Mis-splicing of the ABCC2 gene linked with Bt toxin resistance in *Helicoverpa armigera*

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Toxins from the bacterium *Bacillus thuringiensis* (Bt) are used widely for insect control in sprays and transgenic plants, but their efficacy is reduced when pests evolve resistance. Previous work showed that mutations in a gene encoding the transporter protein ABCC2 are linked with resistance to Bt toxins Cry1Ab, Cry1Ac or both in four species of Lepidoptera. Here we compared the ABCC2 gene of *Helicoverpa armigera* (HaABCC2) between susceptible strains and a laboratory-selected strain with 1,000-fold resistance to Cry1Ac relative its susceptible parent strain. We discovered a 73-base pair (bp) insertion in the cDNA of the resistant strain that generates a premature stop codon expected to yield a truncated ABCC2 protein. Sequencing of genomic DNA revealed that this insertion is an intron that is not spliced out because of a 6-bp deletion at its splicing site. Analysis of progeny from crosses revealed tight genetic linkage between HaABCC2 and resistance to Cry1Ac. These results provide the first evidence that mis-splicing of a gene encoding an ABCC2 protein confers resistance to a Bt toxin.
**Table 1 | Responses to Cry1Ac of two susceptible strains (LF and 96S) and a resistant strain (LF60) of *H. armigera***

| Strain | Form of Cry1Ac | LC50 (95% fiducial limits) (µg Cry1Ac per ml diet) | Resistance ratio* |
|--------|----------------|-----------------------------------------------|------------------|
| LF     | protoxin       | 0.009 (0.002–0.020)                            | 1.0              |
| 96S    | protoxin       | 0.021 (0.008–0.036)                            | 2.3              |
| LF60   | protoxin       | 12.4 (8.0–17)                                  | 1400             |
| LF     | activated toxin| 0.008 (0.002–0.014)                            | 1.0              |
| 96S    | activated toxin| 0.013 (0.007–0.021)                            | 1.6              |
| LF60   | activated toxin| 9.15 (5.8–13)                                  | 1100             |

*LC50 of each strain divided by the LC50 of the susceptible LF strain.

**ABCC2 cDNA in susceptible and resistant strains.** We give the name *HaABCC2* (GeneBank accession no. KF479231) to the gene encoding the ABCC2 protein of *H. armigera*. The cDNA of *HaABCC2* from the susceptible LF strain had 4,017 bp encoding a predicted ABCC2 protein of 1,338 amino acids (Fig. 1). Compared with predicted *HaABCC2* proteins from other Lepidoptera, the predicted *HaABCC2* protein shares 93% amino acid identity with *Heliothis virescens* (GenBank accession nos. AD16740.1) and *Heliothis subflexa* (AD16744.1), 72% from *Bombyx mori* (BAK82126.1) and 66% from *Plutella xylostella* (AE127592.1). Similar to the structures of other lepidopteran ABCC2 proteins (Fig. 1), the proposed structure of *HaABCC2* protein includes twelve transmembrane segments and two ATP-binding domains (Fig. 1).

The *HaABCC2* cDNA from the resistant strain LF60 has a 73-bp insertion between bp 3582 and 3583 that introduces a premature stop codon (Figs. 1 and 2A). As a result, the amino acid sequence encoded by the resistant allele not only lacks the C-terminal 143 amino acids that occur in the susceptible strain (from 1196 amino acid to the end 1338 amino acid), but also has 15 amino acids that are not encoded by the susceptible allele (after amino acid 1196) (Fig. 1). We also found 124 single nucleotide polymorphisms (SNPs) indicating differences between the cDNA from the resistant and susceptible strains (Fig. S1) of which seven cause amino acid changes (Fig. 1). The relative quantity of transcripts of *HaABCC2* did not differ significantly between LF and LF60 (Fig. S2).

We developed an allele-specific RT-PCR gel analysis using specific primers (GF and GR) flanking the 73-bp insertion to distinguish between the transcripts in the susceptible and resistant strains (Fig. 2). As expected, the band amplified was about 70 bp longer in the resistant strain than in the susceptible strain (Fig. 2B).

**ABCC2 gDNA in susceptible and resistant strains.** Alignment of the gDNA and cDNA sequences of *HaABCC2* from the two strains revealed that the 73-bp insertion in the cDNA sequence of the resistant allele is an intron, which was spliced out in the transcript from the susceptible strain but not from the resistant strain (Fig. 2C). Comparison of the gDNA sequences indicates that LF60 lacks the first 6 bp ("GTACGC") of this intron, including the 5’ “GT” dinucleotide intron signature (Fig. 2C), which leads to failure to splic e this intron. In 100 sequenced gDNA samples from each strain, all samples in LF60 were homozygous for the presence of this 6-bp deletion and all samples from LF lacked this deletion. All F1 progeny from a cross between LF60 and LF were heterozygous for the deletion.

**Linkage between *HaABCC2* mis-splicing and Cry1Ac resistance.** Sequencing of the gDNA fragment flanking the 6-bp deletion of larvae from 10 backcross families tested in bioassays shows tight genetic linkage between *HaABCC2* and resistance to Cry1Ac (Table 2). For the 1200 larvae genotyped by sequencing gDNA (120 per backcross family), all 600 survivors on treated diet were homozygous for the 6-bp deletion, but of the 600 larvae on untreated diet, 48% were homozygous for the 6-bp deletion and 52% were heterozygous (Table 2, Fisher’s exact test, P < 0.0001 for each family).

Results from larvae genotyped by RT-PCR with cDNA (Fig. 3) confirm the results of the linkage analysis based on sequencing of gDNA (Table 2). The RT-PCR results show that all of the F1 larvae tested were heterozygous for the 73-bp insertion (Fig. 3A). In backcross families A1 and B6 analyzed by RT-PCR, 20 larvae that survived on treated diet were homozygous for the 73-bp insertion (Fig. 3D and 3E), whereas 20 larvae from untreated treated diet were either heterozygous or homozygous for the insertion in equal proportions (Fig. 3B and 3C).

**Discussion**

The results here showing tight genetic linkage between a mutation in the *ABCC2* gene of *Helicoverpa armigera* (*HaABCC2*) and resistance to Cry1Ac confirm the importance of the *ABCC2* protein in toxicity of Cry1Ac against lepidopteran larvae. Previous work revealed linkage between the *ABCC2* gene and resistance to Cry1Ab, Cry1Ac, or both in strains of four lepidopteran species19–21. In *Bombyx mori*, a mutation in the gene encoding ABCC2 was associated with reduced binding of Cry1Ab in transformed S9 cells25, but not in brush border membrane vesicles isolated from midguts of resistant and susceptible larvae26. It has been hypothesized that for some Bt toxins, binding to ABC22 proteins is essential for toxicity; thus disruption of this binding causes resistance25,26. However, as far as we know, direct evidence of binding of Bt toxins to ABC22 proteins has not been reported.

In *H. armigera*, the association between toxin binding and the ABC22 mutation remains to be determined. In resistant strain LF60 of *H. armigera*, the ABC22 gene is missing six base pairs in one of its intron-exon boundaries, including the conserved 5’ splicing dinucleotide “GT” (Fig. 4A). Because of this mutation, an intron is not spliced out of the ABC22 transcript, which introduces a stop codon. This premature codon is predicted to cause the loss of 143 amino acids, including a C motif and Walker B sequence, which could affect the protein’s conformation (Fig. 4B), potentially interfering with binding of ATP as well as with Bt toxins.

If the mutation in *HaABCC2* does disrupt ATP binding, this might reduce the fitness of resistant insects relative to susceptible insects. However, the lack of a significant defect of resistant homoyzoytes in the backcross larvae fed untreated diet (Table 2) indicates that a major fitness cost affecting survival was not detected in our experiments. Major fitness costs affecting survival also were not detected in similar backcross experiments on diet with *H. virescens*, *P. xylostella* and *T. ni*, but this does not exclude the possibility of fitness costs affecting survival or other traits when larvae develop on plants in the field19,25.

Mis-splicing of cadherin genes is associated with resistance to Cry1Ac in *Pectinophora gossypiella* and *H. armigera*27–29. As far as we know, the results reported here are the first showing that mis-splicing of an ABC22 gene is associated with resistance to a Bt toxin. Based on the results here and previous reports, the mechanisms of resistance to Cry1Ac in laboratory- and field-selected strains of *H. armigera* are diverse, including the recessive mutation disrupting *ABCC2* found here, recessive and dominant mutations disrupting
Figure 1 | Predicted amino acid sequences of the ABCC2 proteins from a susceptible strain (LF) and a resistant strain (LF60) of *H. armigera* compared with susceptible strains from *B. mori* (BmAABCC2, GenBank BAK82126.1) and *H. virescens* (HvABCC2, GenBank ADH16740.1). Walker A and B sequences, C motifs, and transmembrane (TM) domains were predicted by TMHMM ver.2.0 (http://www.cbs.dtu.dk/services/TMHMM/). Arrows show seven amino acid differences between LF and LF60.
the extracellular and intracellular domains of cadherin, reduced transcription of a protease that converts protoxin to toxin, reduced transcription of membrane-bound alkaline phosphatase, reduced activity of aminopeptidase N, elevated immune response, and non-recessive mutations in unidentified genes. This diversity will continue to provide challenges for understanding, monitoring, and managing resistance of \textit{H. armigera} to Bt cotton.

**Methods**

**Insects.** We used three laboratory strains of \textit{H. armigera}: susceptible strains 96S and LF, and resistant strain LF60. The 96S strain was started with 20 pairs of adults collected from conventional cotton in Xinxiang, Henan Province, China in 1996. The LF strain was started with about 100 third to sixth instars collected from Bt cotton in Langfang, Hebei Province, China in 1998. As expected, lanes 2 and 3 from the susceptible strains show a band of about 130 bp, while lane 4 from the resistant strain shows a band of about 200 bp, which reflects the 73-bp insertion. (C) Alignment of gDNA and cDNA sequences. The blue line indicates the intron sequence and the red line indicates the deletion in the splicing site in LF60.

**Bioassays.** We used diet incorporation bioassays to evaluate susceptibility to Cry1Ac protoxin and Cry1Ac activated toxin. Cry1Ac protoxin was extracted and purified from the HD73 strain of \textit{B. thuringiensis} subsp. \textit{kurstaki} by the Biotechnology Group in Institute of Plant Protection, Chinese Academy of Agricultural Sciences, Beijing, China. To obtain activated Cry1Ac toxin, Cry1Ac protoxin was incubated for more than a decade with progressively increasing concentrations: 1, 5, 10, 30 and 60 μg Cry1Ac protoxin per g diet. Insects were reared at 27 ± 2°C and 75 ± 10% relative humidity with a photoperiod of 14L:10D.
with a 25:1 ratio of trypsin (Sigma) to protoxin, and the soluble trypsinized toxin was purified by a Superdex 200 HR 10/30 column (Amersham Biosciences) on a fast protein liquid chromatography (FPLC) system.

Various concentrations of Cry1Ac protoxin and activated toxin were added and thoroughly mixed with diet to obtain the desired concentrations. After mixing, the diet solidified and we put pieces of solid diet (1 g) into each well of a 24-well plate. We

Figure 3 | PCR analysis of HaABCC2 cDNA amplified by allele-specific primers GP-F and GP-R. (A) F1 (rs) progeny from a cross between the resistant LF60 strain and the susceptible LF strain. (B and C) Progeny from backcross larvae reared on untreated diet; genotypes are either rs (two bands) or rr (one band). (D and E) Progeny from backcross larvae reared on diet treated with Cry1Ac; all are rr (one band). Backcrosses were done with single-pair crosses either with a resistant male and an F1 female (B and D) or an F1 male and a resistant female (C and E).

Figure 4 | Summary of differences between ABCC2 in a susceptible strain (LF) and a resistant strain (LF60) of H. armigera. (A) gDNA. E: exon, I: intron, red arrow shows 6-bp deletion in LF60. (B) Schematic structure of HaABCC2 protein showing location of predicted loss of amino acids in the ATP-binding domain 2 of LF60.
put one first instar in each well of a 24-well plate for each replicate, with three replicates per treatment (total n = 72 per treatment). Larvae were considered dead if they died or did not reach third instar after 7 days.

Cloning and sequence analysis of HaABCC2. We analyzed 20 larvae (5th instars, 12 days old), 10 from LF and 10 from LF60. Midguts were dissected under a dissecting microscope, washed with cold 0.7% physiological saline, briefly dried on filter paper, pooled with 1 ml of Tris-EDTA buffer (1 mg/l), frozen in liquid nitrogen, and stored at −80°C for subsequent RNA extraction.

Total RNA was extracted from the sample homogenates according to the standard TRIzol reagent protocols (Invitrogen). RNA purity was evaluated by 260/280 and 260/230 ratios measured in a NanoDrop 3300 (Thermo). Genomic DNA was eliminated from the samples by treatment with DNaseI (Fermentas). Total RNA (4 μg) was reverse transcribed into cDNA in a 20-μl reaction using SuperScript® III First-Strand Synthesis kit (Invitrogen) according to the manufacturer’s instructions. The resultant cDNA was stored at −20°C for subsequent homology-based cloning of ABCC2 cDNA sequences from the two H. armigera strains, and then quantitative polymerase chain reaction (qPCR) analysis of ABCC2 expression in the two strains.

Cloning of H. armigera ABCC2 (named HaABCC2) cDNA sequence was initiated with PCR amplification of a partial fragment using the cDNA sample from LF as the template and the primers VP-F and VP-R (Table S1) designed based on the nucleotide sequence got from data RNA sequence experiment. The PCR conditions were 4 min denaturation at 95°C, followed by 35 cycles of 30 sec denaturation at 94°C, 30 sec annealing at 60°C and 30 sec extension at 72°C, and a final 10 min extension at 72°C. The amplified partial fragment was eluted from the agarose gel, cloned into the pMD20-T Vector (TaKaRa), and sequenced by Sangon Biotech Company.

Based on the sequence of the partial fragment, four gene-specific primers (5′-RACE-R1, 5′-RACE-R2, 3′-RACE-F1 and 3′-RACE-F2) of Table S1 were designed to nest PCR amplify the 5′ (5′-RACR-R1 and 5′-RACE-R2) and 3′ ends (3′-RACE-F1 and 3′-RACE-F2) of HaABCC2 DNA with four general primers (5′ RACE outer primer, 5′ RACE inner primer, 3′ RACE outer primer and 3′ RACE inner primer TaKaRa) to amplify the cDNA as the template and the 5′- and 3′-Full RACE Kit (TaKaRa) following manufacturer’s instructions. The resultant 5′ and 3′ ends were cloned, sequenced, and aligned with the partial fragment to yield the full-length cDNA of HaABCC2 from LF. We then designed a pair of primers HaABCC2 F and HaABCC2 R based on the above full-length cDNA of HaABCC2 to PCR-amplify the ORF (open reading frame) of HaABCC2 from LF and LF60 respectively. The ORF PCR conditions were 4 min denaturation at 95°C, followed by 35 cycles of 30 sec denaturation at 94°C, 30 sec annealing at 55°C and 3 min extension at 72°C, and a final 10 min extension at 72°C.

The programs ClustalX and DNA MAN were used to align the sequences. TMHMM ver 2.0 (http://www.cbs.dtu.dk/services/TMHMM/) was used to predict the HaABCC2 domains.

qPCR analysis of HaABCC2. Three cDNA samples per strain were prepared as above from 3 pools of 10 larval midguts each were use as the templates to compare the expression level of HaABCC2 between LF and LF60. qPCR. Two H. armigera housekeeping genes, beta actin (β-actin) and glyceraldehyde-3-phosphate dehydrogenase (GAPDH), were used as the dual reference genes to normalize the expression of HaABCC2. qPCR (TaqMan) of the three genes were individually performed in a 20-μl reaction containing 2X Maxima probe/ROX qPCR Master Mix (10 μl), cDNA template corresponding to 50 ng, for-forward and reverse primer (0.6 μl each), 0.1X probe (0.4 μl), 2 μl LF or LF60 template cDNA, and 6.4 μl nuclease-free water. All qPCR reactions were performed in 96-well optical plates using the Applied Biosystems 7500 FAST qPCR system (ABI7500 Fast) under the following conditions: 20°C, 2 min; 95°C, 10 min; 40 cycles: 95°C, 15 second, 60°C, 1 min. qPCR of each of the three genes were repeated three times per cDNA sample and no-template nuclease-free water control. Data were processed using the 2-ΔΔCt method. All the primers and FAM probes used for the three genes are shown in Table S1.

Allele-specific RT-PCR. Based on a 73-bp insertion in the ORF of HaABCC2. The resultant PCR products from the gDNA samples were sequenced directly. The gDNA sequences obtained were aligned to locate the gDNA mutations responsible for the 73-bp insertion in cDNA. Genetic linkage between HaABCC2 and resistance to Cry1Ac. To test for genetic linkage between HaABCC2 and Cry1Ac resistance, we generated F1 progeny from a single-pair cross between a male from the susceptible LF strain and a female from the resistant strain. We generated 10 backcross families of two different types: backcross families A1-A5 were each produced by a single-pair cross of a female F1 with a male from the resistant strain, and families B6-B10 were each produced by a single-pair cross of a male F1 with a female from the susceptible strain. From each of the 10 backcross families, 120 larvae were tested using the bioassay method described above; 60 from each family on diet that had no toxin (control) and 60 from each family on diet treated with 2 μg Cry1Ac per ml activated toxin to kill susceptible larvae. We used the allele-specific PCR described above to determine the genotype of larvae from backcross families. We also sequenced the HaABCC2 gDNA fragment flanking the 73-bp insertion. A total of 40 larvae from families A1 and B6 were analyzed with PCR; 10 from each family fed diet without toxin (control) and 10 from each family fed diet treated with 2 μg Cry1Ac activated toxin per ml diet to kill susceptible larvae. Ten F1 larvae were also analyzed with PCR.

Statistical analysis. Data processing system (DPS) software package on analysis of statistical data. Zhejiang University. China was used to get LCLCs. For each of ten backcross families, we used a separate Fisher’s exact test (http://graphpad.com/quickcalcs/contingency1.cfm) to determine if the proportion of genotypes (rr and Rr) differed significantly between larvae reared on treated versus untreated diet and to determine if the proportion of the genotypes (rr and Rr) for larvae fed untreated diet differed significantly from the expected proportion of 0.5 for each genotype.

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**Author contributions**

K.W. and Y.X. designed the study. Y.X., T.Z. and C.L. performed the experiments. K.W., Y.X. and B.E.T. analyzed the data. K.W., Y.X. and B.E.T. wrote the manuscript. All authors have read and approved the manuscript for publication.

**Additional information**

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