Identification and functional analysis of diet-responsive genes in *Spodoptera litura* (Fabricius)

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Abstract

**Background:** *Spodoptera litura* is one of the most devastating agricultural pests with a wide range of host plants. To study larval performance on different diets and midgut adaptation at transcriptional levels, feeding assay and RNA-Seq experiments were conducted. RNA interference technology was used to explore the detoxification and metabolism of two cytochrome *P450* genes.

**Results:** The bioassay data showed that *Spodoptera litura* larvae developed more quickly when fed on cabbage than when fed on soybean, corn and cotton, tannin can inhibit the growth of *Spodoptera litura*. The result of RNA-Seq indicated that *Spodoptera litura* midgut modified gene expression levels to accommodate different diets, and the most differentially expressed genes were detoxification-related and digestion-related genes. Further analysis showed that the glutathione metabolism pathway was the common detoxification pathway in *Spodoptera litura*. The expression of cytochrome *P450* genes showed a clear response to different plant hosts, and these differences may play key functions in primary detoxification of secondary metabolites from host plants. Meanwhile, the digestive enzymes of proteinases, lipases, and carbohydrases in midgut showed special responses to different plant hosts. After injection of dsRNA of *CYP321A19* and *CYP6AB60*, the expression level of target gene were decreased, and the sensitivity of insect to plant allelochemicals increased and the weight increase significantly slowed.

**Conclusion:** In this study, genes involved in detoxification were identified, and the results demonstrate the genes and pathways *Spodoptera litura* utilize to detoxify specific plant-host allelochemicals. These results may also provide a theoretical basis for *Spodoptera litura* management.

**Background**

*Spodoptera litura* Fabricius (Lepidoptera: Noctuidae) is one of the most devastating agricultural pests, with a distribution in tropical and subtropical regions of the world [1, 2]. The insect has a wide host range and can feed on soybean, cotton, corn, tobacco, flax, tea, and some common vegetables [3, 4], which impacts crops can result in economic losses [5, 6]. In China, soybean, cotton, corn, and various vegetables are planted widely, and *S. litura* can feed on those crops and cause serious losses. Host plants biosynthesize a broad range of secondary metabolites, and these can influence the
growth, survival, and reproduction of herbivorous insects [7, 8]. In cotton plants, terpenoid gossypol [9, 10], tannins [11] and flavonoids [12] are the major classes of secondary metabolites. Glucosinolates are particularly abundant and they are important plant secondary metabolites in the Brassicaceae plant of Chinese cabbage [13, 14, 15, 16]. Fabaceae plants, including soybean (Glycine max (L.) Merr.), are rich in isoflavones, that act as part of the plant’s defense response [17, 18]. Derivatives of 1, 4-benzoxazin-3-one (BX) are the most common secondary metabolites in the plants of Poaceae, including maize (Zea mays) [19, 20].

Insect pests have developed various defense systems to detoxify secondary compounds found in host plants [21, 22]. A series of detoxification-related enzymes, including cytochrome P450 monooxygenases (P450s), glutathione S-transferases (GST) and UDP-glucosyltransferases (UGT), and ATP binding cassette (ABC) transporters, participate in the detoxification and metabolism of plant secondary metabolites [23]. P450s are an important group of enzymes involved in detoxification of host secondary metabolites in insects [24].

The insect midgut is regarded as the center of detoxification metabolism [25]. The expression of detoxification related genes found in gut tissues are expressed in the presence of plant secondary metabolites. Liu et al confirmed that cytochrome P450 of H. armigera played an important role in the metabolism of plant secondary substances [26]. Studies have shown that CYP6AE14 in the midgut of H. armigera was highly expressed when gossypol was added in the diet [27]. Gene slgste1 was up-regulated in the midgut of S. litura at the transcriptional and protein levels when the insect fed on Brassica juncea or diet containing phytochemicals [2].

The objective of the current study was to determine global changes in the midgut gene expression of S. litura larvae fed on different diets, as well as to compare the performance of S. litura larvae when fed on these diets. To investigate these transcriptional responses, we used a replicated RNA-Seq approach. In our analyses of the transcriptional responses of S. litura larval midgut, we focused mainly on the digestion-related and detoxification-related genes, with a particular emphasis on cytochrome P450 genes. In addition, two P450 genes were selected in the transcriptome, and the function was verified by RNA interference technology.
Results

Growth Rate On Different Diets

The larvae grew differently on the different diets. The initial weight of S. litura transferred to cabbage was significantly higher than those fed on the soybean and Tannin1 ($T_0$).

Two days later ($T_1$), the weight of S. litura fed on cabbage was 61.02 mg, which was significantly heavier than those fed on other diets, and especially for Tannin1 (29.91 mg).

At the last time point ($T_3$), the average weight of S. litura fed on cabbage (520.37 mg) was heaviest. Those larvae fed on soybean (365.38 mg), cotton (272.09 mg), and artificial diet (193.25 mg) had a middle range weight, and those fed on corn (91.01 mg), Tannin2 (49.24 mg), and Tannin1 (36.98 mg) were significantly lighter than the others (Fig. 1). Larvae fed on cabbage showed the highest growth rate at time points 1 and 2, and the lowest growth rate was found for the larvae fed on the artificial diet with 15 g of tannin at any time point (Table 1).

S. litura larvae had a higher growth rate and a higher final weight when fed on cabbage. S. litura larvae fed on Tannin1 had growth rates and final weights that were significantly lower.

| Treatment | $T_1$ | $T_2$ | $T_3$ |
|-----------|-------|-------|-------|
| Cotton    | 0.35 ± 0.01 de | 0.23 ± 0.00 a | 0.16 ± 0.00 a |
| Cabbage   | 0.40 ± 0.00 a  | 0.24 ± 0.00 a | 0.16 ± 0.00 a |
| Corn      | 0.37 ± 0.01 cd | 0.21 ± 0.00 b | 0.15 ± 0.00 a |
| Soybean   | 0.39 ± 0.01 ab | 0.23 ± 0.00 a | 0.16 ± 0.00 a |
| AD        | 0.38 ± 0.00 bc | 0.23 ± 0.00 a | 0.16 ± 0.00 a |
| Tannin1   | 0.33 ± 0.01 f  | 0.17 ± 0.01 d | 0.12 ± 0.00 a |
| Tannin2   | 0.35 ± 0.02 e  | 0.19 ± 0.01 c | 0.13 ± 0.00 a |

Those in the same column (mean ± SD) followed by different letters show significant difference at the P < 0.05 level by Duncan's multiple range test. AD: artificial diet. T: timepoint.

Illumina Sequencing, Sequence Assembly, And Gene Identification

Transcriptome analysis of 21 samples was completed, and clean reads were obtained. The percentage of Q30 bases was not less than 91.22%. Clean reads of each sample were sequenced with the designated reference genome of S. litura [28], and the alignment efficiency ranged from 83.84–89.66%. A total of 570 new genes were identified by filtering out sequences that were either too short (less than less than 150 nucleotide ORFs) or contained only a single exon. Sequence alignment of the new genes was conducted using BLAST software with the databases of NR, swiss-prot, GO, COG, KOG,
Pfam, and KEGG. KEGG Orthology results of the new genes were obtained using KOBAS2.0 [29]. After the prediction of amino acid sequences of the new genes, the HMMER [30] software was used to compare our results with the database of Pfam to obtain the annotation information of the new genes (Table 2). The transcriptome sequences had submitted to NCBI database and the SRA accession number was PRJNA528696

| Annotated databases | New Gene Number |
|---------------------|-----------------|
| COG                 | 27              |
| GO                  | 91              |
| KEGG                | 69              |
| KOG                 | 100             |
| Pfam                | 136             |
| Swiss-Prot          | 87              |
| eggNOG              | 223             |
| nr                  | 345             |
| All                 | 349             |

Differentially Expressed Genes Between Treatments

Gene expression was analyzed based on the comparison of results. Differentially expressed genes were identified according to their expression levels in different samples, and functional annotation and enrichment analyses were performed. S. litura midgut transcriptomes were studied by RNA-seq to determine changes in expression level in response to feeding from different diets. Gene expression levels for each replicate were assessed using principal component analysis (PCA), and the results revealed obvious differences from different diets. The gene expression in animals fed on artificial diets that contained 15 g/L and 7.5 g/L tannin clustered together and were far from the other treatments based on sample scores for the first (PC1) principal components. These results indicated that S. litura midgut modified gene expression levels in response to different diets (Supplement Fig. 1A).

The purpose of this study was to identify the key genes of S. litura that enable the insect to adapt to plant secondary metabolite. In order to investigate this, we carried out a pair-wise comparison of S. litura fed on other diets against artificial diets was carried out. Samples fed on cabbage, corn, cotton, soybean, and artificial diet (including tannin) exhibited 1,912 (964 up and 948 down), 1,395 (769 up and 626 down), 2,069 (1025 up and 1044 down), 2,998 (1516 up and 1482 down), and 1533 (838 up and 695 down) differentially expressed genes, respectively. Samples grown on artificial diet contain
15 g/L and 7.5 g/L tannin displayed similar gene expression, so those samples were combined as one sample set (Supplement Fig. 1B).

We next used a venn diagram to analyze the distribution of differentially expressed genes. Only 45 up-regulated and 64 down-regulated differentially expressed genes were involved in all treatments, and there were 238 up-regulated and 244 down-regulated differentially expressed genes when tannin treatments were excluded from venn diagram analysis (Supplement Fig. 1 C-F). The unique differentially expressed gene numbers were higher in cotton and soybean treatment samples, both in up-regulated and down-regulated genes analysis.

Identification Of Putative Pathway Related To Diet Adaption

The differentially expressed genes in pair-wise comparison of S. litura fed on various plant diets against artificial diets were subjected to KEGG pathway database analysis to discovery any significant changes to metabolic pathways genes. In top 5 up-regulated pathways, three enriched pathways (Metabolism of xenobiotics by cytochrome P450, Drug metabolism - cytochrome P450, and Pentose and glucuronate interconversions) were present across four treatments. Two enriched pathways (Glutathione metabolism and Ascorbate and aldarate metabolism) were present across three treatments. In the top 5 down-regulated pathways, one enriched pathway (Neuroactive ligand-receptor interaction) was shared across four treatments, and 5 enriched pathways (Endocytosis, Phosphatidylinositol signaling system, N-Glycan biosynthesis, Alanine, aspartate and glutamate metabolism and other glycan degradation) were shared across the treatments (Supplement Fig. 2).

The three KEGG pathways of metabolism of xenobiotics by cytochrome P450, drug metabolism cytochrome P450, and glutathione metabolism were mostly up-regulated in cabbage, corn, cotton, and soybean feed samples as compared to artificial diets, but were down-regulated in the artificial diet with tannin feed samples compared to the artificial diet alone.

The Character Of Genes Involved In Enriched Pathway

In up-regulated pathway, there were many similar genes involved in different pathways. Glutathione S-transferase and UDP-glucuronosyltransferase were two important genes present in up-regulated pathway. Both of these genes are important secondary metabolism detoxification enzymes in insects.
Glutathione S-transferase was present in three pathways out of the top 5 pathways determined in our analysis. We observed metabolism of xenobiotics by cytochrome P450, Drug metabolism-cytochrome P450 and Glutathione metabolism, and because metabolism of xenobiotics by cytochrome P450 and drug metabolism - cytochrome P450 were common in vertebrate animals and few reports in insect, and no cytochrome P450 genes were involved in those pathways, it indicated that glutathione metabolism was the main pathway, in which Glutathione S-transferase was involved.

A total of 17 glutathione S-transferase genes, 1 gamma-glutamyltransferase (GGT), 3 glutathione peroxidases (GPx), and 1 isocitrate dehydrogenase (IDH) gene were involved in the glutathione metabolism pathway. Generally, no special glutathione S-transferase genes were be found when insects were fed with different host-plants, which indicated that Glutathione metabolism pathway were the common detoxification pathway in S. litura. Meanwhile, elevated expression of genes involved in glutathione disulfide produced in feed on cabbage and cotton samples was observed, and gamma-glutamyltransferase genes were elevated when insects were fed on cotton (Supplement Fig. 3).

Identification Of Cytochrome P450s Related To Detoxification

As our focus was primarily on the response of detoxification-related genes of S. litura fed on various diets, we paid special attention to the cytochrome P450 gene family, which is involved in primary detoxification metabolism. The main cytochrome P450 gene involved in detoxification metabolism in insects is typically the special cytochrome P450. However, in the enriched up-regulated pathways from our analysis, no cytochrome P450 genes were present.

A total of 24 cytochrome P450 genes of which FPKM > 100 were chosen in S. litura midgut fed on plant hosts. Of these, 19 out of 24 cytochrome P450 genes belonged to the CYP6 family. Considering the cytochrome P450 genes found in the CYP family: 2 belonged to CYP4, 2 belonged to CYP9, and 1 belonged to CYP12. Unlike in Glutathione metabolism pathway, the expression of cytochrome P450 genes showed a clear response to different plant hosts. There were more induced cytochrome P450 genes when S. litura fed on cabbage and cotton than on other diets (12 genes associated with cabbage and 9 genes associated with cotton). Only 3 cytochrome P450 genes were higher expressed
when fed on artificial diet, 2 genes when fed on soybean, and 1 gene when fed on corn. Artificial diets containing tannin induced 2 cytochrome P450 genes to be expressed in insects, but suppressed the expression of 2 cytochrome P450 genes which had higher expression when feed on artificial diets alone (Supplement Fig. 4).

Expression pattern of CYP321A19 and CYP6AB60 in different developmental stages and tissues
In order to obtain the expression profile of CYP321A19 and CYP6AB60, RT-qPCR analysis showed that CYP321A19 and CYP6AB60 transcript was detected in all tissues and age. For CYP6AB60 gene, it is highly expressed at the 4th and 6th instar larva, and the expression level is lower at the 1st instar larva and pupa (Fig. 2A), and the expression levels were significantly higher in the midgut and fat body (Fig. 2B). Similarly, CYP321A19 was also highly expressed in 4th instar larvae (Fig. 2C), with the highest expression in fat body and midgut (Fig. 2D).

Expression of CYP321A19 and CYP6AB60 was induced by plant allelochemicals
The expression of CYP321A19 in the midgut and fat body of the larvae was increased and showed a significant difference with control artificial diet, when fed with an artificial diet containing quercetin (Fig. 3A). Similarly, when coumarin or soy isoflavones were added to the artificial diet, the expression levels of CYP6AB60 increased significantly compared with the control group (Fig. 3B, C).

Functional analysis of CYP321A19 and CYP6AB60 by RNAi
To evaluate the role of CYP6AB60 in the detoxification of plant allelochemicals, CYP321A19 and CYP6AB60 were silenced by RNAi technique in 4 instar larvae. Compared to the control group (injection of dsGFP), transcriptional levels of CYP321A19 were significantly decreased by 83.9% and 66.2% in the midgut and fat body at 72 h (Fig. 4A, B). Similarly, in both midgut and fat body tissues, CYP6AB60 transcript levels were significantly reduced following dsRNA injection (Fig. 4C-F).

When the larvae were exposed to the plant allelochemicals, the net weight gain on day 5 was lower in the treatment group than in the control group (CYP321A19: 0.57 g vs. 0.70 g) (Fig. 5A). Similarly, daily weight gain was lower in the treatment group than in the control group (Fig. 5B). Thus, CYP321A19 silenced larvae showed both net weight gain and daily growth significantly lower than the control group. In addition, larvae injected with dsCYP321A19 and fed with coumarin and soy
isoﬂavones, exhibited signiﬁcantly lower weight gains than dsGFP-injected controls exposed to the same allelochemicals (Fig. 5C-F).

**Identification Of Digestive Enzymes Related To Diet Adaption**

When *S. litura* fed on different diets, it faced different secondary metabolism stresses when deal with different nutrients. Proteinases, lipases, and carbohydrates make up the main digestive enzymes of insects [31]. In our transcriptome data, digestive enzymes in midgut were identiﬁed, and included proteinases (trypsin and chymotrypsin), lipases, and carbohydrates (alpha-amylase and glucosidase). A total of 34 trypsin genes and 4 chymotrypsin genes were found to be more highly expressed in *S. litura* midgut (FPKM > 100). We found more than 10 induced trypsin genes when *S. litura* fed on cotton, soybean, and artiﬁcial diet, but few trypsin genes were induced when fed on cabbage and corn. Most of the high expression trypsin genes were uniquely induced by diets. We found 8, 11, and 6 unique high expression genes when fed on artiﬁcial diet, soybean, and cotton, respectively, and 5 higher expression trypsin genes were induced when fed on cotton and soybean. Most high expression chymotrypsin genes were only detected in samples fed on artiﬁcial diets, cotton, and soybean.

Considering the lipid digestion and absorption process in the midgut, 13 higher expressing triacylglycerol lipase genes were (FPKM > 100) were found. The triacylglycerol lipase genes were induced when insects were feed on corn, cotton, and soybean, and had the highest induced gene numbers when fed on soybean (triacylglycerol lipase genes). When fed on cotton, a total of 6 triacylglycerol lipase genes were induced. All the corn-induced triacylglycerol lipase genes were the same as those induced by soybean, except LOC111355064. There were 2 cotton-induced triacylglycerol lipase genes that were the same as soybean-induced genes, but there were no shared triacylglycerol lipase genes with corn.

During carbohydrate digestion and absorption process in the *S. litura* midgut, amylases and glucosidases were the main observed differentially expressed genes. A total of 2 amylase and 12 glucosidase genes showed higher expression (FPKM > 100). We found that 1 alpha-amylase was induced in corn and cotton fed insects. However, there were no observed induced alpha-amylase genes in other diet treatments. In corn and cotton fed samples, there was 1 alpha-amylase gene with
higher induced expression. No other diets showed induced alpha-amylase gene expression (Supplement Fig. 5).

Quantitative real-time PCR validation

To verify the transcriptome data, we examined the relative expression levels of P450 (LOC111350062, LOC111358240, LOC111351731), UDP-glucosyltransferase (LOC111348983, LOC111348863, LOC111348860) and GST (LOC111354038, LOC111351682, LOC111352663, LOC111351550). The qRT-PCR of these unigenes showed that the results were consistent with the DGE results (Figure 6).

Discussion

Insect herbivores can feed on their host plants for development and survival. Due to host plant nutrition and allelochemicals, polyphagy insects show differential fitness to the plant-hosts [32]. Studies have shown that plant secondary metabolites can have positive effects on the survival and growth rates of insects [33, 34, 35]. Our larval development assays suggested that cabbage is the best host plant for S. litura, with a higher growth rate and final weight than other host plants. S. litura fed on artificial diets with tannin had a lower growth rate and obtained a lighter final weight in our study.

In the study, we chose host-plants that contain various kinds of secondary metabolites. Gossypol, tannin, and flavonoids are the major secondary metabolites in cotton plants, and glucosinolates and isoflavones are rich in Chinese cabbage and soybean. Derivatives of 1, 4-benzoxazin-3-one are the common secondary metabolites found in maize plants. The explanation for the differences of larval development on different diets could be due to the effect of differential secondary metabolites that the diets contain. A major question in plant-insect interactions is how insect herbivores cope with secondary metabolites compounds in diverse host plants [36]. As insect midgut is the main location to digestive food and detoxification [37], and the mechanisms S. litura use to cope with toxic compounds in diverse host plants are not well understood [38], we used a feeding assay and RNA-Seq of S. litura larval midgut to determine the genes used by S. litura to cope with secondary metabolites associated with different diets.

Based on S. litura genomic data [28], high-throughput sequencing was an efficient research tool to
better understand the molecular mechanisms behind adoption of host-plants. Our results demonstrated that S. litura could develop on different diets and the transcriptional responses of midgut were related to the host diet that S. litura was fed on. Generally, glutathione S-transferase genes and Cytochrome P450 genes were the most differential expressed genes involved in detoxification, while, proteinases, lipases, and carbohydrases were the most differentially expressed genes involved in the digestive system.

A total of 47 Glutathione S-transferase genes were identified in S. litura genomic data [28]. In this study, 17 glutathione S-transferase genes were found to be high expressing in S. litura midgut when fed on different host-plant. It has been demonstrated that glutathione S-transferase can detoxify many plant allelochemicals, and can be induced by plant allelochemicals [21, 39]. In this study, the expression of some glutathione S-transferase genes showed high FPKM valves, but no special response glutathione S-transferase genes were be found when fed with different diets, which indicated that glutathione metabolism pathways were the common detoxification pathway in S. litura. At same time, genes in glutathione disulfide produced in feed on cabbage and cotton samples also showed high FPKM valves. One possible reason for this is that more glutathione disulfide was needed for allelochemical detoxification. As the gamma-gluamyltransferase genes, which are located at last step of glutathione metabolism pathway, were higher expressed in insects fed on cotton, cotton allelochemical detoxification may require more steps than detoxification in S. litura midgut fed on other host-plants.

In insects, cytochrome P450 genes are an important gene family in the detoxification of exogenous or endogenous compounds, including plant secondary metabolites [40, 41]. A total of 138 cytochrome P450 genes were identified in S. litura genomic data [28]. In this study, 17 glutathione S-transferase genes were highly expressed in S. litura midgut when fed on different host-plants. 21 out 61 Clan 3 cytochrome P450 genes were highly expressed in S. litura midgut when fed on different host-plants, and most of these genes belong to the CYP6 gene family. In insect, the CYP6 gene family is mostly involved in plant secondary metabolites detoxification. Here, we selected two P450 genes from the transcriptome and named them CYP6AB60 and CYP321A19. They all showed the highest expression
levels in the 4th instar larva, and the highest expression levels in the 4th instar larva midgut and fat body. When the larvae were exposed to an artificial diet containing quercetin or coumarin and soy isoflavones, the expression levels of the selected genes were significantly up-regulated. Similar results have been reported that CYP6AE14 and CYP6AE11 were significantly up-regulated when H. armigera fed on artificial diet with higher concentration of gossypol [42]. CYP6B46 can be induced when Manduca sexta larvae feed on Nicotiana wild-type plants, which can produce nicotine [43]. The expression of CYP314A1, CYP315A1, CYP18A1, CYP307A1, and CYP306A1 were found to be induced by 2-tridecanone [44]. The xanthotoxin can induce the expression of CYP9A genes from larval Manduca sexta midgut [45]. The above research results are similar to the results of this study.

In the detoxification of plant secondary metabolites, cytochrome P450 can be classed into specialists and generalists [21]. In this study, most high expressing cytochrome P450 genes may be specialists to different host-plant, and those cytochrome P450 gene family may be involved in primary detoxification metabolism. As specialists have a highly efficient and specialized detoxification system, S. litura has a wide host range and significantly impacts agricultural production. The specialist cytochrome P450 genes may represent a potential target site for the development of pest controls.

RNAi technology has been widely used to reveal the role of cytochrome P450 in drug resistance, secondary metabolites and pesticide detoxification [46]. Studies have shown that when Manduca sexta larvae are fed plant material expressing CYP6B46-specific dsRNA, the level of this transcription decreases and larval growth is hindered [47]. In addition, studies have shown that RNAi silences CYP307A1 and blocks molting steroid synthesis, suggesting that this gene is required for molting steroid biosynthesis in anopheles gambiae. Silencing CYP6B7 alone or CYP6B7 in combination with CPR and/or Cyt-b5 increased the sensitivity of bollworm to fenvalerate, indicating that CYP6B7, CPR and cyt-b5 were synergistic in the metabolic enhancement of fenvalerate and played an important role in the resistance of bollworm to fenvalerate [48]. After RNAi silencing of CYP321E1, Plutella xylostella has increased sensitivity to chloroaniline, with a mortality rate of up to 70% 49 (Hu et al., 2014). In our study, RNA interference (RNAi) was used to investigate the function of selected target genes. After injection of dsCYP6AB60 and dsCYP321A19, the tolerance of the 4th instar larva of
Spodoptera litura to plant allelochemicals (quercetin, coumarin, soy isoflavones) was significantly reduced.

**Conclusion**

In our study, some P450 genes of S. litura fed on different host plants and artificial diets with tannin were up-regulated compared with those fed on artificial diets, but all glutathione S-transferase genes were down-regulated. The bioassay data showed that tannin can inhibit the growth of S. litura, which indicated that when S. litura faced harmful allelochemicals, its primary detoxification metabolism optoins were narrowed to Cytochrome P450 genes, and the detoxification metabolism of glutathione metabolism was less utilized.

In addition of detoxification metabolism, digestion was another important function in S. litura. In this study, the expression of digestive enzymes genes was assessed, and we found that, in general, the expression of genes encoding proteinases, lipases, and carbohydrases corresponded to food nutritional composition.

**Methods**

**Insect Rearing**

Insects were purchased from Henan Jiyuan Baiyun Industry Co., Ltd, and then S. litura were fed on artificial diet: 16.7 g of agar, 100 g of soybean flour, 100 g of wheat germ flour, 100 g of oatmeal, 60 g of yeast powder, 40 g of sucrose, 6 g of ascorbic acid, 2 g of sorbic acid, 2 g of methyl paraben, 3 g of sodium benzoate, 8.35 mL of KOH (4M), 15 mL of decavitamin (1.528 g of nicotinic acid, 1.525 g of calcium pantothenate, 0.764 g of riboflavin, 0.382 g of niacin thiamine, 0.382 g of pyridoxine hydrochloride, 0.382 g of folic acid, 0.305 g of biotin and 0.003 g of cyanocobalamine per 3.3 L of water) per 1 L water. Insect rearing took place in an artificial climate chamber (26 ± 1 °C, 65 ± 5% relative humidity, Light : Dark = 14:10) and mixed populations were used in all experiments. The insects were continuous feeding for 3 generations and then the larvae were used for the feeding assays and RNA-Seq.

**Feeding Assays**

The following plant species were used for our experiments: cotton (zhong 49, provided by institute of
cotton research of CAAS), corn (Yuyu 22, Charoen Pokphand Group), soybean (Zhonghuang 35, Beijing huinong fumin technology co. LTD) and cabbage (Jingfeng 1, Fuyichun seed sales co. LTD). All plants were planted in a climate chamber maintained at (26 ± 1) °C, 65 ± 5% humidity, and a 14:10 (Light : Dark) photoperiod. Plants that were 1 month old were used for feeding assays. Additionally, 15 g and 7.5 g tannin were added to artificial diets as treatment 1 (Tannin1) and 2(Tannin2), respectively. After that, three treatments of quercetin, coumarin and soybean isoflavone artificial feed (1 mg/g) were set up respectively to verify the function of P450 gene.

S. litura larvae were reared on an artificial diet until they grew to the third instar (L3), then they were transferred to host plants (cotton, corn, soybean or cabbage) and Tannin1 or Tannin2. Each leaf or small piece of artificial diet contained one larva; there were 3 replicates for each treatment, and each treatment included 20 larvae. Larvae were allowed to feed on the excised leaves of plants or the artificial diet for 6 consecutive days. For the duration of the experiment, larval weight was recorded and fresh leaves were replaced every second day. Growth rates [23] were calculated per unit time for each treatment.

Midgut Tissue Collection

Newly hatched larvae of S. litura were transferred to living plants of cotton, corn, soybean, cabbage, and artificial diet containing tannin. Midguts were dissected after the larvae developed to fourth instar. Insects were dissected in 0.7% saline solution. Dissected tissues without contents were quickly frozen in liquid nitrogen and kept at -80 °C prior to RNA isolation. There were 3 replicates for each treatment and each replicate included 12 larvae.

RNA Isolation And Illumina Sequencing

RNA-Seq experiments were carried out with RNA isolated from larvae reared on different diets. Total RNA was extracted by TRizol® Reagent (Life Technologies, USA) according to the manufacturer's instructions. The quantity and quality of the RNA were assessed by 1.0% agarose gel electrophoresis and absorbance at 260 nm on a NanoDrop 2000c spectrophotometer (Thermo Scientific, USA). Then samples were sent to Beijing Genomics Institute (Shenzhen, China) for cDNA library construction and Illumina sequencing. Sequencing was performed using an Illumina Hi-Seq 4000 sequencer (San Diego,
RNA-seq Data Analysis

Raw reads of fastq format were initially processed using in-house perl scripts. The clean reads were then obtained by removing reads containing adapter, poly-N, and low quality reads from raw reads. The Q20, Q30, GC-content, and sequence duplication level of the clean reads were then calculated. These clean reads were then mapped to the reference genome sequence using hisat2 tools software. The databases of Nr, Nt, Pfam, KOG/COG, Swiss-Prot, KO, and GO were used for gene function annotation. Gene expression levels were estimated by fragments per kilobase of transcript per million fragments mapped. Differential expression analysis of the two groups was performed using the DESeq R package (ver. 1.10.1). The resulting P values were adjusted using Benjamini and Hochberg’s approach for controlling the false discovery rate. Genes with an adjusted P-value < 0.05 found by DESeq were assigned as differentially expressed. Gene Ontology (GO) enrichment analysis of the differentially expressed genes (DEGs) was implemented by the GOseq R package based on Wallenius non-central hyper-geometric distribution [50]. KOBAS [51] software was used to test the statistical enrichment of differentially expressed genes in KEGG pathways.

Quantitative Real-time PCR Validation

Total RNA was extracted as described above and Reverse Transcription System (Takara) was used for cDNA synthesis. The relative expression levels of P450, UDP-glucosyltransferase and GST were conducted by quantitative real-time PCR (qRT-PCR).

Two differentially expressed genes, LOC111355823 and LOC111355088, were identified from the transcriptome. We named them separately CYP321A19 and CYP6AB60. To clarify the expression pattern of the CYP321A19 and CYP6AB60 sequences, four tissues (midgut, fat body, cuticle, and hemolymph) were dissected from 1-day-old 4th instar larva of S. litura. For analyses of the CYP321A19 and CYP6AB60 expression pattern at different development stages, we collected the samples which were at day 2 of each development stage, including eggs (100 eggs per RNA extraction), first to sixth instar larvae (6 larvae per RNA extraction) and pupae (6 pupae per RNA extraction) of S. litura (6 replicates for each sample above). RNA extraction and cDNA synthesis as
described above. To elucidate the expression profiles of CYP321A19 sequences when exposed to plant allelochemicals. Newly molted fourth instar larvae of S. litura were fed on artificial medium containing quercetin. Dissecting the midgut and fat body from surviving larvae after fed on 72 h. The samples were then immediately frozen in liquid nitrogen and stored at −80 °C to extract RNA and further expression analysis. Twenty-four individuals were used for each treatment. Similarly, the midgut and fat bodies of 4th instar larvae that ate coumarin and soybean isoflavones were taken to analysis the expression profile of CYP6AB60. Three independent RNA extractions (representing three biological replicates) were performed for all treatments. Subsequent real-time quantitative PCR (RT-qPCR) were equivalent to those described above.

Beacon Designer 7.7 was used for designing qRT-PCR primers of the selected unigenes. Unigenes and primers were listed (Table S1). GAPDH and RpL10 were chosen for reference genes [52]. Cycle threshold (CT) values were collected after completion of reaction and the relative expression levels were analyzed using the $2^{-\Delta\Delta CT}$ method [53]. All reactions were performed in triplicate (technical repeats) with three independent biological replicates.

dsRNA Synthesis

Using the cloned CYP321A19 and CYP6AB60 sequences as templates, an in vitro transcription reaction template was obtained by PCR amplification. Two pairs of primers (Table S1) were used to amplify CYP321A19 and CYP6AB60. The amplification conditions comprised 5 cycles of 95 °C for 10 s, 50 °C for 15 s, 72 °C for 30 s, 20 cycles of 95 °C for 10 s, 50 °C for 15 s (Increase 0.5 °C by each cycle), 72 °C for 30 s, 10 cycles of 95 °C for 10 s, 60 °C for 15 s, 72 °C for 30 s. PCR products were purified using the Promega SV Gel and PCR clean up system (Promega, Madison, WI, USA) and DNA concentrations were determined using UV-visible spectrophotometry. According to the instructions, dsRNAs (dsCYP321A19 and dsCYP6AB60) corresponding to CYP321A19 and CYP6AB60 were synthesized using the T7 RiboMAX ™ Express RNAi System (Promega, Madison, WI, USA). In addition, dsRNA of the GFP gene used as a negative control was synthesized by the same method as above, and primers used were listed (Table S1). The obtained dsRNA was detected by ultraviolet spectrophotometry and the purity and integrity were evaluated by agarose gel electrophoresis. The
final concentrations of dsRNA were adjusted to 1.5 µg·µL⁻¹ by RNase-free water and kept at -80 °C. For RNAi bioassays, the final concentrations of dsRNA were adjusted to 1.5 µg/µL using DEPC-treated (RNase-free) water prior to use. All dsRNA injection experiments used 4th instar larvae (day 1 and hunger for 4 h) of S. litura, with 2µL (3.0 µg) of dsRNA injected into the distal second segment of the abdomen by manual microinjector, while the control group was injured with an equivalent volume of dsGFP. The treated larvae were fed with artificial feed supplemented with plant allelochemicals (The larvae injected with dsCYP321A19 were fed with quercetin, and the larvae injected with dsCYP6AB60 were fed with coumarin and soybean isoflavones).

For the RNAi efficiency evaluation, the midguts and fat bodies of the surviving larvae (6 larvae) were taken at 24 h, 48 h and 72 h after the last treatment. Calculate the net weight increase (weight of each day minus weight on day 0) and weight increase per day (weight of each day minus weight of the previous day). All experiments were performed in triplicate. For each replicate, midguts and fat bodies from six larvae of S. litura were collected for total RNA extraction. The RNA extraction and RT-qPCR procedures were described above.

Data analysis

qRT-PCR data are presented as mean ± standard error (SE). Statistically significant differences (p < 0.05) of bioassay data and qRT-PCR data were identified by one-way ANOVA followed by Duncan's multiple range test.

Declarations

Ethics approval and consent to participate:

This article does not contain any studies with human participants or animals (others than insects) performed by any of the authors.

Consent for publication:

Not applicable.

Availability of data and materials:

All data generated or analysed during this study are included in this published article and its supplementary information files.
Competing interests:
The authors declare that they have no conflict of interest.

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Authors' contributions:
LW and PZ are co-first authors, acquired, analyzed, and interpreted the data and drafted the manuscript. JJC and SZ conceived and designed the study. SZ, JYL, CYW, XZZ, LJZ, JCJ, KXZ, DYL and JJC were involved in data acquisition and interpretation and in manuscript revision. All authors read and approved the final manuscript.

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**Supplementary Figure & Table Captions**

**Supplement figure 1** (A) Principal component analysis (PCA) for gene expression levels of each treatment. (B) The up-regulated and down-regulated gene numbers of samples feed with cabbage, corn, cotton, soybean, and artificial diet with tannin compared with artificial diet. (C) Venn diagram showing the number of up-regulated and down-regulated genes of samples fed with cabbage, corn, cotton, soybean, and artificial diet with tannin compared with artificial diet. C and E respresnt up-regulated differentially expressed genes; D and F represent down-regulated differentially expressed genes. G0: Artificial diet vs Cabbage; G1: Artificial diet vs Corn; G2: Artificial diet vs Cotton; G3: Artificial diet vs Soybean; G4: Artificial diet vs Artificial diet with tannin.

**Supplement figure 2** The up-regulated and down-regulated pathways of samples fed with cabbage, corn, cotton, soybean, and artificial diet with tannin compared with artificial diet alone.

**Supplement figure 3** The metabolic pathway of Glutathione. Gene expression of glutathione S-transferase when S. litura fed with different diets. Values are based on log2-transformed FPKM values relative to the median intensity of all contigs (red= up-regulation; green = down-regulation).

**Supplement figure 4** Gene expression of cytochrome P450 when S. litura was fed with different diets. The plot is based on log2-transformed FPKM values relative to the median intensity of all contigs (red= up-regulation; green = down-regulation).

**Supplement figure 5** Gene expression of digestive enzymes when S. litura were fed with different diets. Plots are based on log2-transformed FPKM values relative to the median intensity of all contigs (red= up-regulation; green = down-regulation).

**Table S1** Primers used in this study.

Figures
Figure 1

The weight of *S. litura* larvae fed on different diets. The data are mean ± SD and different letters indicate significant difference at the 0.05 level by the Duncan's multiple range test.

AD: artificial diet. T: timepoint.
Expression levels of CYP6AB60 in different developmental stages (A), in different tissues (B), and CYP321A19 in different developmental stages (C), in different tissues (D). All larvae were fed an artificial diet. RT-qPCR analysis was used to determine the relative transcript levels for each gene. Data shown as means ± SE derived from three biological replicates. Different letters above bars indicate significant differences (p < 0.05) according to the Duncan’s multiple range test (same as below).
Figure 3

Expression levels of CYP321A19 sequences when exposed to quercetin (A), and CYP6AB60 sequences exposed to coumarin (B), soy isoflavones (C). Data shown as means ± SE derived from three biological replicates (Student’s t-test, ** p < 0.01, * p < 0.05, same as below).
Figure 4

Effect of CYP321A19 and CYP6AB60 silencing on S. litura resistance to allelochemicals. Knockdown reduction rates of CYP321A19 in midgut (A) and fat body (B) after injection of dsRNA. Relative expression of CYP6AB60 in the midgut and fat bodies of larvae exposed to coumarin (C, D) and soy isoflavones (E, F) after dsRNA injection. Control larvae were injected with dsGFP.
Figure 5

Changes in larval body weight of S. litura following smear of dsRNA. After dsCYP321A19-injected larva, the net weight increased (A) and the weight increased per day (B) after feeding with quercetin. After dsCYP6AB60-injected larva, the net weight increased (A) and the weight increased per day (B) after feeding with coumarin (C, D) and soy isoflavones (E, F).
Figure 6

Gene expression verification of 10 genes of P450, UDP-glucosyltransferase and GST. Red, qRT-PCR results; Blue, DGE data.

Supplementary Files

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