Profile of small interfering RNAs from cotton plants infected with the polerovirus Cotton leafroll dwarf virus

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Abstract

Background: In response to infection, viral genomes are processed by Dicer-like (DCL) ribonuclease proteins into viral small RNAs (vsRNAs) of discrete sizes. vsRNAs are then used as guides for silencing the viral genome. The profile of vsRNAs produced during the infection process has been extensively studied for some groups of viruses. However, nothing is known about the vsRNAs produced during infections of members of the economically important family Luteoviridae, a group of phloem-restricted viruses. Here, we report the characterization of a population of vsRNAs from cotton plants infected with Cotton leafroll dwarf virus (CLRDV), a member of the genus Polerovirus, family Luteoviridae.

Results: Deep sequencing of small RNAs (sRNAs) from leaves of CLRDV-infected cotton plants revealed that the vsRNAs were 21- to 24-nucleotides (nt) long and that their sequences matched the viral genome, with higher frequencies of matches in the 3′-region. There were equivalent amounts of sense and antisense vsRNAs, and the 22-nt class of small RNAs was predominant. During infection, cotton Dcl transcripts appeared to be up-regulated, while Dcl2 appeared to be down-regulated.

Conclusions: This is the first report on the profile of sRNAs in a plant infected with a virus from the family Luteoviridae. Our sequence data strongly suggest that virus-derived double-stranded RNA functions as one of the main precursors of vsRNAs. Judging by the profiled size classes, all cotton DCLs might be working to silence the virus. The possible causes for the unexpectedly high accumulation of 22-nt vsRNAs are discussed. CLRDV is the causal agent of Cotton blue disease, which occurs worldwide. Our results are an important contribution for understanding the molecular mechanisms involved in this and related diseases.

Background

The RNA silencing pathway controls important biological processes in plants, including regulation of gene expression during development, heterochromatin formation, hormone signaling, metabolic processes, and stress responses, as well as being an important antiviral defense mechanism [1]. In plants, antiviral silencing can be triggered by the presence of viral double-stranded RNAs (dsRNA), which are generated by the viral RNA polymerase as an intermediate in genomic replication and transcription, or are predicted to form as secondary structures along single stranded viral genomic RNA (ssRNA) [2]. Both structures are recognized by Dicer-like (DCL) ribonucleases and are processed into virus-derived small interfering RNAs (vsRNAs) that vary in length from 21 to 24 nucleotides (nt) [3-5]. These vsRNAs are then loaded into Argonaute (AGO)-containing complexes known as RNA-induced silencing complexes (RISCs), which promote the degradation of both genomic and subgenomic viral RNAs [6,7].

DCL ribonucleases are present in both monocot and dicot plants. Arabidopsis thaliana contains four DCLs (AtDCLs1-4) [8], while the Populus and rice genomes encode five and six DCLs, respectively [9]. The diversity associated with Dicer ribonucleases, as well as other silencing-related proteins such as AGO, strongly suggest that several silencing pathways have evolved in plants.
Correspondingly, in Arabidopsis, at least six silencing pathways have been identified, and the four DCLs involved are known to act hierarchically. For example, there are 21-nt vsRNAs and other small RNAs (sRNAs) associated with post-transcriptional silencing of endogenous genes generated by AtDCL4. In the absence of AtDCL4, 22-nt vsRNAs are produced by AtDCL2, and in the absence of both AtDCL4 and AtDCL2, 24-nt vsRNAs are produced by AtDCL3 [10-12]. Thus, AtDCL2-4 play essential roles in mediating the antiviral defenses of Arabidopsis. In contrast, AtDCL1 is mainly associated with the production of regulatory microRNAs derived from hairpin-like endogenous transcripts [13].

All of the four Dicer proteins expressed in Arabidopsis are usually present in other plants, also [14] (Additional file 1, figure s1). Correspondingly, 21-, 22-, and 24-nt vsRNAs have been detected in many plant hosts following infection [15]. However, based on the hierarchical roles of DCL4 and DCL2 in antiviral silencing, 21-nt vsRNAs are by far the most abundant class of sRNA found in plants infected with RNA or DNA viruses, followed by 22-nt vsRNAs [15-18]. Previous studies have shown that the accumulation of vsRNAs is affected by viral suppressors of gene silencing [11,19].

Suppressor proteins can directly bind vsRNAs [20-23], or inhibit key proteins of the gene-silencing pathway [24-26]. For example, the Polerovirus P0 protein and the P38 protein from Turnip crinkle virus (TCV) targetAGO1, an important antiviral Argonaute protein [11,24,27-30]. Studies have shown that P0 preferentially targets AGO1, leading to its degradation, but does not affect the sRNA-RISC complex [26]. A similar action has been suggested for the P38 protein, which binds to AGO1 and may prevent the assembly of RISC. However, unlike P0, P38 does not affect the stability of AGO1 [30]. By preventing the association of AGO1 with RISC, P38 has the potential to destabilize a complex homeostatic network involving AGO1, microRNAs, and the four Dicer proteins. This would be consistent with the preferential accumulation of 22-nt vsRNAs observed following infection with TCV [30].

In this study, vsRNAs derived from cotton plants (Gossypium hirsutum) infected with Cotton leafroll dwarf virus (CLRDV) (genus, Polerovirus; family, Luteoviridae) were deep-sequenced and characterized. CLRDV is transmitted by the aphid, Aphis gossypii, and is the causal agent of cotton blue disease [31], which occurs in cotton crops world-wide. Consistent with other members of the same family, CLRDV is phloem-restricted and its genome consists of a single strand, positive sense, non-polyadenylated RNA (5.8 kb) containing six open reading frames (ORFs) [32]. This is the first report of vsRNAs derived from a member of the family Luteoviridae and the first report of vsRNAs in cotton plants.

**Results**

**Characterization of CLRDV-derived sRNAs**

To characterize the vsRNAs produced during CLRDV infection, sRNAs obtained from cotton-infected and uninfected plants were cloned and deep-sequenced using the Illumina platform. A total of 10,566,377 and 9,480,917 reads were sequenced from systemic leaves harvested at 5 dpi from infected and uninfected plants, respectively (Figure 1A). Reads ranging from 18 to 26 nt were mapped in sense and antisense orientations to the viral genome. Only sequences showing no mismatches were regarded as CLRDV vsRNAs in the infected library. In total, 640,325 viral-derived sRNA reads were identified, covering almost the entire sequence of the genome. In the uninfected library, only 1,967 reads matched with the CLRDV genome (corresponding to 0.025% of the 18-26 nt reads sequenced). Of the vsRNA reads identified in infected plants, 51,607 were unique (Figure 1A). Equivalent amounts of sense and antisense vsRNAs were found in the CLRDV-infected cotton library, suggesting that vsRNAs derived from the cleavage of dsRNA are processed by Dicer ribonucleases (Figure 1B). To further characterize the vsRNA library, the frequencies of redundant and unique CLRDV-derived sRNAs ranging from 18 to 26 nt were analyzed (Figure 1). In both sets of analyses, 22-nt vsRNAs were the most abundant. For example, 22-nt redundant vsRNAs (Figure 1C) represented 36.61% of the total vsRNAs sequenced, followed by 23-nt and 21-nt vsRNAs (21.22% and 15.53%, respectively). For the unique vsRNAs (Figure 1D), 22-nt represented 17.74% of the total vsRNAs sequenced, followed by 21-nt and 23-nt vsRNAs (15.42% and 15.23%, respectively). These data were confirmed through the deep sequencing, in an independent channel, of an additional cotton-infected RNA sample (Additional file 2 figure s2), generating 15,415,107 reads, of which 498,367 matched with the CLRDV genome. Together, these results suggest that the cotton homologue of DCL2 (GhDCL2) may be the predominant Dicer ribonuclease involved in their biogenesis.

To determine whether the 22-nt reads are predominant only in the pool of vsRNAs or whether this is the dominant class among small RNAs of cotton, we compared the overall profile of small RNAs between the infected and the uninfected libraries. In both libraries there was an abundance of the 24-nt class among the endogenous sRNAs, followed by the 21-nt class (Figure 2A and 2B). However, in infected plants, there were decreased levels of the 24- and 21-nt classes (Figure 2B) indicating that CLRDV infection may decrease the production of endogenous sRNAs.
The high accumulation of the endogenous 24-nt sRNAs, followed by 21-nt sRNAs, are consistent with sRNA profiles in other plants (33,34). These findings indicate that the cotton RNA silencing machinery responsible for biogenesis of endogenous or viral sRNAs does not tend to produce 22-nt sequences. Therefore, the high levels of 22-nt CLRDV-vsRNA seem to be a result of the antiviral RNA silencing mechanism or a specific CLRDV-host interaction.

In *Arabidopsis*, the 5- terminal nucleotide partially determines the preference of sRNAs for AGO proteins. Therefore, the distribution of 5- terminal nucleotides was determined for the sequenced vsRNAs (Figure 3). For all three types of CLRDV-vsRNAs characterized (i.e., 21-, 22- and 23-nt), cytosine was the most commonly occurring nucleotide at the 5- terminus (32.09%, 42.27%, and 36.65%, respectively), while guanine was the least common (14.98%, 9.62%, and 8.12%, respectively). In contrast,
24-nucleotide CLRDV-vsRNAs often had adenine at the 5'-terminus (53.24%), or guanine (23.26%). For comparison, Arabidopsis AGO1 has a 5'-nucleotide preference for uracil, AGO2 and AGO4 have a preference for adenine, and AGO5 preferentially loads sRNAs with cytosine at the 5'-terminus [33-35]. Therefore, the results of the present study suggest that CLRDV-vsRNAs can be potentially loaded by multiple AGO-containing complexes.

![Figure 2 Size distribution of CLRDV-vsRNA and endogenous sRNAs](image1)

Figure 2 Size distribution of CLRDV-vsRNA and endogenous sRNAs. Histograms compare the distribution of 18-26-nt total sRNA reads with vsRNAs obtained from uninfected (A) and CLRDV-infected (B) cotton plants. Only 0.025% of total sRNA obtained from uninfected plant matched to the CLRDV genome (data not shown).

![Figure 3 Characterization of vsRNAs 5'-terminal nucleotide](image2)

Figure 3 Characterization of vsRNAs 5'-terminal nucleotide. Relative abundance of four distinct 5'-terminal nucleotides in 21-24-nt vsRNAs in CLRDV-infected plant.
However, 21-23 nt CLRDV-viRNAs may be preferentially loaded by AGO5, while 24-nt CLRDV-vsRNAs would be loaded by AGO2 and/or AGO4.

**Distribution of vsRNA abundance**

To examine the spatial distribution and sequence diversity of the viral sRNAs identified, redundant (Figure 4B) and unique (Figure 4C) 21-24 nt vsRNA sequences were mapped to the CLRDV genome. For both classes of sequences, the distribution of CLRDV-vsRNAs along the genome was non-uniform, with most of the vsRNAs accumulating in the 3- region of the genome. This part of the genome encodes structural proteins and proteins that assemble subgenomic RNAs during infection [31]. The greatest numbers of redundant reads were associated with position 5049-5070, with 398,897 reads identified. Overall, this region of the genome was highly represented. In addition, a large number of reads mapped to ORF5, a region encoding an aphid-transmission protein. However, after the unique vsRNAs were aligned with the genome (Figure 4C), there were regions associated with an absence of vsRNA reads. When the same alignment was performed and up to two mismatches were allowed, only positions 4150-4170 did not match with any vsRNA reads (data not shown). Therefore, it is possible that this region may have a structural characteristic that makes it less accessible to Dicer ribonucleases, although sequencing artifacts cannot be ruled out. Furthermore, the hot-spots for vsRNAs that were found to be associated with the 3- region of the CLRDV genome might be generated by the massive accumulation of viral subgenomic RNAs (sgRNA) observed in leaves [15,36].

An analysis of the unique reads that mapped to the CLRDV genome indicated that all Dicer ribonucleases were able to access the entire viral genome. Similar amounts of 21-24 nt vsRNAs corresponding to sense and antisense strands of viral RNA (Figure 4C) were present in our library, reinforcing that virus-derived dsRNAs are the main source of vsRNAs. Furthermore, peaks of both the abundance (Figure 4B) and diversity (Figure 4C) of 21-24 nt reads showed similar patterns of distribution along the genome. These results suggest that all DCL ribonucleases contribute to the generation of vsRNAs with similar substrate affinities and target the same regions of the genome. However, the fact that the 22-nt class of vsRNAs was the most prominent class supports the hypothesis that GhDCL2 may play a role in the generation of CLRDV-vsRNA.

**Expression of Cotton DCL ribonucleases during infection**

Assays of TCV infection have detected a high abundance of 22-nt vsRNAs [11,19,30] associated with the silencing suppressor protein, P38. During infection, P38 inactivates AGO1 by down-regulating miR162. As a result, low levels of miR162 directly and/or indirectly affect transcript levels of Dcl1, 3, and 4 [30]. To determine whether a similar mechanism might be activated in cotton-infected plants, we analyzed the expression levels of mature miR162 and Dcls in CLRDV-infected and uninfected plants.

In contrast with TCV infection of Arabidopsis, qPCR experiments detected almost three-fold higher levels of Gh-miR162 in infected versus uninfected cotton plants (Figure 5A). Furthermore, *in silico* analysis of miRNAs in the deep-sequencing libraries from infected versus uninfected plants showed similar results, with Gh-miR162 levels slightly up-regulated during CLRDV infection (data not shown). The levels of mRNAs for cotton DCLs were assayed, and there were no significant differences in the levels of Dcl1 (GhDcl1) and Dcl3 (GhDcl3) transcripts (Figure 5B) between uninfected and infected plants. In contrast, GhDcl4 was up-regulated during infection, while GhDcl2 was down-regulated (Figure 5B). Taken together, these results suggest that the high levels of 22-nt vsRNAs produced in CLRDV-infected plants may represent a mechanism distinct from that previously described for TCV infections.

**Discussion**

This is the first report of the characterization of small RNAs produced from a member of the genus *Polerovirus*, family *Luteoviridae*. The profile of vsRNAs generated in cotton plants infected with CLRDV revealed some interesting features regarding their biogenesis. For example, both sense and antisense orientations of CLRDV-derived sRNAs accumulated to similar levels (Figure 1B). However, several other studies have found that sense vsRNAs accumulate to higher levels in some hosts [16,17,23]. In those cases, strand biases are usually attributed to preferential processing of highly structured single-stranded genomic viral RNAs by Dicer ribonucleases [15,23]. Despite these differences and considerable experimental efforts, however, the existence of a direct correlation between vsRNA hot-spots and structured regions of genomic viral RNAs has never been proven [15]. The accumulation of equivalent amounts of sense and antisense CLRDV-vsRNAs observed in the present study supports the hypothesis that CLRDV-dsRNAs, which are generated by viral RNA polymerases during genome replication or by the activity of host RNA-dependent RNA polymerases [2], are the main substrates for Dicer ribonucleases. Since the P0 silencing suppressor protein from *Polerovirus* was already shown to inhibit production of secondary vsRNAs in 35S-promoter-driven agroinfiltration assays [27,28,37], it may be speculated that the main substrate of cotton DCLs during CLRDV infection is probably the replicative
Figure 4 Genome view of CLRDV-derived sRNAs recovered from infected cotton plants. (A) Schematic representation of CLRDV genome. ORF0 = silencing suppressor protein; ORF1 and ORF2 = virus RNA polymerase; ORF3 = capsid protein; ORF4 = movement protein; ORF3 and ORF5 = readthrough transmission protein. Distribution of total (B) and unique (C) vsRNAs along viral genome, in either sense (above x axis) or antisense (below x axis) configuration. Figure 4B was scaled to 10000 reads to facilitate the visualization of the obtained data, thus, peaks that eventually were major than 10000 reads were indicated by asterisks.
intermediate forms of viral genomic RNAs. However, the mechanism of P0 protein action in the formation of the secondary siRNAs during virus infection remains unclear.

Overall, the distribution profile of CLRDV-vsRNAs within the genome varied considerably. High vsRNA densities were identified in regions coding for structural proteins, especially in the ORF5 region (Figure 4). Previously, it was shown that genes encoding structural proteins in the family Luteoviridae are expressed from subgenomic RNAs (i.e., sgRNA1 and sgRNA2) [38]. Moreover, studies of the Polerovirus, Potato leafroll virus (PLRV), also identified two sgRNAs associated with the 3'-block of the viral genome [39]. The transcription of sgRNA1 provides for expression of ORFs 3, 4, and 5, while that of sgRNA2 (~800 nt) encodes two proteins located within the 3'-proximal half of ORF5. Since sgRNAs are highly expressed during the infection cycle, an over-accumulation of vsRNAs derived from this region of the genome might be due to a greater availability of dsRNA intermediate templates for processing. Accordingly, the hot-spot of vsRNAs mapped to ORF5 might be due to the expression of sgRNA2, which is also derived from this region of the genome. Although the synthesis of sgRNA2 by CLRDV has not previously been reported, the ACAAAA motif present at the 5'-end of sgRNA1 and sgRNA2 from other Poleroviruses [40] is also present in the ORF5 of CLRDV (position 4821-4828) (data not shown). Based on these results, it is possible that sgRNA2 is produced by CLRDV.

Depending on their length and 5'-identity, sRNAs are selectively loaded into multiple AGO complexes [33,34]. Previous studies have shown that plant virus-specific sRNAs beginning with uracil or adenine are preferentially loaded into AGO1, AGO2, and AGO4 [15-17]. In fact, AGO1 and AGO2 are required for the anti-viral silencing pathway in Arabidopsis [41-43]. However, 21-23 nt CLRDV-vsRNAs usually have a cytosine at the 5'-terminal position (Figure 3), indicating that they may be loaded into a cotton homologue of AGO5. Although the AtAGO5 has no detectable anti-viral function against Cucumber mosaic virus (CMV) [8,43,44], CMV-vsRNAs have been detected in AtAGO5 immunoprecipitates, indicating that the protein may act in the biogenesis of secondary vsRNAs [35]. Moreover, a predominance of 5'-terminal cytosines has been observed for some viroid-derived sRNAs [45]. In contrast, most 24-nt CLRDV-vsRNAs have adenine at the 5'-terminal (Figure 3), indicating that they can be loaded into cotton AGO2 and AGO4 homologues. In Arabidopsis, the association between 24-nt sRNAs and AGO4 has been well-characterized as a mediator of transcriptional silencing for transposons and repeated sequences [41]. In addition, the decreased number of vsRNAs that start with guanine is correlated with the absence of AGO proteins that might otherwise have an affinity for those sRNAs.

The balance between antiviral silencing and suppression mechanisms can directly influence the accumulation of vsRNAs within infected plants. While the functions of the four DCL proteins present in Arabidopsis are well characterized, Dicer ribonucleases from other species, including cotton, remain largely unstudied. However, if the mechanism(s) associated with DCL ribonucleases is conserved between cotton and Arabidopsis, then the

Figure 5 Quantitative RT-PCR analysis. Expression levels of Gh-miR162 (A) and cotton Dcl mRNAs (B) in infected and uninfected plants. Relative expression levels were estimated using polyubiquitin and catalytic subunit of protein phosphatase 2A genes as internal controls. Error bars indicate standard deviations. P values were calculated by t-tests with a significance level of P < 0.05. Asterisks indicate significant differences.

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predominance of 22-nt vsRNAs associated with CLRDV infection would be hypothesized to be the result of GhDCL2 activity. Although 22-nt CMV-vsRNAs produced by AtDCL2 are poor effectors of antiviral defense in Arabidopsis [43], other studies have detected a predominant population of 22-nt vsRNAs following infection with certain plant viruses and viroids [11,15,19,36,45]. For example, Cymbidium ring spot virus (CymRSV) and TCV infections are associated with an abundance of 22-nt vsRNAs, which seem to be related to the activity of the suppressor proteins P19 and P38 [11,15,19]. P19 can specifically sequester 21-nt duplex sRNAs [46], while P38 can indirectly block AtDCL4 activity by suppressing AGO1 function [30]. During TCV infection in Arabidopsis, AtDCL1 levels are indirectly increased due to the P38-mediated down-regulation of microRNAs, including miR162, a negative regulator of AtDcl1 transcripts [47]. Since AtDCL1 negatively regulates AtDcl4 and AtDcl3 [48], over-accumulation of AtDCL1 generates a deficit in the levels of AtDCL4 and AtDCL3, leaving dsRNAs more accessible to AtDCL2 [30]. The Polerovirus P0 suppressor protein is also able to destabilize AGO1 [24,27,28]. Although the activity of CLRDV P0 has not yet been tested, the F-Box-like domain necessary for silencing that is conserved among P0 sequences from other members of the genus is also conserved in CLRDV (data not shown). Thus, CLRDV P0 has the potential to similarly affect cotton Dicer ribonucleases during the infection process. However, in this study, there were no significant changes in the levels of GhDcl1, GhDcl2, and GhDcl3 transcripts in infected plants (Figure 5B). Furthermore, GhDcl4 transcripts and Gh-miR162 were up-regulated (Figure 5A and 5B). The up-regulation of Dcl4 has been observed in other viral infections [49], but the levels of mature miR162 are inconsistent with what was observed during TCV infection [30]. It is possible that differences in tissue tropism between TCV and CLRDV, and/or differences in the silencing machinery of the host, account for the observed differences between the two viruses.

Members of the genus Polerovirus are restricted to the phloem cells of their hosts. Therefore, DCL activity in response to viral dsRNA may be cell-type dependent. Small RNAs derived from Hop stunt viroid (HSVd) infections in cucumber plants showed different sizes in different tissues [45]. For example, most of the sRNAs from infected whole leaves were 21-nt long, while those derived from phloem-sap were more frequently 22 nt in length. Although transgene-induced silencing in phloem cells of Arabidopsis is triggered by AtDCL4 [50], a difference in the affinity or expression of Dicer ribonucleases, or other silencing-related proteins such as dsRNA-binding proteins in companion cells, could possibly explain the tissue-dependent shift in sRNA size.

The production of vsRNAs following virus infection can vary depending on the host. For example, sRNAs derived from Bamboo mosaic virus are mainly 21 nt in length in Arabidopsis, but 22 nt in Nicotiana benthamiana. Therefore, these data suggest that DCL recruitment for vsRNA production is a host-dependent process [36]. This is the first report of a sRNA profile for cotton virus-infected plants. Further research is required to confirm whether the vsRNA profile observed here results from a viral silencing suppressor protein, or from factors such as phloem-restriction or cotton-specific factors that can activate an anti-viral silencing pathway.

Conclusions
This is the first high-throughput sequencing of a member of the Luteoviridae family, CLRDV, from virus-infected cotton plants. This study shows that RNA silencing systems against CLRDV result in the production of 22-nt sRNAs as the predominant sRNA size class. All vsRNAs, independently of the size, and that these are derived mainly from the 3′ region of the viral genome. The sequence data of sense and antisense vsRNAs strongly suggest that dsRNA molecules are the main source of the vsRNAs. During CLRDV infection, we observed up-regulation of GhDcl4 and down-regulation of GhDcl2 transcripts, which are the major DCLs in antiviral defense in the model plant Arabidopsis. There is still much to learn about the molecular mechanisms underlying the prevalence of the 22-nt CLRDV-vsRNAs.

Methods
Sample preparation and sequencing
Fifty-day-old cotton (Gossypium hirsutum) plants (cultivar FM966; Fibermax966) that are susceptible to cotton blue disease were infected with CLRDV using the viruliferous aphid, Aphis gossypii. Aphids were placed on older true leaves and removed 24 h after infestation. Systemic leaves (i.e., representing the youngest completely expanded leaves) were harvested 5 days post-infection (dpi). The same leaves were harvested from mock-infected plants as the control. Total RNAs were extracted from systemic leaves using the Invisorb Spin Plant RNA Mini Kit (Invisorb®).

The quantity and quality of RNA samples obtained were determined by spectrophotometry (Nanodrop ND-1000, Thermo Fisher Scientific) and agarose gel electrophoresis (Additional file 3, figure s3), respectively. Systemic infections were confirmed using nested (RT)-PCR assays to detect the viral capsid protein-encoding gene as previously described [51]. RNA samples were precipitated in ethanol and sequenced at the Fasteris Life Science Co. (Geneva, Switzerland) with an Illumina Genome Analyzer (Illumina, San Diego, USA). Small
| Gene       | Forward primer sequence (5'-3') | Reverse primer sequence (5'-3') | Amplicon size (bp) | Efficiency ± SD* | R2    | Locus accession number |
|------------|---------------------------------|---------------------------------|--------------------|------------------|-------|------------------------|
| DCL1       | AACCTGGGGTGTTGCTCCCTG           | ATGCCCTTTTGGCTTGCTC             | 132                | 0.928647 ± 0.0293724 | 0.9858 | ES804646.1             |
| DCL2       | GATGCTATCATGCTTCCGCAG           | TGGGAACCAAAGACACAGGAA          | 81                 | 0.99522 ± 0.00361129 | 0.9903 | DW488144               |
| DCL3       | ATGTCACATGCCCCCTGGAGCT         | GGCCAACATTAAGGACTCCAGCCG       | 113                | 0.997162 ± 0.0071986 | 0.9646 | DR462994               |
| DCL4       | GCTCCAAGCGGCAACAGCATT          | AGGATGCAAAATCGCCCTGAAGGAG      | 186                | 0.992603 ± 0.0051324 | 0.9706 | DT568872               |
| Gh-miR162  | GCGGGGAGCTATTGGAGACG           | GTGGGCTCTGGTGCAOGGTCGGAGTATCCGGCAT | 72                 | 0.884982 ± 0.032819 | 0.8519 | -                      |
| XTH        | GGAAAGGGTGACAGGGAAAGA         | GGCTGGAGTTTGGGTATGGG        | 173 and 392**     | -                | -     | AY180971.2             |

*Efficiency ± standard deviation (SD) generated by the Miners software. ** in DNA sample.
RNA libraries were prepared according to a modified Illumina protocol. Briefly, small RNAs of 15-30 nt were purified on an acrylamide gel; the 3- IDT miRNA cloning linker (Integrated DNA Technologies, San Diego, USA) and then the 5- Illumina adapters were single-stranded ligated with T4 RNA ligase. The constructs were purified again on an acrylamide gel to remove empty adapters and then reverse-transcribed and PCR-amplified. The primers used for cDNA synthesis and PCR were designed to insert an index in the 3- adapter. The libraries were quality controlled by cloning an aliquot into a TOPO plasmid and capillary sequencing 4-8 clones. High-throughput sequencing was performed on a Genome Analyzer GAIIx for 38 cycles plus 7 cycles to read the indexes. After demultiplexing and adapter removal, 10.5 million pass filter reads were obtained in the library.

All the deep sequencing libraries obtained are deposited at GEO (Gene Expression Omnibus) under the number GSE311062 http://www.ncbi.nlm.nih.gov/geo/info/submission.html

Data mining of the sRNA pool
CLRDV-derived sRNAs sequences were identified using a local BLAST database of the CLRDV-PV1 isolate genomic sequence (accession number HQ827780). Library characterization and mapping to the viral genome were performed using locally developed Perl scripts. Further calculations and statistical analyses were performed using R 2.7.1 software (R Foundation for Statistical Computing).

Real-time analyses
Primers used to amplify the DCLs genes and Gh-miR162 are listed in Table 1.

To measure expression levels of mature Gh-miR162, a stem-loop quantitative RT-PCR technique was used as previously described [52].

Complementary DNA was produced using the Rever-tAid First Strand cDNA Synthesis Kit (Fermentas) and 0.5 μg of total RNA previously treated with DNase I (Fermentas). cDNAs of the cotton DCL genes were synthesized by adding 100 μM Oligo (dT24V) primer. For synthesis of Gh-miR162 cDNA, 100 μM specific primer was added (Table 1). The presence of residual genomic DNA in the RNA samples was verified by PCR of the control gene xyloglucan endotransglycosylase (XTH) (accession number AY189971.2), using primers spanning two exons and RNA samples that were not reverse-transcribed (RT) (Additional file 4, figure s4).

Synthesized cDNAs were diluted 50 times and 2.5 μL of these dilutions were analyzed by quantitative PCR (qPCR). Assays were performed using a 48-well plate on an Step One Real-Time PCR system (Applied Biosystems) with Maxima™ SYBR Green/ROX qPCR Master Mix (Fermentas), following the manufacturer’s instructions. The cycling conditions were as follows: 10 min at 95°C for initial denaturation, followed by 40 cycles of denaturation at 95°C for 15 s and annealing/extension at 60°C for 30 s. Results were normalized against cotton genes for polyubiquitin (accession number DW505546) and the catalytic subunit of phosphatase 2A (accession number DT545658) [53]. The reference genes were validated experimentally in specific CLRDV-infected samples (Additional file 5, figure s5). All reactions were performed using two independent biological samples and each sample was analyzed in triplicate wells. The mean value of each Ct triplicate was used for further calculations by the 2^-ΔΔCt method. Each PCR run included a no-template control containing water instead of cDNA.

The efficiency values of the DCLs and Gh-miR162 primers sets were estimated for each experimental set by Miner software [54], and are listed in Table 1. Amplification of a specific transcript was confirmed by the appearance of a single peak in the melting curve followed by agarose gel electrophoresis (Additional file 6, figure s6). The correlation coefficient (R2) was calculated for each transcript (Table 1). The values shown are averages obtained from three biological replicates, and relative expression levels were obtained by comparing infected plants with uninfected plants.

Additional material

Additional file 1: Phylogenetic relationship between cotton Dicer ribonucleases and their homologues in other species A. B. C. and D. Unrooted Neighbor-joining tree constructed with DCL1, DCL2, DCL3, or DCL4 homologue sequences, respectively. Species used in the phylogeny were as follows: Arabidopsis thaliana (At), Gossypium hirsutum (Gh), Medicago truncatula (Mt), Oryza sativa (Os), Physcomitrella patens (Pp), Populus trichocarpa (Pv) and Vitis vinifera (Vv). Dashes below each tree represent amino acid regions used in alignment. Arrows represent fragments analyzed by qPCR. Bootstrap values from 1,000 replicates were used to assess the robustness of the trees. All DCL sequences, except cotton DCLs, were downloaded from Phytozome 6.0 http://www.phytozome.net/. ESTs from G. hirsutum containing incomplete DCL sequences were obtained from the NCBI database. The GhDCL1 consensus sequence was constructed with ESTs DT564382.1 (Helicase domain), and ES904646.1, together with DW238156.1 (two RNAse III and one Double stranded RNA binding (dsRB) domain). The GhDCL2 consensus sequence was constructed from two ESTs: DW484144 (DEAD-like helicases superb-family (DEAD) domain) and ES906737 (second RNAse III domain). The GhDCL3 sequence was constructed from the ESTs DW477937 and CRA62994 (PAZ and RNAse III domains, respectively). The GhDCL4 consensus sequence was constructed with ESTs ES941096 (PAZ domain) and DT58872 (RNAse III domain). Smart database [55] was used to identify DCL domains from their amino acid sequences.

Additional file 2: Analysis of biological duplicates of CLRDV-vsRNA populations. Histogram showing total (A) and unique (B) viRNA reads in each size class. Biological duplicates were subjected to deep sequencing in independent channels.

Additional file 3: Total RNA quality check. Quality and integrity of each RNA sample was checked by electrophoresis on 0.8% non-denaturing agarose gels, as well as by absorbance at 260 and 280 nm.
Additional file 4: Confirmation of DNA-free status of RNA samples. DNA contamination was checked by gel electrophoresis of products obtained in the xylloglucan endotransglycosylase (XTH) gene amplification reaction. Different sized fragments are amplified from genomic DNA (392 bp) and mRNA transcripts from cDNA (173 bp) with the designed primers. Before reverse transcription (-RT) reactions, RNA samples were used for PCR reactions and showed no amplification. Infected RNA samples from plants independently infected with CLRDV. Uninfected, two biologically independent RNA samples from uninfected plants. DNA; genomic DNA amplification (positive control).

Additional file 5: Determination of reference genes for use in these experimental conditions. Expression stability values of polyubiquitin (UBI), the catalytic subunit of phosphatase 2A (PP2A), and 18S ribosomal RNA (18S) candidate reference genes obtained by different algorithms. (A) NormFinder. (B) Delta CT method. (C) BestKeeper. (D) GeNorm. In all cases, the inclusion of an additional reference gene is not required [56]. All analyses were performed via the Cotton EST Database http://www.leonxie.com/index.php.

Additional file 6: Test of specificity of RT-qPCR primers. (A) Melting curves of the four GhDCLs and Gh-miR162 sequence-related RNAs after RT-qPCR using SYBR-green. (B) Non-denaturing agarose (2.0%) gel electrophoresis showing amplification of single products with the expected size for each of the GhDCL gene transcripts and Gh-miR162. M represents O’GeneRuler 100 bp DNA Ladder (Fermentas).

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TFS, RLC, EAACR, and MSFV conceived and designed the experiments. TFS, LF, MM, and CD performed the experiments. TFS, RRSA, EAACR, and CEGs carried out bioinformatics analysis and analyzed the data. TFS, RLC, LF and MSFV contributed reagents, materials and analytical tools. TFS, RLC and MSFV wrote the paper. All authors read and approved the final manuscript.

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