

SCIENTIFIC REPORTS



OPEN

Molecular evolutionary mechanisms driving functional diversification of α -glucosidase in Lepidoptera

Received: 05 December 2016

Accepted: 01 March 2017

Published: 12 April 2017

Xiaotong Li*, Liangen Shi*, Yanyan Zhou, Hongqing Xie, Xiangping Dai, Rongqiao Li, Yuyin Chen & Huabing Wang

The digestive tract of lepidopteran insects is unique given its highly alkaline pH. The adaptive plasticity of digestive enzymes in this environment is crucial to the highly-efficient nutritional absorption in Lepidoptera. However, little is known about the molecular adaptation of digestive enzymes to this environment. Here, we show that lepidopteran α -glucosidase, a pivotal digestive enzyme, diverged into sucrose hydrolase (SUH) and other maltase subfamilies. SUH, which is specific for sucrose, was only detected in Lepidoptera. It suggests that lepidopteran insects have evolved an enhanced ability to hydrolyse sucrose, their major energy source. Gene duplications and exon-shuffling produced multiple copies of α -glucosidase in different microsyntenic regions. Furthermore, SUH showed significant functional divergence (FD) compared with maltase, which was affected by positive selection at specific lineages and codons. Nine sites, which were involved in both FD and positive selection, were located around the ligand-binding groove of SUH. These sites could be responsible for the ligand-binding preference and hydrolytic specificity of SUH for sucrose, and contribute to its conformational stability. Overall, our study demonstrated that positive selection is an important evolutionary force for the adaptive diversification of α -glucosidase, and for the exclusive presence of membrane-associated SUHs in the unique lepidopteran digestive tract.

The Lepidoptera (butterflies and moths) is one of the most widespread and widely recognisable insect orders in the world. This order contains approximately 180,000 described species in 126 families and 46 superfamilies¹. The larvae of many lepidopteran species are major pests and are considered to be the most economically damaging pests in agriculture. The digestive system of Lepidoptera is quite different from that of other insects and is more complex². All the digestive enzymes of Lepidoptera, other than those for initial digestion, are immobilised at the surface of the midgut cells². In addition, the digestive tract of Lepidoptera is unique because of its extremely alkaline pH^{3,4}, and the pH values measured in particular compartments of the larval digestive tract span a range between 9 and 11^{2,5}. The lepidopteran gut is highly alkaline due to specific dietary preferences^{6–8}, such as feeding on tannin-rich leaves⁹. The digestive enzymes, which have evolved into a specific pH optimum, should match the midgut condition for maximum efficiency. However, the molecular mechanisms of the phylogeny and adaptation of lepidopteran digestive enzymes are still poorly understood.

Sucrose is one of the main products of photosynthesis and the most common transported sugar in plants, and it is also an easily assimilated macronutrient that provides a carbon or energy source for insects. Insect sucrases catalyse the hydrolysis of sucrose into its constituent monosaccharides, which can be used by insects as a food source. Insect sucrase activity is generally thought to depend mainly on α -glucosidase (EC 3.2.1.20). However, sucrose hydrolases in the larval midgut of Lepidoptera have three distinct forms: an α -glucosidase, also known as maltase; a β -fructofuranosidase, which is acquired via horizontal gene transfer (HGT) from bacteria; and a sucrose hydrolase (SUH), which displays specificity for sucrose^{10–13}. Unlike typical α -glucosidase and β -fructofuranosidase, the SUH, which is associated with the midgut membrane, displayed measurable activity only against sucrose and showed a very broad range of pH optima, ranging from approximately pH 6 up to 11.

College of Animal Sciences, Zhejiang University, Hangzhou 310058, China. *These authors contributed equally to this work. Correspondence and requests for materials should be addressed to L.S. (email: slgsilk@zju.edu.cn) or H.W. (email: wanghb@zju.edu.cn)

Recently, sucrose hydrolases categorised into α -glucosidases were found in three lepidopteran species, *Bombyx mori*, *Trilocha varians* and *Samia cynthia ricini*, and were named as BmSUH, TvSUH and ScSUH, respectively¹⁴. Recent genome sequencing projects have shown that SUH sequences are present in several lepidopteran species and absent from other insect orders. Although SUHs belong to α -glucosidases¹⁵, the SUHs were clearly distinct from other α -glucosidases, suggesting that SUHs have diverged from other α -glucosidases during the evolution of lepidopteran insects. Therefore, the evolution of α -glucosidases in Lepidoptera is highly unusual.

Insect α -glucosidases have been studied extensively in Brachycera and Nematocera, and are likely results of an ancient series of duplications^{15–17}. α -glucosidase family underwent a complicated evolutionary history in Diptera, and some amino acid sites might be subjected to the pressure of positive selection or force of functional divergence (FD), which is expected to result in diversified functions. Although a large variety of α -glucosidase genes have been recently identified in Diptera^{15–18}, only a few lepidopteran α -glucosidases were characterised, and not much is known about how such essential enzymatic functions evolve and diversify¹⁴. The recent sequencing of lepidopteran genomes has made the exploration of the molecular functions and evolutionary adaptations of SUH and other α -glucosidases in Lepidoptera possible.

In this study, we provide the first comprehensive analysis of the evolution of the α -glucosidase family in Lepidoptera. We found that α -glucosidase could be divided into two subfamilies, SUH and lepidopteran maltase (LMal), approximately 232 MYA in seven lepidopteran species. SUH and LMal were duplicated to several homologues subsequently, and the splice sites were conserved in each subfamily but differ from each other. FD and selection pressure analyses indicated that SUH or LMal fall under the effects of FD and positive selection, and we detected some amino acid residues that are prone to being influenced. In addition, we found that nine sites, which were involved in both FD and positive selection, were located around the ligand-binding groove of SUH. This study provides an example of how distinct selective pressure, along with ancestral gene differentiation and subsequent gene duplications, can contribute to lineage-specific adaptations and the divergence of lepidopteran digestive enzymes.

Results

Phylogenetic relationship of LMal and SUH in Lepidoptera. In this study, we confirmed the evolutionary relationship between the maltase of Lepidoptera and those of other insect orders. We also determined whether SUH emerged before or after the divergence of LMal and other insect maltases. We retrieved and aligned 62 sequences of the insect α -glucosidase family from the NCBI and Ensembl databases using DmMal1A and BmSUH as query sequences (Supplementary Table S1). After reconstructing the phylogenetic tree by maximum likelihood (ML) and Bayesian methods, we found that the phylogenetic relationship of the insects α -glucosidases was well defined. Each subfamily represented a distinctive monophyletic clade, as supported by the high confidence values observed (Fig. 1). SUH, a subfamily of maltase from Lepidoptera, split from the maltase clade of lepidopteran and dipteran insects (Fig. 1). LMal was duplicated at least thrice to form a large gene family. By contrast, the SUH subfamily duplicated into the *SUH1* and *SUH2* clusters. Moreover, SUH and LMal shared conserved structural characteristics that harboured an α -amylase catalytic (α -amy) domain and seven conserved sequence regions (CSRs) (Supplementary Fig. S1). In addition, SUH possessed an N-terminus hydrophobic amino acid sequence (except AtSUH) (Supplementary Fig. S1). Recent studies have proved that the genes in this distinct clade, specifically, *BmSUH*, *TvSUH* and *ScSUH*, encode membrane-associated sucrose-specific hydrolases at the molecular level¹⁴.

To ascertain the genomic relationship of *SUH1* and *SUH2*, we performed syntenic analyses around the region of *SUH1* and *SUH2* on the chromosomes of seven lepidopteran species (Fig. 2). Only one copy of *SUH* was detected in the genome of *B. mori*, *H. melpomene* and *A. transitella*. However, the *SUH* gene was duplicated to double adjacent copies, *SUH1* and *SUH2*, in *D. plexippus*, *P. xuthus*, *P. machaon* and *P. polytes*. Conserved gene organisation up- and downstream of *SUH* was also found in seven lepidopteran species. RT-PCR shows that *PxSUH1* and *PxSUH2* are mainly expressed in the midgut of fifth-instar larvae of *P. xuthus*, and *PxSUH1* showed a more abundant expression compared with *PxSUH2* (Supplementary Fig. S2). This conserved syntenic relationship indicated that the *SUH* gene was conserved and emerged early during lepidopteran evolution, and its duplication likely occurred specifically among many Rhopalocera species, but not in moth.

Evolutionary history and gene structure of LMal and SUH in Lepidoptera. To shed light on the evolutionary history of LMal and SUH, we estimated the divergence time of particular gene duplications from the ML tree using RelTime method^{19,20} (Fig. 3). With two calibration points of divergence time between *DmMalB1* and *B2* (84 MYA) as well as *DmMalA2* and *DmMalA3/A4/A5* (155 MYA)¹⁵, we were convinced that the divergence time of *SUH/maltase* was approximately 232.07 MYA. After the differentiation of the common ancestral gene from which *SUH* and *maltase* are derived, *SUH* was divided into *SUH1* and *SUH2* approximately 111.62 MYA. Moreover, LMal and DMal (*maltase* of Diptera) were diverged approximately 208.68 MYA, and LMal subsequently was divided into *LMal1*, *LMal2*, *LMal3* and *LMal4* 98.87, 73.16, 60.42 and 48.41 MYA, respectively. Furthermore, the *LMal3* gene was further diverged into *LMal3-1* and *LMal3-2* in several lepidopteran species, and this duplication event occurred only 29.22–35.96 MYA. In addition, we conducted molecular dating by Bayesian inference methods with BEAST software (Supplementary Fig. S3). The BEAST estimation was similar with the divergence time calculated by RelTime method. Therefore, the evolutionary history analysis indicated that *SUH* and *LMal* were diverged from the last common ancestral gene a very long time ago, and some subsequent gene duplication episodes occurred specifically in certain lepidopteran species, which proved that the α -glucosidase family has experienced a complicated evolutionary process in Lepidoptera.

Structural divergences are prevalent in duplicated genes and, in many cases, lead to the generation of functionally distinct paralogs²¹. Splice-site analysis was conducted to investigate the gene structures of *SUH* and *LMal* (Fig. 4). *SUH2* genes have minimal numbers of exons compared with *SUH1* and *LMal*, whereas the exons of *Mali*

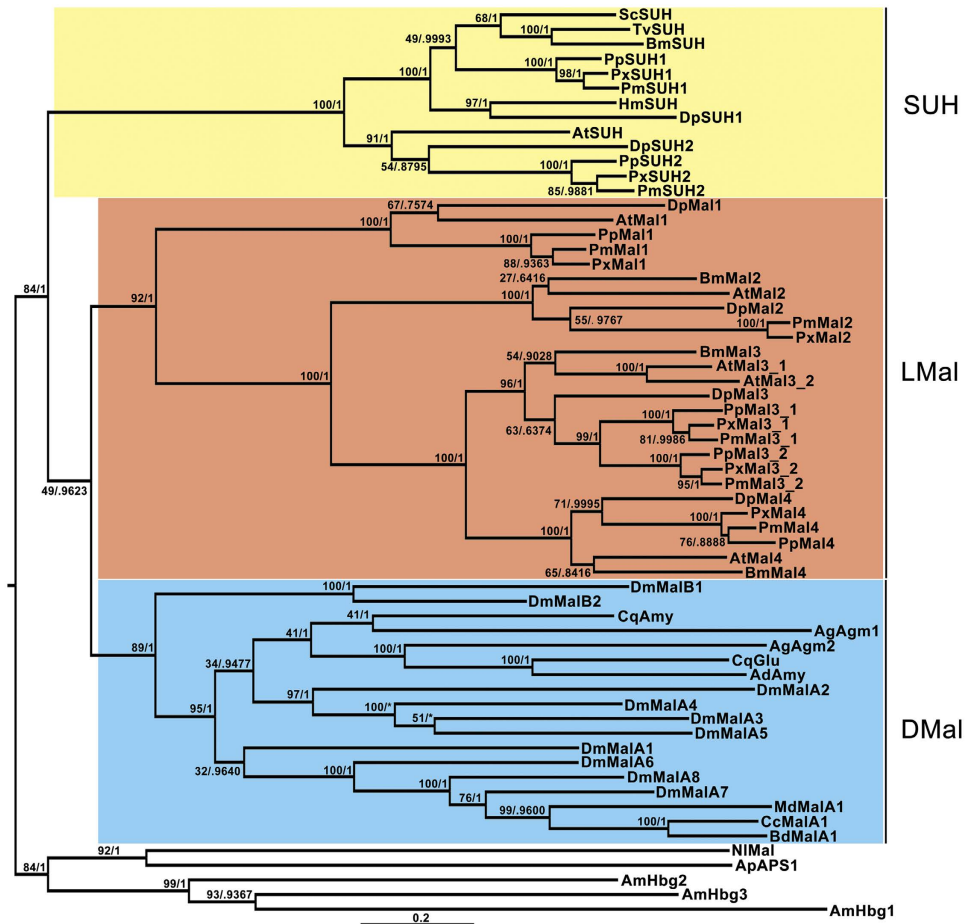


Figure 1. Reconstruction of the phylogeny of insect α -glucosidases. The ML tree depicts the evolutionary relationships among 62 sequences from species representing distinct insect lineages. Statistical supports corresponding to ML LRT and BA posterior probability are shown next to the corresponding nodes at relevant clades. Branch lengths in the tree are proportional to evolutionary distances between nodes, with the scale bar indicating the number of inferred amino acid substitutions per site. SUH, LMal and DMal are short for sucrose hydrolases, lepidopteran maltase, dipteran maltase, respectively.

are the largest. Moreover, the length of exons is conserved in *SUH* and *LMal*, and all genes harbour an exon with a length of 180 nt except *HmSUH*. Every subfamily of maltase has their own characteristics. *SUH* genes share linked exons of length 180-164 (or 158, 167)-386-202-357-239 nt except for *DpSUH2*, which lost the 239 nt exon. *LMal1* genes, which have the most conserved exon structure in maltase family, possess nine identical exons. Moreover, *BmMal2* and *DpMal2* share a very similar structure, and the length of each exon is identical except for the exon at the 5' end. The splice sites of *PxMal2* and *PmMal2* are also analogous on their chromosomes, and eight exons share a consistent length. However, *LMal1* and *LMal2* were lost in the *B. mori* and *D. plexippus*, respectively. *LMal3* is duplicated in three *Papilio* species, and their structures are conserved after gene duplication, which contain linked-exons with lengths of 180-158-582 (579)-108-151-196-138 nt. *LMal4* genes possess a linkage region with eight exons, which is 180-158-126-456-108-151-196-138 nt in length.

Compared with *LMal1* and *LMal2*, the *LMal3* and *LMal4* homologues contain fewer exons, but they gain a longer exon with length of 582 (579) and 456 nt, respectively. We found that this larger exon could be generated by a duplication-induced exon-shuffling event. In *B. mori* and *D. plexippus*, the lengths of the fourth, fifth and sixth exons of *BmMal2/DpMal2* are 126, 257 and 199 nt, but the three exons were non-existent and a novel exon with a length of 582 nt appeared in *BmMal3/DpMal3*. Interestingly, the length of this larger exon was precisely equivalent to the sum of the fourth, fifth and sixth exons. A similar exon-shuffling event was also detected in two *Papilio* species, although the length of the larger exon (582 or 579 nt) of *Mal3* was not completely consistent with the sum of three small exons (585 nt). In *LMal4*, the fifth and sixth exons of *LMal2* were reconstructed into a larger exon of 456 nt. Moreover, *LMal2*, *LMal3* and *LMal4* are tandemly arranged in lepidopteran genomes, which suggest that *LMal3* and *LMal4* were generated by gene duplication of *LMal2*, and then exon-shuffling events occurred to form the contemporary gene structures. Splice-site analyses revealed that gene structure was conserved among the *SUH* subfamily but was not well conserved in the *LMal* subfamily, which underwent changes and exon-shuffling events.

FD between SUH and maltase. Structural divergence occurred between SUH and LMal after gene duplications, and we wonder whether these two clusters underwent FD during evolution. To detect FD after gene

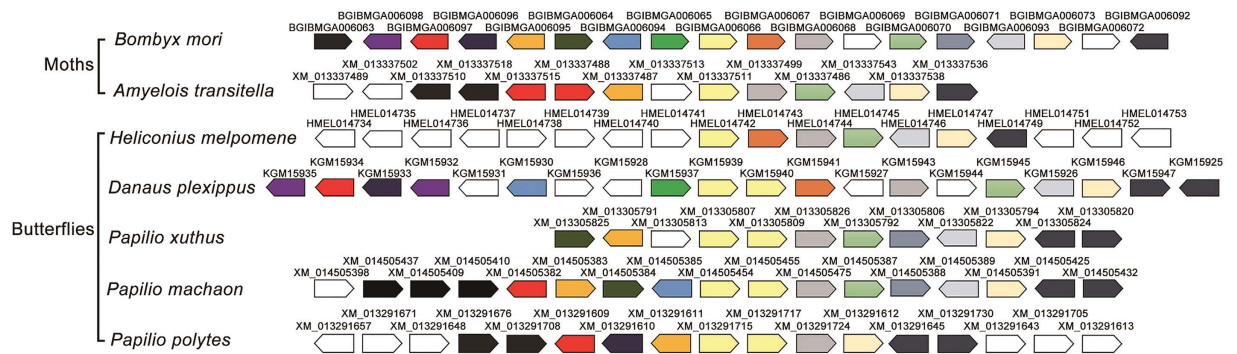


Figure 2. Conserved syntenic analysis of *SUH* and its surrounding genes. Using silkworm *BmSUH* as an anchor site, homologues of genomic genes linked to *SUH* were founded in six other lepidopteran chromosomes. *SUH*s of Lepidoptera are placed in the middle and colored in yellow. Homologues are represented by the same color, and genes with no homologue were indicated by blank boxes. The direction of genes is represented by block arrows. The positions of genes on chromosomes are not drawn to scale.

duplications, we conducted analyses of type I functional divergence (FD I) between *SUH* and other maltases using Diverge 3.0²². By comparing *SUH* and maltase (LMal+DMal), *SUH* and LMal, and *SUH* and DMal, the coefficients of FD I (θ) were 0.3232 ± 0.0577 , 0.3935 ± 0.0630 and 0.2826 ± 0.0549 , respectively (Table 1). This result indicated that functional constraint was altered significantly in *SUH*, and a remarkable FD between *SUH* and other maltases occurred. When comparing LMal versus DMal and *SUH1* versus *SUH2*, the coefficients of FD I were clearly less, with values of 0.1980 ± 0.0516 and -0.4862 ± 0.023 , respectively, which demonstrated that FD was weaker in these cases. Moreover, a total of 34, 30 and 25 critical amino acid sites, which likely are responsible for FD I, were also detected between *SUH* and LMal, *SUH* and DMal, and *SUH* and maltase, respectively. Most of the above-detected critical amino acid sites were located in the α -amy domain and these sites might be crucial to the changes in the enzyme properties and catalytic capability. The FD analyses indicated that the functions of *SUH* may be significantly changed in comparison with other maltases, which was consistent with the biochemical evidence that *SUH* is specific to sucrose hydrolysis but lost maltose digestion activity¹⁴.

Detection of positive selection for *SUH* and LMal sequences. To understand the evolutionary basis of FD between *SUH* and LMal, we estimated the rates of nonsynonymous to synonymous nucleotide substitution (d_N/d_S or ω) under different codon substitution-based evolutionary models. We employed likelihood ratio tests (LRT) with a site-specific model in the CodeML program of PAML4.8²³. Under the most basic model M0 (assuming that ω is invariable among sites and branches), the value of ω was 0.04 among the whole maltase family, which indicated that most sites represent convincing purifying selection during maltase evolution. More realistic conditions allow ω to vary among sites following a β -distribution (models M7 and M8). M8 (β and $\omega > 1$) model was a significantly better fit for the sequences in the ML tree, compared to the M7 (β) model ($2\Delta L = 20621.34$, $p < 0.001$, Table 2). The value of ω was calculated as 2.06636 for the whole α -glucosidase family (Table 2). Most amino acid residues were under purifying selection, as a total of 70 sites, mainly in the α -amy domain, were identified as subject to positive selection under M8 using Bayes empirical Bayes (BEB) analysis with posterior probabilities ≥ 0.95 (Table 2)^{24,25}. Among these sites, 251Y, a site that has been proven to determine the substrate specificity for maltose or sucrose in *A. mellifera*, was also detected under positive selection²⁶.

To test whether certain lineages in *SUH* and LMal are under positive selection, a branch-specific model implemented in CodeML of PAML 4.8 was used to explore lineage-specific variation in selection pressure. The one-ratio model (H_0) assumes a single ω for all lineages in the phylogenetic tree²⁷. When we conducted free-ratio model (H_1) analyses, which assumes a different ω parameter for each branch in the phylogenetic tree, H_1 was found to fit significantly better to the data than did H_0 ($2\Delta L = 499.04$, $p < 0.001$, Table 2), suggesting that α -glucosidases are subjected to different selective pressures²⁷. Positive selection was detected in the lineages of *SUH* ($\omega = 17.14$), *SUH2* ($\omega = 4.94$), maltase ($\omega = 9.01$), and ancestral lineage leading to LMal2, LMal3 and LMal4 ($\omega = 15.49$) (Table 2), which reflected that positive selection occurred when gene duplications happened or when new members arose in α -glucosidase family.

Given that positive selection often affects a small subset of sites along particular lineages, the branch-site model was then used to detect positive selection in individual codons along *SUH* or maltase lineages, allowing ω to vary among both sites and branches²⁸ (Table 2). Ratios of ω in different *SUH* and maltase lineages were varied, and lineages of *SUH1*, *SUH2*, LMal, LMal1, LMal3, LMal4 and ancestral lineages leading to *SUH* and LMal/DMal, LMal2 and LMal3/4, and LMal3 and LMal4 were significantly under positive selection. Moreover, some amino acid sites, which underwent positive selection, were also detected. We detected 16 and 4 sites subjected to significant positive selection from the entire *SUH* lineage (not significant) and *SUH2* lineage, respectively. To the maltase of Lepidoptera, one, eight, one and six sites under positive selection were found in the lineages of LMal, LMal1, LMal2 and LMal4, respectively. In addition, another three or four sites were identified to bear positive selection when the divergence occurred between LMal2 and LMal3/4 or LMal3 and LMal4, respectively. Moreover, two additional sites were also detected to be under positive selection in the ancestral lineage, leading to *SUH* and maltase, and no site was detected under positive selection in the branches of *SUH1* and LMal3. Among

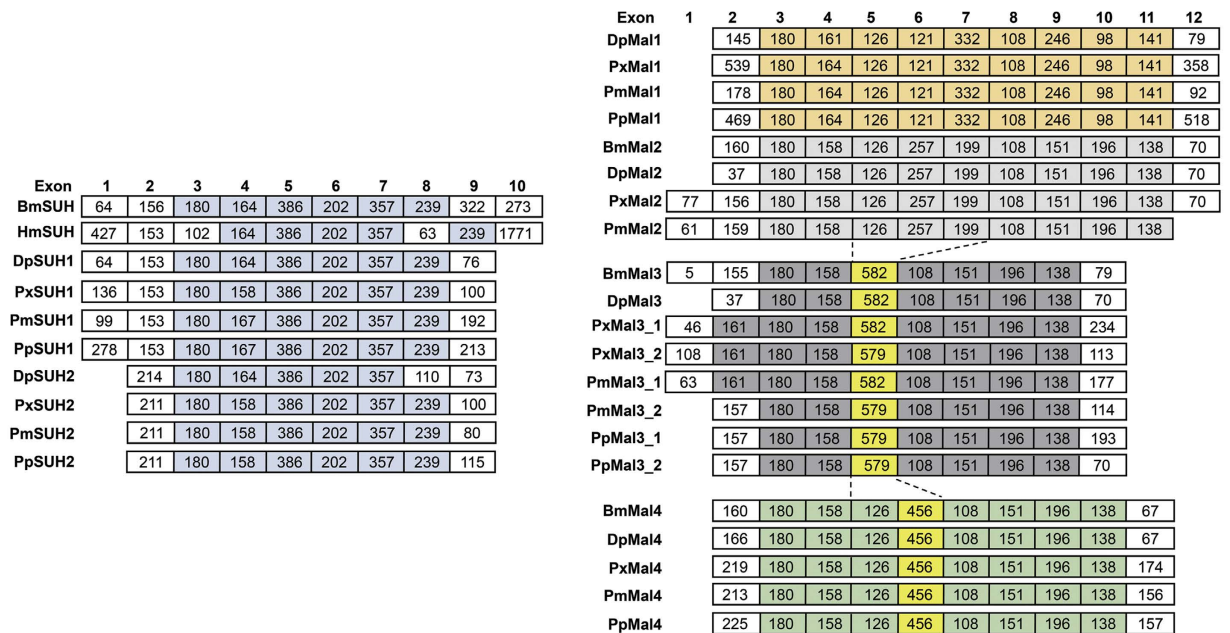


Figure 4. Exon/intron structures of *SUH* and *LMal* genes. The length of each exon is represented by the number in the box. Highly similar exon regions among each subfamily are indicated by the same color, and exons that may be generated by exon-shuffling are colored in yellow. Exon sizes are not drawn to scale.

FD	Subfamilies	Coefficient $\theta \pm SE$ (P)	Critical Amino Acid sites
Type I FD	SUH vs. LMal	0.3232 ± 0.0577 ($P < 0.01$)	101, 137, 164, 180 , <u>212</u> , 230, 232, 237, 250, 269, 279 , 281 , 284 , 288 , 292 , 296, 300, 307, 325, 326 , 329, 330, 331, 332 , 345, 350 , 360, 366 , 368, 369 , 397, 398, 415, 503
	SUH vs. DMal	0.3935 ± 0.0630 ($P < 0.01$)	141, 150, 154 , 161, 165, 191 , <u>212</u> , 222* , 237, 269 , 281 , 295, 329, 330, 332, 345, 350 , 360, 362* , 363, 364, 365, 366* , 370* , 373*, 374, 405, 415, 458, 468
	LMal vs. DMal	0.1980 ± 0.0516 ($P < 0.01$)	148, 150* , 165, 229, 247, 265, 321, 362* , 365*, 369* , 370* , 373, 374, 398, 402, 454, 483
	SUH vs. Mal	0.2826 ± 0.0549 ($P < 0.01$)	101, 154 , 161, <u>212</u> , 222 , 237, 269, 281 , 284 , 295, 296, 325, 326 , 329, 330, 331, 332 , 345, 350* , 360, 362 , 364, 366 , 368, 373, 374, 415, 458
	SUH1 vs. SUH2	-0.4862 ± 0.023	287*, 380* , 393*, 496*

Table 1. Type I functional divergence (FD) of α -glucosidase family of insect. Functional divergences (Coefficient θ) for pairwise comparisons within the α -glucosidase family of insect are shown as value \pm standard error. Critical amino acid sites detected as relating to FD with $P > 70\%$ ($>90\%$, indicated with asterisks) are listed. Numbering refers to the positions in the alignments of protein sequences generated by MAFFT alignment. Residues that also under positive selection and relative to ligand-binding are presented by bold and underline, respectively.

of α -helix and β -strand, respectively, whereas 3% of this protein is made up of transmembrane helix. Moreover, the 3D modelling showed that BmSUH contains three domains (Domains A, B and C), which are similar to other α -glucosidases (Fig. 5A).

The ligand-binding sites are important in determining the interaction between protein and its ligand, and the 3DLigandSite web server was used to predict potential binding sites³⁰. A total of 16 amino acid sites were identified to be crucial for substrate binding (Fig. 5A). For the maltase family, an active site cleft usually exists between Domains A and B, and a triad of catalytic residues (Asp, Glu and Asp) are responsible for the catalytic reaction³¹. The result showed that 16 sites form a binding pocket to the substrate, and three catalytic residues are included in them. The 212A site is one of the potential binding sites. However, this site was also detected to be under positive selection by site-specific model that was responsible for the FD of SUH and maltase. Moreover, 191Q, another potential binding site, was also identified to be subjected to the positive force in LMal1 and LMal4 by branch-site model. These amino acid sites might be responsible for the functional differentiation and specific evolutionary adaptations during the evolution of α -glucosidase family in Lepidoptera. When the sites involved in both positive

Model	Foreground branch	-lnL	2lnL	P level	Parameter Estimates	Positive sites
Site Model						
M7		39876.24			p=0.67039, q=12.21042	not allowed
M8		50195.31	20621.34	<0.01	p0=0.99999, p=0.36811, q=1.82680 (p1=0.00001), ω=2.06636	97E**, 154I**, 158A**, 159R*, 180G**, 181V**, 202K**, <u>212A**</u> , 213I**, 224A**, 234K**, 251Y**, 263R**, 272L**, 275F**, 277S**, 280L**, 281G** , 283T**, 284I** , 291L**, 309N**, 310K*, 311N**, 326N** , 327V**, 328S**, 332L** , 340A**, 341I**, 350D* , 355L*, 357S**, 358K**, 362R** , 366Y** , 367I**, 370R** , 377Y**, 379G**, 380I** , 391N*, 399H**, 400D**, 409N**, 411L**, 427R*, 429G, 466N**, 467S**, 468T** , 476T**, 477N**, 484E**, 487Q**, 488E**, 489I**, 495K**, 496E** , 497T**, 499R**, 507A**, 510K**, 513K**, 585E**, 587T**, 588S**, 589S**, 590Q**, 591L**
Branch-specific model						
M0		41427.55			ω=0.04	not allowed
Free-ratio model		41178.03	499.04	<0.01	ω _{suh} =17.14, ω _{suh1} =0.10, ω _{suh2} =4.94, ω _{LMal1} =0, ω _{LMal2} =0.14, ω _{LMal3} =0.18, ω _{LMal4} =0.88, ω _{LMal} =9.01, ω _{LMal234} =15.49, ω _{LMal34} =0.39	not allowed
Branch-site model						
Ma0	SUH1	41314.68	4.24	<0.05	ω0=0.04, ω1=1.00, ω2=1.00	not allowed
Ma		41312.56			ω0=0.04, ω1=1.00, ω2=999.00	none
Ma0	SUH2	41320.28	17.24	<0.01	ω0=0.04, ω1=1.00, ω2=1.00	not allowed
Ma		41311.66			ω0=0.04, ω1=1.00, ω2=21.51	184 P**, 195 S*, 222 N*, 288 T*
Ma0	Ancestral SUH	41305.76	0.96	>0.05	ω0=0.04, ω1=1.00, ω2=1.00	not allowed
Ma		41305.28			ω0=0.04, ω1=1.00, ω2=2.51	108S**, 150S** , 152Y**, 158A*, 181V*, 186S**, 191Q** , 279Q* , 280L*, 283T**, 292I** , 294L**, 357S*, 369L* , 407I**, 409N*
Ma0	Ancestral SUH and Mal	41313.43	13.67	<0.01	ω0=0.04, ω1=1.00, ω2=1.00	not allowed
Ma		41306.60			ω0=0.04, ω1=1.00, ω2=999.00	162G**, 483A*
Ma0	Ancestral LMal	41320.28	36.48	<0.01	ω0=0.04, ω1=1.00, ω2=1.00	not allowed
Ma		41302.04			ω0=0.04, ω1=1.00, ω2=999.00	199W*
Ma0	LMal1	41306.11	6.40	<0.05	ω0=0.04, ω1=1.00, ω2=1.00	not allowed
Ma		41302.91			ω0=0.04, ω1=1.00, ω2=10.07	181V*, 191Q* , 224A*, 226K*, 251Y**, 276E*, 366Y* , 588S*
Ma0	LMal2	41313.39	0.96	>0.05	ω0=0.04, ω1=1.00, ω2=1.00	not allowed
Ma		41312.91			ω0=0.04, ω1=1.00, ω2=2.14	341I*
Ma0	LMal3	41311.50	9.46	<0.01	ω0=0.04, ω1=1.00, ω2=1.00	not allowed
Ma		41306.77			ω0=0.04, ω1=1.00, ω2=999.00	None
Ma0	LMal4	41300.05	14.14	<0.01	ω0=0.04, ω1=1.00, ω2=1.00	not allowed
Ma		41292.98			ω0=0.04, ω1=1.00, ω2=999.00	147S**, 191Q* , 279Q* , 366Y* , 400D*, 585E*
Ma0	Ancestral LMal2, LMal3 and LMal4	41320.28	16.50	<0.01	ω0=0.04, ω1=1.00, ω2=1.00	not allowed
Ma		41312.03			ω0=0.04, ω1=1.00, ω2=119.75	161P* , 420Q*, 479S*
Ma0	Ancestral LMal3 LMal4	41320.05	9.56	<0.01	ω0=0.04, ω1=1.00, ω2=1.00	not allowed
Ma		41315.27			ω0=0.04, ω1=1.00, ω2=54.58	156S*, 203R**, 362R* , 371W**

Table 2. Tests of positive selection on Lepidopteran α-glucosidase family with site-specific, branch-specific and branch-site models. The ω represents for d_N/d_S . * Significant at $p < 0.05$, ** Significant at $p < 0.01$. The site number was mapped to BmSUH after alignments. 2lnL, log-likelihood difference between compared models. Amino acid residues that also involved in FD I and ligand-binding were presented by bold and underline, respectively.

selection and FD were mapped to the structure of BmSUH, we found that these nine sites were mainly on the α-helix of the molecular surface, and were precisely located around the ligand-binding groove (Fig. 5B). This result indicated that specific ligand-binding sites would not be major targets for adaptive changes in the SUH family. However, the ordered distribution of sites, which involved both positive selection and FD, reflected that they could have effects on the discrepancy of ligand-binding and conformational stability.

Discussion

The lepidopteran digestive system is characterised by two derived features, including developing extremely alkaline midguts and losing the midgut caeca³². These characters reflect the divergent selective pressure may have been imposed on the evolution of lepidopteran digestive system. In the present study, we found that the SUH subfamily of α-glucosidase is only detected in Lepidoptera, which showed high-alkaline adaptability. SUH and other LMal were diverged by ancestral gene expansion events, and their functions showed differentiation in subsequent

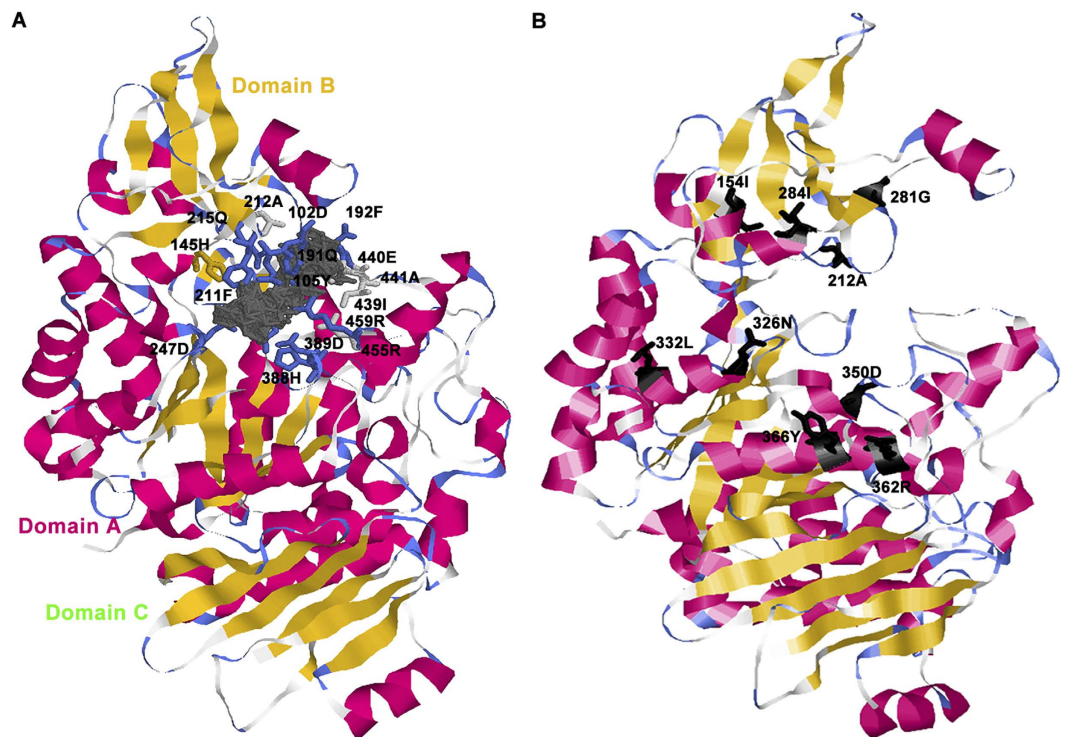


Figure 5. 3D architecture of BmSUH showing positive selection and functional divergence residues. (A) Tertiary structure of BmSUH that binds to sucrose (complex colored in grey). Critical sites that predicted to be involved in ligand binding are mapped onto the structure and are represented as stick model. (B) Nine sites both contributed to positive selection and FD I are mapped onto the tertiary structure of BmSUH with black sticks. α -Helices, β -sheets and turns are shown in magenta, yellow and pale blue, respectively. All other residues are shown in white.

evolutionary process. This differentiation may be caused by various selection pressures, which are exerted in different subfamilies. Adaptive selection pressure led to the exclusive presence of SUH1 in the highly alkaline digestive tract of Lepidoptera. Moreover, nine sites subjected to both positive selection and FD were located around the ligand-binding groove. The sites may contribute to the catalytic specificity to substrates and the stability of molecular conformation. The emergence of *SUH* and its subsequent duplications reflect effective adaptations to the specific diets and digestive environment of Lepidoptera³².

The exon structures of *SUH* are conserved in Lepidoptera, but they significantly changed compared with *LMal* (Fig. 4). Unlike *LMal*, *SUH* possessed an N-terminal hydrophobic amino acid sequence (except *AtSUH*), which could potentially function as a membrane association region, explaining why *SUH*s are associated with membrane¹⁴. The gene structures appear to have great variety between *SUH* and *LMal* after the ancestral gene differentiation event ages ago, which may contribute to the functional diversification and differences in membrane-spanning domains and substrate specificity. The *LMal* subfamily, underwent a more complicated evolutionary process, with at least three rounds of gene duplication, whereas *SUH* was only duplicated once (Fig. 6). In addition, *Mal1* is lost in *B. mori*, which may be an outcome of gene deletion or genome rearrangement. Recent works have started providing strong evidence for the functional diversification of α -glucosidase in Diptera and Hymenoptera, such as maltose hydrolysis (*Agm1* and *Agm2* of *A. gambiae*)³³, sucrose degradation (*HBG1* and *HBG3* of *A. mellifera*), a receptor for Bin toxin (*Cpm1* of *A. gambiae*)³⁴, and heteromeric amino acid transporters (*hcHATs* proteins)³⁵. Compared with Diptera species, the gene expansion of lepidopteran α -glucosidase is much simpler, as *Drosophila* experienced eight rounds of duplications and developed ten α -glucosidase genes¹⁶.

Synteny conservation analysis was performed to confirm the results of the phylogenetic analysis. *SUH* and its surrounding genes were tandemly arrayed in lepidopteran genomes (Fig. 2), but *SUH2* only emerged near *SUH* in several butterfly species^{36,37}. Moreover, RT-PCR analysis showed a weaker expression of *PxSUH2* than *PxSUH1* in the midgut of *Papilio* species (Supplementary Fig. S2). *SUH2* may be generated by the duplication of *SUH1*, and this duplication event occurred in a few species not long ago. The emergence of *SUH2* reflects deep adaptation to the dietary habit or digestive needs of butterfly species, and further biochemical characterisation of *SUH2* will be of great interest. *LMal2*, *LMal3* and *LMal4* were also tandemly arrayed on the chromosome, but *LMal1* was located in separate chromosomal regions. This distribution model was similar to that of some Diptera species, as their α -glucosidase family is also located in two or three chromosomal regions¹⁵. In addition, gene structure analyses indicate that *LMal3* and *LMal4* were generated by exon-shuffling of *LMal2*. In this way to generate new genes was first observed during the evolution of α -glucosidase (Fig. 4). This result demonstrates that gene duplication and exon-shuffling contribute much to the *maltase* gene family expansion in Lepidoptera.

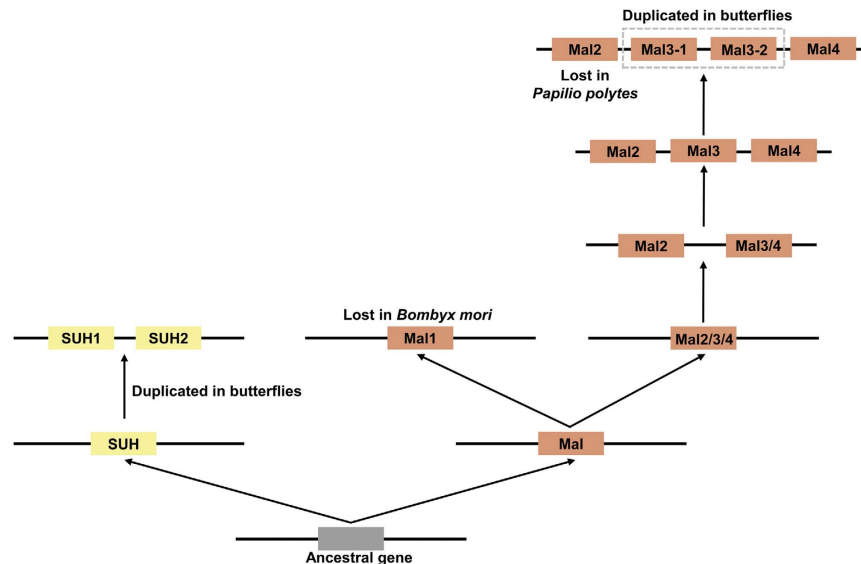


Figure 6. A proposed scenario of evolution and gene duplications of α -glucosidases in Lepidoptera.

SUH and *LMal* were descended from the common ancestor, then went through genes gain and loss events in Lepidoptera. Members from *SUH* and *LMal* are distinguished by different colors.

The subsequent divergence after gene duplication plays an important role in the evolution of novel gene function³⁸. Many residues, including several functionally determined sites (212A, 251Y), were detected to be under positive selection, and key residues affected by diversified natural selection may result in the functional changes. 251Y/H has been previously found to be important in substrate preferences for sucrose or maltose²⁶. This residue differs in *SUH1* and *SUH2*, as *SUH1* mainly harbours Y, whereas *SUH2* harbours H. Interestingly, the residues corresponding to 251Y are in conserved sequence region II of the GH-13 enzyme. Region II has been noted as a determinant of the substrate specificity of GH-13 enzymes²⁶. This result opens exciting avenues for future research where functional changes are caused by Y251H substitution in region II. In addition, strong signals of positive selection were detected during ancestral *SUH* divergence and in the *SUH2* lineage, suggesting that the *SUH* subfamily has evolved an enhanced ability for sucrose digestion in response to the Lepidoptera-specific feeding habits and gastrointestinal circumstance. The evolution of the *SUH* subfamily was concordant with the theory that random mutations were fixed in one daughter gene under relaxed purifying selection, which occurred by the reduced functional constraint provided by genetic redundancy^{39,40}. Compared with the ancestral gene, *SUH2* showed a weaker expression, and may undergo neofunctionalisation or subfunctionalisation during evolution. Moreover, four sites of *SUH2*, which were detected under positive selection by the branch-site model (Table 2), might contribute to the functional change. For *LMal*, positive selections had an effect on the leading branch of the whole *LMal* and ancestral branch of *LMal2* and *LMal3&4*, but not on individual lineages of *LMal* immediately after gene divergence (Table 2). This result suggested that diversifying selection only acted upon the process of *LMal* gene divergence, but not on novel genes after duplication. Moreover, many positively selected sites in the core domain were detected from the whole α -glucosidase family of insect by the site-specific model, which indicated that the α -glucosidase family underwent a changeable evolutionary course. The α -glucosidase family should have been adaptively modified to recognise and bind different substrates and ensure the digestibility of varied diets.

If positive selection largely influenced the evolution of *LMal* and *SUH*, then how many changes occurred in the functions of these genes? To answer this question, we measured the FD I and critical sites involved in it by Diverge3 software, which demonstrated that altered functional constraints may occur after duplication, when *SUH* was compared with maltase, *LMal* or *DMal*. However, it suggested a functional constraint between *SUH1* and *SUH2* (Table 1). Critical amino acid residues, which may contribute to FD, were also detected, and all these sites were located in the α -amy domain when compared *SUH* with maltase (Table 1). Our results are consistent with previous studies, which have shown that *BmSUH* had substrate transformation to sucrose, unlike conventional maltases with maltose specificity¹⁴.

Although the structures of sucrose complexes with acid-base mutants of the GH13 enzymes have been examined, no 3D structures of the enzyme proteins in a complex with sucrose have yet been determined²⁶. In this study, we predicted the tertiary structure and sucrose binding sites of *BmSUH* using Phyre2 and 3DLigandSite software^{29,30}. Our predicted result was concordant with the estimation of Seddigh, who conducted homology modelling of α -glucosidase, such as Dm-NP610382 (*D. melanogaster*), Am-XP006560868 (*A. mellifera*), At-NP196733 (*A. thaliana*), Hs-NP937784 (*H. sapiens*) and Mt-YP007966392 (*M. tuberculosis*)⁴¹. This similarity of tertiary structure prediction analysis indicated that the 3D structures of α -glucosidase are conserved during evolution. Moreover, *BmSUH* harbours the potential 16 binding sites and forms a substrate-binding groove to bind and catalyse sucrose (Fig. 5A). When mapping nine sites, which were detected in both site-specific model analyses and FD analyses of *SUH* versus maltase, onto the modelled protein structure of *BmSUH*, we found that these

sites were mainly located around the substrate-binding groove in the α -helices of Domain A (Fig. 5B). These sites might help to stabilise the protein conformation and assist ligand binding. Among the nine sites, the 212A site was identified to involve FD and positive selection, and also a site that is predicted to participate in the sucrose-binding reaction. Therefore, the nine sites, especially 212A, could be inferred as key sites during the evolution and functional formation of SUH, and they contributed to the recognition mechanism of substrate specificity. We propose these residues as targets for further experimental study of SUH functions. Daimon had proven that a β -fructofuranosidase (SUC), which is originally known as an ‘anomalous’ enzyme that had been believed to be absent in the animal kingdom, serves as a sucrose-digesting enzyme in the silkworm physiology¹³. Moreover, previous studies have shown that organisms, which access sucrose as a major food source, can acquire invertases from bacteria via horizontal gene transfer (HGT) to ensure the efficient utilisation of sucrose, such as plant-parasitic nematodes⁴². Recent genome sequencing projects have shown that SUC and SUH are present in lepidopteran insects, suggesting that Lepidoptera has evolved an enhance ability of digesting sucrose. The evolution of SUH, as a specific sucrose hydrolysis enzyme, reflects that lepidopteran insects can adapt to specific environments and diets by altering their original physiological characteristics.

Materials and Methods

Sequences collection and phylogenetic analyses. A comprehensive search by BLASTp and PSI-BLAST were performed in NCBI, Ensembl and FlyBase using DmMal1A and BmSUH as the query sequences⁴³. After removing the partial sequences and redundant sequences, the final data set included 62 complete maltase and SUH sequences (Supplementary Table S1). All sequences were revised for errors in accession numbers and nomenclature. Multiple sequence alignments of these sequences were generated with MAFFT software⁴⁴. According to the Akaike Information Criterion (AIC) for small sample size, MrModelTest2.3 revealed General Time Reversible model incorporating invariant sites and a gamma distribution (GTR+ I+ G) as the best model of molecular evolution with the best fit to our data⁴⁵. Maximum-likelihood (ML) tree was reconstructed with RAxML-HPC BlackBox (8.2.8) on the CIPRES web portal (<https://www.phylo.org/portal2>) based on the GTR+ I+ G model^{46,47}.

The Bayesian analyses were carried out using Markov chain Monte Carlo (MCMC) sampling in MrBayes3.2.1 with the same model described above, and data sets ran for 300,000 generations until they reached congruence⁴⁸. The Bayesian tree was sampled every 100 generations, and the first 25% of the trees were discarded as burnin. Phylogenetic trees were visualized with FigTree 1.4.2.

Estimation of evolutionary divergence times. To obtain temporal information on the divergence events, we implemented two methods to conduct molecular dating analyses. Frist, Reltime method of MEGA7 was used to infer the time tree by ML approach based on the GTR+ I+ G model. This method allows rates to vary from branch to branch without pre-specification of statistical distribution of lineage rates^{19,20}.

Second, we estimated divergence times using Bayesian approach implemented in BEAST 1.83 with a relaxed molecular clock, which is determined by likelihood ratio test (LRT) of the molecular clock hypothesis ($P < 0.01$)^{49,50}. Uncorrelated lognormal relaxed clock was chosen to estimate the evolutionary rate variations, and Yule speciation process was employed to model tree prior⁵¹. We set the number of generations to 10,000,000 with 10% burnin in MCMC analyses. Moreover, the maximum clade credibility (MCC) chronogram was summarized by TreeAnatorator with posterior probability limit to 0.5. Two calibration constraints, divergence times of DmMal2-DmMal345 (84 MYA) and DmMalB1-DmMalB2 (155 MYA)¹⁶, were applied to date the divergence times of internal nodes within the phylogenetic tree. These analyses involved 62 nucleotide sequences described above.

Expression analysis of SUH genes in *P. xuthus* by RT-PCR. Total RNA from the 3rd day of the fifth instar larvae of *P. xuthus* was used in the RT-PCR analysis. One microgram of total RNA was used to synthesize first-strand cDNA using PrimeScript RT reagent Kit with gDNA Eraser (Takara) according to the manufacturer's instructions. The data were normalized by determination of the amount of gene encoding *ribosomal protein (rpl)* in each sample to eliminate variations in mRNA and cDNA quality and quantity. Gene-specific primers were deposited in Supplementary Table S2.

Conserved synteny analyses. The syntenic relationship of SUH and its up- and downstream genes on lepidopteran genomes were revealed by the Genomics 30.01⁵² from Ensembl 31 database with BmSUH as the query gene. For genomes that not available on Ensembl, we searched genes around its corresponding orthologue of BmSUH from NCBI genome database manually⁵³, and checked the result by reciprocal BLAST.

Splice site and gene structure analyses. The Ensembl Metazoa genome browser release 31 and NCBI database were used to infer the exon boundaries of the coding regions of SUH and LMal genes. The accurate length (nt) of every exon was also determined.

Analyses of type I functional divergence. Type I FD represents amino acid patterns that are highly conserved in one duplicate cluster but shows great variation in the other, which resulted in altered selective constraints between duplicated genes. The DIVERGE version 3.0 software was employed to test Type I FD after gene duplication⁵⁴. The coefficient of FD (θ) is an indicator of the level of type I FD among two homologous gene clusters. The posterior probabilities (Q_k) were also estimated to indicate amino acid sites to be responsible for FD. A value of $Q_k > 0.7$ was chosen as a cutoff to measure the degree of FD at the amino acid level, and $Q_k > 0.9$, which marked with an asterisk, was significant.

Detection of positive selection. To measure the strength and mode of natural selection during the evolution of SUH and LMal gene subfamilies, the ratio of non-synonymous (d_N) to synonymous substitutions (d_S)

($\omega = d_n/d_s$) was calculated by the CodeML program implemented in the PAML 4.8 package²³. The phylogenetic tree was built by the ML method described above, and the alignment of sequences was achieved by MAFFT software. They were used to conduct CodeML analyses.

We employed three model, site-specific model, branch-specific model and branch-site model, to detect relative positive forces during the evolution of SUH and LMal. In the site-specific model, the M7 (β model) and M8 (β and $\omega > 1$ model) were compared to identify the sites which under positive selection. The M7 model uses the flexible β distribution to indicate the difference of ω (value from 0 to 1) among different sites, whereas M8 allows several amino acid sites be under positive selection ($\omega > 1$)⁵⁵. If M8 fits the data of SUH and maltase better than M7, and detects sites under positive selection, we could accept the assumption of certain sites subjected to positive selection. In the branch-specific model, the significance of variation in the ω for each lineage was examined by free-ratio model²⁵. If free-ratio model fits significantly better than the one-ratio model, which assigns a constant ω for all of branches in the tree, the alternative model could be accepted. Branch-site model, in which ω can vary in particular sites along particular branches, was used to test whether only several sites were under positive selection along foreground branch²⁸. The null model, which caps $\omega = 1$, was served to compare with the former model in the significance of fitness to the data. The likelihood ratio test (LRT) was used to test whether alternative model better fitted our data than null model significantly^{25,56}.

Protein structure and ligand-binding prediction. The Phyre2 structure prediction server (<http://www.sbg.bio.ic.ac.uk/phyre2>) was used to conduct tertiary structure prediction and alignment²⁹. Phyre2 uses the alignment of hidden Markov models for homology-based protein modelling. It also incorporates the *ab initio* folding simulation to model regions with no detectable homology to known structures. Moreover, FD sites and positive selected sites were mapped to the predicted model. Predicted tertiary structures and relevant sites were visualized by RasMol 2.7.5⁵⁷. The prediction of ligand-binding sites was employed with 3DLigandSite server, which based upon the modelled structure of query protein (<http://www.sbg.bio.ic.ac.uk/3dligandsite>)³⁰.

References

- Capinera, J. L. *Encyclopedia of entomology*. Vol. 4 (Springer Science & Business Media, 2008).
- Terra, W. R. & Ferreira, C. *11-Biochemistry and Molecular Biology of Digestion* (2012).
- Berenbaum, M. Adaptive Significance of Midgut pH in Larval Lepidoptera. *The American Naturalist* **115**, 138–146 (1980).
- Van Wielendaele, P., Badisco, L. & Vanden Broeck, J. Neuropeptidergic regulation of reproduction in insects. *Gen Comp Endocrinol* **188**, 23–34 (2013).
- Pytelková, J. *et al.* Digestive α -amylases of the flour moth *Ephesia kuehniella*—adaptation to alkaline environment and plant inhibitors. *FEBS Journal* **276**, 3531–3546 (2009).
- Brune, A. & Kuhl, M. pH profiles of the extremely alkaline hindguts of soil-feeding termites (Isoptera: Termitidae) determined with microelectrodes. *Journal of Insect Physiology* **42**, 1121–1127 (1996).
- Harrison, J. F. Insect acid-base physiology. *Annual Review of Entomology* **46**, 221–250 (2001).
- Clark, T. M. Evolution and Adaptive Significance of Larval Midgut Alkalinization in the Insect Superorder Mecoptera. *Journal of Chemical Ecology* **25**, 1945–1960 (1999).
- Appel, H. M. & Martin, M. M. Gut redox conditions in herbivorous lepidopteran larvae. *Journal of Chemical Ecology* **16**, 3277–3290 (1990).
- Motoyuki, S., Yuan, X. L. & Fujiyoshi, M. Sucrase activity and its kinetic properties in peritrophic membrane, and in membrane-bound and soluble fractions of midgut in the silkworm, *Bombyx mori*. *Comparative Biochemistry and Physiology Part A: Physiology* **108**, 255–264 (1994).
- Terra, W. R. & Ferreira, C. Insect digestive enzymes: properties, compartmentalization and function. *Comparative Biochemistry and Physiology B* **109**, 1–62 (1994).
- Sumida, M., Yuan, X. L. & Matsubara, F. Purification and some properties of soluble β -fructofuranosidase from larval midgut of the silkworm, *Bombyx mori*. *Comparative Biochemistry and Physiology B* **107**, 273–284 (1994).
- Daimon, T. *et al.* Beta-fructofuranosidase genes of the silkworm, *Bombyx mori*: insights into enzymatic adaptation of *B. mori* to toxic alkaloids in mulberry latex. *J Biol Chem* **283**, 15271–15279 (2008).
- Wang, H., Kiuchi, T., Katsuma, S. & Shimada, T. A novel sucrose hydrolase from the bombycid silkworms *Bombyx mori*, *Trilocho varians*, and *Samia cynthia ricini* with a substrate specificity for sucrose. *Insect Biochem Mol Biol* **61**, 46–52 (2015).
- Gabrisko, M. Evolutionary history of eukaryotic alpha-glucosidases from the alpha-amylase family. *J Mol Evol* **76**, 129–145 (2013).
- Gabrisko, M. & Janecek, S. Characterization of maltase clusters in the genus *Drosophila*. *J Mol Evol* **72**, 104–118 (2011).
- Zhang, Q., Hua, G., Bayyareddy, K. & Adang, M. J. Analyses of α -amylase and α -glucosidase in the malaria vector mosquito, *Anopheles gambiae*, as receptors of Cry11Ba toxin of *Bacillus thuringiensis* subsp. *jegathesan*. *Insect Biochemistry and Molecular Biology* **43**, 907–915 (2013).
- Ferreira, L. M., Romao, T. P., de-Melo-Neto, O. P. & Silva-Filha, M. H. The orthologue to the Cpm1/Cqm1 receptor in *Aedes aegypti* is expressed as a midgut GPI-anchored alpha-glucosidase, which does not bind to the insecticidal binary toxin. *Insect Biochem Mol Biol* **40**, 604–610 (2010).
- Tamura, K. *et al.* Estimating divergence times in large molecular phylogenies. *Proc Natl Acad Sci USA* **109**, 19333–19338 (2012).
- Kumar, S., Stecher, G. & Tamura, K. MEGA7: Molecular Evolutionary Genetics Analysis version 7.0 for bigger datasets. *Mol Biol Evol* (2016).
- Xu, G., Guo, C., Shan, H. & Kong, H. Divergence of duplicate genes in exon-intron structure. *Proc Natl Acad Sci USA* **109**, 1187–1192 (2012).
- Gu, X. *et al.* An update of DIVERGE software for functional divergence analysis of protein family. *Molecular Biology & Evolution* **30**, 1713–1719 (2013).
- Yang, Z. PAML 4: phylogenetic analysis by maximum likelihood. *Mol Biol Evol* **24**, 1586–1591 (2007).
- Yang, Z. & Nielsen, R. Estimating synonymous and nonsynonymous substitution rates under realistic evolutionary models. *Molecular biology and evolution* **17**, 32–43 (2000).
- Yang, Z. Likelihood ratio tests for detecting positive selection and application to primate lysozyme evolution. *Molecular biology and evolution* **15**, 568–573 (1998).
- Ngiwsara, L. *et al.* Amino acids in conserved region II are crucial to substrate specificity, reaction velocity, and regioselectivity in the transglucosylation of honeybee GH-13 alpha-glucosidases. *Biosci Biotechnol Biochem* **76**, 1967–1974 (2012).
- Yang, Z. & Nielsen, R. Codon-substitution models for detecting molecular adaptation at individual sites along specific lineages. *Molecular biology and evolution* **19**, 908–917 (2002).

28. Zhang, J., Nielsen, R. & Yang, Z. Evaluation of an improved branch-site likelihood method for detecting positive selection at the molecular level. *Mol Biol Evol* **22**, 2472–2479 (2005).
29. Kelley, L. A., Mezulis, S., Yates, C. M., Wass, M. N. & Sternberg, M. J. The Phyre2 web portal for protein modeling, prediction and analysis. *Nat Protoc* **10**, 845–858 (2015).
30. Wass, M. N., Kelley, L. A. & Sternberg, M. J. 3DLigandSite: predicting ligand-binding sites using similar structures. *Nucleic Acids Res* **38**, W469–473 (2010).
31. Watanabe, K., Hata, Y., Kizaki, H., Katsube, Y. & Suzuki, Y. The refined crystal structure of *Bacillus cereus* oligo-1,6-glucosidase at 2.0 Å resolution: structural characterization of proline-substitution sites for protein thermostabilization 1. *Journal of Molecular Biology* **269**, 142–153 (1997).
32. Terra, W. R. Evolution of digestive systems of insects. *Annual review of entomology* **35**, 181–200 (1990).
33. Zheng, L., Whang, L. H., Kumar, V. & Kafatos, F. C. Two genes encoding midgut-specific maltase-like polypeptides from *Anopheles gambiae*. *Experimental parasitology* **81**, 272–283 (1995).
34. Opota, O., Charles, J.-F., Warot, S., Pauron, D. & Darboux, I. Identification and characterization of the receptor for the *Bacillus sphaericus* binary toxin in the malaria vector mosquito, *Anopheles gambiae*. *Comparative Biochemistry and Physiology Part B: Biochemistry and Molecular Biology* **149**, 419–427 (2008).
35. Gabriško, M. & Janeček, S. Looking for the ancestry of the heavy-chain subunits of heteromeric amino acid transporters rBAT and 4F2hc within the GH13 α -amylase family. *FEBS journal* **276**, 7265–7278 (2009).
36. Li, X. *et al.* Outbred genome sequencing and CRISPR/Cas9 gene editing in butterflies. *Nat Commun* **6**, 8212 (2015).
37. Nishikawa, H. *et al.* A genetic mechanism for female-limited Batesian mimicry in *Papilio* butterfly. *Nature Genetics* **47**, 405–409 (2015).
38. Innan, H. & Kondrashov, F. The evolution of gene duplications: classifying and distinguishing between models. *Nat Rev Genet* **11**, 97–108 (2010).
39. Zhang, J. Evolution by gene duplication: an update. *Trends in Ecology & Evolution* **18**, 292–298 (2003).
40. Zhang, J., Rosenberg, H. F. & Nei, M. Positive Darwinian selection after gene duplication in primate ribonuclease genes. *Proceedings of the National Academy of Sciences* **95**, 3708–3713 (1998).
41. Seddigh, S. & Darabi, M. Structural and phylogenetic analysis of α -glucosidase protein in insects. *Biologia* **70** (2015).
42. Danchin, E. G., Guzeeva, E. A., Mantelin, S., Berepiki, A. & Jones, J. T. Horizontal Gene Transfer from Bacteria Has Enabled the Plant-Parasitic Nematode *Globodera pallida* to Feed on Host-Derived Sucrose. *Mol Biol Evol* **33**, 1571–1579 (2016).
43. Altschul, S. F. *et al.* Gapped BLAST and PSI-BLAST: a new generation of protein database search programs. *Nucleic acids research* **25**, 3389–3402 (1997).
44. Katoh, K., Kuma, K.-i., Toh, H. & Miyata, T. MAFFT version 5: improvement in accuracy of multiple sequence alignment. *Nucleic acids research* **33**, 511–518 (2005).
45. Nylander, J. MrModeltest v2. Program distributed by the author. *Evolutionary Biology Centre, Uppsala University* **2** (2004).
46. Miller, M. A., Pfeiffer, W. & Schwartz, T. In *Proceedings of the 2011 TeraGrid Conference: extreme digital discovery*. 41 (ACM).
47. Stamatakis, A., Hoover, P. & Rougemont, J. A rapid bootstrap algorithm for the RAxML web servers. *Systematic biology* **57**, 758–771 (2008).
48. Huelsenbeck, J. P. & Ronquist, F. MRBAYES: Bayesian inference of phylogenetic trees. *Bioinformatics* **17**, 754–755 (2001).
49. Felsenstein, J. Phylogenies from Molecular Sequences: Inference and Reliability. *Annual Review of Genetics* **22**, 521–565 (2003).
50. Drummond, A. J. & Rambaut, A. BEAST: Bayesian evolutionary analysis by sampling trees. *BMC Evolutionary Biology* **7**, 214–214 (2007).
51. Drummond, A. J., Ho, S. Y. W., Phillips, M. J. & Rambaut, A. Relaxed Phylogenetics and Dating with Confidence. *PLOS Biology* **4** (2006).
52. Louis, A., Muffato, M. & Crollius, H. R. Genomicus: five genome browsers for comparative genomics in eukaryota. *Nucleic acids research*, gks1156 (2012).
53. Kersey, P. J. *et al.* Ensembl Genomes 2016: more genomes, more complexity. *Nucleic acids research* **44**, D574–D580 (2016).
54. Gu, X. *et al.* An update of DIVERGE software for functional divergence analysis of protein family. *Mol Biol Evol* **30**, 1713–1719, doi: 10.1093/molbev/mst069 (2013).
55. Jeffares, D. C., Tomiczek, B., Sojo, V. & dos Reis, M. A Beginners Guide to Estimating the Non-synonymous to Synonymous Rate Ratio of all Protein-Coding Genes in a Genome. *Parasite Genomics Protocols*, 65–90 (2015).
56. Anisimova, M., Bielawski, J. P. & Yang, Z. Accuracy and power of the likelihood ratio test in detecting adaptive molecular evolution. *Molecular biology and evolution* **18**, 1585–1592 (2001).
57. OS, R. Recent changes to RasMol, recombining the variants. *Biophys. Res. Commun* **266**, 284–289 (2000).

Acknowledgements

We thank Professor Toru Shimada for access to unpublished gene expression data. We are grateful to Professor Jian-Hong Xu for his helpful comments and suggestions on this manuscript. Our special thanks go to Dr. Yunpeng Zhao and Fangluan Gao for the assistance in analytical methods. This work was supported by the National Natural Science Foundation of China (Grant No. 31572321; 31602010; 31572462; 31272375). The work was also supported by a grant from the Science Foundation of Zhejiang Province of China (No. LY15C170001).

Author Contributions

X.L., H.W. and L.S. conceived this study. X.L., Y.Z. and H.X. collected data. Y.C. contributed analysis tools. X.L., X.D., and R.L. carried out experiments and data analyses. X.L., H.W. and L.S. wrote the manuscript. All authors read and approved the manuscript.

Additional Information

Supplementary information accompanies this paper at <http://www.nature.com/srep>

Competing Interests: The authors declare no competing financial interests.

How to cite this article: Li, X. *et al.* Molecular evolutionary mechanisms driving functional diversification of α -glucosidase in Lepidoptera. *Sci. Rep.* **7**, 45787; doi: 10.1038/srep45787 (2017).

Publisher's note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.



This work is licensed under a Creative Commons Attribution 4.0 International License. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license, users will need to obtain permission from the license holder to reproduce the material. To view a copy of this license, visit <http://creativecommons.org/licenses/by/4.0/>

© The Author(s) 2017