1 Melanism patches up the defective cuticular morphological traits through promoting the up-regulation of cuticular protein-coding genes in Bombyx mori 2 3 Liang Oiao*1, Ri-xin Wang\$, You-jin Hao*, Hai Hu\$, Gao Xiong\$, Song-zhen He\$, 4 Jiang-bo Song[§], Kun-peng Lu[§], James Mallet[†], Ya-qun Xin[‡], Bin Chen^{*}, Fang-yin 5 Dai^{§1} 6 7 8 * Chongqing Key Laboratory of Vector Insects, Institute of Entomology and 9 Molecular Biology, College of Life Sciences, Chongqing Normal University, 10 Chongqing, 401331, China 11 § State Key Laboratory of Silkworm Genome Biology; Key Laboratory of Sericultural 12 13 Biology and Genetic Breeding, Ministry of Agriculture; College of Biotechnology, Southwest University, Chongqing 400716, China. 14 15 † Department of Organismic and Evolutionary Biology, Harvard University, Cambridge, MA 02138, USA 16 ‡Department of Evolutionary Ecology, LMU Munich Großhaderner Str. 2, 82152, 17 18 Germany 19 20 21 22

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Abstract

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Melanin and cuticular proteins are important cuticle components in insect. The cuticle defects caused by the loss function of cuticular protein-encoding genes could hinder melanin deposition. However, the effects of melanin variation on cuticular protein-encoding genes and the corresponding morphological traits associated with these genes remain largely unknown. Using Bombyx mori as a model, we showed that the level of melanism during larval cuticle pigmentation correlated positively with the expression of cuticular protein-encoding genes. This correlation stemmed from the simultaneous induction of these genes by the melanin precursors. More importantly, the effect of the melanism background on the cuticle induced the up-regulation of other functionally redundant cuticular protein-encoding genes to rescue the morphological and adaptive defects caused by the dysfunction of some mutated cuticular proteins, and the restorative ability increased with increasing melanism levels, which gives a novel evidence that melanism enhances insect adaptability. These findings deepen our understanding of the interactions among cuticle components, as well as their importance in the stabilization of the normal morphology and function of the cuticle.

Introduction

The prerequisite for the benefits of melanism to insect is not only the integrity of the melanin biosynthesis and regulatory pathway (Wilson *et al.* 2001; Liu *et al.* 2015; Mallet and Hoekstra 2016), but also the normal presence of the platform it relied on

67 (Wittkopp et al. 2003; Wittkopp and Beldade 2009; Andersen 2010; Moussian 2010; Van Belleghem et al. 2017). For the insects, the most important fundamental platform 68 69 for the color pattern drawing is the cuticle (Hopkins and Kramer 1992; Andersen 2010; 70 Moussian 2010). During shaping the cuticle, the maintenance and stability of the 71 cuticle features depends on normal functional cuticular proteins and the interactions 72 with other components (Hopkins and Kramer 1992; Guan et al. 2006; Suderman et al. 2006; Andersen 2010; Moussian 2010; Chaudhari et al. 2011; Noh et al. 2016). Due 73 74 to the crucial roles of cuticular proteins on cuticle development, when their coding 75 genes are loss of function, the abnormal or defective cuticle will likely affect the 76 deposition and attachment of melanin, which is not conducive to the performance of 77 color pattern (Kanekatsu et al. 1988; Oota and Kanekatsu 1993; Arakane et al. 2012; 78 Jasrapuria et al. 2012; Wang 2013; Noh et al. 2015). Yet little is known about the 79 corresponding response mode of cuticular protein-encoding genes via the alteration of 80 the melanin biosynthesis or regulatory pathway. 81 Recently, several high throughput expression surveys showed that the abundance of 82 cuticular protein-encoding genes in different colored integuments varied in some insect species, especially with evidently up-regulated in the melanic regions 83 84 (Futahashi et al. 2012; He et al. 2016; Wu et al. 2016; Tajiri 2017), and some of those 85 shared very similar expression patterns and functions (Nakato et al. 1994; Nakato et 86 al. 1997; Shofuda et al. 1999; Okamoto et al. 2008; Liang et al. 2010; Tang et al. 87 2010; Qiao et al. 2014). These studies implied that there are probably some relationships between the promotion of melanism and the expression of cuticular 88

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protein-encoding genes. Prior to this study, further exploring and understanding of the potential relationships were unclear. Additionally, when melanism-promoting instructions and defective cuticle proteins occur simultaneously, what are the effects of their relationships on the presence of the corresponding morphological traits? In the Lepidoptera model, Bombyx mori, an intriguing phenomenon has been reported that a larval melanic mutant, Striped $(p^S, 2-0.0)$ is able to rehabilitate the malformed body shape, as well as the adaptability defects of the stony mutant (st, 8-0.0) (Xiang 1995; Banno et al. 2005). A recent study clarified that a transcription factor, Apontic-like, which boosts the expression of melanin synthesis genes in epidermal cells, is responsible for the p^{S} mutant (Yoda et al. 2014). Besides this, there are also multiple alleles with different melanism degrees at the p locus, including p^{B} , p^{M} , etc (Xiang 1995; Banno et al. 2005; Yoda et al. 2014). And the stony mutant (st, 8-0.0) which precisely caused by the deletion of a RR1-type larval cuticular protein-encoding gene, BmLcp17 (or BmorCPR2) showed hard and tight touch feeling, uncoordinated ratios of the length of the internodes and the intersegment folds (I/IF), bulges at intersegment folds, and severely defective locomotion and behavioral activities in the larval stage (Qiao et al. 2014). In addition, the similarity of the gene expression patterns and functional characteristics among some members of the RR1-type larval cuticular protein-encoding gene family, such as BmLcp18, BmLcp22, BmLcp30 also suggest that they may play very similar roles as BmLcp17 in shaping the larvae cuticle (Nakato et al. 1994; Nakato et al. 1997; Shofuda et al. 1999; Okamoto et al. 2008; Liang et al. 2010; Tang et al. 2010; Qiao et al. 2014). These

dispersed findings are linked through the epistasis of p^{S} to stony, then provide the breakthrough and the exceptional genetic resources for exploring the interactions between melanin and cuticular protein-encoding genes. Here we illustrated that the transcripts levels of four cuticular protein-encoding genes were positively correlated with the melanism degree of larval cuticle, which were due to the simultaneous induction these genes by the intracellular melanin precursors. Moreover, by importing melanism-promoting instruction into the stony mutant, the cuticle deficiency rescued through functionally redundant compensation by some other up-regulated cuticular protein-encoding genes, which a new evidence that melanism as a beneficial trait. These findings deepen our understanding of the interactions among genetic factors which shape morphological features in lepidopteran, and emphasize the ecological and evolutionary significance of these mutual interactions.

Materials and Methods

Silkworm strains

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Wild-type strains Dazao ($+^p$) and melanic mutant strains p^M , p^S and p^B (Xiang 1995; Banno *et al.* 2005; Yoda *et al.* 2014) were analyzed in this study. The darkness of pigment was measured as mean OD value using Image J (https://imagej.nih.gov/ij/). In terms of melanism degree, the body color of an individual that is homozygous at one of the aforementioned melanic loci is darker than that of a heterozygous individual (Xiang 1995; Banno *et al.* 2005). The albinism mutant *albino* (*al*) (Banno

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et al. 2005; Fujii et al. 2013), non-diapause wild-type strain N4 (used for melanin inhibition treatment) and BmLcp17 deletion strain Dazao-stony (near isogeneic line of Dazao, which have been backcrossed with Dazao over 26 generations) were supplied by the Silkworm Gene Bank in Southwest University. The al mutant was fed with artificial die at 28°C, and the others were fed fresh mulberry leaves under a $12 \Box h/12 \Box h$ light/dark photoperiod at $24 \Box ^{\circ}C$. Chemicals and cell lines L-dopa (D9628), dopamine (H8502), tetrahydrofolic acid (BH₄) (T3125) and 2,4-Diamino-6-hydroxypyrimidine (DAHP) (D19206) were purchased from Sigma-Aldrich (St. Louis, MO, USA). *BmNs* cell line was kept in our laboratory. Mating combinations and progeny phenotypes identification The p^{S} and p^{M} strains were crossed with the Dazao-stony strain to generate the F_{1} generation, respectively. The F_2 generation were produced by an F_1 self-cross, and individuals of day 5 of the fifth-instar were collected to further use. The p^B strain was crossed with the Dazao-stony strain to generate F₁ progeny, which mated with the Dazao-stony strain to generate the BC₁ generation, and fed them until at day 5 of the fifth-instar. Firstly, individuals of the F2 or BC1 generations were separated by their cuticle pigmentation. Subsequently, their phenotypes were further classified by morphological characteristics, touch feeling, and the ratios between the number of internodes and intersegmental folds in the second, third, and fourth abdominal segments based on a earlier method (Qiao et al. 2014). Genotyping

Because the p^S , p^M and p^B mutations are multiple alleles at the p locus, they should be located in proximity each other on the chromosome 2 (Xiang 1995; Banno *et al.* 2005; Yoda *et al.* 2014). Based on the reported sequence of the gene corresponding to the p^S allele, a polymerase chain reaction (PCR)-based molecular marker within of the genomic region of the *Apontic-like* was designed. PCR screening were performed in the p^X (X=M, S and B) and Dazao-*stony* to obtain the polymorphism molecular marker for p locus. Similarly, molecular marker was also designed within genomic region of the *BmLcp17* for polymorphism screening of the *stony* locus. The primers used in this study are listed in Table S1.

Association analysis of gene expression, phenotype and genotype

Day 5 fifth-instar larvae of the Dazao or Dazao-*stony* strains were selected for cuticle dissection. The cuticles of the semi-lunar marking region and the non-melanic portion between the paired semi-lunar marking were finely dissected. Then, gene expression levels of BmLcp17, BmLcp18, BmLcp22 and BmLcp30 in these regions were compared. Gene expression patterns in the dorsal epidermis of abdominal segments (from semi-lunar marking to star marking) from day 5 of fifth instar larvae (Dazao, $p^S/+$, $p^M/+$, $p^B/+$). Similarly, day 1 of second instar larvae of the al and Dazao strains were also investigated. In addition, the same regions of dorsal epidermis were collected from F_2 generation individuals with the p^S/p^S , st/st, and $p^S/+$, st/st genotypes, as well as the p^M/p^M , st/st and $p^M/+$, st/st genotypes for gene expression analyses. For all genotyped individuals, the ratios of the length of the intersegment and the intersegment fold were also analysed using image J.

Melanin precursors-promoting and inhibition treatments

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The preparation and concentration of L-dopa and dopamine solutions were slightly modified according to the description by Futahashi (Futahashi and Fujiwara 2005), and filtered with a 0.22 µm filter prior to use. Cells were washed three times with Grace medium without melanin precursors to remove metabolites and other contaminants on the cell surface. Then, 0.8mL medium containing L-dopa or dopamine was added separately into each well, and the medium without melanin precursors was used as the control. The plate was sealed with tape, wrapped with foil and incubated at 28°C for 24 h for a gene expression analysis. In BH₄ feeding assays, the 30mM working solution was prepared by dissolving tetrahydrofolic acid into ddH_2O , and then smeared on the artificial diet to feed the *al* mutant, and ddH_2O was used as the control. Phenotype were observed and recorded from the second instar and the expression of cuticular protein-encoding genes was also analysed. For the melanism-inhibition experiment, the wild-type strain N4 was selected. Newly-hatched larvae were divided into treatment and control groups. Individuals in the treatment group fed with DAHP dissolved in 0.1M NaOH (15g/L), and individuals in the control group fed 0.1M with NaOH. Phenotypes observation and gene expressions detection were performed on day 1 of the second-instar larvae.

Quantitative RT-PCR

Total RNA extraction, reverse transcription and qRT-PCR conducted as described previously (Qiao *et al.* 2014). Three biological replicates were prepared for each condition, and *BmRPL3* was used as the internal control. Primers are listed in Table

199 S1. 200 201 **Results** Entirely distinct expression patterns of cuticular protein-encoding genes and the 202 203 cuticle appearance between melanic and non-melanic regions 204 The expression patterns of BmLcp17, BmLcp18, BmLcp22 and BmLcp30 in melanic or non-melanic epidermal regions are shown in Figure 1. These results, together with 205 206 earlier studies (Futahashi et al. 2012; He et al. 2016; Wu et al. 2016), revealed that the 207 gene expressions were significantly higher in melanic parts of the cuticle than in 208 non-melanic parts. Moreover, micro protrusions were more intensive in the melanic 209 regions than in the non-melanic regions, and accompanied by a higher chitin content 210 (another important cuticle component (Hopkins and Kramer 1992; Moussian et al. 211 2006; Andersen 2010; Moussian 2010; Chaudhari et al. 2011; Qiao et al. 2014), and 212 the reduction chitin content was reported to impede cuticle melanism (Moussian et al. 213 2005)) (Figure S1A). These results showed that, regardless of the different genetic 214 basis of the melanic mutants or the melanic markings in the non-melanic strains, excessive accumulation of melanin in the cuticle (accompany by the changes in the 215 216 surface structure and the chitin content of the melanic cuticle) was closely related to 217 the high expression levels of the cuticular protein-encoding genes. 218 Expression level of larval cuticular protein-encoding genes positively correlated 219 with the degree of cuticle melanism 220 To obtain further insights into the relationships between the accumulation of

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melanin and the expression of cuticular protein-encoding genes, we investigated gene expression patterns using four different mutations at the p locus (Dazao $(+^p)$, p^M , p^S and p^{B}), which showed gradually increasing melanin accumulation (Figure 2A). Our results revealed that the gene expression levels were gradually and significantly up-regulated with the increase in the degree of melanism of the cuticle (Figure 2B). They further showed that the expression levels correlated positively with the degree of melanism. Thus, the quantities of cuticular proteins with similar or redundant functions could be increased greatly by the increasing the degree of melanism. Typical stony phenotyped individuals masked after the introduction of melanic loci into the BmLcp17 defection strain We assessed the effects of modulating the melanic background on the phenotypic defects caused by the deletion of BmLcp17. After mating the p^B and stony parental, the percentage of BC₁ individuals with melanism and the normal body shape in the backcrossed population of the $p^B \times stony$ cross was 52 % (290/553; and theoretically, it was 25 %), yet individuals with the melanism cuticle and stony-type body shape were not found (theoretically should be almost equivalent to the number of individuals with melanism cuticle and normal body shape) (Figure 3). In the cross of $p^{S} \times stony$, ~10.8% (36/331) of F₂ progeny had an lighter melanism body color and smaller body size (Figure 3). These individuals exhibited a little hard and tight touched body, but the body was much softer than that of the stony mutant. Their intersegment folds bulged slightly, and the length were significantly shorter than that of the internodes; accordingly, their phenotypes slightly resembled the morphological features of the

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stony mutant (Figure 3 and 4A). Even so, We did not find individuals with the typical stony-type body shape and also defective adaptability under the melanism background (theoretical ratio is 3/16). (Figure 3). Similarly, in the $p^{M} \times stony$ corss, only approximately ~11.7% (51/437) of the individuals of the F₂ population were very lightly melanic, but they exhibited obviously unusual morphological features (Figure 3, Figure 4A). When compared with the ~10.8% F₂ individuals (aforementioned) from the $p^S \times stony$ cross, the touch feelings of individuals from the $p^M \times stony$ cross were tighter and firmer, and the intersegment folds bulged more obviously and had a higher proportion among the segments, meaning that their body features were closer to the phenotype of the *stony* mutant (Figure 3 and 4A) (Qiao *et al.* 2014). Nevertheless, melanic individuals showing the typical stony-type body features and defective adaptability still did not appear (theoretical ratio is 3/16) in the progeny from the p^{M} × stony cross. Therefore, induction of the melanic mutation into individuals with a defective cuticular protein-encoding gene could mask the adverse phenotypes, and the masking effect was more remarkable with the increasing degree of melanism. Other larval cuticular protein-encoding genes up-regulated evidently in the melanic and non-stony phenotypic, but with mutated BmLcp17 genotypic offspring Using the molecular markers (closely linked to the p and st loci, respectively), we further genotyped the progenies with melanic color pattern and non-stony (including ambiguous stony-like) body shape. The results revealed that approximately 49% of the individuals showing a melanic color and normal body shape in the BC₁ population from the $p^B \times stony$ cross was the $p^B/+p^B$, st/st genotype (Figure 4A). The ratios of the

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length of the internodes and the intersegment folds (I/IF) were ~4, which is very similar to that in $p^B/+p^B$, $+s^t/st$ individuals, and also no significant differences as that in the wild-type individuals (Figure 4B) (Qiao et al. 2014). In the F₂ generation from the $p^S \times stony$ cross, approximately 9.3% of the individuals with the p^S/p^S , st/st genotype and very few individuals (~1%) with the genotype $p^S/+p^S$, st/st exhibited a melanic color and the normal body shape (Figure 4A). For p^S/p^S , st/st genotyped individuals, the I/IF value was approximately 3.3 (Figure 4B). Although the I/IF value was lower than that in $p^{S}/_{-}$, $+^{st}/_{-}$ individuals (approximately 4, Figure 4B), it was much higher than that in the stony mutant (approximately 1.6 (Qiao et al. 2014)). Despite the slightly smaller body size of the p^S/p^S , st/st individuals, their body shape was essentially normal (Figure 4A). The genotypes of those lightly melanic individuals, whose body shape was slightly like that of the stony phenotype (mentioned in Result 3), were all the $p^S/+p^S$, st/st, and the I/IF value of these individuals was approximately 2.7 (Figures 4A and 4B). In progeny of the $p^{M} \times stony$ cross, ~10.5% of the individuals with the p^{M}/p^{M} , st/st genotype and ~0.7 % of the individuals with the $p^{M}/+p^{M}$, st/stgenotype showed a melanic color and subtle stony features (just very slight bulges) (Figure 4A). The body size of the p^{M}/p^{M} , st/st individuals were smaller than those of the $p^{M}/_{-}$, $+^{st}/_{-}$ individuals. They exhibited a certain sense of hardness, and the I/IF ratio was approximately 2.8, which is in good agreement with their phenotypes (Figure 4B). Additionally, individuals showing very slight melanism and morphological features that were more similar to that of the stony mutant (mentioned in Result 3) were all the $p^M/+p^M$, st/st genotype, and their I/IF values were

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approximately 1.8, which is closer to that of the stony mutant (Figures 4A and 4B). In addition, the expression of cuticular protein-encoding genes in p^S/p^S , st/st individuals were significantly higher than that in $p^S/+p^S$, st/st individuals (Figure. 4C); a similar result was also obtained from the p^{M}/p^{M} , st/st and $p^{M}/+p^{M}$, st/st individuals (Figure 4C). Taken together, these results revealed that more cuticular proteins with similar functions were accumulated in the cuticle of melanic homologous individuals at the p locus. Based on the comprehensive and association analysis, we infer that melanic background effectively drove the expressions of cuticular protein-encoding genes with similar expression patterns and redundant functions, which compensated for the morphological and adaptability defects caused by the dysfunctional BmLcp17 gene; and the law of compensatory abilities was $p^B/+p^B$, $st/st > p^S/p^S$, $st/st > p^M/p^M$, st/st < $p^{S}/+^{pS}$, $st/st > p^{M}/+^{pM}$, st/st, which corresponds well with the gradual weakening of the degree of melanism (Figures 4 and Figure S3). Content variations of melanin precursors affect the transcript amount of the cuticular protein-encoding genes Due to the crucial material basis for the cuticle melanism (no matter what kind of genetic basis caused by) is the extensive accumulation of melanin precursors in the epidermal cells; thus, the essence that melanism tend to increase the expression of some cuticular protein-encoding genes should be driven by the relations between the accumulation of melanin precursors and the transcripts amount of the cuticular protein-encoding genes. The basal expressions of four cuticular protein-encoding genes were detected in BmNs cells (Figure S5), indicating that there are regulatory

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pathways controlling the expression of cuticular protein-coding genes in this cell line; therefore, this cell line can be used to examine the effect of melanin precursors on gene expressions. After incubating BmNs cells with melanin precursors, the expression of cuticular protein-encoding genes was significantly higher in cells treated either by L-dopa or dopamine, compare with that in the control group (Figure 5 top (left)). In addition, when treated with BH₄, the second-instar al mutant (which is characterized by albinism and a porous cuticle due to a mutation of sepiapterin reductase, which leads to the insufficient synthesis of the co-factor BH₄ during the synthesis of melanin precursors (Banno et al. 2005; Fujii et al. 2013) was rescued to a melanic body color (Lab unpublished contribution) (Fujii et al. 2013), and the gene expressions were obviously higher than that in the control group (Figure 5 top (right)). These results suggest that the expression of cuticular protein-encoding genes can be induced by increasing amounts of melanin precursors. Furthermore, adding DAHP (an inhibitor of guanylate cyclase hydrolase (GTPCHI), which is an important rate-limiting enzyme in the synthesis of BH₄ (Hamadate et al. 2008), inhibited the synthesis of BH₄ in wild-type second-instar larvae by blocking the synthesis of melanin precursors in epidermal cells, which caused individuals to lose their original melanic body color (Lab unpublished contribution). The gene expressions were also significantly reduced when compared with that in the control group (Figure 5 bottom). Thus, the content and variation of the intracellular melanin precursors are important factors regulating the expression of cuticular protein-encoding genes. We concluded that the inducing effect of the melanin precursors on the expression of cuticular

protein-encoding genes is the basis for melanism promoting genes' transcription.

Discussion

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Melanin is deposited in the cuticle, in which many of cuticular proteins also occur; thus, overlapping distribution of cuticular proteins make it possible for them to interact with melanin, thereby contributing to the phenotype of the cuticle (Hopkins and Kramer 1992; Suderman et al. 2006; Andersen 2010; Moussian 2010; Hu et al. 2013). We clearly observed that extensive star-like protrusions were present on the cuticle when a large amount of melanin accumulated (Figure S1). Similar correlations between cuticle structure and body color have been reported multiple times (Futahashi et al. 2012; Noh et al. 2016; Tan et al. 2016), suggest that there should be some interactions between cuticular proteins and melanin. Although the exact interaction pattern between these two cuticular components is unknown, microscopic observation clearly shows that the deposition of an excessive amount of melanin affected the cuticle characteristics. The expression profile of cuticular protein-encoding genes in melanic silkworm mutants and the black markings of Papilio larvae supported the view that overexpressed cuticular proteins probably participate in the transport or maintenance of the corresponding pigments in a specific painted cuticle (Figures 1, 2 and Figure S1) (Futahashi et al. 2012; He et al. 2016; Wu et al. 2016). Yet over-expression of cuticular protein-encoding genes in vivo cannot trigger (or induce) the formation of the melanic cuticle (Tajiri et al. 2017). Therefore, we reasoned that once the

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instructions for melanin accumulation are included in the developmental program (melanism being the original factor), other cuticle features and structures should adapt to the level of melanin accumulation, regardless of genetic background. The up-regulation of cuticular protein-encoding genes should be a necessary adaptation for the maintenance and stability of the structural characteristics and physical properties of melanic cuticles. The interactions of melanin and cuticular proteins at the ultrastructural level deserves special attention in follow-up studies. Despite the elaborate regulatory mechanism by which melanin precursors affect the expression of cuticular protein-encoding genes is unclear, this study really revealed the existence of this regulatory phenomenon (Figure 5). Cuticle formation is regulated stringently in temporal and spatial patterns, the accumulation and oxidization of melanin precursors, and the interactions (such as cross-linking) among other components should occur after the formation of the initial formation of the cuticle (Moussian 2010; Sobala and Adler 2016; Tajiri 2017). Therefore, we propose that the cuticular proteins induced by melanin precursors are used to set up a platform for further accumulation and oxidization of melanin precursors. When melanin-associated protein-encoding genes have similar expression patterns and functions (Nakato et al. 1994; Nakato et al. 1997; Shofuda et al. 1999; Okamoto et al. 2008; Liang et al. 2010; Tang et al. 2010; Qiao et al. 2014), the production of a large amount of functionally similar cuticular proteins would be driven by the melanic background to guarantee construction and stability of the melanic cuticle. During this process, melanism unlocks the complementary features of melanin-associated cuticular proteins (such as

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BmLcp18, BmLcp22, BmLcp30), rescued cuticular malformation caused by the loss of function of some cuticular proteins (such as defected *BmLcp17* in *stony* mutant). Because it appears that a special cuticle-forming pattern is regulated by melanin accumulation, melanism may enhance insect adaptability to avoid the impairment of survival caused by the mutation and/or functional loss of some cuticular proteins, and these results also add new evidence to explain how melanism can be a beneficial trait (Wilson et al. 2001; True 2003; Wittkopp et al. 2003; Wittkopp and Beldade 2009; Liu et al. 2015; Mallet and Hoekstra 2016). As far as we known, as structural proteins, there is no evidence to suggest that the four larval cuticular protein-encoding genes and their orthologous can enter the nucleus and regulate gene expression by acting as transcription factors. In addition, our findings and some other studies showed that changes in the expression of one cuticular protein-encoding genes of R&R family did not affect the expression of other members (Figure S6, S7) (Arakane et al. 2012; Noh et al. 2015). Moreover, several studies revealed that organisms optimize resources use at gene or protein expression level (Liebermeister et al. 2004; Dekel and Alon 2005); thus, the expression of melanin associated cuticle protein-encoding genes might be appropriately and simultaneously coordinated with the sufficient accumulation of intracellular melanin precursors, as a relatively direct, economical, efficient and convenient strategy. Furthermore, DAHP inhibits GTPCHI activity, but it does not directly impair the extracellular accumulation of melanin and protein-protein interactions in the cuticle. Finally, there is some evidence that melanin precursors regulate gene expressions via

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the receptors (Konradi et al. 1996; Berke et al. 1998; Westin et al. 2001). In summary, we hypothesized that the up-regulation of the four BmLcp genes were induced simultaneously by excessive amounts of melanin precursors, but should not be the interaction among these four genes on regulation level. To test our hypothesis, further analyses will be performed to determine the expression of the remaining BmLcp genes in some BmLcps mutant (such as defective BmLcp17) cell line by increasing or decreasing the content of melanin precursors. The markings in the *stony* mutant were lighter than those in the wild-type strain, and accompanied by the down-regulation of melanin synthase genes (Figure S4). If the dysfunction of some cuticular proteins cannot be effectively supplemented, abnormalities of the cuticle structure and interactions of various cuticular components may occur, which probably result in barriers for melanin deposition and metabolism. Perhaps this is the reason why the $p^S/+p^S$, st/st (or $p^M/+p^M$, st/st) genotypes are lighter than the $p^S/+p^S$, $+s^t/$ (or $p^M/+p^M$, $+s^t/$) genotypes, besides to their different body shape. Because the homozygous of p^B mutation is lethal (Xiang 1995; Banno *et al.* 2005), we were unable to obtain F_2 progeny with the p^B/p^B , st/st genotype. However, it is worth noting that the body shape and the melanism degree of $p^B/+^{pB}$, $+^{st}/st$ and $p^B/+^{pB}$, st/stgenotype individuals were almost same (Figure 4A). Therefore, during cuticle formation, if the signal promoting melanism is sufficiently strong, plenty of melanin precursors should be generated. The contents of functionally redundant proteins induced by melanin precursors are enough to fill the gap generated by the dysfunction of some cuticular proteins in theoretically, which will be able to guarantees the normal

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accumulation of melanin. Therefore, we believe that there should be a threshold of the melanism-promoting effect. When the accumulation of melanin precursors spans this threshold, the requirements for cuticular proteins with similar function are not weakened even when one of them loses the function, which ensuring the formation of a normal cuticle structure the later pigmentation. We proposed a possible regulatory model as follows: when large amounts of melanin precursors induced by endogenous- and/or exogenous melanism-promoting factors, the melanin pathway may directly or indirectly induce the up-regulation of some cuticular protein-encoding genes to guarantee the formation of normal structural features of the melanic cuticle. During this process, if some cuticular proteins lose their functions, other functionally redundant cuticular proteins induced by melanin precursors can compensate for the functional defects. The compensatory intensity is increased with increasing melanin accumulation. When melanin accumulation spans a certain threshold, this compensation can totally mask the defective phenotype caused by the malfunctioning genes, which adds to growing evidence that melanism may have pleiotropic effects that enhance fitness over and above the effect of melanin accumulation itself (Figure 6). Due to the coexistence of excess melanin and cuticular proteins is common in other insects, and homologues the four BmLcp gene are widely distributed in the Lepidoptera (Table S2), we presume that the above reciprocal action and its corresponding biological significance are conserved in other Lepidoptera insects to that in *Bombyx mori*. In summary, we used crucial components of the cuticle to elucidate the mutual

- 441 effects among some key genetic factors, and the physiological significance of these
- 442 mutual effects during the development of the morphological features. Our findings not
- only contribute a realistic basis to in-depth study of the interaction patterns of melanin
- and cuticular proteins, but they will also inspire relevant studies of other Lepidopteran
- insects or other insect species.

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Acknowledgments

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Figure legends

- 596 Figure 1. Expression of four larval cuticular protein-encoding genes in melanic and
- 597 non-melanic integuments. A and B represent the gene expressions between the

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semi-lunar marking (black box) and the non-melanic region (between the semi-lunar marking, red box) in Dazao or Dazao-stony, respectively. Scale bar: 2 mm. C. Comparative analysis of gene expressions in the dorsal side of abdominal segments (from the third to the fourth segment, red box) between the p^{S} and Dazao strains. Scale bar: 1 cm. D. Comparison of gene expressions between the second-instar al mutant and the Dazao strain (melanic). The red hashtag symbol indicates the Fig. 1 D we are showing is cited from the previous study of our lab group (Min et al. 2016) with modification. Scale bar: 2 mm. t-test, n=3; * p < 0.05; ** p < 0.01; *** p < 0.01; 0.001. Figure 2. Expression of cuticular protein-encoding genes in integuments showing different degree of melanism. A and B display comparisons of degree of melanism and cuticular protein-encoding gene expression levels among four strains with mutant alleles at the p locus. Scale bar: 1 cm. Ratios represent the ratios of gene expression levels between two strains. Symbols (-, +, ++, and +++) represent the increment of the degree of melanism. Star represents the melanin-associated cuticular protein-encoding genes. *t*-test, n=3; * p < 0.05; ** p < 0.01. Figure 3. Segregation patterns of the phenotypes of progenies from different crosses of melanic mutant strains and the *stony* strain. In the segregated progenies, the first item in the list (B,N), (B,st), (S,N), (S,st), (M,N), (M,st), (N,N) and (N,st) represents p^B -, p^S - p^M - and Normal color patterns, respectively. The second item indicates body

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shape features marked with non-stony type (N) and the stony type (st). It is noteworthy that in (S, st-am+) and (M, st-am+), the second item represents the ambiguous stony body shape. The size of "+" symbol represents the corresponding degree of the ambiguous stony body shape. Superscript "T"s represent theoretical values. Superscript "A"s represent actual values. Backslashes indicates that a value was not obtained. Chi-squared test, * p < 0.05; ** p < 0.01. Figure 4. Association analysis of the genotypes, phenotypes and gene expression levels in segregated progenies from different crosses. A. Correlation analysis between the genotypes and phenotypes in self-crossed or backcrossed progenies. Scale bar: 1 cm. White and red stars represent polymorphic bands at the $+^p$ and p^X loci (X = B, S) or M), respectively. Red and white hash-tag represent polymorphic bands at the st and $+^{st}$ locus, respectively. Solid and dotted red arrows indicate the relative degree of bulging (solid > dotted), respectively. Slashes show the genotypes within one phenotypic category. The thickness of the slash represents the proportion of the corresponding genotypes. Dotted backslashes indicate that the number of individuals with the corresponding genotype is quite low. B. Ratios of the length of internodes and intersegmental folds in the second, third, and fourth abdominal segments of individuals with different genotypes (showing melanic body color) in the self-crossed or backcrossed progenies. $n \ge 3$, t-test, ** p < 0.01. C. Gene expression analysis of cuticular protein-encoding genes in homogeneous and heterogeneous individuals at the p locus from self-crossed progenies of $p^S \times stony$, and $p^M \times stony$ under the

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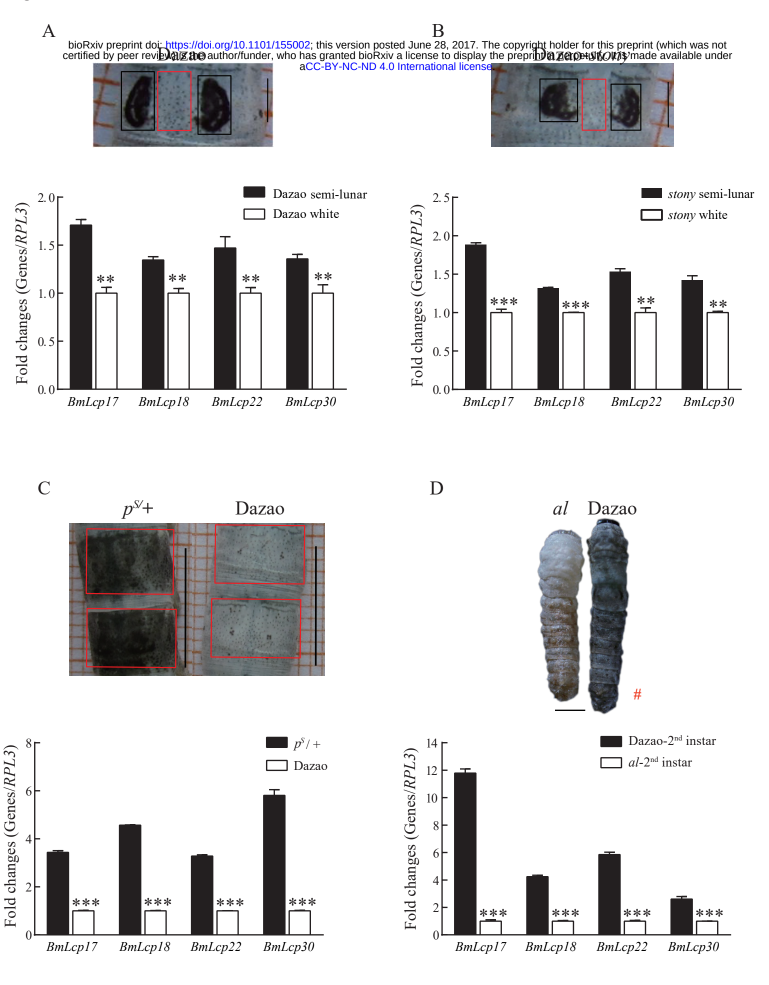
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condition that the cuticle was melanic and the genotype was homozygous recessive at st locus. n = 3, t - test, * p < 0.05; ** $p \le 0.01$. Figure 5. Effect of melanin precursors (top left: in cells) and BH₄ (top right: in vivo) treatments on the expression of cuticular protein-enconding genes (t-test; n = 3, * p <0.05; ** p < 0.01), and variations of gene expression levels in larval cuticle treated with the inhibitor DAHP (bottom). n = 3, t-test, ** p < 0.01. **Figure 6.** Schematic overview of the effect of melanin precursors on the expressions of cuticular protein-encoding genes. Black solid circles represent the melanin. Triangles represent the BmLcps with similar expression patterns and functions. Solid and hollow triangles represent the normal and defective functions, respectively. Red rhombi represent other components (such as chitin) in the cuticle. Solid and dotted double-headed arrows indicate probable interactions and loss of interactions due to the functional deficiency of some BmLcps (such as dysfunctional BmLcp17), respectively. Purple arrows show the direction in which the melanin precursors or cuticular proteins flow from the haemolymph to the epidermal cells as well as from the epidermal cells to the cuticle. Red arrows indicate the increased contents of melanin precursors or the up-regulation of cuticular protein-encoding genes. Purple polyline arrows indicate melanism-promoting factors produced by other genetic instructions. Double dovetail arrows indicate the effect of melanin precursors on cuticular protein-encoding genes. The question mark indicates that the details of induction (directly or indirectly) are unclear. Here the cuticle especially means the exo-cuticles in which the melanin and cuticular proteins deposited in (Hopkins and Kramer 1992; Suderman *et al.* 2006; Andersen 2010; Moussian 2010).



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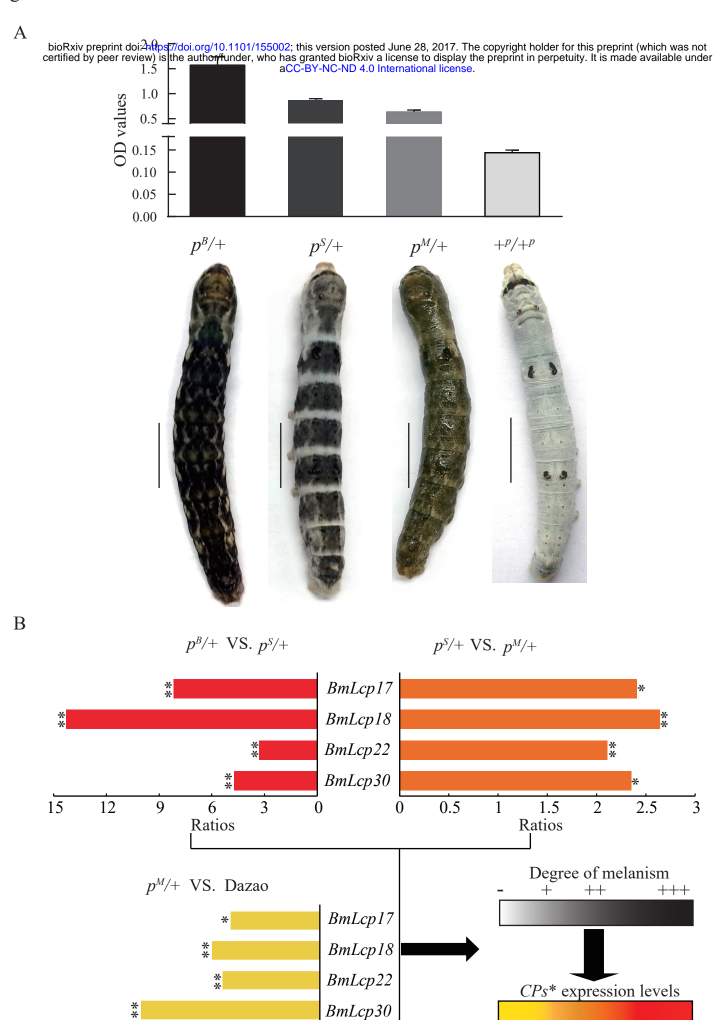


Figure 3

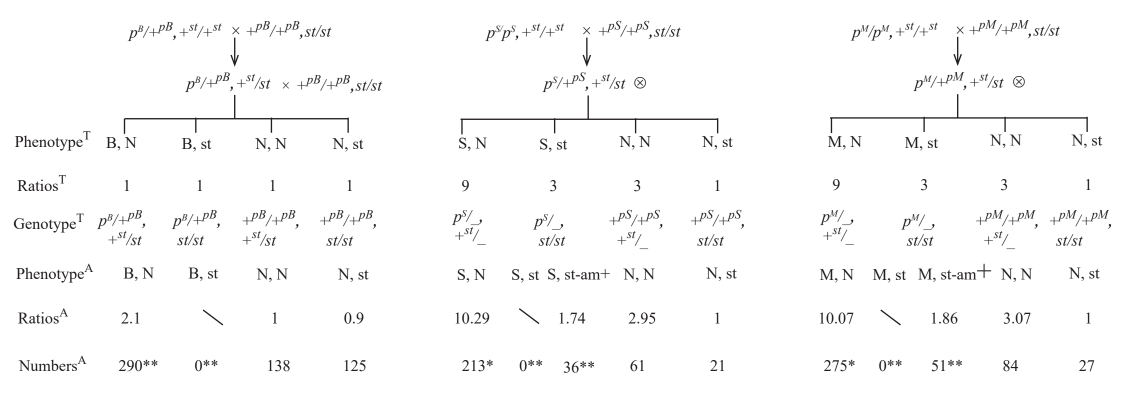
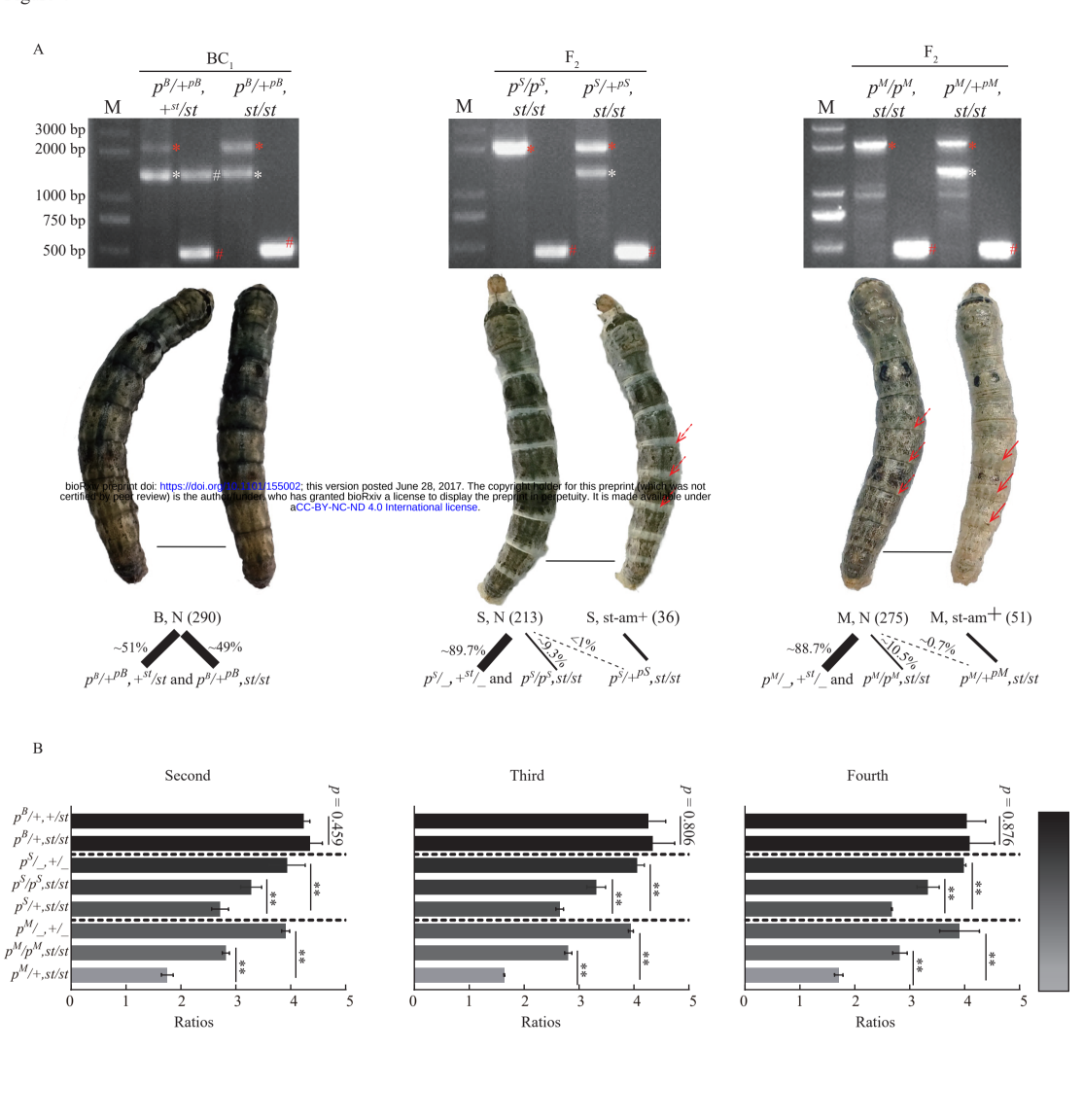
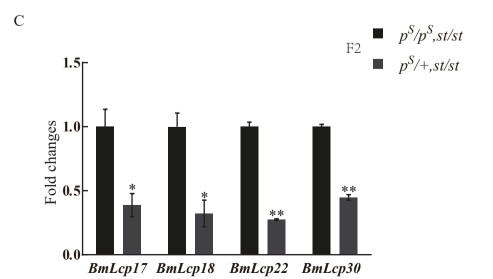


Figure 4





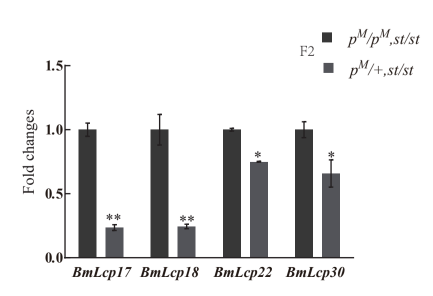


Figure 5

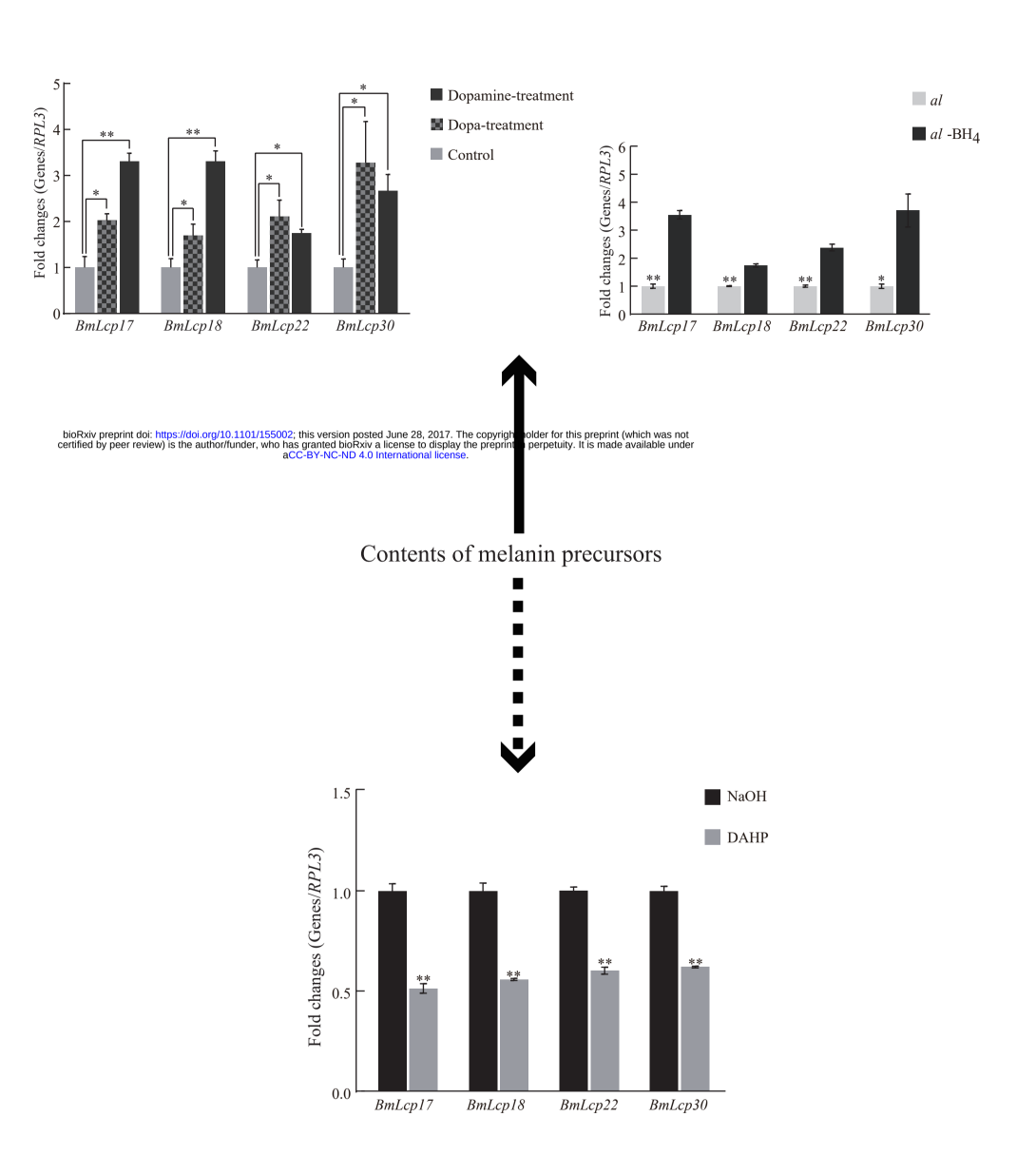


Figure 6

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