Genome-wide characterization and expression profiling of immune genes in the diamondback moth, *Plutella xylostella* (L.)

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The diamondback moth, *Plutella xylostella* (L.), is a destructive pest that attacks cruciferous crops worldwide. Immune responses are important for interactions between insects and pathogens and information on these underpins the development of strategies for biocontrol-based pest management. Little, however, is known about immune genes and their regulation patterns in *P. xylostella*. A total of 149 immune-related genes in 20 gene families were identified through comparison of *P. xylostella* genome with the genomes of other insects. Complete and conserved Toll, IMD and JAK-STAT signaling pathways were found in *P. xylostella*. Genes involved in pathogen recognition were expanded and more diversified than genes associated with intracellular signal transduction. Gene expression profiles showed that the IMD pathway may regulate expression of antimicrobial peptide (AMP) genes in the midgut, and be related to an observed down-regulation of AMPs in experimental lines of insecticide-resistant *P. xylostella*. A bacterial feeding study demonstrated that *P. xylostella* could activate different AMPs in response to bacterial infection. This study has established a framework of comprehensive expression profiles that highlight cues for immune regulation in a major pest. Our work provides a foundation for further studies on the functions of *P. xylostella* immune genes and mechanisms of innate immunity.

The innate immune system, which consists of cellular and humoral responses, is the first line of defense against bacterial and fungal infections in insects¹. The humoral immune response includes the production of antimicrobial peptides (AMPs) that protect against a broad array of infectious agents, such as bacteria, fungi, viruses and even eukaryotic parasites²–⁵. In insects, the genetic mechanisms involved in these defence reactions have been most thoroughly studied in *Drosophila melanogaster*. In this model organism, genes encoding AMPs are activated by NF-κB transcription factors in response to infection through the Toll and immune deficiency (IMD) signaling pathways. Other immune pathways in insects are the JNK and JAK-STAT pathways, which participate in cell stress or wound response⁴–⁶. The JAK-STAT pathway also participates in antiviral response⁷. Different pattern recognition receptors (PRRs), such as

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peptidoglycan recognition proteins (PGRPs) and beta-1, 3-glucan recognition proteins (βGRPs), detect microbial infection and trigger the signaling cascades to activate the Toll and IMD pathways.

The innate immune system is known to be highly conserved in the animal kingdom and evolutionarily very stable. It is, however, suggested that as insects are exposed to a multiplicity of continuously evolving pathogens, genes involved in signaling pathways may be under selection, leading to diversification. Casanova-Torres and Goodrich-Blair compared the genetic mechanisms related to the immune system of D. melanogaster (Order: Diptera) and Bombyx mori and Manduca sexta, two representatives of Order: Lepidoptera. They showed, for example, that while some genetic components of the Toll signaling pathway may be similar between Lepidoptera and Diptera, some other genes differ. Gaps remain in some of these pathways and underline the need for further studies to better understand the differences between the Toll and IMD signaling pathways in these two insect orders. With greater availability of genetic tools and published genome sequences in Lepidoptera, it is possible to acquire a better understanding of the genetics of the immune system in other insect species such as P. xylostella, a notorious pest attacking many economically important food and oil crops in Brassicaceae.

Recent studies of P. xylostella immunity have mainly focused on individual immune genes and aspects such as production and characterization of cecropin10, the influence of entomopathogenic nematodes (Xenorhabdus nematophila), and immune suppression by the parasitoid Cotesia plutellae polydnaviral gene11–13. Additionally, expressed sequence tags and cDNA microarrays have been used to analyze immune-inducible genes14. Whilst these studies of specific aspects are enlightening, there is a more fundamental need to develop an understanding of immune related genes based on genome-wide analysis, especially since the mechanisms of P. xylostella immunity at the molecular level are poorly understood.

Understanding the genetics and regulatory mechanisms of immune systems of insect pests, such as P. xylostella, has emerged as an important field to facilitate the development of effective biological control tactics. Comparative analyses of immune genes at the genome wide level in insects is the first step to define the candidate genes and functional networks associated with immune regulation. Thus, the aim of this study was to identify the immune signaling pathways, along with the pathogen recognition proteins and immune effectors, from the recently sequenced P. xylostella genome15. A total of 112 Serine proteases (SPs) and 102 serine protease homologs (SPHs) (9 SPs and 2 SPHs contain clip domains), as well as 26 serine protease inhibitors (Serpin) were also identified in the P. xylostella genome (data not shown), the detailed information on these components will be addressed in other paper. The expression patterns of immune genes that identified in this study were examined in P. xylostella at various development life stages, in different tissues and in susceptible and insecticide-resistant lines. Finally, P. xylostella immune response to bacterial pathogens was studied.

Results

Classification of P. xylostella immune genes. A total of 149 immune genes were identified in the P. xylostella genome, and classified into 20 families and 4 functional groups: immune recognition, signaling pathways, immune effectors, and others. The immune recognition genes included PGRPs, βGRPs, galectins, C-type lectins, fibrinogen-related proteins (FREPs), and scavenger receptors (SCRs). The signaling pathways genes were further classified into the pathway-related categories of TOLL, IMD, JNK and JAK-STAT. The immune effectors comprised prophenoloxidase (PPO), thioester-containing proteins (TEPs), and AMPs. Others included the enzyme superoxide dismutase (SOD), catalases and peroxidases that participate in detoxification of reactive oxygen species (ROS) (Table S1). The total number of P. xylostella immune genes was less than that of the other four insect species, D. melanogaster, Anopheles gambiae, B. mori, and Tribolium castaneum. The βGRPs were strikingly expanded, while AMPs were relatively reduced in P. xylostella.

Immune recognition families. Peptidoglycan recognition proteins (PGRPs). In this study, we identified 9 PGRP genes in P. xylostella, fewer than the 13 for D. melanogaster, 11 in A. gambiae, and 12 in B. mori (Table S1). Phylogenetic analysis showed that the 9 P. xylostella PGRPs were distributed in different branches and were homologous to Danaus plexippus and B. mori PGRPs, suggesting conservation of this protein family in Lepidoptera (Fig. 1). The 9 P. xylostella PGRPs were located in four different scaffolds (Table S2). Previous studies have shown that the Toll pathway is initiated by PGRP-SA 16,17, thus the two PGRP-SAs (Px015207 and Px015209) in the same scaffold may function in the Toll pathway for the surveillance of Gram-positive bacteria. The IMD pathway, a conserved pathway that mainly defends against poly-dnaviral gene, is triggered through PGRP-LC 18,19. Additionally, 4 βGRPs in A. gambiae, and 4 βGRPs in B. mori, we found 18 βGRP genes in P. xylostella, indicating a dramatic
expansion of this gene family in *P. xylostella* (Table S1). The 18 βGRPs of *P. xylostella* were clustered into three clades in the phylogenetic tree (Fig. 2), suggesting that the functions of this protein family might be diverse. Multiple sequence alignments and domain architecture analysis indicated that the conserved domains of glycosyl-hydrolase family 16 were present in *P. xylostella* βGRPs except for Px009706, which might have been an incomplete sequence or a pseudogene (Fig. S2). Previous studies indicated that the N-terminal regions of GNBPs/βGRPs also participate in carbohydrate recognition. Our results showed that the N-terminal regions of *P. xylostella* βGRPs were more diverse than the C-terminal regions (Fig. S2), suggesting their importance in functional divergence. The 18 βGRPs were located in 7 scaffolds. The sequences with higher identities that looked like duplications in the phylogenetic tree, such as Px008677 (GRP7) and Px015064 (GRP16), Px001058 (GRP1) and Px009702 (GRP11), were not located in the same scaffolds (Table S2-3). The genes located in the same scaffold were those that were more
diverse and distributed in different branches of the phylogenetic tree (Fig. 2), suggesting that there was no tandem duplication of βGRP genes in *P. xylostella* genome.

**Galectins.** Galectins are a family of lectins that contain evolutionary conserved carbohydrate-recognition domain (CRD) specifically for β-galactoside sugar. Four galectins were identified in the *P. xylostella* genome (Table S2), and phylogenetic analysis showed that the *P. xylostella* galectins are homologous to those in other Lepidoptera insects such as *Danaus plexippus* and *B. mori* (Fig. S3). The domain architecture analysis (Fig. S4) suggests that all the *P. xylostella* putative galectins possess conserved CRD domains. Previous studies indicated that galectins function in innate immunity of *D. melanogaster* and *A. gambiae*, such as in microbial recognition and phagocytosis. Therefore, the complete CRD domains in the sequences suggest a possible immune function in *P. xylostella*.

**Fibrinogen-related proteins (FREPs).** FREPs are a family of proteins containing fibrinogen domains in the C-terminal region and they function in recognizing bacteria and parasites in invertebrates. Two fibrinogen-related proteins were identified in the *P. xylostella* genome (Table S2). This family is large in *A. gambiae* with 61 genes in its genome, moderate in *D. melanogaster* with 14 genes and in *T. castaneum* with 7 genes, but quite small in the Lepidoptera (3 genes in *B. mori* and 2 genes in *P. xylostella* (Table S1). Phylogenetic analysis showed that the *P. xylostella* FREPs were most homologous to those of *B. mori*, and the genes in *D. melanogaster*, *A. gambiae*, and *T. castaneum* were more species-specific (Fig. S5). The domain architecture analysis (Fig. S6) indicated that all the putative *P. xylostella* FREPs possess conserved fibrinogen-related domains at the C-terminus, suggesting their possible full functions in the *P. xylostella* immunity.

**C-type lectins.** C-type lectins contain a wide variety of soluble and membrane-bound proteins with calcium-dependent carbohydrate-recognition domains (CRD). Previous studies argue that invertebrate C-type lectins play an important role in immune responses, such as activating PPO cascade, participating in hemocyte nodule formation, and recognizing microorganisms to enhance microbial clearance. Seven C-type lectins were identified in the *P. xylostella* genome, which is much fewer than in *D. melanogaster*, *A. gambiae* and *B. mori* (Table S1). This might result from the incomplete information of the current *P. xylostella* genome. It is also possible that this is a species-specific characteristic of *P. xylostella* since the βGRP family is expanded in *P. xylostella*. Phylogenetic analysis showed that the *P. xylostella* C-type lectins were homologous to those of *B. mori* and *D. plexippus*, and that there existed common 1:1 orthologs among the six species, indicating that the C-type lectins are conserved in insects (Fig. S7). The domain architecture analysis (Fig. S8) suggests that the putative *P. xylostella* C-type lectins possess one or two conserved carbohydrate-recognition domains which might function for the sugar-binding activity.

**Scavenger receptors.** The family of scavenger receptors was documented to contain multidomains and function as pattern recognition receptors in innate immunity. This family can be divided into three subfamilies, scavenger receptors A (SCRAs), scavenger receptors B (SCRBs) and scavenger receptors C (SCRCs), based on their functional domains. The SCRAs have been implicated in the host defense by binding polyanionic ligands such as lipoteichoic acid (LTA) or lipopolysaccharide (LPS). The scavenger receptor Cysteine-Rich (SRCR) domain is usually located in some members of this subfamily, and contributes to binding to Gram-positive and Gram-negative bacteria. One SRCR was identified in the *P. xylostella* genome (Table S2), and a SRCR domain is located at the middle of the sequence (Fig. S9). Phylogenetic analysis indicated the 1:1 ortholog between *P. xylostella* and *B. mori* (Fig. S10). SCRMs are thought to be a novel class of scavenger receptors characterized by a CD36 domain, they participated in the phagocytosis of microbes and binding to apoptotic cells. Thirteen SCRCs were identified in the *P. xylostella* genome (Table S2), and they all contain a CD36 domain with one or two transmembrane domain(s) (Fig. S9). Phylogenetic analysis suggests that this subfamily in *P. xylostella* was more homologous to that in *D. plexippus* than in *B. mori* (Fig. S10). The SCRCs were previously identified to function as PRRs in phagocytosis and innate immunity. This subfamily contains multidomains including two complement-control protein (CCP), one Meprin A5 antigen and RPTP Mu (MAM), and one somatomedin-B-like (BO). One SCRC was identified in the *P. xylostella* genome (Table S2) with 1 MAM, 1 BO, and a transmembrane domain in the sequence (Fig. S9). Phylogenetic analysis showed that this subfamily in *P. xylostella* was homologous to that in *D. plexippus* (Fig. S10).

**Immune signaling pathways.** The Toll pathway. Spätzle (SPZ) is a ligand of the Toll receptor, and forms a complex with Toll receptor resulting in activation of the Toll signaling pathway. There are six SPZ genes in *D. melanogaster* and *A. gambiae*. In *P. xylostella* genome, five SPZ genes were identified (Table S1). Common 1:1 orthologs or paralogs of SPZ genes were present in *P. xylostella*, *D. plexippus* and *B. mori* (Fig. S11), suggesting the conservative nature of this gene family in lepidopteran insects. Nine Toll receptor genes were identified in *P. xylostella* genome, with a gene duplication of Toll9. Phylogenetic analysis revealed gene duplication of Toll receptors in each of the five insect species (Fig. S12). The domain architecture analysis showed that six Toll receptors contain the functional domain of leucine rich repeats (LRRs), transmembrane domain (TM) and Toll/interleukin-1 receptor (IL-1R) homologous region (TIR) (Fig. S13), indicating that they may function in signal transduction. When Toll receptors
are activated, they can bind to cytoplasmic MyD88 and, as a result, MyD88, tube and pelle form a complex to phosphorylate cactus, leading to degradation of cactus and release of Dorsal/Dif, which then translocate to the nucleus to activate antimicrobial peptide genes\(^\text{36,39}\). The Toll pathway genes of tube, pelle, cactus, TRAF6 and dorsal/dif but MyD88 were identified in \(P.\) \(xylostella\). TRAF6 protein functions as an adapter in the shrimp \(Litopenaeus\) \(vannamei\) (Decapoda: Penaeidae) to regulate AMP gene expression\(^\text{36}\). However, function of \(P.\) \(xylostella\) TRAF6 needs to be further validated because \(D.\) \(melanogaster\) TRAF6 is not implicated in immune signaling\(^\text{36,39}\). The absence of MyD88 gene in \(P.\) \(xylostella\) might be due either to the homologue search method based on sequence similarity or incomplete genome information. It is also possible that the function of MyD88 is substituted by other adaptors. Further experiments would be needed to identify the cause.

### The IMD pathway

The IMD pathway is a conserved pathway that is mainly activated by Gram-negative bacteria\(^\text{16,36}\). Genes in the IMD pathway, including IMD, Dredd, FADD, TAK1, TAB2, IKK-\(\beta\) and IKK-\(\gamma\), were all identified in \(P.\) \(xylostella\) genome. Interestingly, TAK1 is not only a downstream component of IMD, but also a protein kinase that triggers the JNK pathway to activate gene expression in response to cell stress or wound\(^\text{28,35}\). JNK, C-Jun and Kay genes in the JNK pathway were all identified in \(P.\) \(xylostella\) genome. The common 1:1 orthologs of genes in the IMD and JNK pathways of \(P.\) \(xylostella\) and other insects indicated that these pathways were complete and conserved (Table S1-2). Three inhibitors of apoptosis 2 (IAP2) were also identified in \(P.\) \(xylostella\) genome (Table S1-2). IAP2 might be involved in Relish nuclear localization as evident in \(Drosophila\)\(^\text{39}\).

### The JAK-STAT pathway

In \(Drosophila\), only one cytokine receptor (Domeless), one Janus kinase (Hopscotch), and one transcription activator (STAT) have been identified in the JAK-STAT pathway\(^\text{26,40}\). However, only one Domeless, one Hopscotch and two STATs homologues were identified in \(P.\) \(xylostella\) genome. In addition, only one TEP, which is under the control of the JAK-STAT pathway\(^4\), was identified in \(P.\) \(xylostella\) genome compared to other insect species (Table S1).

### Immune effectors

**Antimicrobial peptides (AMPs).** AMPs are evolutionarily conserved proteins/peptides involved in innate immune responses. These peptides have small molecular weights and broad-spectrum of activities against bacteria, fungi and viruses\(^\text{27,41}\). Three AMP gene families, two cecropins, three moricins and two gloverins, were identified in \(P.\) \(xylostella\) genome. The seven AMP genes were located in 3 scaffolds, and AMPs of the same family were located in the same scaffold (Table S2). Cecropins are commonly found in insects, and are active against both Gram-positive and Gram-negative bacteria\(^42\). Phylogenetic analysis showed that \(P.\) \(xylostella\) cecropins were similar to those of \(B.\) \(mori\) and \(D.\) \(plexippus\), and they formed a cluster in the tree (Fig. S14A). Moricin and gloverin have been found only in Lepidoptera thus far. Moricin acts against both Gram-positive and Gram-negative bacteria\(^\text{43}\). Although phylogenetic analysis showed that \(P.\) \(xylostella\) moricins were more homologous with \(B.\) \(mori\) moricins, different branches were formed in the tree and no 1:1 orthologs were found between the two moth species (Fig. S14B). Gloverin is a glycine-rich protein, which was first identified in \(Hyalophora\) pupae and was active against \(E.\) \(coli\)\(^\text{44}\). \(B.\) \(mori\) gloverins also exhibited activity against \(E.\) \(coli\) with rough lipopolysaccharide\(^\text{45}\), but studies of \(M.\) \(sexta\) gloverin revealed that it is active against both Gram-positive and Gram-negative bacteria, as well as fungi\(^\text{46}\). The gloverin family tended to be species-specific in the phylogenetic tree, and \(P.\) \(xylostella\) gloverins appeared to be more ancient in the tree (Fig. S14C). Lysozyme comprises a protein family that defends against bacteria by attacking peptidoglycans in cell walls, especially of Gram-positive bacteria\(^\text{47}\). Two lysozymes were identified in \(P.\) \(xylostella\) genome, and they are homologous to lysozymes in \(D.\) \(plexippus\) and \(B.\) \(mori\).

### Enzymes in reactive oxygen species (ROS) detoxification

Reactive oxygen species contribute to defense against invading microbial pathogens, but over-production can also harm host cells. Thus, ROS production must be tightly controlled. The concentration and conversion of ROS can be regulated by superoxide dismutases (SODs), peroxidases and catalases. SODs convert superoxide radical (\(O_2^-\)) into a less toxic product, hydrogen peroxide (\(H_2O_2\)). \(H_2O_2\) is converted to water and oxygen by catalases. Peroxidase also scavenges \(H_2O_2\) and converts it to hydroperoxide\(^\text{48,50}\). SODs are classified into two sub-families in \(D.\) \(melanogaster\), with one family (Cu-Zn SOD) located in the cytosol and the other (Mn-Fe SOD) in the mitochondria. Seven SOD genes were identified in \(P.\) \(xylostella\) genome compared with 4 in \(D.\) \(melanogaster\), 5 in \(A.\) \(gambiae\) and 6 in \(B.\) \(mori\). Five of the 7 SODs in \(P.\) \(xylostella\) were Cu-Zn SOD, and the other two were Mn-Fe SODs, while three Cu-Zn SOD and one Mn-Fe SOD are present in \(D.\) \(melanogaster\)\(^\text{36}\). Orthologs were common in the SOD family (Fig. S15). Similarly, in the peroxidase family, orthologs were also common as shown in the phylogenetic tree (Fig. S16), but the catalase family was more species-specific (Fig. S17). The peroxidase family was expanded, with 17 peroxidase genes in \(P.\) \(xylostella\), 20 in \(D.\) \(melanogaster\), 26 in \(A.\) \(gambiae\) and 23 in \(B.\) \(mori\). Thirteen catalase genes were identified in \(P.\) \(xylostella\), significantly more than the 2 in \(D.\) \(melanogaster\), 1 in \(A.\) \(gambiae\), and 7 in \(B.\) \(mori\). Previous studies have shown that plants attacked by insects may increase ROS production, resulting in oxidative damage to the insect midgut\(^\text{51}\). SODs, peroxidases and the expansion of catalases in \(P.\) \(xylostella\) genome is likely to reflect their roles in ROS detoxification and the wider co-evolution of this herbivore with cruciferous plants.
Prophenoloxidase (PPO). PPO is an important enzyme for melanization in invertebrates to defend against pathogens and for wound healing. PPO is cleaved by a serine protease cascade and converted to functional active enzyme phenoloxidase (PO). PO catalyzes the conversion of monophenols to quinones, thus contributing to melanin synthesis to defend against pathogens. There are 3 PPO genes in *D. melanogaster*, 9 in *A. gambiae*, and 2 in *B. mori*. We identified only 1 PPO gene in the *P. xylostella* genome, which is highly supported by TblastN and manual NCBI blast annotation, and *P. xylostella* PPO formed a cluster with *B. mori* and *D. plexippus* PPOs in the phylogenetic tree (Fig. S18). The expansion of PPO is not common in insects, and there are only 2 PPOs in *M. sexta* and 3 in *T. castaneum*. We also predicted another 8 PPO-like sequences with high TblastN scores; seven of them were located in the same scaffold. Manual annotation showed that these PPO-like sequences consist of three conserved domains, Hemocyanin_N, Hemocyanin_C and the copper-containing Hemocyanin_M. But NCBI blast results suggested that they were more homologous to hexamerin storage protein, which are hemocyanin-derived proteins with functions in amino acid storage, juvenile hormone (JH) binding and in reproduction and metamorphosis. Functional analysis needs to be performed to identify the actual roles of these proteins in *P. xylostella*.

Expression analysis of immune genes in *P. xylostella*. Stage-specific expressions of immune genes. Expression profiles of immune genes were determined by RNA-seq from multiple life stages to show regulation patterns of different immune genes in *P. xylostella* (Fig. 3). PGRP and most βGRP genes were strongly expressed in pupae. The Toll pathway genes were highly expressed in eggs, 1st-instar larvae, pupae, and adults. The IMD pathway genes were highly expressed in pupae and adults whilst all AMP genes were highly expressed in pupae. The expression profiles of AMP genes at different developmental stages were also validated by qRT-PCR (Fig. S19). Our results indicated high expression levels of Toll and IMD pathways genes in pupae, resulting in high expression of AMPs. The JNK and JAK-STAT pathways genes were also highly expressed in pupae and adults, indicating their possible roles in cell stress or wound response.

Immune gene expression in the midgut and head. Much research has demonstrated that insect guts are related to immunity, such as the activation of host defense in the *Drosophila* gut in response to bacterial infection and the activation of AMPs by the IMD pathway in epithelia in response to infection. Although the head is not generally considered an immune organ in insects, previous work has suggested that the honey bee head exhibits differential expression of proteins, including those that participate in signal transduction, in response to a bacterial challenge. More generally, the head has many chemoreceptors, particularly associated with the mouthparts and antennae; is fundamental to feeding, the center for the nervous system and associated processing of various environmental signals including those associated with mating. Accordingly, the present study focused on the immune protection system in the *P. xylostella* head as well as the midgut to enrich the availability of information on insect immunity. Tissue-specific immune genes were differentially expressed in the heads of 4th-instar larvae, male and female adults, as well as in the midgut of 4th-instar larvae in *P. xylostella*. The results showed that four PGRP genes (Px008495, Px001312, Px008494 and Px004941) were strongly expressed in the 4th-instar larvae, pupae, and adults. The IMD pathway genes were highly expressed in pupae and adults whilst all AMP genes were highly expressed in pupae. The expression profiles of AMP genes at different developmental stages were also validated by qRT-PCR (Fig. S19). Our results indicated high expression levels of Toll and IMD pathways genes in pupae, resulting in high expression of AMPs. The JNK and JAK-STAT pathways genes were also highly expressed in pupae and adults, indicating their possible roles in cell stress or wound response.

Strain-specific expressions of immune genes. AMP genes were all down-regulated in the chlorpyrifos and fipronil insecticide-resistant *P. xylostella* (CRL and FRL) compared to the susceptible strain (SS) (Fig. S21), which were confirmed by qRT-PCR (Fig. S22). Genes involved in the Toll pathway were affected by the insecticide resistance status of the *P. xylostella* strains; up-regulated in CRL but down-regulated in FRL compared to SS. However, the IMD pathway genes were down-regulated in both CRL and FRL, similar to the expression patterns of AMP genes. Thus, we hypothesize that repeated insecticide application may decrease expression of IMD pathway genes, resulting in down-regulation of AMP expression. A similar expression pattern was also found for lysozyme, which was down-regulated in the insecticide-resistant lines. In contrast to these down-regulated genes, those participating in the JNK and JAK-STAT pathways were all up-regulated in the resistant lines compared to the susceptible lines. Although it has been shown that the JNK pathway also regulates AMP production, our finding of up-regulation of JNK pathway genes and down-regulation of AMPs suggest that the JNK pathway may not regulate AMP expression, at least in the insecticide-resistant lines. Previous studies showed that the JNK and JAK-STAT pathways participated in the cell stress or wound response. The expression patterns of the JNK and JAK-STAT...
pathways genes suggest that the two pathways may respond to insecticide stress. Additionally, PPO genes were also up-regulated in the resistant lines, indicating their possible functions in response to insecticides (Fig. S21). Previous work indicated that *P. xylostella* phenoloxidase may play an important role in the increasing resistance to butane-fipronil65.

**Induced expression of AMPs by bacterial challenge.** Results from the bacteria-feeding experiment suggest that *P. xylostella* AMPs could be induced by bacterial infection (Fig. 4). After feeding on the Gram-negative bacterium *Enterobacter* sp., cecropin and moricin expressions were up-regulated significantly. Feeding *Serratia* sp. also significantly up-regulated moricin, but cecropins and moricins were not induced by the Gram-positive bacterium *Enterococcus* sp. Gloverin and lysozyme were up-regulated by *Enterococcus* sp., gloverin was also up-regulated by *Enterobacter* sp. These results indicate that *P. xylostella* may activate the expressions of specific immune effectors such as AMPs to defend against different types of pathogens. In *P. xylostella*, gloverin and lysozyme may mainly function in defense against...
Gram-positive bacteria, while cecropin and moricin may mainly defend against Gram-negative bacterial infection. Although Mackintosh et al.\textsuperscript{66} reported that gloverin isolated from the old world bollworm Helicoverpa armigera is active against Gram-negative bacteria (\textit{Escherichia coli} and \textit{Acinetobacter calcoaceticus}), but not active against Gram-positive bacteria (\textit{Arthrobacter globiformis} and \textit{Bacillus thuringiensis}), another study\textsuperscript{46} has reported that gloverin from \textit{M. sexta} is active against Gram-positive bacteria (\textit{Bacillus cereus}) but almost inactive against Gram-negative bacteria (\textit{E. coli}). These studies suggest that gloverin in different insects may have species-specific functions in defense against pathogens.

**Discussion**

By comparative analysis of immune genes among five insect species, 20 gene families were identified in the \textit{P. xylostella} genome, including components of the conserved immune signaling pathways (Toll, IMD, JNK and JAK-STAT), pathogen recognition and immune effectors. Based on the identified genes (Table S1-2) and their functions documented in other insect species\textsuperscript{35,36,54,67}, we propose a model of the potential immune pathways in \textit{P. xylostella} (Fig. 5) that need to be experimentally validated. The Toll and IMD pathways regulate synthesis of immune responsive effectors such as AMPs. The Toll pathway is also known to participate in \textit{Drosophila} embryonic development\textsuperscript{16,68,69}. Based on our current expression pattern of the Toll pathway genes, we believe that it might also play a similar role in \textit{P. xylostella}. The JNK pathway is for transcriptional activation of defense genes or may participate in the synthesis of AMP, whilst the JAK-STAT pathway is for transcriptional activation of stress response genes like TEPs. As to the pathogen recognition gene families, we observed substantial expansion of βGRPs in \textit{P. xylostella}, but also noted a large number of common 1:1 orthologs among the genes involved in intracellular signal transduction pathways related to immune responses. Previous studies have shown that \textit{D. melanogaster} DmGNBP1 has a high binding affinity to lipopolysaccharide (LPS) and beta-1,3-glucan from bacteria and fungi, and can trigger the Toll pathway\textsuperscript{70}. DmGNBP3 not only triggers the Toll pathway during fungal infection, but also activates the prophenoloxidase cascade\textsuperscript{71}. In \textit{M. sexta} larval plasma, β-1,3-glucanase-related protein may stimulate prophenoloxidase activation\textsuperscript{72,73}. The large expansion of βGRPs in \textit{P. xylostella} genome suggests that βGRPs may be involved in effective defense against Gram-negative bacteria and fungi. These results also suggest that while recognition proteins in the \textit{P. xylostella} immune system may be diverse, the intracellular signal transduction pathways may be more conserved. Previous studies in \textit{D. melanogaster}\textsuperscript{36}, \textit{A. gambiae}\textsuperscript{36}, and \textit{B. mori}\textsuperscript{15} all suggested that the pathogen recognition receptors are more expanded and diverse than the genes participating in the intracellular signal transduction pathways. The
expansion of diverse recognition receptors enables insects to recognize different pathogens and to trigger
immune pathways that respond effectively to pathogens. This study also provides a foundation for future
functional studies of βGRPs, which could be unique to P. xylostella.

In contrast with βGRPs, only seven AMP genes in three families were identified in P. xylostella
genome (Table S1-2). These results are consistent with those of previous studies using molecular cloning techniques10, EST and microarray to analyze immune-inducible genes14, or transcriptome analysis based on deep sequencing74. It remains unclear why there are fewer AMPs in P. xylostella than in other insects, such as D. melanogaster36, A. gambiae36 and T. castaneum54. Because our results were obtained from homologous search, we might have missed some AMP genes with sequences that are likely divergent from typical AMPs. It is also possible that reduction in common AMPs can be compensated by other unknown P. xylostella-specific AMP genes.

Gene expression profiles showed that most immune genes were highly expressed at the pupal stage
and in the head tissue. Previous work showed that the bacterial pathogen Campylobacter jejuni could be
transferred between life stages of Musca domestica (larva-pupa), but the number of C. jejuni declined
during pupal development, coinciding with the increased expression of AMPs, indicating effective innate
immunity at the pupal stage75. Thus, high expression of immune genes in the pupal stage of P. xylostella
may be key to defense against pathogens and cell stress in this important developmental stage. On the
other hand, since the head is the center for nervous system and feeding in insects with many important
sense organs to perceive various environmental signals63, it might require a higher level of immune sur-
veillance for adequate protection.

All the P. xylostella immune genes studied here were identified based on bioinformatics analysis.
Their in vivo functions still need to be validated by molecular studies, such as gain-of-function or
loss-of-function analysis. Nevertheless, through this study, we have generated information about the
genetic composition and regulation of the immune system of an important agricultural pest. Our work
will help drive future studies focusing on identifying molecular functions and mechanisms of immune
genes and pathways of P. xylostella and other arthropods.

Methods

Identification and classification of immune genes in P. xylostella. The sequences of immune
genes in D. melanogaster and Anopheles gambiae were downloaded from the immunology database at
http://cegg.unige.ch/Insecta/immunodb. The Bombyx mori and Tribolium castaneum immune genes
were edited in previous studies35,54, and were downloaded as queries. These immune-related sequences
were used to search for immune genes in P. xylostella genome database (DBM-DB, http://iae.fafu.edu.
cn/DBM/)76. A local TblastN search with an E-value of 10^-6 was performed to collect putative immune
genes from *P. xylostella* genome, and the default E-value of 1.0 was used to search for antimicrobial peptides (AMPs) that are 12-50 amino acids long and readily filtered out by high stringency conditions. As some AMPs, such as gloverin and moricin, could only be found in Lepidoptera, the relevant sequences from *D. plexippus*, *M. sexta* and *B. mori* were downloaded from the NCBI GenBank as queries to search for AMPs in *P. xylostella* database. The selected immune-related *P. xylostella* genes were then manually edited by comparing with *P. xylostella* transcriptome database (http://iae.fafu.edu.cn/DBM/) and the Fgenesh program (http://linux1.softberry.com/berry.phtml). Finally, the edited sequences were manually confirmed against the NCBI curated and conserved domains (CDD) database by blastX. Domain architecture was analyzed by the SMART (http://smart.embl-heidelberg.de/), NCBI CDD database (http://www.ncbi.nlm.nih.gov/Structure/cdd/docs/cdd_search.html) and PROSITE (http://au.expasy.org/prosite/). Transmembrane domains were analyzed by TMHMM server v. 2.0 (http://www.cbs.dtu.dk/services/TMHMM/). Signal peptide was analyzed by SignalP3.0 (http://www.cbs.dtu.dk/services/SignalP/).

**Phylogenetic analysis.** The phylogenetic relationships of immune genes were compared among six insect species: *P. xylostella*, *D. plexippus*, *B. mori*, *D. melanogaster*, *A. gambiae*, and *T. castaneum*. Multiple sequence alignments were performed with ClustalW and neighbor-joining method was used to construct phylogenetic trees by MEGA5.0 with 1000 bootstrap replicates.

**Expression patterns of immune genes in *P. xylostella***. RNA sequencing of different strains, developmental stages and tissues were performed previously in our laboratory[15,59]. Based on the *P. xylostella* transcriptome database (DBM-DB, http://iae.fafu.edu.cn/DBM/), expression profiles of immune genes were analyzed and clustered by Cluster3.0. The strain- (susceptible and insecticide resistant), stage- (E: Eggs; L1-L4: 1 st, 2 nd, 3 rd and 4 th instar larvae; P: Pupae; Ma: Male adults; Fa: Female adults) and tissue- (LAEH: 4 th instar larval head; MAH: Male adult head; FAH: Female adult head; L4Mg: 4 th instar larval midgut) specific patterns of expression were then visualized using Java TreeView.

Insect rearing. A susceptible strain of *P. xylostella* (Fuzhou-S, hereafter abbreviated as SS) was collected in July 2004 from a vegetable field in Fuzhou (26.08°N, 119.28°E), Fujian province, southeastern China. All necessary permits were obtained from the Institute of Plant Protection at the Fujian Academy of Agricultural Sciences. Two insecticide resistant lines, chlorpyrifos- and fipronil-resistant lines (CRL and FRL), were selected previously[15]. The median lethal concentrations (LC50) for CRL and FRL were 574-fold (51,500.00 mg·L−1 vs. 89.79 mg L−1) and 72-fold (16.85 mg L−1 vs. 0.23 mg L−1) higher than the SS line, respectively. The three lines were reared on radish seedlings at 25 ± 2°C, 70-80% RH and a 16 h light/8 h dark photoperiod without exposure to insecticides. Adults were fed with 10% honey solution and held in 500 mL plastic bottles for mating and oviposition. The newly hatched larvae were able to pass through holes on the bottom of bottles and drop onto radish plants underneath. Individuals at various developmental stages were collected for quantitative real-time PCR (qRT-PCR) analysis. The insects used for expression analysis of pathogen-induced immune gene were reared on radish seedlings to the third instar. The Gram-negative bacteria *Enterobacter* sp. (GenBank Accession Number: JQ396388) and *Serratia* sp. (JQ396393), and the Gram-positive bacterium *Enterococcus* sp. (KC150018) which were isolated from *P. xylostella* midgut by our lab were cultured in Luria Bertani (LB) media (10 g tryptone, 5 g yeast extract, 10 g NaCl, in one liter of distilled water, pH 7.0). The 3 rd-instar larvae were placed in Petri dishes and starved for 12 hours. Cabbage leaves, dipped in a suspension of *Enterobacter* sp., *Serratia* sp. or *Enterococcus* sp. at OD600 = 1.0, were then added to the Petri dishes as diet. Cabbage leaves dipped in sterilized double distilled water were used as controls. *P. xylostella* larvae were allowed to feed on the cabbage leaves for 12 hours, and then collected for further analysis.

**Sample Collection for qRT-PCR.** To investigate the changes in gene expression profiles during various life stages, newly laid *P. xylostella* eggs (~200), the 3 rd-instar larvae, 4 th-instar larvae, pupae, and adults (10 individuals for each of the developmental stages) were collected from the SS strain, samples were surface-sterilized with 75% ethanol for 60 sec, and then rinsed with DEPC water. The 3 rd-instar larvae from CRL and FRL lines, as well as SS fed with the bacteria-dipped cabbage leaves were also collected (10 larvae per group). Total RNA was extracted from the whole bodies of each group using TRIzol (Takara Biotechnology (Dalian) Co., Ltd. (Takara Dalian)). cDNA was synthesized from 2 μg total RNA according to the instructions of GoScript™ Reverse Transcription System (Promega, USA).

qRT-PCR. qRT-PCR was performed to validate the expression profiles based on *P. xylostella* transcriptome, and evaluate immune response to bacterial infection. Primers for the tested genes are listed in Table S4. The ribosomal protein gene (RISC) was used as a control to calibrate the relative abundance of the representative genes. qRT-PCR contained 10 μL of GoTaq™qPCR Master Mix (Promega, USA), 7.2 μL of nuclease-free water, 2 μL of cDNA template from the representative samples (100 ng/μL final concentration), and 0.4 μL of each primer (10 mmol/L). qRT-PCR was performed in triplicate for each of three biological repeats in a BIO-RAD C1000 Touch™ thermal cycler with cycling parameters as follows: initial denaturation at 95°C for 3 min, followed by 35 cycles of 10 sec at 95°C and 30 sec at 58°C. To determine whether gene expressions in different lines, various stages or in response to bacterial infection...
were significantly different, data were analyzed by one-way ANOVA followed by LSD post hoc test using IBM SPSS Statistics 19.

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**Author Contributions**

X.X. and M.Y. designed the research; X.X., M.X., L.Y., H.L. and J.L. performed the research; X.X., M.Y., X.Y., S.B., G.M.G. and L.V. wrote and revised the manuscript.

**Additional Information**

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