Abstract

Lepidopteran insect pests are the main class of pests causing significant damage to crop plant yields. Insecticidal scorpion peptides exhibit toxicity specific for insects. Here, we report that a peptide LMX, optimized from the insect-specific scorpion neurotoxin \( \text{LqhIT2} \), showed high levels of activity against rice leaf folder \( \text{in vitro} \) and \( \text{in planta} \). Oral ingestion of LMX protein led to a significant decrease in feeding on rice leaves, repression of larval growth and development, delay in molting, and increase in larval lethality. Compared with \( \text{LqhIT2} \) protein, the stability and insecticidal efficacy of LMX was better. Meanwhile, biochemical analysis showed that LMX protein ingestion dramatically decreased ecdysone content in rice leaf folder larvae, and down-regulated enzymatic activities of the detoxification system (\( \text{Naphthyl acetate esterase and glutathione S-transferase} \)), the digestive system (trypsin and chymotrypsin), and the antioxidant system (catalase). These changes were tightly correlated with the dosage of LMX protein. Transgene analysis showed that the rate of leaf damage, and the number of damaged tillers and leaves in the transgenic line were greatly reduced relative to wild type plants and empty vector plants. Based on these observations, we propose that the insect-specific scorpion neurotoxin peptide LMX is an attractive and effective alternative molecule for the protection of rice from rice leaf folder.

Introduction

Rice (\( Oryza sativa \)) acts as a staple food for much of the global population [1], particularly in Asia where about 90% of people live on rice [2]. Thus, rice plays a critical role in safeguarding the food security of the world. However, in commercial production, rice productivity and quality are adversely affected by many biotic stresses, particularly insect pests.

Rice leaf folder (\( \text{Cnaphalocrocis medinalis} \) Guenee) is a migratory insect which causes significant damage to rice yields [3]. Rice leaf folder larvae damage rice crops by folding leaf blades and feeding inside the rolled leaves. A single larva can damage a number of rice leaves, disturbing photosynthesis and reducing the rice yield [4]. Rice leaf folder is widely distributed around the rice-growing areas of the world. Since the middle of the 1960s, rice leaf folder has become one of the most severe paddy field pests and outbreaks of rice leaf folder have been reported in many Asian countries [5]. In China, since 2003, the annual average area damaged by rice leaf folder was more than 20 million \( \text{hm}^2 \) and the grain yield loss was up to 760 million kg every year. In 2011, rice leaf folder damaged 22 million \( \text{hm}^2 \) of rice in China [5].

To avoid rice yield loss caused by leaf folder infestation, conventional means rely on the extensive use of chemical pesticides. However, chemical pest control is expensive, environmentally unfriendly and pest-resurgence often occurs [6]. With wide use of modern biotechnology in rice breeding programs, development of pest-resistant plants through the introduction of foreign insect-resistance genes offers a potentially desirable and effective way to accomplish rice pest control [7]. To date, the most widely used pest-resistant gene is \( \text{Cry} \), isolated from \( \text{Bacillus thuringiensis} (\text{Bt}) \), which has shown good performance in public and private sector rice breeding programs [8]. However, long-term use of \( \text{Bt} \) genes will increase the risk of narrowing the insect-resistant spectrum of transgenic plants [9]. Thus, to meet this challenge, there is an urgent need to explore economically and ecologically sound alternatives so as to enrich genetic diversity.

Scorpion venom contains a variety of polypeptides with diverse biological activities [10]. According to their targets, scorpion venom polypeptides are divided into three categories: mammal neurotoxins, crustacean neurotoxins and insecticidal toxins [11]. The insecticidal toxins are further divided into two groups based on molecular size and activity. One is the short group with 30–40 amino acid residues and 3–4 disulphide bridges, which mainly affect conductance of potassium channels [12]. The other is the long group with 60–70 amino acid residues cross-linked by 4 disulphide bridges, which principally affect sodium channels in excitable cells [13]. Recently, a series of scorpion insecticidal peptides including \( \text{LqhIT2} \) [14,15], \( \text{AaHIT} \) [16,17], \( \text{ButaIT} \) [18,19], \( \text{ButaIT} \) [20] and \( \text{AaHIT} \) [21] have been isolated and tested for enhancement of protection against lepidopteran insect pests. It is notable that these scorpion toxins are neuromuscular system-specific in insects and are safe for other animals [22–25]. The specificity to insect pests offers significant potential for the development of safe insecticides with a broad spectrum.

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* Email: shaoqingli@whu.edu.cn
none of these have yet been employed as an insecticide defense against rice leaf folder by introducing the genes into rice.

LqhIT2, an insect-specific scorpion toxin, is purified from the venom of *Leiurus quinquestriatus hebraeus* [26]. It is confirmed that having the effective toxic to lepidopteran insect pests [14,15], however, the insecticide efficacy still have some distance to meet our expectation in the practical use. Here, we designed an optimized peptide named LMX which has more than 85% amino acid and 92% nucleotide identity with LqhIT2 (Figure S1). After optimizing the nucleotide sequence of *LMX* based on rice codon preference, we introduced the gene into rice to improve resistance to rice leaf folder. Our results demonstrated that the *in vitro* expressed LMX fusion protein had high biological activity and toxicity against rice leaf folder, and the toxicity was better than that of LqhIT2 fusion protein. The LMX transgenic rice lines showed high resistance to rice leaf folder. LMX is therefore revealed as a good alternative tool for the further improvement of insect-pest resistance in rice.

**Materials and Methods**

**Ethics Statement**

The experimental field in the town of Huashan, Wuhan city, did not require any specific permission for use in this study. This study did not involve endangered or protected species. The specific location of the experimental field was as follows: 114°47′~114°60′ E; 30°51′~30°59′ N.

**Plant materials**

The Indica rice variety Yuetai (*Oryza sativa* L.), which was supplied by our laboratory, was used in this research. Rice was planted in the experimental field in the town of Huashan, Wuhan city, during the summer season from 2010 to 2012.

**Insect culture**

The third instar larvae of the rice leaf folder were collected from the experimental field in Huashan town, Wuhan city. Rice leaf folder larvae were reared in controlled chambers at 27±1°C and 75±3% relative humidity under a photoperiod of 16-h-light/8-h-dark using fresh leaves of Yuetai [27]. The larvae were fed individually from day 1 of 3rd instar.

**Construction of the expression vector and the binary vector**

The *LMX*-encoding nucleotide sequence was synthesized (Genscript, China) based on plant codon usage bias [28] and cloned into the vector pUC57 (Genscript, China). To construct a plasmid based on plant codon usage bias [28] and cloned E. coli, the complete coding region of the *LMX* gene as a marker. A strain of *Agrobacterium tumefaciens* EHA105 was used for the transformation experiment.

For large-scale preparation of active LMX protein, the soluble cell extracts of 3 L. IPTG-induced culture were collected and purified from clarified bacterial lysate by affinity chromatography on GSTrap FF 1 ml columns (Amersham Biosciences, USA), and then centrifuged at 14,000 g for 30 min at 4°C. The soluble cell extracts and cell pellets were confirmed by 12% SDS-PAGE and western blot.

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**Analysis of protein characters and homologous modeling**

The basic physic-chemical characters of LMX and LqhIT2 were performed by ProtParam online tools in the ExPASy database (Physico-chemical parameters of a protein sequence, http://expasy.org/tools/protparam.html). The homologous modeling of LMX was performed by SWISS-MODEL which used the three-dimensional structure of LqhIT2 as a template.

**Analyses of circular dichroism (CD) spectroscopy**

The CD spectra for LMX and LqhIT2 were carried out by CD spectropolarimeter (Jasco, J-810 (Japan). The acquisition parameters were 0.1 cm optical path, 50 nm/min scan rate, 25°C temperature, 0.1 nm resolution, 2.0 nm bandwidth, 0.5 nm data
pitch, 1 s response and 250~190 nm wavelength range. All treatments were performed in triplicate.

**Analysis of the biological toxicity of the LMX protein**

To investigate toxin effects of LMX protein on rice leaf folders, the leaf disc method [29] was used as a biological toxicity assay. Freshly collected rice leaves from the Yuetai were cut into 1 cm × 10 cm pieces weighing about 0.2 g, placed in petri dishes padded with humid filter paper and pressed with absorbent cotton on the opposite ends of the leaf blade. For the biological toxicity assay, supernatants containing 2.5 µg, 5 µg, 7.5 µg and 10 µg LMX fusion protein were dried under reduced pressure and the dried residues were dissolved in 100 µl PBS buffer. The different dosages of LMX fusion protein were individually daubed on one piece of the leaves. Larva on day 1 of 3rd instar were released and reared individually with one piece of leaf fragment placed in dishes. The dishes were placed in controlled chambers at 27 ± 1°C and 75 ± 5% relative humidity under a photoperiod of 16-h-light/8-h-dark. 100 µl PBS buffer and 10 µg purified GST protein dissolved in 100 µl PBS buffer were used as controls. After 72 h, we recorded the average consumption rates of leaves, the average weight of surviving larvae, and the larvae lethality. Thirty larvae and thirty leaves were used in each treatment group. Three repeats were performed for each treatment, and calculations were done based on the following formula:

\[
\text{Consumed rate} (\%) = \frac{\text{The area of leaf by larva feeding}}{\text{The total area of leaf}} \times 100
\]

\[
\text{The average weight of surviving larva (mg)} = \frac{\text{Total weight of surviving larva}}{\text{Number of surviving larvae}}
\]

\[
\text{Larvae lethality (\%)} = \frac{\text{Number of dead larvae}}{\text{Number of total larvae}} \times 100
\]

**Measurement of enzyme activities of rice leaf folder**

Larvae on day 1 of 3rd instar were released and reared individually with one piece of leaf fragment, which were each applied with a different protein dosage of 2.5, 5 or 7.5 µg of LMX fusion protein. After 72 h, ten larvae as a group were homogenized with PBS buffer, and centrifuged for 30 min at 4,000 × g. The suspension of every group of larvae was collected as a crude enzyme extract for measurement. The measurement of the activities of α-naphthyl acetate esterase (α-NAE), glutathione S-transferase, superoxide dismutase (SOD), catalase (CAT), tryphtase and chymotrypsin were performed with insect enzyme activity assay kits (Nanjing Jiancheng Bioengineering Institute, China) according to the manufacturer's instructions. PBS buffer and 7.5 µg GST protein were used as controls. Thirty larvae were used in each treatment group. Three repeats were performed in each treatment.

**Measurement of rice leaf folder ecdysteroids**

Briefly, larvae on day 1 of 3rd instar were individually fed with one piece of leaf fragment coated with different protein dosages of 2.5, 5 or 7.5 µg LMX fusion protein. After 72 h, ten larvae as a group were weighed and homogenized in 2 ml methanol. Ecdysteroids were extracted according to the method described by Hagele et al. [30]. The ecdysteroid measurement was carried out using High Performance Liquid Chromatography (HPLC) (More Biotechnology, China) based on the following program: Twenty microliters of the solution was injected into a LC-MS instrument (Waters e2695/2495, USA), separations were performed on a Waters Symmetry C18 column (4.6 × 250 mm i.d., 5 µm, Waters, USA) at a flow rate of 0.8 ml/min at 30°C, eluting with acetonitrile, methanol and water in a ratio of 1:2:4. α-ecdysteroid and β-ecdysteroid were used as standards. Thirty larvae were used in each treatment. All treatments were performed in triplicate. PBS buffer and 7.5 µg GST protein were used as controls.

**Genetic transformation and molecular identification of LMX transfectants**

Transformation of Yuetai plant by A. tumefaciens was performed following the method described by Peng et al. [31]. The T0 generation plants were first screened for LMX positive lines using specific PCR primer pairs: LMX-F1 (5-GCTCTAGAATG-GACGGCTACATCCGCAAG-3) and NOS-R1 (5-GGAATTCTGGGTATACCGCAATATATCC-3). PCR was performed based on the following program: 94°C for 10 min followed by 35 cycles of 94°C, 1 min; 58°C, 1 min; 72°C, 1 min; and a final extension of 10 min at 72°C. PCR products were separated by 1% (w/v) agarose gel electrophoresis. The PCR positive transgenic lines were further confirmed by Southern blot. Briefly, 10 µg of genomic DNA per sample was digested overnight with Hind III and separated on a 0.8% agarose gel, then transferred to a nylon membrane and probed with LMX cDNA labeled with biotin-11-dUTP, using the NorthSouth Biotin Random Prime Labeling Kit (Thermo, USA). The blots were hybridized with the labeled probe overnight at 55°C and washed at 55°C for 15 min in 2×SSC and 0.1% SDS three times. The membrane was visualized using the Chemiluminescent Nucleic Acid Detection Module (Thermo, USA).

**Quantitative real time PCR**

For quantitative real time PCR analysis, total RNA was extracted from approximately 100 mg fresh rice leaves from control and positive transgenic rice plants using the Trizol reagent (Invitrogen) according to the manufacturer’s instructions. First-strand cDNA was synthesized using oligo(dT)20 primers with the ReverTra Ace transcriptase kit (Toyobo, Japan) according to the manufacturer’s instructions. Quantitative real time PCR was performed in a 20 µl volume using SYBR Green Master Mix (Roche, Switzerland) and data was analyzed on the iCycler iQ5 Real Time PCR Detection System (Bio-Rad, USA) according to the manufacturer’s instructions. The cDNA was diluted 10-fold and amplified for quantitative real time PCR analysis with the specific primers LMX-F1 (5-ATGGACGGCTACATCCGCAAG-3) and LMX-R1 (5-TTAGCGGTGTTTGGTCTCT-3). Quantitative real time PCR reactions containing 1 µl of cDNA were carried out by the following the program: 1 cycle at 94°C for 5 min followed by 35 cycles each consisting of 10 s at 94°C, 10 s at 58°C and 10 s at 72°C. Each sample was run in triplicate. The endogenous ubiquitin gene was used as the internal control with specific primers (UBI-F1: 5-GTTTCGGCCGTTGGAGTTGATTGC-3) [32].
Detection of the LMX protein in transgenic rice
The homologous T$_2$ transgenic plants were used for Western blotting. Approximately 20 mg of protein extracted from fresh rice leaves was loaded in each lane, separated on a 15% Tricine-SDS-PAGE gel, and transferred onto a PVDF membrane (Amersham). Membranes were blocked with TBST buffer (20 mM Tris-HCl pH 8.0, 150 mM NaCl, 0.05% Tween 20) containing 5% fat-free milk, and incubated 1 h at 37°C. The blot was then incubated with LMX rabbit anti-serum diluted 1:2000 overnight at 4°C, washed three times with TBST at room temperature for 15 min each time. Next the blot was incubated with a secondary antibody goat anti-rabbit IgG (Thermo, USA). The homologous T$_2$ transgenic plants were used for Western blotting. Approximately 20 mg of protein extracted from fresh rice leaves was loaded in each lane, separated on a 15% Tricine-SDS-PAGE gel, and transferred onto a PVDF membrane (Amersham). Membranes were blocked with TBST buffer (20 mM Tris-HCl pH 8.0, 150 mM NaCl, 0.05% Tween 20) containing 5% fat-free milk, and incubated 1 h at 37°C. The blot was then incubated with LMX rabbit anti-serum diluted 1:2000 overnight at 4°C, washed three times with TBST at room temperature for 15 min each time. Next the blot was incubated with a secondary antibody goat anti-rabbit IgG (Thermo, USA).

Evaluation of transgenic rice insect-resistance
To investigate the resistance of transgenic plants expressing LMX to rice leaf folder, we used a leaf disc method to analyze the transgenic plants. Freshly collected rice leaves from the 2-month old homozygous transgenic rice plants expressing LMX were used. The leaves of transgenic plants were cut into fragments about 10 cm$^2$, and placed in petri dishes padded with humid filter paper. Rice leaf folder larvae on day 1 of 3rd instar were released individually with one piece of leaf fragment placed in each dish for 72 h. The plates were kept at a temperature of 27±1°C and relative humidity of 75±5% under a photoperiod of 16-h-light/8-h-dark. The leaves of wild-type plants and transgenic plants expressing the empty vector pCAMBIA1301-rbcS-Nos (rbcS plants) were used as the controls. Ten larvae and ten leaves were placed in each dish, three dishes were used in each treatment, and all treatments were performed in triplicate. The consumption rate of leaves was counted 72 h after release of the insects. The consumption rate was calculated using the following formula:

$$\text{Consumed rate (}) \% = \frac{\text{The area of leaf by larva feeding}}{\text{The total area of leaf}} \times 100$$

For whole plant analysis of pest-resistance, each homozygous transgenic rice plant was placed in a plastic bucket and infested with five rice leaf folder larvae on day 1 of 3rd instar. Then, the plastic buckets were covered with a net. The severity of tillers and leaves damage to the whole plant was scored on day 7 post-infestation. The wild-type plants and the plants transformed with the empty vector (rbcS plants) were used as controls. Twenty-five plants were used. Twenty-five larvae and five plants were used in each treatment and each treatment was repeated three times. Further, we also tested resistance of the whole transgenic plant to rice leaf folder under natural conditions. Each plot consisted of 30 plants grown in three rows. No insecticides were applied during the entire growing season. The damage rate to tillers and leaves of the whole plants were calculated using the following formula:

$$\text{Damage rate of tillers (}) \% = \frac{\text{Number of tillers by larva feeding}}{\text{Number of tillers of the whole plant}} \times 100$$

$$\text{Damage rate of leaves (}) \% = \frac{\text{Number of leaves by larva feeding}}{\text{Number of leaves of the whole plant}} \times 100$$

Statistical analysis
All experimental data are represented as the mean over three independent replicates. Statistical analysis was done using GraphPad Prism 5.0 software. The values shown in the table and figures represent means ± SD of triplicate. Statistical significance was determined as $P \leq 0.05$.

Results
Molecular optimization of LMX polypeptide
In order to enhance the insecticide efficacy of LqhIT2, we changed the amino acid in the non-conservative regions of LqhIT2 to construct the optimized peptide named LMX. The result of amino acid alignment showed that there were nine amino acid sequences differences between LMX and LqhIT2 (Figure S1). The nine amino acid was follow: the lysine, arginine, aspartic acid, valine, alanine, aspartic acid, alanine, tyrosine and glycine at the 5, 6, 8, 12, 13, 22, 27, 28 and 30 were replaced by arginine, lysine, asparagine, isoleucine, serine, asparagine, glycine, phenylalanine and alanine, respectively (Figure S1). The optimized LMX shared has more than 85% amino acid and 92% nucleotide identity with LqhIT2 (Figure S1).

Expression and purification of scorpion polypeptide LMX
To investigate the activity of LMX against rice leaf folder, a GST-LMX recombinant construct was produced using the codon-optimized LMX gene (Figure 1A), and expressed in E. coli. Coomassie staining revealed that an additional band about 32.8 kDa (LMX 6.8 kDa plus GST 26 kDa) was observed both in soluble and insoluble cell fractions of GST-LMX recombinant E. coli, and the GST-LMX fusion protein mainly existed in the soluble cell fraction (Figure 1B). After affinity purification, two bands were visible on a 12% polyacrylamide gel, the biggest one being about 32.8 kDa was GST-LMX fusion protein, and the smaller one around 26 kDa was GST tag protein (Figure 1B). This was further confirmed by Western blot using an antibody specific to the LMX protein (Figure 1C).

Structural analysis of the polypeptide LMX
In order to know the structure of LMX, we analyzed the physico-chemical parameters of LMX protein by using ProtParam online tools in the ExPASy database. The result showed that the instability index and hydropathicity of LMX protein was 30.29 and −0.621 (Table S1). Meanwhile, the instability index and hydropathicity of LqhIT2 protein was 41.89 and −0.651, respectively (Table S1), which means that the hydrophobicity and stability of LMX protein was better than the LqhIT2 protein. Furthermore, we analyzed the secondary structure of LMX and LqhIT2 protein by CD spectra, and the result showed that the spectral curves of LMX and LqhIT2 were highly similar (Figure 2A). Additionally, the homologous modeling showed the three-dimensional structures of LMX and LqhIT2 were also highly similar (Figure 2B). All of these indicated that although there were some differences in physico-chemical parameters between LMX and LqhIT2, the core spatial structure of LMX had no significant change in comparison with LqhIT2 protein.
LMX protein showed high effectively than LqhIT2 in decreases feeding on rice leaves and increase lethality of the rice leaf folder

Some reports have shown that oral ingestion of exogenous neuropeptides can lead to effective toxicity against pests [19,33]. To investigate the toxicity of the LMX fusion protein against rice leaf folder by oral ingestion, 2.5~10 μg dose gradients of LMX fusion protein were applied to Yuetai rice leaves. Compared to a 91.9% damage rate for PBS buffer treated leaves and an 86.6% damage rate for the 10 μg GST tag protein control, LMX fusion protein treatments led to significant reductions in damage to rice leaves on the 3rd day post-infestation. The damage rates of leaves in the 2.5 μg and 5 μg LMX samples were 45.0% and 20.5%, respectively; and almost no damage was observed on the leaves with 7.5 μg and 10 μg of protein (Figure 3A-B). The damage rate was negatively correlated with the dosage of LMX fusion protein, and the correlation coefficient reached -0.957 (Figure 3B).

Furthermore, we also detected the toxicity of the LqhIT2 fusion protein against rice leaf folder by oral ingestion. The damage rates of leaves in the 2.5, 10 μg dose gradients of LqhIT2 protein were 54.0%, 34.5%, 15.2% and 8.6%, respectively. The comparison result of LMX and LqhIT2 fusion protein showed that the damage rates of leaves in 2.5~10 μg dose gradients of LqhIT2 samples were both higher than those in LMX samples (Figure S2A).

Meanwhile, we investigated the effects of LMX fusion protein on the survival and development of the leaf folder. Results showed that the lethality rates of insects fed on the rice leaves applied with LMX protein were much higher than those of the controls on the 3rd day post-infestation (Figure 3C). The lethality rates of larvae fed with 2.5 μg, 5 μg and 7.5 μg LMX fusion proteins were 12.2%, 45.5% and 68.9%, respectively. When the dosage reached 10 μg, insect mortality was as high as 94.4%. Insect lethality had a strong positive correlation with the dosage of LMX fusion protein, and the correlation coefficient was 0.982. There was no significant difference in insect lethality of controls (larvae ingesting PBS buffer or 10 μg GST tag protein). Control lethality rates were 7.8% and 8.9%, respectively (Figure 3C). Whereas, the lethality rates of larvae fed with 2.5 μg, 5 μg, 7.5 μg and 10 μg LqhIT2 fusion proteins were 11.1%, 35.6%, 58.9% and 78.9%, respectively. Compared with the insect lethality of 2.5~10 μg dose gradients of LMX protein, we also found that the lethality rates of gradients of LqhIT2 protein were much lower (Figure S2B). The results of the damage rates of leaves and the insect lethality rates suggested that the insecticidal efficacy of optimized peptide LMX was better than that of LqhIT2.

LMX protein represses growth and disrupts the equilibrium of ecdysone in larvae of rice leaf folder

Further, we investigated the growth status of the surviving larvae treated with LMX fusion protein. Compared with the controls, average weight of the surviving rice leaf folder larvae in the LMX fusion protein treatment groups was much lower on both the 3rd day and the 5th day post-infestation (Figure 4). As shown in Table 1, the average weight of the surviving larvae ingesting 2.5 μg, 5 μg and 7.5 μg LMX increased 6.1 mg, 3.8 mg and 2.4 mg after 3 days of treatment, respectively. However, the average weight of the larvae in the PBS and GST protein control groups increased 7.4 mg and 6.5 mg, respectively. At the 5th day of treatment, the larval weight in the GST tag protein control group increased 10.4 mg, but the LMX (2.5 μg, 5 μg and 7.5 μg) treatment groups only increased 8.3 mg, 4.7 mg and 2.7 mg, respectively. This reflects a strong inhibition of LMX fusion protein on the growth and development of rice leaf folder.
It is well known that growth and development of the larvae of lepidopteran insects are tightly related to molting. Prohibition of the growth of rice leaf folder fed with LMX fusion protein suggests that molting characteristics of the larvae may also change. Interestingly, compared to the normal molting and pupation of the surviving larvae in controls, we observed that molting of the surviving larvae both in the 5\(\mu\)g and 7.5\(\mu\)g LMX fusion protein treatment groups was retarded (Figure 5). This indicates that LMX fusion protein could postpone and impede molting and development in the rice leaf folder.

\(\alpha\)-ecdysone and \(\beta\)-ecdysone are two major insect hormones which play a critical role in controlling insect growth and development [34]. Hence, we investigated the ecdysone content of larvae in the control and treatment groups fed with LMX fusion protein after 72 h of ingestion. Results showed \(\alpha\)-ecdysone in the 3rd instar larvae fed with 2.5 \(\mu\)g, 5 \(\mu\)g and 7.5 \(\mu\)g LMX decreased 15.8\%, 35.2\% and 42.8\% respectively, compared to the 7.5 \(\mu\)g GST protein control. Meanwhile, \(\beta\)-ecdysone in larvae of the treatment groups was reduced 5.0\%, 31.4\% and 50.3\% relative to the 7.5 \(\mu\)g GST tag protein control (Figure 5). The changes in \(\alpha\)- and \(\beta\)-ecdysone content were both negatively correlated with the dosage of LMX fusion protein, and the correlation coefficients were \(-0.989\) and \(-0.968\), respectively. This reflects that oral ingestion of LMX fusion protein disturbs the metabolism of \(\alpha\)-ecdysone and \(\beta\)-ecdysone in larvae of rice leaf folder.

**LMX protein disturbs digestive, detoxification and antioxidant systems in rice leaf folder larvae**

In insects, tryptase and chymotrypsin, \(\alpha\)-naphthyl acetate esterase (\(\alpha\)-NAE) and glutathione S-transferase (GST), superoxide dismutase (SOD) and catalase (CAT) are the key enzymes of the digestive, detoxification and antioxidant systems, respectively. The activities of these enzymes can directly reflect the physiological state of the rice leaf folder. We measured the enzyme activities in larvae of rice leaf folder from the 1st to the 3rd day after infestation. Similar trends were observed for each of the enzymes during the three days (Table S2 and Table 2). According to these trends, we focused on the enzyme activities in larvae of rice leaf folder from the 1st to the 3rd day after infestation. After 72 h of treatment, the activities of \(\alpha\)-NAE, GST, CAT, POD, tryptase and chymotrypsin in the larvae fed with LMX fusion protein (2.5 \(\mu\)g, 5 \(\mu\)g and 7.5 \(\mu\)g) were reduced. Enzyme activity of \(\alpha\)-NAE was decreased 17.5\%–25.42\%, glutathione S-transferase decreased 65.66\%–76.42\%, CAT decreased 28.71\%–58.63\%, tryptase decreased 28.54\%–40.35\%, and chymo-
trypsin decreased 43.34–63.89% compared to the GST tag protein control (Table 2). In contrast, the activity of SOD was activated, and the increase was 139–149.66% compared to the GST protein control (Table 2).

Expression of the LMX gene improves rice resistance to rice leaf folder

In order to evaluate whether LMX can effectively function at the plant level, a rice genetic transformation experiment was designed. Figure 4. Phenotype of the rice leaf folder larvae fed with LMX protein. The larvae fed on 2.5 – 7.5 μg LMX protein-applied rice leaves were treatment groups, while the larvae fed on PBS buffer-applied rice leaves and 7.5 μg GST tag protein-applied rice leaves were control groups. Pictures were taken at the start (0 days), 3 days, and 5 days.

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Table 1. Weight of surviving larvae of rice leaf folder on the 3rd and 5th day post-infestation (mg/larvae).

| Culture duration | PBS       | GST (7.5 μg) | LMX (2.5 μg) | LMX (5.0 μg) | LMX (7.5 μg) |
|------------------|-----------|--------------|--------------|--------------|--------------|
| 0d               | 21.1±0.31 | 21.4±0.42    | 21.0±0.38**  | 21.6±0.25**  | 21.3±0.32**  |
| 3d               | 28.5±1.20 | 27.9±0.60    | 27.1±0.86**  | 25.4±0.89*   | 23.7±0.87**  |
| 5d               | 32.4±0.45 | 31.8±0.55    | 29.3±0.44*** | 26.3±0.60*** | 24.0±0.21*** |

Weight of surviving larvae was measured after 72 h feeding on 2.5 μg–7.5 μg LMX protein-applied rice leaves. The larvae fed on PBS buffer-applied rice leaves and 7.5 μg GST tag protein-applied rice leaves were used as control groups. Values are presented as means of triplicates ±SD. * denotes p<0.05; ** denotes p<0.01; *** denotes p<0.001. ns denotes no significant difference.

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performed. In total, 73 T<sub>0</sub> independent transgenic plants were generated, of which 63 were identified as positive by PCR amplification (Figure 6B). Furthermore, Southern blot analysis showed that three transgenic plants (LMX-5, LMX-10 and LMX-22) harbored a single copy of the insertion (Figure 6C). Western blot and qPCR analysis indicated successful expression of the LMX gene under the <i>rbcS</i> promoter in these three transgenic rice lines, with the LMX-10 plant having the highest expression level (Figure 6D-E).

Further, we studied the insect resistance of LMX transgenic rice plants using wild type and empty vector transgenic plants (rbcS plants) as controls. Larvae on day 1 of the 3<sup>rd</sup> instar of rice leaf folder were individually fed on one piece of transgenic rice leaf. This showed that the average damage rate to the LMX transgenic rice leaf was about 58.5% after 72 h, whereas the wild type and the rbcS plants were both more than 90% damaged (Figure 7). Meanwhile, we found that LMX transgenic rice plants had significantly less damage than the wild type and rbcS plants after a 7 d infestation in an artificial infestation setting (Figure 7 and Table 3). Under natural field conditions, the average damage rates to plant tillers and leaves of wild type plants over two years (2011, 2012) were about 86.1% and 68%. The damaged tillers and leaves of transgenic lines decreased 30.6%~38.3% and 19.1%~36.9% relative to those of the wild type group, respectively (Table 3). However, no significant differences were observed in the damage rates of tillers and leaves between wild type and rbcS plants (Table 3). These data reveal that LMX can efficiently improve the resistance of rice plants against the rice leaf folder.

### Discussion

LMX has oral activity against rice leaf folder

It has been proved that scorpion toxin LqhIT2 has high preference for insect voltage-gated sodium channels. In the present work, we designed an optimized protein LMX. The CD spectra and homologous modeling analysis showed that substitution amino acids did not affect the structure, and the core spatial structures of LMX and LqhIT2 were highly similar (Figure 2). It is consistent with previous study that most substitutions in non-conservative regions of LqhIT2 could not affect the CD signature [34]. It was indicated that LMX and LqhIT2 has the similar or same model of action and LMX also affects the insect voltage-gated sodium channels.

### Table 2. Enzyme activities in larvae feeding on LMX fusion protein on the 3<sup>rd</sup> day post-infestation.

| Enzymes | PBS | GST (7.5 μg) | LMX (2.5 μg) | LMX (5.0 μg) | LMX (7.5 μg) |
|---------|-----|-------------|-------------|-------------|-------------|
| α-NAE (U/L) | 79.59±8.13 | 81.88±4.39 | 67.56±6.35** | 65.23±5.74* | 61.07±3.78* |
| GST (U/mg.prot) | 452.42±25.25 | 445.31±11.47 | 152.91±8.11*** | 114.75±12.79*** | 104.99±3.71*** |
| SOD (U/mg.prot) | 13.98±0.35 | 14.54±0.57 | 34.75±0.31*** | 33.63±0.43*** | 36.30±0.62*** |
| CAT (U/mg.prot) | 43.95±0.96 | 44.02±0.35 | 31.38±0.42*** | 21.56±0.24*** | 18.21±0.28*** |
| TPS (IU/ml) | 20.58±2.46 | 21.09±1.05 | 15.07±1.98** | 14.79±1.05** | 12.58±1.27*** |
| CTP (U/L) | 36.99±1.22 | 37.61±1.63 | 21.31±1.13*** | 17.3±0.75*** | 13.58±1.91*** |

For the control, the third larvae were fed with PBS buffer and 7.5 μg GST tag protein. α-NAE: α-naphthyl acetate esterase, GST: glutathione S-transferase, SOD: superoxide dismutase, CAT: catalase, TPS: trypstatase, CTP: chymotrypsin. Values are means ± SD of triplicates. * denotes p<0.05; ** denotes p<0.01; *** denotes p<0.001. ns denotes no significant difference.

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It has been suggested that scorpion insecticidal neurotoxins function by affecting conductance of ion channels by direct injection [35,36]. However, very few studies have investigated the oral activity of scorpion peptides. In the present work, oral ingestion of LqhIT2 protein could affect larval feeding and kill larval (Figure S2), and oral ingestion of LMX protein was also shown to decrease larval feeding on leaves, retard larval development and increase larvae lethality (Figure 3–5; Table 1). The insect-feeding trials clearly demonstrated that bacteria-expressed LMX protein was toxic to rice leaf folder larvae, and the effect of insect-resistance was positively correlated with the dosage of LMX protein. Furthermore, the comparison results demonstrated that the hydrophobicity, stability and toxicity of optimized LMX is better than that of LqhIT2 protein (Figure 3 and Figure S2, Table S1). The amino acid Alanine of LqhIT2 which is at 13 site has a unique role in toxin function of LqhIT2 [35]. It could explain that why the toxicity of LMX is better than LqhIT2. It also may be relative with substitution of alanine at 13 site. Ingestion of the LMX protein into the insect gut system is different from the mechanism of direct injection. Though we know little about the exact mechanism of LMX toxicity to the larvae by oral ingestion, we propose two possibilities that may partly explain the toxicity of LMX: (1) The stability of LMX makes LMX peptide have high-levels of resistance to protease. It is likely to be hyper-stable and has a long residence time in the gut of leaf folder larvae. Based on online prediction by the DiANNA 1.1 web server (https://www.predictprotein.org/) LMX has 4 disulfide bonds; C_{10}-C_{60}, C_{14}-C_{35}, C_{21}-C_{42} and C_{25}-C_{44} (Figure S1C). Disulfide bonds can enforce protein structural stability and resistance to proteases [37]. Hence, absorption of a minority of ingested protein may exert some level of oral activity. This is consistent with the observation that the spider peptide H-HXTX-Hvla is orally active against lepidopteran pests when expressed in cotton, poplaf and tobacco plants [38]. (2) The stable LMX protein impedes the physiological and chemical metabolism of leaf folder larvae, which results in disrupted enzyme activities of digestive, antioxidant and detoxification systems and the disequilibrium of ecdysones involved in the process of insect molting, causing the observed delay in larva development (Table 2 and Figure 5). This further inhibits the growth and development of insect larva, and therefore decreases feeding and damage to rice leaves.

**LMX is a valuable insecticidal candidate for rice improvement**

In the last few decades, some valuable resistance genes have been successfully identified. Combined with transgenic techniques, this will make it possible to breed environmentally friendly rice by improving rice resistance to insect pests through bioengineering techniques. This engineered rice will potentially reduce the use of...
pesticide in commercial rice production [39,40]. *Bacillus thuringiensis* (*Bt*) genes are the most prominent examples used for rice genetic improvement in resistance to insects, especially lepidoptera [41]. However, widespread use of a single exogenous *Bt* insecticidal protein will lead to the adaptation of insect pests to this protein. In fact, a number of *BT*-resistant insect strains have been observed under laboratory, greenhouse, and even field conditions [42–44]. This reflects the urgency in exploring new useful insecticidal genes to arm crop plants against challenges from insect pests. Recently, some insect toxin genes, such as proteases [45], protease inhibitors [46–48], and the spider venom toxin gene *Hvt* [49] etc. have been tested for the improvement of plant resistance to insect pests, but the number of available genes conferring effective pest resistance remains limited.

It has been suggested that ectopic expression of insect-selective neurotoxin peptides would be highly advantageous in coping with lepidopteran insect larvae, which attack over 60 crop plants ranging from monocotyledon to dicotyledon [50]. As discussed above, many insecticidal scorpion peptides have been suggested as a promising alternative for insect pest control [17,18,21,51]. Of these, LqhIT2 has been demonstrated to confer resistance to lepidoptera insects [14] without toxicity to mammals [23,24]. Its efficacy in transgenic rice against leaf folder has also been confirmed in our laboratory (data in preparation for subsequent publication). In our study, LMX shares more than 85% amino acid homology with LqhIT2 (Figure S1). Both nurse chamber feeding and natural field investigations confirmed that LMX can effectively improve rice plant resistance to rice leaf folder (Figure 7; Table 3). Therefore, the LMX gene is critical for improving resistance to rice leaf folder.

After oral ingestion, we observed the toxicity of LMX protein to larvae to be positively correlated with protein dosage. 10 μg of LMX protein almost completely repressed larvae feeding on rice leaves. Practically, LMX content in transgenic rice leaves will be far lower than this level. As larvae living on LMX transgenic rice can implement the transit from larvae to pupa, this implies that to some extent these larvae are able to endure the toxicity of LMX on the *in planta* level. If we use only LMX as a resistance gene in rice improvement, it will lead to the leaf folder’s swift adaptation to this toxin [15]. Thus, comprehensive strategies such as combining scorpion peptide LMX with other insecticidal genes like spider venom peptide [49] and plant insecticidal proteins [52], are necessary to prevent the occurrence of pest resistance. In the interest of increasing the security of transgenic plants, as well as the safety of mammals, we propose that insecticidal scorpion peptide LMX may play a useful role in the genetic modification of rice for insect pest resistance.
**Table 3.** Damage analysis of LMX transgenic rice caused by rice leaf folders from 2011 to 2012.

| Years | Treatments     | Genotypes | Damage rate of tillers (%) | Damage rate of leaves (%) |
|-------|----------------|-----------|---------------------------|--------------------------|
| 2011  | Wild type      | 95.0±2.35 | 95.2±1.12                 |
|       | rbcS           | 95.3±1.76 ** | 95.5±1.36 **              |
|       | Artificial infestation | LMX-5 | 71.2±1.59 *** | 62.8±1.61 ***          |
|       | LMX-10         | 65.6±3.16 *** | 56.6±3.20 ***          |
|       | LMX-22         | 78.6±2.0 *** | 74.2±1.58 ***          |
|       | Wild type      | 95.9±4.40 | 86.5±3.10                 |
|       | rbcS           | 95.6±4.70 ** | 86.2±3.70 **             |
|       | Natural infestation | LMX-5 | 54.0±3.68 *** | 59.7±4.46 ***          |
|       | LMX-10         | 45.8±6.50 *** | 50.7±4.00 ***          |
|       | LMX-22         | 62.9±2.51 *** | 67.5±1.99 ***          |

| 2012  | Wild type      | 93.0±0.93 | 94.0±2.08                 |
|       | rbcS           | 93.8±2.28 ** | 93.0±1.49 **             |
|       | Artificial infestation | LMX-5 | 73.1±5.45 *** | 66.5±2.59 ***          |
|       | LMX-10         | 64.3±2.33 *** | 54.5±2.68 ***          |
|       | LMX-22         | 80.3±4.01 *** | 76.5±2.46 ***          |
|       | Wild type      | 76.3±11.80 | 49.5±10.30               |
|       | rbcS           | 79.0±7.50 ** | 50.5±7.30 **            |
|       | Natural infestation | LMX-5 | 34.5±1.66 *** | 21.7±2.44 ***          |
|       | LMX-10         | 29.8±5.70 *** | 11.5±2.60 ***          |
|       | LMX-22         | 48.2±4.65 *** | 30.3±1.13 ***          |

Values are means ± SD of ten different plants. *** denotes P < 0.001.

Conclusions

The results of *in vitro* and *in planta* experiments provide evidence that insect-specific scorpion neurotoxin peptide LMX has high levels of activity against rice leaf folder. LMX represses larval growth and molting, affects enzyme activities, and increases larval lethality *in vitro*. In addition, transgenic rice plants expressing LMX demonstrate enhanced resistance to rice leaf folders. Therefore LMX could be considered as a candidate gene to improve plant resistance to lepidopteran insects.

Supporting Information

**Figure S1** LMX disulfide bonds and sequence alignment. A. Alignment of the amino acid sequences of the mature polypeptide LMX and the LqhIT2 polypeptide. The gray frames indicate the same amino acid for the LMX and LqhIT2 polypeptides in non-conservative region. The purple frames highlight the conservative region between LMX and LqhIT2 polypeptides. B. Alignment of the full-length LMX and LqhIT2 nucleic acid sequence according to plant codon usage. The gray frames indicate the same nucleic acid sequence, shared between LMX and LqhIT2. C. Disulfide bonds of the mature polypeptide LMX.

(TIF)

**Figure S2** The LqhIT2 protein confers resistance to the rice leaf folder. A. Ratio of the damaged leaf area to the whole leaf applied with 2.5 µg−10 µg bacterial-expressed LqhIT2 protein. B. Larvae lethality of the rice leaf folder fed with 2.5 µg−10 µg bacterial-expressed LqhIT2 protein. For the control, the rice leaves were daubed with 1 × PBS buffer and 10 µg GST tag protein. Pictures were taken and ratios of the damaged leaf area were measured after 72 h incubation with the rice leaf folder. Values are means ± SD. *** denotes 0.001 significant difference.

(TIF)

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Author Contributions

Conceived and designed the experiments: SQL YGZ XZTP. Performed the experiments: XZTP. Analyzed the data: XZTP. Contributed reagents/materials/analysis tools: XZTP. Wrote the paper: XZTP SQL.
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