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Title: Preproglucagon (PPG) neurons innervate neurochemically identified autonomic neurons in the mouse brainstem

Article Type: Research Paper

Section/Category: Cellular and Molecular Neuroscience

Keywords: glucagon-like peptide 1;
nucleus of the solitary tract;
tyrosine hydroxylase;
choline acetyltransferase;
serotonin;
green fluorescent protein

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Order of Authors: Ida J Llewellyn-Smith, PhD; Greta J Gnanamanickam; Frank Reimann; Fiona M Gribble; Stefan Trapp, PhD

Abstract: Preproglucagon (PPG) neurons produce glucagon-like peptide-1 (GLP-1) and occur primarily in the nucleus tractus solitarius (NTS). GLP-1 affects a variety of central autonomic circuits, including those controlling the cardiovascular system, thermogenesis, and most notably energy balance. Our immunohistochemical studies in transgenic mice expressing YFP under the control of the PPG promoter showed that PPG neurons project widely to central autonomic regions, including brainstem nuclei. Functional studies have highlighted the importance of hindbrain receptors for the anorexic effects of GLP-1.

In this study, we assessed YFP innervation of neurochemically-identified brainstem neurons in transgenic YFP-PPG mice. Immunoreactivity for YFP plus choline acetyltransferase (ChAT), tyrosine hydroxylase (TH) and/or serotonin (5-HT) were visualised with two- or three-colour immunoperoxidase labelling using black (YFP), brown and blue-grey reaction products. In the dorsal motor nucleus of the vagus (DMV), terminals from fine YFP-immunoreactive axons closely apposed a small proportion of ChAT-positive and rare TH-positive/ChAT-positive motor neurons, mostly ventral to AP. YFP-immunoreactive innervation was virtually absent from the compact and loose formations of the nucleus ambiguus. In the NTS, some TH-immunoreactive neurons were closely apposed by YFP-containing axons. In the A1/C1 column in the ventrolateral medulla, close appositions on TH-positive neurons were more common, particularly in the caudal portion of the column. A single YFP-immunoreactive axon usually provided 1-3 close appositions on individual ChAT- or TH-positive neurons. Serotonin-immunoreactive neurons were most heavily innervated, with the majority of raphé pallidus, raphé obscurus and parapyramidal neurons receiving several close appositions from large varicosities of YFP-immunoreactive axons.

These results indicate that GLP-1 neurons innervate various populations of brainstem autonomic neurons. These include vagal efferent neurons and catecholamine neurons in areas linked with cardiovascular control. Our data also indicate a synaptic connection between GLP-1 neurons and 5-HT neurons, some of which might contribute to the regulation of appetite.

Response to Reviewers: We thank the Reviewers for their thorough evaluation of our manuscript and their helpful suggestions, which have improved the manuscript significantly. We now provide a revised manuscript that incorporates the suggestions of the Reviewers and hope that it is now acceptable for publication in Neuroscience.

Below are our detailed responses to each point raised by the Reviewers:

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1) "The presentation of the data is inadequate. The data are important and the authors stress several times that the powerful immunohistochemistry of YFP has enabled them to view the complete brainstem PPG innervation. If this is indeed the case - and I must admit that the staining shown in Figs 1-3 is really clear - they should help the reader in understanding the full extent of this innervation. This could be done in a series of line-drawings through the caudal brainstem (e.g. using images from mouse brain atlases) illustrating the location of 5HT, ChAT and TH cells (could even be illustrated on the same images) and the degree of innervation by PPG/YFP fibers. It is very difficult from the rather selected illustrations to get a full picture of the brainstem innervation and the extent to which the different phenotypically labeled neurons are contacted by the YFP fibers."

- We have added three new figures to the revised manuscript (Figures 2, 4 and 6). The new figures show maps of the distributions of TH-immunoreactive, ChAT-immunoreactive and 5-HT-immunoreactive neurons in sections through various regions of the medulla. The maps indicate which neurons receive close appositions from YFP-immunoreactive varicosities and which do not and include counts of the neurons with and without appositions. The inclusion of three new figures in the revised manuscript has required us to add a section to Experimental Procedures describing how the maps were constructed and neurons were counted (section 2.2.1, "Mapping neurochemically-identified brainstem neurons that received or lacked YFP-immunoreactive innervation" on pages 10 and 11).

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"The first statement is not supported by the data - and it certainly does not explain the cardiovascular effects of GLP-1 - well it could potentially be a route by which GLP-1 influences the cardiovascular system, but the authors do not know if this is an important pathway."

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We have altered the Discussion accordingly (section 4.4, pages 19 and 20).

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1) “On Page 10 the authors give a long description of how they tested for the specificity of the 5-HT antibody: “We tested the specificity of the anti-5-HT antiserum by adsorbing it with a glutaraldehyde conjugate of 5-HT and keyhole limpet haemocyanin (KLH). To prepare the 5-HT-glutaraldehyde-KLH conjugate, 50 mg of KLH (Catalogue #H-2133, Lot #43H9524;”

“This entire section can be removed. 5HT staining is widely used and the use of a commercially antibody giving the correct pattern of staining is in my view sufficient documentation for specificity.”

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- We have changed the abstract accordingly.

Dear Prof Lisberger,

RE: NSC 12-587

We submit for your consideration for publication in *Neuroscience* a revised version of our article entitled '**Preproglucagon (PPG) neurons innervate neurochemically identified autonomic neurons in the mouse brainstem**' by *Llewellyn-Smith, Gnanamanickam, Reimann, Gribble & Trapp*.

The manuscript has been modified on the basis of the suggestions of the Reviewers. In particular, we have included 3 new figures in addition to the three figures in the original version of the manuscript. The new figures document the distributions in the medulla of cholinergic, catecholaminergic and serotonergic neurons that receive or lack close appositions from YFP-PPG axons.

We have uploaded colour versions of all 6 Figures. If the revised manuscript is accepted for publication in *Neuroscience*, we would like all of the Figures to be published in colour online. However, in print, Figures 2-5 should appear in black and white as colour reproduction is only essential for Figure 1. In the revised manuscript, we have included two versions of the Figure Legends. The Legends that should appear in the online version of the article ("Figure Legends (ONLINE VERSION)", pages 21-24) describe the six figures in colour. The Legends that should appear in the print version of the article ("Figure Legends (PRINT VERSION)", pages 24-26) describe the colour version of Figure 1 and the black and white versions of Figures 2-5.

Thank you very much for your consideration. We look forward to hearing from you about the manuscript in due course.

Yours sincerely,

Dr Stefan Trapp
Department of Surgery and Cancer
Imperial College London
On behalf of the authors

Manuscript NSC-12-587 - Response to Reviewers

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HIGHLIGHTS

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- GLP-1 may influence satiety via heavy GLP-1 innervation of brainstem 5-HT neurons.
- Innervation of dorsal vagal neurons underpins GLP-1's role in control of digestion.
- Central GLP-1 may affect cardiovascular control via GLP-1 inputs to C1 neurons.

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11 **PREPROGLUCAGON (PPG) NEURONS INNERVATE NEUROCHEMICALLY**
12 **IDENTIFIED AUTONOMIC NEURONS IN THE MOUSE BRAINSTEM**
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14 Ida J Llewellyn-Smith¹, Greta J. E. Gnanamanickam¹, Frank Reimann², Fiona M Gribble² &
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30 **Abbreviated title:** GLP-1 innervation of brainstem neurons
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ABSTRACT

Preproglucagon (PPG) neurons produce glucagon-like peptide-1 (GLP-1) and occur primarily in the nucleus tractus solitarius (NTS). GLP-1 affects a variety of central autonomic circuits, including those controlling the cardiovascular system, thermogenesis, and most notably energy balance. Our immunohistochemical studies in transgenic mice expressing YFP under the control of the PPG promoter showed that PPG neurons project widely to central autonomic regions, including brainstem nuclei. Functional studies have highlighted the importance of hindbrain receptors for the anorexic effects of GLP-1.

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KEY WORDS

glucagon-like peptide 1, nucleus of the solitary tract, tyrosine hydroxylase, choline acetyltransferase, serotonin, green fluorescent protein

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LIST OF ABBREVIATIONS

| | | |
|----|-------|---|
| 1 | | |
| 2 | | |
| 3 | | |
| 4 | 5-HT | Serotonin, 5-hydroxytryptamine |
| 5 | | |
| 6 | AP | Area postrema |
| 7 | | |
| 8 | ChAT | Choline acetyltransferase |
| 9 | | |
| 10 | DAB | Diaminobenzidine |
| 11 | | |
| 12 | DMH | Dorsomedial nucleus of the hypothalamus |
| 13 | | |
| 14 | DMV | Dorsal motor nucleus of the vagus |
| 15 | | |
| 16 | GFP | Green fluorescent protein |
| 17 | GLP-1 | Glucagon-like peptide 1 |
| 18 | | |
| 19 | Ig | Immunoglobulin |
| 20 | | |
| 21 | IRT | Intermediate reticular nucleus |
| 22 | | |
| 23 | KLH | Keyhole limpet haemocyanin |
| 24 | | |
| 25 | LPS | Lipopolysaccharide |
| 26 | | |
| 27 | NA | Nucleus ambiguus |
| 28 | | |
| 29 | NHS | Normal horse serum |
| 30 | | |
| 31 | NTS | Nucleus of the solitary tract |
| 32 | | |
| 33 | PPG | Preproglucagon |
| 34 | PPY | Parapyramidal region |
| 35 | | |
| 36 | PVN | Paraventricular nucleus of the hypothalamus |
| 37 | | |
| 38 | RVLM | Rostral ventrolateral medulla |
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| 40 | TH | Tyrosine hydroxylase |
| 41 | YFP | Yellow fluorescent protein |
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INTRODUCTION

Glucagon-like peptide 1 (GLP-1) is an incretin that is released from enteroendocrine cells and facilitates absorption of nutrients (Holst, 2007). Like other gut peptides, such as cholecystokinin, GLP-1 is also synthesized by neurons within the central nervous system. GLP-1 is produced by post-translational processing of proglucagon (PPG) and immunoreactivity for the products of PPG processing is found in many brain regions, with highest levels occurring in the dorsomedial (DMH) and paraventricular nucleus (PVN) of the hypothalamus and lowest levels in the cortex and hindbrain (Jin et al., 1988, Vrang et al., 2007, Tauchi et al., 2008). Somata capable of synthesizing GLP-1, however, are restricted to the lower brainstem. The largest population of GLP-1-containing somata occurs in the caudal nucleus of the solitary tract (NTS) and there are also some cell bodies in the dorsomedial part of the medullary reticular nucleus (Jin et al., 1988, Larsen et al., 1997). Similarly, *in situ* hybridisation has revealed PPG mRNA only in the caudal NTS, the intermediate reticular nucleus (IRT) and the olfactory bulb (Merchenthaler et al., 1999). Retrograde tracing has confirmed that the hypothalamic axons containing GLP-1 arise from the cell bodies in the NTS and IRT (Larsen et al., 1997, Vrang et al., 2007).

Microinjection of GLP-1 or GLP-1 agonists into the brain has a multitude of effects, including suppression of food intake, control of blood glucose levels, nausea, changes in blood pressure and heart rate, as well as neuroprotection and effects on learning and memory (Tang-Christensen et al., 1996, Turton et al., 1996, Van Dijk et al., 1996, Thiele et al., 1998, Kinzig et al., 2002, Yamamoto et al., 2002, Daring et al., 2003, Cabou et al., 2008, Sandoval et al., 2008). Because of the functional importance of GLP-1 in the brain, we recently reinvestigated the distribution of central neurons capable of synthesizing GLP-1 using a transgenic mouse in which GLP-1 neurons express yellow fluorescent protein (YFP) throughout their cytoplasm under the control of the PPG promoter (Reimann et al., 2008). This approach allowed us to visualise PPG neurons and their processes with an unprecedented level of detail. Our study revealed the full distribution of PPG cell bodies and dendrites within the medulla and demonstrated that axons were widespread throughout the brain with the notable exception of cerebellum, hippocampus and cerebral cortex (Llewellyn-Smith et al., 2011). These observations matched well with the distribution of GLP-1 receptors within the brain (Shughrue et al., 1996, Merchenthaler et al., 1999). Notably, we found varicose axons in many central sites that are involved in regulating autonomic functions.

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GLP-1 released in these areas could contribute to the global and coordinated control of food intake, energy balance and maintenance of cardiovascular homeostasis.

Recent functional studies have highlighted the importance of brainstem circuitry and brainstem GLP-1 receptors for physiological function (Yamamoto et al., 2003, Wan et al., 2007a, Wan et al., 2007b, Hayes et al., 2008, Hayes et al., 2009, Holmes et al., 2009, Williams et al., 2009, Barrera et al., 2011) see also (Trapp and Hisadome, 2011) for review). These findings, together with the fact that our immunohistochemical study revealed many more varicose immunoreactive axons in the brainstem than previously reported, prompted us to define the innervation targets of PPG neurons within the brainstem in more detail. Catecholamine neurons in the area postrema (AP) have been shown to express GLP-1 receptors (Yamamoto et al., 2003) and a subpopulation of cholinergic dorsal vagal motor neurons responds electrically to GLP-1 (Wan et al., 2007a, Wan et al., 2007b, Holmes et al., 2009), consistent with these cell types receiving inputs from GLP-1 neurons.

In this study, we used transgenic YFP-PPG mice (Reimann et al., 2008) in order to take advantage of the strong YFP expression that occurs throughout the cytoplasm of PPG neurons, including their terminals (Hisadome et al., 2010, Llewellyn-Smith et al., 2011). To reveal GLP-1 innervation of cholinergic, catecholamine and serotonin neurons in the brainstem, we detected YFP-immunoreactivity in combination with immunoreactivity for choline acetyltransferase (ChAT), tyrosine hydroxylase (TH) or 5-hydroxytryptamine (5-HT), using two-colour or three-colour immunoperoxidase staining.

2. EXPERIMENTAL PROCEDURES

These studies were performed on 11 adult male and 11 adult female mGLU-124 Venus YFP mice (Reimann et al., 2008), which will be referred to here as YFP-PPG mice. Mice were perfused at 12-16 weeks of age and weighed between 25 and 35g with males consistently being heavier than females of the same age. Mice were bred at Imperial College, kept on a 12 hour light:dark cycle and had unlimited access to food and water. All experiments were carried out in accordance with the UK Animals (Scientific Procedures) Act, 1986, with appropriate ethical approval.

Mice under halothane anesthesia were given heparin (500 IU/l), flushed with phosphate-buffered saline to remove blood and perfused transcardially with 60 ml of phosphate-buffered 4% formaldehyde, pH 7.4. Brains were post-fixed intact for 3 days at

1 room temperature on a shaker in the same fixative and then shipped to Flinders for sectioning
2 and immunohistochemical staining.

3
4 Blocks containing the brainstem were trimmed from the post-fixed brains and
5 infiltrated with sucrose. Three series of transverse 30µm cryostat sections were cut from
6 each block. Because we did not use a matrix to block the brains, the dorsoventral tilt of the
7 sections varied among mice. Consequently, Bregma values could not be reliably assigned to
8 the brainstem sections studied here.
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12 13 14 **2.1 IMMUNOHISTOCHEMISTRY**

15
16 Cryostat sections were first washed 3 x 10 min in 10 mM Tris, 0.9% NaCl, 0.05%
17 thimerosal in 10 mM phosphate buffer, pH 7.4, (TPBS) containing 0.3% Triton X-100 and
18 then exposed to TPBS-Triton containing 10% normal horse serum (NHS) for at least 30 min.
19 TPBS-Triton containing 10% NHS was used to dilute primary antibodies; TPBS-Triton
20 containing 1% NHS, to dilute secondary antibodies; and TPBS-Triton, to dilute the avidin-
21 horseradish peroxidase complex. All steps in each protocol occurred on a shaker at room
22 temperature and 3 x 10 min washes in TPBS were done after each incubation in an
23 immunoreagent.
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31 Titration was used to determine the working concentrations of primary antibodies.
32 Optimal dilutions produced the maximum number of immunoreactive structures with
33 minimal non-specific background staining. In the case of the anti-green fluorescent protein
34 (GFP) antibody used to detect the YFP-expressing neurons, more dilute antibody was used to
35 optimally visualize axons than for optimally visualizing cell bodies (see Llewellyn-Smith et
36 al., 2011). We have previously shown that mouse tissue lacking GFP-expressing neurons
37 shows no staining after immunoperoxidase processing to reveal GFP (Llewellyn-Smith et al.,
38 2011).
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46 After completion of the double or triple-labelling immunoperoxidase protocol
47 detailed below, sections were mounted in serial order onto chrome alum-gelatine coated
48 slides, dried and dehydrated. Coverslips were applied with Permaslip mounting medium
49 (Alban Scientific, St Louis MO, USA).
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54 **2.1.1. Two-colour Immunoperoxidase Labelling**

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56 Double immunoperoxidase labelling was used to localise YFP-immunoreactivity
57 with a black reaction product and either ChAT-, TH- or 5-HT-immunoreactivity with a brown
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1 reaction product. After treatment with TBS-Triton and 10% NHS-TBS-Triton, the sections
2 were transferred into 1:50,000 or 1:100,000 chicken anti-GFP (Catalogue #ab13970, Lot
3 #623923; Abcam, Cambridge, UK) for 3-5 days. After washing, the sections were exposed
4 overnight to 1:500 biotinylated donkey anti-chicken immunoglobulin (Ig) Y (Catalogue
5 #703-065-155, ; Jackson ImmunoResearch, West Grove PA) and then to 1:1,500 ExtrAvidin-
6 peroxidase (Sigma-Aldrich, St Louis MO, USA) for 4-6 hours. Structures containing YFP-
7 immunoreactivity were stained black with a nickel-intensified diaminobenzidine (DAB)
8 reaction in which peroxide was generated using glucose oxidase (Llewellyn-Smith et al.,
9 2005). After a second blocking step in 10% NHS-TBS-Triton, sections were exposed to
10 either 1:5,000 goat anti-ChAT (Catalogue # AB144P, Lot # 18110644 or JC1669317;
11 Chemicon, Temecula, CA, USA), 1:5,000 rabbit anti-TH (Catalogue # AB152, Lot #
12 LV1375881 or LV1532665; Chemicon) or 1:5,000 or 1:10,000 rabbit anti-5-HT (Catalogue
13 #8250-0009, Lot #23073052; Biogenesis, Poole UK) for 2-3 days. Next, there was an
14 overnight incubation in 1:500 biotinylated donkey secondary antibody directed against the
15 immunoglobulin of the species in which the primary antibody was raised, i.e., donkey anti-
16 goat IgG (Catalogue #711-705-065-147; Jackson ImmunoResearch Laboratories Inc., West
17 Grove, PA, USA) or donkey anti-rabbit IgG (Catalogue #711-065-152; Jackson
18 ImmunoResearch). After a final 4 hour incubation in ExtrAvidin-peroxidase, immunoreactive
19 neurons were stained brown with an imidazole-intensified DAB reaction (Llewellyn-Smith et
20 al., 2005).

2.1.2 *Three-colour Immunoperoxidase Labelling*

21 Triple immunoperoxidase labelling was used to localise YFP-, TH- and ChAT-
22 immunoreactivity in the same sections. After the two colour protocol was completed using a
23 1:400,000 dilution of anti-GFP and a 1:5,000 dilution of anti-TH, an additional block with
24 10% NHS-TBS-Triton was done and the sections were incubated first in 1:5,000 goat anti-
25 ChAT for 2-3 days, then in biotinylated anti-goat IgG overnight and finally in ExtrAvidin-
26 horseradish peroxidase for 4-6 hours. ChAT-immunoreactivity was localised with a Vector
27 SG kit (Vector Laboratories, Burlingame, CA, USA) with reagents added to generate
28 peroxide with glucose oxidase reaction (Llewellyn-Smith et al., 1999).

2.1.3 *Antibody Characterization*

29 In our previous study on YFP-PPG mice (Llewellyn-Smith et al., 2011), we
30 demonstrated the specificity of the chicken anti-GFP antiserum used here. We showed that
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1 no staining occurred in sections from animals that lacked GFP-expressing neurons and that
2 the anti-GFP antibody stained all of the neurons that fluoresced due to the presence of YFP.

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4 Sections of rat spinal cord stained with the goat anti-ChAT antiserum show the
5 expected distribution of cholinergic neurons in the spinal cord (Fenwick et al., 2006). When
6 we have used recombinant rat ChAT (Chemicon Catalogue #Ag220) to adsorb this antiserum,
7 we found no staining in tissue fixed and processed as here (Llewellyn-Smith et al., 2005).
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10 The supplier of the rabbit anti-TH antibody has used western blotting to check its
11 specificity and obtained selective labelling of a single band at approximately 62kDa
12 (reduced), which corresponds to TH. In sections from rats perfused and
13 immunohistochemically processed as here using this anti-TH antibody, we have found that
14 the medulla contains immunoreactive neurons with the expected morphologies and
15 distributions and that the spinal cord contains axons with the expected distribution. This
16 antibody has been used extensively to study catecholaminergic innervation in the brain,
17 spinal cord and periphery in a variety of species (e.g. (Cano et al., 2008, Nangle et al., 2009,
18 Gnanamanickam and Llewellyn-Smith, 2011)).
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27 We tested the specificity of the anti-5-HT antiserum by adsorbing it with a
28 glutaraldehyde conjugate of 5-HT and keyhole limpet haemocyanin, which we prepared.
29 Adsorbed and unadsorbed antibody solutions were incubated overnight at 4°C and used to
30 immunoperoxidase stain 30 µm cryostat sections of formaldehyde-fixed rat and mouse
31 medulla. Overnight adsorption removed the vast majority of staining but did not completely
32 abolish it. Medulla sections stained with the adsorbed antiserum contained rare, faintly-
33 immunoreactive cell bodies in raphé pallidus and a few immunoreactive axons in the dorsal
34 vagal complex compared to the usual distribution of immunoreactivity obtained with the
35 unadsorbed antiserum.
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45 **2.2. DATA COLLECTION AND ANALYSIS**

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47 Immunoperoxidase-stained sections from the spinomedullary junction to the caudal
48 end of the aqueduct were examined with an Olympus BH-2 brightfield microscope. Cell
49 bodies and proximal dendrites containing immunoreactivity for TH, ChAT or serotonin were
50 examined for the presence of close appositions from YFP-positive axon terminals. An x100
51 oil immersion lens was used to determine whether or not YFP-containing terminals were
52 closely apposed to neurochemically-identified cell bodies and dendrites. YFP terminals lying
53 side-by-side with an immunoreactive neuron were classed as closely apposed when there was no
54 space visible between the terminal and the neuron. In the case of terminals overlying immunoreactive
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neurons, a close apposition was considered to be present when the terminal and the filled neuron were in focus in the same plane.

A SPOT RT colour camera and SPOT RT software version 4.6 (Diagnostic Instruments Inc., Sterling Heights, MI, USA) were used to collect digital images from the sections as TIFF files. The sharpness, brightness, and contrast of the TIFFs were adjusted in Adobe PhotoShop, which was also used to colour match digital images. Montages of micrographs and plates containing micrographs were prepared using PhotoShop.

2.2.1 Mapping neurochemically-identified brainstem neurons that received or lacked YFP-immunoreactive innervation

To construct line drawings, a SPOT RT colour camera and an x10 objective were used to capture overlapping digital images through the dorsal vagal complex or ventral medulla on one side of the brainstem from each mouse. The micrographs from each section were saved as TIFF files, imported into Adobe Photoshop and montaged to create a single image of the region to be mapped. A layer was added to the montage in PhotoShop and the surface of the medulla was traced with the pen tool. Landmarks were traced with the pen tool on another layer. Low magnification line drawings of each section were created from digital images taken with an x2 objective using a similar strategy.

For recording the distribution of the each different neurochemical class of neuron that received or lacked appositions from YFP-immunoreactive varicosities on the line drawings, a separate layer was added to the TIFF file containing the x10 montage of each section. The position of each neuron was marked with a dot using the pencil tool and a different colour to represent each different type of neuron. To count neurons with and without appositions, the layers with coloured dots were saved as individual TIFF files. Each TIFF file was opened in ImageJ (National Institutes of Health) and the image was converted to black and white. Dots were counted using the Analyze Particle function.

Maps of DMV, AP and A2 neurons: In the dorsal vagal complex from one female mouse, TH-immunoreactive neurons with and without YFP appositions and ChAT-immunoreactive neurons with and without YFP appositions were mapped from every third section triple-stained for YFP, TH and ChAT (60 μ m separating mapped sections). A total of 6 sections were mapped. The mapped region included most of the dorsal vagal complex ventral to AP. However, YFP innervation was not assessed in the most caudal portion of the dorsal vagal complex because the dense accumulation of YFP-immunoreactive dendrites

made it impossible to determine whether close appositions were present on TH- or ChAT-immunoreactive neurons.

Maps of A1/C1 neurons: TH-immunoreactive neurons with and without YFP appositions that lay in the A1/C1 column in the ventral medulla from one female mouse were mapped in every sixth section triple-stained for YFP, TH and ChAT (150 μ m separating mapped sections). A total of 13 sections were mapped. The mapped region extended from the pyramidal decussation through the caudal portion of the facial nucleus.

Maps of serotonin neurons: Ventral medullary 5-HT neurons with and without appositions from YFP-immunoreactive varicosities were mapped in every sixth section double-stained for YFP and 5-HT from one male mouse (150 μ m separating mapped sections). A total of 8 sections were mapped. The mapped region extended from approximately the middle of AP to approximately the middle of the facial nucleus.

Plates containing the maps were constructed in Adobe Illustrator. The layer containing the montage was deleted from the TIFF file for each section. The resulting image was imported into Illustrator and the positions of all the neurons were remapped using larger symbols of different colours.

3. RESULTS

YFP-immunoreactive innervation of catecholamine neurons (Figures 1-4), cholinergic neurons (Figures 1 and 2), and serotonin neurons (Figures 5 and 6) in the brainstem was assessed in 11 adult male and 11 adult female YFP-PPG mice. After immunoperoxidase staining, YFP-PPG neurons in the NTS and IRT showed intense YFP-immunoreactivity throughout their cell bodies and dendrites (Figures 1A-C, I and J). In the brainstem, varicose axons arising from the YFP-PPG neurons were also very intensely stained for YFP. The YFP-containing axons had clearly delineated varicosities and fine intervaricose segments (Figures 1 and 3). Some of the axons had large varicosities; others had small fine varicosities and on some axons both large and small varicosities occurred (e.g., Figures 1G, 1J, 3B). All three types of axons made close appositions on neurochemically-identified brainstem neurons.

The distribution of YFP-expressing neurons in the medulla of YFP-PPG mice was the same as reported previously (Llewellyn-Smith et al., 2011). The largest group of YFP-immunoreactive cell bodies occurred in the caudal lateral NTS and extended caudally from approximately mid-AP level past the caudal tip of AP (Figures 1A, F and D). The second

1 major group of YFP-immunoreactive somata was located in the IRT, extending from about
2 mid-AP to roughly where the NTS moved away from the fourth ventricle. Triple labelling
3 for GFP, TH and ChAT revealed that the YFP-PPG neurons in IRT lay dorsomedial to both
4 the cholinergic neurons of the nucleus ambiguus (NA) and the noradrenergic neurons of the
5 A1 cell group (Figure 1B).
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10 **3.1. INNERVATION OF CATECHOLAMINE NEURONS BY YFP-IMMUNOREACTIVE** 11 **PREPROGLUCAGON NEURONS** 12 13

14 Catecholamine neurons from the spinomedullary junction to the caudal end of the
15 aqueduct were revealed by immunoreactivity for TH in transverse sections through the brains
16 of 10 male and 7 female YFP-PPG mice (Figures 1 and 3). In the medulla, TH-
17 immunoreactive neurons occurred in the ventrolateral medulla (A1/C1 cell group), the NTS
18 (A2 cell group), the AP and around the midline in the dorsal medulla (C2 and C3 cell
19 groups). In the pons, TH-immunoreactive cell bodies were located around the exit of the
20 facial nerve (A5 cell group) and in the locus coeruleus (A6 cell group).
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27 In the dorsal medulla, some of the TH-immunoreactive neurons of the A2 cell group
28 in the NTS received close appositions from YFP-immunoreactive axons (Figures 1C-E and
29 2). These appositions were more common in the region where YFP-immunoreactive cell
30 bodies and dendrites formed a dense network throughout the NTS (Figure 2). In AP, a small
31 number of TH-immunoreactive cells were closely apposed by YFP-immunoreactive axons
32 (Figures 1F-H and 2). There were fewer TH-positive neurons with appositions in caudal AP
33 than in rostral AP (Figure 2). Throughout AP, the TH-containing neurons that received close
34 appositions usually occurred near the lateral edge of AP or within the clusters of TH-positive
35 neurons that lay on the dorsal surface of the most rostral portion of AP (Figure 2). TH-
36 positive C2 and C3 neurons near the dorsal medullary midline received little YFP-positive
37 innervation (not shown).
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47 As we have described previously, there was a moderate YFP-immunoreactive
48 innervation of the rostral ventrolateral medulla (RVLM). Boutons of YFP-positive axons in
49 this region closely apposed the somata or proximal dendrites of some TH-immunoreactive C1
50 neurons (Figures 3A-C and 4). Close appositions from YFP-immunoreactive axons also
51 occurred on TH-positive neurons of the A1 cell group in the caudal ventrolateral medulla
52 (Figure 4). Although rare on the most caudal A1 neurons near the spinomedullary junction
53 (not shown), appositions from YFP-positive axons were common on TH-immunoreactive A1
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1 neurons in the intermediate reticular region ventral to AP (Figure 4). Near the caudal tip of
2 the 4th ventricle, TH-positive neurons in the rostroventral respiratory group also received
3 close appositions from YFP-immunoreactive axons.
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5 In the ventral pons, YFP-immunoreactive varicosities closely apposed the somata or
6 proximal dendrites of occasional TH-immunoreactive neurons of the A5 cell group. In the
7 dorsal pons, TH-immunoreactive A6 neurons of the locus coeruleus formed a dense band of
8 cell bodies (Figure 3D-F). Few YFP-immunoreactive axons penetrated the locus coeruleus
9 region and the cell bodies of TH-positive A6 neurons only rarely received close appositions
10 from YFP-positive boutons (Figure 3E, F). Many more varicose YFP-positive axons occurred
11 within the neighbouring and more medially located Barrington's nucleus (Figure 3E). The
12 TH-positive dendrites of A6 neurons penetrated the more heavily innervated area of
13 Barrington's nucleus but close appositions by YFP-immunoreactive varicosities were still
14 rare on the dendrites that originated from the A6 neurons in locus coeruleus.
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23 In all of the TH-immunoreactive cell groups examined, innervated neurons generally
24 received close appositions from 1-3 varicosities on a single YFP-immunoreactive axon.
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28 **3.2. PREPROGLUCAGON INNERVATION OF CHOLINERGIC NEURONS**

29 Cholinergic neurons were revealed by the presence of ChAT-immunoreactivity in
30 their cytoplasm. Transverse sections between the spinomedullary junction and the caudal end
31 of the aqueduct were analysed from 4 male and 9 female YFP-PPG mice. These sections
32 contained major cholinergic cell populations in the hypoglossal nucleus, the dorsal motor
33 nucleus of the vagus (DMV), NA, the facial nucleus and the abducens, pedunculopontine and
34 tegmental nuclei.
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42 A relatively small proportion of the ChAT-immunoreactive DMV neurons were
43 directly innervated by axons containing immunoreactivity for YFP (Figures 1I and J, 2).
44 Near the spinomedullary junction, the DMV contained a moderate supply of varicose YFP-
45 immunoreactive axons. Nevertheless, at this most caudal level of the DMV, only occasional
46 ChAT-positive dorsal vagal motor neurons received close appositions. PPG cell bodies
47 marked by YFP-immunoreactivity were located more rostrally in the NTS beneath AP (see
48 above). At this level, a dense network of YFP-positive dendrites was present along the
49 ventral aspect of the DMV. More DMV neurons in this region than rostrally or caudally
50 appeared to receive close appositions from YFP-positive terminals but the network of YFP-
51 PPG dendrites may also have obscured some appositions. Mapping ChAT-positive DMV
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1 neurons with and without appositions from YFP-immunoreactive varicosities ventral to AP
2 (Figure 2) showed that motor neurons receiving GLP-1 innervation were distributed
3 throughout the subpostremal DMV. Nevertheless, innervated neurons were most often
4 encountered medially and ventrally in this portion of the nucleus. Occasional DMV neurons
5 rostral to AP were also closely apposed by YFP-containing terminals. In sections triple-
6 stained for YFP, TH and ChAT, YFP-immunoreactive appositions were present on a very
7 small number of DMV neurons containing immunoreactivity for both ChAT and TH (not
8 shown). When DMV neurons were innervated, it was usually by a single YFP-
9 immunoreactive axon that had 1-3 fine, small varicosities in close apposition to the ChAT-
10 positive neuron (Figure 1J).
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18 In contrast to the DMV, very few YFP-immunoreactive axons travelled through the
19 hypoglossal nucleus and none closely apposed ChAT-positive hypoglossal motor neurons. A
20 small population of neurons with faint ChAT-immunoreactivity was located ventral to the
21 rostral NA close to the surface of the medulla. YFP-immunoreactive axons with fine
22 varicosities made rare close appositions on these ventral cholinergic neurons. Although YFP-
23 containing axons ran amongst the cholinergic neurons of the loose portion of NA, no close
24 appositions were found. The compact formation of NA was also virtually devoid of YFP-
25 immunoreactive axons.
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32 Few YFP-immunoreactive axons occurred in the facial nucleus. Close appositions
33 were almost never encountered on ChAT-positive neurons in this region. The rare neurons
34 that did receive YFP-immunoreactive input were mostly located in the most medial part of
35 the facial nucleus. Similarly, the nucleus abducens was virtually devoid of YFP-
36 immunoreactive axons and close appositions were never found. Further rostral, cholinergic
37 neurons in the trigeminal motor nucleus did not receive a YFP-positive innervation although
38 YFP-containing axons traversed the lateral edge of the nucleus. However, rare ChAT-positive
39 neurons of the laterodorsal tegmental nucleus received close appositions from YFP-
40 immunoreactive axons. Single close appositions from YFP-positive varicosities also occurred
41 on the somata or proximal dendrites of occasional pedunclopontine tegmental neurons.
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52 **3.3. *PREPROGLUCAGON INNERVATION OF SEROTONIN NEURONS***

53 5-HT-immunoreactive neurons from the spinomedullary junction to the decussation
54 of the superior cerebellar peduncle were revealed in transverse sections from 5 male and 5
55 female YFP-PPG mice. The most caudal 5-HT-immunoreactive cell bodies were located in
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1 raphé pallidus, raphé obscurus, the parapyramidal region (PPY) and along the ventral surface
2 of the medulla (Figures 5 and 6). The 5-HT-immunoreactive neurons on the ventral surface
3 first appeared in the region of AP (beneath caudal AP in some mice and beneath rostral AP in
4 others) and disappeared at about the level of the rostral edge of NA. The 5-HT-positive PPY
5 neurons were located just lateral to the pyramids and were roughly co-extensive
6 rostrocaudally with the motor neurons of NA. The long axes of many of the serotonergic PPY
7 cell bodies were oriented dorsoventrally (Figure 5B). At the level of the caudal facial
8 nucleus, the dorsoventrally-oriented cell bodies characteristic of PPY serotonin neurons
9 transitioned into cell bodies with a predominantly mediolateral orientation, which were the
10 serotonin neurons of raphé magnus. The dorsal raphé and pontine raphé nuclei contained the
11 most rostral 5-HT-immunoreactive neurons examined.
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20 The serotonin-containing neurons in PPY and on the ventral medullary surface were
21 the most densely innervated of any of the neurochemically-identified neurons in this study
22 (Figures 5B, D, E and 6). The cell bodies and proximal dendrites of almost all of these
23 neurons received several close appositions from YFP-positive terminals. In general, close
24 appositions from YFP-immunoreactive axons were somewhat less common on neurons in
25 raphé pallidus than on PPY neurons. However, for both raphé pallidus neurons and PPY
26 neurons, the density of appositions was highest at the caudal end of their distributions and
27 diminished rostrally (Figure 6).
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34 YFP-immunoreactive axons also travelled through raphé magnus and the dorsal and
35 pontine raphé nuclei. In all of these locations, occasional 5-HT-immunoreactive neurons
36 received close apposition from YFP-labelled varicosities but the majority of serotonin
37 neurons in these locations did not receive YFP innervation.
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42 Overall, more 5-HT-immunoreactive neurons in the medulla were innervated than
43 were cholinergic or catecholamine neurons and the majority of caudal medullary 5-HT-
44 positive neurons received close appositions from YFP-immunoreactive varicosities. The YFP
45 innervation of individual serotonin neurons was also denser than the innervation of individual
46 cholinergic or catecholamine neurons. Each innervated ChAT-positive or TH-positive neuron
47 received between one and three close appositions from YFP-containing varicosities. In
48 contrast, 5-HT-immunoreactive somata in the caudal medulla often received at least twice
49 this number of appositions.
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4. DISCUSSION

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3 In this study, we used two- and three-colour immunoperoxidase labelling to assess
4 the GLP-1 innervation of neurochemically-identified brainstem neurons in YFP-PPG mice
5 (Reimann et al., 2008, Llewellyn-Smith et al., 2011). YFP expression in these mice is
6 restricted to cells in which the PPG promoter is active (Reimann et al., 2008, Hisadome et al.,
7 2010). Consequently, in these mice, YFP-immunoreactivity occurs exclusively in cells that
8 synthesize PPG, the precursor for glucagon, as well as GLP-1, GLP-2 and oxyntomodulin
9 (Holst, 2007). As a result, YFP labels only pancreatic α -cells that produce glucagon,
10 enteroendocrine L-cells and populations of central neurons that produce GLP-1 and GLP-2 in
11 equimolar amounts (Reimann et al., 2008). Single-cell RT-PCR has demonstrated that the
12 YFP-expressing cells in YFP-PPG mice still produce PPG mRNA and should therefore
13 synthesize GLP-1 in addition to YFP (Hisadome et al., 2010, 2011). Here, in addition to
14 revealing YFP, we visualized neurons releasing catecholamines or acetylcholine using the
15 presence of immunoreactivity for enzymes involved in the synthesis of these
16 neurotransmitters (TH and ChAT, respectively). Immunoreactivity for 5-HT was used to
17 identify serotonin neurons. We assumed that YFP-containing axon terminals found in close
18 apposition to TH-, ChAT- or 5-HT-immunoreactive cell bodies or dendrites signified
19 innervation. Whilst only electron microscopy can provide the ultimate proof for this
20 assumption, work on other types of central autonomic neurons has shown that at least half of
21 the terminals that form close appositions at the light microscope level make synaptic contacts
22 at the ultrastructural level (Pilowsky et al., 1992).

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25 Of brainstem neurons containing immunoreactivity for TH, 5-HT or ChAT,
26 serotonin neurons received the heaviest YFP innervation. A high proportion of the 5-HT-
27 positive neurons in the parapyramidal region and on the ventral surface of the medulla
28 received many close appositions from YFP-immunoreactive varicosities. Appositions were
29 also common on serotonin neurons in raphé pallidus and occurred on some neurons in caudal
30 raphé obscurus. In contrast, neurons in raphé magnus and the dorsal raphé nuclei were
31 moderately to sparsely innervated. Amongst brainstem catecholamine neurons, a significant
32 fraction of the TH-immunoreactive neurons in the A1/C1 column in the ventrolateral medulla
33 received close appositions, with 1-3 varicosities on the same axon providing innervation to a
34 soma or proximal dendrite. In contrast, only rare TH-immunoreactive neurons in the locus
35 coeruleus received appositions from YFP-positive varicosities. Of brainstem cholinergic
36 neurons, only a subset of ChAT-positive neurons in the DMV received appositions from
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1 YFP-containing axons. A small number of DMV cell bodies with appositions also contained
2 immunoreactivity for TH. Cholinergic DMV neurons with appositions most commonly
3 occurred at the same rostrocaudal level as the cell bodies and dendrites of YFP-PPG neurons.
4 The remaining groups of brainstem cholinergic neurons, including those in NA and the
5 hypoglossal nucleus, essentially lacked YFP innervation. All of the neuronal populations
6 mentioned above as receiving close appositions from YFP-immunoreactive varicosities have
7 been shown to express mRNA for the GLP-1 receptor (Merchenthaler et al., 1999).
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13 **4.1. *INNERVATION OF AREA POSTREMA***

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16 Activation of GLP-1 receptors within the brainstem is associated with a number of
17 physiological and pathophysiological effects. Pharmacological activation of GLP-1 receptors
18 in the brainstem reduces gut motility (Holmes et al., 2009), mediates lipopolysaccharide-
19 induced anorexia (Grill et al., 2004), influences satiety elicited by gastric distension (Hayes et
20 al., 2009) and causes hypothermia (Skibicka et al., 2009). In AP, GLP-1 receptors reside
21 primarily on catecholamine neurons (Goke et al., 1995, Merchenthaler et al., 1999), which
22 exhibit Fos-immunoreactivity after intravenous administration of the GLP-1 receptor agonist
23 exendin-4 (Yamamoto et al., 2002, Yamamoto et al., 2003). Because the AP has an
24 incomplete blood-brain-barrier, it has been suggested that these receptors are a ‘gateway’
25 through which GLP-1 from the circulation can influence central nervous function. However,
26 our previous study demonstrated that YFP-PPG neurons send axons into AP (Llewellyn-
27 Smith et al., 2011) and we suggested that GLP-1 released from these axons, rather than
28 circulating GLP-1, might activate at least some of the receptors in AP. Indeed, our present
29 results demonstrate that YFP-PPG neurons innervate catecholaminergic AP neurons although
30 this innervation is relatively sparse. It is therefore possible that both centrally-derived and
31 peripherally-produced GLP-1 might activate receptors within AP.
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47 **4.2. *GASTRIC AND PANCREATIC FUNCTION***

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50 Neurons in the DMV respond functionally to GLP-1 or its stable analogue exendin-
51 4. *In vivo*, activation of GLP-1 receptors in the DMV has been linked to a reduction in gut
52 motility (Holmes et al., 2009). *In vitro*, both gastric-projecting and pancreas-projecting
53 dorsal vagal motor neurons alter their electrical activity in response to GLP-1 application
54 (Wan et al., 2007a, Wan et al., 2007b, Holmes et al., 2009). In the presence of TTX, about
55 40% of pancreas-projecting DMV neurons maintained their GLP-1 induced depolarisation,
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1 and thus are expected to have GLP-1 receptors (Wan et al., 2007b). In agreement with these
2 pharmacological data, we found that a subset of ChAT-immunoreactive neurons within the
3 DMV had a sparse YFP-immunoreactive innervation. Interestingly, DMV neurons with
4 YFP-positive appositions often occurred near the medial portion of the nucleus, where we
5 have previously identified neurons that express Fos in response to glucoprivation induced by
6 2-deoxyglucose in rats (Llewellyn-Smith et al., 2012). Furthermore, in rats, DMV neurons
7 containing both TH- and ChAT-immunoreactivity project exclusively to the gastric corpus
8 (Guo et al., 2001). In this study, we found that a very small number of DMV neurons with
9 this neurochemistry received YFP-positive innervation. Despite these observations, tracing
10 experiments will be required to determine whether the DMV innervation reported here
11 specifically targets gastric- and pancreas-projecting neurons or neurons supplying other
12 regions of the gastrointestinal tract or other organs. At the level of the DMV, GLP-1 appears
13 to be excitatory (Wan et al., 2007b, Holmes et al., 2009). However, its effect on gastric
14 motility is inhibitory, suggesting that non-adrenergic, non-cholinergic vagal postganglionic
15 neurons are part of the circuitry that mediates the effects of GLP-1 at the level of the stomach
16 (Travagli et al., 2006, Holmes et al., 2009).
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30 **4.3. *CARDIOVASCULAR CONTROL***

31 In this study, we found that YFP-PPG neurons innervated ventral medullary
32 catecholamine cell groups, including A1, A5 and C1 neurons. We found substantial numbers
33 of YFP-positive axons throughout the rostrocaudal extent of the ventrolateral medulla and
34 catecholaminergic A1 and C1 neurons clearly received close appositions. C1 neurons of the
35 RVLM are important for regulation of blood pressure because they mediate sympathetic tone
36 (Abbott et al., 2009, Marina et al., 2011, Schreihofner and Sved, 2011). It seems feasible to
37 assume that the innervation of C1 neurons described here provides an anatomical substrate
38 for at least some of the demonstrated cardiovascular effects of central GLP-1 administration
39 (Yamamoto et al., 2002, Cabou et al., 2008, Hayes et al., 2008). Since catecholamine neurons
40 in the A1 region project to the hypothalamus (Sawchenko and Swanson, 1982, Rinaman,
41 2001), GLP-1 inputs to A1 neurons could influence cardiovascular homeostasis as well as
42 other autonomic functions through an A1-hypothalamus pathway. We also found that A5
43 catecholamine neurons received input from YFP-immunoreactive axons. GLP-1 input to A5
44 neurons may also influence sympathetic control of blood pressure (Guyenet, 2006). In
45 contrast to the C1, A1 and A5 neurons, catecholaminergic A6 neurons of the locus coeruleus
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in the dorsal pons were almost completely without innervation from YFP-immunoreactive axons. This observation suggests that any influence of central GLP-1 on wakefulness and arousal (Sara, 2009) is unlikely to be mediated by locus coeruleus neurons.

The innervation of Barrington's nucleus by YFP-immunoreactive axons could have significance for the control of autonomic function. This nucleus is important for the control of micturition (Sasaki, 2005) and viral tracing studies have shown that Barrington's nucleus is also part of the circuitry that controls sympathetically innervated targets such as the kidney, spleen and brown adipose tissue (Cano et al., 2000, Cano et al., 2003).

4.4. AUTONOMIC FUNCTIONS CONTROLLED BY SEROTONIN NEURONS

We found here that GLP-1 axons heavily innervated the majority of 5-HT neurons in ventral regions of the medulla and also provided input to other serotonergic nuclei in the brainstem. Serotonin (5-HT) is important for the control of a variety of homeostatic processes, including energy balance, respiration and body temperature (Breisch et al., 1976, Ray et al., 2011).

The strong innervation of 5-HT neurons in the ventral medulla might play an important role in the satiating effects of central GLP-1. GLP-1 and 5-HT have both been implicated in the regulation of food intake (Tang-Christensen et al., 1996, Turton et al., 1996, Halford et al., 2010, Lam et al., 2010). A role of 5-HT downstream of GLP-1 is suggested by the finding that 5-HT-receptor 2C knockout mice fail to show satiation after intraperitoneal administration of GLP-1 (Asarian, 2009). Anorexia caused by lipopolysaccharide (LPS) is also likely to involve both GLP-1 signalling and 5-HT receptors (Langhans, 2007). LPS induces Fos-immunoreactivity in both GLP-1 neurons in the NTS (Rinaman, 1999) and in serotonin neurons of raphé pallidus, raphé magnus and dorsal and median raphé nuclei (Kopf et al., 2010). Furthermore, both injection of the GLP-1 antagonist exendin-9 into the 4th ventricle (Grill et al., 2004) and intraperitoneal administration of the brain-penetrant serotonin 2C-receptor antagonist SB 242084 reduce anorexia evoked by LPS (Kopf et al., 2010). Taken together with our observation that 5-HT immunoreactive neurons in raphé nuclei receive close appositions from the axons of GLP-1 neurons, these data might indicate that 5-HT neurons are also a downstream target of GLP-1 neurons involved in LPS-induced anorexia.

GLP-1 might also affect thermoregulatory responses through an input to 5-HT neurons. Pharmacogenetic suppression of electrical activity in serotonin neurons *in vivo*

1 causes hypothermia and a reduction in respiratory chemosensitivity (Ray et al., 2011). The
2 effects of central GLP-1 on chemosensitivity have not been assessed, but activation of
3 brainstem GLP-1 receptors has been shown to cause hypothermia (Hayes et al., 2008).
4 Furthermore, blockade of hindbrain GLP-1 receptor with exendin-9 prevents CART-induced
5 hypothermia (Skibicka et al., 2009). Innervation of raphé pallidus neurons could provide a
6 link between central GLP-1 and thermogenesis, as 5-HT neurons in this region are known to
7 stimulate neurons in sympathetic nuclei of the spinal cord, which in turn promote brown
8 adipose tissue metabolism (Morrison et al., 2008). As the GLP-1 receptor is predominantly
9 coupled to G_s (Thorens and Widmann, 1996), it remains to be determined if raphé pallidus
10 neurons are inhibited by GLP-1 or if the hypothermic effects of central GLP-1 involve
11 alternative neuronal circuits.
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21 **4.5. CONCLUSIONS**

22 In summary, this study provides the first detailed description of the GLP-1
23 innervation of neurochemically-identified monoamine and serotonin neurons in the
24 brainstem. Our observations create an anatomical scaffold for understanding the diverse
25 array of autonomic functions affected by central GLP-1. We expect that the information
26 about the distribution of GLP-1 inputs to brainstem neurons presented here will provide a
27 valuable guide for functional studies probing the roles of central GLP-1 neurons.
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41 Grants 480414 and 1025031) to ILS, and Wellcome Trust Senior Research Fellowships
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43 technical assistance.
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FIGURE LEGENDS (ONLINE VERSION)

FIGURE 1. Three-colour immunoperoxidase labelling for YFP in YFP-PPG neurons (black), tyrosine hydroxylase (TH) in catecholamine neurons (brown) and choline acetyltransferase (ChAT) in cholinergic neurons (blue-grey) in transverse sections through the dorsal medulla of YFP-PPG mice. **A**, Low magnification micrograph showing the cell bodies of black, YFP-immunoreactive PPG neurons in the nucleus of the solitary tract (NTS). These neurons lie lateral to the dorsal motor nucleus of the vagus (DMV) and the majority occur at the level of the area postrema (AP). The dendrites of the YFP-PPG neurons in NTS extend dorsomedially and also run lateral to the hypoglossal nucleus (HGN) and along the border between the DMV and the HGN. cc, central canal. Bar, 100 μ m. **B**, In the intermediate reticular nucleus (IRT), the black, YFP-immunoreactive cell bodies of PPG neurons are located dorsomedial to both the brown, TH-immunoreactive neurons of the A1 cell group and the blue-grey, ChAT-immunoreactive neurons of the nucleus ambiguus (NA). Bar, 100 μ m. **C**, Low magnification micrograph showing the dorsal vagal complex at the level of the most rostral YFP-PPG neurons. A collection of YFP-immunoreactive dendrites travels between the DMV and HGN. Arrows, brown, TH-immunoreactive cell bodies that are shown at higher magnification in D and E. Bar, 250 μ m. **D and E**, Single varicosities (arrows) of black, YFP-immunoreactive axons closely appose brown, TH-immunoreactive cell bodies in the NTS. Bars, 20 μ m. **F**, Low magnification micrograph showing the AP near the level of the most rostral YFP-PPG neurons. Arrow, a brown, TH-immunoreactive cell body near the surface of AP that is shown at higher magnification in G. Bar, 100 μ m. **G**, A single varicosity (arrow) of a black, YFP-immunoreactive axon closely apposes the brown, TH-immunoreactive cell body in AP. Bar, 10 μ m. **H**, At the surface of rostral AP, black, varicose, YFP-immunoreactive axons and black, varicose, YFP-immunoreactive dendrites are intermixed with brown, TH-immunoreactive dendrites arising from AP neurons. Bar, 20 μ m. **I**, Low magnification micrograph showing the DMV near the level of the most rostral YFP-PPG neurons. Arrows J1 and J2, two blue-grey, ChAT-immunoreactive vagal motor neurons that are shown at higher magnification in J. Bar, 100 μ m. **J**, Fine varicosities (arrows) of black, YFP-immunoreactive axons closely appose the blue-grey, ChAT-immunoreactive cell bodies of DMV neurons J1 and J2. Bar, 20 μ m.

FIGURE 2. Maps of sections through the dorsal vagal complex of a YFP-PPG mouse showing the distribution of ChAT-immunoreactive and TH-immunoreactive neurons with and without close appositions from YFP-immunoreactive varicosities. Sections were triple-stained to reveal YFP, TH and ChAT. Every third section was mapped; maps are therefore separated by 60µm. The most caudal section is at the top left of the figure and the most rostral section is at the bottom right. TH-immunoreactive neurons in the nucleus tractus solitarius (NTS) and area postrema (AP) are represented by circles; and ChAT-immunoreactive neurons, predominantly in the dorsal motor nucleus of the vagus (DMV), by diamonds. TH-immunoreactive neurons that receive close appositions from YFP-immunoreactive varicosities are shown in red; those that do not receive appositions are shown in blue. ChAT-immunoreactive neurons that receive close appositions from YFP-immunoreactive varicosities are shown in gold; those that do not receive appositions are shown in green. The cell bodies of YFP-PPG neurons are represented by white stars. Numerical values indicate the proportion of neurons in each region that received close appositions from YFP-immunoreactive varicosities. The percentages of neurons that received YFP-immunoreactive appositions in each region are given in parentheses. The asterisk indicates an area of the DMV in which appositions could not be assessed because the dendrites of YFP-PPG neurons were too dense. cc, central canal; HGN, hypoglossal nucleus.

FIGURE 3. Two-colour immunoperoxidase labelling for YFP in YFP-PPG neurons (black) and tyrosine hydroxylase (TH) in catecholamine neurons (brown) in transverse sections through the rostral ventrolateral medulla (RVLM) and dorsal pons of YFP-PPG mice. **A**, Low magnification micrograph showing the RVLM. Arrows, brown, TH-immunoreactive cell bodies that are shown at higher magnification in D and E. py, pyramidal tract. Bar, 100 µm. **B and C**, Single varicosities (arrows) of black, YFP-immunoreactive axons closely appose brown, TH-immunoreactive cell bodies in the RVLM. Bar in B, 10 µm; bar in C, 20 µm. **D**, Low magnification micrograph showing the region of the locus coeruleus (LC) and Barrington's nucleus (Bar). Left is medial and the ventricle appears at top left. The region in Box E is shown at higher magnification in E. Arrow F, a brown, TH-immunoreactive neuron in the subcoeruleus region (SubC) that is shown at higher magnification in F. Bar, 250 µm. **E**, There are many black, varicose YFP-immunoreactive axons in Bar. However, only a few, black, varicose YFP-immunoreactive axons travel amongst the brown, TH-immunoreactive neurons of LC. Of all of the TH-immunoreactive cell bodies in this region of LC, only 6 received close appositions from YFP-immunoreactive

1 varicosities, one of which (lower right) is marked by an arrowhead. There were no close
 2 appositions on the TH-immunoreactive dendrites of LC neurons that penetrated Bar. Bar, 50
 3 μm . **F**, A single varicosity (arrow) of a black, YFP-immunoreactive axon closely apposes the
 4 brown, TH-immunoreactive cell body in SubC. Bar, 20 μm .
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9 **FIGURE 4.** Maps of sections through the ventral medulla of a YFP-PPG mouse showing
 10 the distribution of TH-immunoreactive neurons with and without close appositions from
 11 YFP-immunoreactive varicosities. Sections were triple-stained to reveal YFP, TH and
 12 ChAT. Every sixth section was mapped; maps are therefore separated by 150 μm . The most
 13 caudal section is at the top left of the figure and the most rostral section is at the bottom right.
 14 TH-immunoreactive neurons that receive close appositions from YFP-immunoreactive
 15 varicosities are represented by red circles; TH-immunoreactive neurons that do not receive
 16 appositions are represented by blue circles. The cell bodies of YFP-PPG neurons are
 17 represented by white stars. Numerical values indicate the proportion of neurons in each
 18 section that received close appositions from YFP-immunoreactive varicosities. The
 19 percentages of neurons that received YFP-immunoreactive appositions in each section are
 20 given in parentheses. The insets show the entire section from which each map was
 21 constructed, allowing the dorsoventral tilt of the section can be assessed. FN, facial nucleus;
 22 IO, inferior olive; NAc, compact formation of the nucleus ambiguus; py, pyramidal tract;
 23 pyx, pyramidal decussation.
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38 **FIGURE 5.** Two-colour immunoperoxidase labelling for YFP in YFP-PPG neurons
 39 (black) and 5-HT in serotonin neurons (brown) in transverse sections through the caudal
 40 ventral medulla of YFP-PPG mice. **A**, Low magnification micrograph showing the ventral
 41 medulla from the midline through the RVLM, Arrows B and C, brown, groups of 5-HT-
 42 immunoreactive cell bodies that are shown at higher magnification in B and C. py, pyramidal
 43 tract. Bar, 100 μm . **B**, In the parapyramidal region (PPY), a cluster of serotonin-
 44 immunoreactive cells bodies receives close appositions from many YFP-immunoreactive
 45 axons. Bar, 20 μm . **C**, The cell bodies of two brown, 5-HT immunoreactive neurons in raphé
 46 pallidus (RPa) receive several close apposition from varicosities (arrows) of black, YFP-
 47 immunoreactive axons. Bar, 10 μm . **D**, A black, YFP-immunoreactive varicosity (arrow)
 48 forms a close apposition on a brown, 5-HT immunoreactive cell body that lies near the
 49 ventral surface (VS) in PPY. Bar, 10 μm . **E**, Two black varicosities on the same YFP-
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immunoreactive axon (arrows) closely appose a brown, 5-HT-immunoreactive cell body in RPa. Bar, 10 μ m.

FIGURE 6. Maps of sections through the ventral medulla of a YFP-PPG mouse showing the distribution of 5-HT-immunoreactive neurons with and without close appositions from YFP-immunoreactive varicosities. Sections were double-stained to reveal YFP and 5-HT. Every sixth section was mapped; maps are therefore separated by 150 μ m. The most caudal section is at the top left of the figure and the most rostral section is at the bottom right. 5-HT-immunoreactive neurons that receive close appositions from YFP-immunoreactive varicosities are represented by red circles; 5-HT-immunoreactive neurons that do not receive appositions are represented by blue circles. The cell bodies of YFP-PPG neurons are represented by white stars. Numerical values indicate the proportion of 5-HT-immunoreactive neurons in different region or the entire mapped population that received close appositions from YFP-immunoreactive varicosities. The percentages of neurons that received YFP-immunoreactive appositions are given in parentheses. The insets show the entire section from which each map was constructed, allowing the dorsoventral tilt of the section to be assessed. FN, facial nucleus; IO, inferior olive; NAc, compact formation of the nucleus ambiguus; py, pyramidal tract; RPa, raphé pallidus; VS, ventral surface..

FIGURE LEGENDS (PRINT VERSION)

FIGURE 1. *Three-colour immunoperoxidase labelling for YFP in YFP-PPG neurons (black), tyrosine hydroxylase (TH) in catecholamine neurons (brown) and choline acetyltransferase (ChAT) in cholinergic neurons (blue-grey) in transverse sections through the dorsal medulla of YFP-PPG mice. A, Low magnification micrograph showing the cell bodies of black, YFP-immunoreactive PPG neurons in the nucleus of the solitary tract (NTS). These neurons lie lateral to the dorsal motor nucleus of the vagus (DMV) and the majority occur at the level of the area postrema (AP). The dendrites of the YFP-PPG neurons in NTS extend dorsomedially and also run lateral to the hypoglossal nucleus (HGN) and along the border between the DMV and the HGN. cc, central canal. Bar, 100 μ m. B, In the intermediate reticular nucleus (IRT), the black, YFP-immunoreactive cell bodies of PPG neurons are located dorsomedial to both the brown, TH-immunoreactive neurons of the AI*

1 cell group and the blue-grey, ChAT-immunoreactive neurons of the nucleus ambiguus (NA).
 2 Bar, 100 μm . **C**, Low magnification micrograph showing the dorsal vagal complex at the
 3 level of the most rostral YFP-PPG neurons. A collection of YFP-immunoreactive dendrites
 4 travels between the DMV and HGN. Arrows, brown, TH-immunoreactive cell bodies that are
 5 shown at higher magnification in **D** and **E**. Bar, 250 μm . **D and E**, Single varicosities
 6 (arrows) of black, YFP-immunoreactive axons closely appose brown, TH-immunoreactive
 7 cell bodies in the NTS. Bars, 20 μm . **F**, Low magnification micrograph showing the AP near
 8 the level of the most rostral YFP-PPG neurons. Arrow, a brown, TH-immunoreactive cell
 9 body near the surface of AP that is shown at higher magnification in **G**. Bar, 100 μm . **G**, A
 10 single varicosity (arrow) of a black, YFP-immunoreactive axon closely apposes the brown,
 11 TH-immunoreactive cell body in AP. Bar, 10 μm . **H**, At the surface of rostral AP, black,
 12 varicose, YFP-immunoreactive axons and black, varicose, YFP-immunoreactive dendrites
 13 are intermixed with brown, TH-immunoreactive dendrites arising from AP neurons. Bar, 20
 14 μm . **I**, Low magnification micrograph showing the DMV near the level of the most rostral
 15 YFP-PPG neurons. Arrows J1 and J2, two blue-grey, ChAT-immunoreactive vagal motor
 16 neurons that are shown at higher magnification in **J**. Bar, 100 μm . **J**, Fine varicosities
 17 (arrows) of black, YFP-immunoreactive axons closely appose the blue-grey, ChAT-
 18 immunoreactive cell bodies of DMV neurons J1 and J2. Bar, 20 μm .

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 35 **FIGURE 2.** Maps of sections through the dorsal vagal complex of a female YFP-PPG
 36 mouse showing the distribution of ChAT-immunoreactive and TH-immunoreactive neurons
 37 with and without close appositions from YFP-immunoreactive varicosities. Sections were
 38 triple-stained to reveal YFP, TH and ChAT. Every third section was mapped; maps are
 39 therefore separated by 60 μm . The most caudal section is at the top left of the figure and the
 40 most rostral section is at the bottom right. TH-immunoreactive neurons in the nucleus tractus
 41 solitarius (NTS) and area postrema (AP) that receive close appositions from YFP-
 42 immunoreactive varicosities are represented by black circles; TH-immunoreactive neurons
 43 that do not receive appositions are represented by grey circles. ChAT-immunoreactive
 44 neurons, predominantly in the dorsal motor nucleus of the vagus (DMV), that receive close
 45 appositions from YFP-immunoreactive varicosities are represented by black diamonds with
 46 white centres; ChAT-immunoreactive neurons that do not receive appositions are represented
 47 by pale grey diamonds. The cell bodies of YFP-PPG neurons are represented by white stars.
 48 Numerical values indicate the proportion of neurons in each region that received close
 49 appositions from YFP-immunoreactive varicosities. The percentages of neurons that
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1 received YFP- immunoreactive appositions in each region are given in parentheses. The
 2 asterisk indicates an area of the DMV in which appositions could not be assessed because the
 3 dendrites of YFP-PPG neurons were too dense. cc, central canal; HGN, hypoglossal
 4 nucleus.
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9 **FIGURE 3.** Two-colour immunoperoxidase labelling for YFP in YFP-PPG neurons and
 10 tyrosine hydroxylase (TH) in catecholamine neurons in transverse sections through the
 11 rostral ventrolateral medulla (RVLM) and dorsal pons of YFP-PPG mice. **A**, Low
 12 magnification micrograph showing the RVLM. Arrows, TH-immunoreactive cell bodies that
 13 are shown at higher magnification in **D** and **E**. py, pyramidal tract. Bar, 100 μ m. **B and C**,
 14 Single varicosities (arrows) of black, YFP-immunoreactive axons closely appose TH-
 15 immunoreactive cell bodies in the RVLM. Bar in **B**, 10 μ m; bar in **C**, 20 μ m. **D**, Low
 16 magnification micrograph showing the region of the locus coeruleus (LC) and Barrington's
 17 nucleus (Bar). Left is medial and the ventricle appears at top left. The region in Box **E** is
 18 shown at higher magnification in **E**. Arrow **F**, a TH-immunoreactive neuron in the
 19 subcoeruleus region (SubC) that is shown at higher magnification in **F**. Bar, 250 μ m. **E**,
 20 There are many black, varicose YFP-immunoreactive axons in Bar. However, only a few,
 21 black, varicose YFP-immunoreactive axons travel amongst the TH-immunoreactive neurons
 22 of LC. Of all of the TH-immunoreactive cell bodies in this regions of LC, only 6 received
 23 close appositions from YFP-immunoreactive varicosities, one of which (lower right) is
 24 marked by an arrowhead. There were no close appositions on the TH-immunoreactive
 25 dendrites of LC neurons that penetrated Bar. Bar, 50 μ m. **F**, A single varicosity (arrow) of a
 26 black, YFP-immunoreactive axon closely apposes a TH-immunoreactive cell body in SubC.
 27 Bar, 20 μ m.
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45 **FIGURE 4.** Maps of sections through the ventral medulla of a female YFP-PPG mouse
 46 showing the distribution of TH-immunoreactive neurons with and without close appositions
 47 from YFP-immunoreactive varicosities. Sections were triple-stained to reveal YFP, TH and
 48 ChAT. Every sixth section was mapped; maps are therefore separated by 150 μ m. The most
 49 caudal section is at the top left of the figure and the most rostral section is at the bottom
 50 right. TH-immunoreactive neurons that receive close appositions from YFP-immunoreactive
 51 varicosities are represented by black circles; TH-immunoreactive neurons that do not receive
 52 appositions are represented by grey circles. The cell bodies of YFP-PPG neurons are
 53 represented by white stars. Numerical values indicate the proportion of neurons in each
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1 section that received close appositions from YFP-immunoreactive varicosities. The
 2 percentages of neurons that received YFP- immunoreactive appositions in each section are
 3 given in parentheses. The insets show the entire section from which each map was
 4 constructed so that the dorsoventral tilt of the section can be assessed. FN, facial nucleus;
 5 IO, inferior olive; NAc, compact formation of the nucleus ambiguus; py, pyramidal tract;
 6 pyx, pyramidal decussation.
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 12 **FIGURE 5.** Two-colour immunoperoxidase labelling for YFP in YFP-PPG neurons and 5-
 13 HT in serotonin neurons in transverse sections through the caudal ventral medulla of YFP-
 14 PPG mice. **A**, Low magnification micrograph showing the ventral medulla from the midline
 15 through the RVLM, Arrows B and C, groups of 5-HT-immunoreactive cell bodies that are
 16 shown at higher magnification in B and C. py, pyramidal tract. Bar, 100 μ m. **B**, In the
 17 parapyramidal region (PPY), a cluster of 5-HT-immunoreactive cells bodies receives close
 18 appositions from many YFP-immunoreactive axons. Bar, 20 μ m. **C**, The cell bodies of two 5-
 19 HT immunoreactive neurons in raphé pallidus (RPa) receive several close apposition from
 20 varicosities (arrows) of black, YFP-immunoreactive axons. Bar, 10 μ m. **D**, A black, YFP-
 21 immunoreactive varicosity (arrow) forms a close apposition on a 5-HT immunoreactive cell
 22 body that lies near the ventral surface (VS) in PPY. Bar, 10 μ m. **E**, Two black varicosities
 23 on the same YFP-immunoreactive axon (arrows) closely appose a 5-HT-immunoreactive cell
 24 body in RPa. Bar, 10 μ m.
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38 **FIGURE 6.** Maps of sections through the ventral medulla of a male YFP-PPG mouse
 39 showing the distribution of 5-HT-immunoreactive neurons with and without close appositions
 40 from YFP-immunoreactive varicosities. Sections were double-stained to reveal YFP and 5-
 41 HT. Every sixth section was mapped; maps are therefore separated by 150 μ m. The most
 42 caudal section is at the top left of the figure and the most rostral section is at the bottom
 43 right. 5-HT-immunoreactive neurons that receive close appositions from YFP-
 44 immunoreactive varicosities are represented by black circles; 5-HT-immunoreactive neurons
 45 that do not receive appositions are represented by grey circles. The cell bodies of YFP-PPG
 46 neurons are represented by white stars. Numerical values indicate the proportion of 5-HT-
 47 immunoreactive neurons in different region or the entire mapped population that received
 48 close appositions from YFP-immunoreactive varicosities. The percentages of neurons that
 49 received YFP-immunoreactive appositions are given in parentheses. The insets show the
 50 entire section from which each map was constructed so that the dorsoventral tilt of the
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section can be assessed. *FN*, facial nucleus; *IO*, inferior olive; *NAc*, compact formation of the nucleus ambiguus; *py*, pyramidal tract; *RPa*, raphé pallidus; *VS*, ventral surface.

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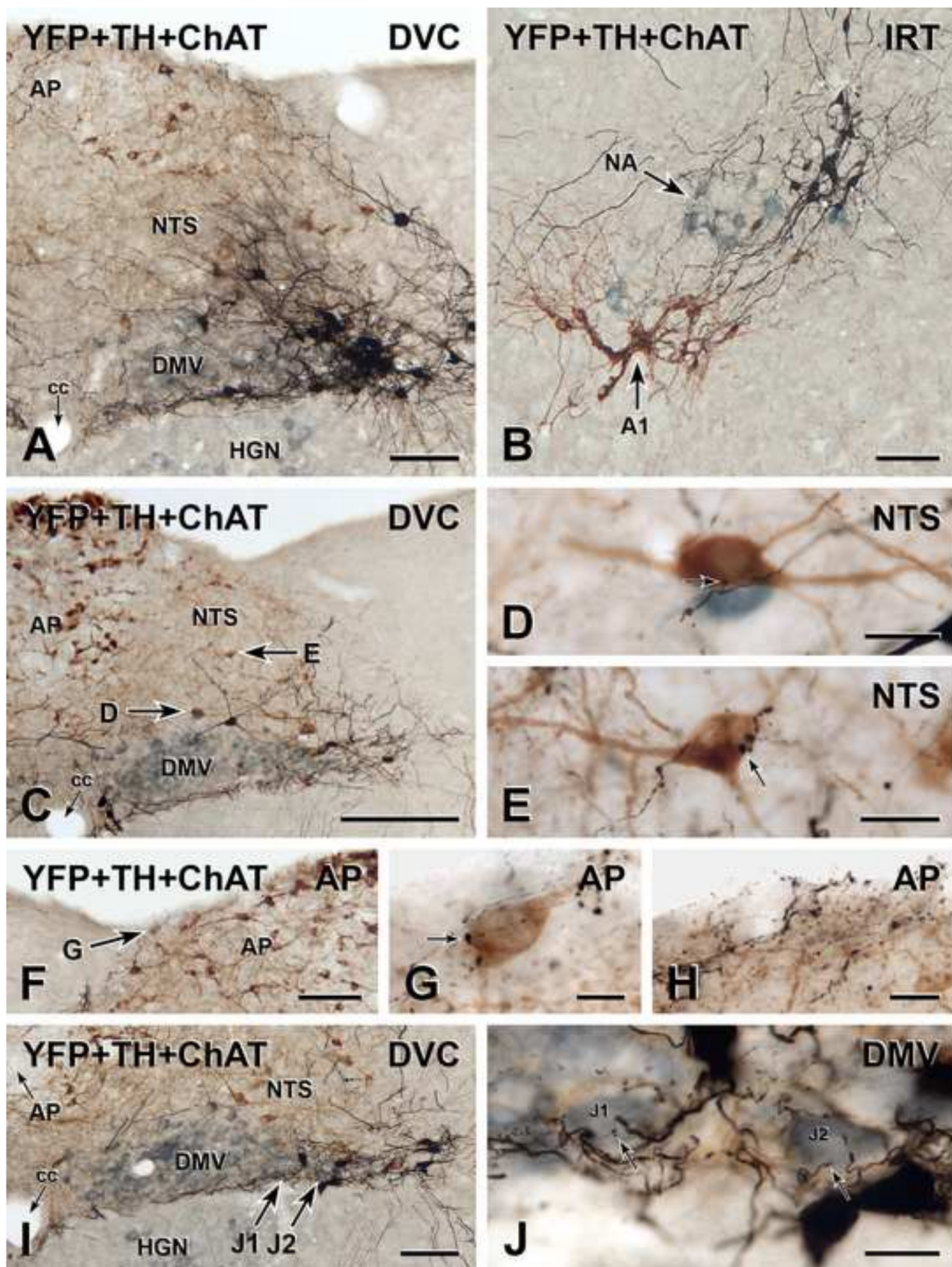
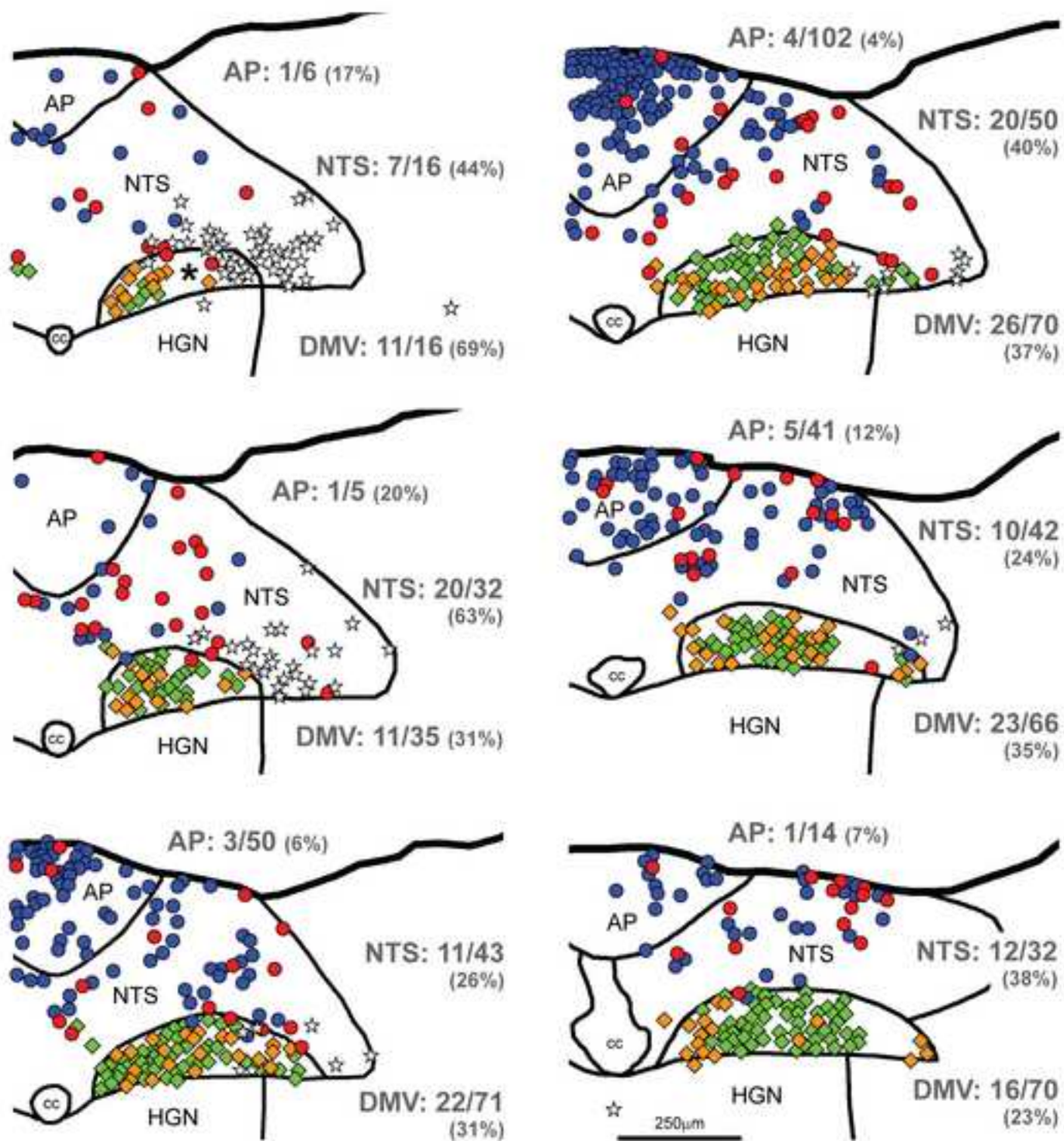


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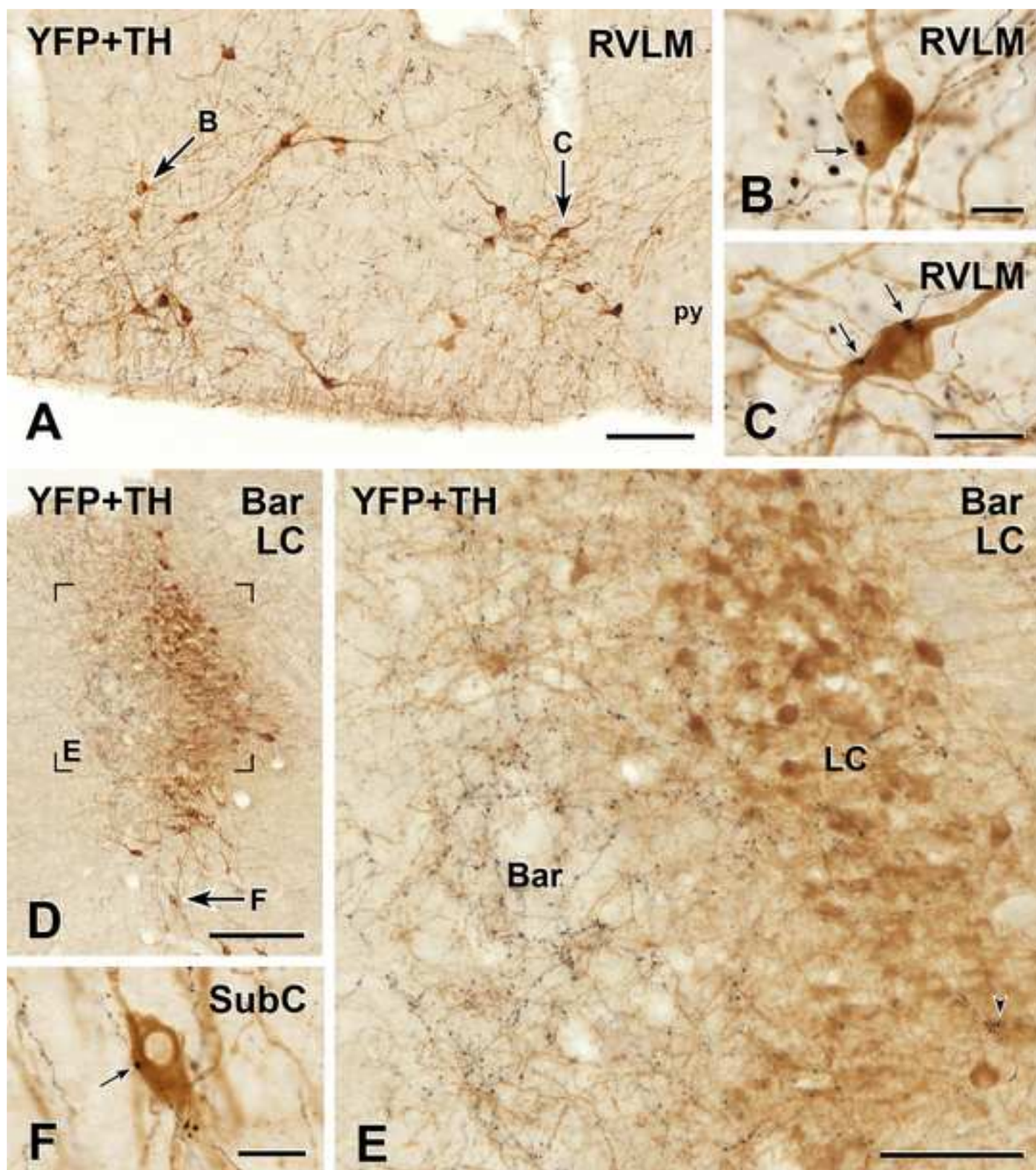


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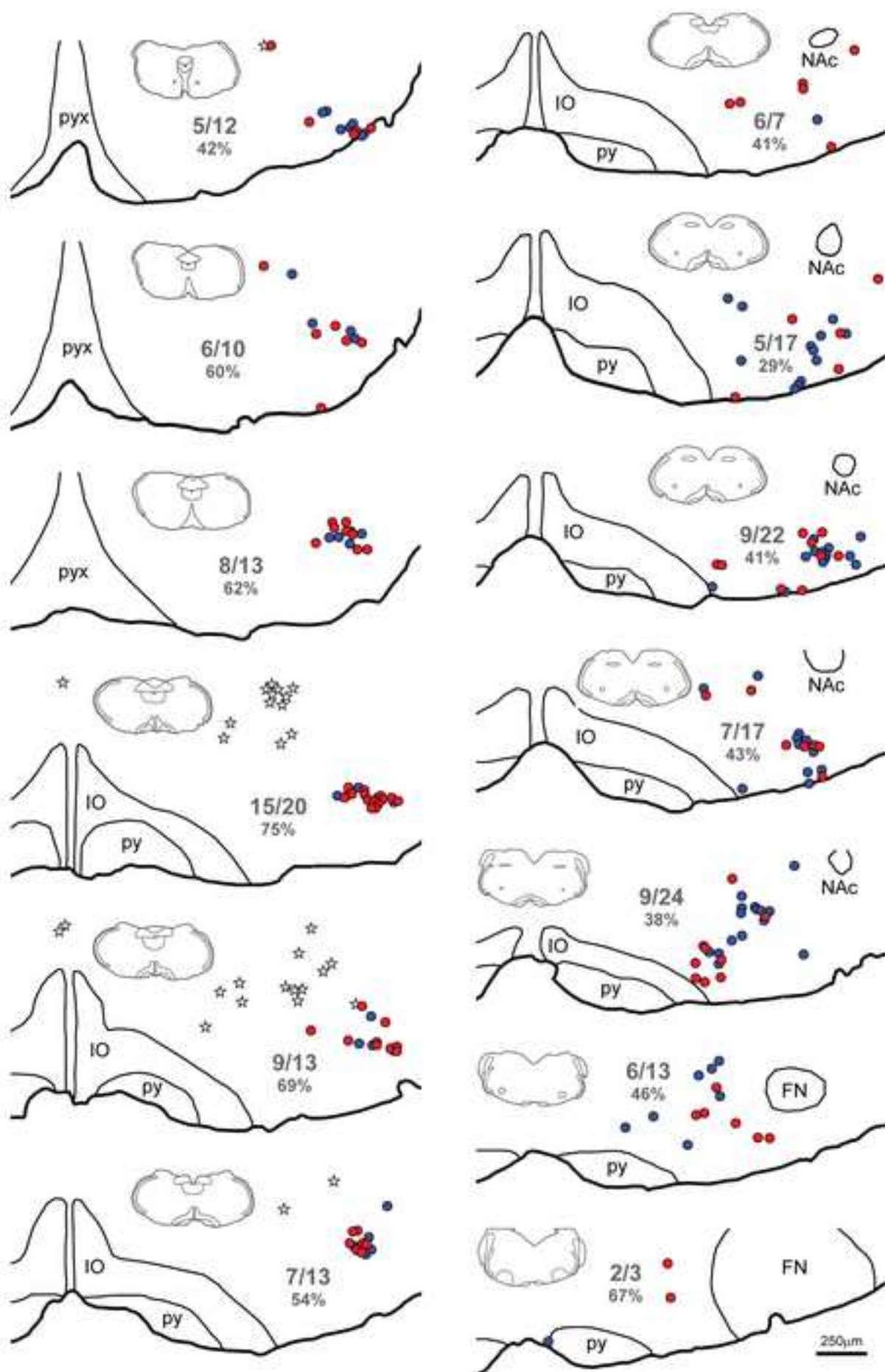
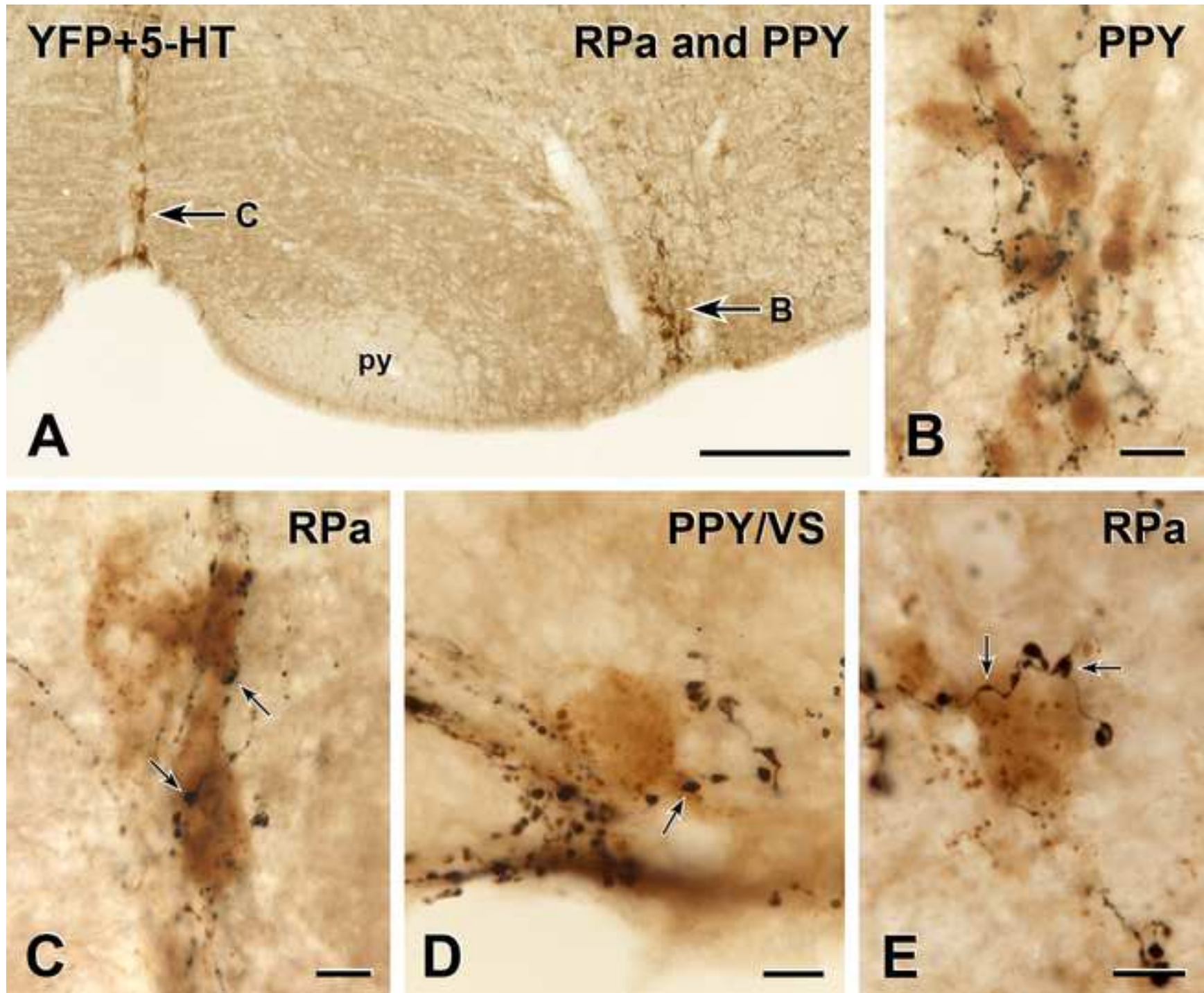


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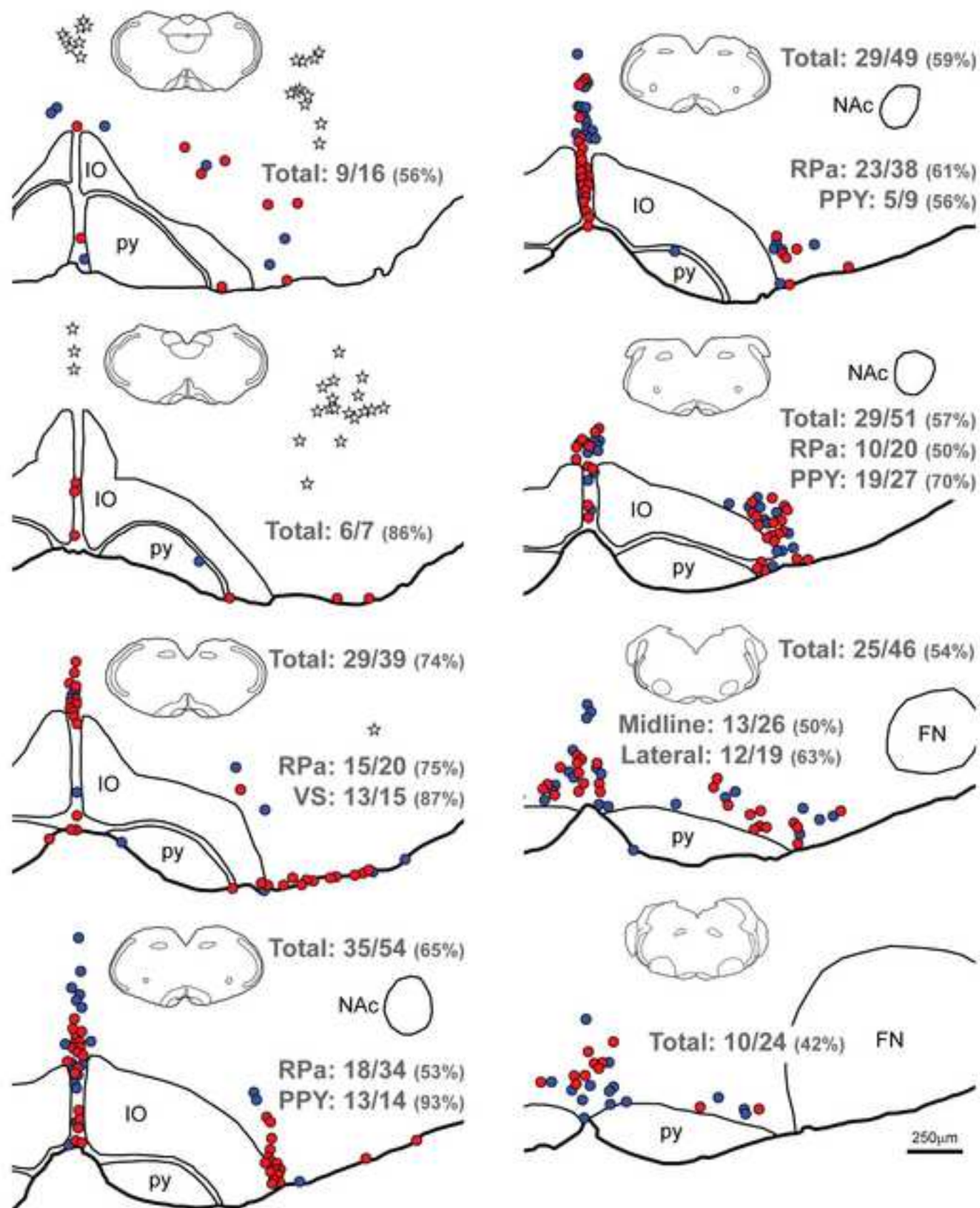
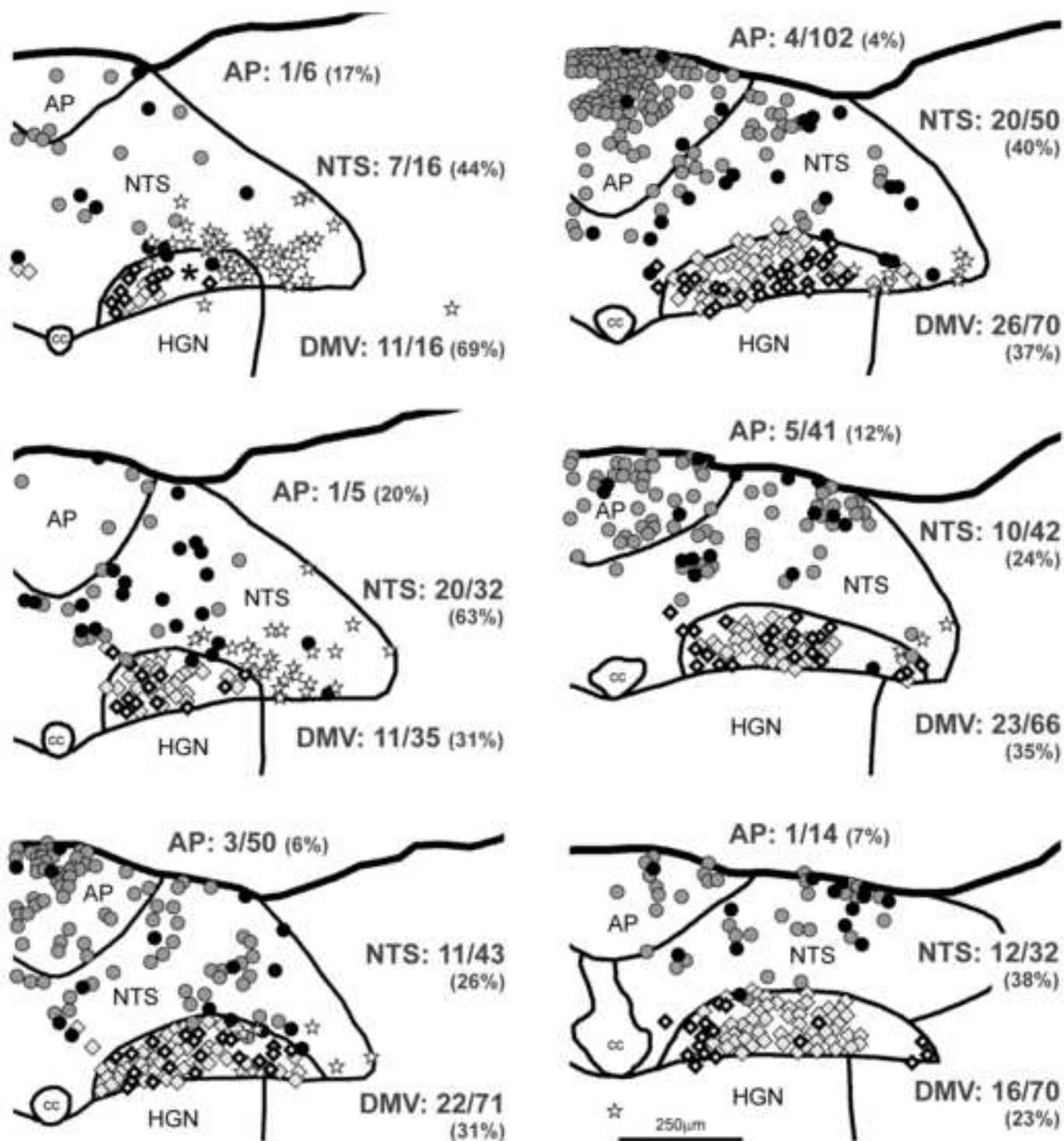


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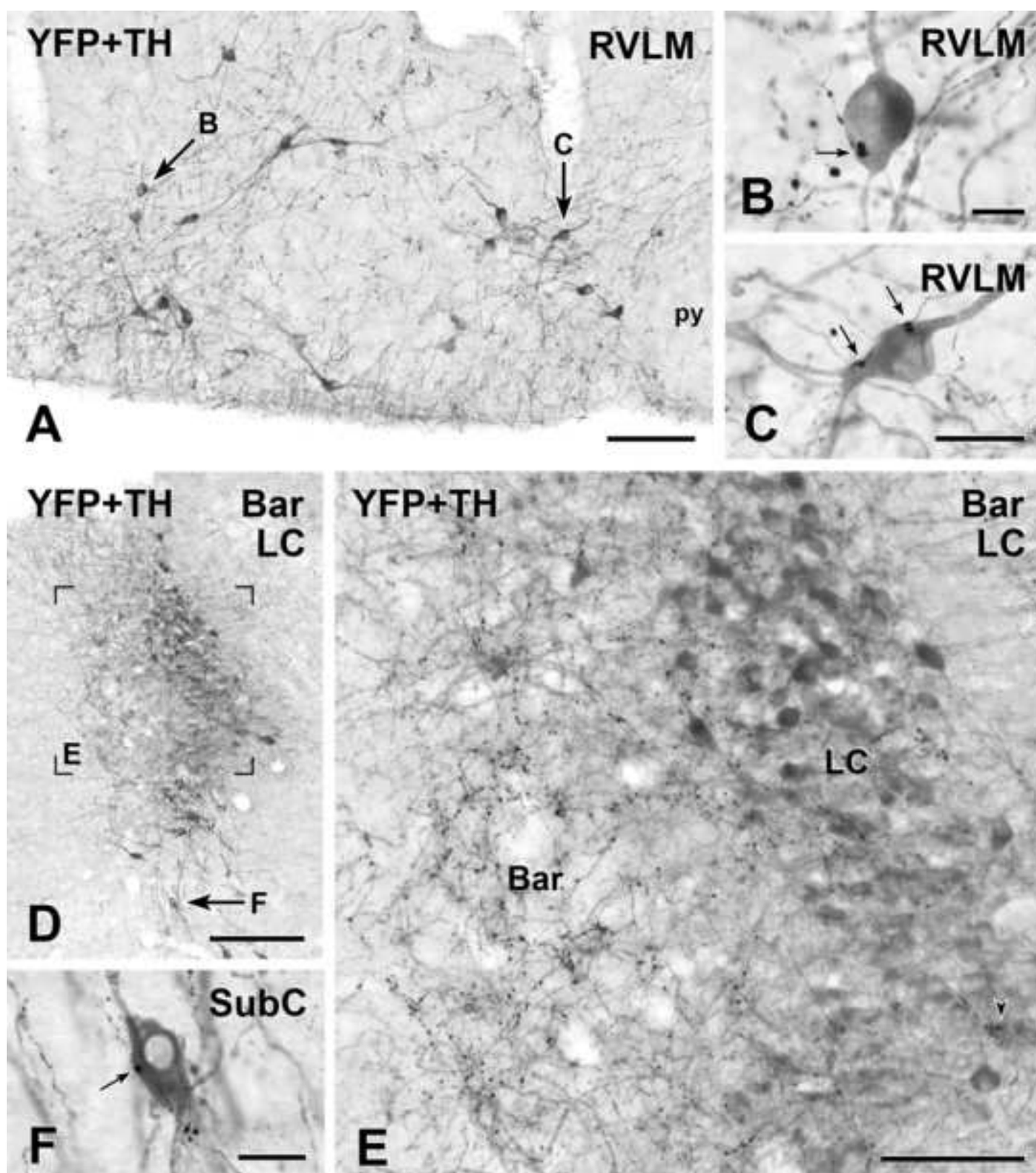


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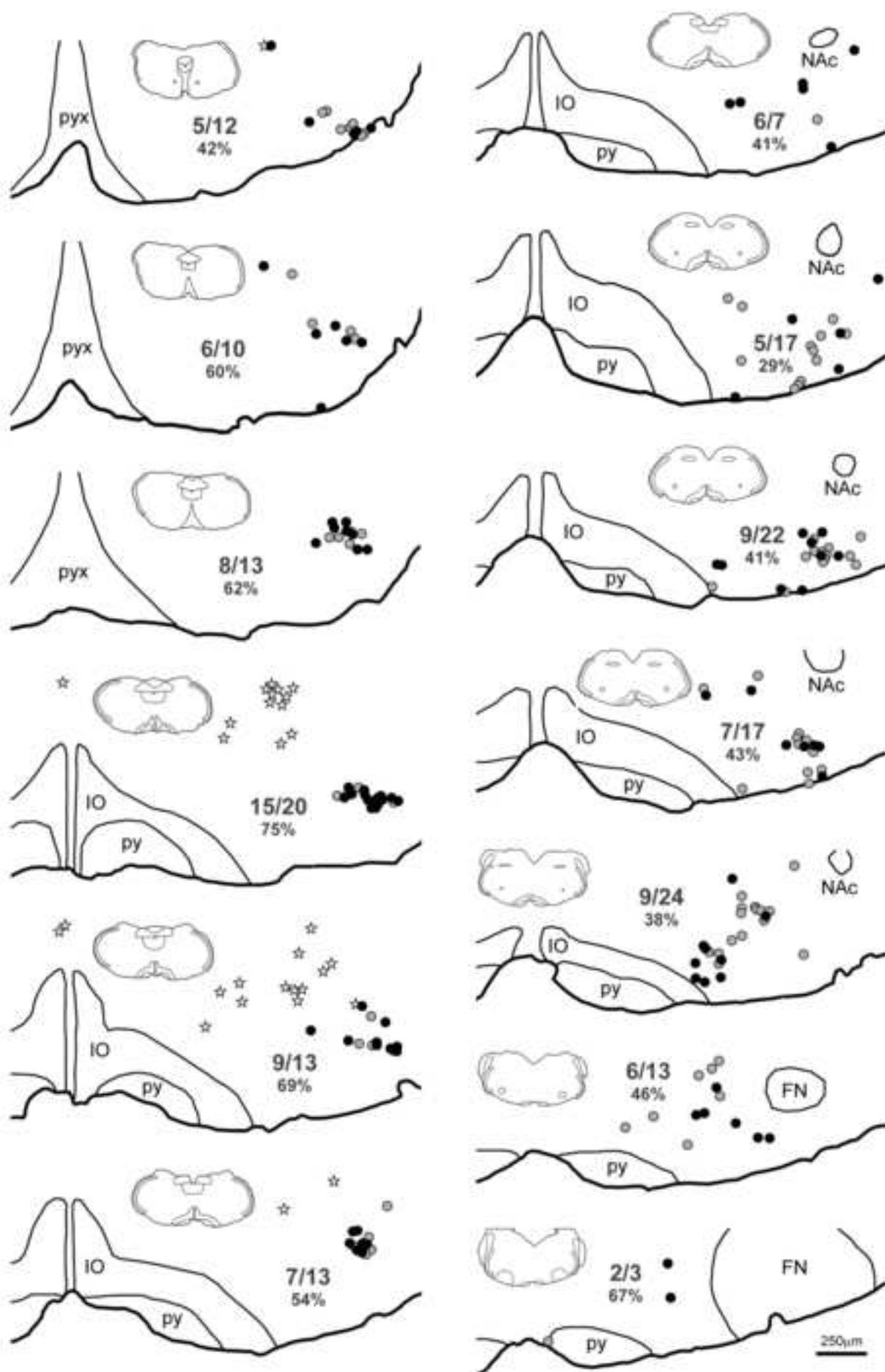


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