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In Silico Identification of Sugarcane Genome-Encoded MicroRNAs Targeting Sugarcane Mosaic Virus

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Abstract: Sugarcane mosaic virus (SCMV) (genus, *Potyvirus*; family, *Potyviridae*) is widespread, deleterious, and the most damaging pathogen of sugarcane (*Saccharum officinarum* L. and *Saccharum* spp.) that causes a substantial barrier to producing high sugarcane earnings. Sugarcane mosaic disease (SCMD) is caused by a single or compound infection of SCMV disseminated by several aphid vectors in a non-persistent manner. SCMV has flexuous filamentous particle of 700–750 nm long, which encapsidated in a positive-sense, single-stranded RNA molecule of 9575 nucleotides. RNA interference (RNAi)-mediated antiviral innate immunity is an evolutionarily conserved key biological process in eukaryotes and has evolved as an antiviral defense system to interfere with viral genomes for controlling infections in plants. The current study aims to analyze sugarcane (*S. officinarum* L and *S. spp.*) locus-derived microRNAs (sof-miRNAs/ssp-miRNAs) with predicted potential for targeting the SCMV +ssRNA-encoded mRNAs, using a predictive approach that involves five algorithms. The ultimate goal of this research is to mobilize the in silico-predicted endogenous sof-miRNAs/ssp-miRNAs to experimentally trigger the catalytic RNAi pathway and generate sugarcane cultivars to evaluate the potential antiviral resistance surveillance ability and capacity for SCMV. Experimentally validated mature sugarcane (*S. officinarum*, 2n = 8X = 80) and (*S. spp.*, 2n = 100–120) sof-miRNA/ssp-miRNA sequences (n = 28) were downloaded from the miRBase database and aligned with the SCMV genome (KY548506). Among the 28 targeted mature locus-derived sof-miRNAs/ssp-miRNAs evaluated, one sugarcane miRNA homolog, sof-miR159c, was identified to have a predicted miRNA binding site, at nucleotide position 3847 of the SCMV genome targeting CI ORF. To verify the accuracy of the target prediction accuracy and to determine whether the sugarcane sof-miRNA/ssp-miRNA could bind the predicted SCMV mRNA target(s), we constructed an integrated Circos plot. A genome-wide in silico-predicted miRNA-mediated target gene regulatory network was implicated to validate interactions necessary to warrant in vivo analysis. The current work provides valuable computational evidence for the generation of SCMV-resistant sugarcane cultivars.

Keywords: *potyvirus*; in silico tools; sugarcane mosaic virus; miRNA; RNA interference

1. Introduction

Sugarcane (*Saccharum officinarum*) is a prolific tropical and subtropical crop that is economically important, has a long life span, serves as a biofuel, is enriched with energy-rich
roughage, and is also a source of agroindustrial residues [1–3]. The genome of octaploid sugarcane (\(S. \text{ officinarum}\) \(2n = 80; x = 10\)) [4,5], also known as “noble” sugarcane, and the genome of sugarcane species and cultivars have been assembled, drafted, and resequenced [6–11]. Sugarcane mosaic virus (SCMV) is a highly transmissible and pathogenic potyvirus that causes sugarcane mosaic virus disease (SCMD) [12,13]. Potyviruses are spread by a common complex of sap-sucking vectors such as aphid species [14]. Innovative approaches are still needed to increase sugarcane productivity [15]. The genome of SCMV consists of a +ss RNA molecule with a length of 9575 nucleotides encoding a single large polyprotein. The genome polyprotein precursor was predicted to be cleaved resulting in ten functional proteins: P1, HC-Pro, P3, 6 K1, CI, 6 K2, VPg, Nla, Nib, and CP [16–19].

In plants, microRNAs (miRNA) are endogenously expressed small (19–25 nucleotides), evolutionarily conserved, non-coding (NC)-ss RNA molecules [20]. In higher plants, the biogenesis and transcription of the miRNA gene (MIR) is controlled by RNA polymerase II, which is then transcribed into single-standard polycistronic primary transcripts (pri-miRNAs). They control a variety of biological processes in plants by regulating gene expression, cell growth, development, differentiation, and host–virus interactions [21,22]. The miRNA-mediated RNAi is a post-transcriptional gene-silencing mechanism that provides antimicrobial innate immunity and regulates host–virus interactions to limit or inhibit viral infection [23].

Artificial miRNA-mediated (amiRNA) technology is an alternative, robust biotechnology based on engineering miRNA genes to control viral infections in plants [24]. RNAi-based amiRNA constructs have been used in research to induce antiviral resistance in plants against plant viruses such as tomato [25,26], cucumber [27], rice [28], and cotton [29]. Mature miRNAs in the sugarcane genome have been predicted, identified, isolated, analyzed, and validated to evaluate host–virus interactions and gene regulation, and they have been associated with abiotic and biotic stresses [30–40]. Recently, experimental validations of 35 conserved mature locus-derived high-confidence sof-miRNAs/ssp-miRNAs in the sugarcane genome and further depositions in the miRBase database were reported.

An integrative multi-network approach based on SCMV infection assessment was used to identify target binding sites of sugarcane genome-encoded sof-miRNAs/ssp-miRNAs in the SCMV genome. The identification of multiple host-derived miRNA binding sites in the SCMV genome for the creation of transgenic sugarcane varieties resistant to SCMV is the main objective of this study. In this study, several miRNA prediction tools were evaluated and used to identify microRNA–mRNA binding sites in the SCMV genome for use in developing transgenic or non-transgenic modified sugarcane plants with resistance to SCMV and, potentially, closely related potyviruses. Potential targets of the most promising sugarcane miRNAs for breeding were also of interest to better understand potyvirus–sugarcane plant interactions during infection. Until now, there have been no reports on the use of an amiRNA-based strategy to develop SCMV tolerance in sugarcane plants, based on the prediction of homologous amiRNAs for silencing SCMV. The predicted locus-derived sof-miRNAs/ssp-miRNAs in the sugarcane genome were further evaluated to understand the complex interactions between sugarcane host planta and SCMV potyviruses and to identify novel antiviral targets.

2. Materials and Methods
2.1. Sugarcane MicroRNAs and SCMV Genome Data Retrieval and Processing

Experimentally validated high-confidence mature sugarcane microRNAs (sof-miRNA1 56-sof-miR11892/ssp-miR156-ssp-1432) (Accession ID: MIMAT0001656-MIMAT0001671/ MIMAT0020291-MIMAT0020290) and Saccharum sp.-microRNAs (ssp-miR166-ssp-miR1432) (Accession ID: MIMAT0030451-MIMAT0020290) (Table S1) were retrieved from the miRNA registry (miRBase, version 22) [41]. The full-length SCMV +ssRNA genome sequence (9575 bases) (Accession number KY548506) was acquired from the NCBI GenBank database [42].
2.2. Potential Targets of Sugarcane MicroRNAs in the SCMV Genome

The prediction of effective microRNA–mRNA binding sites is a first step toward understanding microRNA-regulated gene regulatory networks. The accuracy of miRNA target site prediction can be affected by several factors, such as the specificity and sensitivity of the algorithm, the choice of reference sequence, and the length of the target sequence. Various in silico methods for effective silencing have been developed for the computational prediction of miRNA–mRNA target sites. A computational approach refers to the use of multiple computational methods, algorithms, or tools to analyze and interpret biological data. This approach combines different types of publicly available in silico algorithms, including miRanda [43,44], RNA22 [45,46], TAPIR [47], psRNATarget [48,49], and RNAhybrid [50] (Table 1).

Table 1. Different features and parameters of algorithms applied for miRNA target predictions.

<table>
<thead>
<tr>
<th>Algorithms</th>
<th>Features</th>
<th>Organisms</th>
<th>Parameters</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>miRanda</td>
<td>Seed-based interaction, multiple target sites, free energy of miRNA–mRNA duplex, conservation</td>
<td>Human, rat, fly, and worm</td>
<td>Score threshold = 140, Free energy = −15 Kcal/mol, Gap Open penalty = −9.00, Gap Extend penalty = −4.00</td>
<td><a href="http://www.microrna.org/">http://www.microrna.org/</a> (accessed on 14 August 2019)</td>
</tr>
<tr>
<td>RNA22</td>
<td>Pattern recognition, folding energy, heteroduplex,</td>
<td>Human, mouse, fly, and worm</td>
<td>Number of paired-up bases = 12, Sensitivity (63%), Specificity (61%), Folding energy = −14 Kcal/mol</td>
<td><a href="https://cm.jefferson.edu/rna22/Interactive/">https://cm.jefferson.edu/rna22/Interactive/</a> (accessed on 22 June 2019)</td>
</tr>
<tr>
<td>TAPIR</td>
<td>Sees pairing, target site accessibility, multiple sites</td>
<td>Plants</td>
<td>Free energy ratio = 0.2, Score = 9</td>
<td><a href="http://bioinformatics.psb.ugent.be/webtools/tapir">http://bioinformatics.psb.ugent.be/webtools/tapir</a> (accessed on 25 June 2021)</td>
</tr>
<tr>
<td>psRNATarget</td>
<td>Complementarity scoring, multiple target sites, translation inhibition</td>
<td>Plants</td>
<td>Expectation score = 7, Penalty for G:U pair = 0.5, HSP size = 19, Penalty for opening gap = 2</td>
<td><a href="https://www.zhaolab.org/psRNATarget/analysis?function=2">https://www.zhaolab.org/psRNATarget/analysis?function=2</a> (accessed on 26 May 2022)</td>
</tr>
<tr>
<td>RNAhybrid</td>
<td>Seed pairing and free energy</td>
<td>Any</td>
<td>Free energy = −20 Kcal/mol, Hit per target = 1</td>
<td><a href="http://bibiserv.techfak.uni-bielefeld.de/rnahybrid">http://bibiserv.techfak.uni-bielefeld.de/rnahybrid</a> (accessed on 26 May 2022)</td>
</tr>
</tbody>
</table>

2.3. miRanda

MiRanda is one of the first miRNA target predictors, a highly versatile algorithm based on the seed-based interactions of miRNA target duplexes [43]. It was implemented as a standard tool to detect potential miRNA binding sites. RNA–RNA duplex dimerization and sequence complementarity are features considered by the miRanda algorithm. It considers the cross-species conservation of target sites, which distinguishes it from other algorithms [44]. The miRanda algorithm has been implemented in C, and its first version was published in 2003. The default parameters were selected for the analysis (Table 1).

2.4. RNA22

The RNA22 algorithm has a diverse, web-based application with implemented interactive exploration. It uses a pattern-recognition based approach to serve as a miRNA target discovery tool. It predicts statistically significant target patterns using maximum folding energy (MFE) [45,46]. Site complementarity and non-seed-based interactions are important
features. Its prediction is also based on highly sensitive and significant target patterns. The default parameters were chosen (Table 1).

2.5. TAPIR

The TAPIR algorithm is used to assess the seed-based interactions of plant miRNAs in the target sequence. It is a highly precise plant miRNA target prediction algorithm used to detect target binding sites in the target sequence. It is used to deliver precise miRNA target predictions, including target mimics, with FASTA and RNAhybrid search options [47]. The default parameters were chosen (Table 1).

2.6. psRNATarget

The psRNATarget algorithm is a highly sensitive, newly designed web-based tool developed for plant miRNA prediction. The target binding sites of plant miRNAs are predicted based on complementary scoring schema. The algorithm predicts the inhibition pattern of cleavage [48,49]. The default parameters were chosen (Table 1).

2.7. RNAhybrid

The RNAhybrid is a seed-based scanning algorithm based on intermolecular hybridization used to predict effective binding sites of miRNAs in the target sequence. It predicts target binding sites in a very easy and flexible manner [50]. It is an online available tool. It is used for the rapid prediction of miRNA targets based on the MFE hybridization of mRNAs and miRNAs. The default parameters were chosen (Table 1).

2.8. RNAfold

The RNAfold algorithm is available on a web server implemented in the ViennaRNA package [51].

2.9. Statistical Analysis

The miRNA–mRNA target prediction biological data were further processed. Graphical representations of the miRNA data were prepared using the R language [52].

3. Results

3.1. Prediction and Analysis of Sugarcane MicroRNAs Targeting SCMV Genome

An integrative computational approach for identifying the possible interactions of high-confidence target sites of mature sugarcane miRNAs located in the SCMV positive-sense single-stranded (+ssRNA) genome from among the 28 sugarcane miRNAs (sof-miRNAs/ssp-miRNAs) revealed sof-miRNA/ssp-miRNA-derived MIR genes at a high proportion of sugarcane miRNA gene loci [33,53–56]. The predicted SCMV +ssRNA-encoded mRNA sequences were localized hypothetically by the sugarcane locus-derived sof-miRNA/ssp-miRNAs based on the miRanda algorithm predicted 19 miRNA-mRNA target pairs and RNA22: 15 sugarcane sof-miRNAs/ssp-miRNAs and 20 binding sites i. The TAPIR identified seven binding sites of mature sugarcane locus-derived sof-miRNA/ssp-miRNA target pairs. In total, 16 sugarcane miRNAs targeting 30 cleavable attachments sites in the SCMV genome were identified by the psRNATarget algorithm. RNAhybrid predicted 28 high-probability binding sites of sugarcane miRNAs in the SCMV genomic RNA sequence (Figures 1 and 2, File S1, Table S2).
Figure 1. Five-set Venn diagram representing mutually common binding sites of mature sugarcane miRNAs predicted to potentially target the SCMV genome. The in silico prediction was established using computational tools (miRanda, RNA22, TAPIR, psRNATarget, and RNAhybrid) to identify potential targets of sugarcane-encoded miRNAs. The areas of overlap among computational tools show miRNA binding sites. The high-order intersection of five algorithms revealed the most potent mature sugarcane miRNA—ssp-miR1444c-3p.

3.2. Sugarcane miRNAs Targeting P1

The potyviral first protease protein P1 was encoded by P1 ORF (149–847) (698 bases). The miRanda and RNA22 algorithms predicted the bindings of two sof-miRNAs: sof-miR168 (a, b) at nucleotide positions 547 and 846, respectively, as shown in (Figure 2A,B). The sof-miRNA168a was targeted at nucleotide position 406, using the TAPIR algorithm (Figure 2C). No sof-miRNA/ssp-miRNA was predicted to target the P1 region, by the psRNATarget and RNAhybrid algorithms (Figure 2D,E) (File S1) (Tables S2 and S3).
Figure 2. Individual sugarcane sof-miRNAs/ssp-miRNAs and their predicted high-confidence binding sites in the SCMV genome were predicted based on the ‘five algorithms’ approach. (A) miRNA sites were detected by miRanda. (B) Several miRNA target sites were detected by RNA22. (C) TAPIR identified sugarcane miRNA binding sites. (D) psRNATarget predicted several binding sites of sugarcane miRNAs. (E) The prediction of miRNA sites by RNAhybrid. (F) Union plot representing all predicted binding sites detected by all the algorithms used. Multiple copies of miRNA target binding sites are represented by colored dots. Targeted genes of SCMV are indicated by different colors.

3.3. Sugarcane miRNAs Targeting HC-Pro

The HC-Pro ORF (848–2227 nucleotides) encodes. The miRanda and RNA22 algorithms predicted the target site of sof-miR168 (a, b) at nucleotide position 1827 and sof-miR168a at nucleotide position 1296 (Figure 2A,B).

TAPIR predicted the attachment site of sof-miRNA159e at locus 1159 (Figure 2C). The psRNATarget algorithm detected the binding of ssp-miR444 (a, b, c-3p) at nt positions 1058 and 1763 (Figure 2D). The RNAhybrid algorithm predicted seven sof-miRNAs/ssp-
miRNAs: sof-miR159c, sof-miR168 (a, b), ssp-miR444 (a, b, c-3p), and ssp-miR1432 at nucleotide coordinates 1830, 1296, 1818, 1057, and 1316, respectively (Figure 2E, File S1, Tables S2 and S3).

3.4. Sugarcane miRNAs Targeting P3

The P3 ORF (2228–3268 nt) encodes a membrane-associated P3 protein which is required for SCMV genomic RNA replication, potential cell-to-cell spread (movement and transport) and is also responsible for determining host-range and symptoms [57–59]. The miRanda algorithm predicted the binding of sof-miRNAs: sof-miR168 (a, b) at nucleotide position 2562 (Figure 2A). No sof-miRNA/ssp-miRNA was predicted for targeting the P3 region by the RNA22 and TAPIR algorithms (Figure 2B,C). The potential target sites of sof-miR167 (a, b), sof-miR168a, ssp-miR437c, and ssp-miR444 (a, b, c-3p) at nucleotide positions 2427, 2971, 2981, and 2367, respectively, were detected by the psRNATarget algorithm (Figure 2D). In addition, RNAhybrid identified sugarcane sof-miRNAs/ssp-miRNAs; sof-miR167 (a, b) and ssp-miR437b at nucleotide positions 2699 and 2416, respectively (Figure 2E, File S1, Tables S2 and S3).

3.5. Sugarcane miRNAs Targeting 6K1

The 6K1 ORF (3269–3469 nucleotides) encode a 6K1 protein that functions in viral genome replication. It mediates cell-to-cell movement, controlling defense mechanism and gene regulation. It is a key component of the 6K2-induced viral replication complex (VRC) and regulation [60,61]. The 6K1 had the least number of predicted sugarcane sof-miRNAs/ssp-miRNAs. The ssp-miR444c-3p was predicted to optimally target 6K1 at nucleotide position 3441, according to the psRNATarget algorithm (Figure 2D, File S1, Tables S2 and S3).

3.6. Sugarcane miRNAs Targeting CI

The CI ORF (3470–5383 nt) encodes a multifunctional cylindrical inclusion protein (CI) essential for ATP binding and RNA helicase activity [62–64]. CI was targeted by two miRNAs: sof-miR396 and ssp-miR166 at nt positions 3634 and 4178, respectively, as indicated by the miRanda algorithm (Figure 2A). The RNA22 algorithm predicted two miRNAs: sof-miR159c and ssp-miR444b at nt positions 3730 and 5311, respectively, (Figure 2B). In addition, TAPIR predicted three sugarcane miRNAs: sof-miR159c, ssp-miR437a, and ssp-miR1128 at nucleotide positions 3847, 4869, and 4534, respectively (Figure 2C). The psRNATarget algorithm identified seven miRNAs: sof-miR159 (a, b, c, d, e), ssp-miR444b, and ssp-miR1432 at nt positions 3847, 3992, and 3980, respectively (Figure 2D). Five miRNA-binding sites were detected by RNAhybrid: sof-miR396 (start site 5016), sof-miR480e (3633), ssp-miR166 (3714), ssp-miR437a (4868), and ssp-miR1128 (4533) (Figure 2E, File S1, Tables S2 and S3).

3.7. Sugarcane miRNAs Targeting 6K2

Potyvirus 6K2 (5384–5542 nt) encodes the multifunctional protein 6K2, induces the formation of RE-derived complexes, and develops resistance to drought [65,66]. The RNA22 algorithm identified five sugarcane sof-miRNAs: sof-miR408 (a, b, c, d, and e) at locus position 5538 (Figure 2B, File S1, Tables S2 and S3).

3.8. Sugarcane miRNAs Targeting Nla-VPg

Potyvirus Nla-VPg ORF (5543–6109 nt) encodes a viral genome-linked protein (VPg) that functions as a virulence determinant and genome translator [67–71]. It is also involved in replication, translation, and movement [72–74]. The RNA22 and TAPIR algorithms predicted the binding of ssp-miR444c-3p at locus position 5552 (Figure 2B,C). The psRNATarget algorithm predicted six miRNAs: sof-miR156, sof-miR159 (a, b, c, d), and ssp-miR444c-3p (Figure 2D). No sugarcane sof-miRNA/ssp-miRNA was predicted to target the Nla-VPg region using the RNAhybrid algorithm (Figure 2E, File S1, Tables S2 and S3).
3.9. Sugarcane miRNAs Targeting Nla

Potyvirus Nla ORF (6110–6835 nt) encodes nuclear inclusion protein a (Nla), which is involved in RNA binding and also interacts with Nlb [75,76]. miRanda, RNA22, and RNAhybrid predicted the binding of only one sugarcane miRNA: ssp-miR528, sof-miR396, and ssp-miR827 at nucleotide positions 6376, 6821, and 6338, respectively (Figure 2A,B,E). The psRNATarget algorithm identified three sugarcane miRNAs: sof-miR408e and ssp-miR444 (a, b) at nucleotide positions 6544 and 6641, respectively (Figure 2D). No miRNA target pair was identified to target Nla by the TAPIR algorithm (Figure 2C, File S1, Tables S2 and S3).

3.10. Sugarcane miRNAs Targeting Nlb

Potyvirus Nlb ORF (6836–8398) encodes nuclear inclusion protein b (Nlb), which is involved in translocation activity and also interacts with Nla [77]. It contains nuclear signals and is also referred to as RdRp [78]. The miRanda algorithm detected the binding of two sugarcane ssp-miRNAs: ssp-miR169 and ssp-miR1432 at nucleotide positions 7798 and 7523, respectively (Figure 2A). The psRNATarget algorithm predicted the binding of two sugarcane ssp-miRNAs: sof-miR396 and ssp-miR444b at nucleotide positions 7798 and 7523, respectively (Figure 2D). No miRNA-target pair was identified based on the RNA22, TAPIR, and RNAhybrid algorithms (Figure 2B,C,E, File S1, Tables S2 and S3).

3.10.1. Sugarcane miRNAs Targeting CP

Potyvirus CP ORF (8399–9337) encodes a multitasking coat protein (CP), which is involved in the development of virion assembly. The CP is involved in all steps of the potyviral life cycle [79–81]. The miRanda algorithm predicted the binding of three sugarcane ssp-miRNAs (ssp-miR444 (a, b, c-3p) start site 8501). ssp-miR444c-3p also targeted the CP region at nucleotide position 9268 (Figure 2A). The RNA22 algorithm predicted the binding of the ssp-miRNA444 family at nt positions 8502 and 9181 (Figure 2B). The psRNA1Target algorithm predicted the binding of ssp-miR444c-3p at nt position 9282 (Figure 2D). Potential binding sites of sugarcane miRNAs, sof-miR159 (a, b, d, e), sof-miR408 (a, b, c, d), and ssp-miR169, were detected by the RNAhybrid algorithm at nucleotide positions 8953, 8355, and 8458, respectively (Figure 2E, File S1, Tables S2 and S3).

3.10.2. Sugarcane miRNAs Targeting UTR

The potyvirus 5′ untranslated region (5′ UTR) (1–148 nt) and 3′ UTR (9341–9575 nt) are involved in the replication and translational activities of the ORFs [82,83]. The sof-miR408 (a, b, c, d) was predicted to target the 5′ UTR at nt positions 139 by miRanda (Figure 2A). Similarly, ssp-miR528 was identified to target the 5′ UTR at nt position 122 by TAPIR and RNAhybrid (Figure 2C,E). RNA22 predicted the binding of sof-miR168 (a, b) at nt position 9520 in the 3′ UTR (Figure 2B). RNAhybrid predicted the binding of two sugarcane miRNAs in the 3′ UTR: sof-miR156 and ssp-miR437c at nt positions 9402 and 9395, respectively (Figure 2E, File S1, Tables S2 and S3).

3.11. Identification of Consensual Sugarcane MicroRNAs

The present study was concluded based on the consensus of the genomic target binding sites of sugarcane miRNAs determined by different algorithms. Among them, we selected nine sugarcane miRNAs (sof-miR159c, sof-miR168a, ssp-miR437a, ssp-miR528, ssp-miR444 (a, b), ssp-miR444c-3p), (ssp-miR1128, and ssp-miR1432), which were based on the consensus genomic positions 3847 (target gene CI), 1296 (HC-Pro), 4869 (CI), 122 (5′ UTR), 8502/1058 (CP/HC-Pro), 5583 (Nla-Vpg), 4534 (CI), and 1316 (HC-Pro) detected (Tables 2 and 3). Of the nine consensus locus-derived sof-miRNAs/ssp-miRNAs in the sugarcane genome investigated in this study, only one sof-miRNA (sof-miR159c at nt position 3847 targeting CI) was identified by the union of genomic consensus positions by at least three algorithms (RNA22, TAPIR, and psRNA1Target) (Figure 3, Tables 2 and 3, File S1, Tables S2 and S3).
Table 2. Predicted high-confidence binding sites of consensus sugarcane miRNAs targeting the SCMV genome detected by different computational algorithms.

<table>
<thead>
<tr>
<th>Sugarcane miRNA</th>
<th>Position miRanda</th>
<th>Position RNA22</th>
<th>Position TAPIR</th>
<th>Position psRNATarget</th>
<th>Position psRNAhybrid</th>
<th>MFE * psRNATarget</th>
<th>MFE ** RNA22</th>
<th>MFE Ratio TAPIR</th>
<th>Expectation psRNAhybrid</th>
<th>MFE * RNAhybrid</th>
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<tbody>
<tr>
<td>sof-miR159c</td>
<td>3847</td>
<td>3847</td>
<td>3847</td>
<td>−</td>
<td>−18.00</td>
<td>0.58</td>
<td>5.50</td>
<td>−25.80</td>
<td>−25.80</td>
<td>−21.99</td>
</tr>
<tr>
<td>sof-miR168a</td>
<td>1296</td>
<td>1296</td>
<td>1296</td>
<td>−</td>
<td>−18.70</td>
<td>0.69</td>
<td>2.20</td>
<td>−20.80</td>
<td>−20.80</td>
<td>−19.99</td>
</tr>
<tr>
<td>ssp-miR437a</td>
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<td>4868</td>
<td>4868</td>
<td>−</td>
<td>−18.00</td>
<td>0.60</td>
<td>2.40</td>
<td>−19.60</td>
<td>−19.60</td>
<td>−19.60</td>
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<tr>
<td>ssp-miR528</td>
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<td>121</td>
<td>121</td>
<td>−</td>
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<td>0.69</td>
<td>2.29</td>
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<tr>
<td>ssp-miR444a</td>
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<td>2.70</td>
<td>−20.70</td>
<td>−20.70</td>
<td>−20.70</td>
</tr>
</tbody>
</table>

* MFE: minimum free energy measured in Kcal/ml. ** MFE: maximum folding energy for heteroduplex measured in Kcal/mol.

Table 3. Predicted consensus sugarcane-encoded miRNA target sites localized in the different target genes of SCMV-SO.

<table>
<thead>
<tr>
<th>miRNA ID</th>
<th>Accession ID</th>
<th>Mature Sequence (5′–3′)</th>
<th>Target Genes ORF(s)</th>
<th>Target Binding Locus Position</th>
</tr>
</thead>
<tbody>
<tr>
<td>sof-miR159c</td>
<td>MIMAT0001662</td>
<td>CUUGGAUUGAGGAGCUCUCU</td>
<td>CI</td>
<td>3847–3868</td>
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<tr>
<td>sof-miR168a</td>
<td>MIMAT0001665</td>
<td>UCAGUUGUGUGACAGUGGAC</td>
<td>HC-Pro</td>
<td>1296–1317</td>
</tr>
<tr>
<td>ssp-miR437a</td>
<td>MIMAT0020280</td>
<td>AAAGUAGAGAGAGUAGAGUU</td>
<td>CI</td>
<td>4869–4890</td>
</tr>
<tr>
<td>ssp-miR528</td>
<td>MIMAT0020288</td>
<td>UGGAAAGGAGAGAUGGAGAGAGA</td>
<td>5′UTR</td>
<td>122–143</td>
</tr>
<tr>
<td>ssp-miR444a (1)</td>
<td>MIMAT0020284</td>
<td>UGCAGUUGUGGGCUAAGCUU</td>
<td>CP</td>
<td>8501–8521</td>
</tr>
<tr>
<td>ssp-miR444b (1)</td>
<td>MIMAT0020285</td>
<td>UGCAGUUGUGGGCUAAGCUU</td>
<td>CP</td>
<td>8501–8521</td>
</tr>
<tr>
<td>ssp-miR444c-3p</td>
<td>MIMAT0020286</td>
<td>UGCAGUUGUGGGCUAAGCUU</td>
<td>Nla-VPg</td>
<td>5583–5604</td>
</tr>
<tr>
<td>ssp-miR1128</td>
<td>MIMAT0020289</td>
<td>UACUACUCCUCCUCUGCCAAA</td>
<td>CI</td>
<td>4534–4555</td>
</tr>
</tbody>
</table>

Figure 3. Intersection plot shows the consensus high-confidence binding sites of mature sugarcane miRNAs predicted by at least two computational tools. The colored dots represent sugarcane miRNA binding sites targeting different genes of SCMV.
3.12. Identification of the miRNA–mRNA Regulatory Network

A Circos plot represents the predicted host–virus interactions of sugarcane miRNAs and SCMV target genes. A Circos plot was generated to visualize a comprehensive master miRNA regulatory network with novel antiviral targets (Figure 4). The generation of the miRNA–mRNA regulatory network was conducted using ‘Circos’ software [84].

![Circos plot](image)

**Figure 4.** Integrated Circos plot shows multiple targets of sugarcane-encoded miRNAs. The colored connection lines are targeted genes (ORFs) in the SCMV genome. Construction, exploration, target predictions, and interactions between the sugarcane miRNAs and SCMV genes are mapped.

3.13. RNA Secondary Structures

The computationally predicted locus-derived mature miRNAs in the sugarcane genome were analyzed by generating their secondary structures using the original precursor sequences. Pre-miRNA hairpin sequences were used for manual curation. The main parameters of the predicted stable secondary structures were evaluated (Table 4). The stable secondary structures of the potential consensus sugarcane precursor sequences were predicted by the RNAfold algorithm [51].
Table 4. Features of the predicted precursors of sugarcane were determined.

<table>
<thead>
<tr>
<th>miRNA ID</th>
<th>Accession ID</th>
<th>MFE */ Kcal/mol</th>
<th>AMFE **</th>
<th>MFEI ***</th>
<th>(G + C)%</th>
</tr>
</thead>
<tbody>
<tr>
<td>sof-MIR159c</td>
<td>MI0001760</td>
<td>−110.60</td>
<td>−46.47</td>
<td>−0.87</td>
<td>53.36</td>
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<tr>
<td>sof-MIR168a</td>
<td>MI0001763</td>
<td>−66.20</td>
<td>−63.65</td>
<td>−0.83</td>
<td>75.96</td>
</tr>
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<td>ssp-MIR437a</td>
<td>MI0001763</td>
<td>−57.10</td>
<td>−32.62</td>
<td>−1.29</td>
<td>25.14</td>
</tr>
<tr>
<td>ssp-MIR528</td>
<td>MI0001763</td>
<td>−48.50</td>
<td>−52.71</td>
<td>−0.86</td>
<td>60.84</td>
</tr>
<tr>
<td>ssp-MIR444a</td>
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<td>−57.70</td>
<td>−54.94</td>
<td>−1.28</td>
<td>42.86</td>
</tr>
<tr>
<td>ssp-MIR444b</td>
<td>MI0001763</td>
<td>−63.70</td>
<td>−60.09</td>
<td>−1.38</td>
<td>43.39</td>
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<tr>
<td>ssp-MIR444c</td>
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<td>−57.22</td>
<td>−1.31</td>
<td>43.52</td>
</tr>
<tr>
<td>ssp-MIR1128</td>
<td>MI0001763</td>
<td>−101.70</td>
<td>−36.98</td>
<td>−1.18</td>
<td>31.27</td>
</tr>
<tr>
<td>ssp-MIR1432</td>
<td>MI0001763</td>
<td>−57.10</td>
<td>−64.88</td>
<td>−1.14</td>
<td>56.82</td>
</tr>
</tbody>
</table>

* MFE is minimum free energy. ** AMFE represents adjusted minimum free energy. *** MFEI defines as minimum free energy index.

4. Discussion

The SCMV is a monopartite potyvirus suspected as an etiological agent that has spread to Pakistan and China due to its high transmissibility and has become an increasingly potential long-lasting threat to sugarcane and maize production in the last two decades [13,17,85]. In our previous studies, we have investigated experimentally validated mature locus-derived microRNAs in the sugarcane genome, which were predicted to be targets of SCBGAV, SCYLV, and SCBV based on in silico criteria [37–39]. Several studies have identified complex host–virus interactions and have investigated miRNAs targeting plant viruses using an in silico approach [86–92]. miRNAs have emerged as novel endogenous targets for multiple levels of miRNA gene-level regulation [53,93,94]. Several studies have shown that the efficacy of amiRNA-based RNA interference leads to specific gene silencing in transgenic crops to reduce host plant virus infection [27,28,95–97]. In this computational research, mature sugarcane sof-miRNAs/ssp-miRNAs were aligned with the genomic sequence of the SCMV target to identify miRNA–mRNA binding sites hypothesized to understand complex host–virus specific interactions with the P1, HC-Pro, P3, 6K1, CI, 6K2, NIa-VPg, NIa-Pro, NIb, and CP of SCMV. The P1 is the least-conserved hypervariable that modulates host responses and is essential for the replication of the viral +ssRNA genome [98,99]. Host adaptation is a key process for virus genome evolution [100,101]. P1 is also related to virus–host adaptation [102]. The HC-Pro is a multifunctional, non-structural dimeric helper component proteinase. It has been reported as a viral suppressor. HC-Pro is required to enhance expression via the fusion of P1, symptom development, and viral replication [103–108]. Until now, the potential for exploiting the regulation of sugarcane genome-encoded miRNA to abate infection by SCMV has not been investigated as a strategy for developing tolerant or resistant sugarcane cultivars. The results of this study provide the first computationally based evaluation of mature locus-derived miRNAs in the sugarcane plant genome to enable the prediction of effective miRNA binding sites and provide new tools for better understanding the molecular and omic interactions between sugarcane plant host cells and SCMV-encoded mRNAs/protein.

Based on our findings, the SCMV genome (HC-Pro, CI, NIa-VPg, and CP) is susceptible to nine consensus sugarcane miRNAs. We found that nine miRNAs could theoretically originate from the sugarcane genome (Tables 2 and 3). In silico tools, RNA22, TAPIR, and psRNATarget, identified a genomic consensus base pair complementarity in sof-miR159c at nucleotide position 3847 (Figure 2 and Table 2). The ssp-miR444c-3p was predicted by all five algorithms, making it the only unique sugarcane miRNA identified in this study (Figure 1 and Table S2). We identified the maximum folding energy of the consensus functional miRNA–mRNA target pair, which is −18.00 Kcal/mol, using RNA22. RNA22 is a highly sensitive algorithm that uses a pattern-based approach to target miRNAs. Using the psRNATarget algorithm, we estimated an expectation score of 5.50 for a consensus target pair (Table 2) [109]. The RNA22 and psRNATarget algorithms predicted target sites using a non-seed-based approach. Experimentally determining miRNA–mRNA interactions
can be expensive and time-consuming; making the accurate computational prediction of miRNA targets a high priority. The limitations and bottlenecks of existing algorithms and approaches are interpreted using the union and intersection level of the predictions in this study. The miRNA targeting relies on the base pairing of miRNA–mRNA targets [110]. These results suggest that the predicted consensus miRNA–mRNA duplex represents a ‘true target’. Our results indicate that sugarcane miRNAs likely play a role in the interaction between host and virus. Our results highlight the interaction of SCMV ss-RNA with the sugarcane miRNA target interaction network.

Potyvirus cylindrical inclusion helicase (CI) is required for the initiation of the viral replication mechanism, cell-to-cell movement, and plant–host, protein–virus interactions [62,63,111]. Computational predictions and analyses revealed that the sugarcane consensus sof-miR159c is a high-confidence target site potentially targeting the CI ORF (Table 3). The conserved precursor MIR159 is considered to be controlled by plant growth and fertility [112]. In our previous study, the consensus sof-miR159e (Accession ID: MIR-MAT0001661), predicted to have an effective target binding site at nucleotide position 5535 in the SCBV genome, was identified as the most effective miRNA by the miRanda, RNA22, and RNAhybrid algorithms.

While miRNA–mRNA target pair interactions between locus-derived miRNAs in the sugarcane genome and SCMV have been determined, the development of amiRNA-based constructs and further transformations in sugarcane to control SCMV are not fully understood. We have performed a comprehensive analysis of SCMD-associated Potyvirus for the first time, which is a first step toward the development of miRNA-based antiviral therapy. An amiRNA construct relies on the high-level specificity of a nucleotide base pairing to control deleterious off-target effects. The small size of amiRNA is a unique feature for the development of a single gene expression vector to control multiple potyviruses in transgenic sugarcane. This approach offers specificity and sensitivity and complements existing molecular approaches for analyzing targets for SCMV disease abatement. A number of environmental concerns have been raised regarding the large-scale use of virus-resistant transgenic plants [113–118]. As amiRNAs have high specificity to the designed target gene, detrimental off-target effects can be minimized, permitting their silencing expression to be stably transmitted to future generations [119–123]. The results indicate that the use of in silico tools provides better results than a single algorithm when developing amiRNA-based mdm-miRNA therapeutics to target SCMV and other plant viruses as well. Despite the frequent use of RNAi in biology and agriculture, there are several drawbacks and challenges in designing efficient silencing constructs. Furthermore, the small size of amiRNA permits for the insertion of multiple and distinct amiRNAs within a single gene expression cassette, which can then be transformed to develop transgenic plant resistant to multiple viruses simultaneously [27,95,124]. The in silico analysis was designed for experimental validation to show whether these predicted miRNAs could make the plants resistant to SCMV. Future work will be focused on transiently expressing these miRNAs or injecting RNA hairpins in N. benthamiana to show its efficacy against SCMV.

5. Conclusions and Future Directions

The SCMV, which infects sugarcane crops worldwide, is the most damaging potyvirus pathogen associated with an ongoing SCMD epidemic that reduces yield in all sugarcane cultivars cultivated in China. This study involved in silico tools and approaches to characterize the target binding sites of mature sugarcane locus-derived miRNAs in the SCMV genome. Among the 28 sugarcane miRNAs from the miRBase database, only one, sof-miRNA (sof-miR159c), was identified as the most effective, naturally occurring sof-miRNA biomolecule for targeting the SCMV genome (nucleotide 3847 onward), based on the consensus of multiple algorithms used herein. This approach offers specificity and sensitivity and complements existing molecular approaches for analyzing targets for SCMV disease abatement. The current focus of attention is the development of SCMV-resistant sugarcane plants that abate the effects of SCMD.
Supplementary Materials: The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/microbiolres15010019/s1, Table S1: Mature sugarcane microRNA sequences used for the prediction of binding sites in the SCMV genome; Table S2: Identification of high-confidence binding sites of sugarcane miRNAs in the SCMV; Table S3: Gene-wise prediction; File S1: Prediction results of computational tools.

Author Contributions: M.A.A., W.W. and S.Z. conceived the study. All authors analyzed the computational data. M.A.A., H.G., Z.I., W.u.Z., H.M. and M.Z. made the graphs. M.A.A., W.W. and H.G. wrote the original draft. All authors have read and agreed to the published version of the manuscript.

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