Digital Atlases of the Antennal Lobe in Two Species of Tobacco Budworm Moths, the Oriental Helicoverpa assulta (Male) and the American Heliothis virescens (Male and Female)

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ABSTRACT

The antennal lobe of the moth brain is the primary olfactory center processing information about pheromones and plant odors. We present here a digital atlas of the glomerular antennal lobe structures in the male of Helicoverpa assulta and the male and female of Heliothis virescens, based on synaptic antibody staining combined with confocal microscopy. The numbers of the glomeruli in the three specimens were similar, 65, 66, and 62, respectively. Whereas the male antennal lobe has a macroglomerular complex consisting of three and four units in the two species, the female lobe has two enlarged glomeruli at a corresponding position, near the entrance of the antennal nerve. Another large glomerulus, showing homology in the three specimens, is ventrally located. The small size of the heliothine moths is advantageous for confocal microscopy because the entire brain can be visualized as a single image stack. The maps are freely accessible on the internet, and the digital form of the data allows each atlas to be rotated and sectioned at any angle, providing for the identification of glomeruli in different preparations. J. Comp. Neurol. 446:123–134, 2002.

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Indexing terms: olfactory system; glomeruli; heliothine moths; anatomical reconstruction; confocal microscopy; internet

Olfactory detection in insects is performed by receptor neurons housed in sensilla, most of which are located on the antennae. The sensory neurons send projections directly to the antennal lobe, the primary olfactory center of the insect brain. Here, the terminals make synapses with second-order neurons in spheroidal structures called glomeruli (Rospars, 1988; Boeckh and Tolbert, 1993). Shepherd (1974) defines a glomerulus as “a synaptic complex enclosed in glial membranes or otherwise set apart.” This definition applies to most animal species, vertebrates as well as invertebrates. In some insects, such as lepidopterans, the males possess a specific macroglomerular complex (MGC) in the antennal lobe, forming a first relay for processing information about sexual pheromones and interspecific signals. The functional significance of the individual MGC compartments has been thoroughly investigated in several moth species by functional tracing of the pheromone receptor neurons as well as of the antennal lobe projection neurons (Christensen et al., 1991, 1995;...

All heliothine species so far investigated use a multi-component pheromone blend containing the same chemicals in different compositions. The receptor neurons detecting the “insect-produced” compounds in the various species can be classified into types according to their specificities, each type receiving information about one chemical (Mustaparta, 1997). Because of the detailed knowledge of physiology and behavior involved in intra- and interspecies pheromone communication, the MGC in heliothine moths also offers interesting comparative aspects concerning functional organization of the glomerular units. In the male of the American tobacco budworm moth *Heliothis virescens*, the number of identified receptor neuron types detecting insect-produced substances matches perfectly the number of glomerular units constituting the MGC. Four behaviorally relevant chemicals are detected by four specific receptor neuron types, each projecting in one of four MGC compartments (Hansson et al., 1995; Berg et al., 1998). The two largest units receive input from neurons carrying information used in behavioral attraction and the two smallest from neurons responsible for interruption of attraction. Furthermore, there appears to be a correspondence between the input and the output information of each MGC compartment, supporting the idea that the individual glomeruli are particularly involved in processing information about one of the chemicals (Christensen et al., 1995; Vickers et al., 1998; Berg et al., 1998).

The role of the ordinary glomeruli, which are the targets for the receptor neurons detecting plant odor volatiles, has until now remained relatively unexplored. In contrast to the pheromone system, offering a small number of easily recognizable MGC compartments, the ordinary glomeruli are numerous and very challenging for identification. Another important task is to identify relevant odorants for which the plant odor receptor neurons have evolved. Recently, a major receptor neuron type responding with high sensitivity and selectivity to the sesquiterpene germacrene D has been found in the *H. virescens* female (Rosteljen et al., 2000). The large number of these neurons indicates the significance of the odorant for the detection of plants by female moths. A functionally similar receptor neuron type has also been identified in the closely related species *H. assulta*. An interesting question is whether the axons of all germacrene D receptor neurons project to a single glomerulus and, if so, whether this particular glomerulus can be considered as homologous in different species of heliothine moths.

In the present study, we have mapped the individual glomeruli of the antennal lobes in two heliothine species, the male and female of *H. virescens* and the male of *H. assulta*. As a contribution in a series of digital atlases of the insect antennal lobe that are published on the internet, including the antennal lobe of the honeybee *Apis mellifera* (Galizia et al., 1999) and the fruit fly *Drosophila melanogaster* (Laisseau et al., 1999), we present here the antennal lobes of two moth species. The internet publication of the present data makes these atlases freely available as a tool, allowing other researchers to map physiological responses to morphologically identified glomeruli. The results obtained from studies of the pheromone system in combination with the recent data on the plant odor system make heliothine moths suitable model organisms for investigating the function of individual glomeruli as well as enlightening similarities and differences between related species.

**MATERIALS AND METHODS**

**Antibody staining procedure**

*H. virescens* and *H. assulta* pupae, originating from laboratory cultures, were kindly provided by Dr. Max Angst (Novartis Crop Protection, Basel, Switzerland), Dr. K.S. Boo (Seoul National University, Suwon, Republic of Korea), and Dr. Sadahiro Tatzuki (The University of Tokyo, Tokyo, Japan). Newly eclosed insects were used for the experiments. The brains were dissected from the head and kept overnight in a 4% paraformaldehyde fixation at 4°C. After fixation, the tissue was rinsed (10 × 1 hour) with 0.1 M phosphate-buffered saline (PBS), pH 7.4, with added 1% Triton X-100 (PBST: 8 g NaCl, 0.2 g KCl, 1.44 g Na2HPO4, 0.24 g KH2PO4, 1 ml Triton). The brains were preincubated in PBST, pH 7.4, +10% normal goat serum (NGS; Sigma) for 2 hours and then incubated for 60 hours at room temperature with the primary antiserum SYNORF1 and nc46, each diluted 1:30 in 10% NGS. The SYNORF1 detects a particular part of the synapsin molecule, whereas the antigen of nc46 is not known. The mixture of the two instead of one is shown to be favorable in honeybees. Both primary antibodies stem from a screen of *Drosophila* synapse protein (Reichmuth et al., 1995; Klagges et al., 1996) and were kindly provided by Dr. E. Buchner (Würzburg, Germany). Only one type of secondary antibody was used. After incubation in the primary antiserum, the brains were rinsed (10 × 1 hour) in PBST and then incubated in the secondary antiserum, a Cy3-conjugated mouse anti-rabbit antibody (Jackson Immunoresearch, West Grove, PA; diluted 1:200 in 10% NGS), and kept in darkness for ca. 36 hours at room temperature. After 10 rinses, each for 1 hour, in PBS (in the darkness), the brains were dehydrated through a graded series of alcohols (50%, 70%, 90%, 99%, 2 × 100%, each for 10 min) and cleared in methyl salicylate. The brains were oriented in a frontal position and embedded in Permoun (Sigma, St. Louis, MO) in double-sided custom slides.

**Confocal laser scanning microscopy**

Twenty-eight brains were reconstructed by laser scanning microscopy (LSM). For further analyses, six were selected (two *H. assulta* males, two *H. virescens* males and two *H. virescens* females). One of each of three preparations (all named specimen 1) was used as a model to create the digital atlases. The other three (all named specimen 2) were studied for comparing the number and position of the glomeruli between individuals of each type.

The three brains (specimens 1) used for the creation of the atlases were analyzed with a Leica TCS 4D confocal microscope (DM RBE). Dry immersion objectives were used to scan the specimens, a Leica HC PL APO 20×/0.4 dry objective for the whole brain and a Leica PL APO 150 mW mixed gas laser and detected with a longpass filter (LP590). The brains were scanned in a set of frontal planes. To remove random noise, a scan average of eight lines was applied. The spatial sampling frequency was 0.72 μm/voxel in the image plane (512 × 512 pixels) and
2.01 μm/voxel in the axial direction. The entire stack of images contained 130 sections for the H. assulta brain specimen, 150 for the H. virescens male, and 160 for the H. virescens female. The data were stored on a CD-ROM. The z-axis dimension was corrected by a factor of 1.6 to allow for a z-axis shortening introduced by the refractive index mismatch in the optical pathway when using a dry objective. Under these conditions, movement of the microscope stage by 1.25 μm per slice corresponded to an optical interslice distance of 2 μm.

The three brains (specimens 2) used for comparison of the glomerular pattern were analyzed with a Zeiss LSM 510 LSM. A dry immersion objective, plan-Neofluar 20×/0.5, was used to scan the preparations. An excitation wavelength of 543 nm was used with a He-Ne 1 mW mixed-gas laser. The spatial sampling frequency was similar to that described above, 0.9 μm/voxel in the image plane (512 × 512 pixels) and 1.9 μm/voxel in the axial direction.

Three-dimensional reconstructions and data analyses

The choice of the six scanned preparations used for further analyses was based on the quality of the stainings. The selected brains displayed a satisfactory extent of fluorescence staining in each section of the entire image stack, making it possible to discriminate the glomerular structures from the background throughout the antennal lobe.

To analyze the raw data, professional, general-purpose visualization and 3D reconstruction software were used: Amira 2.1 (Indeed GmbH, Berlin; http://www.amiravis.com) and Zeiss 3D for LSM 510 2.8. The digital atlases were created in Amira, based on the extraction of polygonal surface models from 3D image data by tracing the individual glomeruli section by section. The segments were made in frontally oriented optical slices. Each digital reconstruction was based on optical slices of one specimen (specimen 1). Different tools were used for the segmentation analysis. By adjusting the gray value mapping in each slice, the anatomical structures were discriminable from the background. The simultaneous display of the data sets in the xy-, xz-, and yz-planes made it easier to decide the outlines in the z-direction. For the H. assulta male and the H. virescens female preparation, each image section of the entire stack was analyzed separately; i.e., outlines of the glomeruli were traced every second micrometer. For the H. virescens male, two image sections were put together, so that tracings were made every fourth micrometer. To visualize the raw data from the confocal microscope in 3D, as shown in Figure 1, the “volume rendering” tool of Amira was used. This technique, which gives a 3D impression of the whole data set without segmentation, is based on the attribution of light emission and absorption to every voxel of the volume as a function of its staining intensity, which is subsequently projected onto the viewing plane.

For comparing the glomerular patterns in two individuals of each type, a second reconstruction of one H. assulta male, one H. virescens male, and one H. virescens female was made using the software program Zeiss 3D for LSM 510 2.8. In H. assulta a comparison of the glomerular pattern was carried out by identifying each glomerulus in specimen 2 according to specimen 1 using the two stacks of sequential optical images. In H. virescens, 36 glomeruli in the male specimen 2 and 32 glomeruli in the female specimen 2 were identified by comparing the respective 3D reconstructions of the raw data with the corresponding atlases. In addition, the total number of glomeruli was counted by analyzing the stacks of sequential optical images in the two sexes. To visualize the raw data of each specimen 2 in 3D, the “projection” tool of LSM 510 was used. This function calculates a series of sections after rotating the data package about one of three axes. The final figures were edited with Adobe Pagemaker 6.5. Some of the images were adjusted for brightness and contrast with Adobe Photoshop v. 5.5.

Terminology

The MGC structures in the two male species and the enlarged glomeruli in the female (LFG), positioned at the entrance of the antennal nerve, were easy to recognize. These glomeruli were given names according to previous descriptions for heliothine moths and the sphinx moth Manduca sexta (Christensen et al., 1991; Berg et al., 1998, 1999; King et al., 2000). The remaining ordinary glomeruli were given numbers for identification, roughly following a system in which each glomerulus was given a value according to its position along the z-axis, so that frontally located glomeruli have low numbers, whereas those located posteriorly have high numbers. The landmarks used for orientation of the antennal lobe were the antennal nerve and the two cell soma clusters, the lateral and the medial. These brain structures were included in the atlases for orientation purposes.

RESULTS

Histology

The neuropil-specific monoclonal antibody staining gave rise to an intense labeling of the glomeruli, revealing them as anatomically separable units (Fig. 1A,B). The total assembly of glomeruli forms the outer part of the spheroidal antennal lobe, with the central area devoid of any glomerulus (Fig. 2). The antennal lobes are the most anterior and prominent part of the deutocerebrum. The remaining deutocerebrum, the antennal mechanosensory and motor center, which was also strongly labeled, showed no glomerular organization (Fig. 1A,B). With this technique we also obtained clear labeling of the protocerebral structures, the calyces of the mushroom bodies, located posteriorly in the moth brain, as well as of the subesophageal ganglion and the internal structures of the optic lobes (Fig. 1B,C). Weak staining of the antennal nerve appeared. However, no nerve tracts within the antennal lobe were visible. The labeling of the two main cell clusters, the lateral and the medial, clearly appeared in all preparations.

Comparison of the six analyzed antennal lobes showed minor differences in size. The maximal thickness, i.e., in the anterior–posterior direction (z-axis) of the antennal lobe of specimen 1 was 216 μm, 236 μm, and 264 μm for H. assulta male and H. virescens male and female, respectively, whereas specimen 2 measured 225 μm, 238 μm, and 228 μm, respectively. The high resolution of the raw data allowed individual delineation of and numbering of the glomeruli, as shown in Figures 2 and 3.
Reconstructions of the glomeruli in the *H. assulta* male

**MGC.** Three glomeruli located dorsolaterally in the antennal lobe, clearly separated from the array of ordinary glomeruli, form the MGC of the *H. assulta* male (Figs. 2B–D, 3B–D, 4C,D, 5A,D–F). They have previously been shown to receive information about “insect-produced” signals (Berg et al., 1999). The largest MGC unit, the cumulus, is positioned at the center of the antennal nerve entrance, between the dorsomedial and the ventral compartments. In following the optical sections in the direction of the z-axis from anterior to posterior, the cumulus appeared first, then the dorsomedial compartment, and finally the ventral compartment. The cumulus, appearing in frontal sections as a cone-shaped structure, measured 92 μm in maximal thickness in specimen 1, whereas the dorsomedial and the ventral compartments were 78 and 84 μm, respectively. The maximal thickness measured in each of the three MGC compartments of specimen 2 was similar (Table 1). The position of the MGC was about 50 μm posterior to the most anterior glomerulus.

**Ordinary glomeruli.** Sixty-two ordinary glomeruli were found in the male of *H. assulta* (Figs. 2, 3). Most of them were spheroid or slightly oblong, with a diameter in the range of 40–60 μm. One glomerulus particularly large in size, G52, was recognized as the so-called labial pit organ (LPO) glomerulus (Figs. 2C–F, 3D,E, 4D, 5C,D,G), elaborately described for *M. sexta* on the basis of its position as the most ventral in the antennal lobe (Kent et al.,...
1986). Mass staining of labial palp receptor neurons in the *H. virescens* female has resulted in heavy labeling of a glomerulus located at a corresponding position (Almaas personal communication). Another relatively large compartment, G34, was located adjacent to the LPO glomerulus, at the dorsomedial side (Figs. 2C–E, 3C,D, 4D, 5D). The unit G61 was also noteworthy in its position posterior to the dorsomedial MGC compartment (Figs. 2E,F, 3D,E, 5C). The smallest glomerulus, G16, was located in a frontolateral part of the antennal lobe (Figs. 2A, 5A) and had an axial thickness of only 16 μm (Table 1). Except for two units, all the glomeruli in specimen 2 were identified according to specimen 1 as shown in Figures 2 and 3. The array of ordinary glomeruli was highly stereotyped and showed only minor differences between individuals, for instance, in the relative position of G48 and G49 (Figs. 2F, 3E,F). The unidentified glomeruli were nn1, which was located posteroomedially in the antennal lobe, surrounded by G51, G55, G54, and G53 (Fig. 3E), and nn2, which was located posteriorly to G34 (Fig. 3E,F). Two of the glomeruli in the atlas, the smallest, G16 (Fig. 2A), and the medially located, G57 (Fig. 2D), were not found in specimen 2. Thus, the total number of counted glomeruli in the two specimens was the same, 65. Also by comparing the 3D reconstruction of specimen 2 (Fig. 4D) with the labelled atlas (Fig. 5A), the conformity of the glomerular pattern was demonstrated.

**Reconstructions of the glomeruli in the *H. virescens* male**

The MGC. The MGC of the *H. virescens* male appeared in an image section of 35 μm depth from the most anterior glomerulus, well separated from the array of the ordinary glomeruli (Fig. 4H, Table 2). As shown in Figure 4G, four compartments constitute this particular complex, previously described morphologically and functionally by Vickers et al. (1998) and Berg et al. (1998). The large cumulus is located close to the entrance of the antennal nerve, another large glomerulus dorsomedially of the cumulus, and two smaller compartments ventrally: the ventrolateral and the ventromedial. In the frontally scanned preparations, the four MGC compartments appeared in the following sequence from anterior to posterior: (1) the ventrolateral, (2) the ventromedial (3) the cumulus, and (4) the dorsomedial compartment. The size of the cumulus showed a maximal thickness of 116 μm in specimen 1, whereas the dorsomedial compartment measured 84 μm. The two ventrally located MGC compartments had the
size of the ordinary glomeruli, about 45 μm (Table 2). The maximal thickness measured in each of the four MGC compartments of specimen 2 was slightly smaller for the cumulus and the dorsomedial unit, whereas the two ventral compartments were similar (Table 2). Ordinary glomeruli. In addition to the four MGC compartments, 62 ordinary glomeruli were counted in the H. virescens male. Most of them showed a spherical structure with a diameter of 40–60 μm (Fig. 4E,H). One glomerulus, G54, located adjacent to the dorsomedial MGC compartment, was recognized by its relatively large size. With a maximal diameter of about 80 μm in specimen 1 and 65 μm in specimen 2, this particular unit resembled the large MGC compartments rather than the ordinary glomeruli (Table 2). Another glomerulus, G39, was easily recognized as the LPO unit based on its large size and ventral position in the antennal lobe (Fig. 4F,H). Adjacent and medially to the LPO unit was the relatively large glomerulus, G35 (Fig 4F,H). The smallest glomerulus, G23 (Fig. 4F,H), had a thickness of 28 and 25 μm in specimens 1 and 2, respectively (Table 2).

**Table 1. Position and Size of Eight Selected Glomeruli in the Antennal Lobe of the Helicoverpa assulta Male**

<table>
<thead>
<tr>
<th>Labeling</th>
<th>First sectiona</th>
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<th>z-Axis size3 (sp 1/sp 2)</th>
<th>Max xy size3 (sp 1/sp 2)</th>
<th>Volume4</th>
</tr>
</thead>
<tbody>
<tr>
<td>MGC: cumulus Cumulus</td>
<td>25 (50)</td>
<td>71 (142)</td>
<td>92/85</td>
<td>90/100</td>
<td>304.2</td>
</tr>
<tr>
<td>MGC: dorsomedial Dorosomial</td>
<td>29 (58)</td>
<td>68 (136)</td>
<td>78/79</td>
<td>65/75</td>
<td>135.0</td>
</tr>
<tr>
<td>MGC: ventral Ventrал</td>
<td>33 (66)</td>
<td>76 (150)</td>
<td>84/73</td>
<td>78/83</td>
<td>155.0</td>
</tr>
<tr>
<td>OG: MGC medial G61</td>
<td>70 (140)</td>
<td>98 (196)</td>
<td>56/52</td>
<td>65/50</td>
<td>96.0</td>
</tr>
<tr>
<td>OG: LPO G52</td>
<td>44 (88)</td>
<td>101 (202)</td>
<td>114/70</td>
<td>95/75</td>
<td>349.2</td>
</tr>
<tr>
<td>OG: LPO-medial G34</td>
<td>36 (72)</td>
<td>92 (184)</td>
<td>112/90</td>
<td>75/85</td>
<td>250.8</td>
</tr>
<tr>
<td>OG: smallest glom. G16</td>
<td>30 (60)</td>
<td>38 (76)</td>
<td>16/—</td>
<td>26/—</td>
<td>6.0</td>
</tr>
<tr>
<td>OG: typical G57</td>
<td>55 (110)</td>
<td>78 (156)</td>
<td>46/52</td>
<td>53/45</td>
<td>82.2</td>
</tr>
</tbody>
</table>

1Three MGC compartments (identified by name), four ordinary glomeruli (OG; labeled by number) that were easily recognized, and one randomly chosen, “typical” OG.
2The section numbers refer to the image planes, counted from the anterior to the posterior of the antennal lobe of specimen 1; for each glomerulus, the section of first and last appearance is indicated. The position (in μm) along the z-axes is given in parentheses.
3Thickness along the z-axis and maximum diameter in the x-y-plane, measured in specimen 1 (sp 1) and specimen 2 (sp 2) is given (in μm).
4The volume of specimen 1, given by 10^6 μm³, is calculated by the “tissue statistics” tool of Amira 2.1.

Fig. 3. Helicoverpa assulta male (specimen 2). Images of the antennal lobe in six different planes, showing each of the 63 identified and the two unidentified glomeruli. A: At 43 μm depth, the most anterior glomeruli are seen (the glomerulus G16 is not present). B: At 73 μm depth, two of the three MGC units [the cumulus (C) and the dorsomedial compartment (DM)] appear, residing outside the array of ordinary glomeruli. C: At 103 μm depth, all three MGC compartments [C, DM, and the ventral (V)] are visible. Both groups of cell bodies are present; the large lateral cell cluster (LCCl) and the smaller medial cell cluster (MCCl). D: At 134 μm depth, the LPO glomerulus (G52) can be seen with an adjacent large glomerulus (G34). The G57 is not present. E: At 164 μm depth, the two unidentified glomeruli nn1 and nn2 can be seen. F: At 195 μm depth, the most posterior glomeruli, including the G53 and G54, are shown. The glomeruli G48, G49, and G60 differ slightly compared with specimen 1, as shown in Figure 2F. AN, antennal nerve. Scale bar = 100 μm in A.
Fig. 4. Three-dimensional reconstructions of the antennal lobe of the *Helicoverpa assulta* male (left column), the *Heliothis virescens* male (middle column), and the *Heliothis virescens* female (right column). **A,E,I:** Data from the confocal stacks of each preparation (specimen 1), visualized by using the “volume rendering” tool of Amira. **B,F,J:** The digital atlases of each preparation, obtained by using the segmentation tool of Amira. Each glomerulus is given a distinct value, which is visualized as an individual color (The *H. virescens* atlases are labeled, whereas the *H. assulta* atlas is labeled in Fig. 5). **C,G,K:** Surface views of the male-specific MGCs and the large female glomeruli (LFG) obtained by segmentation. **D,H,L:** Data from the confocal stacks of each second preparation (specimen 2), visualized by using the “projection” tool of Zeiss LSM 510. The cLFG and the mLFG are named M1 and M2, respectively. Scale bars = 200 μm.
frontally located glomerulus, NN, found in specimen 2, is not present in the atlas (Fig. 4H).

Reconstructions of the glomeruli in the *H. virescens* female

Among the 62 glomeruli counted in the *H. virescens* female, two large compartments were found dorsolaterally in the antennal lobe. Their position close to the entrance of the antennal nerve corresponds to the location of the MGC in the male (Fig. 4K). Differently from the MGC, however, which is clearly separated from the ordinary glomeruli, these two compartments form a more integrated part of the shell-like array of ordinary glomeruli. The two units are named the *central large female glomerulus* (cLFG) and the *medial large female glomerulus* (mLFG), corresponding to the lateral and the medial large glomeruli of the *M. sexta* female (Rospars and Hildebrand, 2000). The cLFG, situated in the middle of the antennal nerve entrance, had a maximal thickness of 110 μm in specimen 1, whereas the medial compartment measured 102 μm (Table 3). The large female glomeruli appeared in a section at 125 μm depth from the most anterior glomerulus of the antennal lobe. The position of the two compartments, here appearing much deeper than the MGC of the males, is probably caused by the more ventral orientation of the preparation.

Most of the 60 remaining glomeruli counted in the *H. virescens* female were spheroid or ellipsoid, with a diameter of 40–60 μm (Fig. 4L). Also in the female the large LPO glomerulus, G19, located ventrally in the antennal lobe, was recognized (Fig. 4J,L). A large glomerulus located medially to the LPO compartment was the G20 unit. The smallest glomerulus, G3, measured 40 μm in specimen 1 and 35 in specimen 2 (Table 3). Altogether, specimen 2 contained 65 glomeruli, i.e., three more than the atlas. Thirty-two of them were identified according to the atlas and included, in addition to the two LFG compartments, glomeruli G1–G22, G25, G27, G29, G34, G35, G57, G58, and G60. All of them are shown in Figure 4L. Although the sizes of the glomeruli G19, G20, and G3 were similar in the two specimens, the two LFG compartments (cLFG and mLFG) were considerably smaller in specimen 2, measuring 72 and 68 μm, respectively (Table 3).

Reconstructions of the lateral and the medial cell cluster of the antennal lobe

In all six preparations, the two main cell clusters of the antennal lobe were easily recognized as densely packed somata located at the lobe periphery (Figs. 2, 3). The largest, the lateral cell cluster, appeared in one of the first image planes, continuing in a ventrolateral position throughout the antennal lobe. This large assembly of cell bodies sends its neurites in a bundle penetrating the outer shell of glomeruli. The considerably smaller medial cell cluster appears more posteriorly in the antennal lobe (Figs. 2C, 3C, Table 4).

**DISCUSSION**

“...There is nothing casual in its [the universal organism] life but a single melody and order...” (Armstrong, 1984). This quotation of Plotinus comes to mind when considering the appearance of the discrete neuropil structures forming the glomeruli of the primary olfactory center of most animals. Indeed, the well-organized system of distinct glomeruli in the antennal lobe of the heliothine moth seems to reflect the truth of these words. The use of an antibody recognizing synaptic structures resulted in distinct staining of the glomeruli in the two moth species investigated here. Thus, the glomerular units are visualized as condensed collections of synapses, according to the definition of glomeruli as sites of increased synaptic density (Shepherd, 1974). The remaining area of the deutocerebrum, the antennal mechanosensory and motor center, was also strongly labeled but lacked the glomerular organization. This is in accordance with previous anatomical studies of the two main deutocerebral divisions (Rospars, 1988; Homberg et al., 1989; Kloppenburg et al.,

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**TABLE 2. Position and Size of Nine Selected Glomeruli in the Antennal Lobe of the Heliothis virescens Male**

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<th>Max xy size</th>
<th>Volume</th>
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</thead>
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<tr>
<td>MGC: cumulus</td>
<td>15 (60)</td>
<td>44 (176)</td>
<td>116/83</td>
<td>110/99</td>
<td>481.2</td>
</tr>
<tr>
<td>MGC: dorsomedial</td>
<td>19 (76)</td>
<td>40 (160)</td>
<td>84/71</td>
<td>72/65</td>
<td>229.8</td>
</tr>
<tr>
<td>MGC: ventrolateral</td>
<td>10 (40)</td>
<td>22 (88)</td>
<td>48/42</td>
<td>55/45</td>
<td>75.6</td>
</tr>
<tr>
<td>MGC: ventromedial</td>
<td>12 (48)</td>
<td>22 (88)</td>
<td>40/57</td>
<td>48/40</td>
<td>48.0</td>
</tr>
<tr>
<td>OG: MGC medial</td>
<td>G54</td>
<td>G74</td>
<td>47 (188)</td>
<td>60/58</td>
<td>164.0</td>
</tr>
<tr>
<td>OG: LPO</td>
<td>G39</td>
<td>G59</td>
<td>51 (204)</td>
<td>104/68</td>
<td>389.4</td>
</tr>
<tr>
<td>OG: LPO-medial</td>
<td>G35</td>
<td>G44</td>
<td>42 (168)</td>
<td>64/55</td>
<td>386.0</td>
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<td>OG: smallest glom.</td>
<td>G23</td>
<td>G32</td>
<td>16 (64)</td>
<td>29/25</td>
<td>26.4</td>
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<tr>
<td>OG: typical</td>
<td>G5</td>
<td>G5</td>
<td>13 (52)</td>
<td>44/34</td>
<td>66.0</td>
</tr>
</tbody>
</table>

1 Four MGC compartments (identified by name), three ordinary glomeruli (OG; labeled by number) that were easily recognized, the smallest glomerulus, and one randomly chosen, “typical” OG.

2 The section numbers refer to the image planes, counted from the anterior to the posterior of the antennal lobe of specimen 1; for each glomerulus, the section of first and last appearance is indicated. The position (in μm) along the z-axis is given in parentheses.

3 Thickness along the x-axis and maximum diameter in the xy-plane are given (in μm), measured in specimen 1 (sp 1) and specimen 2 (sp 2).

4 The volume of specimen 1, given by \( \text{Volume} = \text{Max xy size} \times \text{z-Axis size} \), is calculated by the “tissue statistics” tool of Amira 2.1.
The staining of the cell clusters, in addition to the somewhat weaker labeling of the antennal nerve, was beneficial, in that these structures represent reliable landmarks for orientation. Labeling of cell somata and axons is probably due to the presence of synapsin intracellularly in these structures. It has been shown in vertebrates that both structures are stained by high concentrations of synapsin antibodies, whereas lower concentrations stain only the axon terminals.

The small heliothine brain is advantageous for confocal microscopy; the entire antennal lobe, as well as the whole brain, can be visualized as a single stack of sequential optical images. The efficiency of the staining technique and the resolution of the images, showing an interglomerular space of 2 μm in some sections (Fig. 2), made it possible to outline the borders of the individual glomeruli. A similar technique, using a neuropil-specific antibody, has been used for the computerized map of the Drosophila antennal lobe (Laisisse et al., 1999), whereas the honeybee atlas was based on a different staining technique, tracing the sensory axons with a specific dye (Galizia et al., 1999). Labeling of the primary axons, which was also attempted in the present study, was insufficient for segmentation. A combination of different staining techniques may further improve the visualization of the brain structures in heliothine moths. It would be of particular interest to resolve the antennal tracts in order to determine their innervation of the glomeruli. In the honeybee, the antennal nerve splits into four tracts that innervate the antennal lobe. These tracts have been used as the basis for the nomenclature of the glomeruli (Flanagan and Mercer, 1988). For the moth, such a system is not yet available. A list of five antennal tracts has been described for M. sexta, but without mapping their innervation pattern to the glomeruli (Rospars and Hildebrand, 1992). The terminology used in the fly atlas is based on the exact position of the 43 glomeruli, i.e., their position according to all three axes (x,y,z; Stocker et al., 1990; Laisisse et al., 1999). Because of the relatively large numbers of units and the lack of visualized tracts, we found that it was appropriate in this study of heliothine moths to use the more incidental system of labeling the glomeruli by numbers according to their position along the z-axis.

Each of the three digital atlases presented here is based on the reconstruction of one antennal lobe. Analysis carried out by comparing a second specimen with the original indicates only small variations of the glomerular pattern within each of the three types. The number of counted glomeruli was identical in the two male individuals of *H. assulta* (65) as well as of *H. virescens* (66) and differed only by three in the two female individuals of *H. virescens* (62/65). A relatively consistent glomerular pattern was also found, showing only minor differences concerning one or two glomeruli in each type. Interestingly, the smallest glomerulus, G16, in the *H. assulta* male atlas was not found in specimen 2. The results presented here are in accordance with previous findings showing that the number and position of the glomeruli are relatively invariant across individuals of the insect species (Rospars and Hildebrand, 1992, 2000; Galizia et al., 1999). Whereas small variations in the array of the numerous ordinary glomeruli were present, the number, position, and shape of the MGC compartments were strictly constant within the species, which is also consistent with previous findings in heliothine moths (cf. Mustaparta, 1997). It is interesting that the number of counted glomeruli in the six specimens is relatively constant, 62, 65, 65, 65, 66, and 66. This is in the range of the number found in other moth species, for instance, *M. sexta*, with 63 glomeruli (Rospars and Hildebrand, 2000), and *Mamestra brassicae*, with 67 male and 68 female glomeruli (Rospars, 1983).

Some glomeruli can be considered as homologous in different species. This particularly applies to the two MGC compartments, the cumulus and the dorsomedial unit, in males of *H. virescens* and *H. assulta* as well as of other heliothine species (Christensen et al., 1991; Vickers et al., 1997).
1998; Berg et al., 1998; T.J. Almaas, unpublished observations). In addition to the anatomical correspondence, the functional aspects show interesting similarities. The cumulus, located close to the entrance of the antennal nerve, always receives input about the major pheromone component (Berg et al., 1998). It is the same chemical, cis-11-hexadecenyl, in all heliothine species studied so far (Arn et al., 1992), except for H. assulta, which uses cis-9-hexadecenyl (Cork et al., 1992). The dorsomedial compartment always receives input about another compound, cis-9-tetradecenyl. However, in this case, the message differs, being a prerequisite for pheromone attraction in H. virescens (Roelofs et al., 1974) and an interruptant of the attraction in H. assulta (Boo et al., 1995), Helicoverpa zeae (Shaver et al., 1982), and Helicoverpa armigera (Kehat and Dunkelblum, 1990). Thus, the dorsomedial compartment has the same location and receives input information about the same chemical in all heliothine species investigated but differs with respect to the behavior it elicits when excited by an olfactory stimulus.

Homologous glomerular structures may also be present in the heliothine females. In the one female investigated here, of H. virescens, two enlarged compartments were shown at the entrance of the antennal nerve, in a position corresponding to the male-specific MGC, the largest residing centrally at the antennal nerve entrance and a somewhat smaller unit in the dorsomedial position. Large female glomeruli have previously been described for the silk moth Bombyx mori (Koontz and Schneider, 1987) as well as for M. sexta (Rössler et al., 1998). It has recently been reported that one of the two units identified in the latter mediates information specifically about the plant volatile linalool (King et al., 2000). The function of the large female glomeruli in H. virescens is not yet clarified.

Among the ordinary glomeruli, the LPO compartment was particularly easy to recognize in all six specimens (H. assulta male: G52 (Figs. 2, 3, 4D, Table 1), H. virescens male: G39 (Fig. 4F,H, Table 2), H. virescens female: G19 (Fig. 4J,L, Table 3)). This is the largest ordinary glomerulus, residing most ventrally in the antennal lobe. Another large glomerulus, found in all specimens, was located adjacent to the LPO unit, on the medial side (H. assulta male: G34 (Figs. 2, 3, 4D, Table 1), H. virescens male: G35 (Fig. 4F,H, Table 2), H. virescens female: G20 (Fig. 4J,L, Table 3)). A glomerulus situated adjacent to the dorsomedial MGC compartment, in the posterior part of the antennal lobe, was found in the four male preparations (H. assulta male: G61 (Figs. 2, 3, Table 1), H. virescens male: G54 (Table 2)). Particularly in H. virescens this unit, G54, was conspicuous by its large size. Interestingly, a large glomerulus occupying a location next to the MGC has also been identified in both sexes of M. sexta. It is noteworthy that the number of ordinary glomeruli in the male and female of H. virescens was similar (62 and 62/65, respectively). According to intersexual homology, a recent study of the antennal lobe anatomy of adult male and female M. sexta has shown that 60 ordinary glomeruli are isomorphic in the two sexes, whereas three are dimorphic (Rospars and Hildebrand, 2000).

The manner in which the glomeruli in general operate as functional units is not yet fully understood for any organism. However, results from various studies indicate that they may represent a physical basis for a functional map of odor qualities. The use of transgenic D. melanogaster has recently shown that receptor neurons expressing one type of receptor molecules project exclusively to one (or sometimes two) homologous glomerulus in each antennal lobe (Gao et al., 2000; Vossshall et al., 2000). These and other data from molecular biological studies (Ressler et al., 1994; Vassar et al., 1994; Mombaerts et al., 1996; Wang et al., 1998) are in accordance with the results obtained for the pheromone system in the heliothine moth, showing that neurons of one functional type project to one MGC unit (Hansson et al., 1992; Berg et al., 1998). In addition, data from calcium imaging of the antennal lobe of the honeybee have demonstrated that different odor qualities induce a specific pattern of glomerular activation (Joerges et al., 1997; Galizia and Menzel, 2000). By using the same imaging technique, a pattern of glomerular activation of the MGC that corresponds to the functional organization demonstrated by tracing of receptor neurons has recently been shown for H. virescens (Galizia et al., 2000). Also in this case a specific activation pattern of the ordinary glomeruli was obtained by stimulation with different plant odors. To study in detail which of the 60–62 ordinary glomeruli are activated by the various biologically relevant plant odors in the antennal lobe of heliothine moths, it is necessary to have an atlas of the glomeruli in these species.

ACKNOWLEDGMENTS

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LITERATURE CITED


APPENDIX. Nomenclature of Individual Glomeruli (The Value of Each Voxel Gives the Identity of the Glomerulus)

<table>
<thead>
<tr>
<th>Helicoverpa assulta, male</th>
<th>Heliosis virescens, male</th>
<th>Heliosis virescens, female</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Ordinary glomeruli</strong></td>
<td><strong>Ordinary glomeruli</strong></td>
<td><strong>Ordinary glomeruli</strong></td>
</tr>
<tr>
<td>MGC/LFG</td>
<td>G1-062 as 1–82</td>
<td>G1–G60 as 1–60</td>
</tr>
<tr>
<td></td>
<td>Cumulus: 102</td>
<td>Cumulus: 101</td>
</tr>
<tr>
<td></td>
<td>Dorsomedial: 102</td>
<td>Dorsomedial: 102</td>
</tr>
<tr>
<td></td>
<td>Ventral: 103</td>
<td>Ventral: 104</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Ventrataleral: 104</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Ventromedial: 105</td>
</tr>
<tr>
<td>Medial cell cluster</td>
<td>201</td>
<td>201</td>
</tr>
<tr>
<td>Lateral cell cluster</td>
<td>202</td>
<td>202</td>
</tr>
<tr>
<td>Antennal nerve</td>
<td>203</td>
<td>203</td>
</tr>
</tbody>
</table>

Format of the atlas on the internet: http://www.neurobiologie.fu-berlin.de/mothatlasy/ or http://kocz.chemio.ntna.no/mothAL2. Each atlas is a volumetric representation of a right antennal lobe. It consists of a series of sections. In each section, the position belonging to a particular glomerulus is identified by its color code (or gray-level value). By using any software capable of 3D graphics, it is possible to resolve and/or rotate the atlas. Most software packages have a tool giving the color/table value of a chosen voxel (= pixel in three dimensions). This value reveals the identity of the glomerulus, following the key given in the Appendix. On the web site, examples of the atlas as well as the raw confocal data can be seen as Quicktime movies. To work with the atlas, it is best to download the data onto a local computer. The atlas can then be used with one’s own software. Basically, each atlas consists of a series of 2-axis sections, where each glomerulus is identified by a value (color). Because not all software can read all data formats, we provide the identical data in different formats. It is not necessary to download all the material. Rather, find out what your software can read, and then download that particular format.

We have included, for each species, the following formats: three-dimensional TIFF file, compressed and not compressed; three-dimensional GIF file; Amira-mesh file (readable only with the amira software); individual TIFF files for each layer; individual GIF files for each layer; individual BMP files for each layer. The atlas will soon be accessible also from the formation of a precise topographic map. Cell 79:1245–1255.


