A comprehensive anatomical map of the peripheral octopaminergic/tyraminergic 3 system of *Drosophila melanogaster*

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33 Abstract

34 The modulation of an animal's behavior through external sensory stimuli, previous 35 experience and its internal state is crucial to survive in a constantly changing 36 environment. In most insects, octopamine (OA) and its precursor tyramine (TA) 37 modulate a variety of physiological processes and behaviors by shifting the organism 38 from a relaxed or dormant condition to a responsive, excited and alerted state. Even though OA/TA neurons of the central brain are described on single cell level in 39 40 Drosophila melanogaster, the periphery was largely omitted from anatomical studies. 41 Given that OA/TA is involved in behaviors like feeding, flying and locomotion, which 42 highly depend on a variety of peripheral organs, it is necessary to study the peripheral 43 connections of these neurons to get a complete picture of the OA/TA circuitry. We here 44 describe the anatomy of this aminergic system in relation to peripheral tissues of the 45 entire fly. OA/TA neurons arborize onto skeletal muscles all over the body and innervate 46 reproductive organs, the heart, the corpora allata, and sensory neurons in the antenna, 47 wings and halteres underlining their relevance in modulating complex behaviors. 48

49 Introduction

50 The adrenergic system of mammals influences various aspects of the animal's life. Its 51 transmitters/hormones, adrenaline and noradrenaline, modulate a variety of 52 physiological processes and behaviors. They are secreted into the bloodstream by the 53 adrenal glands in response to stress. In addition, they are synthesized and released by 54 axonal terminals in the central nervous system (CNS) as well as sympathetic fibers of 55 the autonomic nervous system. Adrenaline and noradrenaline have been described as 56 modulators to shift the organism from a relaxed or dormant state to a responsive, 57 excited and alerted state ¹. Stressful stimuli induce a metabolic and behavioral 58 adaptation, leading to enhanced energy supply, increased muscle performance, 59 increased sensory perception and a matched behavior. This so-called "fight or flight" 60 response can be seen in vertebrates and invertebrates. In insects, the stress response 61 is mediated - among others - by octopamine (OA) and its precursor tyramine (TA) ²⁻⁴. TA 62 is synthesized from tyrosine by the action of a tyrosine decarboxylase enzyme (Tdc) and 63 functions as an independent neurotransmitter/-modulator as well as the intermediate 64 step in OA synthesis. For this, TA is catalyzed by the tyramine-ß-hydroxylase (TßH).

65 Similar to the vertebrate adrenergic system, OA and TA act through specific G-protein 66 coupled receptors. Besides structural similarities between OA/TA and 67 adrenaline/noradrenaline and the corresponding receptors, functional similarities are illustrated by the action of these transmitters/hormones in the regulation of physiological 68 69 processes and behaviors. OA and TA are known to modulate muscle performance, 70 glycogenolysis, fat metabolism, heart rate, and respiration in insects (reviewed by: ⁵). 71 While the role of TA as an independent signaling molecule was underestimated for a 72 long time, OA has been extensively studied and was shown to have effects on almost 73 every organ, sensory modality and behavior in a great variety of insects. The most 74 intensively studied peripheral organs regarding the modulatory role of OA are muscles 6-75 ¹⁰. Here, OA is thought to not exclusively modulate muscle performance or motor 76 activity. OA rather modulates muscle action according to metabolic and physiological 77 processes, for example by promoting energy mobilization directly from the fat body, or 78 indirectly by promoting the release of adipokinetic homones (AKH) from neuroendocrine 79 cells in the corpora cardiaca (CC, a homolog of the vertebrate anterior pituitary gland and an analog of mammalian pancreatic alpha cells) ^{11,12}. In addition to the impact of 80 81 OA/TA on muscles, fat body and AKH cells, OA is shown to modulate the heart, trachea 82 and air sacs, gut, hemocytes, salivary glands, Malpighian tubules and ovaries in insects, 83 mainly to induce a general stress or arousal state. However, in total OA seems to 84 modulate a vast number of behaviors, which are not necessarily coupled to stress responses. The OA/TA system is shown to also act on i.a. learning and memory, sleep, 85 feeding, flight, locomotion, and aggression ^{8,10,12–35}. 86

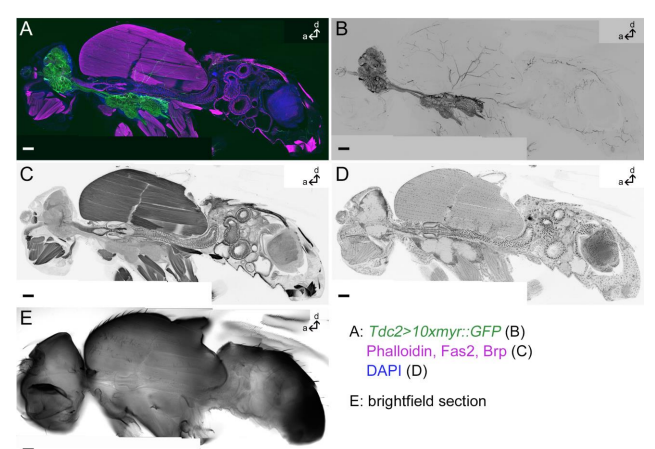
As mentioned above, OA and TA act as neurotransmitters and neuromodulators, 87 88 allowing them to act in a paracrine, endocrine or autocrine fashion. In the fruit fly 89 Drosophila, huge efforts were made to describe OA/TA neurons (OANs/TANs) in the brain and ventral nervous system (VNS) down to the single cell level ^{8,16,36–40}. 90 91 Nevertheless, although our knowledge about physiological processes and behaviors 92 modulated by the OA/TA system in the brain is rich, less is known about how OA and TA 93 reach all its target organs and tissues in the periphery (exceptions: reproductive organs ^{36,40–43} and muscles ^{8,44–46}). 94

95 Here we use the genetically tractable fruit fly Drosophila melanogaster to describe the 96 arborizations of Tdc2-Gal4-positive, and therefore OANs and TANs in the periphery, as 97 the Drosophila Tdc2 gene is expressed neurally ⁴⁰. We found that OANs/TANs are 98 widespread distributed throughout the fly's body with innervations in the skeletal 99 muscles, reproductive organs, corpora allata, antenna, legs, wings, halteres and the 100 heart. This diverse innervation pattern reflects the modulatory role of OA/TA in many 101 different behaviors and physiological processes. Our results provide, for the very first 102 time, a complete and comprehensive map of the OA/TA circuitry in the entire insect 103 body. This map allows assumptions about the type of OA/TA signaling (paracrine or 104 endocrine) to a specific organ and, at the same time, it provides a deeper understanding 105 to what extend the OA/TA-dependent activity of peripheral organs is altered, for example 106 by genetically manipulating Tdc2-Gal4-positive neurons in the brain and VNS.

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108 Results

109 The OANs/TANs of the brain and ventral nervous system (VNS) are described in detail even on single cell level in *Drosophila*^{10,37–39,44}. In contrast little is known about their 110 111 peripheral arborizations. We used the well-characterized Tdc2-Gal4 line to deepen our knowledge about the OA/TA system in the entire body of *Drosophila* ^{38–41,47}. Nearly all 112 113 *Tdc2-Gal4*-positive cells in the brain are stained by a Tdc2 antibody ⁴⁷. In the VNS all of 114 the *Tdc2-Gal4*-positive cells were labeled by a TβH antibody and therefore have to be 115 Tdc2-positive ³⁹. We here expressed myristoylated GFP, enhanced by GFP antibody 116 staining, to label the membranes of Tdc2-Gal4-positive neurons from the soma to its fine 117 endings in the periphery. The peripheral organs, tissues and cells are visualized by 118 fluorescent markers for cell bodies (DAPI binds to DNA), muscles (Phalloidin binds F-119 actin) and antibodies against the synaptic protein Bruchpilot and the cell adhesion 120 molecule Fasciclin 2 (Fig. 1).



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Fig. 1: Tdc2-Gal4-positive arborizations in a whole fly section. Projection of one
medial sagittal agarose section of 80μm thickness labeled by anti-GFP to visualize
membranes of Tdc2-Gal4-positive neurons (green in A, black in B); Phalloidin, antiFasciclin2 (Fas2) and anti-Bruchpilot (Brp) to visualize muscles, cells and synapses,
respectively (magenta in A, black in C) and DAPI to mark cell bodies (blue in A, black in
D). E: A single optical section showing the bright-field picture. Scale bars = 50 μm.

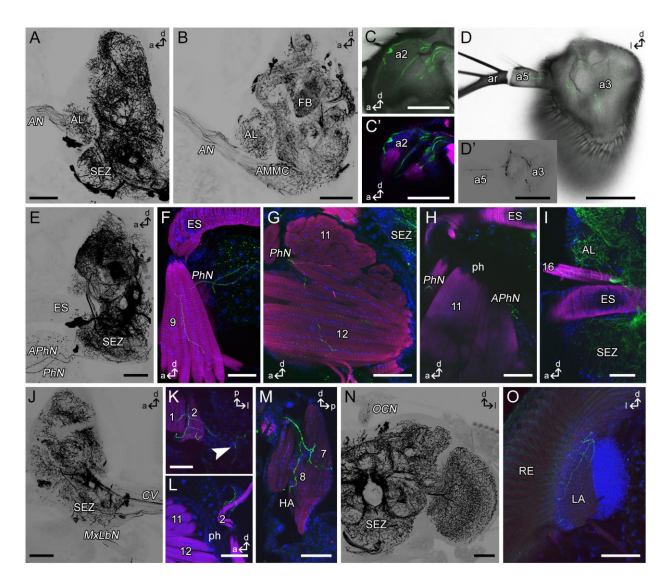
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129 *Tdc2-Gal4*-positive arborizations in the head

130 *Tdc2-Gal4*-positive neurons (Tdc2N) project through all peripheral nerves of the brain:

131 antennal nerve (AN), ocellar nerve (OCN), pharyngeal nerve (PhN) and accessory PhN

- 132 (APhN), maxillary-labial nerve (MxLbN), corpora cardiaca nerve (NCC), and the cervical
- 133 connective (CV) (Fig. 2). Tdc2Ns in the antennal nerve are connected to the antennal
- 134 lobe and the antennal mechanosensory and motor center (ammc), respectively, and give
- rise to staining in the pedicle- the Johnston's organ (JO)-, funiculus and arista of the
- antenna (Fig. 2A-D). While no cell bodies are visible in the third to fifth segment of the
- 137 antenna, the JO contains stained cell bodies indicating that the *Tdc2-Gal4* line includes
- 138 mechanosensory neurons (Fig. 2C).



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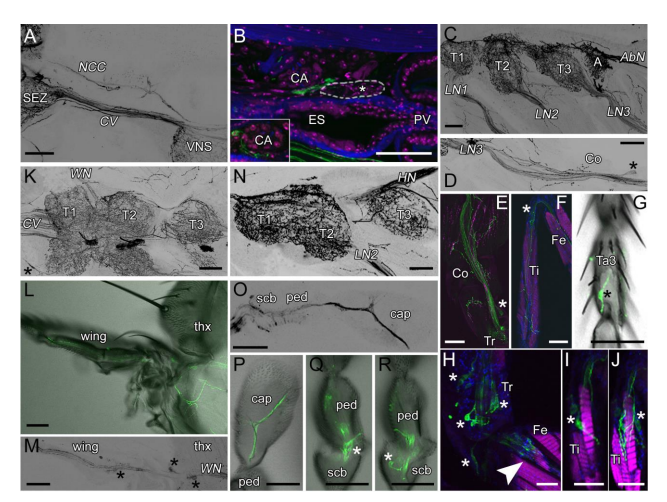
140 Fig. 2: Tdc2-Gal4-positive arborizations in the fly's head. Projections of sagittal (A-141 C,E-J,M) or frontal (D,K,O) or horizontal (L) optical sections visualizing the arborization 142 pattern of Tdc2-Gal4-positive neurons (Tdc2N; black or green) in the head. A-D: Tdc2Ns 143 run through the antennal nerve (AN) and project in antennal segments a2, a3 and a5. 144 Mechanosensory neurons of the Johnston's organ are visible (C). E-H: Efferent Tdc2Ns 145 of the pharyngeal (PhN) and accessory pharyngeal nerve (APhN). F-G: Cells of the PhN 146 innervate muscles 9, 11 and 12, H: Bouton-like structures of APhN neurons beside the 147 pharynx (ph). I: Innervation of muscle 16. J-M: Tdc2Ns of the maxillary-labial nerve 148 (MxLbN) project along muscle 1 and 2 (K, L) in the haustellum (HA) and seem to 149 innervate muscles 7 and 8 (M). Arborizations from the MxLbN reach the lateral brain 150 (arrowhead in K). N: Tdc2Ns arborize in the ocellar nerve (OCN). O: Ramifications in the 151 lateral lamina (LA) close to the retina (RE). a, anterior; AL, antennal lobe; AMMC, 152 antennal mechanosensory and motor center; ar, arista; d, dorsal; CV, cervical 153 connective; FB, fan-shaped body; I, lateral; p, posterior; SEZ, subesophageal zone. 154 Scale bars = $50 \ \mu m$.

156 The Tdc2-Gal4-positive efferent nerves of the subesophageal zone (SEZ) arborize in the 157 rostrum of the proboscis mainly onto muscles (Fig. 2E-M). Cells leaving the brain via the PhN innervate muscles 9, 10, 11 and 12 (nomenclature after ⁴⁸; Fig. 2F,G). Cells 158 159 projecting towards the APhN build bouton like structures beside muscle 11 ventral to the 160 pharynx (ph; Fig. 2H). Cells of the MxLbN arborize along muscles 1 and 2 (Fig. 2J-L). It 161 seems as if also muscles 7 and 8 of the haustellum are innervated (Fig. 2M). We only 162 observed this staining in two different specimens. Due to our cutting technique we 163 probably lost these parts of the haustellum frequently. In addition to the innervation of 164 the proboscis muscles we observed arborizations in the ventrolateral head arising from 165 the MxLbN (arrowhead Fig. 2K). The ocellar nerve, which connects the ocellar ganglion 166 with the brain, contains fibers arising from the brain (Fig. 2N). The central brain and optic 167 lobes were shown to contain a dense network of OANs/TANs ^{37,38}. In addition, we 168 identified arborizations in the distal part of the lamina by Tdc2Ns (Fig. 20). Muscle 16, 169 which is located dorsal to the esophagus, is innervated via ascending Tdc2Ns from the 170 thorax (Fig. 2I).

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172 *Tdc2-Gal4*-positive arborizations in the thorax

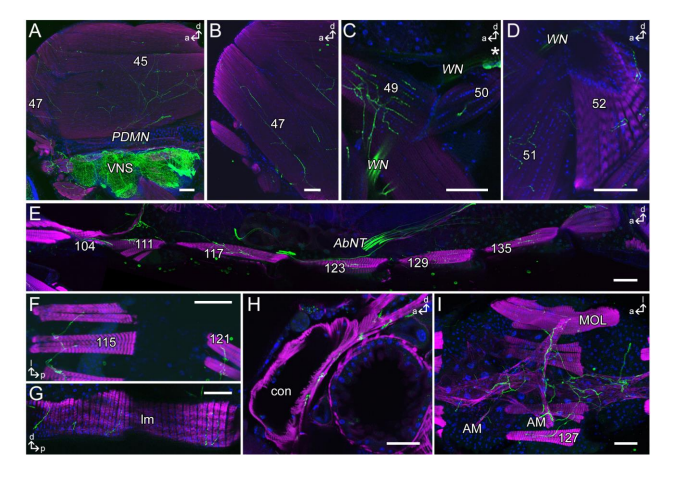
173 OANs/TANs form connections between the head and thorax via the CV and NCC (Fig. 174 3A). Tdc2Ns running through the NCC arborize close to the corpora allata (CA; Fig. 3B) 175 and anterior stomatogastric ganglion, while no staining is visible in the corpora cardiaca 176 (CC: asterisk Fig. 3B). The CV connects the brain and VNS and contains many Tdc2Ns 177 (Fig. 3A). All peripheral nerves of the thoracic ganglion seem to contain Tdc2-Gal4-178 positive axons. Most prominent are the paired leg nerves of each thoracic neuromere (LN1-3 Fig. 3C; ProLN, MesoLN, MetaLN after ⁴⁹), the paired wing (WN Fig. 3K; ADMN 179 180 after ⁴⁹) and posterior dorsal mesothoracic nerve (*PDMN*; Fig. 4A) of the mesothoracic 181 neuromere and paired haltere nerves of the metathoracic neuromere (HN Fig. 3N: 182 DMetaN after ⁴⁹). Interestingly, all these nerves, with the exception of *PDMN*, seem to 183 contain efferent Tdc2Ns innervating mainly muscles as well as afferent Tdc2-Gal4-184 positive sensory neurons.



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187 Fig. 3: Tdc2-Gal4-positive arborizations in the thorax. Projections of optical sections 188 visualizing the arborization pattern of Tdc2-Gal4-positive neurons (Tdc2Ns: black or 189 green) in the thorax. A: Tdc2Ns run through the cervical connective (CV) and corpora 190 cardiaca nerve (NCC). B: Tdc2Ns arborize close to the corpora allata (CA) and the 191 anterior stomatogastric ganglion (white-rimmed). C-F: Tdc2Ns project along the legs and 192 innervate leg muscles. G: An afferent sensory neuron in the third segment of the tarsus. 193 H-J: Cell bodies of sensory neurons (asterisks) of the trochanter (Tr; H) and tibia (Ti; I,J). 194 H: Neurons of the chordotonal organ in the femur (Fe; arrowhead). K-M: Tdc2Ns project 195 along the wing nerve (WN). K: Innervation of the thoracic chordotonal organ (asterisk). 196 L,M: Tdc2Ns run along the L1 wing vein. Cell bodies of sensory neurons are visible 197 (asterisks). N-R: Tdc2Ns in the haltere nerve (HN). Tdc2-positive cells project to the 198 distal part of the capitellum (cap). Sensory neurons of the pedicellus (ped) and 199 scabellum (scb) are labeled by Tdc2-Gal4. A-J,N-R: sagittal sections; K: horizontal 200 sections; L.M: frontal sections. A. abdominal segment; Co. coxa; ES. esophagus; Fe. 201 femur; LN, leg nerve; PV, proventriculus; SEZ, subesophageal zone; T1-3, thorax 202 segment1-3; Ta, tarsus; thx, thorax; Ti, tibia; Tr, trochanter. Scale bars: A-G,K-R = 50 203 μm ; H-J = 25 μm .

205 The efferent Tdc2Ns in LN1-3 arborize on the leg muscles down to the tibia (Fig. 3E,F). 206 while afferent fibers originate from sensory neurons of all leg segments (asterisks Fig. 207 3D-J), including i.a. mechanosensory neurons of the chordotonal organ of the femur 208 (arrowhead Fig. 3H) and campaniform sensilla of the tarsus (asterisk Fig. 3G). The WN 209 contains Tdc2Ns arborizing on indirect and direct flight muscles (Fig. 4B-D) and afferent 210 axons from sensory neurons of the proximal wing (asterisks Fig. 3M, 4C). Moreover, 211 efferent Tdc2Ns running to the PDMN innervate all six longitudinal indirect flight muscles 212 (45a-f; Fig. 4A) and the posterior dorsal-ventral indirect flight muscles (46a-b). Tdc2Ns 213 project along the L1 wing vein (Fig. 3L). Tdc2-positive cells innervating the haltere 214 project to the most distal tip of the capitellum (cap; Fig. 30,P). Additionally, Tdc2-Gal4 215 includes sensory neurons of campaniform sensilla of the pedicellus (ped; Fig. 3Q,R) and 216 scabellum (scb; Fig. 3R). Additionally, it seems that Tdc2-Gal4 labels sensory neurons 217 of the chordotonal organs of the haltere and wing.

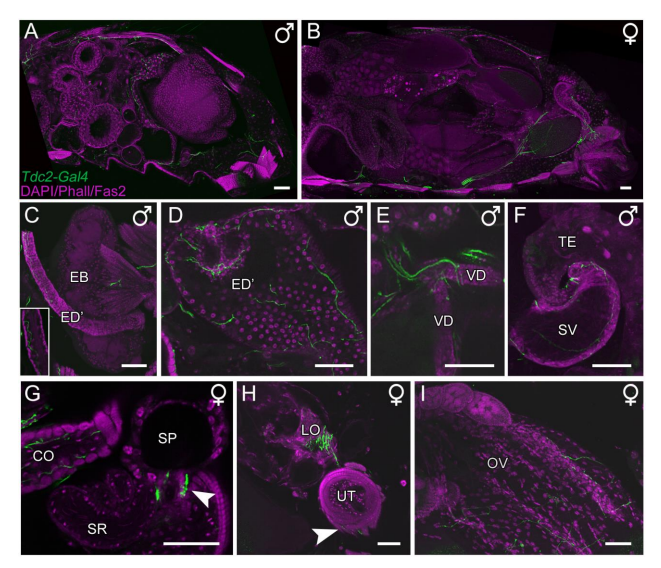


220 Fig. 4: Tdc2-Gal4-positive innervation of skeletal muscles. Projections of optical 221 sections visualizing the innervation pattern of Tdc2-Gal4-positive neurons (Tdc2Ns; 222 black or green) on skeletal muscles of thorax and abdomen. A-D: Innervation of the 223 indirect flight muscles (A,B) and direct flight muscles (C,D). E-H: Tdc2Ns innervate the 224 ventral (E), dorsal (F) and lateral (G) abdominal body wall muscles. H,I: The longitudinal 225 (H) as well as the alary muscles (I) of the heart are innervated by Tdc2Ns. I: In males 226 arborizations on the muscle of Lawrence (MOL) in segment 5 are visible. A-E,G,H: Sagittal sections; F.I: dorsal sections. a, anterior: AbNT, abdominal nerve trunk; AM, 227 228 alary muscle: con: conical chamber: d. dorsal: l. lateral: lm. lateral muscles: MOL. 229 muscle of Lawrence; p, posterior; PDMN, posterior dorsal mesothoracic nerve; VNS, 230 ventral nervous system; WN, wing nerve. Scale bars = $50 \mu m$.

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232 Tdc2-Gal4-positive arborizations in the abdomen

233 Tdc2Ns innervate all ventral (111,117,123,129,135; Fig. 4A) and dorsal skeletal muscles 234 (109,115,121,127,133; Fig. 4B) of abdominal segments 2-6 as well as lateral muscles (Fig. 4C). Additionally, Tdc2-Gal4-positive ramifications on the male specific "muscle of 235 236 Lawrence" in segment 5 are visible (Fig. 4E). Beside the body wall muscles, the ventral 237 longitudinal and the alary muscles of the heart are innervated (Fig. 4D,E). Tdc2Ns 238 running along the abdominal nerve trunk (AbNT) innervate the female and male 239 reproductive organs, respectively (Fig. 5). In males, as described before ⁴¹ the anterior 240 ejaculatory duct, the vas deferens and seminal vesicle are innervated, while the 241 ejaculatory bulb itself is not innervated but its muscles (Fig. 5C-F). The innervations of 242 the female oviducts, uterus and spermathecal duct have been described in previous publications ^{36,40–44}. 243



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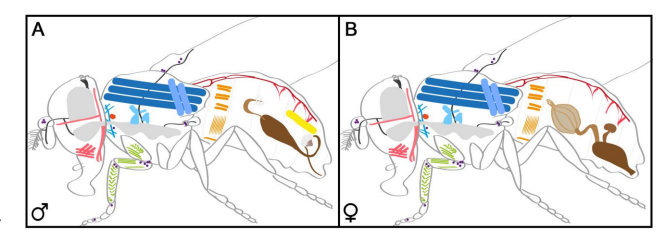
Fig. 5: Tdc2-Gal4-positive innervation of the reproductive organs. Projections of 246 247 optical sections visualizing the innervation pattern of Tdc2-Gal4-positive neurons 248 (Tdc2Ns; green) of reproductive organs visualized by DAPI, Phalloidin and Fasciclin2 249 staining (magenta). A-B: Tdc2N arborization pattern in a sagittal section of a male (A) 250 and female (B) abdomen, respectively. C-F: The anterior ejaculatory duct (ED'), muscles 251 of the ejaculatory bulb (EB), the vas deferens (VD) and seminal vesicle (SV) are innervated by Tdc2Ns. G-I: Tdc2Ns arborize onto the common and lateral oviduct (CO, 252 253 LO), spermathecal duct (arrowhead in G), uterus (UT) muscles (arrowhead in H) and the 254 ovaries (OV). SP, spermatheca; SR, seminal receptacle; TE, testes. Scale bars = 50 255 μm. 256

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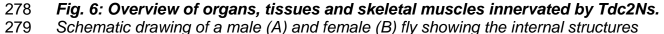
261 Discussion

262 Here we show a comprehensive description of the innervation of OANs/TANs in the 263 periphery of Drosophila. For this, we used the Tdc2-Gal4 line, allowing Gal4 expression 264 under the control of a regulatory sequence of the tyrosine decarboxylase enzyme ⁴⁰. As 265 this enzyme is essential for the synthesis of TA from tyrosine, the Tdc2-Gal4-line labels 266 both TANs and OANs. Within the Drosophila brain, Tdc2-Gal4 labels in total about 137 cells, while additional 39 cells are located in the VNS ^{38,39}. The small number of Tdc2Ns 267 268 lead to arborizations in large parts of the central brain, optic lobes and the thoracic and abdominal ganglion ^{37–40}. Based on the profound innervation of Tdc2Ns in the brain and 269 270 VNS, the variety of behaviors modulated by the OA/TA system including learning and 271 memory, feeding, vision, and sleep, are not surprising. Beyond the brain and VNS, 272 OANs and TANs massively innervate regions within the periphery of the fly. Here, we 273 described arborizations on most skeletal muscles, the antenna, wings, halteres and 274 reproductive system and parts of the circulatory system and stomodaeal ganglion (Fig. 275 6; Table 1).

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innervated by Tdc2Ns: proboscis/head muscles (rose); CNS and VNS (grey); peripheral

281 nerves targeting the antenna, ocelli, wings or halters (dark grey); neck, direct and

282 indirect flight muscles (blue); corpora allata (orange); leg muscles (green); heart and

alary muscles (red); abdominal skeletal muscles (ocher); muscle of Lawrence (yellow);

- reproductive organs (brown). Sensory neurons labeled by the Tdc2-Gal4 line are shown as purple dots.
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287 Our findings are in line with previous reports focusing on the expression of different OA 288 and TA receptors in the fly ^{50,51}. Accordingly, the OA receptor OAMB is expressed in 289 reproductive organs (in both male and female flies) and muscles, which are directly 290 innervated by Tdc2Ns. Additionally, the midgut and trachea contain OA and TA receptors ^{50,51}, but do not seem to be innervated by Tdc2Ns, even though axons run in 291 292 close vicinity to these organs. Likewise, the OA receptor Octß2R is expressed in the fat body, salivary glands and Malpighian tubules, tissues that seem not to be innervated by 293 294 Tdc2Ns, while the expression of Octß1R and Octß3R is more specific ^{50,51}. The three 295 tyramine receptors TyrR, TyrRII and TyrRIII show a broad expression in the periphery. 296 Interestingly, TyrR seems to be the only receptor expressed in the heart, suggesting that 297 only TA modulates heart function ⁵⁰. Contrary, OA has a modulatory effect on the heart 298 of other insect species including honeybees, olive fruit flies and cockroaches ⁵². This is 299 also in line with a previous report providing evidence that OA modulates the heart rate of the Drosophila fly and pupa, but not the larva ⁵³. 300

301 OA-dependent modulation of organs and tissues is mainly elicited through muscle 302 action, especially in terms of its impact on the "fight or flight" response. In line with this, 303 we observed *Tdc2-Gal4*-positive arborizations on nearly all skeletal muscles and many 304 visceral muscles (Table 1). In both Drosophila and desert locusts OA and TA is 305 expressed in type II terminals of skeletal muscles ⁴⁵. OA has an excitatory effect on 306 Drosophila flight muscles, while TA was shown to inhibit excitatory junction potentials, and thereby reduce muscle contractions and locomotion at least in the larva 6,8,54,55. In 307 308 addition, flies lacking OA show severe deficits in flight initiation and maintenance ^{10,45}. 309 Interestingly, in an antagonistic effect to serotonin, OA reduces crop muscle activity 310 presumably via Octß1R, suggesting that OA has different effects on muscle activity 311 dependent on the type of muscle ^{50,56}. However, our data do not provide any evidence of 312 a direct innervation of Tdc2Ns of the crop, even though many fibers run in close vicinity, 313 suggesting that OA might target the crop by volume transmission.

Furthermore, OA modulates ovulation and fertilization in insects ^{57–61}. Flies lacking OA
 display a severe egg-laying phenotype. Remarkably, within the female reproductive

316 organ two different OA receptors, OAMB and Octß2R, are necessary. Again, OA has a

317 strong impact on muscle activity within the reproductive system. Octß2R is expressed in

318 the visceral oviduct muscle and elicits muscle relaxation through an increase of 319 intracellular cAMP levels ⁵⁷. Such an OA-dependent modulation appears to be 320 conserved as OA is found in dorsal unpaired median neurons of locusts innervating oviduct muscles through the oviducal nerve ⁶². However, our data suggest that OA-321 322 positive fibers not only innervate oviduct muscles, but also enter the organs themselves. 323 The OAMB receptor is expressed in ephitelial cells inducing fluid secretion through 324 increasing intracellular Ca²⁺ levels ⁵⁷. Thus, OA affects different processes within the 325 female reproductive organ due to the expression of different receptors and their coupled 326 signaling pathways, which may be a general mechanism of the OA/TA system to fulfill 327 an extensive modulatory function ⁶³.

328 OA does not exclusively modulate muscle activity, but also sensory neurons of external 329 tissues like the antenna, halteres and wings. OA has also been shown to increase the 330 spontaneous activity of olfactory receptor neurons (ORN) ^{64,65}. The modulation of ORNs 331 allows OA to modulate the innate response to attractive stimuli like fruit odors or 332 pheromones ^{66,67}. Further, this modulation helps nestmate recognition in ants ⁶⁸. In 333 addition to Tdc2-Gal4-positive arborizations in the funiculus, we found Tdc2-Gal4-334 positive sensory neurons in the Johnston's organ, a chordotonal organ sensitive to 335 mechanosensory stimuli and thus important for hearing in insects. In mosquitos, OA 336 modulates auditory frequency tuning and thereby affects mating behavior ⁶⁹. In locusts, 337 OA similarly modulates the response of chordotonal neurons in the legs to encode proprioceptive information ⁷⁰. Our data suggest that chordotonal neurons in the leg, 338 339 wings, halteres and thorax are included in the Tdc2-Gal4 line suggesting a conserved 340 modulatory role of OA/TA for insect proprioception.

341 Taken together, our study suggest that the OA/TA system massively modulates various 342 organs and tissues in the periphery of *Drosophila*. Through distinct receptors and 343 coupled signaling pathways OANs/TANs mainly induce "fight or flight" responses by 344 modulating muscle activity, proprioception, and heart rate. As a result, the innervation 345 pattern in the periphery supports the idea that the OA/TA system is crucial for insects to 346 switch from a dormant to an excited state, by a positive modulation of muscle activity, 347 heart rate and energy supply, and a simultaneous negative modulation of physiological 348 processes like e.g. sleep.

349 Table 1: Organs, tissues, visceral and skeletal muscles innervated by Tdc2-Gal4-

350 positive neurons (Tdc2Ns).

Organs, tissues, visceral muscles	Tdc2N staining
Digestive system	
Esophagus (ES)	0
Crop	?
Hindgut	?
Circulatory system	
Corpora allata (CA)	0
Ventral longitudinal muscles of heart	x
Alary muscles of heart	х
Nervous system	
Brain	х
Ventral nervous system (VNS)	х
Stomodaeal ganglion	х
Corpora cardiaca (CC)	
Ocellar nerve (OCN)	х
Sensory organs	
Antenna	
Pedicle (a2)	х
Funiculus (a3)	х
Arista (a4,a5)	х
Wings	х
Halteres	х
Male reproductive system	
Muscles of Ejaculatory bulb (EB)	х
Anterior ejaculatory duct (ED')	х
Accessory gland (AG)	?
Vas deferens (VD)	х
Seminal vesicle (SV)	х
Female reproductive system	
Ovary (OV)	х
Lateral oviduct (LO)	х
Common oviduct (CO)	х
Spermathecal duct (SPd)	х
Spermatheca (SP)	

Skeletal muscles (after ⁴⁸)	Tdc2N staining
Head	
1	х
2	х
7	(x)
8	(x)
10	х
11	x
12	х
16	х
Thorax	
Leg muscles (after ⁷¹)	
Coxa trdm, trlm, trrm	x,x,x
Trochanter fedm, ferm	x,x
Femur Itm2, tilm, tidm, tirm	?,x,x,x
Tibia ltm1, talm, tadm, tarm	x,x,x,x
Flight muscles	
indirect	
45,46,47,48	x,x,x,x
direct	
49,50,51,52,54, 55,56,57,58	x,x,x,x,x, (x),?,x,?
Cervical muscles D 20,21,22,23, L 24; V 25,26,27	x,?,x,x, x; x,x,x
Mesothorax muscles 59,60,61,62	x,?,?,x
Metathorax muscles 77,78,79	?
Abdomen Segment 1	

Uterus (UT)	х	D 98,99,100,101,102
Jterus muscles	х	L 103; V 80,81,104
Sensory organ (SO)	Tdc2N cell body	Segment 2-6
Chordotonal organ (CO) antenna	x	D 109,115,121,127,133 L 110,116,122,128,134
CO thorax	х	V 111,117,123,129,135
CO leg	х	
SO leg	х	
SO wing	x	

х

x, staining; o, encircled by staining; (x), not enough samples; ?, ambiguous results or not investigated; D,
 dorsal; L, lateral; V, ventral

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- 354

355 Methods

SO haltere

356 Fly strains and fly rearing

357 All flies were cultured according to standard methods. In short, vials were kept under

358 constant conditions with 25°C and 60% humidity in a 12:12 light:dark cycle. Flies

359 carrying the Tdc2-Gal4 (⁴⁰, Bloomington Stock Center) and 10xUAS-IVS-myrGFP (⁷²,

360 Bloomington Stock Center) constructs were used for immunohistochemistry. To control

361 for an unspecific expression of the UAS construct, we stained 10xUAS-IVS-myrGFP

362 alone. No GFP staining was detected.

363

364 Immunocytochemistry

To visualize the arborisations of *Tdc2-Gal4*-positive neurons in the periphery whole body sections, as well as sections of the head, thorax and abdomen, were performed, respectively (see ⁷³). In short, the cuticle of 4 to 7 days old flies was opened in phosphate buffered saline (PBS, 0.1M) to ensure that the fixative is able to penetrate into the tissue. Whole flies were fixated with 4% paraformaldehyde in PBS for two hours

at room temperature and afterwards washed three times with PBS. Subsequently flies

were embedded in hot 7% Agarose low EEO (A2114; AppliChem). After hardening, the

372 flies were cut with a vibratome (Leica VT1000S) into 80-100 µm sections. Staining of the

373 sections was continued after washing in PBS containing 0.3% Triton-X100 (PBT) and 374 blocking in 5% normal goat serum in PBT. Rabbit anti-GFP (A6455, Molecular Probes) 375 in combination with mouse anti-Synapsin (3C11; ⁷⁴; 1:50) or mouse anti-Bruchpilot 376 (nc82; ⁷⁵) and mouse anti-Fasciclin 2 (1D4; DSHB; 1:100) were used as primary 377 antibodies. After one night at 4°C the specimens were washed six times in PBT and 378 incubated in secondary antibody solution for a subsequent night at 4°C. As secondary 379 antibodies goat anti-rabbit Alexa488 (Molecular Probes; 1:200) and goat anti-mouse 380 DyLight649 (Jackson ImmunoResearch; 1:200) were used. 4',6-Diamidino-2-phenylindol 381 Dihydrochlorid (DAPI: Sigma-Aldrich: 1:1000) and Alexa Fluor 633 Phalloidin (Molecular 382 Probes; 1:400) were used to visualize DNA and actin, respectively.

383

384 Confocal microscopy and data processing

Confocal images were taken with a Leica TCS SP8 microscope (Leica Microsystems,
Germany) with a 20x high aperture objective. Labelled specimens were scanned with a
step size of 1.0 µm to 1.5 µm. Image processing and alignment was performed using Fiji
(⁷⁶), Amira 5.3 (Visage Imaging, Berlin, Germany and Adobe Photoshop CS6 (Adobe
Systems, USA).

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- and M.S. prepared the figures. All authors reviewed the manuscript.