Virus-derived small RNAs: molecular footprints of host–pathogen interactions

Eric Roberto Guimarães Rocha Aguiar, Roenick Proveti Olmo and João Trindade Marques*

Viruses are obligatory intracellular parasites that require the host machinery to replicate. During their replication cycle, viral RNA intermediates can be recognized and degraded by different antiviral mechanisms that include RNA decay, RNA interference, and RNase L pathways. As a consequence of viral RNA degradation, infected cells can accumulate virus-derived small RNAs at high levels compared to cellular molecules. These small RNAs are imprinted with molecular characteristics that reflect their origin. First, small RNAs can be used to reconstruct viral sequences and identify the virus from which they originated. Second, other molecular features of small RNAs such as size, polarity, and base preferences depend on the type of viral substrate and host mechanism of degradation. Thus, the pattern of small RNAs generated in infected cells can be used as a molecular footprint to identify and characterize viruses independent on sequence homology searches against known references. Hence, sequencing of small RNAs obtained from infected cells enables virus discovery and characterization using both sequence-dependent strategies and novel pattern-based approaches. Recent studies are helping unlock the full application of small RNA sequencing for virus discovery and characterization. © 2016 Wiley Periodicals, Inc.

INTRODUCTION

Viruses are intracellular obligatory parasites that repurpose the host cell machinery to replicate. Viruses can create modified intracellular compartments referred to as viral factories where they concentrate resources required for replication and are shielded from host antiviral mechanisms.1,2 However, because viruses do not encode their own ribosomes, viral messenger RNAs (mRNAs) require access to cellular ribosomes to be translated. When viral mRNAs traffic to cellular ribosomes, they are exposed to host defense mechanisms more than other products generated during virus replication. Viral genomic RNAs are also excellent targets of host surveillance mechanism as they can be significantly different from cellular molecules. Thus, viral nucleic acids are a common target of several host antiviral mechanisms.3–6 Targeting of viral RNAs often results in the generation of virus-derived small RNAs (vsRNAs) that can be detected during infection in fungi, plants, arthropods, and mammals.7–13 Although vsRNAs are commonly observed in different organisms, their relative abundance can vary substantially.8 In insects, for example, viral sequences are 10-fold enriched in the small RNA fraction compared to what was observed in the long RNA pool from infected cells.10 In contrast, viral sequences in lungs of infected mice were underrepresented by a factor of 100 in the pool of small compared with long RNAs.10 This relative enrichment or depletion is likely caused by

*Correspondence to: jtm@ufmg.br
Department of Biochemistry and Immunology, Instituto de Ciências Biológicas, Universidade Federal de Minas Gerais, Belo Horizonte, Brazil

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differential targeting of viruses by RNA surveillance mechanisms. Nevertheless, targeting of viral transcripts and genomes is a common feature of different antiviral mechanisms and the production of vsRNAs can be highly informative. An analogy can be made about the indirect information about viruses obtained from vsRNAs and the mammalian antibody response. As initially proposed by Rivers and coworkers, virus-specific antibody responses can be indicative of an infection without requiring direct detection of the virus.14 Similarly, vsRNAs generated by host pathways can also provide information about the infection without the requirement for direct detection of the virus.

**FIGURE 1** Legend on next page.
HOST PATHWAYS THAT GENERATE vsRNAs

Viruses have evolved efficient mechanisms to escape host defense and ensure their survival. However, viral transcripts and genomes are often significantly different from cellular molecules, which allows them to be targeted by host pathways. Viral RNAs can be recognized by different RNA surveillance mechanisms, which often lead to their degradation and the generation of vsRNAs. RNA interference (RNAi), RNA decay, and RNase L pathways are good examples of such RNA surveillance mechanisms that have been extensively reviewed elsewhere. Here, we provide a few examples of how RNA degradation pathways may generate small RNA products with distinct molecular characteristics (Figure 1).

RNA Decay

RNA decay mechanisms play an important role in the quality control and turnover of cellular transcripts. Cellular transcripts are characterized by bona fide stability determinants such as the 5’ cap and 3’ poly(A) tails that are required to optimize translation. Transcripts that lack canonical features are targeted by several RNA decay mechanisms. In the deadenylation-dependent mRNA decay, the poly(A) tail of transcripts is shortened by the action of different deadenylases generating an unprotected 3’ end that can be degraded 5’–3’ by a large complex of exonucleases known as exosome. After poly(A) shortening, the transcript can also be degraded by removal of the 5’ cap by mRNA-decapping enzymes (DCPs), which allows the transcript to be targeted by XRN1, an exoribonuclease that carries out 5’–3’ degradation. In addition, nonsense-mediated decay (NMD) is a highly conserved deadenylation-independent mechanism capable of detecting aberrant mRNAs with short open reading frames and premature termination codons. NMD triggers endonucleolytic cleavage of target transcripts that is followed by degradation promoted by XRN1 and the exosome complex. RNA degradation by XRN1 and the exosome generates not only nucleosides but also small RNA fragments with 3’ OH and/or 5’ monophosphate groups depending on the structure and secondary modifications found in the substrate. These features help to explain how viral RNAs are often detected as aberrant transcripts by RNA decay mechanism.
decay mechanisms help monitor proper translation in infected cells, thus playing important roles in the antiviral response. In addition to RNA decay, other specialized antiviral pathways can also target viral RNAs and generate vsRNAs in eukaryotes.

RNA Interference

RNAi is a conserved mechanism of sequence-specific regulation of gene expression found in most eukaryotes. Initiation of RNAi requires recognition of an RNA substrate by specialized nucleases that are often type III RNases such as Dicers responsible for the biogenesis of small RNAs. During the effector phase of RNAi, these small RNAs associate with Argonaute proteins to form the RNA-induced silencing complex (RISC). RISC will utilize the small RNA sequence to select target RNAs via Watson–Crick base pairing, which will be degraded by Argonaute-mediated endonucleolytic cleavage.\(^{20}\) RNA fragments generated by RISC cleavage are short lived likely because they are further degraded by regular RNA decay mechanism. Small RNAs generated by both Dicer and Argonaute have 5′ monophosphates and 3′ OH groups. However, small RNAs generated during the initiation phase that associate with Argonaute proteins may be further stabilized by secondary modifications such as 2′O-methylation.

There are different RNAi pathways in eukaryotes that differ in their mechanism of activation, small RNA biogenesis, and action.\(^{22,23}\) In animals, there are at least three separate RNAi mechanisms that can generate vsRNAs during viral infection: the microRNA (miRNA), piwi-interacting RNA (piRNA), and small interfering RNA (siRNA) pathways (Figure 1). Each different RNAi pathway requires specific types of Argonaute–small RNA complexes.\(^{32}\)

The miRNA pathway is triggered by the recognition of short hairpins (~65 nt) formed by secondary structures found in long single-stranded RNAs.\(^{33}\) These internal hairpins are processed in two steps by two double-stranded RNA (dsRNA)-specific ribonucleases, Drosha and Dicer, to generate small RNA duplexes of ~20–23 nt that will be subsequently loaded onto an Argonaute protein to form miRISC.\(^{34}\) This complex will target complementary regions in the 3′ UTR of cellular mRNA leading to translation inhibition and mRNA destabilization.

Activation of the siRNA pathway is triggered by Dicer-mediated recognition of long dsRNA that is processed progressively to generate phased duplex small RNAs of ~20–23 nt.\(^{35}\) siRNA duplexes are loaded onto specialized Argonaute proteins to generate siRISC that remains associated with one of the strands.\(^{36}\) Mature siRISC is a multiple turnover enzyme that can efficiently catalyze endonucleolytic cleavage of RNA targets.\(^{37,38}\)

Unlike miRNA and siRNA, the piRNA pathway is triggered by the recognition of single-stranded RNA precursors seemingly independent on the presence of secondary structures.\(^{39}\) This single-stranded RNA precursor is processed by an endonuclease that generates phased single-stranded small RNAs that are ~24–30 nt long.\(^{40}\) The generation of these primary piRNAs requires a distinct subgroup of animal Argonautes known as PIWI proteins.\(^{40}\) Primary piRNAs can also trigger the production of secondary piRNAs by a self-amplifying cycle dependent on PIWI proteins known as the *ping-pong* mechanism.\(^{41,42}\) The complex formed by piRNAs and PIWI proteins known as piRISC can mediate transcriptional silencing as well as target RNA degradation.\(^{39}\) In contrast to siRNAs and miRNAs, piRNAs are commonly enriched in the germline and reproductive tissues of different animals.\(^{31,43-45}\)

RNAi pathways can have different roles during viral infection. The siRNA pathway is directly involved in the antiviral immunity in most eukaryotes including fungi, animals, and plants; although in mammals, this seems to be restricted to undifferentiated cells.\(^{12,46}\) In contrast, production of virus-derived miRNAs and piRNAs during viral infection appears to be more restricted. Activation of the piRNA pathway by virus infection has been reported in insects, although its antiviral role remains unclear.\(^{10,13,47-51}\) Generation of virus-derived miRNAs seems to be restricted to some animal viruses that exploit this pathway to control the expression of viral and cellular transcripts and help their replication.\(^{24,25}\)

Mammalian RNase L

Mammals have developed an antiviral system based on the degradation of viral RNAs by a specialized ribonuclease known as RNase L\(^{19}\) (Figure 1). RNase L is found in a latent state in healthy cells and can be rapidly activated during virus infection.\(^{19}\) The RNase L pathway is initiated by a group of enzymes known as 2′-5′-oligoadenylate synthetases (OAS) that can bind to dsRNA generated during viral infection. Once activated by dsRNA, these enzymes catalyze the polymerization of cellular ATP to generate 2′-5′-oligoadenylates (2-5A) that act as second messengers.\(^{52}\) RNase L directly binds to 2-5A, which induces its dimerization and activation.\(^{53,54}\) Active RNase L will degrade single-stranded RNAs within
infected cells to generate small RNAs of varied sizes that contain 5'-hydroxyl and 2',3'-cyclic phosphoryl group characteristic of metal ion-independent ribonucleases. Although both cellular and viral RNAs can be targeted by RNase L, there is evidence that the enzyme has some capacity to selectively target viral molecules. In addition to its direct antiviral activity, RNase L products may also work in a positive feedback loop where the RNA helicase RIG-I is activated by small RNA products of RNase L degradation. The RNase L system plays an important role in the antiviral response against several viruses in mammals.

WHAT WE CAN LEARN FROM vsRNAs

Different RNA surveillance mechanisms contribute to the degradation of viral molecules and generation of vsRNAs in infected cells. Small RNAs have molecular characteristics that reflect the virus and host pathways from which they originated. Thus, the pattern of vsRNAs can be analyzed and provide extensive information about their source.

Reconstruction of Viral Sequences From vsRNAs

Small RNAs derived from viral transcripts and genomes can be used to reconstitute the sequences from which they originate. Thus, the presence of viral sequences in small RNA libraries prepared from an organism provides evidence for the presence of viruses. However, owing to their limited size, vsRNAs require prior assembly into longer contiguous sequences (contigs) before they can be identified by comparisons to known viral sequences present in reference databases (Figure 2). This strategy has been successfully applied to identification and characterization of viruses in plants, fungi, and animals. Despite these successful efforts using small RNAs, sequencing of long RNAs has been a preferred strategy in studies aiming at identification of viruses within infected hosts. As large-scale sequencing of small and long RNAs have both been used to identify viruses within infected hosts, our group has directly compared strategies for virus identification. In Aedes aegypti mosquitoes, we observed that the identification of viruses by small RNA sequencing performed significantly better than long RNAs. Small RNAs provided more sequence coverage of the viral genome in less processing time. Efficient generation of small RNAs by RNAi pathways has been proposed as the main reason for the optimized detection of viruses by small RNA sequencing. Indeed, we observed that small RNA products from siRNA and piRNA pathways tended to optimize the assembly of longer viral contigs likely due to extensive overlap between sequences. Viral sequences also showed a 10-fold natural enrichment in the small RNA fraction compared to long RNAs in the same mosquito sample. Interestingly, small RNAs from lungs of coronavirus-infected mice, where no activation of RNAi pathways was observed, showed a 100-fold depletion of viral sequences relative to long RNAs. Nevertheless, even in this situation, viral contigs represented 30% of contigs assembled from small RNAs compared to 5% of long RNAs. Thus, small RNAs seem to favor assembly of viral contigs even when they represent a small fraction of the total. This can help to save computational time used for the analysis of large-scale sequencing data that can be the bottleneck for many metagenomic studies. Thus, some advantages of virus detection using small RNAs compared to long RNAs are not restricted to situations where the RNAi pathway is activated.

Construction and sequencing of small RNA libraries requires little sample manipulation and processing steps, which can be simpler compared to long RNAs. Long RNA libraries often require extensive depletion of ribosomal RNAs that may otherwise represent ~80% of all RNA species within an organism. Long RNA libraries may also utilize polyA-enriched RNA without the need for prior depletion of ribosomal RNA but this can compromise the detection of viruses whose RNAs are not polyadenylated. Notably, some insects, such as Drosophila melanogaster and Lutzomyia longipalpis, encode an abundant small RNA corresponding to the 2S ribosomal RNA that requires depletion before preparation of small RNA libraries. Even in this case, this corresponds to a single sequence that can be depleted with a single complementary probe more efficiently than large ribosomal RNA subunits that require multiple probes. It is noteworthy that the preparation of small RNAs libraries directly from total RNA could interfere with the detection of viruses. Hence, sequencing of small RNAs allows efficient detection of viral sequences although more studies are required to determine the power and extent of its application.

Virus Discovery Using vsRNA Patterns

Different host pathways generate vsRNAs with unique molecular characteristics such as size, polarity, and base enrichment. The siRNA pathway is
likely the best example on how molecular patterns of small RNAs can be explored. Viruses often produce dsRNA molecules during their replication cycle, which is considered a hallmark of viral infection (Figure 1). In most eukaryotes, the siRNA pathway is activated when Dicer directly recognizes and processes viral dsRNA into 20–23 nt long duplex small RNAs. These siRNAs usually do not have strong nucleotide preferences and cover symmetrically both strands of the dsRNA trigger. Canonical siRNAs also have 3’ 2-nt overhangs and ~19–21 nt overlap between the two strands. Thus, these molecular characteristics are a signature of activation of the siRNA pathway that has been consistently detected for a variety of viruses in fungi, animals, and plants (Figure 3(a)). As the activation of the siRNA pathway is so strongly associated with viral infection, the detection of small RNAs containing molecular patterns consistent with siRNAs could indicate a viral origin (Figure 2(b)). We and others have used this premise to discover novel viruses based on the ability of their sequences to generate siRNAs. Importantly, this strategy has enabled the discovery of divergent viral sequences with no similarities to known viruses in reference databases. The detection of sequences that generate siRNAs can be an efficient strategy to find viruses although there are potential limitations that can affect its sensitivity and specificity.
Activation of the small interfering RNA (siRNA) pathway is a common and specific response to virus infection. (a) Virus-derived small RNAs in different animals often show a profile consistent with the activation of the siRNA pathway. Virus-derived siRNAs range from ~20 to 23 nt and are symmetrical in polarity and base preferences. (b) Endogenous viral elements (EVEs) represent remnants of virus sequences integrated into animal genomes and also generate small RNAs. EVE-derived small RNAs are often ~24–30 nt, asymmetrical in polarity, and base preferences that is consistent with a piRNA signature. (c) Size, polarity, and nucleotide preferences were determined for small RNAs derived from EVEs and active viruses. These molecular footprints were then used to perform hierarchical clustering of different EVEs and viruses. The small RNA pattern clearly separates clusters containing viruses (in red) and EVEs (in black). Notably, within the cluster of viruses, we observe two small subclusters. Virus grouped in the larger subcluster show a classical siRNA signature even when the pathway is partially inhibited such as for FHV and DCV in Drosophila. The second subcluster contains the vsRNA profile of coronavirus in mouse lungs and PCLV in mosquitoes, both of which show a more divergent profile from the siRNA signature. The data in this figure were obtained from the analysis of small RNA libraries from published studies (accession numbers: SRR1803378, SRR1803382, ERR555100, ERR274423, ERR654010, SRR1803383, GSM792688, GSM792692, SRR452408, and SRR640612).
First, there are endogenous sources of dsRNA capable of generating siRNAs. Endogenous siRNAs arise from regions of convergent transcription, structured RNAs, or repetitive elements. In each of these cases, it would not be difficult to ascertain the origin of siRNAs because these are easily differentiated from viruses. Convergent transcription and structured RNAs arise from annotated genes and repetitive elements can be identified using repeat filters. However, animal genomes have also integrated endogenous viral elements (EVEs) that represent remnants of viral sequences. We analyzed the small RNA profile observed for EVEs in different organisms compared to active viruses. EVE-derived small RNAs were observed but show clearly different molecular patterns compared to viruses in the same organism (Figure 3). Interestingly, EVEs seemed to favor the generation of small RNAs with molecular characteristics of piRNAs rather than siRNAs.

Second, some viruses have developed viral suppressors of the siRNA pathway (VSRs) that can significantly affect the pattern of vsRNAs. VSRs allow for accumulation of viral RNA that can be degraded by other host mechanisms. Patterns of vsRNA distinct from canonical siRNAs have been observed upon suppression of the siRNA pathway by VSRs. For example, Drosophila adults infected with Flock house virus (FHV) show accumulation of small RNAs in the positive strand with a broad size distribution and only a very small peak at 21 nt in the negative strand (Figure 3(a)). FHV encodes a potent VSR known as B2 that successfully protects its dsRNAs from access by Dicer-2 and prevents activation of the siRNA pathway. The viral RNA genome accumulates at higher levels than the antigenome and is likely degraded by other nucleases, which explains the bias observed in small RNAs covering the positive strand and the broad size profile. Accordingly, in the absence of B2, the pattern of small RNAs observed in FHV-infected flies is consistent with the activation of the siRNA pathway (Figure 3(a)).

Lastly, technical artifacts from small RNA sequencing strategies could interfere with the detection of siRNA signatures. Biased cloning due to inefficient adaptor ligation or sequestration of antisense small RNAs by the sense target can result in a distortion of vsRNA profile. However, it is unclear how much the detection of a canonical siRNA profile can be compromised by technical biases. In arthropods, for example, a canonical siRNA profile is observed more often than not, even using standard strategies for small RNA library construction (Figure 3(a)).

Hence, small RNA sequencing is a promising strategy to identify viruses, despite potential limitations that require further analysis.

**Using vsRNA Patterns to Extract Information About the Infection**

The pattern of vsRNAs is a molecular footprint that can be explored beyond the activation of the siRNA pathway. Each RNA surveillance mechanisms may show cell-type- or tissue-specific expression and differentially target viral RNA intermediates. Thus, virus tropism and the abundance of different viral RNA targets will have a direct influence on the pattern of small RNAs produced by host pathways. Consequently, the pattern of vsRNAs generated during infection can provide information about their origin, which reflects both virus and host features (Figure 4).

Some viruses have developed VSRs to escape the siRNA pathway as we described for the B2 protein of FHV. Interestingly, another insect virus, Drosophila C virus (DCV), encodes a VSR that binds long dsRNA and prevents Dcr-2-mediated processing. The profile of vsRNAs observed in DCV-infected flies shows a peak in 21 nt long coming from both strands but also accumulation of small RNAs with smaller sizes derived from the coding strand. The VSR encoded by DCV is not as potent as B2 and presumably still allows detectable activation of the siRNA pathway. This situation generates superimposed vsRNAs pattern combining products of different host pathways that could still allow virus identification based on a canonical siRNA pattern. Nevertheless, it is interesting to speculate that once a virus is identified, the absence of virus-derived siRNAs in infected hosts may be interpreted as active inhibition by VSRs as shown for FHV (Figure 4(a)). In addition, the pattern of vsRNAs may also provide insights into the mechanism of action and potency of the VSRs considering the differences we observed between the small RNA profile generated by DCV and FHV (Figure 3(a)).

RNAi mechanisms other than the siRNA pathway could also be explored. The piRNA pathway, for example, is not broadly activated during viral infections. At the moment, virus-derived piRNAs have only been observed in insect cell lines including Drosophila ovary, culicoides, and mosquito cell lines. In vivo, virus-derived piRNAs have only been observed in A. albopticus and A. aegypti mosquitoes. Molecular characteristics of piRNAs are quite distinct from siRNAs, as the former are ~24–30 nt long and show enrichment for U at the 5’
of antisense RNAs and A in 10th position of sense RNAs. piRNAs in opposite strands may also show an overlap of 10 nt between their 5' ends, which is signature of the ping-pong amplification mechanism. In terms of their origin, piRNAs may show asymmetrical coverage of the viral genome or more homogenous distribution similar to siRNAs. It is unclear why the activation of the piRNA pathway is restricted to certain viruses and how it is initiated during viral infection. Indeed, viruses that generate virus-derived piRNAs have RNA genomes and replicate in the cytoplasm while the endogenous piRNA pathway is initiated from DNA-dependent transcripts in the nucleus, at least in Drosophila. Our own data suggest that a strong activation of the piRNA pathway requires that the virus infects reproductive tissues where components of this pathway are commonly enriched in insects. We observed that the same A. aegypti mosquitoes infected by two different viruses, Phasi Charoen like-virus (PCLV) and Humaita-Tubiacanga virus (HTV), only the former showed abundant production of piRNAs which correlated with ovary infection (Figure 3(a)). In contrast, A. albopictus mosquitoes infected with Chikungunya virus showed production of virus-derived piRNAs in the head and
Virus-derived small RNAs can also be observed during infection but are usually produced by the virus to regulate the expression of viral or cellular genes. miRNAs are distinct from siRNAs and piRNAs because they originate uniquely from specific regions of the virus genome and are restricted to just one of the strands. In terms of molecular features, miRNAs are ~20–24 nt long with a strong enrichment for U at 5’ end. The production of virus-derived miRNAs has mostly been reported for animal DNA viruses. This means that the generation of miRNAs is a very broad distribution as this endonuclease targets mostly been reported for animal DNA viruses. Thus, in a somewhat more limited manner, the detection of these molecular patterns arising from regions of the viral genome could be used to infer whether a virus is capable of generating its own miRNAs (Figure 4(c)).

Non-RNAi pathways, such as RNA decay and RNase L, also leave molecular footprints on vsRNAs that could be potentially explored. Indeed, these nucleases have substrate specificities that generate nonrandom patterns. Ribonucleases from RNA decay pathways such as XRN1 and the exosome attack from their substrates from the extremities, which tends to reduce the RNA to a few nucleotides. However, these exonucleases can be impaired by highly structured regions or internal modifications within the target. As a result, RNA decay nucleases often produce vsRNAs of different sizes that tend to accumulate near the end of the substrate or close to regions with secondary structure. In the case of RNase L, small RNA products also have a very broad distribution as this endonuclease targets U-rich sequences whose abundance may vary within target RNAs. In addition, RNA decay and RNase L pathways may act together to generate complex patterns of vsRNAs in infected cells. In human embryonic kidney 293 cells and green monkey Vero cells, the size profile of small RNAs derived from *Sindbis virus* was remarkably consistent even in the absence of detectable activation of RNAi pathways. RNase L was required for the generation of some vsRNAs although they did not seem to be direct products of this nuclease. Indeed, the generation of Sindbis vsRNAs required the joint action of RNase L and other nucleases such as XRN1. In this case, vsRNAs seemed to accumulate in regions of the Sindbis genome containing posttranscriptional modifications that inhibited further degradation. Similarly, in human HeLa cells, the degradation of the poliovirus RNA is further targeted by RNase L together with other nucleases, which resulted in a complex pattern of vsRNAs generated across the virus genome. Thus, small RNAs generated by RNA decay and RNase L seem to reflect structured or modified regions within viral RNAs (Figure 4(d)). Indeed, the consistency of vsRNA profiles observed in different mammalian cells even without activation of RNAi suggests that the pattern does reflect stable virus characteristics.

Hence, complex vsRNA patterns can be very informative even when they result from the combined action of different nucleases. We are only beginning to understand how to decipher this information.

**CONCLUSIONS**

In the arms race between host and viruses, different pathways have evolved to target viral RNAs. These pathways contribute to the antiviral response by degrading viral RNAs, which results in the generation of vsRNAs. This common targeting of viral RNAs helps to explain why viral sequences are often abundant within the pool of small RNAs in infected cells. In addition, these tiny molecules are imprinted with molecular characteristics that reflect the host pathway and viral RNA from which they originate. Indeed, information can be extracted from these molecular footprints to trace back their viral origin. Directly, small RNA from infected hosts can be assembled into longer contiguous sequences and used to detect viruses by homology searches against known viral references. Indirectly, molecular patterns of small RNAs that are consistent with the activation of the siRNA pathway can be used as a signature to suggest a viral origin to novel assembled sequences. This latter strategy can help overcome a great limitation of virus discovery by metagenomic strategies, because it does not require sequence similarity searches against known references. Pattern analysis of vsRNAs allows the discovery of more divergent viral sequences with no similarities to known viruses present in reference databases. We also showed here that the pattern of small RNAs is able to differentiate between sequences derived from viruses and integrated EVEs, which can be a powerful tool to
indicate whether viral sequences originate from an active infection. Furthermore, the pattern of vsRNAs generated by host mechanisms other than the siRNA pathway can also provide information about viruses such as their tissue tropism, coding of inhibitors to the siRNA pathway, and structured regions of the viral genome. Although it might not be universally applicable, these advantages make small RNA sequencing a great strategy to identify and characterize viruses.

The potential use of vsRNA patterns is just beginning to be unlocked. We have recently showed that it is possible to use patterns of small RNAs different from the canonical siRNA signature to detect novel viral sequences. As a general idea, if the profile of small RNAs is known for any segment of a viral genome, this can be used as a reference to find other potential sequences from the same virus that show a similar pattern. This broadens the diversity of vsRNA patterns that can be utilized beyond the dependence on activation of the siRNA pathway. The use of additional tools could significantly improve our capacity to explore small RNAs patterns. Small RNAs generated by RNA decay, RNAi, and RNase L pathways have clear molecular features that can be utilized to differentiate specific products. Indeed, unique molecular features found in small RNAs generated by RNAi pathways such as 5′ monophosphate or 2′O-methylation have been extensively explored. More recently, characteristics generated by RNase L have also been explored to identify specific products of this nuclease in infected cells. These strategies allow more specific analysis of the products from each pathway and can be combined to provide a broader view of small RNAs within infected cells. This expansion in known vsRNA patterns would certainly broaden our references for pattern searches and improve our ability to identify and characterize viruses.

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