

Research Paper

Candidate chemosensory Genes in Female Antennae of the Noctuid Moth *Spodoptera littoralis*

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Abstract

Chemical senses are crucial for all organisms to detect various environmental information. Different protein families, expressed in chemosensory organs, are involved in the detection of this information, such as odorant-binding proteins, olfactory and gustatory receptors, and ionotropic receptors. We recently reported an Expressed Sequence Tag (EST) approach on male antennae of the noctuid moth, *Spodoptera littoralis*, with which we could identify a large array of chemosensory genes in a species for which no genomic data are available.

Here we describe a complementary EST project on female antennae in the same species. 18,342 ESTs were sequenced and their assembly with our previous male ESTs led to a total of 13,685 unigenes, greatly improving our description of the *S. littoralis* antennal transcriptome. Gene ontology comparison between male and female data suggested a similar complexity of antennae of both sexes. Focusing on chemosensation, we identified 26 odorant-binding proteins, 36 olfactory and 5 gustatory receptors, expressed in the antennae of *S. littoralis*. One of the newly identified gustatory receptors appeared as female-enriched. Together with its atypical tissue-distribution, this suggests a role in oviposition. The compilation of male and female antennal ESTs represents a valuable resource for exploring the mechanisms of olfaction in *S. littoralis*.

Key words: Olfactory receptor, Gustatory receptor, Odorant-binding protein, Expressed sequence tag, Lepidoptera, *Spodoptera littoralis*.

Introduction

The sense of smell is highly important for most animals to detect chemical information regarding various fitness-related resources. For many decades,

moths have proven to be important models for studying the physiological and molecular bases of olfactory detection [1]. First, their nocturnal life has led to a

highly developed and sensitive olfactory system well suited to describe general paradigms. Second, moths include diverse and important pests of crops, forests and stored products, and studying their sense of smell offers the possibility to develop olfactory-based strategies to perturb critical behaviors, such as sex pheromone-mediated reproduction, host selection and oviposition.

The noctuid moth *Spodoptera littoralis* is one of the major Lepidoptera models used in olfaction research. This polyphagous noctuid species is an important cotton pest, and extensive research has led to a comprehensive insight into its chemical communication system. The sex pheromone and plant volatiles activating olfactory sensory neurons (OSN) have been identified [2, 3], various functional types of antennal olfactory sensilla have been characterized [3, 4], and the organization of the primary central olfactory system, the antennal lobe, as well as higher areas as the mushroom bodies has been described [5-7]. From a molecular point of view, we recently sequenced an Expressed sequence tag (EST) library prepared from male *S. littoralis* antennae, leading to the identification of a partial repertoire of genes putatively involved in odorant and pheromone detection [8]. Among them, we described members of crucial gene families involved in the olfactory process, such as odorant-binding proteins (OBPs), olfactory and gustatory receptors (ORs, GRs), and ionotropic receptors (IRs). OBPs are soluble proteins proposed to bind odorant molecules and transport them through the aqueous sensillar lymph, allowing hydrophobic molecules to reach OSN dendrites [9, 10]. Another family of binding proteins, the chemosensory protein (CSP) family, groups soluble proteins whose function is unclear. Although some exhibit binding activity towards odorants and pheromones [11, 12], they may participate in other physiological processes beyond chemoreception [13, 14]. ORs and IRs constitute two families of chemosensory receptors located in the dendritic membrane of OSNs, whose activation upon ligand binding leads to the generation of an electrical signal that is transmitted to the brain [15, 16]. ORs are multi-transmembrane domain ionotropic receptors, functioning as ion channels via heterodimerization with a subunit conserved within insects, referred to as ORco [17-19], while the variable part of the dimer defines ligand specificity [20, 21]. This variable part also couples to G proteins, inducing upon activation a complementary metabotropic signaling [20]. IRs are related to ionotropic glutamate receptors but with a divergent ligand-binding domain [16]. Although they appear to be far more ancient than ORs, as they are found across protostomians, they were only recently

discovered as key actors in odorant detection [22, 23]. GRs are transmembrane domain receptors distantly related to insect ORs and expressed in gustatory receptor neurons. Recent studies suggest that GRs present a similar membrane topology to ORs [24] and act as ionotropic channels [25].

Research in the molecular field of moth olfaction has, until recently, largely been restricted to the silkworm *Bombyx mori*, due to the availability of genomic data. Our EST strategy has been demonstrated to be well suited to identify a wide array of olfactory genes in a given species and was particularly relevant for the identification of divergent chemosensory receptors in a species for which no genomic data is available [8]. Similar strategies have been followed to identify broad or partial repertoires of chemosensory receptors in some other moth species, such as *Epiphyas postvittana* [26], *Manduca sexta* [27] and *Cydia pomonella* [28], but the picture of the moth OR and IR gene families is still incomplete.

Here, we report the sequencing and analyses of a female *S. littoralis* antennal EST library that, together with our previous male EST data, allows us to extend our view of the *S. littoralis* antennal transcriptome and to significantly increase the number of identified olfactory genes in *S. littoralis*. Both male and female ESTs were assembled, leading to a total of 13,685 unigenes whose gene ontology (GO) annotation revealed enrichment in binding and catalytic activities. These data allowed us to identify new *S. littoralis* olfactory genes, including binding proteins and chemosensory receptors. Further qPCR experiments were conducted to study tissue- and sex-distribution of the newly identified ORs and GRs, revealing one female-enriched transcript possibly involved in oviposition.

Materials and Methods

Insect rearing and female antennae cDNA library construction

Insects were from a laboratory strain of *S. littoralis* that originated from an Egyptian population. Insects were reared on semi-artificial diet [29], under 23°C, 60-70% relative humidity and 16:8 light:dark cycle. Pupae were sexed and males and females were kept separately. Antennae were collected from 1-2 day old naïve adult females and stored at -80°C. Total RNA were isolated using the TriZol reagent (Invitrogen, Carlsbad, CA, USA), quantified in a spectrophotometer, and the quality verified by agarose gel electrophoresis. A custom normalized cDNA library was created by Evrogen (Russia) in the pAL 17.3 vector, using 1 mg of total RNA as starting material, without

any amplification. The normalization procedure was used to minimize EST redundancy and to enrich the library for rare and low abundant genes, to allow new gene discovery.

EST sequencing

The library was plated, and 20,000 clones were randomly picked. Their 5' ends were sequenced using REV primer by the Genoscope (Evry, France). The plated library was arrayed robotically and bacterial clones had their plasmid DNA amplified using phi29 polymerase. The plasmids were end-sequenced using BigDye Termination kits on Applied Biosystems 3730xl DNA Analysers. Adaptor and vector were localized using cross_match (<http://www.phrap.org/>) using default matrix (1 for a match, -2 penalty for a mismatch), with mean scores of 6 and 10, respectively. Sequences were then trimmed following three criteria: vector and adaptor, poly(A) tail or low quality (defined as at least 15 among 20 bp with a phred score below 12). We finally obtained the 5' end sequences of 18,342 ESTs.

Sequence processing, assembly, unigene and peptide generation

Female ESTs were assembled with previously obtained male ESTs using the TGI Clustering tools (TGICL, <http://compbio.dfci.harvard.edu/tgi/software>) using the default parameters (minimum percent identity for overlaps: 94, minimum overlap length: 30 bp, maximum length of unmatched overhangs: 30), generating unigenes and singletons. Peptides were extracted from the unigenes using FrameDP 1.03 [30], with three training iterations and using Swissprot (398,181 entries, August 2009) as reference protein database.

Gene identification and functional annotation

The newly identified unigenes were compared to the NCBI non redundant protein database (June 24th 2011 version), using BLASTX, with a 1e-8 e-value threshold. The Gene Ontology mapping and distribution were done with the help of BLAST2GO (GO association done by a BLAST against the NCBI NR database) [31]. Finally, the functional domain protein profile and domain were predicted by queries against InterPro using InterProScan, [32], running a batch of analyses (BLASTProDom, Coil, FprintScan, Gene3D, HMMPanther, HMMPfam, HMMPPIR, HMMSmart, HMMTigr, PatternScan, ProfileScan, RNA-BINDING, Seg and Superfamily) on the predicted ORFs.

GO-term enrichment

The analysis of the enrichment in GO-term be-

tween male and female transcriptomes was performed with the BLAST2GO application using GOSSIP [33]. GO-terms in female data (female singletons + contigs assembled from female ESTs only) (test group) were tested for enrichment compared to male data (male singletons + contigs assembled from male ESTs only) (reference group) using Fisher's exact test with multiple testing correction.

Identification of odorant-binding proteins and chemosensory receptors

The *S. littoralis* antennal unigenes were searched with *B. mori* OBPs, CSPs, chemosensory receptors and available insect ORs retrieved from Swissprot as queries using TBLASTN [34]. Additionally, the Interproscan results were scanned for the Interpro accession IPR006170 (Pheromone/general odorant-binding protein, PBP/GOBP Molecular Function: odorant binding GO:0005549) and IPR004117 (Molecular Function: olfactory receptor activity GO:0004984). *S. littoralis* putative chemosensory receptor sequences were in turn employed in searches to find more genes in an iterative process. OR transmembrane domains were predicted using the TMHMM server v.2.0 [35]. OBPs and CSPs were searched for the occurrence of a signal peptide using SignalP 4.0 [36], secondary structures were predicted using the Psipred server [37], and logos were generated using WebLogo [38].

Phylogenetic analyses

We built OR and GR neighbor-joining trees based on Lepidoptera data sets. The OR data set contained 64 amino acid sequences from *B. mori* [39], 18 sequences from *H. virescens* [40, 41], 45 sequences from *M. sexta* [27] and the three OR sequences characterized in *Epiphyas postvittana* [26]. The GR data set contained 65 amino acid sequences from *B. mori* [42], three, two and one sequences from *H. virescens* [41], *M. sexta* [27] and *Papilio xuthus* [43], respectively. Amino acid sequences were aligned using ClustalW2 [44]. Unrooted trees were constructed using the BioNJ algorithm with Poisson correction of distances, as implemented in Seaview v.4 [45]. Node support was assessed using a bootstrap procedure based on 1000 replicates. Images were created using the iTOL web server [46].

Quantitative real-time PCR

Naïve males and females in the middle of their second scotophase were used in the following experiments: female antennae, brains, proboscis, legs (mixture of front, middle, and hind legs), thorax, ovipositors and male antennae total RNAs (one sample per tissue) were extracted with the RNeasy® Mi-

croKit (Qiagen, Hilden, Germany) that included a column DNase treatment. For each tissue, single-stranded cDNA was synthesized from 1 µg of total RNA with 200 U of M-MLV reverse transcriptase (Clontech, Mountain View, CA, USA) using buffer and protocol supplied in the Advantage® RT-for-PCR kit (Clontech). Gene-specific primers for *S. littoralis* chemosensory receptors and the endogenous control *rpL8* were designed using the Beacon Designer 4.0 software (Bio-Rad, Hercules, CA, USA), yielding PCR products ranging from 100 to 250 bp (Table 1). qPCR mix was prepared in a total volume of 20 µl with 10 µl of Absolute QPCR SYBR Green Mix (ThermoFisher Scientific, Epsom, UK), 5 µl of diluted cDNA (or water for the negative control or RNA for controlling for the absence of genomic DNA) and 200 nM of each primer.

qPCR assays were performed on *S. littoralis* cDNAs using a MJ Opticon Monitor Detection System (Bio-Rad). The PCR program began with a cycle at 95 °C for 15 min, followed by 40 cycles of 20 s at 95 °C, 15 s at 53 to 62 °C (depending on the primer pair) and 20 s at 72 °C. To assess the purity of the PCR reactions, a dissociation curve of the amplified product was performed by gradual heating from 50°C to 95°C at 0.2 °C/s. Standard curves were generated by a five-fold dilution series of a cDNA pool evaluating primer efficiency E ($E=10^{-(1/\text{slope})}$). All reactions were performed in duplicate. Chemosensory receptor expression levels were calculated relative to the expression of the *rpL8* control gene and expressed as the ratio = $E_{\text{SlitCR}}^{(\Delta\text{CT SlitCR})}/E_{\text{rpL8}}^{(\Delta\text{CT rpL8})}$ [47].

Table 1. Forward and reverse primer sequences used in real-time PCR, annealing temperatures, resulting amplicon lengths and PCR efficiencies.

Unigenes	qPCR Forward primer sequences (5' to 3')	qPCR Reverse primer sequences (5' to 3')	Annealing T (°C)	Amplicon length (pb)	Efficiency
SlitOR7	CCITCCTATCGATGGCTCTG	CCCAGGTACCACTTGCAGTT	60	115	2.1
SlitOR10	TTGCACTTTATGGGCAATGA	GAAGAGGAAAAAGCGCTGATG	62	188	1.9
SlitOR12	TGGCCTTGGGTGTATCTTC	AAACGGCCACAAGTCTCATC	62	171	2.1
SlitOR19	AAACGTGACTCCGTGAGCTT	CCGCCATCAACGTATTTTCT	62	148	2.3
SlitOR24	CGCATCCGTTTATCGACTTT	CAAACCAGACCACAAGAGCA	60	116	2.1
SlitOR32	GGTACTAAGGCGGTGGATGA	CCAATCCACAACCAAAATCC	58	192	2.2
SlitOR34	CGCAATATGGGTGTCTTCCT	CATGTGCTCGATTCCCTTT	62	178	2.3
SlitGR1	CGACATTTACCGGAATTTT	TTGGGACGAGCCTCAATTAC	60	114	2.1
SlitGR2	GCCGGTGTCCAAGATACACT	CATGCTGATTGCCGAAGTAA	62	168	2.2
SlitGR4	ATGCTGCGTCACACGACTAC	CCAACGGGAACATCTTCAAT	58	115	2.2
SlitGR5	GTTGTGTTGCTGGTGATGG	TTCGAGGCTAGGATCAAGGA	62	159	1.9
<i>S. littoralis</i> RpL8	ATGCCTGTGGGTGCTATGC	TGCCTCTGTGCTTGATGGTA	58/62	189	1.90

Results and discussion

Complete repertoires of moth olfactory genes have been established only in *B. mori*, thanks to the availability of its sequenced genome. To date, it remains the only moth genome available. RNA sequencing appears to be a good alternative to identify partial olfactory gene repertoires in other species, and previous studies have proved it to be efficient [8, 27]. In particular, the high sequence divergence observed within insect ORs has long served as a brake in chemosensory receptor identification. These receptors are very divergent both within and across insect species, precluding their identification through classical homology-based approaches [48].

EST statistics and unigene prediction

A total of 18,342 ESTs (mean length: 664.9 bp, median length: 665.1 bp, max length: 888 bp, min length: 72 bp, Table 2) were obtained from female antennae of *S. littoralis*. These ESTs have been deposited in the European Molecular Biology Laboratory database (EMBL) [EMBL:FQ958213-FQ976554]. The ESTs were processed and assembled with 20,760 male ESTs previously obtained [8]. 30,986 ESTs were assembled into 5,623 contigs, 8,062 ESTs were singletons (20.6 % of total ESTs). On average, each contig was assembled from 18 ESTs. Among the singletons, 4,361 corresponded to male and 3,701 to female ESTs. The assembly led to a total of 13,685 unigenes (mean length: 980.9 bp, median length: 832 bp, max length: 4,100 bp, min length: 40 bp, Table 2) that putatively

represent different transcripts. Compared to the 9,033 unigenes we previously identified in male antennae [8], this led to the identification of 4,652 new unigenes from *S. littoralis* antennae. It must be pointed out that we did only 5' end sequencing that, together with splice variants, polymorphism or reverse transcriptase errors, may have led to under-assembly and thus over-estimation of unigene counts. Indeed, by using the mean coding sequence size of 1.1–1.2 kb in the *B. mori* genome as reference [49], we could estimate the number of genes expressed to be 11,000–12,000 (13,685 unigenes with an average size of ~ 981pb). This is higher than the estimation made in *M. sexta*, for which 7,000–8,000 genes were estimated to be expressed [27].

Identification of putative ORFs

Among the 13,685 unigenes, 8,449 presented a coding region (62 %, mean length: 205.6 aa, median length: 203 aa, max length: 922 aa, min length: 30 aa, Table 2). Protein sequences translated from the predicted open reading frame (ORF) set were compared to the non-redundant protein database (NR, version October 26th 2011). Most of the sequences (80.3 %) translated from predicted ORFs showed similarity to known proteins. 1,663 ORFs showed no similarity at all.

Gene identification and functional annotation

Fig. 1 illustrates the distribution of the *S. littoralis* unigene set in GO terms. Among the 13,685 *S. littoralis* unigenes, 3,376 correspond to at least one GO-term. As observed in the antennal transcriptome of *M. sexta* [27], a large number of transcripts could not be associated with a GO-term (75 %). Among those associated to a GO-term, 2,944 were assigned to a molecular function (87.2 %), 2,147 to putative biological processes (63.6 %), and 1,673 to a cellular component (49.6 %) (Fig. 1). In the molecular function category, the terms “binding” and “catalytic activity” were the most represented (54% and 47%, respectively), as already observed through the analyses of the male ESTs [8]. When we compared the GO-term distribution of “male only” sequences (male singletons + contigs assembled from male ESTs only) with “female only”, differential distribution ($P < 0.05$) was observed for some GO-terms, such as “signal transduction” enriched in male antennae (Fig. 2). Interestingly, numbers of GO-terms were similar in the two datasets, suggesting a similar complexity of male and female antennae (Supplementary material S1). This is different from what has been observed in the analysis of *M. sexta* male and female transcriptomes [27], where the number of terms appearing for biological process and molecular function in female antennae was approximately twice the number as for male antennae.

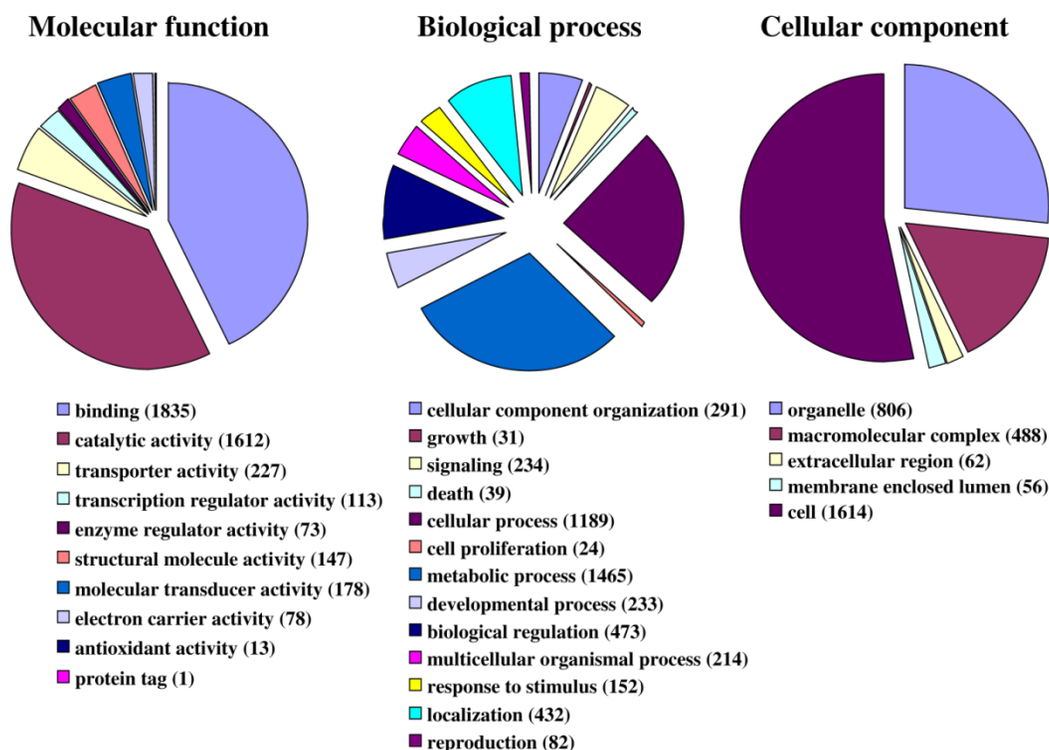


Figure 1. Distribution of *S. littoralis* unigenes annotated at GO level 2.

Table 2. Data summary.

	Counts (total nb)	Min. length (bp/aa)	Average length (bp/aa)	Max length (bp/aa)	Median length (bp/aa)	Accession numbers
Male ESTs	20760	40	958.1	1525	820	FQ0142366-FQ032656 GW824594-GW826804 HO118288-HO118415
Female ESTs	18 342	72	664.9	888	665.1	FQ958213-FQ976554
Unigenes	13685	40	980.9	4,100	832	
Coding regions	8449	30	205.6	922	203	

Differential GO-term distribution (% of sequences)

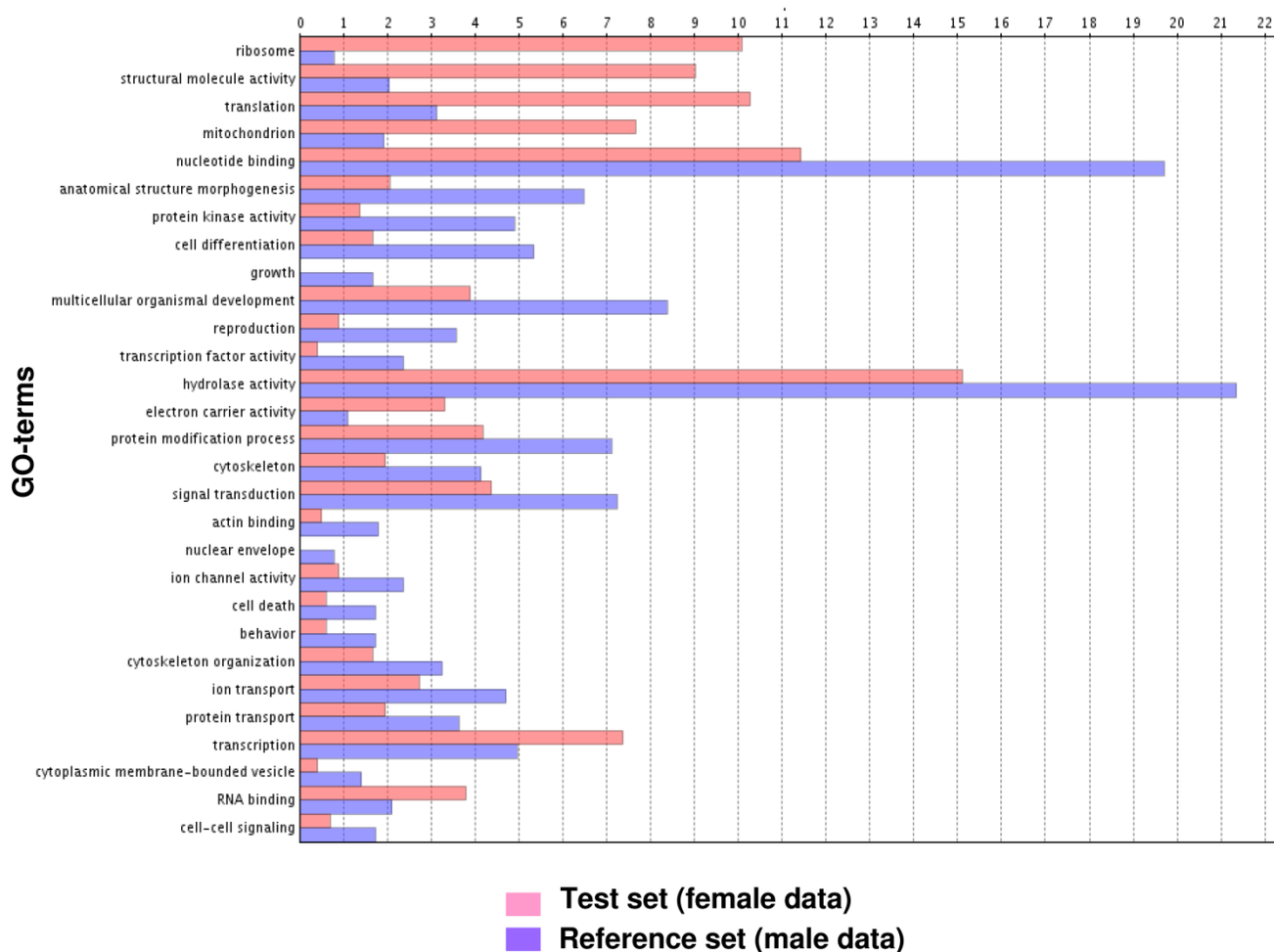


Figure 2. GO-terms (GO-Slim) differentially distributed between male and female transcriptomes (Fisher's exact test, $P < 0.05$). Female data: female singletons + contigs assembled from female ESTs only (test group). Male data: male singletons + contigs assembled from male ESTs only (reference group).

Olfactory and gustatory receptors

In our previous study, which focused on male antennal ESTs, we annotated 29 ORs and 2 GRs. Thanks to the female antennal sequences, we could

extend five of the previously characterized SlitORs (OR4, 5, 8, 21, 29, former accession numbers FQ018861, EZ982621, FQ030158, EZ983645, EZ981024) and we identified seven new candidate SlitORs (OR7, 10, 12, 19, 24, 32, 34). In addition, one GR (GR2, former

accession number GW825869) could be extended and three new candidate GRs were annotated (GR1, 4, 5). This led to a total of 36 ORs and 5 GRs identified in *S. littoralis* antennae (Table 3, fasta format file in Supplementary material S2). For convenience, SlitORs and GRs were numbered according to their closest

homologs –when possible – from *H. virescens*, *M. sexta* or *B. mori* in the phylogenetic analyses (Fig. 3 & 4). Note that SlitOR9 is missing because the corresponding sequence was first annotated as a candidate OR but was later eliminated due to the presence of a stop codon within the predicted ORF.

Table 3. List of *S. littoralis* unigenes putatively involved in chemosensory reception. Transmembrane domains (TM) were predicted using TMHMM version v.2.0.[35].

Name	Length (amino acid)	TM nb	BlastP hit	E value
SlitOR1	298	4	ref NP_001116817.1 olfactory receptor-like [Bombyx mori]	1e-118
SlitOR2	473	7	gb ABQ82137.1 chemosensory receptor 2 [Spodoptera littoralis]	0.0
SlitOR3	270	4	gb AEF32141.1 odorant receptor [Spodoptera exigua]	8e-174
SlitOR4	386	6-7	ref NP_001166616.1 olfactory receptor 54 [Bombyx mori]	2e-101
SlitOR5	397	6	ref NP_001103623.1 olfactory receptor 33 [Bombyx mori]	8e-99
SlitOR6	263	3	emb CAG38117.1 putative chemosensory receptor 16 [Heliothis virescens]	5e-111
SlitOR7	216	4	emb CAD31853.1 putative chemosensory receptor 7 [Heliothis virescens].	6e-98
SlitOR8	258	4	emb CAD31949.1 putative chemosensory receptor 8 [Heliothis virescens]	2e-110
SlitOR10	212	3	gb ACC63238.1 olfactory receptor 10 [Helicoverpa armigera]	1e-70
SlitOR11	223	3	gb ACS45305.1 candidate odorant receptor 2 [Helicoverpa armigera]	1e-139
SlitOR12	224	3	emb CAG38113.1 putative chemosensory receptor 12 [Heliothis virescens].	8e-79
SlitOR13	299	4	dbj BAG71423.2 olfactory receptor [Mythimna separata]	5e-130
SlitOR14	238	4	ref NP_001155301.1 olfactory receptor 60 [Bombyx mori]	8e-126
SlitOR15	390	7	ref NP_001091789.1 olfactory receptor 15 [Bombyx mori]	1e-156
SlitOR16	410	7-8	emb CAG38117.1 putative chemosensory receptor 16 [Heliothis .virescens]	0.0
SlitOR17	391	5	emb CAG38118.1 putative chemosensory receptor 17 [Heliothis virescens].	0.0
SlitOR18	328	5	gb ACL81189.1 putative olfactory receptor 18 [Spodoptera littoralis]	0.0
SlitOR19	216	4	tpg DAA05980.1 TPA: TPA_exp: odorant receptor 22 [Bombyx mori]	1e-101
SlitOR20	331	5	emb CAD31949.1 putative chemosensory receptor 8 [Heliothis virescens]	4e-108
SlitOR21	215	4	emb CAG38122.1 putative chemosensory receptor 21 [Heliothis virescens]	5e-112
SlitOR22	280	5	dbj BAH66361.1 olfactory receptor [Bombyx mori]	5e-07
SlitOR23	422	6	gb EHJ75140.1 olfactory receptor [Danaus plexippus]	4e-63
SlitOR24	211	5	ref NP_001166621.1 olfactory receptor 64 [Bombyx mori]	4e-42
SlitOR25	131	1	ref NP_001166621.1 olfactory receptor 64 [Bombyx mori]	8e-58
SlitOR26	391	5	ref NP_001091790.1 candidate olfactory receptor [Bombyx mori]	0.0
SlitOR27	429	6-7	ref NP_001166607.1 olfactory receptor 44 [Bombyx mori]	0.0
SlitOR28	453	5	gb ABQ84982.1 putative chemosensory receptor 12 [Spodoptera littoralis].	0.0
SlitOR29	326	5	ref NP_001166894.1 olfactory receptor 29 [Bombyx mori]	9e-169
SlitOR30	239	4	dbj BAH66327.1 olfactory receptor [Bombyx mori]	9e-57
SlitOR31	400	5	dbj BAH66346.1 olfactory receptor [Bombyx mori]	4e-93
SlitOR32	205	4	ref NP_001104832.2 olfactory receptor 16 [Bombyx mori]	2e-101
SlitOR33	250	4	ref NP_001091785.1 olfactory receptor 19 [Bombyx mori]	7e-47
SlitOR34	141	2	gb ADM32898.1 odorant receptor OR-5 [Manduca sexta]	8e-13
SlitOR35	408	6	ref NP_001103476.1 olfactory receptor 35 [Bombyx mori]	1e-156
SlitOR36	245	4	ref NP_001103476.1 olfactory receptor 35 [Bombyx mori]	5e-89
SlitOR37	266	5	gb EFN70678.1 Putative odorant receptor 13a [Camponotus floridanus].	0.90
SlitGR1	120	1	gb EHJ69979.1 putative gustatory receptor candidate 59 [Danaus plexippus].	1e-22
SlitGR2	244	4	gb EHJ68848.1 putative Gustatory receptor 21a [Danaus plexippus].	3e-155
SlitGR3	213	4	gb EHJ78216.1 gustatory receptor 24 [Danaus plexippus]	4e-93
SlitGR4	128	1	ref NP_001091791.1 candidate olfactory receptor [Bombyx mori]	2e-26
SlitGR5	205	4	emb CAD31947.1 putative chemosensory receptor 5 [Heliothis virescens]	2e-73

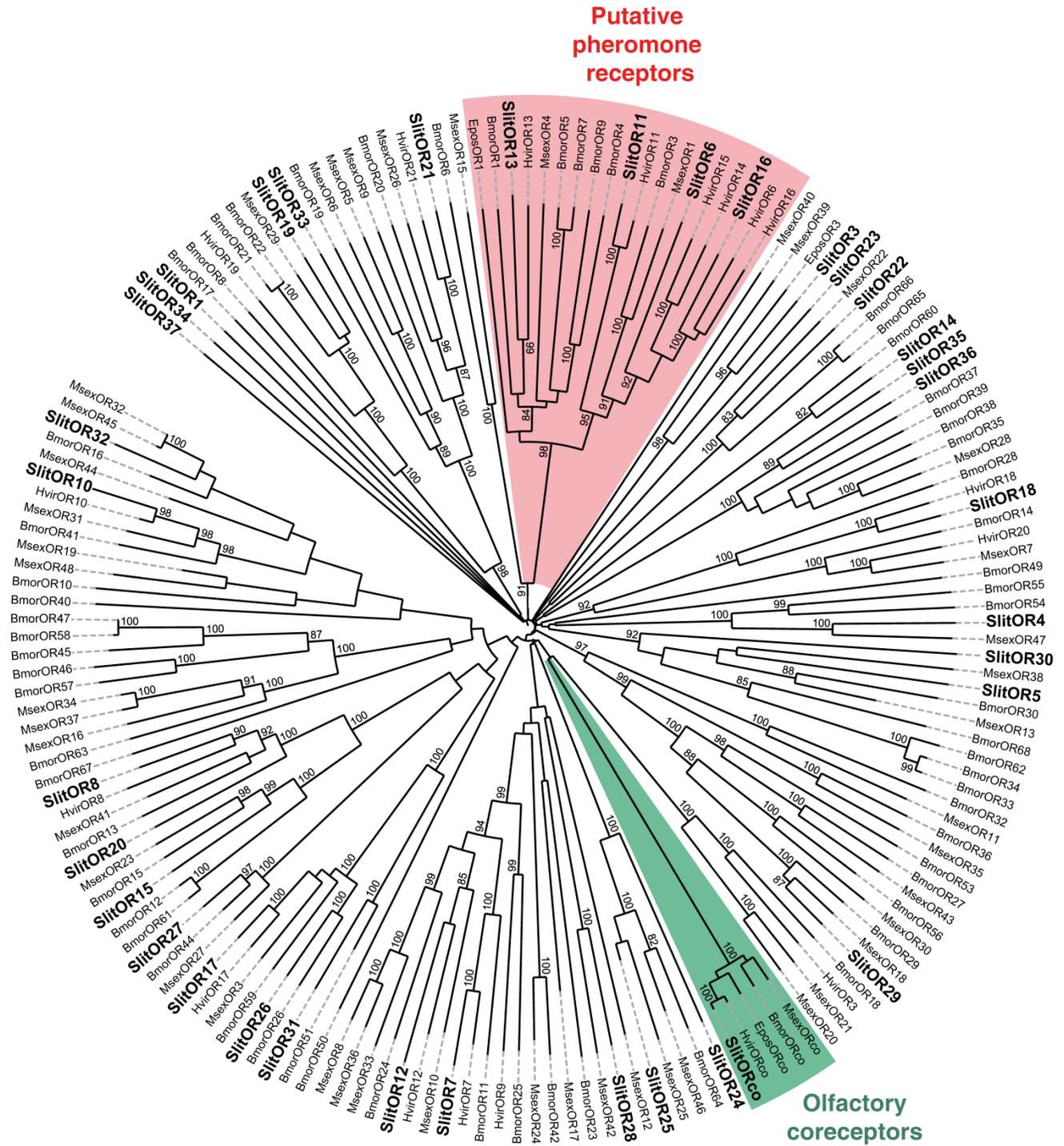


Figure 3. Neighbor-joining tree for candidate olfactory receptors (ORs) from *S. littoralis* and other Lepidoptera. The tree was drawn with iTOL, based on an unrooted tree constructed using the BioNJ algorithm in Seaview v.4, which was made based on a sequence alignment using ClustalW2. Bmor, *B. mori* [39]; Epos, *E. postvittana* [52], Hvir, *H. virescens* [40, 41]; Msex, *M. sexta* [27]; Slit, *S. littoralis* [8](this paper).

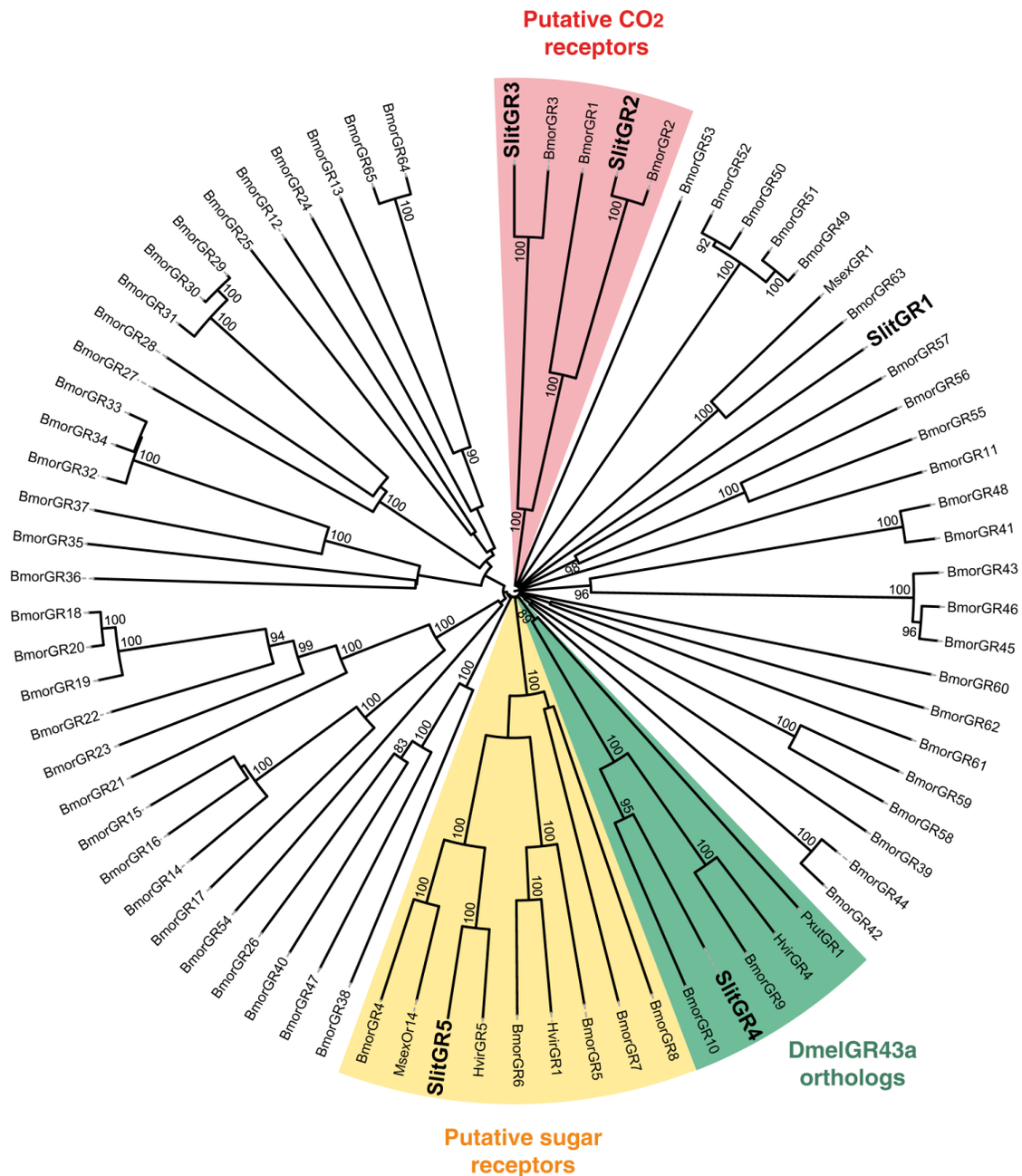


Figure 4. Neighbor-joining tree for candidate gustatory receptors (GRs) from *S. littoralis* and other Lepidoptera. The tree was drawn with iTOL, based on an unrooted tree constructed using the BioNJ algorithm in Seaview v.4, which was made based on a sequence alignment using ClustalW2. Bmor, *B. mori* [42]; Hvir, *H. virescens* [41]; Msex, *M. sexta* [27]; Pxut, *P. xuthus* [43]; Slit, *S. littoralis* [8](this paper).

In *S. littoralis*, 63 glomeruli have been identified in the antennal lobe [6]. Considering the one receptor-one glomerulus paradigm [50, 51], by which the number of expected ORs in a given species should correlate with the number of glomeruli in the antennal lobe, the 36 candidate OR genes identified may not represent the entire repertoire of adult *S. littoralis* ORs. Indeed, 47 and 43 ORs were identified through adult antennal transcriptome sequencing in *M. sexta* [27] and *C. pomonella* [28], respectively. In *B. mori*, 66 ORs

were annotated from genome analyses [39]. However, expression studies revealed that *B. mori* adult antennae express only 35 ORs out of these 66 [39], a number that is very close to the 36 ORs we annotated in *S. littoralis*. Some glomeruli are also very likely innervated by OSNs expressing other classes of chemoreceptors, such as ionotropic receptors and gustatory receptors.

To complete our previously work [8], we conducted a preliminary study of the tissue-distribution

of the newly identified *S. littoralis* candidate chemosensory receptors by qPCR, and we built neighbor-joining trees with all identified SlitORs and GRs. The seven new candidate ORs were clearly antenna enriched, with antennal expression in both sexes (Fig. 5AB), supporting our annotation. In the OR phylogenetic analysis (Fig. 3), no homologs of the *B. mori* cis-jasmone receptor (BmorOR56) or the linalool/linalyl acetate receptor (BmorOR42) were found, but homologs of BmorOR29 (whose known ligands are linalool, citral and several acetates) and of the *Epiphyas postvittana* citral receptor (EposOR3) were present [39, 52]. Neither were homologs of the female-enriched *B. mori* ORs (OR30, 45, 46, 47, 48, 50 [53, 54]) found, but SlitOR33 clustered with a *B. mori* female-specific receptor, BmorOR19, which binds linalool [53, 54]. SlitOR33 was, however, equally expressed in male and female antennae of *S. littoralis* [8]. None of the new sequences clustered in the pheromone receptor clade (Fig. 3).

In the GR phylogeny (Fig. 4), one of the new GR candidates (SlitGR4) grouped in the *D. melanogater* GR43a ortholog subgroup, which includes the newly

characterized *B. mori* fructose receptor (BmorGR9) [25] and the *P. xuthus* synephrine receptor (PxutGR1). The latter is expressed in female tarsi and necessary for correct oviposition behavior in this butterfly [43]. SlitGR4 appeared in our preliminary qPCR analysis as enriched in female antennae compared to male (Fig. 5B), and was also expressed in other female body parts such as proboscis, legs and ovipositors (Fig. 5A). Together with the fact that sugars and other carbohydrates are known to influence host preference and oviposition in female moths [55], these data suggest that SlitGR4 may fulfill an important function in *S. littoralis* female oviposition. SlitGR5 grouped in the putative sugar receptor subfamily (Fig. 4), which includes the newly characterized *B. mori* inositol receptor (BmorGR8) [24]. This is in concordance with previous electrophysiological studies that revealed that moth antennae are involved in sugar detection [56]. SlitGR5 was found to be expressed not only in antennae but also in female proboscis and legs (Fig. 5A). Taken together, our results suggest that sugar/oviposition site detection may involve different sensory organs. Another SlitGR (SlitGR1) did not

cluster with any identified moth candidate GRs (Fig. 4). Although found in antennae, this GR was mainly expressed in ovipositors and legs (Fig. 5A), which the female may use to explore oviposition sites.

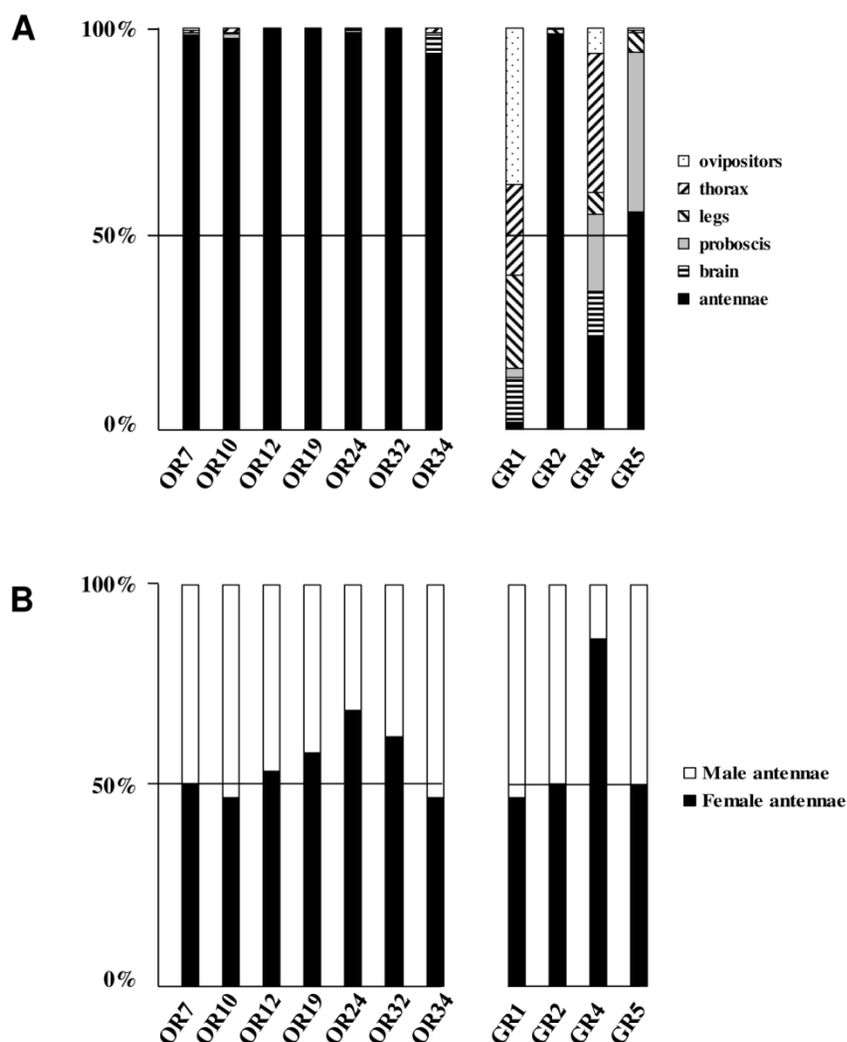


Figure 5. Distribution patterns of the newly identified *S. littoralis* candidate olfactory and gustatory receptors in different female tissues (A) and in male and female antennae (B). Gene expression levels were determined by real-time PCR and calculated relative to the expression of the rpL8 control gene and expressed as the ratio = $E_{\text{SlitCR}}^{\Delta\text{CT SlitCR}}/E_{\text{rpL8}}^{\Delta\text{CT rpL8}}$.

The male EST analyses allowed us to classify one candidate SlitGR (GR2, formerly named GW825869) as a putative CO₂ receptor [8]. Here we could extend its sequence, confirming its high identity (83%) with one of the three *B. mori* candidate CO₂ receptors, BmGr2NJ [42]. Using qPCR, we revealed an unexpected antennal-specific expression of this gene (Fig. 5A). The construction of the GR tree revealed that another SlitGR, GR3, also clustered in the CO₂ receptor family (Fig. 4), this one being equally expressed in antennae and proboscis [8]. Up to now, moth sensory neurons specific for CO₂ have been described only on labial palps [57]. The annotation of two candidate CO₂ receptors being expressed in antennae support our hypothesis that moths may also detect CO₂ *via* their antennae.

Ionotropic receptors

We previously identified 12 candidate IRs in male antennal ESTs [58]. The sequencing of female ESTs revealed that at least seven IR genes were also expressed in female antennae (IR76b, 25a, 75q.1, 75q.2, 41a, 87a, 25a), in concordance with our previous RT-PCR analyses [58]. No additional IR sequences could be identified in female ESTs. Thus, we yet lack the SlitIR8a sequence, a gene suspected to be expressed in insect antennae and, like IR25a, encoding a co-receptor for other IRs [23]. Both IR8a and IR25a

could be identified in the transcriptomes of *M. sexta* [27] and *C. pomonella* [28].

Odorant-binding proteins

A total of 35 sequences showing similarities with Lepidoptera OBPs were identified. Comparisons with the 17 OBP dataset obtained after analysis of the male data revealed that 9 female sequences were new OBPs and one female sequence extended an existing OBP sequence. This led to a total of 26 OBPs identified in *S. littoralis* antennae (FASTA format file in Supplementary material S2). Almost all the identified genes have the characteristic hallmarks of the OBP gene family: the presence of a signal peptide, the six α -helix pattern, and the highly conserved six cysteine profiles (Table 4, Fig. 6). However, despite the highly conserved secondary structure of OBP proteins, the SlitOBPs are highly divergent (average amino acid identity 18 %, min 6 %, max 65%) and exhibit a wide range of protein lengths (up to 266 amino acids) and cysteine number (up to 12, Table 4). For convenience, SlitOBPs were numbered according to their closest homologs. The number of candidate SlitOBPs identified is far less than the 46 annotated OBPs found in the genome of *B. mori* [59, 60] but more than the 18 putative OBPs identified in the transcriptome of *M. sexta* [27].

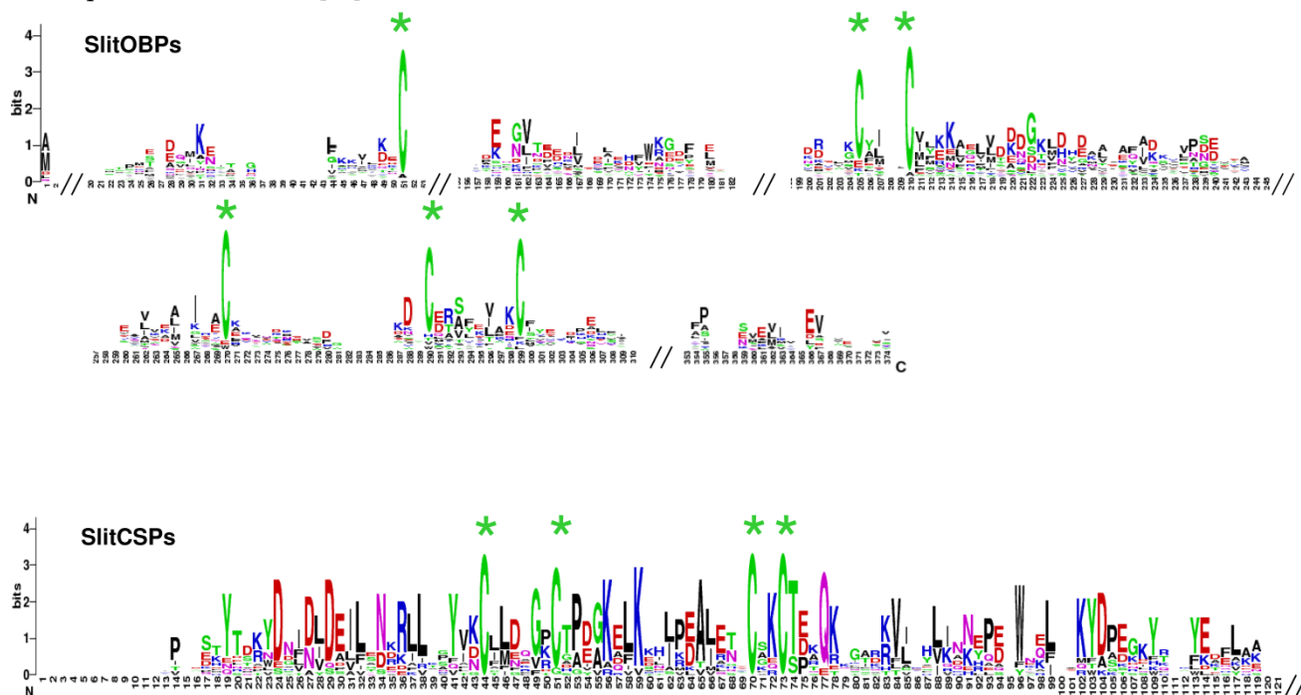


Figure 6. SlitOBP and CSP sequence logo. Degree of amino acid sequence conservation [38] along the primary sequence axis of odorant-binding proteins (OBPs) and the chemosensory proteins (CSPs) of *S. littoralis*. Depicted amino acid character size correlates to relative conservation across aligned sequences. Green asterisks indicate the conserved six and four cysteine motifs of OBP and CSP, respectively.

Table 4. List of *S. littoralis* unigenes putatively involved in odorant binding. Signal peptides were determined using SignalP 4.0 [36] and α -helices structures were predicted using the Psipred server [37].

Name	Length (amino acid)	Signal peptide	C nb	α -helice nb	BlastP hit	E value
SlitPBP1	164	Yes	6	7	ABQ84981.1 pheromone-binding protein 1 [Spodoptera littoralis]	7e-120
SlitPBP2	162	Yes	6	7	AAS55551.2 pheromone binding protein 2 [Spodoptera exigua]	2e-115
SlitPBP3	164	Yes	6	7	ACY78413.1 pheromone binding protein 3 [Spodoptera exigua]	4e-104
SlitGOBP1	163	Yes	7	7	ABM54823.1 general odorant-binding protein GOBP1 [Spodoptera litura]	9e-104
SlitGOBP2	151	No	6	7	ABM54824.1 general odorant-binding protein GOBP2 [Spodoptera litura]	1e-104
SlitOBP1	194	Yes	14	5	EHJ77172.1 odorant binding protein [Danaus plexippus]	1e-59
SlitOBP2	184	Yes	8	6	CAX63249.1 odorant-binding protein SaveOBP4 precursor [Sitobion avenae]	1e-44
SlitOBP3	129	Yes	4	6	ADY17884.1 odorant binding protein [Spodoptera exigua]	8e-72
SlitOBP4	134	Yes	4	6	AAL60426.1 antennal binding protein 8 [Manduca sexta]	6e-74
SlitOBP5	158	Yes	7	6	ADY17882.1 odorant binding protein [Spodoptera exigua]	1e-108
SlitOBP6	62	No	3	3	NP_001140187.1 odorant-binding protein 3 precursor [Bombyx mori]	4e-15
SlitOBP7	141	Yes	7	6-7	AEB54588.1 OBP13 [Helicoverpa armigera]	5e-78
SlitOBP8	252	Yes	2	2	BAH79159.1 odorant binding protein [Bombyx mori]	1e-121
SlitOBP9	258	Yes	6	5	ADQ01713.1 odorant binding protein 29 [Anopheles funestus]	1e-14
SlitOBP10	174	No	5	4	EFA09155.1 odorant binding protein 22 [Tribolium castaneum]	4e-10
SlitOBP11	147	Yes	6	6	AAR28762.1 odorant-binding protein [Spodoptera frugiperda]	8e-94
SlitOBP12	147	Yes	6	6	ADY17881.1 antennal binding protein [Spodoptera exigua]	2e-93
SlitOBP13	217	Yes	12	6	AAL60414.1 twelve cysteine protein 1 [Manduca sexta]	2e-21
SlitOBP14	142	Yes	6	6	AEB54586.1 OBP2 [Helicoverpa armigera]	2e-87
SlitOBP15	131	No	7	6	AAL60415.1 antennal binding protein 4 [Manduca sexta]	7e-53
SlitOBP16	122	No	9	5	AAL60414.1 twelve cysteine protein 1 [Manduca sexta]	0.040
SlitOBP17	144	Yes	6	6	AAL60415.1 antennal binding protein 4 [Manduca sexta]	5e-75
SlitOBP18	151	Yes	6	6	AEB54589.1 OBP8 [Helicoverpa armigera]	1e-83
SlitOBP19	239	Yes	4	2	NP_001157372.1 odorant binding protein fmxg18C17 precursor [Bombyx mori]	1e-73
SlitOBP20	137	Yes	6	6-7	CAA05508.1 antennal binding protein X [Heliothis virescens]	5e-75
SlitOBP21	126	no	8	6	AAL60413.1 antennal binding protein 3 [Manduca sexta]	1e-49
SlitCSP1	128	Yes	4	6	ACX53804.1 chemosensory protein [Heliothis virescens]	1e-71
SlitCSP2	120	Yes	4	6	ACX53800.1 chemosensory protein [Heliothis virescens]	2e-75
SlitCSP3	75	No	3	4	ABM67686.1 chemosensory protein CSP1 [Plutella xylostella]	8e-20
SlitCSP4	148	Yes	5	5	EHJ76401.1 chemosensory protein CSP1 [Danaus plexippus]	2e-51
SlitCSP5	128	Yes	4	6	ABM67688.1 chemosensory protein CSP1 [Spodoptera exigua]	7e-85
SlitCSP6	123	Yes	4	6	ACX53806.1 chemosensory protein [Heliothis virescens]	3e-72
SlitCSP7	108	Yes	4	5	BAF91720.1 chemosensory protein [Papilio xuthus]	1e-56
SlitCSP8	128	Yes	4	6	ABM67689.1 chemosensory protein CSP2 [Spodoptera exigua]	5e-86
SlitCSP9	127	Yes	6	6	AAAY26143.1 chemosensory protein CSP [Spodoptera litura]	1e-89
SlitCSP10	266	Yes	4	6-7	NP_001037069.1 chemosensory protein 9 precursor [Bombyx mori]	2e-89
SlitCSP11	113	Yes	4	6	AAK14793.1 sensory appendage protein-like protein [Mamestra brassicae]	5e-44
SlitCSP12	124	Yes	4	6	ACX53817.1 chemosensory protein [Heliothis virescens]	4e-58
SlitCSP13	122	Yes	4	6	ACX53813.1 chemosensory protein [Heliothis virescens]	4e-75
SlitCSP14	127	Yes	4	6	BAF91712.1 chemosensory protein [Papilio xuthus]	1e-70

Sensory neuron membrane proteins and chemosensory proteins

Sensory neuron membrane proteins (SNMPs), located in the dendritic membrane of primarily

pheromone-specific OSNs, are thought to trigger ligand delivery to the receptor [61]. Moths usually express two SNMPs, and the *D. melanogaster* SNMP1 homolog has been demonstrated to play a role in pheromone detection [62]. The two SlitSNMPs were

previously identified in the male ESTs, and were also represented in female ESTs, revealing that these proteins are expressed in both sexes. The SlitSNMP1 sequence was complete thanks to the male data. The female data allowed us to complete the SlitSNMP2 sequence.

Chemosensory proteins are soluble proteins expressed in a wide range of tissues and a chemosensory function has in fact been demonstrated for only a restricted number of such proteins [10], the others acting as general carriers for hydrophobic ligands in the insect body. Here, we identified 17 sequences expressed in female antennae showing similarities with Lepidoptera CSPs. Comparison with the 9 CSP annotated in the male ESTs revealed that five female sequences were new CSPs, leading to a total of 14 CSPs identified in *S. littoralis* antennae (FASTA format file in Supplementary material S2). Almost all deduced protein sequences have the characteristic hallmarks of CSPs: the presence of a signal peptide, the six α -helix pattern, and the highly conserved four cysteine profiles (Table 4, Fig. 6). 22 putative CSPs have been annotated in *B. mori* [60, 63] and 21 in *M. sexta* [27], while we identified only 14 candidate CSPs. Nevertheless, our data confirmed that Lepidoptera express a higher number of CSPs than other insect families, such as Diptera [60]. Although their role in chemoreception should be taken with caution, at least one SlitCSP (SlitCSP3) may be involved in contact chemoreception since it is homologous to *Plutella xylostella* CSP1 that binds non volatile oviposition deterrents [64].

LepidoDB implementation

Lepido-DB (<http://www.inra.fr/lepidodb>) is a centralized bioinformatic resource for the genomics of major lepidopteran pests [65]. This Information System was designed to store, organize, display and distribute various genomic data and annotations. All the data, unigenes, ORFs and their annotation generated in this project have been included in LepidoDB. As a result, from the project page http://www.inra.fr/lepidodb/spodoptera_littoralis, one can retrieve the whole sequence set, query with a keyword and retrieve the corresponding sequences.

Conclusion

The main objective of this study was to investigate the antennal transcriptome of the noctuid moth *S. littoralis*. Thanks to the sequencing of female ESTs, we annotated a total of 26 candidate OBPs, 14 CSPs, 36 ORs and 5 GRs in the antennae of *S. littoralis*. This strategy appears to be particularly relevant for the identification of new insect chemosensory receptors in a species for which no genomic data are available. The

availability of this large antennal transcriptome constitutes a valuable resource for studies of insect olfaction.

Although the generation of gender-specific transcriptomes did not highlight strong differences between sexes, we evidenced male-specific ORs as candidate pheromone receptors and a female-enriched GR as a candidate oviposition stimulant receptor. It has to be noticed that we focused our study on naïve insects, whereas mating is known to induce big changes in the olfactory behaviors. For example, *S. littoralis* females switch their olfactory response from food to egg-laying cues following mating [66]. This switch may be associated with regulation in the transcriptome expression. Comparison of antennal transcriptomes from males and females that encountered diverse experiences would then lead to identification of more regulated genes, as candidate genes involved in gender-specific behaviors.

Supplementary Material

S1: Comparison of male and female gene expression using GO categorization for biological processes (upper) and molecular function (lower), both level 3.

S2: Fasta amino acid sequences of annotated *S. littoralis* OBPs, CSPs, ORs and GRs.

<http://www.biolsci.org/v08p1036s1.pdf>

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Competing Interests

The authors have declared that no competing interest exists.

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