siRNA Machinery in Whitefly (*Bemisia tabaci*)

Santosh Kumar Upadhyay1*, Sameer Dixit2, Shailesh Sharma1, Harpal Singh2, Jitesh Kumar1, Praveen C. Verma2*, K. Chandrashekar2,3*

1 National Agri-Food Biotechnology Institute, (Department of Biotechnology, Government of India), Mohali, Punjab, India, 2 CSIR-National Botanical Research Institute, Council of Scientific and Industrial Research, RanaPratapMarg, Lucknow, India, 3 Indian Agricultural Research Institute-Regional Station, Agricultural College Estate, Shivaji Nagar, Pune, Maharashtra, India

**Abstract**

**Background:** RNA interference has been emerged as an utmost tool for the control of sap sucking insect pests. Systemic response is necessary to control them in field condition. Whitefly is observed to be more prone to siRNA in recent studies, however the siRNA machinery and mechanism is not well established.

**Methodology/Principal Findings:** To identify the core siRNA machinery, we curated transcriptome data of whitefly from NCBI database. Partial mRNA sequences encoding Dicer2, R2D2, Argonaute2 and Sid1 were identified by tblastn search of homologous sequences from *Aphis glycines* and *Tribolium castaneum*. Complete encoding sequences were obtained by RACE, protein sequences derived by Expasy translate tool and confirmed by blastp analysis. Conserved domain search and Prosite-Scan showed similar domain architecture as reported in homologs from related insects. We found helicase, PAZ, RNaseSIIa, RNaseSIIb and double-stranded RNA-binding fold (DSRBF) in Dicer2; DsRBD in R2D2; and PAZ and PIWI domains in Argonaute2. Eleven transmembrane domains were detected in Sid1. Sequence homology and phylogenetic analysis revealed that RNAi machinery of whitefly is close to Aphids. Real-time PCR analysis showed similar expression of these genes in different developmental stages as reported in *A. glycines* and *T. castaneum*. Further, the expression level of above genes was quite similar to the housekeeping gene actin.

**Conclusions/Significance:** Availability of core siRNA machinery including the Sid1 and their universal expression in reasonable quantity indicated significant response of whitefly towards siRNA. Present report opens the way for controlling whitefly, one of the most destructive crop insect pest.

Citation: Upadhyay SK, Dixit S, Sharma S, Singh H, Kumar J, et al. (2013) siRNA Machinery in Whitefly (*Bemisia tabaci*). PLoS ONE 8(12): e83692. doi:10.1371/journal.pone.0083692

Editor: Subba Reddy Palli, University of Kentucky, United States of America

Received June 10, 2013; Accepted November 6, 2013; Published December 31, 2013

Copyright: © 2013 Upadhyay et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Funding: This work is supported by CSIR-Empower project grant and DST-INSPIRE Grant. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: The authors have declared that no competing interests exist.

* E-mail: santoshnbri@hotmail.com (SKU); praveencverma@yahoo.com (PCV); kc_shekar2001@yahoo.com (KC)

**Introduction**

Transgenic crops expressing δ-endotoxins of *Bacillus thuringiensis* (Bt) provide incredible control of chewing type lepidopteran and coleopteran pests [1]. However, they are completely failed against sap sucking hemipteran insects like aphids, mealybugs, whiteflies and others [2,3]. These sucking pests are now emerged as major pests in crop field. Some of the plant lectins are reported to be effective against these insects, but none of them are toxic to whiteflies. RNA interference (RNAi) has been reported as good alternative to combat these issues. Although most of the RNAi studies are focused on regulation, expression of target genes and mechanism of small RNA in the insects [4,5]; yet the development of insect resistant transgenic plants expressing dsRNA/siRNA is becoming more popular due to their target specificity [6–9].

RNAi can be triggered by both exogenous and endogenous dsRNA/siRNA, which silences the endogenous target gene having similar sequence. RNAi has been described in various insect orders including hemiptera [10]. siRNA and miRNA pathways are reported as two overlapping pathways for RNA mediated gene silencing. Both siRNA and miRNA pathways use related but discrete protein molecules at each step of their activity. Dicer1, Loquacious and Argonaute1 are involved in miRNA pathway in *Drosophila*, while Dicer 2, R2D2 and Argonaute2 function in siRNA mediated pathway [11–15]. *Tribolium castaneum* is reported as model organism among insects for systemic silencing by RNAi [16–18] because *Drosophila melanogaster* do not show systemic RNAi response due to the absence of gene called systemic RNA interference deficient-1 (Sid1) [19,20]. Sid1 is accountable for scattering the intensified signal for RNAi [21]. A few reports are available for Sid1 in insects and further identification and characterization of homologous sequences is in progress from other insects [18,22,23]. siRNA mediated control of insects like whitefly can be accelerated by understanding the pathway and mechanism of systemic silencing. High level expression of Sid1 in target insects might act as an indicator for systemic RNAi response.

Whitefly (*B. tabaci*) is reported as a serious pest of several crops. Further, none of the reported insecticidal proteins are significantly effective against them. However, in our previous study, we have shown RNAi as a good tool for the control of whitefly [24]. Although we found that some of critical gene targeted by siRNA molecules (like vATPaseA, RPL 9) are very effective [24], but the
RNAi machinery and their mechanism in whiteflies are still unknown. Recently, availability of transcriptome data of whitefly on NCBI [25,26,27] opens the possibility of exploring RNAi machinery. Present study aimed to identify and characterize major components of siRNA machinery like Dicer2, R2D2, Argonaute2 and Sid1 in whiteflies. We found the presence of complete siRNA machinery in whitefly and significant expression level in different developmental stages.

Materials and Methods

Mining of cDNA encoding Dicer2, R2D2, Argonaute2 and Sid1 from transcriptome data

To identify the Dicer2, R2D2, Argonaute2 and Sid1 encoding cDNA from whitefly transcriptome data, homologous proteins sequences (AFZ74931, AFZ74932, AFZ74933 and AFZ74934 for Dicer2, R2D2, Argonaute2 and Sid1 of *Aphis glycines*; and NP_001107840.1, NP_001128425.1, NP_001107842.1 and NP_001099012.1 for Dicer2, R2D2, Argonaute2 and Sid1 of *T. castaneum*, respectively) were downloaded from NCBI database [28]. These sequences were used for blastn search against transcriptome data (i.e. TSA) of *B. tabaci* and matching TSA sequences were retrieved. To further confirm the identity, TSA sequences were analysed using blastx search against NCBI non-redundant proteins database.

Cloning of full length genes by RACE

Sequences identified in above blast analysis were used for primer designing to obtain complete gene sequence by RACE. Sequences used were as follows- E2956195.1 for R2D2, HP663253.1 for Argonaute2 and EZ964892.1 for Sid1. In case of Dicer2, we designed primers from two TSA sequences (EZ956963.1 and EZ954838.1) to ease the amplification, cloning of Dicer2, we designed primers from two TSA sequences (EZ956195.1 for R2D2, EZ954838.1 for Sid1 of *Aphis glycines*; and NP_001107840.1, NP_001128425.1, NP_001107842.1 and NP_001099012.1 for Dicer2, R2D2, Argonaute2 and Sid1 of *T. castaneum*, respectively) were downloaded from NCBI database [28]. These sequences were used for blastn search against transcriptome data (i.e. TSA) of *B. tabaci* and matching TSA sequences were retrieved. To further confirm the identity, TSA sequences were analysed using blastx search against NCBI non-redundant proteins database.

Sequence analysis

To get the open reading frame, each gene sequence was subjected to ORF finder (http://www.ncbi.nlm.nih.gov/orffig.cgi) at NCBI database. ORFs of all the genes were further confirmed by blast against NCBI database. Encoding protein sequences were derived by Expasy translate tool (http://web.expasy.org/translate/). Theoretical molecular mass and PI of translated sequences were determined by Expasy MW/PI tool (http://web.expasy.org/compute_pi/).

To analyse the domain architecture of whitefly Dicer2, R2D2, Argonaute2 and Sid1; derived protein sequences were subjected to Scan-Prosite (http://prosite.expasy.org/scanprosite/), a database of protein families and domains [29]. It contains pattern and profile specific for thousands of protein families or domains. Sid1 sequence was analysed by TMHMM server v2.0 (http://www.cbs.dtu.dk/services/TMHMM/) and InterProScan (http://www.ebi.ac.uk/Tools/pfa/iprscan/) to detect the transmembrane helices. TMHMM is a server which predicts transmembrane protein topology with hidden Markov model. Gene sequences encoding *B. tabaci* Dicer2, R2D2, Argonaute2 and Sid1 were submitted to NCBI database (Table 2).

Multiple sequence alignments (MSA) and phylogenetic analysis

Multiple sequence alignments were performed with the well-known insect sequences to analyse the homology, and presence of conserved domains and amino acids sequences. Sequences used for alignment are given in table S1. Phylogenetic analyses were performed by MEGA 5.2.1 software. Conserved domains used in phylogenetic analysis were RNAseIIIa and b of Dicer2 and DsRBD of R2D2. Further full length protein sequences of Dicer2, R2D2, Argonaute2 and Sid1 were also used in phylogenetic analysis. Sequence alignments were performed using Muscle. Neighbour joining analysis was performed with boots trapping test using 10,000 replicates. Maximum likelihood analysis [30] was also performed for the same alignments; however both the analysis showed similar relationship.

Expression analysis of core components of siRNA pathway in different developmental stages of whitefly

For experimental purpose, we reared whiteflies in control condition on cotton plants as described earlier [24]. Total RNA was isolated from egg, nymph and adult insects (~10 mg each) using Tri reagents (Sigma, USA). cDNA was synthesized from 2 μg of total RNA using first strand cDNA synthesis kit (Invitrogen, USA). Quality of cDNA was analysed by PCR amplification of *actin* gene. cDNA from different stages of insects

### Table 1. Primer used in RACE of siRNA components of whitefly.

| Gene  | 5’ RACE (5’-3’) | 3’ RACE (5’-3’) |
|-------|----------------|----------------|
| Dicer2 | GSP1 GTTGATCGAAGTCGTGATCCAC | GGAAGGAGCAGCCTACAGGT |
|       | NSGP1 CTCCAGCTGGCCTGGTCTCT | CCGGGAGTACCAAACATTGATCATG |
| R2D2  | GSP1 CTCCAGCTGGCCTGGTCTCT | CCGGGAGTACCAAACATTGATCATG |
|       | NSGP1 CTCCAGCTGGCCTGGTCTCT | CCGGGAGTACCAAACATTGATCATG |
| Argonaute2 | GSP1 TTTCATCGATTTGGGAGCGCG | GAACTCAAGAACGACTGGGCAGC |
|       | NSGP1 TGGCGAGGGTATTGGTCTGCT | GACGACTGGAGCTCTCAG |
| Sid1  | GSP1 GTATATATTGGCATGTATCCATCTG | GAGCCCTACTGAGCCCCTACCA |
|       | NSGP1 ACTATATATTGGCATGTATCCATCTG | GAGCCCTACTGAGCCCCTACCA |

doi:10.1371/journal.pone.0083692.t001
was used for expression analysis of Dicer2, Argonaute2, R2D2 and Sid1 by real time PCR on GeneAmp 5700 (Applied Biosystems, USA) using SYBR Green detection dye (InvivoGen, USA). Primers used for real time PCR is provided in Table 3. Amplification of actin gene was used as control. Expression analysis experiment was performed in triplicates.

**Results and Discussion**

**Identification and cloning of core components of siRNA pathway in whitefly**

Usually the core components of siRNA machinery are highly conserved within species, however the depth of conservation often differs between the species. Further the efficiency of RNAi and degree of systemic response also varies from species to species. In certain organisms like *C. elegans* and *Tricholium*, injection of a small amount of dsRNA induces significant systemic response [17,21]. However, some lepidopteran insect do not show such kind of response [31]. Therefore, understanding of molecular machinery of RNAi is pre-requisite in different insects. Presence and absence of the components of RNAi machinery (especially Sid1 protein), in an organism might be an indicator for their response. Therefore, we surveyed for the presence of core components of siRNA machinery in whitefly, a devastating insect pest of several crops.

We mined the transcriptome data of whitefly for the presence of Dicer2, Argonaute2, R2D2 and Sid1 by using homologous sequences from *A. glycines* and *T. castaneum* [28]. In this process, we identified mRNA sequences EZ956963.1 and EZ954838.1 for Dicer2, EZ956195.1 for R2D2, HP663253.1 for Argonaute 2 and EZ964892.1 for Sid1; which were used in primer designing. Gene specific primers (Table 1) were designed and complete gene sequences obtained by RACE (File S1). Sequences were submitted to NCBI, accession numbers, molecular weight, pI and highly homologous protein to each sequence is given in Table 2. Whitely Dicer2, R2D2, Argonaute2 and Sid1 showed homology with *Blattella germanica* (accession number CCF23094.1), *Acromyrmex echinatior* (EGH7607.1), *N. lugens* (AGH30327.1) and *Locusta migratoria* (AFQ00936.1) protein sequences, respectively.

**Dicer**

Dicer is a multi-domain protein basically involved in generation of small RNA molecules (siRNA, miRNA) [32,33]. A typical Dicer contains two N-terminal helicase domains, one PAZ domain, tandem RNAseIII domains and a C-terminal dsRNA binding domain (Figure 1). In case of *C. elegans*, single Dicer protein is responsible for both miRNA and siRNA pathway [33–35]. However, both the pathways are governed by two different Dicers (Dm-Dcr1 and Dm-Dcr2) in *Drosophila* [12]. Dm-Dcr2 is involved in siRNA pathway, whereas Dm-Dcr1 in miRNA pathway. Similar kind of gene duplication is also reported in aphid *Acyrthosiphon pisum* [36].

We retrieved whitely TSA sequences (accession numbers - EZ956963.1, EZ957381.1, EZ955681.1, EZ954014.1, EZ954838.1, EZ954888.1, EZ939818.1, HP647437.1, EZ942655.1, EZ942655.1, EZ947031.1) showing significant similarity in tblastn search with *A. glycines* and *T. castaneum* Dicer2 protein sequences. Blastx search of retrieved sequences at NCBI-nr protein database indicated the presence of two Dicer (Dicer1 and Dicer2) proteins in whiteflies, as reported in case of other insects [12,36]. It is possible that Dicer1 is involved in miRNA pathway and Dicer2 in siRNA pathway. Since, we were focussing on characterization of siRNA machinery; we amplified the dicer gene only. Scan-Prosite search of whitely Dicer2 protein sequence showed the domain organization similar to *T. castaneum* Dicer2 (Figure 1). C-terminus double stranded RNA binding domain (DspRD, PS50137) was absent in whitely, as reported in *T. castaneum*. However, other domains like helicase, double-stranded RNA-binding fold (DSRBF), PAZ and RNAseIII were similar to other analysed insects. Whitely Dicer2 contains two helicase, one DSRBF, one PAZ and two RNAseIII (a and b) domains. Scan-Prosite analysis showed significant score for helicase I (21.6), II (12.8), PAZ (16.1), RNAseIIIa (14.5) and b (34.0) domains (Table 4), which were similar to other insects. However, whitely and *T. castaneum* lacks the C-terminus DSRBD domain and *D. melanogaster* lacks full-length PAZ domain.

Multiple sequence alignments and phylogenetic analysis of Dicer2 were performed using full length protein as well as RNAseIIIa and b domains sequences from different insects (Table S1, File S2, Figure 2a b and c). Insect Dicer2 proteins were clustered in two groups apart from *D. melanogaster*. Full length whitely Dicer2 clustered with aphids. However RNAseIIIa and b domains are clustered with aphids and *B. germanica*, respectively.

Multiple sequence alignment results also supported the phylogenetic results. Whitely RNAseIIIa showed ~45% homology with aphids; however RNAseIIIb showed ~63% with *B. germanica* (File S2). Results indicated that the two domains might evolve independently during evolution.

**Argonaute**

Argonaute is a core component of miRNA and siRNA pathways [37,38]. It contains two distinctive domains i.e. PAZ and PIWI [38]. Besides these, DUF1785 domain is also reported, however its function is still unknown. PAZ domain is responsible for siRNA binding at 2 nucleotide 3’ overhang, while PIWI domain shows RNAseH like activity. Argonaute is reported as large family of protein in *C. elegans* and *Drosophila* with different functions [11,39]. Five different Argonautes are reported in *Tricholium* and *Drosophila* [18], in which Argonaute1 and 2 are involved in miRNA and siRNA pathway, respectively [11]. Similar kind of gene diversification is also observed in case of *A. pisum* [36]. However, single Argonaute is reported from *A. glycines* [28].

**Table 2.** Details of whitefly Dicer2, R2D2, Argonaute2 and Sid1 sequences.

| Gene       | Accession Number | Orf length | Protein length (AA) | Molecular mass (kDa) | pI     | Closest homolog                        | % Similarity |
|------------|------------------|------------|---------------------|----------------------|--------|----------------------------------------|--------------|
| Dicer2     | KF740508         | 4944       | 1647                | 189.0                | 5.91   | *Blattella germanica* CCF23094.1       | 38           |
| R2D2       | KF740509         | 759        | 252                 | 27.7                 | 6.98   | *Acromyrmex echinatior* EG67607.1     | 40           |
| Argonaute2 | KF192313         | 2505       | 834                 | 95.1                 | 9.48   | *Nilaparvata lugens* AGH30327.1       | 51           |
| Sid1       | KF192314         | 2181       | 726                 | 83.1                 | 5.79   | *Locusta migratoria* AFQ00936.1       | 51           |

doi:10.1371/journal.pone.0083692.t002
Argonaute2 [14,43,44]. forms the RISC loading complex, and enhances siRNA transfer to Dcr2 [13,14,42]. Dcr2/R2D2 complex binds to duplex siRNA, however R2D2 is not directly involved in siRNA producing activity of Dcr1 by increasing the affinity toward pre-miRNA, ing activity of Dcr1 by increasing the affinity toward pre-miRNA, Dicer complex [45]. We observed that three non-bridging oxygen atoms at 5
phosphate of siRNA involve in interaction with several amino acids of PIWI domain [41]. We observed the presence of these amino acids in whitefly and fount that they were highly conserved in different organisms (File S3).

Phylogenetic analysis of whitefly Argonaute2 was performed with selected insects (Figure 3c). It was grouped with N. lugens and aphids. Multiple sequence alignment of PIWI domain showed high homology with N. lugens and aphids sequences followed by L. migratoria and T. castaneum (File S3). These results indicate close homology of whitefly Argonaute2 with other related insects.

**R2D2**

R2D2 and Loquacious are family of dsRNA-binding proteins and function in tandem with specific RNaseIII enzymes. There are two dsRNA-binding domains in R2D2 and three in Loquacious. Two distinct Dicer complexes, Dcr1/Loquacious and Dcr2/R2D2 are reported in Drosophila, which produce miRNA and siRNA, respectively [13,14,15,42]. Loquacious enhances miRNA producing activity of Dcr1 by increasing the affinity toward pre-miRNA, however R2D2 is not directly involved in siRNA producing activity of Dcr2 [13,14,42]. Dcr2/R2D2 complex binds to duplex siRNA, forms the RISC loading complex, and enhances siRNA transfer to Argonaute2 [14,43,44].

**Table 3. Primers used in real time PCR of siRNA components of whitefly.**

| Gene   | Forward primer (5’-3’) | Reverse Primer (5’-3’) |
|--------|------------------------|------------------------|
| Dicer2 | CAGCCTCAGGATTACTCT     | CCGTCTCCTGTACCAAG      |
| R2D2   | GTCCGCTAGTACCTGGTAC    | GGAGCAACCACTTCCTC      |
| Argonaute2 | GGCCCCAGGCGAGCCTAAT  | CCGTGATGCAAGCATTCTA    |
| Sid1   | CACACTTCCAGGGCAGACC   | TTGGTGGATGATGGGTGATG   |
| Actin  | GAGCCGACCAACTTCAACAG  | CTTTTGTCAGGTAGGTCTTCAGT |

doi:10.1371/journal.pone.0083692.t003

Phylogenetic analyses were performed with full length as well as DSRBD1 and DSRBD2 domains sequences of whitefly R2D2 with other insect sequences. Full length R2D2 and DSRBD1 were clustered with aphids. DSRBD2 was clustered with B. mori, however closely followed by aphids (Figure 4b, c and d). Multiple sequence alignment also supports the phylogenetic results. Overall, we found both R2D2 and Loquacious in B. tabaci; which might be involved in two parallel siRNA and miRNA pathways, respectively.

**Sid 1**

It is the best known protein for systemic RNAi in C. elegans [21,45] and insects [18]. It comprises tandem repeats of transmembrane domains along with long N-terminus extracellular domain. Transmembrane domains form channel for the movement of dsRNA molecules [21,45]. Sid1 is reported from several insects like T. castaneum, A. mellifera, A. glycines, B. mori and others, and involved in systemic spreading of RNAi [18,28]. However, it is absent in Drosophila which lacks the systemic RNAi response. Tomoyasu et al. [18] performed robust analysis of Sid1 gene from several insects genome including 11 Drosophila species and tried to correlate the presence and absence of Sid1 gene with RNAi response, however it is still under debate [22,23,46,47,48].

Expression of C. elegans Sid1 in Drosophila culture cells enables them to uptake dsRNA from media, which confirmed the role of Sid1 in dsRNA uptake [45]. However, Luo et al [23] reported that Sid1 is not required for systemic RNAi in the migratory locust Locusta migratoria. This showed that role of insect Sid1 in systemic silencing is still a matter of profound investigation.

After deep analysis of transcriptome data available at NCBI [25,26,27], we found that at least one Sid1 gene is present in whitefly, as observed in case of aphids [28]. Full length gene was obtained by RACE, which encodes for 726 amino acids residue long protein (Table 2, File S1). Blastp analysis at NCBI-nr protein database confirmed that the cloned gene was Sid1. Domain architecture of Sid1 was analysed by TMHMM server version 2.0 and InterProScan, which showed the presence of 11 transmembrane domains separated by extra and intracellular domains (Figure 5a). Besides this, a long extracellular domain was located at N-terminus. Similar kind of domain organization has been observed with aphid A. glycines, followed by P. humanus and A. mellifera.

Phylogenetic analyses of Sid1 were also found highly conserved (File S3). PIWI domain sequence was analysed for the presence of signature residues involve in binding with siRNA/miRNA. It is reported that three non-bridging oxygen atoms at 5’ phosphate of siRNA involve in interaction with several amino acids of PIWI domain [41]. We observed the presence of these amino acids in whitefly and fount that they were highly conserved in different organisms (File S3).

**Table 3. Primers used in real time PCR of siRNA components of whitefly.**

| Gene   | Forward primer (5’-3’) | Reverse Primer (5’-3’) |
|--------|------------------------|------------------------|
| Dicer2 | CAGCCTCAGGATTACTCT     | CCGTCTCCTGTACCAAG      |
| R2D2   | GTCCGCTAGTACCTGGTAC    | GGAGCAACCACTTCCTC      |
| Argonaute2 | GGCCCCAGGCGAGCCTAAT  | CCGTGATGCAAGCATTCTA    |
| Sid1   | CACACTTCCAGGGCAGACC   | TTGGTGGATGATGGGTGATG   |
| Actin  | GAGCCGACCAACTTCAACAG  | CTTTTGTCAGGTAGGTCTTCAGT |

doi:10.1371/journal.pone.0083692.t003

Phylogenetic analyses were performed with full length as well as DSRBD1 and DSRBD2 domains sequences of whitefly R2D2 with other insect sequences. Full length R2D2 and DSRBD1 were clustered with aphids. DSRBD2 was clustered with B. mori, however closely followed by aphids (Figure 4b, c and d). Multiple sequence alignment also supports the phylogenetic results. Overall, we found both R2D2 and Loquacious in B. tabaci; which might be involved in two parallel siRNA and miRNA pathways, respectively.

**Sid 1**

It is the best known protein for systemic RNAi in C. elegans [21,45] and insects [18]. It comprises tandem repeats of transmembrane domains along with long N-terminus extracellular domain. Transmembrane domains form channel for the movement of dsRNA molecules [21,45]. Sid1 is reported from several insects like T. castaneum, A. mellifera, A. glycines, B. mori and others, and involved in systemic spreading of RNAi [18,28]. However, it is absent in Drosophila which lacks the systemic RNAi response. Tomoyasu et al. [18] performed robust analysis of Sid1 gene from several insects genome including 11 Drosophila species and tried to correlate the presence and absence of Sid1 gene with RNAi response, however it is still under debate [22,23,46,47,48].

Expression of C. elegans Sid1 in Drosophila culture cells enables them to uptake dsRNA from media, which confirmed the role of Sid1 in dsRNA uptake [45]. However, Luo et al [23] reported that Sid1 is not required for systemic RNAi in the migratory locust Locusta migratoria. This showed that role of insect Sid1 in systemic silencing is still a matter of profound investigation.

After deep analysis of transcriptome data available at NCBI [25,26,27], we found that at least one Sid1 gene is present in whitefly, as observed in case of aphids [28]. Full length gene was obtained by RACE, which encodes for 726 amino acids residue long protein (Table 2, File S1). Blastp analysis at NCBI-nr protein database confirmed that the cloned gene was Sid1. Domain architecture of Sid1 was analysed by TMHMM server version 2.0 and InterProScan, which showed the presence of 11 transmembrane domains separated by extra and intracellular domains (Figure 5a). Besides this, a long extracellular domain was located at N-terminus. Similar kind of domain organization has been observed with aphid A. glycines, followed by P. humanus and A. mellifera.
Whitefly Dicer2 showed the presence of all the domains except DSRBF as reported in *T. castaneum*.  
doi:10.1371/journal.pone.0083692.g001

### Table 4. Scan-Prosite score for common domains of Dicer2 protein in selected insects.

| Dicer2 | HELICASE 1 | HELICASE 2 | PAZ | RNAse III (a) | RNAse III (b) | DSRBD |
|--------|------------|------------|-----|---------------|---------------|-------|
| *B. tabaci* | 21.6 | 12.8 | 16.1 | 14.5 | 34.0 | Absent |
| *A. glycines* | 19.4 | 13.9 | 17.3 | 20.0 | 35.5 | 9.9 |
| *A. pisum* | 19.6 | 13.6 | 18.4 | 19.6 | 33.9 | 9.6 |
| *B. mori* | 21.3 | 12.6 | 11.9 | 21.3 | 34.6 | 9.0 |
| *C. elegans* | 22.7 | 14.6 | 23.5 | 23.7 | 40.6 | 11.9 |
| *D. melanogaster* | 18.6 | 12.1 | 8.7 | 18.4 | 31.0 | 9.6 |
| *T. castaneum* | 22.1 | 12.0 | 17.1 | 23.8 | 36.8 | Absent |

doi:10.1371/journal.pone.0083692.t004

Figure 2. Phylogenetic analysis of whitefly Dicer2 protein with other insects and *C. elegans*. Phylogenetic trees were constructed from amino acid sequences of (A) full length protein and (B) RNAseIIa and (C) RNAseIIIb domains of whitefly Dicer2 clustered with aphids.  
doi:10.1371/journal.pone.0083692.g002
(44%). In phylogenetic analysis, whitefly Sid1 was clustered with aphids and result was in agreement with the multiple sequence alignment (Figure 5b). Further, B. mori Sid 1, 2 and 3 were clustered together, and closer to the T. castaneum.

Expression analysis of siRNA components

Expression analysis of dicer2, r2d2, argonaute2 and sid1 genes of whitefly was performed in egg, nymph and adult insects by real time PCR. Expression level was compared with the actin gene. We found that all the genes were expressed at each developmental stage (Figure 6). Significant transcript abundance was observed for each gene which was almost equal to the expression level of actin. All the genes expressed at nearly similar level in all developmental stages. Similar result has been reported in case of A. glycines and T. castaneum [18,28]. Significant expression of siRNA components in whitefly indicated the possibility of massive siRNA response, and creates a hope for the use of this technique in insect control.

Conclusion

We observed that the siRNA machinery of whitefly showed significant sequence homology with aphids and other insects. Further, transcript abundance of each component was also significant. These results indicated the possibility of massive siRNA response in whitefly. However, the previous reports with whitefly and other insects like A. pisum with similar domain organization and expression show inconsistent siRNA response with different target genes. In earlier study we observed that feeding of equal quantity siRNA targeting different genes (actin ortholog, ADP/ATP translocase, α-tubulin, ribosomal protein L9 and V-ATPase A subunit) in whitefly showed diverse kind of responses [24]. Ribosomal protein L9 and V-ATPaseA targeting siRNA caused significant mortality of whitefly in comparison to others. In A. pisum, only transient reduction in gene expression is reported after dsRNA injection and feeding [36,49]. However, injection of siRNAs targeting coo2 gene of aphid salivary protein showed strong systemic response in A. pisum [50]. But similar response was not observed in green peach aphid M. persicae for the same gene when delivered through transgenic plants [51]. These variations in RNAi responses might be due to the difference in importance of genes, method of delivery, different role of same gene in various insects and others so many unknown regions. Therefore, future studies regarding the insect control can target multiple genes at a time to get significant response. We have observed in our earlier experiment that the feeding of dsRNA through artificial diet offers the best option for the screening of
target gene in insects [24]. Moreover, translation of such technology efficiently in the field by using transgenic plants is necessary [7,8]. In this process we have developed the transgenic plants expressing the most effective dsRNA (V-ATPase A, which was earlier analysed by feeding in artificial diet) [24] and found similar effect [unpublished data]. Further, present study of characterization and gene expression analysis of siRNA machinery

![Figure 4. Domain architecture and phylogenetic analysis of whitefly R2D2 protein.](/images/figure4.jpg)

**Figure 4.** Domain architecture and phylogenetic analysis of whitefly R2D2 protein. (A) Comparative domain architecture of whitefly R2D2 with other insects and C. elegans. Figure shows two DSRBD in whitefly R2D2 as reported in other insects and C. elegans. (B) (C) and (D) Phylogenetic analysis of full length, and DSRBD1 and DSRBD2 domains of whitefly R2D2, respectively. DSRBD1 clustered with aphids; however DSRBD2 clustered with B. mori, indicating independent evolution of both the domains.

doi:10.1371/journal.pone.0083692.g004

**Table 5.** Scan-Prosite score for common domains of Argonaute2 protein in selected insects.

| Argonaute2 | PAZ | PIWI |
|------------|-----|------|
| B. tabaci  | 18.2| 46.8 |
| A. glycines| 15.8| 44.3 |
| A. pisum   | 16.7| 44.9 |
| B. mori    | 11.9| 34.8 |
| C. elegans | 31.1| 50.8 |
| D. melanogaster | 14.1| 42.2 |
| T. castaneum| 12.7| 38.2 |

doi:10.1371/journal.pone.0083692.t005

**Table 6.** Scan-Prosite score for common domains of R2D2 protein in selected insects.

| R2D2 | DSRBD 1 | DSRBD 2 |
|------|---------|---------|
| B. tabaci | 14.8 | 15.4 |
| A. glycines| 14.4 | 13.7 |
| A. pisum  | 14.8 | 13.9 |
| B. mori   | 15.8 | 15.2 |
| C. elegans| 10.9 | 9.5  |
| D. melanogaster | 12.4| 11.7 |
| T. castaneum| 14.4| 14.5 |

doi:10.1371/journal.pone.0083692.t006
supports our earlier results and opens a new way for the presumption of insect responses towards RNAi.

Sid1 has been reported from diverse groups of insects except some dipterans like Drosophila and correlated with the systemic RNAi responses [52,53]. Further it is highly conserved among different taxa even when they are discrete from each other [18]. Lack of Sid1 in dipterans is astonishing and therefore very deep analysis is required regarding the molecular evolution of Sid1 by wide sampling of insect orders including diptera. Moreover, Luo et al [23] reported that Sid1 is not required for systemic RNAi in the migratory locust Locusta migratoria. These reports indicated that wide analysis of different insect is required to reach the base of RNAi.

Systemic and vigorous RNAi response is pre-requisite for the RNAi based pest control using transgenic crops. Knowledge of siRNA machinery and their detailed characterization not only explains the molecular mechanism of RNAi, but also indicates the probable response of target insects before developing the transgenic plants.

Supporting Information

File S1 Complete nucleotide and protein sequences of core RNAi components of whitefly (B. tabaci). (A) Dicer2, (B) R2D2, (C) Argonaute and (D) Sid1. Important domains are highlighted by different colours.

File S2 Sequence alignment of RNaseIIIa (A) and RNaseIIIb (B) of Dicer2.

File S3 Sequence alignment of PIWI domain of Argonaute2. Triangle indicates the residues interact with oxygen molecules of 5’P of miRNA/siRNA [ref. 14].
File S4: Multiple sequence alignment of R2D2.

File S5: Alignment of Sid1 sequences. Black line denotes the conserved region in N-terminus extracellular domains. Blue lines denote the trans-membrane helix.

Table S1: Sequences used for various analyses during study.

Acknowledgments

Authors are thankful to CSIR-National Botanical Research Institute and National Agri-Food Biotechnology Institute for providing research facility.

References

1. Sanahuja G, Banakar R, Twyman RM, Capell T, Christou P (2011) Bacillariar Thuringiensis: a century of research, development and commercial applications. Plant Biotechnol 9: 203–300.
2. Dutt U (2007) Mealy bug infestation in Punjab: Bt cotton falls flat Environment News Service, 21 August (countercurrents.org).
3. Faria CA, Wackers FL, Pritchard J, Barrett DA, Turlings TC (2007) High susceptibility of br maize to aphids enhances the performance of parasitoids of lepidopteran pests. PLoS One 2: e600.
4. Ghanim M, Kontsedalov S, Czosnek H (2007) Tissue-specific gene silencing by RNA interference for the control of whiteflies (Bemisia tabaci). Genetica. Insect Biochem Mol Biol 37: 732–739.
5. Zhang H, Li HC, Miao XX (2013) Feasibility, limitation and possible solutions of RNAi-based technology for insect pest control. Insect Sci 20: 12–30.
6. Gordon KHJ, Waterhouse PM (2007) RNAi for insect proof plants. Nature Biotech 25: 1231–1232.
7. Baum JA, Roggen T, Clinton W, Heck GR, Feldmann P, et al. (2007) Control of coleopteran insect pests through RNA interference. Nature Biotech 25: 1322–1326.
8. Mao YB, Cui WJ, Wang JW, Hong GJ, Tao XY, et al. (2007) Silencing a cotton bollworm P450 monooxygenase gene by plant-mediated RNAi impairs larval tolerance of gossypol. Nature Biotech 25: 1307–1315.
9. Chen J, Zhang D, Yao Q, Zhang J, Dong X, et al. (2010) Feeding-based RNA interference of a trehalose phosphate synthase gene in the brown planthopper, Nilaparvata lugens. Insect Mol Biol 19: 777–786.
10. Hovenne H, Smagghe G (2010) Mechanisms of dsRNA uptake in insects and potential of RNAi for pest control: A review. J Insect Physiol 56: 227–235.
11. Okamura K, Ishizuka A, Siomi H, Siomi MC (2004) Distinct roles for Argonaute proteins in small RNA-directed RNA cleavage pathways. Genes Dev 18: 1655–1666.
12. Lee YS, Nakahara K, Pham JW, Kim K, He Z, et al. (2004) Alignement of Sid1 sequences. Black line denotes the conserved region in N-terminus extracellular domains. Blue lines denote the trans-membrane helix.
13. Liu Q, Rand TA, Clinton W, Heck GR, Feldmann P, et al. (2007) Parental RNAi in absence of transitive and systemic pathways allows cell-specific and isoform-specific RNAi in Tribolium castaneum. Genome Biol 9: R10.
14. Lee YS, Nakahara K, Pham JW, Kim K, He Z, et al. (2004) Distinct roles for Drosophila Dicer-1 and Dicer-2 in the siRNA/miRNA silencing pathways. Cell 117: 69–81.
15. Saito K, Ishizuka A, Siomi H, Siomi MC (2005) Processing of pre-micro RNAs by the Dicer-1-Loquacious complex in Drosophila melanogaster. Mol Biol Evol 22: 1561–1568.
16. Liu Q, Rand TA, Akahisa S, Du F, Kim HE, et al. (2003) R2D2, a bridge between the initiation and effector steps of the Drosophila RNAi pathway. Science 301: 1921–1925.
17. Forstermann K, Tomari Y, Du T, VaginVV, Denli AM, et al. (2005) Normal micro RNA maturation and germ-line stem cell maintenance requires Loquacious, a double-stranded RNA-binding domain protein. PLoS Biol 3: e236.
18. Richards S, Gibbs RA, Weinstock GM, Brown SJ, Demell R, et al. (2006) The genome of the model beetle and pest Tribolium castaneum. Nature 432: 949–955.
19. Bucher G, Scholten J, Klingler E, Muller G (2002) Parental RNAi in Tribolium (Coleoptera). Cur Biol 12: R53–R56.
20. Tomoyasu Y, Miller SC, Tomita S, Schoppmeier M, Grossmann D, et al. (2008) Exploring systemic RNA interference in insects A genome-wide survey for RNAi genes in Tribolium. Genome Biol 9: R10.
21. Reiganant JV, Carre C, Magat B, Szymczak D, Lespamant JA, et al. (2003) Expansion of transitive and systemic pathways allows cell-specific and isoform-specific RNAi in Drosophila. RNA 9: 299–306.
22. Price DRG, Gatehouse JA (2000) RNA-mediated crop protection against insects. Trends in Biotech 20: 393–400.
23. Winston WM, Molodovitch C, Hunter CP (2002) Systemic RNAi in C. elegans requires the putative transmembrane protein SDF-1. Science. 295: 2456–2459.
24. Anwesin K, Pasik B, Saldizar E (2006) Sid-1 is implicated in systemic gene silencing in the honey bee. J Apicultural Res 45: 29–24.
25. Luo Y, Wang X, Yu D, Kang L (2012) The SID-1 double-stranded RNA transporter is not required for systemic RNAi in the migratory locust. RNA Biol 9: 665–671.
26. Upadhyay SK, Chandrashekar K, Thakur N, Verma PC, Borgis JF, et al. (2011) RNA interference for the control of whiteflies (Bemisia tabaci) by oral route. J Biosci 36: 153–161.

SKU is thankful to Department of Science and Technology, India for DST-INSPIRE faculty fellowship. JK and HS are grateful to CSIR for senior research fellowship.

Author Contributions

Conceived and designed the experiments: SKU KC. Performed the experiments: SKU SS HS JK. Analyzed the data: SKU SS KS PCV. Contributed reagents/materials/analysis tools: SKU KC PCV. Wrote the paper: SKU HS PCV KC. Arranged for the sequencing of clones: SKU KC SJ HK HS. Insured the availability of instruments: SKU KC SD JK HS.
49. Shakesby AJ, Wallace IS, Isaacs HV, Pritchard J, Roberts DM, et al. (2009) A water-specific aquaporin involved in aphid osmoregulation. Insect Biochem Mol Biol 39: 1–10.

50. Mutti NS, Park Y, Reese JC, Reeck GR (2006) RNAi knockdown of a salivary transcript leading to lethality in the pea aphid, *Acyrthosiphon pisum*. J Insect Sci 6: 1–7.

51. Pitino M, Coleman AD, Maffei ME, Ridout CJ, Hogenhout SA (2011) Silencing of aphid genes by dsRNA feeding from plants. PLoS ONE 6: e25709.

52. Gu L, Knipple DC (2013) Recent advances in RNA interference research in insects: Implications for future insect pest management strategies. Crop Prot 45: 36–40.

53. Xu W, Han Z (2008) Cloning and phylogenetic analysis of sid-1-like genes from aphids. J Insect Sci 8:30.