.

Highlights

- Metatranscriptome search yields a 5-fold increase in viroid-like circular RNA diversity
- Distant relatives of hepatitis delta virus and diverse viroid-like viruses are discovered
- Viroid-like circular RNAs are targeted by endogenous CRISPR systems
- Diverse ribozymes and ribozyme combinations identified in viroid-like RNAs



Resource

Mining metatranscriptomes reveals a vast world of viroid-like circular RNAs

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SUMMARY

Viroids and viroid-like covalently closed circular (ccc) RNAs are minimal replicators that typically encode no proteins and hijack cellular enzymes for replication. The extent and diversity of viroid-like agents are poorly understood. We developed a computational pipeline to identify viroid-like cccRNAs and applied it to 5,131 metatranscriptomes and 1,344 plant transcriptomes. The search yielded 11,378 viroid-like cccRNAs spanning 4,409 species-level clusters, a 5-fold increase compared to the previously identified viroid-like elements. Within this diverse collection, we discovered numerous putative viroids, satellite RNAs, retrozymes, and ribozyme-like viruses. Diverse ribozyme combinations and unusual ribozymes within the cccRNAs were identified. Self-cleaving ribozymes were identified in ambiviruses, some mito-like viruses and capsid-encoding satellite virus-like cccRNAs. The broad presence of viroid-like cccRNAs in diverse transcriptomes and ecosystems implies that their host range is far broader than currently known, and matches to CRISPR spacers suggest that some cccRNAs replicate in prokaryotes.

INTRODUCTION

Viroids, which cause several economically important diseases in agricultural plants, are the smallest and simplest among the known infectious agents.^{1–3} Viroids are small, covalently closed circular (ccc) RNA molecules of 220–450 nucleotides that encode no proteins and consist largely of RNA structures that are required for replication or viroid-host interaction. In contrast to viruses, which hijack the host translation system to produce proteins encoded in virus genes, viroids take advantage of the host transcriptional machinery. Specifically, viroids hijack the host plant's DNA-dependent RNA polymerase II to transcribe their RNA and thus catalyze viroid replication.^{4–6} Viroids utilize the rolling circle replication (RCR) mechanism, producing multimeric intermediates that are cleaved into genome-size monomers by ribozymes that are present in both polarities of viroid RNAs or by recruited host RNases.^{7,8} The resulting linear monomers are then ligated by a host DNA ligase to form the mature cccRNA.^{9,10}

Since the discovery of viroids in 1971,¹¹ about 50 viroid species were identified in plants and classified into two families, Avsunviroidae and Pospiviroidae. Members of the Avsunviroidae family use viroid-encoded autocatalytic hammerhead (HHR) ri-

bozymes to process replication intermediates into unit length viroid genomes.^{12,13} Members of the family Pospiviroidae lack ribozymes and instead rely on conserved sequence motifs that serve as recognition and cleavage sites for host RNase III.¹⁴ The members of the two viroid families adopt distinct RNA structures: a branched RNA conformation is predominant in avsunviroids,¹⁵ in contrast to the typically rod-shaped conformation of pospiviroids.¹⁶

In addition to viroids, several other groups of infectious agents also possess genomes consisting of cccRNA.¹⁷ Many plant viruses support the replication of small (about 300 nt) circular satellite RNAs (satRNAs) that closely resemble viroids¹⁸ and also replicate via RCR.¹⁹ The satRNAs differ from viroids in that they are replicated by the RNA-dependent RNA polymerase (RdRP) of the helper virus and are encapsidated in that virus's capsid.^{20,21} Thus, satRNAs are effectively encapsidated viroids. Unlike viroids, satRNAs encode both HHRs and hairpin ribozymes (HPRs).²²

Another viroid-like agent is the retroviroid, carnation small viroid-like RNA (CarSV), which, unlike viroids, does not appear to transmit horizontally among plants.²³ CarSV, the only currently known retroviroid, is a cccRNA that is similar to viroids in size and contains HHRs in both strands. However, in contrast to the



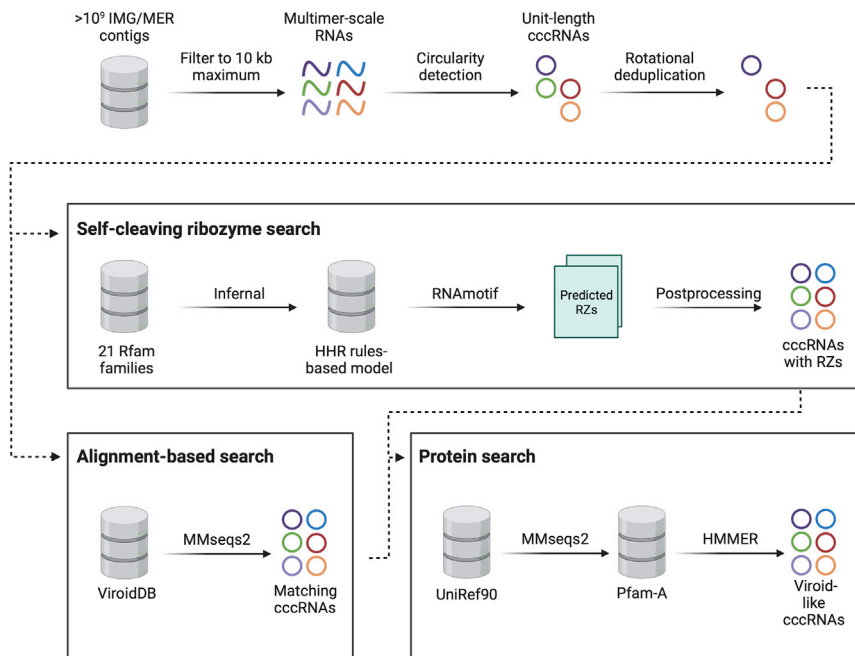


Figure 1. Viroid-like cccRNA detection pipeline

row host range of viroids, which so far have been reported only in plants, is poorly compatible with this evolutionary scenario. Instead, given the similarities between retrozymes and avsunviroids, it has been suggested that avsunviroids descended from retrozymes.^{27,39}

Given the ultimate structural simplicity of viroids and related cccRNAs and the universality of the DNA-dependent RNA polymerases involved in their replication across life forms, the current narrow spread and limited diversity of parasitic cccRNAs appear puzzling. Furthermore, this apparent paucity of viroid-like agents is in stark contrast to the burgeoning diversity of RNA viruses, many thousands of which include numerous, distinct groups have been discovered by metatranscriptome analyses.^{40–43} At present,

viroids, an extrachromosomal DNA copy of CarSV has been discovered and shown to integrate into the plant genome with the help of a pararetrovirus.^{24,25}

A recently discovered group of cccRNA agents are retrozymes, retrotransposons that propagate via circular RNA intermediates of about 170–400 nucleotides. The retrozymes are viroid-like in that they do not encode any proteins but contain self-cleaving HHRs.^{26,27} However, unlike viroids, the retrozymes are neither infectious nor autonomous but rather hijack the replication machinery of autonomous retrotransposons. Resembling satRNAs and avsunviroids, retrozyme cccRNAs adopt a branched conformation.

A conspicuous group of viroid-like agents is the viral realm *Ribozyviria*²⁸ that includes deltaviruses, such as hepatitis delta virus (HDV), an important human pathogen. Similarly to pospiviroids, ribozyviruses possess rod-shaped cccRNA genomes that replicate via the RCR mechanism and encode distinct ribozymes, unrelated to those of viroids, that autocatalytically process multimeric replication intermediates.^{29–31} Ribozyviruses have substantially larger genomes than other viroid-like agents (about 1.7 kb) and encode their own nucleocapsid protein. The reproduction of ribozyviruses relies on a helper virus (hepatitis B virus in the case of HDV), which provides the envelope protein for ribozyviruses. For years, HDV remained the only known deltavirus. Recently, however, viruses more distantly related to HDV have been discovered in various vertebrates and invertebrates,^{32–36} suggesting a considerable uncharacterized diversity of ribozyviruses.

Viroids and viroid-like cccRNAs are minimal replicators, or ultimate parasites, that lack genes and effectively consist only of RNA structures required for replication. This extreme simplicity of viroids triggered speculation on their direct descent from primordial RNA replicators.^{37,38} However, the apparent nar-

row host range of viroids, which so far have been reported only in plants, is poorly compatible with this evolutionary scenario. Instead, given the similarities between retrozymes and avsunviroids, it has been suggested that avsunviroids descended from retrozymes.^{27,39}

Given the ultimate structural simplicity of viroids and related cccRNAs and the universality of the DNA-dependent RNA polymerases involved in their replication across life forms, the current narrow spread and limited diversity of parasitic cccRNAs appear puzzling. Furthermore, this apparent paucity of viroid-like agents is in stark contrast to the burgeoning diversity of RNA viruses, many thousands of which include numerous, distinct groups have been discovered by metatranscriptome analyses.^{40–43} At present,

there are at least three orders of magnitude more known RNA viruses than there are viroids and viroid-like cccRNAs.

We were interested in investigating the global diversity of viroids and viroid-like agents. To this end, we performed an exhaustive search for cccRNAs in a collection of 5,131 diverse metatranscriptomes that have been recently employed for massive RNA virus discovery⁴³ and additionally searched 1,341 plant transcriptomes.⁴⁴ This search yielded more than 10,000 viroid-like cccRNAs that represent an about 5-fold increase of the known diversity of viroid-like agents. Further analysis of these cccRNAs led to the identification of numerous putative viroids, satRNAs, retrozymes, and ribozy-like viruses.

RESULTS

Computational approach for the discovery of viroid-like cccRNAs

We developed an integrated, scalable computational pipeline for the discovery and analysis of viroids and viroid-like cccRNAs directly from assembled transcriptomes and metatranscriptomes (Figure 1). The pipeline starts with the reference-free and *de novo* identification of cccRNAs or RCR intermediates, capitalizing on the fact that assemblies of both complete circular monomers and multimeric linear intermediates contain head-to-tail repeats.⁴⁵ The identified sequences are then cleaved *in silico* to unit length and deduplicated, taking circularity into account. Starting from the set of detected cccRNAs, the pipeline performs both alignment-free and alignment-based searches. The primary approach for the identification of viroid-like agents among the cccRNAs is the prediction of self-cleaving ribozymes using RNA sequence and secondary structure covariance models.⁴⁶ Assuming that the diversity of ribozymes in viroid-like RNAs could be greater than so far uncovered, we curated a database

of known self-cleaving ribozyme models from Rfam,⁴⁷ of which only a minority were detected in viroids and viroid-like RNAs. We supplemented this model database with the pospiviroid RY motif⁴⁸ to enable detection of potential pospiviroids, which lack ribozymes. The pipeline also performs direct sequence similarity searches against reference databases such as ViroidDB.⁴⁹

Ribozyme-containing cccRNA sequences were classified as symmetric or asymmetric depending on whether they contained predicted ribozymes in both or only one RNA polarity, respectively, reflecting the RCR mode these cccRNAs are likely to undergo. However, it cannot be ruled out that some apparently asymmetric cccRNAs actually contain a second ribozyme distinct from the currently known ones.

We validated this method by demonstrating its ability to recover known viroid-like RNAs in transcriptomes and metatranscriptomes (Table S1). For the transcriptomic validation, we processed and searched the 1,000 Plant transcriptome (1KP) dataset.⁴⁴ We chose this dataset due to the known presence of all types of viroid-like cccRNAs except ribozyviruses in plant transcriptomes. Assembling the raw reads of 1,344 transcriptomes resulted in 103,139,086 contigs, of which 163,970 were predicted to be circular. Of these putative cccRNAs, 42 were identified as viroid-like via ribozyme search (15 sequences), sequence search against ViroidDB (33 sequences), or both (6 sequences).

To verify the efficacy of the detection method, we performed a direct search of all contigs against ViroidDB and identified 12 contigs that matched a known viroid sequence with at least 50% target coverage. The detection pipeline found four of these potentially complete viroid contigs. Of the rejected contigs, four were much larger than typical viroids (>1,000 nt) and contained major ambiguous regions. The other four were low-coverage fragments that could not be verified as circular being smaller than unit length. Iresine viroid 1, Citrus exocortis viroid, and a *Coleus blumei* viroid (CbVd) were successfully retrieved. While the former two were nearly identical to the corresponding reference sequences, the CbVd-like sequence was not. At 350 nt, this sequence differed in length from all known coleoviruses and in the terminal conserved region, which was identical to that of Dahlia latent viroid, suggesting an origin of this viroid by recombination, as reported for other CbVd species.^{50,51} At 85% identity, this CbVd-like sequence falls below the species membership threshold for coleoviruses.⁵² The relatively low number of viroids identified in plant transcriptomes is likely due to selection of healthy plants for RNA isolation and transcriptome analysis.

A 5-fold expansion of the known diversity of viroid-like cccRNAs

After testing the pipeline on plant transcriptomes, we applied it to a set of 5,131 diverse metatranscriptomes totaling 1.5 billion metatranscriptomic contigs (708 Gbp) after size filtration. We identified 10,183,455 putative cccRNAs with a median contig length of 269 nt. After removing overlapping regions and eliminating rotationally identical sequences, the median length of the 8,748,001 resultant monomers was 165 nt. Of these, 2,791,251 were within the known size range of viroids (200–400 nt), including 11,378 we classified as viroid-like because they contained a confidently predicted self-cleaving ribozyme

in at least one RNA polarity. No metatranscriptomic cccRNAs matched the pospiviroid RY motif. Among the viroid-like cccRNAs, 10,181 were detected by alignment-free methods, that is, showed no detectable sequence similarity to known viroids. The remaining 1,197 sequences shared significant nucleotide sequence similarity with known viroid-like RNAs spanning the entire gamut from viroids to satRNAs to retrozymes (Table S2). 907 sequences were identified as viroid-like by both the ribozyme detection and alignment-based search approaches. Among the 10,181 ribozyme-containing viroid-like cccRNAs unrelated to known viroids, 3,434 were symmetric, that is, contained predicted ribozymes in both polarities. Of the 5,131 metatranscriptomes searched, 1,841 contained at least one viroid-like cccRNA.

Of the sequences aligning to viroid-like agents, the majority only contained short (<40 nt) alignable regions, generally localized to the ribozyme motifs. However, 33 sequences yielded long (>100 nt) alignments. These cccRNAs aligned to Tobacco ringspot virus satRNA (satTRSV), Lucerne transient streak virus satRNA (satLTSV), citrus dwarfing viroid (CDVd), and two retrozymes. For satTRSV and satLTSV, the range of identity between the recovered cccRNAs and the reference sequence ranged 80%–98% and 81%–99%, respectively. The match to CDVd was 80% identical to the nearest reference sequence. In all cases, the cccRNAs were similar in length and structure to the reference sequences.

We clustered the viroid-like cccRNAs identified here to estimate the increase in diversity compared to previously known viroid-like RNAs (Figure 2). Aligning cccRNAs poses a challenge due to the variation in the rotation of the sequences. Two identical cccRNAs could appear to have only half the bases aligning if rotated completely out of phase. Therefore, we took special care to compensate for the circularity of the sequences during the postprocessing of the pairwise nucleotide search results (see STAR Methods). To validate our clustering method, we tested it on ViroidDB. Previously, we identified 458 clusters at the average nucleotide identity (ANI) 90% level in ViroidDB using a method that was not circular-aware.⁴⁹ Using the improved method, we identified 50 clusters in ViroidDB, generally corresponding to individual species.

In the combined metatranscriptomic, transcriptomic, and reference datasets, we identified 4,823 ANI90 clusters of which 4,121 did not include any sequences from the reference datasets and thus were considered distinct, a 5.9-fold increase in viroid-like RNA diversity. Of the remaining 702 clusters containing at least one known sequence, 288 (41%) were expanded by at least one distinct sequence. Notably, 39 distinct clusters were represented in plant transcriptomes, of which 8 were symmetric.

The relative abundance of HHR types in the cccRNAs varied significantly from what would be expected given the sequence and species count. Within Rfam, HHR1 swamps HHR3 by two orders of magnitude by sequence count (190,679 versus 538 sequences). However, among the cccRNA cluster representatives, HHR3 was found in 94% and was two orders of magnitude more common than HHR1 (1,952 versus 32). Given the dominance of HHR3 in known viroids,^{8,18} this overabundance of HHR3 is suggestive of the presence of numerous viroid-like cccRNAs.

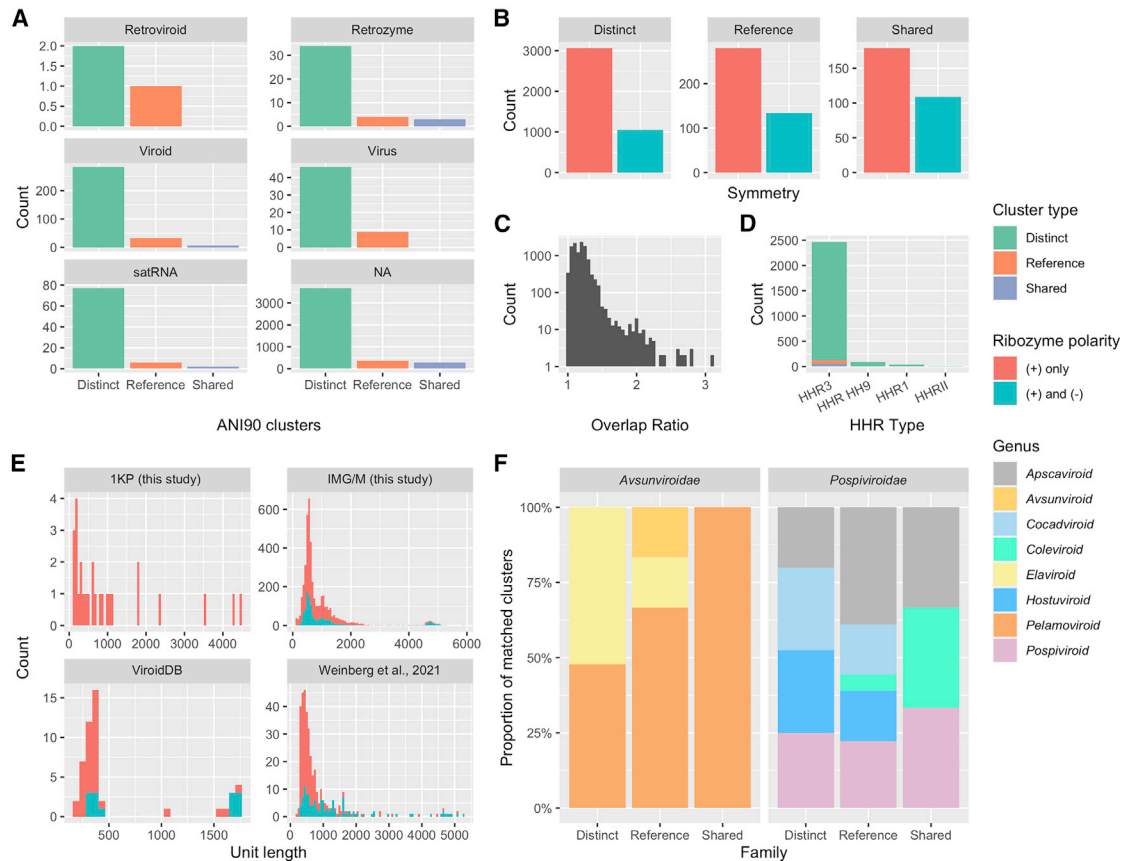


Figure 2. Viroid-like cccRNAs identified in metatranscriptomes

(A) Number of ANI90 clusters with most significant matches to given viroid-like cccRNA agent types, which are either “distinct” (derived exclusively from transcriptome and metatranscriptome analysis in this work), “reference” (no members distinct from previously identified ones), or “shared” (containing at least one of both types of sequence).

(B) Comparative distribution of inferred ribozyme architectures by cluster type.

(C) Plot of overlap ratios in cccRNAs, defined as the assembled length divided by the monomer length, from IMG and 1KP.

(D) Counts of HHR types in representative clusters.

(E) Length distributions of cluster representatives in the present analysis (transcriptomes and metatranscriptomes), ViroidDB, and a previous study.⁵³

(F) Relative abundance of clusters matching different genera within each viroid family by cluster type.

Putative distinct viroid-like cccRNAs

We briefly describe the 5 largest distinct ANI90 clusters (denoted 1 to 5, in the descending order of the cluster size) derived from the metatranscriptomic data to exemplify the type of findings obtained in this work. All these clusters included members with symmetric, matched ribozymes. The cccRNAs in four of these clusters contained matched HHR3s, whereas those in the fifth cluster contained twister-P1 ribozymes.

The largest, cluster 1, consisted of 149 sequences with a mean length of 562 nt (± 9.0 nt). During circularity detection, an average of 18% of the monomer was removed, although one member of the cluster yielded a contig with a 60% (341 nt) overlap. The cccRNAs in this cluster are predicted to adopt a rod-shaped conformation with 74% of the bases (on average) paired in both polarities (Figure 3A). Most members of this cluster ($n = 137$) contain symmetric HHR3s, whereas for the remaining 12 members, only one ribozyme was predicted, suggesting the presence of a divergent HHR3. Among the members of this clus-

ter, 38 sequences yielded a short (26–37 nt) alignment to the HHR3 of Eggplant latent viroid or Grapevine hammerhead viroid-like RNA. However, the cccRNAs comprising this cluster are substantially longer than the respective viroids (334 and 375 nt, respectively). The majority of the cluster members ($n = 133$) were found in terrestrial metatranscriptomes from 11 distinct locations, four members were identified in three distinct freshwater locations, and one member was found in a spruce rhizosphere sample.

The cccRNAs in cluster 2 (68 members) contained twister-P1 ribozymes in both polarities. All but four of these cccRNAs are 494 nt in length and form a branched structure with a mean of 70% base complementarity (Figure 3B). These cccRNAs were found in 10 locations in terrestrial ecosystems, such as soil and plant litter, with nearly half ($n = 31$) identified in switchgrass phyllosphere samples. Cluster members were repeatedly found during sampling of the switchgrass phyllosphere over the course of a year at a sample site in Michigan, USA.

Like cluster 1, cluster 3 (61 members) consisted of comparatively large (605 nt), rod-shaped cccRNA with symmetric HHR3s (Figure 3C). However, unlike the largest cluster, no member shows a detectable direct nucleotide match to any ViroidDB sequence, ribozyme, or otherwise. In the majority ($n = 40$) of the members, symmetric ribozymes were not detected, but remaining ones were symmetric, again, suggesting the presence of divergent HHRs. Between 73% and 83% of the bases in these cccRNAs were paired in the (+) polarity and between 74% and 82% of bases were paired in the (–) polarity. On average, 28% of the monomer's length was cleaved during circularity detection although one member was sequenced at 2.78 unit length (almost a complete head-to-tail trimer). The largest two members of the cluster (1,693 and 1,640 nt originally, 1,224 nt after cleavage) were not correctly monomerized by the circularity detection procedure due to mismatches in the seed sequence (see STAR Methods). Manual monomerization of these sequences showed they both were 612 nt, resulting in overlap ratios of 2.76 and 2.67, respectively. Alignment results in 99.0% and 99.5% identity between monomers within each dimer, approximately the error rate of RNA polymerase II when using an RNA template.^{54–56} Almost all members of this cluster were identified in soil samples from six locations around the world (including Colombia, Czech Republic, Germany, and USA), and two were found in creek freshwater samples.

Cluster 4 ($n = 61$) is similar to clusters 1 and 3 in that its members consist of 615 nt and are predicted to form rods containing HHR3s in both polarities (Figure 3D). The secondary structure results in a high mean self-complementarity of 75% of bases. However, these cccRNAs contain no alignable regions to the other large clusters or to any ViroidDB sequence. As with many other HHR3 rods, the majority ($n = 39$) contain only one HHR3 above the significance threshold. All but two members of the cluster were derived from eight soil locations, and the remaining ones were found in the spruce rhizosphere.

Among the largest clusters we examined, cluster 5 (55 members) is the only one that consists of cccRNAs with predicted quasi-rod-shaped structures (Figure 3E), with 64% bases paired on average. At 454 nt mean length (± 8.9 nt), these cccRNAs are the smallest among the five largest clusters. In this case, all members contained two HHR3s, but no nucleotide matches to ViroidDB were detected. Most members were found in terrestrial samples including soil ($n = 38$) and plant litter and peat ($n = 2$). As in the case of cluster 2, 14 members of this cluster were also found over the span of a year in the phyllosphere of switchgrass, and one member was found in a freshwater sample. Altogether, nine distinct locations contained members of this cluster.

In summary, analysis of these five largest clusters showed that they consisted of cccRNAs endowed with all the hallmarks of viroids including symmetric ribozymes (HHR3s, with one notable exception), an extensive branched or (quasi) rod-shaped secondary structure, and evidence of multimeric intermediates. Furthermore, the members of these clusters were independently identified in diverse samples, primarily those from soil, indicating they are widespread and are likely to be infectious agents.

Virus-like elements blurring lines between riboviruses, ribozviruses, and viroids

Among the cccRNAs containing symmetric HHR3s, we identified rod-shaped sequences up to 4,705 nt, far outside the length range of viroids and ribozviruses. We hypothesized that these cccRNAs could be previously uncharacterized ribozy-like viruses. To perform a comprehensive search for potential ribozy-like viruses, all open reading frames (ORFs) longer than 75 codons present in cccRNAs were translated, and the resulting sequences of putative proteins were clustered by amino acid sequence similarity and compared to protein sequence databases (STAR Methods; Table S3).

Almost all reliable matches were to virus proteins (Table S3). One protein cluster showed significant sequence similarity to the predicted RdRPs of a distinct group of ssRNA viruses, ambiviruses, that were recently discovered in fungal isolates and transcriptomes.^{57–59} Ambiviruses have RNA genomes of approximately 4 kb that encompass bidirectional ORFs, one of which encodes a predicted RdRP. To date, ambiviruses have not been reported to be circular.⁵⁸ In the IMG (integrated microbial genomes) metatranscriptomes, 163 ANI90 clusters (274 cccRNAs total) were found to encode ambivirus-like RdRP (E-values between $1.3e-229$ and $8.7e-04$). Notably, these clusters of cccRNAs were also predicted to contain HHR3, HPR-meta1, and CEPB3 ribozymes, including symmetric sequences with different ribozymes in the two RNA polarities. All these sequences were predicted to adopt a rod-like structure in which the two ORFs encoding, respectively, the RdRP and an uncharacterized protein are arranged along the rod in the opposite strands (Figure 4). These sequences showed varying degrees of terminal overlap, with a median trimmed repeat length of 123 nt. Three representative sequences were recovered with $>2,000$ nt overlaps, of which one was an almost-complete dimer, suggestive of RCR. Three of the ambi-like clusters were detected at very low levels in 10 plant transcriptomes.

We then ran the detection pipeline on the 30 ambivirus and ambivirus-like sequences from GenBank and found significant ribozyme matches in 15 of these sequences of which 13 contained two predicted ribozymes. Of the remaining 15 sequences, 11 showed ribozyme matches in the expected locations that failed to pass the significance threshold (Table S4). As in the IMG data, the HHR3 and HPR-meta1 ribozymes are present in both matched and mismatched combinations. Similarly, three of the published genomes (MT354566.2, MN793994.1, and MT354567.1) contain terminal overlaps of 160–250 nt, suggestive of circularity. Furthermore, all known ambivirus and ambivirus-like sequences were predicted to adopt a rod-shaped conformation. Taken together, these observations strongly suggest that ambiviruses comprise a distinct group of ribozy-like viruses that encode a RdRP homologous to the RdRPs of riboviruses.

Three cccRNA clusters with significant mitovirus RdRP matches were detected, including two with symmetric ribozymes. The symmetric singletons are 3,283 and 3,058 nt in size and contain matched twister-P1 ribozymes and an HHR3/twister-P1 combination, respectively. The HHR3 aligns to ELVd with 96% identity. A third cccRNA cluster with three members encoding putative mitovirus-like RdRP, of 3,363 nt, contains a

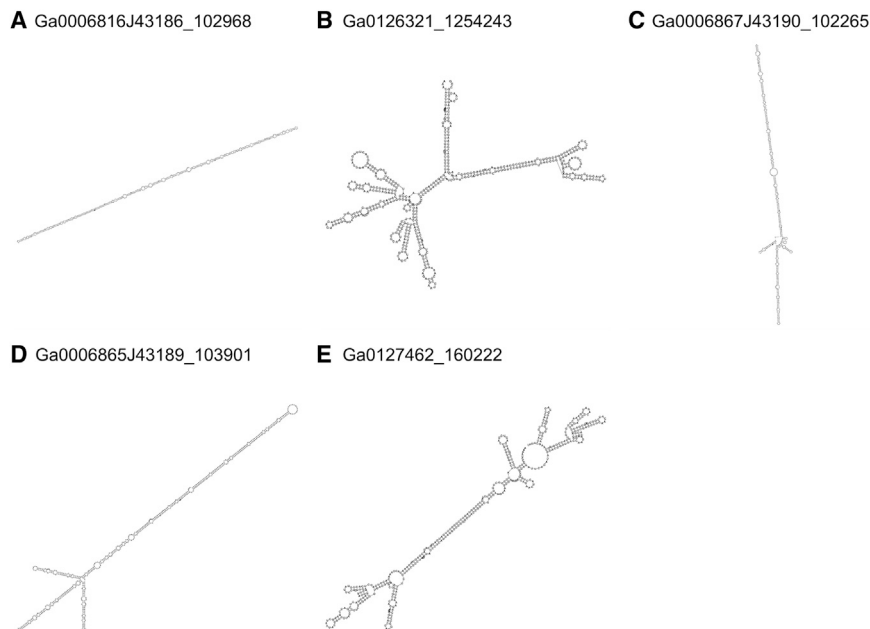


Figure 3. Predicted secondary structures of representatives of the five largest clusters of distinct viroid-like cccRNAs

Structures were predicted using ViennaRNA's RNAfold program configured to operate on circular sequences. Representative predicted structures are shown for clusters 1(A), 2 (B), 3 (C), 4 (D) and 5 (E). Sequence data and metadata are available in [Table S1](#).

similar match to ELVd (including the HHR conserved core) that was not identified as an HHR by either detection method. This cccRNA lacks the HHR core in the opposite polarity but shows weak similarity (E-value = 0.19) to twister-P1. All cccRNAs were detected with >100 nt overlaps and are predicted to adopt a branched conformation with between 63% and 66% of bases paired in both polarities. These three genomes have a low (36%–40%) GC content, a hallmark of mitoviruses.⁶⁰ Searching the predicted RdRPs against the protein sequence databases yielded the most significant matches for the three cccRNAs to Grapevine-associated mitovirus 13, Grapevine-associated mitovirus 14, and *Fusarium asiaticum* mitovirus 8. Upon closer inspection of Grapevine-associated mitoviruses 11 and 13, we found that they also contained HHR3s in both polarities and a twister-P1/HHR3 combination, respectively. Ribozyme searches of all available *Lenarviricota* (taxid 2732407) and unclassified *Riboviria* (taxid 2585030) sequences did not yield other matches besides the ambiviruses and these few mitoviruses.

Apart from the RdRps, we identified 135 sequences comprising 53 ANI90 clusters with significant similarity to capsid proteins of single-stranded (ss) DNA viruses, in particular, CRESS viruses.⁶¹ The sequences in 50 of these clusters contained predicted ribozymes, and 13 contained HHR3s in both polarities. Two clusters contained paired HHR3 and twister-P1 ribozymes, whereas two other clusters contained symmetric HP-meta1 ribozymes. 26 clusters contained a single HHR3, three a twister-P1, and four an HPR-meta1. 21 clusters, including all three without ribozyme profile matches, produced a nucleotide alignment to a known viroid's ribozyme, ranging in length from 25 to 50 nt at 83%–96% identity. The cccRNAs in these clusters varied between 1,092 and 1,632 nt in length, with a mean of 1,317 nt and GC content with the mean of 44%. Four cccRNAs were sequenced as complete head-to-tail

dimers. The secondary structures of these cccRNAs included, on average, 66% paired bases. Given the strong evidence of circularity, extensive self-complementarity resulting in predicted branched structure and confident prediction of ribozymes, these cccRNAs most likely represent a distinct group of ribozyme-like satellite viruses.

Diverse deltavirus-like viruses

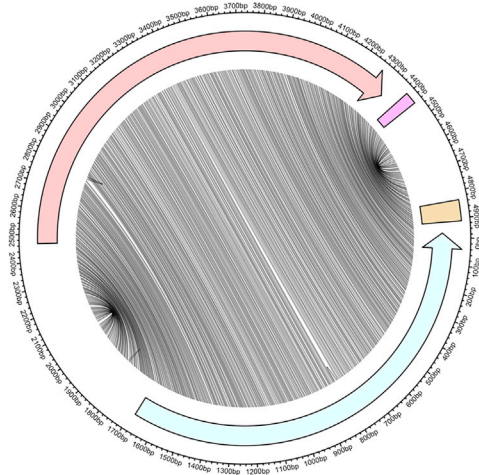
Apart from the viruses that resembled ribozyviruses conceptually, that is, were identified as protein-coding viroid-like

cccRNA but encoded proteins unrelated to HDV antigen (HDVAg), we searched for actual relatives of the deltaviruses. To this end, clusters of ORFs from the identified cccRNAs were compared to the sequences of the HDVAg and its homologs from other ribozyviruses. A total of 12 ORF clusters were identified above the HDVAg Pfam profile's gathering threshold; additional 21 representative ORFs were significant at the E-value < 1e-03 level and 34 at E-value < 1e-02. Of these clusters, only one showed a significant nucleotide alignment to a known HDV-like virus. The other clusters were found in a variety of environments ranging from soil to wastewater to coastal wetland sediment. Samples with matching cccRNAs were collected from as far north as Alaska to as far south as Florida. All 69 members of the clusters encompassing ORFs with HDVAg profile matches (E-value < 1e-02) were predicted to adopt a rod shape in both polarities. The genome size of ribozyviruses in ViroidDB ranges from 1,547 to 1,735 nt. However, among the HDV-like clusters identified in metatranscriptomes, the size ranged from 1,019 nt to 1,757 nt, with a median length of 1,317 nt.

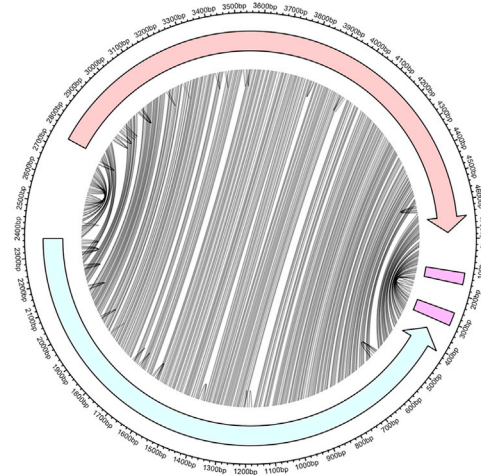
Clustering the HDV-like sequences in combination with the known ribozyviruses in ViroidDB produces no ANI80 clusters with both reference and distinct members. Each of the 26 HDV-like clusters falls below the species demarcation criterion for ribozyviruses (80% nucleotide identity).²⁸ Clustering the ORFs from both the detected HDV-like sequences and reference ribozyviruses with 60% minimum identity (the genus demarcation criterion) using CD-HIT resulted in 36 clusters, of which 10 consisted entirely of reference sequences whereas 26 contained only sequences discovered here.

Clustering of the HDVAg sequences and their homologs from other animals and metatranscriptomes with a permissive threshold showed that all previously known HDVAg homologs formed a single tight cluster whereas the metatranscriptome sequences

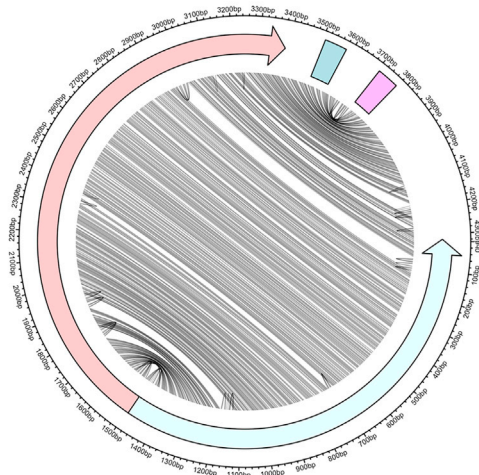
A MW423804.1



B MN793991.1



C Ga0181510_1007696



D Ga0180113_1206164

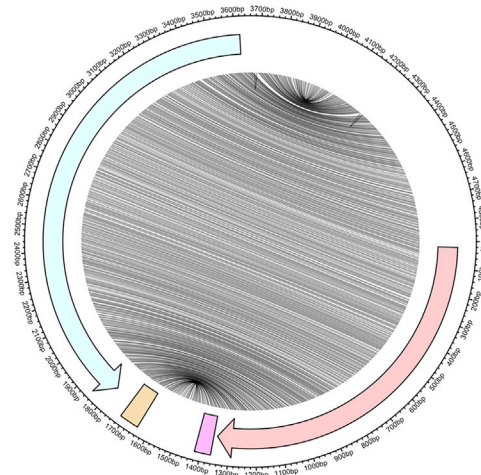


Figure 4. Covalently closed circular RNA genomes of ambi-like viruses

Depicted are the genomic and secondary structures of *Armillaria borealis* ambi-like virus 1 (A), *Tulasnella ambivirus* 1 (B), and ambi-like sequences discovered here (C and D). Red and blue denote (+) and (–) polarities, respectively. Lines connect bases in the genome that are paired in the predicted secondary structure. Arrows represent ORFs and rectangles represent self-cleaving ribozymes. The (+) and (–) ribozymes are HHR3 and HPR-meta1 in (A), HHR3 and HHR3 in (B), CPEB3 and HHR3 in (C), and HHR3 and HPR-meta1 in (D). In all cases, the ribozymes are located outside the ORFs at the end of the rod.

formed multiple smaller clusters and singletons distant from each other and from HDV (Figure 5A). The conservation profile of the multiple alignment of the HDVAg homologs showed that the dimerization region and one of the RNA-binding regions were prominently conserved whereas the second RNA-binding region was not (Figures 5B and 5C). The sequences of the distant HDVAg homologs from metatranscriptomes showed low sequence similarity to the previously known HDVAg, far below the similarity among the latter, with the distributions of percent identities almost non-overlapping (Figure 5D). Finally, in the phylogenetic tree of the HDVAg homologs, all previously known sequences formed one

compact clade, whereas the homologs from metatranscriptomes identified here comprised several remaining clades, with a much greater phylogenetic depth (Figure 5E).

The nucleotide sequences of these HDV-like cccRNAs formed 26 ANI90 clusters, none of which contained confidently predicted self-cleaving ribozymes. However, 13 of these clusters produced weak ribozyme matches (E-value < 1e-01), and eight of these were symmetric. Both HHR-like (n = 36) and HDV-like (n = 26) ribozymes were detected although no clusters contained ribozymes of both types. Of the HDV-like ribozymes detected, only five most closely matched the canonical HDV ribozyme.

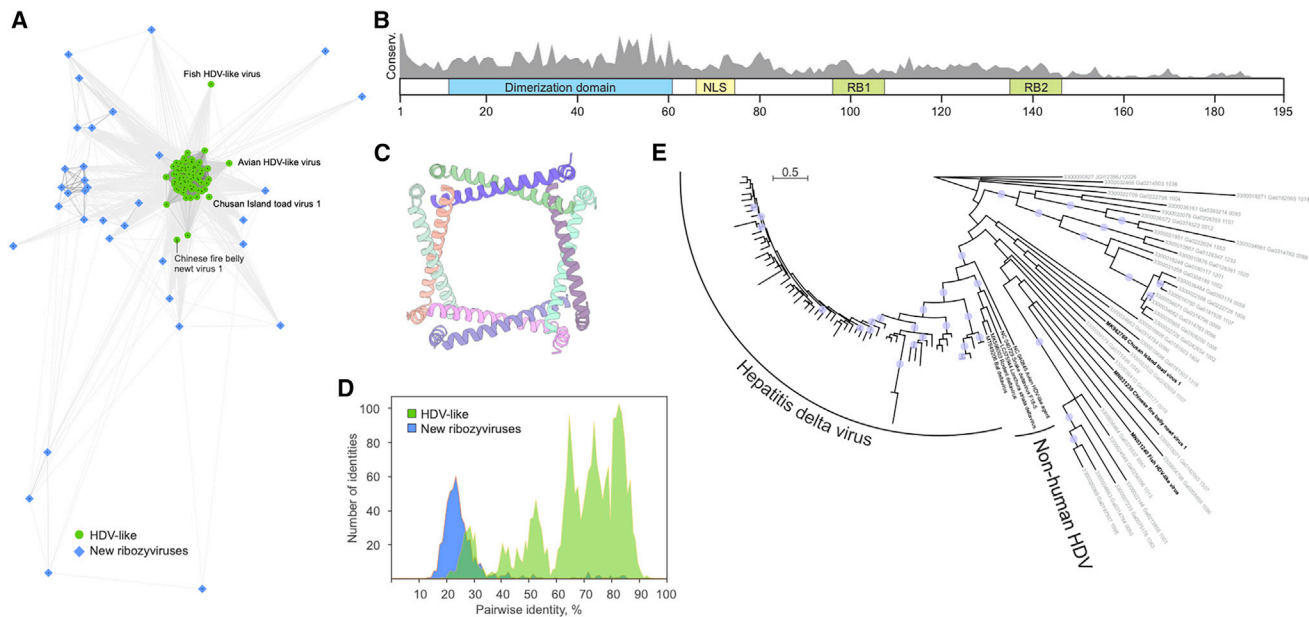


Figure 5. Diversity of HDV antigen-like proteins in known ribozyviruses and ribozyme-like viruses identified in metatranscriptomes

(A) Clustering of the HDV antigen (Ag)-like protein homologs based on their sequence similarity. Lines connect nodes (sequences) with p value $< 1e-05$. Reference HDVAg-like sequences from GenBank are shown as green circles, whereas those detected in metatranscriptomic datasets are shown as blue diamonds. Some of the divergent reference sequences are labeled.

(B) Schematic representation of the HDVAg with functionally important regions indicated with colored boxes. RB1 and RB2, RNA-binding sites 1 and 2, respectively; NLS, nuclear localization signal. Gray histogram shows the sequence conservation (percent identity) of HDVAg-like sequences from metatranscriptomic datasets.

(C) Octameric structure of the conserved dimerization domain of HDVAg. PDB ID: 1A92.⁶² Each protein molecule is shown with a different color.

(D) Comparison of the sequence conservation among reference HDVAg from GenBank (green) and those from metatranscriptomic datasets (blue).

(E) Maximum likelihood phylogeny of HDVAg-like sequences. The tree was constructed with IQ-TREE.⁶³ Circles at the nodes represent SH-aLRT support higher than 90%. The scale bar represents the number of substitutions per site.

10 putative ribozymes showed the strongest similarity to the HDV ribozyme (HDVR) found in the genome of *Faecalibacterium prausnitzii*,⁶⁴ seven were most similar to the HDV-like ribozyme found in the genome of *Anopheles gambiae*,⁶⁴ and four were most similar to the mammalian CPEB3 ribozyme.^{65,66} The causes of these similarities remain to be investigated; given the small size of the ribozyme, convergence cannot be ruled out.

The limited number of significant ribozyme matches among the HDV-like sequences posed an opportunity for detecting distinct ribozymes or divergent variants of known ones. For example, we examined an HDV-like cccRNA cluster (representative member 3300009579_Ga0115599_1049451) with no predicted ribozymes. However, upon closer examination, sequences from this cluster were shown to contain the conserved HHR core in both polarities in the expected locations, a recently discovered ribozyme configuration.⁶⁷ Some clusters entirely lacked the HHR core in either polarity, suggesting the use of alternative, yet-unknown ribozymes.

Diverse ribozymes and ribozyme combinations

Almost all viroid-like RNAs described to date contain the same type of ribozyme in both polarities, with the exception of some satRNAs. Surprisingly, many viroid-like cccRNAs identified here were predicted to contain ribozyme combinations that have not been so far reported in replicating cccRNAs (Figure 6).

Specifically, we identified numerous cccRNAs containing twister ribozymes, a recently described ribozyme motif that so far has only been found in combination with the HP-meta1 ribozyme. Both symmetric ($n = 381$) and asymmetric ($n = 930$) variants are present in the metatranscriptomic cccRNA clusters. Most symmetric twister clusters contained matched twister ribozymes (218 clusters) in both polarities, an unexpected ribozyme combination. In 87 clusters including mitovirus-like and satellite-like cccRNAs, we found another distinct combination of ribozymes, with HHRs opposite twister ribozymes. The unusual twister ribozyme is widespread in plant transcriptomes, with 59% of the transcriptomes containing reads mapping to a twister-bearing cccRNA. Indeed, we recovered three asymmetric cccRNA clusters from plants that contained a twister-P1 ribozyme.

In addition to the twister combinations, we identified several other unusual ribozyme combinations in symmetric cccRNAs. Previously, HHR3s have been found in conjunction with HDVAg⁶⁷ but have not been reported to be paired with HDV ribozymes. We identified three clusters in which HHR3s were paired with HDV-type ribozymes, namely, CPEB3 and HDVR *F. prausnitzii*. The CPEB3-HHR3 combination was found in an ambivirus-like sequence, and the two HDVR *F. prausnitzii*-HHR3 clusters were both predicted to adopt rod-shaped structures. One of these, a 1,052 nt singleton, did not match any sequences in ViroidDB, nt, or UniRef90, but in the other cluster

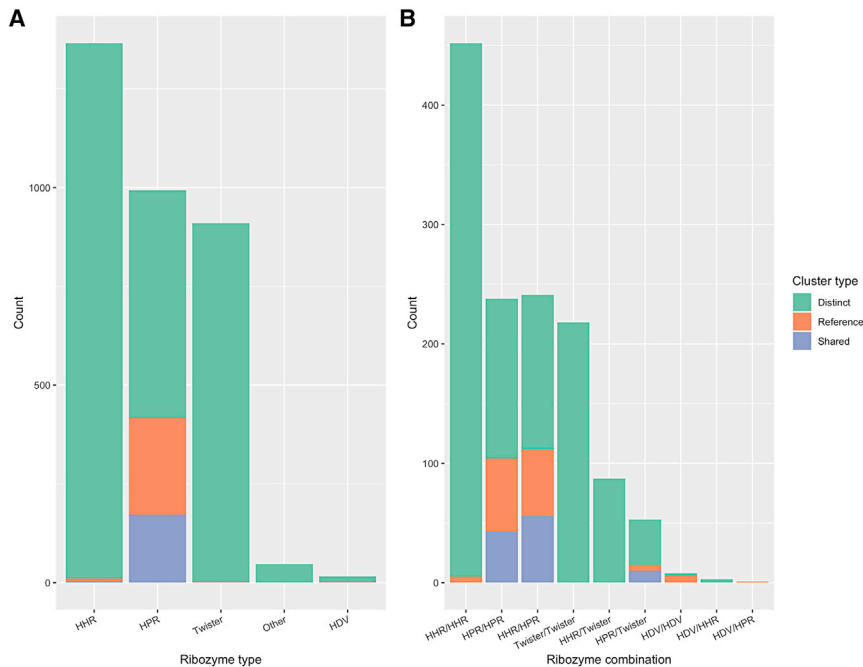


Figure 6. Ribozyme diversity in viroid-like cccRNAs

(A) Distribution of ribozyme types in asymmetric clusters.

(B) Ribozyme co-occurrence within the symmetric viroid-like cccRNA cluster representatives derived from metatranscriptomes.

(978 nt, two members), the HHR was closely similar to that of *Cryphonectria parasitica* ambivirus 1 (44/50 nt identical), whereas the HDV *F. prausnitzii* motif (32/33 nt and 41/46 nt) aligned to two chromosomes of the *Vanessa atalanta* butterfly.

Among the asymmetric cccRNAs, we identified two additional types of self-cleaving ribozymes. The hatchet ribozyme was found in 34 ANI90 clusters that ranged from 357 nt to 567 nt in length and came primarily from aquatic metatranscriptomes, in contrast to the general trend among the cccRNAs that derived from soil metatranscriptomes (Figure 7B). For example, the most diverse of these clusters (440 nt) contained 22 members from aquatic (almost all freshwater) metatranscriptomes sampled from around the United States. For the hatchet clusters, the Rfam profile matches were the strongest among all detected ribozymes (median E-value 1.4×10^{-8}) and, unusually for viroid-like cccRNAs, the GC content was low (median 35%). The predicted structures of these sequences varied from branched to quasi-rod shaped, with a mean of 62% of the bases paired.

The pistol ribozyme was identified in asymmetric cccRNAs. Like the clusters containing the hatchet ribozyme, clusters with the pistol ribozyme were found primarily (9/13) in marine metatranscriptomes ranging from the Antarctic Ocean to the Baltic Sea. The clusters have a slightly lower median profile match E-value (E-value = 6.4×10^{-5}) compared to the hatchet ribozyme but, unlike the hatchet ribozyme, have GC content ranging from 33% to 59% (median 49%), more characteristic of viroid-like RNAs. The predicted secondary structures of the pistol-containing cccRNAs were branched, often with several long hairpin structures.

CRISPR spacers matching cccRNAs

CRISPR spacer matches provide one of the most reliable means for assigning hosts to viruses and other mobile ge-

netic elements in prokaryotes and for differentiating prokaryote-infecting from eukaryote-infecting viruses.^{68,69} Our recent search for riboviruses in the same set of metatranscriptomes that is analyzed here identified multiple spacers matching RNA viruses, resulting in the assignment of several groups of viruses to bacterial hosts including several previously thought to infect eukaryotes.⁴³ To identify viroid-like agents that potentially might replicate in prokaryotes, we searched the viroid-like cccRNA sequences identified here against the IMG CRISPR spacer database,⁷⁰ and detected 89 spacers

with significant matches to viroid-like cccRNAs from nine clusters (Table S5).

One spacer was an identical match of 37 nt to a member of a cluster of 16 cccRNAs with prominent viroid-like features. The cccRNAs of this cluster are 315 nt long, contain symmetric HHR3s, and are predicted to adopt a rod shape, with 73% of the bases paired. Unusual for viroids, the cccRNAs comprising this cluster were found in Mushroom Spring, a hot spring at Yellowstone National Park. The matching spacer was also detected in a 60°C hot spring, Great Boiling Spring, albeit more than 800 km away.⁷¹ The repeats in this CRISPR locus were identical to those in the type III CRISPR locus of *Roseiflexus* sp. RS-1, an anoxygenic filamentous bacterium of the phylum *Chloroflexota* that was itself identified in Yellowstone hot springs.^{72,73} Searching for spacers matching all 16 cluster members with more relaxed criteria (i.e., more than one mismatch but with E-value < 1×10^{-5}), we identified a further 13 nearly identical matching spacers from eight Yellowstone hot springs samples collected between 2007 and 2017. One spacer (35/37 identities, E-value = 3×10^{-6}) included a precise match to the HHR core motif. The repeats from this expanded set of loci all matched those from *Roseiflexus* sp. RS-1. Previously, we identified multiple spacers in *Roseiflexus* sp. RS-1 that matched a group of partiti-like riboviruses that were accordingly assigned to this bacterial host.⁴³ Apparently, the type III CRISPR system of *Roseiflexus* sp. that encompasses a reverse transcriptase actively incorporates spacers from multiple RNA replicons.

The cluster with the most spacer matches—57 matches spanning 26 metagenomes, largely from sludge and bioreactor samples—includes a single cccRNA of 606 nt (recovered as 841 nt) with an asymmetric twister-P5 ribozyme and a predicted branched conformation with 63% of bases paired. Eight

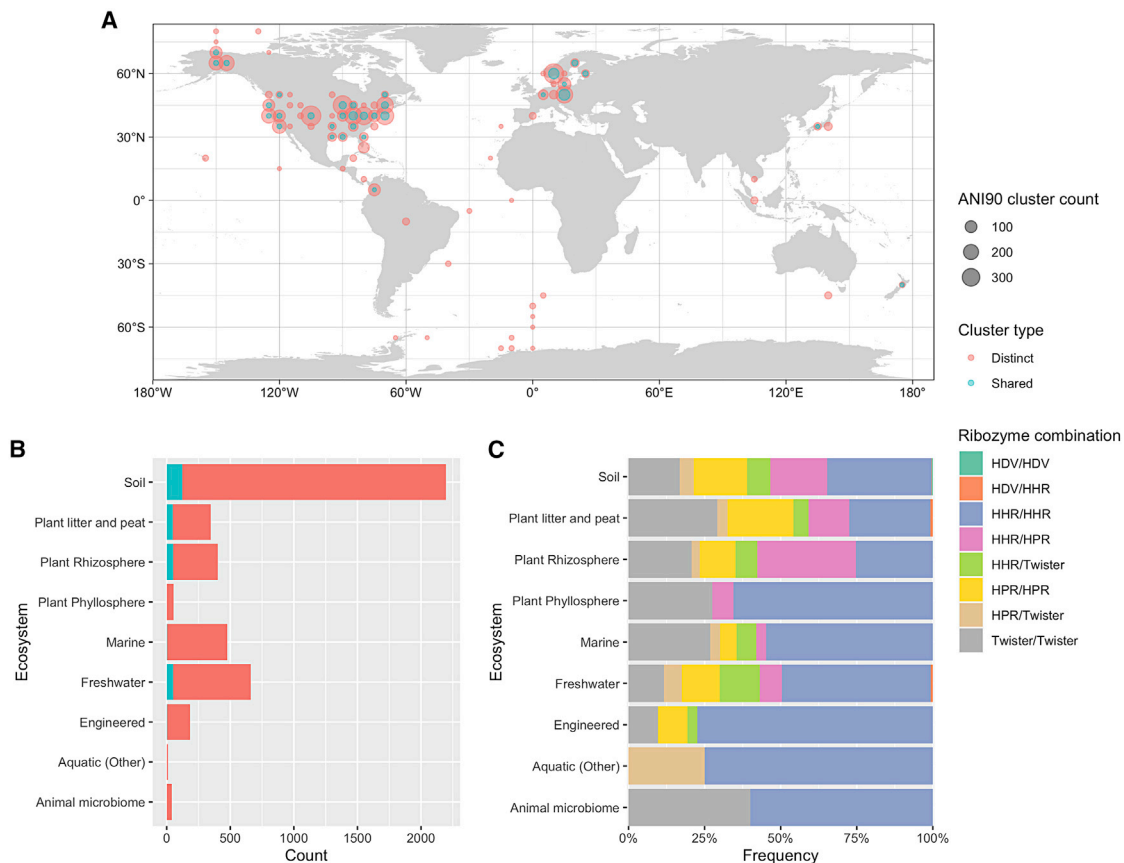


Figure 7. Global distribution and habitats of viroid-like cccRNAs found in metatranscriptomes

(A) Map of sample locations from which viroid-like cccRNAs were detected. The size of each circle corresponds to the number of clusters identified in each location (grouped to the nearest 5° of latitude and longitude) while the color represents the fraction of distinct clusters (blue shows reference clusters, and red shows distinct clusters).

(B) Reference (blue) and distinct clusters of viroid-like cccRNAs in different types of ecosystems.

(C) Relative frequencies of ribozyme combinations within symmetric cccRNA clusters in each ecosystem type.

spacers, detected in seven metagenomes, covered the predicted ribozyme region.

The second most frequently matched cluster, also a singleton, was associated with 13 spacers from 10 metagenomes, all from the same location in the Southern Indian Ocean. This cccRNA is 286 nt long (recovered with a 123 nt overlap) and contains a predicted HHR_{II} in one polarity only. Like the most-matched singletons, this sequence also is predicted to adopt a branched conformation, with 57% of bases paired. The spacers collectively covered 33% of the sequence but do not include the HHR region.

Geographic and ecological distribution of viroid-like cccRNAs

We examined the global distribution of the cccRNA clusters. Distinct clusters were found throughout the world (Figure 7A) and in all types of ecosystems (Figure 7B). Soil samples were the primary source of both distinct and shared clusters, reflecting both the greater number of such samples (twice as many as the next most common sample type) and the apparent greater sequence diversity in soils.

The viroid-like cccRNAs displayed non-uniform ribozyme distribution among ecosystems (Figure 7B). Mismatched HPR/HHR ribozymes were especially prevalent among samples from plant rhizospheres, whereas matched HHRs were notably more abundant in engineered ecosystems, such as bioreactors, than in soil environments.

Among the 10 most geographically dispersed, distinct clusters, eight included symmetric ribozymes, of which six were matched HHR₃s. The other two symmetric clusters contain HPR-meta1/twister-P1 ribozymes and the two asymmetric clusters contain HHR₃s. These widely dispersed clusters ranged in length from 372 nt to 1,039 nt and were predicted to adopt either a rod-like shape (the six HHR₃ clusters) or branched conformations.

We examined the coverage depth of the viroid-like cccRNAs relative to the other contigs within the same metatranscriptomes. The absolute coverage depth is not directly informative due to the variation between sequencing methods used, but the ranked coverage depth percentile can serve as a proxy for the abundance of a sequence within the sample. In the 747 samples for which coverage information was available, viroid-like cccRNAs

ranged from the most covered to least covered contigs. Notably, 150 viroid-like cccRNAs were within the top percentile (Table S7). Among these highly covered contigs are known viroid-like cccRNAs, such as satLTSV.

Identifying potential hosts of the viroid-like agents within these ecosystems remains a challenge. Based on the IMG annotation pipeline, the majority of the analyzed metatranscriptomes were dominated by prokaryotic sequences⁴³ but still contained at least 1% of contigs affiliated to eukaryotes (Table S6). Nonetheless, 187 metatranscriptomes in which viroid-like cccRNAs were detected contained <0.1% of eukaryotic contigs, suggesting that these elements replicate in either rare and undetected eukaryotes or in some of the much more abundant prokaryotic hosts. Notable among these datasets were the hot spring metatranscriptomes in which CRISPR spacers targeting viroid-like cccRNAs were identified (see above). The apparent lack of eukaryotic RNA in these samples strengthen the hypothesis of prokaryotic hosts. Additionally, we found 104 symmetric clusters in marine samples that are far beyond the habitation range of the known hosts of viroids and satRNAs. These findings combined with the clusters from prokaryote-dominated samples suggest that the ecological and host ranges of viroid-like agents are far broader than currently appreciated.

DISCUSSION

Viroids and viroid-like cccRNAs, such as satRNAs and ribozyviruses, are the smallest, simplest known replicators that hijack either a host DNA-dependent RNA polymerase or a virus RdRP for their replication. Given the universality of the cellular transcription machinery across all life forms and the enormous diversity and near ubiquity of RdRP-encoding riboviruses, the narrow diversity and host range of the known viroid-like agents appeared puzzling. We suspected that viroid-like agents could actually be far more common than presently known and, with this motivation, searched a collection of more than 5,000 metatranscriptomes for viroid-like cccRNAs.

We identified millions of putative cccRNAs by searching for signatures of circularity or RCR, namely, the presence of head-to-tail repeats in assembled contigs. Because reads spanning the origin cannot be reconciled with a linear sequence, the assembler produces contigs with the same subsequence repeated at both the end and the beginning.⁴⁵ Alternatively, when linear replication intermediates containing head-to-tail repeats are sequenced, the ends of the sequences are also repeated. After compensating for the low fidelity of RNA polymerase II by allowing for up to 5% mismatches in the repeated regions, testing this method on assembled plant transcriptomes demonstrated that known viroids were reliably recovered in the absence of major assembly errors. However, the extensive secondary structure of many viroid-like cccRNAs and the use of poly-A enrichment during RNA isolation prior to sequencing⁷⁴ makes it likely that many viroid-like cccRNAs either were not sequenced at all or were grossly misassembled and thus could not be recognized as circular. Even under a conservative approach, where only predicted cccRNAs containing confidently identified ribozymes counted as “viroid-like,” this search resulted in an approximately 5-fold increase in the diversity of

viroid-like agents. This is most likely a substantial underestimate of the true span of the viroid-like domain of the replicator space because among the millions of the predicted cccRNAs, in which no ribozymes were confidently identified, some, and possibly many, could be viroid-like agents containing unknown ribozymes or lacking ribozymes altogether, like pospiviroids. Although perhaps only a coincidence, it is worth noting that a recent analysis of the same collection of metatranscriptomes also yielded an approximately 5-fold increase in the diversity of riboviruses.⁴³ It is further notable that a substantial number of viroid-like cccRNAs are among the most abundant sequences in the respective metatranscriptomes, emphasizing the prominence of these agents in diverse ecosystems.

The majority of the detected viroid-like cccRNAs possessed characteristic features of viroids including the presence of HHR, often in both polarities, and predicted rod-like or extensive branched conformation. However, the search resulted not only in quantitative expansion of viroid diversity but also in qualitatively distinct findings, in particular, unexpected ribozyme combinations, such as those including twister, and ribozymes not previously found in viroid-like RNAs, such as hatchet and pistol. There is little doubt that additional ribozymes and ribozyme combinations in viroid-like cccRNAs remain to be discovered.

Another key finding is the discovery of diverse groups of ribozy-like viruses. Even if perhaps not unexpected, it is notable that the diversity of the ribozyvirus sequences discovered in metatranscriptomes far exceeds that of the previously known HDV relatives including the recently discovered non-mammalian ones. Moreover, many of the identified ribozyviruses lack the HDV ribozyme or even any known ribozymes, suggesting distinct replication mechanisms. The host range of the discovered ribozyviruses remains to be explored but, probably, includes non-animal hosts (see discussion below).

In contrast, the demonstration that ambiviruses are actually viroid-like agents and the discovery of viroid-like mitoviruses and satellite viruses was surprising. These three groups of viroid-like agents resemble ribozyviruses in that these are relatively large, protein-coding viroid-like cccRNAs. However, unlike HDV and its relatives, these viroid-like agents are clearly linked to riboviruses through the RdRPs encoded by ambiviruses and mitoviruses, and capsid proteins encoded by satellite viruses. These findings show that combinations of viroid-like cccRNA and protein-coding genes emerged on multiple occasions during evolution. These ribozy-like viruses unrelated to deltaviruses likely evolved through recombination between typical riboviruses and viroids. The implications for virus taxonomy, in particular, whether such viruses should be classified into the existing realm *Ribozyviria* or into the respective divisions of the realm *Riboviria*,⁷⁵ or perhaps, into a separate realm, remain to be sorted out.

One of the most interesting but also most challenging problems is the host range of the expanded diversity of viroid-like agents. There is currently no direct computational approach for connecting viroid-like RNAs with specific hosts. Nevertheless, it appears exceedingly unlikely that all or even the majority of the viroid-like cccRNAs discovered in metatranscriptomes are parasites of plants. Indeed, we identified orders of magnitude more viroid-like cccRNAs in metatranscriptomes than in plant transcriptomes, and most of the analyzed metatranscriptomes

are dominated by bacteria followed by unicellular eukaryotes. Furthermore, ambiviruses were isolated from fungi,^{57–59} and the demonstration that these are viroid-like agents expands the host range of the latter. A potential prokaryotic connection of viroid-like cccRNAs through CRISPR spacer matches is particularly notable. The detected spacer matches were not numerous but reliable, in particular, because multiple spacers matching the same cccRNA were identified in diverse metagenomes. At least, the typical viroid-like cccRNAs that matched spacers from the reverse-transcribing type III CRISPR system of *Roseiflexus* sp. appear to be strong candidates for distinct bacterial parasites. These viroid-like cccRNAs that likely replicate in bacteria merit further, dedicated metatranscriptome and metagenome searches as well as experimental investigation. These findings echo the recent expansion of the bacterial RNA virome through the search of the same metatranscriptome collection and suggest that bacteria might support a much greater diversity of RNA replicators than previously suspected.⁴³

LIMITATIONS OF THE STUDY

This work targets low hanging fruits in the search for viroid-like agents, being limited to the cccRNAs that contain reliably identifiable, known ribozymes or align directly to known viroid-like agents. This conservative approach was adopted purposefully in order to avoid potential artifacts resulting from erroneous identification of cccRNA or contamination with DNA-encoded sequences or other sources. A potential opportunity for the discovery of a far greater diversity of viroid-like agents and a challenge for further research is a comprehensive analysis of the massive set of predicted cccRNAs that lack known ribozymes. Computational methods for *de novo* prediction of ribozymes need to be developed to advance such analyses. Furthermore, there are numerous additional metatranscriptomes, in particular, those recently analyzed by the Serratus⁴¹ and Tara⁴² teams, as well as numerous animal metatranscriptomes, that should be searched for viroid-like cccRNAs. To make a comprehensive search practicable, more efficient algorithms for circularity detection are required. A technical limitation is that some of the software used to assemble metatranscriptomes (Table S5) would cut circular molecules to unit length excluding terminal repeats. Although the method we used for cccRNA detection took into account even short repeats, this feature of some assemblers could yield false negatives. Evidently, the computational approaches applied in this work only identify candidates for viroid-like agents. Experimental validation is needed and is especially important in the case of putative cccRNAs lacking known ribozymes.

STAR★METHODS

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SUPPLEMENTAL INFORMATION

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AUTHOR CONTRIBUTIONS

E.V.K. accepted the project; B.D.L. and E.V.K. designed research; B.D.L. and U.N. compiled the datasets; B.D.L., U.N., S.R., A.P.C., Y.I.W., and M.K. analyzed the data; P.S., U.G., N.K., V.V.D., and E.V.K. supervised research; B.D.L. and E.V.K. wrote the manuscript that was read, edited, and approved by all authors.

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The authors declare no competing interests.

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STAR★METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Deposited data		
All original code and data for this paper	This paper	https://doi.org/10.5281/zenodo.6859104
The viroid-like cccRNA detection pipeline	This paper	https://github.com/Benjamin-Lee/vdsearch
Software and algorithms		
CD-HIT v4.8.1	Li and Godzik ⁷⁶	https://github.com/weizhongli/cdhit
HMMER v3.3.2	Eddy ⁷⁷	https://github.com/EddyRivasLab/hmmer
Infernal v1.1.4	Nawrocki and Eddy ⁴⁶	https://github.com/EddyRivasLab/infernal
python-igraph v0.9.10	Gábor and Nepusz ⁷⁸	https://github.com/igraph/python-igraph
scikit-bio v0.5.6	Caporaso et al. and Knight et al. ^{79,80}	https://github.com/biocore/scikit-bio
MMseqs2 v13.45111	Steinegger and Söding ⁸¹	https://github.com/soedinglab/mmseqs2
R v4.2.0	R Foundation for Statistical Computing	https://www.r-project.org
Python v3.8.3	Python Software Foundation	https://www.python.org
Nim v1.6.2	Rumpf ⁸²	https://nim-lang.org
ViennaRNA v2.5.0	Lorenz et al. ⁸³	https://github.com/ViennaRNA/ViennaRNA
SeqKit v2.1.0	Shen et al. ⁸⁴	https://github.com/shenwei356/seqkit
Pandas v1.2.0	McKinney ⁸⁵	https://github.com/pandas-dev/pandas/
ggplot2 v3.3.6	Wickham ⁸⁶	https://github.com/tidyverse/ggplot2/
orfiy v0.0.4	Singh et al. ⁸⁷	https://github.com/urmi-21/orfiy
fastp v0.20.1	Chen et al. ⁸⁸	https://github.com/OpenGene/fastp
Snakemake v6.10.0	Mölder et al. ⁸⁹	https://github.com/snakemake/snakemake
circlize v0.4.15	Gu et al. ⁹⁰	https://github.com/jokerghoo/circlize
RNAmotif v3.1.1	Macke et al. ⁹¹	https://github.com/dacase/rnamotif
maSPAdes v3.14.1	Bushmanova et al. ⁹²	https://github.com/ablab/spades
MinCED v0.4.2	Bland et al. ⁹³	https://github.com/ctSkennerton/minced
IQ-TREE v1.6.12	Nguyen et al. ⁶³	https://github.com/iqtree/iqtree2
CLANS	Frickey and Lupas ⁹⁴	http://protevo.eb.tuebingen.mpg.de/download
Sequence Demarcation Tool (SDT) v1.2	Muhire et al. ⁹⁵	https://github.com/brejnev/SDTv1.2
PROMALS3D	Pei and Grishin ⁹⁶	http://prodata.swmed.edu/PROMALS3D
ITOL v6.5.8	Letunic and Bork ⁹⁷	https://itol.embl.de
bowtie2 v2.4.2	Langmead and Salzberg ⁹⁸	http://bowtie-bio.sourceforge.net/bowtie2/index.shtml
BLAST + suite	Altschul et al. and Camacho et al. ^{99,100}	https://blast.ncbi.nlm.nih.gov/
EMBOSS etandem v6.0.0	Rice et al. ¹⁰¹	http://emboss.sourceforge.net

RESOURCE AVAILABILITY

Lead contact

Further information and requests for resources and data should be directed to and will be fulfilled by the lead contact, Benjamin Lee (benjamin.lee@chch.ox.ac.uk).

Materials availability

This study did not generate unique reagents. All data used as inputs for the pipeline are listed in the “data acquisition” section. All output of analyses are listed in the “data and code availability” section below.

Data and code availability

- This paper analyzes existing, publicly available data. These accession numbers for the datasets are listed in the [key resources table](#). Data generated during downstream analysis have been deposited at Zenodo and are publicly available as of the date of publication. DOIs are listed in the [key resources table](#).
- All original code has been deposited at Zenodo and is publicly available as of the date of publication. DOIs are listed in the [key resources table](#).
- Any additional information required to reanalyze the data reported in this paper is available from the [lead contact](#) upon request.

METHOD DETAILS

Data acquisition

The search for cccRNAs was performed on a collection of 5,131 assembled metatranscriptomes sourced from the IMG/MER database. In addition to the IMG metatranscriptomes, we searched the complete set of transcriptomes of the 1,000 Plants (1KP) project, totalling 1,344 paired-end sequencing runs. Before applying the search pipeline to 1KP, we filtered the raw reads for quality using fastp⁸⁸ and assembled them using rnaSPAdes⁹² using default parameters. We also included the 2021-09-07 release of ViroidDB and the set of HPR sequences identified by.⁵³

Circularity detection

We identified cccRNAs using a modified and improved version of the reference-free and *de novo* Cirit algorithm^{45,102} implemented in the Nim programming language. This method relies upon assembly errors for circular sequences resulting in terminal repeats. Detecting cccRNAs via this method requires searching forward for the last several bases of the sequence (the seed) and, if a match is found, comparing backwards to the start of the sequence. If the start and end of the sequence “overlap”, this repetitive region is then trimmed off. However, the existing implementation was unable to monomerize multimeric transcripts resulting from rolling circle replication among known viroid-like agents due to its single pass design and requirement of exact sequence identity within repetitive regions. Our reimplemention solves these problems by reiteratively attempting to monomerize putative cccRNAs while allowing for a configurable minimum identity within repeats. For this study, we required a minimum of 95% identity with no insertions or deletions within the overlapping region. In addition, the ratio of the length of the contig and computed monomers is reported. Sequences with monomer lengths below a threshold of 100 nt were excluded.

cccRNA deduplication

Standard approaches to sequence deduplication are insufficient for cccRNAs. Most modern approaches rely on hashing for memory efficiency. Such approaches are effective for linear sequences but circular sequences pose a challenge due to the arbitrariness of their start position. To enable deduplication of putative cccRNAs, we define a sequence’s canonical representation as the alphabetically earlier of the lexicographically minimal rotations of the sequence and its reverse complement. This approach, drawn from k-mer counting methods^{103,104} and, if further optimized, would be able to be computed in linear time and with constant memory for even greater scalability.¹⁰⁵

Ribozyme-based filtering

To identify sequences likely to replicate via ribozyme-catalyzed rolling circle replication, we searched the cccRNAs for the presence of known self-cleaving ribozymes using Infernal.⁴⁶ In each polarity, we identified ribozymes above Rfam’s curated gathering cutoff or with E-values <0.1. Sequences with ribozymes in both polarities that met these criteria were considered viroid-like. Alternatively, we considered sequences as viroid-like with one ribozyme with an E-value <0.01 or a score above the gathering cutoff. For each polarity, we considered only the most significant (by E-value) ribozyme. To identify more divergent ribozymes that were not detected using Infernal, we searched sequences containing one significant (E-value <0.01) using RNAmotif.⁹¹

Sequence search

We searched all cccRNAs against ViroidDB using MMseqs2 easy-search (version 13.45111)⁸¹ with the highest available sensitivity (-s 7.5). For each sequence, we considered only the most significant match as determined by bit score. In addition to searching ViroidDB, we also searched the cccRNAs identified by metatranscriptome mining against the set of cccRNAs recovered from plant transcriptomes using the same method.

RNA secondary structure prediction

We predicted the secondary structures of all viroid-like cccRNAs for both polarities using the ViennaRNA package.⁸³ For each predicted structure, we computed the percentage of bases paired and the number of hairpins present. We used a temperature of 25°C and the circular prediction mode.

Clustering

We performed several types of clustering including both alignment-based and alignment-free methods. To produce the alignment-based clustering, we performed an all-versus-all search using MMseqs2 on the sequences of ViroidDB, the HPR dataset of,⁵³ and this study. For this method, each sequence was concatenated to itself to compensate for potential variation in the sequence relative to otherwise-similar sequences due to their circular nature. Next, we executed MMseqs2 (v13.45111, easy-search -s 7.5 -min-seq-id 0.40 -search-type 3 -e 0.001 -k 5 -max-seqs 1000000) and computed the pairwise average nucleotide identity (ANI) between sequences by taking the alignment identity of the best hit for each pair. We computed the ANI for two self-concatenated sequences by taking the length of the smaller sequence and dividing by two. We then cap the computed alignment length the length of the now-monomerized smaller sequence. The ANI is then defined as the percent identity within the aligned region times the alignment length divided by the smaller sequence monomer length. Similarly, we defined the alignment fraction as the smaller of the doubled query coverage, doubled target coverage, or one.

To cluster the viroids based on their pairwise ANI, we build a graph by connecting pairs of sequences where the alignment covers at least 25% of the shorter sequence with 40% identity within the alignment. We then weighted the connections between the sequences by the ANI and employed the Leiden algorithm (as implemented in the igraph Python library, version 0.9.10) to delineate communities of similar sequences.¹⁰⁶ The clustering granularity was optimized by iterating over the resolution parameter space until the difference between average intra-cluster ANI and the target ANI began to increase.

ORF prediction

To find ORFs present within the sequences, we used orfipy⁸⁷ configured to operate on circular sequences. Specifically, we searched sequences concatenated to themselves to ensure ORFs spanning the origin were detected. Only ORFs longer than 100 amino acids and using the standard genetic code were considered for each cccRNA.

Protein searches

We searched all viroid-like cccRNAs for matches to known proteins. The primary search method we used was by performing translated searches (BLASTX-style) against the UniRef90 protein database¹⁰⁷ using MMseqs2.⁸¹ For each cccRNA, we considered only the best match by E-value.

As a second approach, we also searched the ORFs from all cccRNAs, viroid-like or not, using HMMER.⁷⁷ We searched both the full Pfam-A profile database using hmmscan as well as a curated subset (the profiles for RdRP clan combined with the HDVAg profile) using hmmsearch.

HDV antigen analysis

Sequences were clustered using CLANS with BLASTP option (BLOSUM62 matrix, E-value cutoff of 1e-03).⁹⁴ Sequence similarity among reference HDVAg from GenBank and those from metatranscriptomic datasets was analyzed with the Sequence Demarcation Tool.⁹⁵ For phylogenetic analysis, HDVAg-like sequences were aligned using PROMALS3D.⁹⁶ Due to the short length of the sequences, the alignment was not further processed. Maximum likelihood phylogenetic analysis was performed using IQ-TREE.⁶³ The best fitting model was selected by IQ-TREE and was VT + F + R4. The tree was visualized with iTOL.⁹⁷

Read mapping

We used bowtie2 to perform read mapping from the 1KP transcriptomes to the entire viroid-like cccRNA dataset in parallel. We configured bowtie to use its most sensitive setting (-very-sensitive) and ignore unaligned reads.

CRISPR spacer analysis

Viroid-like sequences were compared to predicted CRISPR spacers from prokaryotic (meta)genomes to identify potential cases of spacer acquisition from, and possible defense against, viroids by prokaryotes. The full set of 22,109 viroid and viroid-like sequences, including all reference sequences and sequences identified in this work, was compared to 1,961,109 CRISPR spacers predicted from whole genomes of bacteria and archaea (vJune2022) and 61,658,467 CRISPR spacers predicted from metagenomes in the IMG database⁷⁰ using BLASTN v2.9.0 with options -dust no -word_size 7. To minimize the number of false-positive hits due to low-complexity and/or repeat sequences, CRISPR spacers were excluded from this analysis if (i) they were encoded in a predicted CRISPR array including 2 spacers or less, (ii) less than 66% of the predicted repeats were 100% identical to each other, (iii) the spacers were at most 20 bp, or (iv) they included a low-complexity or repeat sequence as detected by dustmasker (v1.0.0)¹⁰⁸ (options -window 20 -level 10) or a direct repeat of at least 4 bp detected with etandem¹⁰¹ (options -minrepeat 4 -max-repeat 15 -threshold 2). To initially link viroid-like sequences to CRISPR spacers, only hits with 0 or 1 mismatch over the entire spacer were considered (Table S5). To find additional spacer matches, we searched all members of the clusters with a spacer match against the IMG public metagenomic spacer data (dated 2022-06-18) set using IMG's workspace BLAST with a minimum E-value of 1e-05. We also extracted the repeats matching loci using MinCED⁹³ and searched them against nt using BLASTN v2.13.0.^{99,100}

QUANTIFICATION AND STATISTICAL ANALYSIS

No statistical analysis was involved in this study apart from the sequence similarity searches. For ribozyme detection, significance was defined by a single ribozyme match with E-value <0.01 or a score above the gathering threshold. Additionally, two ribozymes were considered significant if both had E-values <0.1 . For CRISPR spacer detection, the BLASTN cut-off was set at $E < e^{-05}$. For protein sequence similarity, the BLASTX cut-off was set at $E < 0.01$. The rest of the search parameters are indicated in the respective section of [STAR Methods](#).