The Key Role of Fatty Acid Synthase in Lipid Metabolism and Metamorphic Development in a Destructive Insect Pest, *Spodoptera litura* (Lepidoptera: Noctuidae)

Yan Song 1, Fengming Gu 1, Zhixiang Liu 1, Zongnan Li 1, Fu’an Wu 1,2 and Sheng Sheng 1,2,*

1 School of Biotechnology, Jiangsu University of Science and Technology, Zhenjiang 212100, China
2 The Key Laboratory of Silkworm and Mulberry Genetic Improvement, Ministry of Agriculture and Rural Affairs, Sericultural Research Institute, Chinese Academy of Agricultural Science, Zhenjiang 212100, China
* Correspondence: parasitoids@163.com

Abstract: Fatty acid synthase (FAS) is a key enzyme in the lipid synthesis pathway, however, its roles in insects remain largely unknown. Here, we firstly identified two FAS genes from the transcriptome dataset of the general cutworm *Spodoptera litura*, which is a destructive insect pest of many crops. Both *SIFAS1* and *SIFAS2* were highly expressed in third instar larvae and in their fat bodies. Then, we successfully silenced *SIFAS1* in third instar larvae and the content of α-linolenic acid and triglyceride was significantly decreased. Besides that, the effect of FAS on the metamorphic development in *S. litura* was evaluated. The results indicate that after silencing *SIFAS1*, the survival rates of *S. litura* larvae decreased significantly compared to the control groups. Silencing *SIFAS1* in fifth instar larvae resulted in more malformed pupae and adults, and the emergence rates were significantly reduced. Furthermore, the ecdysone content in the haemolymph of fifth instar larvae was significantly decreased after silencing *SIFAS1*. In addition, knocking down *SIFAS1* significantly alters the expression of other key genes in the lipogenesis pathway, implying that FAS has an impact on the lipogenesis pathway. The present study deepens the understanding of FAS in insects and provides novel potential targets for managing insect pests.

Keywords: fatty acid synthase (FAS); *Spodoptera litura*; lipid metabolism; metamorphosis; RNA interference

1. Introduction

Lipids are essential in organisms and play multiple roles such as energy storage, signal transduction, hormone synthesis, and cell membrane component formation, and they are vital for maintaining normal life activities. Lipid synthesis is a highly complex process catalysed by a battery of enzymes, among which fatty acid synthases (FASs) are key and multifunctional enzymes in fatty acid synthesis and lipid metabolism. There are two types of FASs existing in organisms. Type I FASs are predominantly found in animals and fungi. The type I FASs of fungi are encoded by two genes that assemble an α6β6-heterododecamer [1], but mammalian type I FASs are expressed as a single protein and assemble as a homodimer [2]. In contrast, type II FASs are common in bacteria and eukaryotic organelles, found especially in chloroplasts and mitochondria, and are expressed as discrete proteins in the cytosol [3]. The general FAS monomer contains seven functional domains including β-ketoacyl synthase (KS), malonyl/acetyl transferase (MAT), β-hydroxyacyl dehydratase (DH), enoyl reductase (ER), β-ketoacyl reductase (KR), acyl carrier protein (ACP), and thioesterase (TE) [4].

Until now, the identification, as well as the function, of FAS genes has only been studied in a narrow range of insect species [5,6]. The first reported insect FAS was purified from the fat body of *Drosophila melanogaster* and there are three FAS genes (*FASCG3523, FASCG3524, FASCG17374*) identified from *D. melanogaster* [7]. *FASCG3523* is expressed in the fat body while *FASCG3524* and *FASCG17374* are expressed in oenocytes. Silencing *FASCG3524*...
and FAS<sup>CG17374</sup> can induce the lethal phenotype [8]. Generally, FAS genes are mainly expressed in the fat body and oenocytes, and some others are also expressed in different tissues with different biological functions [9]. In the fat body, FASs are responsible for the biosynthesis and storage of lipids, mainly deposited in the form of triglyceride, and transport it to different tissues in the form of diacylglycerols through the haemolymph [10]. In Aedes aegypti, the expression level of FAS1 was highest in the fat body after 48 h, but highest in the ovary 72 h after blood feeding. Besides that, triglyceride and phospholipid levels of A. aegypti adults were significantly reduced after silencing FAS1 [11].

Triglyceride is the main component of lipids and FAS regulates de novo lipogenesis by converting acetyl-CoA to palmitate, further leading to the production and storage of triglyceride [12]. The fatty acid synthesis pathway also involves other enzymes, such as acetyl-CoA carboxylase (ACC), desaturases (Desat), and lipase. Furthermore, considering the importance of triglyceride in insect development, the exact roles of FASs still receive less attention. Previous research revealed that silencing GpylFAS1 in a mulberry pest, Glyphodes pyloalis, significantly reduced the content of α-linolenic acid, resulting in more malformed pupa and lower emergence rates [13]. Studies have shown that two hormones in insects, juvenile hormone and ecdysone, also participate in the regulation of triglyceride metabolism [14]. However, it remains unknown whether FAS regulates the dynamics of juvenile hormone and ecdysone during metamorphic development.

The general cutworm, Spodoptera litura (Fabricius) (Lepidoptera: Noctuidae) is a serious worldwide polyphagous pest. It causes heavy damage to dozens of crops, such as potato, tomato, corn, and mulberry trees every year [15]. At present, studies on lipid metabolism in S. litura have been reported [16], but there is no reference to the role of FASs in regulating lipid metabolism in S. litura.

In the present study, we obtained two FAS genes from the transcriptome of S. litura and focused on the role of SIFASs in the synthesis of lipids and in the development and metamorphosis of S. litura larvae. The present study reveals the molecular mechanism of FAS during the development of S. litura, providing a novel potential target for the integrated biological control of pests.

2. Results

2.1. Gene Identification and Sequence Analysis of FAS Genes in S. litura

Two FAS genes were identified from our previously constructed S. litura larvae transcriptome database and named SIFAS1 (LOC111359194) and SIFAS2 (LOC111359981). The sequences of SIFAS1 and SIFAS2 contained complete ORFs of 6240 and 7269 bp which encoded 2079 and 2422 amino acid residues, respectively. The protein sequences of SIFAS1 have eight functional domains, including Ketoacyl-synt (ks, pfam00109), Ketoacyl-synt_C (KsC, pfam02801), KAsynt_C_assoc (KCa, pfam16197), Acyl_transf_1 (At1, pfam00698), PS-DH (PF14765), PKS_ER (smart00829), PKS_KR (smart00822), and PP-binding (Pb, pfam00550). However, SIFAS2 contains nine functional domains, with an additional Thioesterase domain (Th, pfam00975), compared to SIFAS1 (Figure 1). Sequence alignment showed 35.63% amino acid identity between SIFAS1 and SIFAS2 (Figure S1). Phylogenetic analysis of FASs revealed that SIFAS1 and FASs in Agrotis ipsilon and Trichoplusia ni were clustered into a subcluster, and SIFAS2 was clustered with FASs in Spodoptera frugiperda into a subbranch (Figure 2).
Figure 1. Domain schematic of SlFAS1 (A) and SlFAS2 (B) in *S. litura*. KS, Ketoacyl-synt (pfam00109); Ks C, Ketoacyl-synt_C (pfam02801); KC a, KAsynt_C_assoc (pfam16197); At1, Acyl_transf_1 (pfam00698); PS-DH (PF14765); PKS_ER (smart00829); PKS_KR (smart00822); Pb, PP-binding (pfam00550); Th, Thioesterase (pfam00975).

Figure 2. Phylogenetic tree of FAS amino acid sequences in different species. *Spodoptera litura* (Sl), *Bombyx mori* (Bm), *Drosophila melanogaster* (Dm), *Spodoptera frugiperda* (Slf), *Helicoverpa zea* (Hz), *Trichoplusia ni* (Tn), *Ostrinia furnacalis* (Of), *Helicoverpa armigera* (Ha), *Plutella xylostella* (Px), *Agrotis ipsilon* (Ai).

2.2. Expression Patterns of SIFAS1 and SIFAS2

*SIFAS1* had the highest expression level in adults, and the expression level in males was significantly higher than that in females. In the immature stages, the expression level of *SIFAS1* was higher in third instar larvae (Figure 3A). In contrast, *SIFAS2* had the highest expression level in third instar larvae compared to other stages followed by first and fourth instar larvae (Figure 3B). Besides that, both *SIFAS1* and *SIFAS2* were expressed at the highest levels in the fat body and the lowest in the midgut (Figure 3C,D).
Gas chromatography determination showed that the content of methyl α-linolenate was significantly reduced 48 h after silencing SIFAS1, but no obvious change was observed
at 24 h compared to the dsGFP-injection group (Figure 5A). Moreover, after silencing SIFAS1, the content of methyl palmitate, methyl oleate, and methyl linoleate did not change significantly compared to the control cohorts at both 24 and 48 h (Figure 5B–D).

Figure 5. Effect of silencing SIFAS1 on the fatty acid content in S. litura larvae. (A) Effect of silencing SIFAS1 on methyl α-linolenate content. (B) Effect of silencing SIFAS1 on methyl palmitate content. (C) Effect of silencing SIFAS1 on methyl oleate content. (D) Effect of silencing SIFAS1 on methyl linoleate content. Significant differences are indicated by asterisks (* p < 0.01; ns: no significant differences; non parametric Mann–Whitney test).

2.4. Effect of Silencing SIFAS1 on Lipid Accumulation in S. litura Larvae

To understand the effect of silencing SIFAS1 on lipid accumulation in third instar S. litura larvae, we detected the triglyceride content after dsSIFAS1 injection. Nile red staining revealed that the density of lipid droplets was significantly reduced in the fat body at 24 and 48 h compared to the control group after dsRNA injection, respectively (Figure 6A). Besides that, the triglyceride content of the fat body decreased significantly at 24 and 48 h after dsSIFAS1 injection compared to the dsGFP group, respectively (Figure 6B).
At the same time, we also determined the content of juvenile hormone and ecdysone in the haemolymph of fifth instar larvae after silencing SlFAS1. More morphologically abnormal adult moths were observed in the dsSlFAS1-injection group than the dsGFP-injection group (Figure 8C, D, G). At the same time, we also determined the content of juvenile hormone and ecdysone in the haemolymph of fifth instar larvae after silencing SlFAS1; the results showed that the content of ecdysone decreased significantly 24 h after silencing SlFAS1, but no significant change was observed 48 h post knocking. The emergence rate of pupae derived from the dsSlFAS1-injection group was significantly lower than that of the dsGFP-injection group (Figure 8C, D, G). Subsequently, the emergence rate of pupae derived from the dsSlFAS1-injection group was significantly lower than that of the dsGFP-injection group (Figure 7A). In addition, compared to the dsGFP-injection group, the growth trajectory in terms of body weight at each time point between the dsSlFAS1-injection and dsGFP-injection groups was not significantly different (Figure 7B).

### 2.5. Effect of Silencing SlFAS1 on the Development of S. litura Larvae

To study the effect of SlFAS1 on the development of S. litura larvae, we analysed survival rates and the growth trajectory of body weight after dsSlFAS1 injection in third instar larvae. The results showed that the survival rates of dsSlFAS1-injection larvae were significantly reduced by approximately 45% compared to the dsGFP control (Figure 7A). In addition, compared to the dsGFP-injection group, the growth trajectory in terms of body weight at each time point between the dsSlFAS1-injection and dsGFP-injection groups was not significantly different (Figure 7B).

After the dsSlFAS1 injection in the fifth instar larvae, the proportion of abnormal pupae was significantly higher compared to the dsGFP-injection groups (Figure 8A, B, E). Subsequently, the emergence rate of pupae derived from the dsSlFAS1-injected fifth instar larvae was significantly lower than that of the dsGFP control group (Figure 8F). Meanwhile, more morphologically abnormal adult moths were observed in the dsSlFAS1-injection group than in the dsGFP-injection group (Figure 8C, D, G). At the same time, we also determined the content of juvenile hormone and ecdysone in the haemolymph of fifth instar larvae after silencing SlFAS1; the results showed that the content of ecdysone decreased significantly 24 h after silencing SlFAS1, but no significant change was observed 48 h post knocking.
down SlFAS1 (Figure 9A). By contrast, the content of juvenile hormone was changed neither 24 h nor 48 h after silencing SlFAS1 (Figure 9B).

Figure 7. Effect of SlFAS1 on the development of S. litura...content in the haemolymph of fifth instar S. litura larvae. Significant differences were indicated by asterisks (* p < 0.05; ** p < 0.01; *** p < 0.001); scale bars in A and B: 2 mm; scale bars in C and D: 2 mm.

Figure 8. Morphology of pupae and adults after knocking down SlFAS1 in fifth instar S. litura. (A) Normal pupae after dsGFP injection. (B) Abnormal pupae after dsSlFAS1 injection. (C) Normal adults after dsGFP injection. (D) Abnormal adults after dsSlFAS1 injection. (E) Effect of silencing SlFAS1 on the proportion of abnormal pupae in S. litura. (F) Effect of silencing SlFAS1 on emergence rate in S. litura. (G) Effect of silencing SlFAS1 on the proportion of abnormal adults in S. litura. Significant differences were compared using a one-way analysis of variance (ANOVA). Significant differences were indicated by asterisks (* p < 0.05; ** p < 0.01; *** p < 0.001); scale bars in A and B: 2 mm; scale bars in C and D: 2 mm.

Figure 9. Effect of knocking down SlFAS1 on the content of juvenile hormone and ecdysone in the haemolymph of fifth instar larvae. (A) Effect of silencing SlFAS1 on ecdysone content in the haemolymph of fifth S. litura larvae. (B) Effect of silencing SlFAS1 on juvenile hormone content in the haemolymph of fifth instar S. litura larvae. Significant differences were compared using a one-way analysis of variance (ANOVA). Significant differences were indicated by asterisks (** p < 0.01; ns, no significant differences).
2.6. Effect of Silencing SIFAS1 on the Expression of other Key Genes in the Lipogenesis Pathway

To explore the effect of silencing SIFAS1 on the lipogenesis pathway, the expression levels of other key genes in the fatty acid synthesis pathway after dsSIFAS1 injection were validated by qRT-PCR. The expression levels of SIFAS2 and SIdesat were significantly upregulated 24 h after dsSIFAS1 injection, and the expression of SIdesat was also increased 48 h after dsSIFAS1 injection. The expression levels of SIACC and SILIPase were significantly upregulated 48 h after dsSIFAS1 injection; in addition, SILIPase was significantly downregulated 24 h after silencing SIFAS1 (Figure 10).

![Graphs A, B, C, D](image)

**Figure 10.** Effect of silencing SIFAS1 on the expression levels of other key genes in the lipid synthesis pathway in S. litura larvae. (A) SIFAS2 expression levels after dsSIFAS1 injection. (B) SIdesat expression levels after dsSIFAS1 injection. (C) SIACC expression levels after dsSIFAS1 injection. (D) SILIPase expression levels after dsSIFAS1 injection. One-way analysis of variance (ANOVA) with Tukey’s post hoc test was used to compare significant differences in the expression levels of each target gene. Significant differences were indicated by asterisks (*p < 0.05; **p < 0.01; ***p < 0.001; ns, no significant differences).

3. Discussion

Lipid is important for energy homeostasis in insect development, reproduction and hormone synthesis [17]. The synthesis of lipids is an intricate process regulated by multiple enzymes. Among these, FAS is a vital participant mainly contributing to fatty acid synthesis and its functional characterization has been reported in bacteria [18] and fungi [19]. However, the identification and characterization of FAS in insects have still received less attention.

In the present study, we firstly identified two FAS genes from our previously constructed transcriptome database. The number of FAS genes identified here was similar to that present in other insects, such as Colaphellus bowringi (two FAS genes) [20]. The relatively small number suggests the conservative function of FAS. On the other hand, sequence analysis showed that SIFAS1 and SIFAS2 shared only 35.63% amino acid identity...
with each other, and conserved domain prediction revealed that unlike SIFAS2, SIFAS1 lacks a Thioesterase domain (pfam00975), implying the potential functional divergence of these two genes.

To explore the function of the SIFASs, we analysed their expression patterns in S. litura larvae by qRT-PCR. Although SIFAS1 was expressed highest in adults, it was also expressed higher in third instar larvae, suggesting that SIFAS1 still plays a role in S. litura larvae. Meanwhile, SIFAS2 was highly expressed in the third instar larvae of S. litura. Previous studies have reported this similar development stage-biased expression pattern. The expression levels of MpulFAS1 and MpulFAS2 reach a peak at the adult stage but the MpulFAS3 and MpulFAS4 expressions peaks at the larval stage in Meteorus pulchricornis [21]. The divergence of the FAS expression at various different developmental stages indicates that FAS plays distinct functions at different time periods and its role is more complicated. For tissue expression, both SIFAS1 and SIFAS2 were highest in fat bodies. This fat body-biased expression is also reported in other insect species and attributed to the main role of the fat body in lipid metabolism [22]. In Rhodnius prolixus, FASI is predominantly expressed in the fat body [23]. A high transcript abundance of FAS2 has been observed in the fat body of diapaused female Calophellus boweringi [20]. Meanwhile, a higher expression of SIFASs in other tissues such as the epidermis needs to be noted in future studies for revealing the potential role of FASs in insects.

FAS is a central enzyme in de novo lipogenesis, which catalyses the production of C14, C16, and C18 fatty acyl-CoAs, and produces multiple fatty acids [24,25]. Although this catalysis is well summarized in model organisms, there is still a lack of sufficient evidence to support whether FAS undertakes a consistent mission in insects. In the present study, changes in the content of four major fatty acids were examined after silencing SIFAS1, and the GC results showed that the content of methyl α-linolenate was decreased significantly only at 48 h after silencing SIFASI. However, no significant changes in the other three fatty acids were observed. This result strongly suggests that SIFASI is involved in α-linolenic acid synthesis in third instar S. litura larvae. Wang et al. [13] reported that the content of methyl α-linolenate, as well as methyl palmitate, methyl oleate, and methyl linoleate, is decreased significantly after silencing GpylFAS1 in the pupae of the lepidopteran pest Glyphodes pyloalis. Similarly, knocking down Bgfas1 significantly reduces the cuticle free fatty acid content in Blattella germanica [26,27]. Our results not only show that SIFASI solely affects α-linolenic acid, but also indicate that the synthesis of insect fatty acids may be regulated by diverse FAS genes or even other enzymes, and further research is needed to confirm this hypothesis.

The fat body is a crucial biosynthetic and metabolic factory in insects [28], and lipids are mainly stored in the fat body in the form of triglyceride. In the present study, Nile red staining indicated that lipid droplets in the fat body were significantly reduced at 24 and 48 h after dsSIFAS1 injection compared to the dsGFP group. Meanwhile, the triglyceride content was significantly decreased compared to the dsGFP control group at 24 and 48 h. These results reveal a close relationship between SIFASI and lipid synthesis in S. litura larvae, and this correlation has been reported in a limited number of species. For example, the level of triglyceride was severely decreased after knocking down FAS1 in the mosquito Aedes aegypti [11]. In Serratia-infected Acyrthosiphon pisum, the triglyceride content was reduced by feeding with dsFASN1 compared with the control [29]. Therefore, the present study demonstrates that FAS is not only responsible for fatty acid synthesis, but also affects the lipid dynamic in insects.

It is well accepted that lipid accumulation affects insect growth and development [30]. In the present study, our results show that body weight in each developmental time point was not significantly different between the dsSIFASI-injection larvae and dsGFP-injection cohorts. However, the survival rates of the dsSIFASI-injection third S. litura larvae were reduced sharply compared to the dsGFP-injection group. In the model organism C. elegans, silencing FASN-1 at the larval stage ceases normal development and juvenile nematodes are not able to reach adulthood successfully [31]. In a recent study, Yang et al. [32] reported that
the injection of dsLmFAS1 and dsLmFAS3 caused about 80% mortality in Locusta migratoria. Therefore, FAS may be essential for insect development.

For insects, pupation and emergence are key scenarios of metamorphosis [33,34]. Knocking down GpylFAS1 or GpylDesat5, which are essential for metamorphosis in G. pyloalis, leads to more abnormal pupae and adults, and lower emergence rates [13]. Analogously, our results showed that silencing SIFAS1 can induce many more abnormal pupae and adults, and the emergence rates were decreased significantly. Similarly, knocking down LdHAMT, which is an essential metabolism-related enzyme, causes pupation failure and lower emergence rates in the potato beetle Leptinotarsa decemlineata [35]. Furthermore, it is well demonstrated that the hormones in insect larvae significantly shape the process of metamorphosis, and this regulation can be easily measured by monitoring the content of hormones or the expression of hormone genes. Silencing juvenile hormone acid methyltransferase (JHAMT) in the beetle Tribolium castaneum reduces juvenile hormone levels, resulting in the deceleration of lipid catabolism [36]. In the mosquito A. aegypti, 20-hydroxyecdysone (20E) can induce the expression of the transcription factor hepatocyte nuclear factor 4 (HNF4), which accelerates β-oxidation and produces the substances and energy required for oocyte development [37]. Interestingly, our results showed that the content of juvenile hormone in the haemolymph of fifth instar S. litura was not significantly changed after SIFAS1 injection, while the content of ecdysone was significantly decreased at 24 h, suggesting a positive linkage between SIFAS and the ecdysone. Tan et al. [20] reported that the transcription of FAS2 is suppressed by the juvenile hormone and the juvenile hormone receptor methoprene-tolerant. It can be speculated that FAS can regulate the ecdysone metabolism and eventually affect insect metamorphosis.

To further explore the effect of silencing SIFAS1 on other genes in the lipid synthesis pathway, we detected alterations in the expression of SIFAS2, SISdesat, SIACC, and SILIPase by qRT-PCR. Our results suggest that the expression of these genes changes to varying degrees after knocking down SIFAS1, indicating that these genes may be connected with SIFAS1 to jointly regulate the synthesis of lipids in the body. Yang et al. [38] revealed that the glucose-6-phosphate (G6P) metabolism of the host D. melanogaster parasitized by Pachycrepoides vindemmiae is affected by PvG6PDH; the expression levels of the host G6P-metabolism-related genes changed at different time points after the host was injected with PvG6PDH, suggesting the expression of nutrition–related genes can be affected by other genes. In a recent study, Wang et al. [13] also reported that silencing GpylFAS1 affected the expressions of the other fatty acid synthesis-related genes in G. pyloalis. Taken together, it can be concluded that lipid metabolism-related genes can be regulated by FAS and future studies are required to reveal the detailed molecular mechanism of this regulation.

4. Materials and Methods

4.1. Insects

The S. litura larvae were collected from the mulberry fields at the campus of Jiangsu University of Science and Technology, Zhenjiang City, Jiangsu Province, China, and reared in the insectary [26 ± 2 °C, 60–80% relative humidity, and photoperiod of 14:10 (L:D) h]. The adult moths were fed with 10% (w/w) honey solution and provided with strips of paper as the substrate for egg deposition in organza-covered cages (23 cm × 23 cm × 21 cm). After hatching, the newly hatched S. litura larvae were fed with artificial diets daily under the same conditions.

4.2. Bioinformatics Analysis of FAS Genes in S. litura

The sequences of FASs were identified from the previously constructed S. litura transcriptome database (BioProject Acc. in NCBI: PRJNA810583). An open reading frame (ORF) finder (https://www.ncbi.nlm.nih.gov/orffinder/) (accessed on 21 February 2021) was used to predict the ORFs of putative SIFAS1 and SIFAS2 genes. The NCBI Conserved Domain Search website (https://www.ncbi.nlm.nih.gov/Structure/cdd/wrpsb.cgi) (accessed on 21 February 2021) was used to predict the protein functional domains. The Simple
Modular Architecture Research Tool (SMART) online software (Version: 3.3.2, EMBL, Heidelberg, Germany) (http://smart.embl-heidelberg.de/) (accessed on 21 February 2021) was used to predict the conserved motifs. DNAMAN 8.0 (Lynnon Corporation, Quebec City, QC, Canada) was used for multiple alignments of various protein sequences. Phylogenetic analysis was conducted using the neighbour-joining method by Molecular Evolutionary Genetic Analysis 6.0 (MEGA 6.0, Mega Limited, Auckland, New Zealand) software with 1000 bootstrap replications. The Interactive Tree Of Life (iTOL) online tool (https://itol.embl.de) (accessed on 21 February 2021) was used to generate the circular phylogenetic tree. SIFAs homologous amino acid sequences of *Homo sapiens*, *Spodoptera litura*, *Bombyx mori*, *Drosophila melanogaster*, *Spodoptera frugiperda*, *Helicoverpa zea*, *Trichoplusia ni*, *Ostrinia furnacalis*, *Helicoverpa armigera*, *Plutella xylostella*, and *Agrotis ipsilon* were downloaded from GenBank (http://www.ncbi.nlm.nih.gov/) (accessed on 21 February 2021).

4.3. RNA Isolation and Quantitative RT-PCR Validation

Total RNA from the whole body or different tissues (head, midgut, haemolymph, fat body, epidermis) was extracted using a TRIzol reagent (Invitrogen, Carlsbad, CA, USA) according to the manufacturer’s recommendations. The concentration of RNA samples was evaluated according to the absorbance at 260 nm and the purity of RNA was confirmed by the OD260/280 ratio using a 2100 Bioanalyzer (Agilent Technologies, California, CA, USA). Then, the RNA integrity was verified by using 1% agarose gel electrophoresis. Subsequently, 1 µg of total RNA was used to synthesize the first-strand cDNA with a PrimeScript® RT reagent kit (Takara, Dalian, China) following the manufacturer’s instructions. The total qRT-PCR reaction volume was 20 µL, containing 10 µL of 2 × iQ™SYBR® Green I, 1 µL of 10 µM primer of each of the forward and reverse primers, 2 µL of cDNA template, and 6 µL of RNA-free enzyme water. The qRT-PCR validation was conducted using a Roche LightCycler 96 (Roche, Switzerland) and the programme was as follows: 95 °C for 5 min and 40 cycles of 95 °C for 15 s and 60 °C for 31 s. Each sample was repeated three technical times. Meanwhile, the no-template controls (NTCs) of each primer were negative with non-detection of Cq value. The *elongation factor-1 alpha (EF1)* (GenBank accession: DQ192234.1) and *glyceraldehyde-3-phosphate dehydrogenase (GAPDH)* genes (GenBank accession: MZ393966.1) were used as the reference genes to normalize the mRNA expression levels. LightCycler® 96 software (Roche, Switzerland) was used to analyse the qRT-PCR results. The relative expression levels were calculated using the 2−△△Ct method [39]. The primers were designed using the Primer-BLAST online programmer (https://www.ncbi.nlm.nih.gov/tools/primer-blast/) (accessed on 21 February 2021) (Table S1). Each sample was run in triplicate for technical repeats, and three biological replicates were performed simultaneously.

4.4. RNA Interference in *S. litura* Larvae

The oligonucleotide sequences were designed using BLOCK-iTTM RNAi Designer (https://maidesigner.thermofisher.com/) (accessed on 21 February 2021) (Table S2). *dsSIFAS1* and *dsSIFAS2* were synthesized in vitro using a Transcription T7 kit (Taktableara Biotechnology Co. Ltd., Dalian, China) following the manufacturer’s protocol, and the dsRNA of the green fluorescent protein (GFP) gene was synthesized as the negative control. A NanoDrop 2000 spectrophotometer was used to detect the concentration and purity of synthesized dsRNA, and 1% agarose gel electrophoresis was used to examine the RNA integrity.

*dsSIFAS1* and *dsSIFAS2* (500 ng) were injected into the third abdominal segment of 1-day-old third instar *S. litura* larvae using a microsyringe (Drummond Scientific, Broomall, PA, USA), respectively. The samples were collected 24 and 48 h after *dsGFP-* and *dsSIFAS-* were injected, and then RNA samples were extracted and the cDNA was synthesized to verify interference efficiency using qRT-PCR. These procedures were the same as described above.
4.5. Determination of Fatty Acid Content

The fatty acid content was determined according to the method described by Wang et al. [13] with some minor modifications. In brief, third instar *S. litura* larvae were collected 24 and 48 h after dsRNA injection, and dried at 65 °C for fatty acid determination. Each individual was weighed, and transferred to a 10 mL centrifuge tube, and then mashed with a plastic pestle to which 2 mL of n-hexane was added. The homogenate was centrifuged at 10,000 rpm for 10 min, and then 500 µL of supernatant was transferred to a new 10 mL centrifuge tube, adding 1.5 mL of n-hexane and 2 mL of 0.5 mol L\(^{-1}\) KOH-CH\(_3\)OH solution. The mixture was reacted at 60 °C for 1 h. After the mixture was cooled to room temperature, 4 mL of ultrapure water was added, and the resulting mixture was centrifuged at 10,000 rpm for 10 min. The supernatant was removed to 2 mL Eppendorf (EP) tubes for gas chromatography (GC) analysis [40]. The content of methyl α-linolenate, methyl palmitate, methyl oleate, and methyl linoleate was detected using an Agilent 6820 gas chromatograph (Agilent, Santa Clara, CA, USA) equipped with a Supelco capillary column (HP-INNOWax, 30 m × 0.25 mm, i.d. = 0.20 µm; Agilent), a split injection port, and a flame ionization detector. The initial oven temperature was 200 °C, held for 1 min, then increased to 240 °C at 1.5 °C min\(^{-1}\) and held for 1 min. The detector was set at 280 °C and the injector was set at 250 °C. Nitrogen was used as the carrier gas at a flow rate of 1 mL min\(^{-1}\). The split ratio was 50:1 and the sample size was 1 µL [40].

The content of four fatty acids was calculated according to the standard curves for methyl palmitate, methyl oleate, methyl linoleic, and methyl linoleate, respectively. The content of fatty acids was measured based on content per capita (total content divided by individual weight). For each treatment, 15–20 individuals were determined.

4.6. Measurements of Triglyceride Content

The triglyceride content of the fat body was measured using a triglyceride ELISA kit (BYabscience, China) 24 and 48 h after dsRNA injection, following the manufacturer’s instructions. Briefly, the fat body of the larvae was collected and then homogenized in 1 × phosphate-buffered saline (PBS, PH = 7.4). The samples were centrifuged for 10 min at 12,000 rpm at 4 °C, and the supernatants were removed to new 1.5 mL EP tubes for detection. Subsequently, the supernatant was transferred into a 96-well plate, incubated with the reaction mix at room temperature in the dark for 30 min, and measured using a spectrophotometer (Thermo 1500, Waltham, MA, USA). The experiments were repeated three times.

4.7. Nile Red Staining

The lipid of fat bodies in third instar larvae was observed by staining with Nile red [30]. The fat bodies of *S. litura* larvae were dissected and washed with 1 × PBS (pH = 7.4) three times, each washing procedure lasting 5 min; subsequently, the fat bodies were fixed with 4% paraformaldehyde on a glass slide for 30 min at 4 °C. After fixation, the samples were washed with 1 × PBST (1 × PBS containing 0.1% Triton and 0.05% Tween20) three times for 5 min each. For lipid staining, the samples were incubated for 30 min in a 1:1000 dilution of 1 mg ml\(^{-1}\) Nile red solution (Aladdin, Shanghai, China) in 1 × PBST, and then rinsed three times with 1 × PBS (pH = 7.4). The samples were visualized using a fluorescent microscope (LSM 710, Carl Zeiss, Germany) at Ex 543/Em 626 nm.

4.8. Development of *S. litura*

To investigate the effect of SIFASs on the growth and development of *S. litura* larvae, we measured the weight and survival rate of third instar *S. litura* larvae and the emergence rate of fifth instar *S. litura* larvae after silencing the SIFAS, respectively. One-day-old third instar *S. litura* larvae were injected with *dsSIFAS* or *dsGFP* and weighed daily until pupation. A total of 30 third instar *S. litura* larvae were tested. To test the effects of silencing SIFAS on the metamorphosis of *S. litura*, the morphology and the emergence of the pupae were
observed after injecting dsSlFAS1 into fifth instar S. litura larvae. Each treatment was tested on 30 individuals and the dsGFP-injection groups were taken as the control.

4.9. Content of Juvenile Hormone and Ecdysone in Haemolymph of Fifth Instar S. litura after Silencing SlFAS

To determine the effects of silencing SIFAS on the dynamics of the juvenile hormone and ecdysone in the haemolymph of fifth instar S. litura larvae, the content of the juvenile hormone and ecdysone in larvae haemolymph was measured using an insect juvenile hormone and ecdysone ELISA kit (Rui Fanbioon, Shanghai, China) according to the manufacturer’s protocols. In brief, the haemolymph was collected from the fifth instar S. litura at 24 and 48 h after dsSlFAS1 injection and centrifuged at 1000 × g for 5 min at 4 °C. The supernatant was transferred into a new centrifuge tube. Subsequently, 10 µL of supernatant was added to a 96-well plate and incubated with the reaction mixture for 15 min at 37 °C. The optical density value was read at 450 nm using a spectrophotometer (Thermo1500, Waltham, MA, USA). Each sample contained three biological replicates.

4.10. Effect of Silencing SlFAS1 on the Expression of Other Key Genes in the Lipid Synthesis Pathway

To explore the effects of SIFAS1 on other genes in the lipid synthesis pathway in the third instar S. litura larvae, the expression levels of acetyl-CoA carboxylase (ACC), Desaturase (Desat), and Lipase, which were key genes in the lipid synthesis pathway, were validated 24 and 48 h after dsSlFAS1 or dsGFP injection by qRT-PCR; the procedures were the same as mentioned above. The primers of each target gene are listed in Table S1.

4.11. Statistical Analysis

Mann-Whitney tests were used to analyse the significant differences in fatty acid content; data analysis was performed using R software 3.4.0 (R Development Core Team, Vienna, Austria) [41]. One-way analysis of variance (ANOVA) followed by Tukey’s post-hoc test was used to compare the differences in the relative expression levels and the triglyceride content, as well as the content of juvenile hormone and ecdysone.

5. Conclusions

In the present study, we studied the function of FAS genes in a major pest, S. litura, and found that silencing SIFAS1 of the third instar S. litura larvae can depress the synthesis of fatty acids and the content of triglycerides in the fat body and, furthermore, affect development. Meanwhile, we also demonstrated that SIFAS1 is necessary for the metamorphosis of S. litura, and ecdysone alteration may be involved in this process. This study provides a new insight into lipid synthesis in insects, especially in insect pests, and would provide novel targets in pest management.

Supplementary Materials: The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/ijms23169064/s1.

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