Advance towards Host Mediated RNA Interference Insect Pest Management

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Abstract

Gene suppression via RNA interference (RNAi) provides an alternative strategy for insect pest management. Insect pests cost billions of dollars in the form of crop losses and insecticides, and farmers face an ever-present threat of insecticide resistance, fueling a continual search for alternative pest-control strategies. Of late, novel insect pest management strategies like transgenic plants expressing insecticidal crystal protein genes from the soil bacterium, *Bacillus thuringiensis* (Bt) are effective in managing the insect pests that belong to *Lepidoptera* and *Coleoptera*. But success of the above technology is threatened by accelerated development of resistance, which is also true with chemical insecticides. Therefore, there is a need for identifying some additional effective pest management strategies, which could also augment integrated pest management (IPM). In this scenario, RNA interference (RNAi) offers a great deal of hope in successful mitigation of various insect pests. This review will feature late endeavors to comprehend the boundaries to RNA conveyance in headstrong insect species, depict late advances in the commercial development of insect-protected crops and biological insecticides using RNAi, and talk about this methodology with regards to an integrated pest management approach.

Keywords: RNAi; Pest control; IPM; Biotechnology

RNA Interference-Introduction and Mechanism

![RNA interference- Introduction and Mechanism.](image_url)
RNA interference (RNAi) is a potent method using only a few double stranded RNA (dsRNA) molecules per cell to silence the expression. RNA mediated silencing of specific gene(s) in pest insects through plant delivered RNA, offers the possibility to target genes necessary for their development, reproduction, or feeding success. Principally, this technology enables engineering of a new generation of pest-resistant GM crops. RNA interference (RNAi) is the specific downregulation of gene expression by double-stranded RNA (dsRNA) involving degradation of a target mRNA through the production of small interfering RNAs (siRNAs) from the dsRNA, which is cleaved by dsRNA-specific endonucleases referred to as dicers (Figure 1). One strand of the siRNA is assembled into an RNA-induced silencing complex (RISC) in conjunction with the Argonauts multi-domain protein, which contains an RNase H-like domain responsible for target degradation. The basis of this effect is thought to lie in the presence of an RNA-dependent RNA polymerase (RdRP) that is able to interact with the RISC complex and generate new dsRNA based on the partially degraded target template by using the hybridized siRNA strands as primers. The synthesized dsRNA is then acted on by the dicer enzymes to generate new siRNAs (secondary siRNAs), thus acting as an amplification step. In this way, once a dsRNA is introduced into a cell, its effect can persist over development; in addition, the ds RNAs can be exported to neighboring cells and thus spread the gene knockout effect through the organism [1].

**Mechanism of dsRNA Uptake**

**Transmembrane channel-mediated uptake mechanism**

The gene identified, systemic RNA interference deficient-1 (sid-1), is essential and sufficient to mediate systemic RNAi effect in *C. elegans* [5]. Two proteins involved. As (i) SID-2 modifies the SID-1 molecule to activate the transport, or (ii) It binds the dsRNA from the environment and delivers it to SID-1, or (iii) It induces the endocytosis pathway of the dsRNA, in which case SID-1 delivers the dsRNA to the cytoplasm [6].

**Endocytosis-mediated uptake mechanism**
Apart from genes involved in cell-autonomous RNAi, a gene with a role upstream of the intracellular silencing events was identified: the clathrin heavy chain gene. It is known as a component of the endocytosis machinery. The role of vacuolar H+ ATPase was confirmed by Saleh et al. [6], and their pharmacological tests also showed that a combination of scavenger receptors participated in dsRNA uptake. An active control role for vacuolar H+ ATPase is proposed in breaking the normal endocytosis pathway to induce RNAi silencing in cells (Figure 3).

**Link with immunity response**

Link between antiviral immunity and RNAi was recently demonstrated in *D. melanogaster*. In this model infected insect cells release viral dsRNA which is taken up by uninfected cells through the dsRNA uptake pathway. Subsequently, an antiviral RNAi response, limiting virus replication, is elicited in those cells. Therefore it was concluded that immunity in multicellular organisms requires both cell-autonomous and systemic mechanisms to create pre-existing immunity to protect uninfected cells [7].

**Factors influencing the silencing effect**

A. Concentration of dsRNA

B. Nucleotide sequence

C. Length of the dsRNA fragment

D. Persistence of the silencing effect

E. Life stage of the target organism

**Production of dsRNA from an RNAi vectors**

Vectors containing inverted repeats separated by an intron produce double-stranded RNA (dsRNA) or hairpin RNA (hpRNA) which effectively silences genes of interest. Producing dsRNA in RNase III deficient *E. coli* strains and extracting that product from RNase III deficient *E. coli* strains and extracting that product to treat plants as an alternative to RNAi transgenic plants has recently been accomplished. The key to an RNAi commercial product will be targeting genes that potently kill pests or inhibit toxin resistance [8]. Specific promoter, inverted repeats, and an intron spacer are used to produce the best RNAi effects. Insect systemic RNAi was first documented in another model insect *Tricholium castaneum* (hour beetle) and multiple genes such as *Tc-ASH*, Distalless, maxillopedia and proboscipedia were targeted by injection of specific dsRNA [9]. Turner offered experimental validation of this strategy on the larval stage of the light brown apple moth (*Epiphyas postvittana*). Transcript level of a larval gut-expressed gene (*EposCXE1*) and adult antennae expressed gene (*EposPBP1*) were reduced by feeding specific dsRNA [10]. Microinjection of dsRNA into adults and larvae has been used to silence genes in *Tricholium castaneum*. Similarly, dsRNA induced an RNAi response in Lepidopteran, such as *Spodoptera litura* (*Fabri cius*) (*Lepidoptera: Noctuidae*) and *Epiphyas postvittana*.

No effective Bt toxins are known against sap-sucking homopteran pests such as aphids, leafhoppers, etc (Figure 4).

**Delivery of dsRNA in plants**

**Biologist and agroinfection delivery**: When plant tissues are bombarded with gold or tungsten particles that have been coated with DNA or RNA, the nucleic acid is released and expressed in the cells where the particles come to rest. Bombarding cells with double-stranded (ds) RNA, siRNA, or DNA constructs that encode hairpin RNA can produce transient silencing of target reporter genes.

A. Virus induced delivery

B. Stable transformation with transgenes

**Application of RNAi in Pest Control**

A serious problem for insecticides is that they can kill non-targeted animals. To address this issue, the possibility of using RNAi to kill only the target animals by down-regulating essential gene functions in insects has been recognized for many years [1].

**Cell line experiments**

Cell line experiments can give additional information specific for the target gene in a simplified model. They can also inform on the possibility and effect of environmental RNAi when the dsRNA is applied through soaking of the cells in a dsRNA-enriched medium. This way the link between Cry1Ac insecticidal protein and an aminopeptidase N in the gut of larvae of the cotton bollworm (*Helicoverpa armigera*) (*HaAPN1*) was shown with RNAi cell line experiments. Sf21 cells were modified to express *HaAPN1* and 48h after *Haapn1* dsRNA treatment the expression of *Haapn1* mRNA and its protein were reduced 70% compared to the controls, resulting in decreased sensitivity of the *HaAPN1*-expressing cells to Cry1Ac protein [11].
ds RNA feeding and in planta experiments

Silencing a cotton bollworm, *P450* monooxygenase gene: Mao et al. [12] identified a *P450* monooxygenase from the cotton bollworm (*Helicoverpa armigera*), it was named *CYP6AE14* and was involved in detoxification of the otherwise toxic allelochemical an indiscriminately toxic compound called gossypol-produced by cotton plant. This gene is induced by gossypol and its suppression reduced the larval tolerance to gossypol. The researchers transferred hairpin RNA constructs directed against *CYP6AE14* into plant and fed cotton bollworm with the plant material. The model plants *Nicotiana tabacum* and *Arabidopsis thaliana* were modified with the cytochrome *P450* gene of *H. armigera*. When the cotton bollworm larvae were fed transgenic leaves, levels of cytochrome *P450* mRNA were reduced and larval growth retarded. As a glutathione-S-transferase gene (GST1) is silenced in GST1 dsRNA-expressing plants, feeding insects plant material expressing dsRNA may be a general strategy to trigger RNA interference and could find applications in entomological research and field control of insect pests.

Control of coleopteran pests through RNAi

Baum et al. [13] utilized a screening approach where genes from Western corn root worm were identified in cDNA libraries, and genes encoding polypeptides predicted to provide an essential biological function were classified as ‘targets’. dsRNAs directed against three target genes (*b-tubulin, V-ATPase A and V-ATPase E*) demonstrated an effective RNAi response in WCR that resulted in high larval mortality. Transgenic corn expressing one of these dsRNAs, targeted to a subunit of the midgut enzyme vacuolar ATPase (V-ATPase), showed protection against WCR infestation comparable to that provided by a *Bt* transgene. When dsRNAs designed to target WCR genes were tested in larvae from two other insect pests, mortality declined with decreasing sequence identity between the WCR genes and their orthologs in the other species, indicating that gene silencing was potentially very selective (Table 1) [13-22].

Table 1: Use of RNAi in insects with dsRNA being applied through feeding.

| Organism                  | Target Gene                  | Stage & Location      | Application Method    | mRNA Silencing            | Reference |
|---------------------------|------------------------------|-----------------------|-----------------------|---------------------------|-----------|
| Coleoptera Diabrotica virgifera | Vascular ATPase-Vacuolar ATPase subunit A & E Tubulin/Arginine kinase | Neonates Neonates Adults | Artificial diet Transgenic plant Artificial diet | Drastic reduction Gradual decrease Significant decrease | [13] |
| Diptera Glossina morsitans | Midgut protein               | Midgut                | Blood meal            | 30-55%                    | [15] |
| Hemiptera Acyrthosiphon pismus | Water specific aquaporin Nitropin 2 | 6 day aphids - Gut 2nd instar- Salivary glands | Artificial diet Artificial diet | 50% 54-59% | [16,17] |
| Lepidoptera Epiphysis postvittana | Larval gut Carboxylesterase | 3rd instars - Gut | Droplet               | 60%                        | [10] |
| Lepidoptera Helicoverpa armigera | Pheromone binding protein | 3rd instars Gut | Transgenic plant Decrease | [12] |
| Lepidoptera Plutella xylostella | Acylcholinesterase | Neonates Artificial diet | Drastic reduction | [18] |
| Lepidoptera Spodoptera frugiperda | Cytochrome P450 (CYPBG1) | Midgut + carcass Droplet | 98% | [19] |
| Hymenoptera Apis mellifera | Toll-related receptor Vitellogenin | Whole organism Fat body carcass | Soaking Natural diet | 60% 90% | [21,22] |
| Isoptera Reticulitermes flavipes | Cellulose enzyme Caste regulatory hemocerin storage protein | Saliva gland Workers Fat body | Artificial diet Artificial diet | 60% 50-70% | [14] |

RNAi for the control of whiteflies by oral route

RNA interference mediated gene silencing was explored for the control of sap-sucking pest *Bemisia tabaci*. dsRNAs and siRNAs were synthesized from five different genes, actin ortholog, ADP/ATP translocase, α-tubulin, ribosomal protein L9 (RPL9) and V-ATPase A subunit. A simplified insect bioassay method was developed for the delivery of ds/siRNA through the oral route, and efficacy was evaluated. ds/siRNA caused 29–97% mortality after 6 days of feeding. Knocking down the expression of RPL9 and V-ATPase A caused higher mortality with LC50 11.21 and 3.08μg/ml, respectively, as compared to other genes [23].

Other applications

RNAi-based strategy being explored to fight IAPV infection is to feed bees siRNAs targeting specific IAPV sequences such that, following viral entry into bee cells, translation of viral proteins is blocked [24]. Kumar et al. [18] studied the effect of siRNA on larval development by selective targeting of the
acetylcholinesterase (AChE) gene of *Helicoverpa armigera*. Chemically synthesized siRNA molecules were directly fed to *H. armigera* larvae along with the artificial diet. The siRNA treatment resulted in specific gene silencing of AChE and consequently brought about mortality. Growth inhibition of larvae, reduction in the pupal weight, malformation and drastically reduced fecundity as compared to control larvae. Using chitin synthase gene A (SeCHSA) as the target gene, which is expressed in the cuticle and tracheae of the lepidopteran pest *Spodoptera exigua*, showed that the growth and development of *S. exigua* larvae fed *Escherichia coli* expressing dsRNA of SeCHSA was disturbed, resulting in lethality [25]. Whyard et al. [26] harnessed the sequence specificity of RNAi to design orally-delivered dsRNAs that selectively killed target insects. They found that *D. melanogaster*, *T. castaneum*, pea aphids (*Acyrthosiphon pisum*), and tobacco hornworms (*Manduca sexta*) were selectively killed when fed species-specific dsRNA targeting vacuolar-type ATPase transcripts. For the aphid nymphs and beetle and moth larvae, dsRNA could simply be dissolved into their diets. However, to induce RNAi in the drosophilid species, the dsRNAs needed to be encapsulated in liposomes to help facilitate the uptake of the dsRNA [27]. Overcoming the specific delivery of dsRNA or siRNA into the cytoplasm of target cells is still an important issue for the use of RNAi for insecticides.

**Bio-safety**

A key advantage of RNAi-mediated resistance is that dsRNA has no inherent translational ability to produce a functional protein. This means that non-target effects should be minimal and lower than those of even highly specific transgenic proteins. Protection of non-target species that also feed on the plant can be provided by using dsRNA that has no effect when ingested by other invertebrates such as insects. One of the key advantages of this technology could be its high species specificity for the target pests. If the target sequence is carefully chosen, a specific gene or genes can be silenced. RNAi can also be used to achieve varying levels of gene silencing, using the same ihpRNA construct in different lines. This allows for selection of lines with varying degrees of gene silencing. In addition to this, the timing and extent of the gene silencing can be controlled, so that genes that are essential will only be silenced at chosen stages of growth or in chosen plant tissues. So, RNAi provides us with a great degree of flexibility in the field of functional genomics.

**Limitations**

There are also limitations however to RNAi. Unlike in insertional mutagenesis, for the use of RNAi the exact sequence of the target gene is required. Once this sequence information is available, the rest of the process is however relatively fast. Secondly, delivery methods for the dsRNA is a limiting step for the number of species which RNAi based approaches can be used easily. Due to this, improvement and further research into the kinds of vectors that can be used safely and reliably is needed. There have also been some reports that it has been difficult to detect mutants in which there has been subtle changes in gene expression. In plants, marker genes are being developed that will indicate if there has been a change in gene expression.

**Conclusion and Future Perspectives**

The RNA silencing pathways seem to have diversified during the evolution of eukaryotes, but the phenomenon of RNAi that regulates gene expression appears to be common among eukaryotic organisms. Therefore, once exogenous elicitor (dsRNA or siRNA) production from transgenic plants is delivered into pest cells, the endogenous cognate transcripts would be degraded. RNAi mediated plants resistance offer several advantages over conventional bio-engineering crops resistance. Firstly, in engineering broad-spectrum plant resistance even distinct lineages of plants pathogens have undergone convergent evolution and sharing homologues. Therefore multiple pathogens resistance can be achieved by silencing conserved and essential genes. Secondly, the resistance has the potential to be more durable. The RNAi-mediated resistance is based on RNA hybridization rather than protein-protein interaction; the molecular hybridization cannot be inhibited by minority nucleic acid mutation. Therefore, the possibility of the pests overcoming the resistance are likely difficult. In addition, this biotechnology represents a flexible means of developing pest resistant crops. In theory, all the pests’ genes showing detrimental knockdown phenotype can be considered as potential targets. So this strategy would not be limited by the scarcity of resistance genes. Nevertheless, RNAi has opened up a new line of thinking in designing a futuristic approach which could result in paradigm shift in insect pest management strategies.

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