RNAi-mediated knockdown of the voltage gated sodium ion channel TcNa\textsubscript{v} causes mortality in Tribolium castaneum

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The voltage-gated sodium ion channel (VGSC) belongs to the largest superfamily of ion channels. Since VGSCs play key roles in physiological processes they are major targets for effective insecticides. RNA interference (RNAi) is widely used to analyse gene function, but recently, it has shown potential to contribute to novel strategies for selectively controlling agricultural insect pests. The current study evaluates the delivery of dsRNA targeted to the sodium ion channel paralytic A (TcNa\textsubscript{v}) gene in Tribolium castaneum as a viable means of controlling this insect pest. Delivery of TcNav dsRNA caused severe developmental arrest with larval mortalities up to 73% post injection of dsRNA. Injected larvae showed significant (p < 0.05) knockdown in gene expression between 30–60%. Expression was also significantly (p < 0.05) reduced in pupae following injection causing 30% and 42% knockdown for early and late pupal stages, respectively. Oral delivery of dsRNA caused dose-dependant mortalities of between 19 and 51.34%; this was accompanied by significant (p < 0.05) knockdown in gene expression following 3 days of continuous feeding. The majority of larvae injected with, or fed, dsRNA died during the final larval stage prior to pupation. This work provides evidence of a viable RNAi-based strategy for insect control.

Insect pest control in agriculture is predominantly based on the use of synthetic chemical pesticides\textsuperscript{1–3}. Despite their effectiveness at controlling pest insects, there is a real need to develop alternative approaches with lower environmental and non-target impacts\textsuperscript{4}.

Current insecticides most commonly target components of the insect nervous system\textsuperscript{5}, often targeting the ion channels responsible for perpetuating the action potential along neurons and the enzymes of the synaptic cleft responsible for the degradation of neurotransmitters. Of these, the voltage-gated sodium channel (VGSC) is the primary target of pyrethroids\textsuperscript{6–7}. VGSCs are part of a super family of ion channels that includes the voltage-gated potassium channel, the voltage-gated calcium channel, TRP-related channels and cyclic nucleotide gated channels\textsuperscript{8}. The correct functioning of these channels is essential for normal transmission of nerve impulse and any inhibition of the action potential as a result of pesticide binding will lead to paralysis and eventual death\textsuperscript{9}. Insect VGSCs were first cloned in the late 1980s from Drosophila melanogaster\textsuperscript{10}. Although initially it was believed that D. melanogaster possessed two distinct isoforms of sodium ion channels, the DSC-type and the para-type, Zhou \textit{et al.}\textsuperscript{11} latterly showed DSC1 to be a Ca\textsuperscript{2+}-selective cation channel. Thus, it is the para-type that are involved in the nervous system, acting as voltage-gated sodium transporters\textsuperscript{12–16}. Whilst chemicals such as pyrethroids are very effective at killing insect pests they are neither targeted, nor impervious to loss of function from evolution of insect resistance\textsuperscript{17}. The lack of specificity of pyrethroids is partly a consequence of the VGSCs being highly conserved, not only across different insect orders but across animal species\textsuperscript{18}. Whilst this confers broad spectrum effects of insecticidal molecules targeting these particular ion channels, it results in limited specificity and, of more concern, non-target effects. Thus there is a need to develop novel insecticides which are of comparable...
efficacy but demonstrate higher specificity. The use of a molecular approach such as RNA interference provides a platform enabling a highly specific and targeted strategy to insect control.

RNA interference (RNAi) represents a unique form of post-transcriptional gene silencing (PTGS); it is a recognized cellular mechanism for defence against viral invasion and post-transcriptional regulation of mRNA. Further, it is the specific downregulation of gene expression mediated by an artificial double-stranded RNA (dsRNA) molecule where one strand of the dsRNA corresponds to part or all of a specific gene transcript. This sequence-based interaction confers an extremely high level of specificity. Although first described in plant and animal viruses, this evolutionarily conserved phenomenon of RNA silencing has now been observed in a range of organisms from Neurospora (an ascomycete fungus) to much more complex organisms including insects and mammals.

RNAi-based gene silencing thus has the potential to represent a novel insecticide technology, since it is theoretically possible to protect plants against insects by down regulating the expression of essential genes in the pest. Furthermore, this technology should also allow non-conserved sequences to be specifically targeted, thus conferring a high degree of specificity. The red flour beetle, Tribolium castaneum (Tc), is a major global pest. Furthermore, this technology should also allow non-conserved sequences to be specifically targeted, theoretically possible to protect plants against insects by down regulating the expression of essential genes in the pest. Furthermore, this technology should also allow non-conserved sequences to be specifically targeted, thus conferring a high degree of specificity. The red flour beetle, Tribolium castaneum (Tc), is a major global pest and it has become the genetic model for agriculturally important coleopteran species, representing an ideal system for the identification of novel pesticide targets.

The present study demonstrates that RNA interference can be used to knockdown the expression of the D. melanogaster DmNav1 homologue in T. castaneum, resulting in high insect mortality. The results provide proof of concept for this approach to be used in the sustainable control of insect pests, since this strategy is both highly specific and effective towards the targeted species, and is therefore unlikely to effect beneficial insects such as pollinators and those involved in biological control, as there is no homology between the dsRNA fragment and off target sequences.

Materials and Methods
Insects. A culture of Tribolium castaneum was obtained from Blades Biological Ltd, Kent TN8 7DX and reared at 30°C, 16:8 (L:D) on organic whole flour supplemented with 5% brewer’s yeast. Flour was replaced every 2–4 weeks.

Design of dsRNA. Selection of the target sequence used in the present study was made using the latest version of the E-RNAi web tool (http://www.dkfz.de/signaling/e-rnai3//). Output from E-RNAi selected a region of TC004126 transcript that had no similarities with other transcripts or low-complexity regions in the T. castaneum genome. The same process was employed to select a region of the kanamycin resistance gene (npt II), JN638547 (synthetic construct) from the cloning vector pSC-A-amp/kan (Stratagene) to be used as a control to assess the effect of injecting and feeding target-less dsRNA.

Total RNA isolation and cDNA synthesis. Total RNA was isolated from 4th instar larvae using TRIzol® Plus RNA Purification Kit (Ambion, TRI reagent, #12183-555) following the manufacturer’s protocol. RNA integrity was evaluated on 1.5% agarose gels as described in Sambrook and Russell, and quantified spectrophotometrically (NanoDrop, Labtech, ND-1000). cDNA synthesis was performed by reverse transcribing RNA using the i-Script™ reverse transcription supermix for RT-qPCR kit (BIO-RAD, 170-8841); 1000 ng of the extracted total RNA was used per each reaction.

Synthesis of dsRNA molecules. PCR reactions were performed in a thermal cycler (Applied Biosystems, GeneAmp PCR system 2700), in 50 μl reaction volumes using (Thermo scientific PCR master mix (2x), #K0171). The final concentration of reagents used for the majority of PCRs was as follows: dNTP 0.4 mM, Magnesium Chloride 4 mM, primer concentrations 0.2 μM (Table 1), Taq DNA polymerase, 1.25 Units/reaction. The basic cycling parameters used were as follows: Denaturation step at 95°C for 3 minutes, followed by a cycle of denaturation at 95°C for 30 s, annealing at 57°C for 30 s and elongation step at 72°C for 15 s. This cycle was repeated 35 times.

### Table 1. Primers used in PCR, synthesizing dsRNA, and performing qRT-PCR.

| Application Of Primers | Sequence (5′-3′) | Amplicon size |
|------------------------|-----------------|--------------|
| PCR                    |                 |              |
| F                      | AAGGCAAGGACATATTCCGA |              |
| R                      | TCAAACCTGTAAGATGCCAGTG | 217 bp |
| dsRNA synthesis        |                 |              |
| F<sub>Tc</sub>         | TAAATACGACTCACTATAGGGAAGGCAAGGACATATTCCGA | 217 bp |
| R<sub>Tc</sub>         | GAAAAAGATTCAAGGTAGATGCCAGTG |              |
| qRT-PCR                |                 |              |
| Tc<sub>paraA</sub>     | F               | 100 bp       |
| R                      | GATTCCCCAAAGCCCCCATC |              |
| Tc<sub>RPS6</sub>      | F               | 92 bp        |
| R                      | GTTTCTTTTCACGGCATCACC |              |
| Kanamycin resistance   | F               | 468 bp       |
| R                      | TAATAGGAAGCTCATATAGGGAAGGCAAGGACATATTCCGA |              |

Further, it is the specific downregulation of gene expression mediated by an artificial double-stranded RNA (dsRNA) molecule where one strand of the dsRNA corresponds to part or all of a specific gene transcript. This sequence-based interaction confers an extremely high level of specificity. Although first described in plant and animal viruses, this evolutionarily conserved phenomenon of RNA silencing has now been observed in a range of organisms from Neurospora (an ascomycete fungus) to much more complex organisms including insects and mammals.

RNAi-based gene silencing thus has the potential to represent a novel insecticide technology, since it is theoretically possible to protect plants against insects by down regulating the expression of essential genes in the pest. Furthermore, this technology should also allow non-conserved sequences to be specifically targeted, thus conferring a high degree of specificity. The red flour beetle, Tribolium castaneum (Tc), is a major global storage pest of grain, legumes and cereal products both for human consumption and animal feed. It has been demonstrated that T. castaneum is readily adaptable to all currently available classes of chemical insecticide. However, it is also particularly amenable to RNAi. In addition, there are many genetic and genomic tools available for this insect and it has become the genetic model for agriculturally important coleopteran species, representing an ideal system for the identification of novel pesticide targets.

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followed by a final elongation cycle of 10 minutes. Following electrophoresis, amplified bands were gel purified using QIAquick MinElute Gel Extraction kit (Qiagen, #28604), cloned into Stratagene vector pSC-A-amp/kan (Stratagene, #240205) and then transformed into Stratagene SoloPack competent cells. Recombinant plasmids were sequenced to verify the cloned insert. The dsRNA was synthesized using replicator RNAi kit (Thermo Scientific, #F-610) and stored at −80 °C prior to injection.

**Delivery of the double-stranded RNA to larvae and pupae.** Larvae and pupae (reared on organic whole flour wheat as above) were injected as described by Tomoyasu and Denell37, using a dissecting stereomicroscope. The dsRNA was injected into the dorsal side between the 1st and 2nd abdominal segments of 6th (final) instar larvae, and laterally between the 2nd and 3rd abdominal segments for pupal stages, using a NanojectII™ injector (Drummond Scientific Company). Three biological replicates (5 larvae or pupae/replicate) were injected from one dsRNA preparation (780 ng/μl in RNAs free water, equivalent to (53.6, 106.7, 160 ng/larva and 160 ng/pupa); 48 h post injection gene knockdown was measured by RT-qPCR using gene-specific primers. For the longer-term injection bioassays, 6th instar larvae (3 biological replicates, 15 larvae/rep) for each dose (53.6, 106.7, 160 ng/larva) were injected, returned to the diet and monitored on a daily basis until adult eclosion. As described previously, all injected controls received RNAase free water at the same volumes as experimental conditions; a second control of dsRNA towards the microbial kanamycin resistance gene (nptII) was also included in addition to the non-injected insect group.

The dsRNA was also delivered orally via flour disks at a range of concentrations (0, 50, 100, 150 ng dsRNA/μl diet) prepared as described by Xie et al.30, in 96-well-flat-bottom well microtiter plates; one larva (6th instar) was added to each well. Three biological replicates, each of 15 larvae/rep were carried out for each dose; for all feeding assays controls were used as described above. Survival of larvae was monitored until adult eclosion. To monitor gene knockdown, three biological replicate (5 larvae/rep/dose) were allowed to feed for 72 h on flour disks as described above at the same doses, after which time larvae were snap frozen in liquid nitrogen prior to assessing gene knockdown by RT-qPCR.

Since preliminary studies with different doses showed no difference in either control mortality or expression of target gene30, subsequent controls were only carried out at a single (highest) dose for each treatment.

**Gene expression studies with quantitative reverse transcription PCR (RT-qPCR).** cDNA, prepared as shown above, was used as the template; RT-qPCR was performed on the template in a final volume of 25 μl. Each reaction contained: 12.5 μl 2x Rotor-gene SYBR green PCR master mix solution (Qiagen Co.), forward and reverse primers (Table 1) were added in a final concentration of 0.5 mM, 9 μl nuclease free water and 1 μl of undiluted cDNA. To validate the primers, a standard curve based on a serial dilution of cDNA was carried out to determine the primer annealing efficiency, the presence of primer dimers and the production of a single PCR product. RT-qPCR conditions were as follows: 95 °C for 5 min, followed by 40 cycles of 95 °C for 15 sec, 55 °C for 30 sec and 60 °C for 15 sec. Amplifications were carried out using three biological replicates of cDNA, each one from RNA obtained from five individuals, and the mean values of three technical replicates were analyzed. The efficiency of the primers were equivalent and the relative transcript quantity was calculated according to the delta-delta Ct method41, with Ct values of the respective target gene compared to those of the reference gene TcRPS6 (GenBank XP_968395.1); this gene is a reliable reference gene for RT-qPCR in T. castaneum42, and hence used to normalize gene expression.

**Statistical analysis.** Experimental data and qPCR results were analysed by ANOVA followed by Tukey Kramer Multiple Comparison; statistical differences are shown as different letters. Insect mortalities were analysed by Kaplan–Meyer survival analysis, SIGMAPLOT program, version 12.5, and larval mortality was corrected according to Abott’s formula43, with percent adult emergence calculated as previously described44,45.

**Results**

**Bioinformatic analysis of targeted sequence.** Homology of the TcNa v dsRNA fragment designed by the E-RNAi web tool to the other 744 insect Na v sequences, including representatives from important pollinator species, held by NCBI was investigated using the MegaBLAST algorithm. No homology was reported compared as shown above, was used as the template; RT-qPCR was performed on the template in a final volume of 25 μl. Each reaction contained: 12.5 μl 2x Rotor-gene SYBR green PCR master mix solution (Qiagen Co.), forward and reverse primers (Table 1) were added in a final concentration of 0.5 mM, 9 μl nuclease free water and 1 μl of undiluted cDNA. To validate the primers, a standard curve based on a serial dilution of cDNA was carried out to determine the primer annealing efficiency, the presence of primer dimers and the production of a single PCR product. RT-qPCR conditions were as follows: 95 °C for 5 min, followed by 40 cycles of 95 °C for 15 sec, 55 °C for 30 sec and 60 °C for 15 sec. Amplifications were carried out using three biological replicates of cDNA, each one from RNA obtained from five individuals, and the mean values of three technical replicates were analyzed. The efficiency of the primers were equivalent and the relative transcript quantity was calculated according to the delta-delta Ct method41, with Ct values of the respective target gene compared to those of the reference gene TcRPS6 (GenBank XP_968395.1); this gene is a reliable reference gene for RT-qPCR in T. castaneum42, and hence used to normalize gene expression.

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**Temporal changes in TcNa v gene expression.** Relative abundance of TcNa v mRNA at specific life stages was estimated using qRT-PCR to ensure that subsequent studies were carried out on appropriate stages. Transcripts were detected in all life stages investigated. Relative expression of TcNa v, was more abundant in the late pupal stage compared to either the adult or any of the larval stages, although these differences were not significant (p > 0.05; Fig. 1). Expression in the late pupal stages was 1.6 fold greater relative to the adult stage, but lower in the larval stages, with transcript levels in 6th, 4th and 2nd instar larvae being decreased (0.9, 0.6 and 0.4-fold, respectively). Based on these relative expression levels, both pupal stages (early and late) and 6th (final) instar larvae were selected as suitable stages for subsequent RNAi studies.

**Physiological and molecular response of T. castaneum to TcNa v dsRNA.** Effects on changes in expression levels of TcNa v transcripts and changes in phenotype (survival and adult emergence) were investigated using both oral delivery of dsRNA and via injection.
Delivery of dsRNA via injection. The dose response effects of dsRNA directed to TcNav, on survival and subsequent development into adulthood of final (6th) instar larvae were investigated. Larvae were monitored daily post injection until pupation, and surviving insects recorded at eclosion. Injection of dsRNA caused a significant dose dependant decrease in survival ($p < 0.05$), (Fig. 2a). Six days post injection, larval mortality was 35.2% ± 1.7, 62.23% ± 2.4 and 73% ± 5.9 relative to the –ve control (injected with RNase free water), when injected with dsRNA at concentrations of 53.6, 106.7 and 160 ng/larva, respectively (Fig. 2a). Over this same time period, no mortality occurred in the non-injected control larvae; the 2 control groups (RNase free water and kanamycin resistance dsRNA, –ve control = larvae injected with RNAase free water).

Knockdown of TcNav expression following injection of dsRNA was demonstrated using RT-qPCR (Fig. 3). Analysis showed a direct correlation between the amounts of dsRNA injected with a subsequent decrease in the abundance of TcNav mRNA transcript. Gene knockdown at 48 h post injection of 6th instar larvae was shown to be significant ($p < 0.05$) between the different treatments, with expression of TcNav, being reduced 1.43, 2.0 and 2.5-fold in response to increasing amounts of dsRNA corresponding to 53.6, 106.7 and 160 ng/larva, respectively.
relative to larvae injected with RNase free water. Furthermore, qPCR analysis revealed that mRNA transcript levels in early stage and late stage pupae injected with 160 ng/pupae dsRNA were also significantly (p < 0.05) down regulated at 48 h post injection, with mRNA levels for TcNav being 1.43 and 1.73 fold lower, respectively (Fig. 4a,b) compared to their appropriate controls, which were injected with RNase free water. The differences between the two pupal stages were not significant (α = 0.05). The injection of naked off target dsRNA into either larvae or pupae did not affect expression of Na\(_V\) in \(T.\ castaneum\). This dsRNA molecule designed towards a 468 bp fragment of the microbial kanamycin resistance gene (nptII) was synthesised and injected at 160 ng/larva. RT-qPCR revealed that expression of TcNav in these individuals was 0.98-fold (p > 0.05) that of the control group that were injected solely with RNase free water.

**Oral delivery of dsRNA.** Induction of RNA interference via oral delivery was carried out by restricted feeding of individual 6\(^{th}\) instar larvae on flour disks containing the dsRNA at a range of different concentrations (0, 50, 100 and 150 ng/mg of diet). Relative to the control diet, which fed on flour disks containing kanamycin dsRNA, feeding of the dsRNA-containing diet for 6 days resulted in a significant (p < 0.05) impact on larval survival, with larval mortality of 19\% ± 6.6, 37.83\% ± 0.72 and 51.34\% ± 6.8 respectively, for the 3 doses, 50, 100 and 150 ng/mg (Fig. 5a). Probit analysis showed this to be a linear dose response, with an L\(C_{50}\) of 150.23 ng dsRNA/mg of diet. As with the injection of dsRNA, oral delivery also resulted in a dose dependent response. At termination of the bioassay, reduction in adult emergence from insects reaching pupation correlated with the administered dose, with the higher doses causing greater reduction in development to adulthood, with reductions of

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**Figure 3.** Transcript levels of TcNav mRNA after normalization with the TcRPS6 as an endogenous control 48 h post larval injection. Mean ± SEM of three replications are shown. Means with the different letters are significantly different (p = 0.05; ANOVA; with Tukey Kramer Multiple Comparison). Knockdown of transcript level is indicated for each dose (KD%) with respect to the control group, +ve control = larvae injected with Kanamycin resistance dsRNA, −ve control = larvae injected with RNAase free water.

**Figure 4.** Transcript levels of TcNav mRNA after normalization with the TcRPS6 as an endogenous control 48 h post pupal injection. (a) Early stage pupae, (b) Late stage pupae. Mean ± SEM of three replications are shown. Means with the different letters are significantly different (p = 0.05; ANOVA; with Tukey Kramer Multiple Comparison). Knockdown of transcript level is indicated for each dose (KD%) with respect to the control group, +ve control = larvae injected with Kanamycin resistance dsRNA, −ve control = larvae injected with RNAase free water.
9.9% ± 13.2, 24.35% ± 6.2 and 51.26% ± 3.8 at 50, 100 and 150 ng/mg, respectively (Fig. 5b). Although the effects of the lowest dose were not significant (p > 0.05), the two higher doses did have significant (p < 0.05) effects on adult emergence.

Not only did oral delivery cause a reduction in larval survival, resulting in reduced adult emergence, there was also evidence of gene knockdown. Oral delivery of dsRNA caused significant reductions in transcript levels of TcNav. RT-qPCR analysis revealed that after 72 hours feeding by final (6th) instar larvae, transcript levels were significantly (p < 0.05) down regulated by 1.67 and 1.75-fold for the different treatments (100 and 150 ng dsRNA/mg diet) respectively, compared to the control group, which was not challenged with the TcNav dsRNA, but fed on kanamycin dsRNA flour disks (Fig. 6). However, oral delivery of dsRNA at 50 ng dsRNA/mg diet did not result in a significant decrease in gene TcNav gene expression (1.47-fold reduction, p > 0.05). Insects feeding on flour disks containing 150 ng/mg dsRNA towards the kanamycin gene did not show any significant reduction in expression of TcNav at any of the concentrations tested (0.83-fold of control, p > 0.05) compared to the group consuming control diet.

Discussion and Conclusions
The function of the voltage-gated sodium ion channels (VGSCs) is to initiate and then propagate action potentials in the nervous system and other excitable cells. The transmembrane region S4 of each domain contains 3–5 positively charged residues that are essential for the channel to respond to changes in membrane potential. In response to membrane depolarization, electrostatic interactions between positively charged residues and the depolarized cytoplasm force the S4 region towards the extracellular surface causing a conformational change in S5–S6 that opens the channel pore and allows sodium influx. These ion channels are the primary target for pyrethroid insecticides which act by preventing the closure of the VGSC. The pyrethroid group of insecticides
has proven to be extremely successful in the broad spectrum control of common agricultural insect pest species. However target-site resistance to these insecticides has been demonstrated in several insect orders and is based on single point mutations within the DmNa, orthologues.

RNA interference for suppression of specific transcripts is proving to be a powerful tool in several insect species, and, in contrast to many current chemical formulations, may provide a highly specific control strategy. Hence, such an approach has the potential to significantly reduce non-target effects, including those on beneficial insects such as parasitoid wasps and pollinators. T. castaneum has been shown to exhibit a robust systemic RNAi response in the postembryonic stages, thus it has become a model insect in which RNAi proof of concept can be demonstrated. As far as the authors are aware, this is the first study to show RNAi-induced knockdown of VGSC gene expression in insects, resulting in a dose-dependent, fatal phenotype. In the present study RNAi was induced following administration of dsRNA corresponding to the DmNa, orthologue from T. castaneum (TcNa, gene bank accession number NM_001165908.1 and TC004126 at Beetle-Base), with gene knockdown being observed following both injection into the insect haemocele and via oral uptake. Prior to delivery of the dsRNA it was first necessary to determine transcript abundances of this gene at different developmental stages to select for appropriate stages for study. Although not statistically significant, the results showed that transcript levels peaked during the late pupal stage followed by the adult and 6th instar larval stages. It is perhaps not surprising to find that TcNa transcripts were more abundant in late pupal stages, as it is during this time that axons and dendrites are remodelled to generate an adult-specific arbor during metamorphosis prior to adult eclosion. These results are consistent with those reported for two acetylcholinesterase genes (TcAce1 and TcAce2), where the transcript levels were more abundant in 6-day old pupae followed by 2-day old adult T. castaneum. Furthermore, whilst transcript levels of three genes encoding diuretic hormones were present in all developmental stages of this insect, relative expression levels were similarly dependent upon the actual development stage. In contrast, analysis of the developmental stage-specific expression pattern of a lethal giant larval gene (TcLgl1) in T. castaneum showed constitutive expression.

The injection of TcNa dsRNA into the haemocele of larval T. castaneum induced a quantifiable RNAi effect in a dose dependent manner resulting in reduction of gene expression after 2 days. Similar observations in T. castaneum have been reported by Lu et al., where transcript levels for two acetylcholinesterase genes (TcAce1 and TcAce2) were reduced by 92.3 and 95.2%, respectively, compared to the control 4 days post injection of corresponding dsRNA fragments. Similarly, studies with the crustacean cardioactive peptide (CCAP) showed that loss of CCAP activity can be induced through RNAi with dsRNA towards TcCcapr-2. Such gene knockdown resulted in no visible abnormal phenotype but demonstrated that TcCcapr-2 is essential for cardioacceleratory activity. Furthermore, Xiao et al. demonstrated that T. castaneum larvae injected with the double stranded RNA of TcLgl (lethal giant larva) at each of the three doses (100, 200, and 400 ng/larva) resulted in significant and dose dependent suppression of TcLgl transcripts 2 days post injection.

As with larvae, suppression of TcNa expression levels were also observed both in the early and late pupal stages, again showing consistency with those reported by where injection of 1-day old pupae with dsTcLgl caused significant suppression of the corresponding transcript on days 2, 4, 6 and 8. Furthermore, studies demonstrated that freshly moulted pupae of T. castaneum injected with dsRNA led to a specific knockdown of the respective transcripts with the exception of one gene only. This dsRNA-mediated silencing caused a loss of function phenotype.

The present study demonstrates, irrespective of the method of delivery, significant and severe developmental arrest (in terms of larval mortalities and reduced pupation) following knock-down of the VGSC, the severity being dependent on the dose administered. Injection of dsRNA into Locusta migratoria showed that locusts also have a sensitive and systemic dose-dependent RNAi response. Similar responses to oral delivery of dsRNA have also been reported in T. castaneum.

Data presented by Clemens and Elia, Oates et al. and Geiss et al. confirmed that the interferon-regulated innate immunity pathway that protects vertebrates from invasion of long dsRNA is absent in insects. It is thus feasible and preferable to deliver long dsRNAs to insects to ensure a maximal RNAi effect. Furthermore, Whyard et al. observed that it is possible to deliver relatively short (29–40 nt) dsRNAs designed towards insect vATPase transcripts whilst maintaining levels of induced mortality. This observation will prove advantageous if RNAi is to be developed as a viable tool in crop protection. Several T. castaneum genes such as heat shock protein 90, chitin synthase, the segmentation gene hairy, and a matrix metalloprotease have been shown to be down regulated by RNAi, and therefore may represent good candidate genes to target in future RNAi-based crop protection strategies. The high degree of specificity of the dsRNA molecules used in the present study towards their individual targets, as shown by the MegaBLAST homology searches, lend support for the use of this technology in crop protection. Use of this novel technology is further supported by the lack of any appreciable dsRNA uptake from the diets of higher animals. Therefore, dietary dsRNA is not anticipated to represent any hazard or risk to humans, mammals, or other vertebrates. For these reasons transgenic crops expressing dsRNA are currently under investigation as a new tool for targeted and specific protection of crops from insect pests.

The findings of the present study provide the first demonstration of the silencing of a voltage-gated sodium ion channel gene (TcNa,) via RNAi in T. Castaneum. It provides evidence to suggest that this gene is a feasible candidate to target for the control of this insect pest using RNAi. Furthermore, the findings presented here support the temporal expression of the genes during development, which will help in targeting the key developmental stage if this technology is to be used in crop protection.

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Author Contributions

H.M.A.E.H., M.G.E., E.Y.T.N. and A.M.R.G. designed the overall experiment. H.M.A.E.H., M.G.E. and M.S.A. designed the dsRNA fragment and performed the molecular cloning from T. castaneum. H.M.A.E.H., M.G.E., E.Y.T.N. and M.S.A. performed the insect injections and the RT-qPCR. B.M.H.A. performed the additional bioassays and expression analysis requested after initial review. H.M.A.E.H., M.G.E. and A.M.R.G. the interpreted of results. H.M.A.E.H., M.H.A. and E.M.S. wrote the manuscript. A.M.R.G. and M.G.E. corrected the manuscript for final submission. All of the authors reviewed and contributed to the manuscript.

Additional Information

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