Behavioral/Systems/Cognitive

Single Nigrostriatal Dopaminergic Neurons Form Widely Spread and Highly Dense Axonal Arborizations in the Neostriatum

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The axonal arbors of single nigrostriatal dopaminergic neurons were visualized with a viral vector expressing membrane-targeted green fluorescent protein in rat brain. All eight reconstructed tyrosine hydroxylase-positive dopaminergic neurons possessed widely spread and highly dense axonal arborizations in the neostriatum. All of them emitted very little axon collateral arborization outside of the striatum except for tiny arborization in the external pallidum. The striatal axonal bush of each reconstructed dopaminergic neuron covered 0.45-5.7% (mean \pm SD = $2.7 \pm 1.5\%$) of the total volume of the neostriatum. Furthermore, all the dopaminergic neurons innervated both striosome and matrix compartments of the neostriatum, although each neuron's arborization tended to favor one of these compartments. Our findings demonstrate that individual dopaminergic neurons of the substantia nigra can broadcast a dopamine signal and exert strong influence over a large number of striatal neurons. This divergent signaling should be a key to the function of the nigrostriatal system in dopamine-based learning and suggests that neurodegeneration of individual nigral neurons can affect multiple neurons in the striatum. Thus, these results would also contribute to understanding the clinicopathology of Parkinson's disease and related syndromes.

Key words: dopamine; nigrostriatal; Parkinson's disease; reinforcement; striatum; tyrosine hydroxylase

Introduction

Dopamine-containing neurons are distributed in the retrorubral field (RRF), substantia nigra pars compacta (SNc), and ventral tegmental area (VTA), which correspond to cell groups A8, A9, and A10, respectively, in the mammalian mesencephalon (Dahlstroem and Fuxe, 1964). These dopamine-containing neurons have a strong influence on emotion, motivation and cognitive processes mainly by the projection from the VTA to limbic forebrain areas, and on motor control chiefly by the projection from the SNc and RRF to the neostriatum (for review, see Joel and Weiner, 2000; Björklund and Dunnet, 2007). It has also been suggested that dopamine signal serves as a reinforcement signal in those regions (for review, cf. Schultz, 1998). The neostriatum,

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being a main input nucleus of the basal ganglia, receives massive cortical and thalamic inputs and sends inhibitory projections to the external segment of the globus pallidus (GPe), entopeduncular nucleus [internal segment of the GP (GPi)] and substantia nigra pars reticulata (SNr), the latter two structures being output nuclei of the basal ganglia circuit. Furthermore, the neostriatum is known to have a mosaic organization and be composed of patch/striosome and matrix compartments which are distinguishable from each other by the expression of neurochemical markers and by input-output organization (Graybiel and Ragsdale, 1978; Bowen et al., 1981; Herkenham and Pert, 1981; Gerfen, 1984, 1985, 1989; Gerfen et al., 1985, 1987a,b; Donoghue and Herkenham, 1986; Jimenez-Castellanos and Graybiel, 1987; Langer and Graybiel, 1989; Kaneko et al., 1995). In particular, neurons in the striosome compartment are known to send their inhibitory axons selectively to the SNc (Gerfen, 1985), whereas those in the matrix compartment project directly or indirectly to the SNr and GPi (Alexander et al., 1990).

Recently, we have developed a new recombinant viral vector which labels the infected neurons in a Golgi-stain-like manner (Furuta et al., 2001). This vector is based upon a replicationdefective Sindbis virus and designed to express green fluorescent protein (GFP) with a membrane-targeting palmitoylation signal of GAP-43 (palGFP) under the control of a powerful subgenomic promoter of the virus. Sindbis virus is known to have high infec-

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Figure 1. Single-neuron labeling with palGFP Sindbis virus vector. A nigral neuron showing green fluorescence of palGFP (A, A') was immunolabeled for TH (A, A''), and further stained black by the immunoperoxidase method with anti-GFP antibody and diaminobenzidine/nickel reaction (DAB/Ni) (B). Double arrowheads in A and B point to the same small vessel. Another example for palGFP-labeled TH-positive neurons is indicated by arrowheads in C-C''. The black-stained axon fibers of palGFP-labeled neurons densely innervated the striatum (D, E), being distributed both in the striosome/patch and matrix compartments (D) or with preference for the striosome compartment (E). MOR immunoreactivity was used to label the striosome compartments (violet-red areas indicated by arrows in D, E), palGFP-labeled axon fibers (F) were compared with GFP-labeled fibers (G) at a higher magnification. Scale bar in C'' applies to A-C'', that in E applies to D, E, and that in G applies to F, G.

tion efficiency to a broad range of hosts, including neural cells (for review, cf. Xiong et al., 1989; Schlesinger, 1993; Piper et al., 1994; Strauss and Strauss, 1994; Lundstrom, 1999). Furthermore, since a large amount of palGFP is produced and distributed not only on the somatodendritic membrane but also on the axonal membrane of infected neurons (Furuta et al., 2001), the vector has been used as a highly sensitive anterograde tracer in the CNS (Nakamura et al., 2002, 2004, 2005; Ito et al., 2007). In the present study, we used the recombinant Sindbis virus vector as a tool for single-neuron labeling of dopaminergic neurons. We determined the dopaminergic characteristics of the infected neurons immunocytochemically, and visualized highly extensive axonal arborization of single nigrostriatal dopamine neurons. The axonal arborization was further examined in association with the striosome and matrix compartments of the neostriatum to clarify whether single dopamine signals exerted a selective influence upon the two different compartments in the neostriatum.

Materials and Methods

Animals. Forty-two adult male Wistar rats, weighing 250–310 g, were used. Experiments were conducted in accordance with the guidelines of the Committee for Animal Care and Use of the Graduate School of Medicine, Kyoto University, and those for Recombinant DNA Study, Kyoto University. All efforts were made to minimize the suffering and number of animals used in the present study.

Injection of Sindbis virus vector and fixation. Rats were anesthetized by ether inhalation and intraperitoneal injection of chloral hydrate (35 mg/100 g body weight). Five to 10 infectious units of palGFP- or GFP-expressing Sindbis virus vector (Furuta et al., 2001) in 0.5 µl of PBS containing 2% bovine serum albumin were injected bilaterally into the SNc (5.3-5.5 mm posterior to the bregma, 2.2–2.4 mm lateral to the midline, and 7.2-7.7 mm deep from the brain surface) by pressure through a glass micropipette attached to Picospritzer III (General Valve Corporation). After survival for 36-42 h, the rats were reanesthetized by intraperitoneal injection of chloral hydrate (70 mg/100 g body weight), and perfused transcardially with 200 ml of PBS [0.9% (w/v) saline buffered with 5 mM sodium phosphate, pH 7.4], followed by 200 ml of 3% (w/v) formaldehyde, 75% saturated picric acid and 0.1 M Na₂HPO₄ (adjusted with NaOH to pH 7.0) over 30 min. The brains were then removed, cut at the midline into two hemispheric blocks, and postfixed for 4 h at 4°C with the same fixative. After cryoprotection with 30% (w/w) sucrose in PBS, the blocks were cut into 40-µm-thick parasagittal sections on a freezing microtome, and the sections were collected serially in PBS.

Immunofluorescence labeling for tyrosine hydroxylase. The sections including the injection site were observed under epifluorescent microscope Axiophot 2 (Zeiss) with an appropriate filter set (excitation, 450–490 nm; emission, 515–565 nm) for fluorescence of GFP to find SN neurons infected with the virus. All the fol-

lowing incubations described hereafter were performed at room temperature and followed by a rinse with PBS containing 0.3% (v/v) Triton X-100 and 0.02% (w/v) merthiolate (PBS-X). The sections containing palGFP-positive SN neurons were incubated overnight with 1:200diluted mouse ascites containing anti-tyrosine hydroxylase (TH) monoclonal IgG₁ (clone LNC1; Millipore) in PBS-X containing 0.25% (w/v) λ -carrageenan, 0.02% sodium azide and 1% (v/v) donkey serum (PBS-XCD), and then for 4 h with 1 µg/ml AlexaFluor 594-conjugated anti-[mouse IgG] goat antibody (Invitrogen) in PBS-XCD. Under the epifluorescence microscope with appropriate filter sets for GFP fluorescence and AlexaFluor 594 (excitation, 530–585 nm; emission, ≥615 nm),

TH-positive neuron	Position (tier)	Length of axon fibers (in μ m	Striosome		
		In matrix (x)	In striosome (y)	Total (z)	rate (%) ^b
1	Dorsal	638,332	141,601	779,933	18.2
2	Dorsal	402,242	63,660	465,902	13.7
3	Dorsal	130,981	7512	138,493	5.4
4	Dorsal	245,004	33,982	278,986	12.2
5	Dorsal	415,437	19,072	434,509	4.4
6	Ventral	438,992	86,407	525,399	16.4
7	Ventral	288,105	216,055	504,160	42.9
8	Ventral	475,978	131,291	607,269	21.6
Average \pm SD		379,384 ± 155,772	87,448 ± 71,599	466.831 ± 195.198	16.8 ± 12.1

^aThe length of axon fibers was estimated by multiplying the length of axon fibers projected onto a parasagittal plane by $\pi/2$ (when straight fibers having unit length and distributed in three-dimensionally random direction are projected to a plane, the mean length of projected fibers was $2/\pi$).

 $^{b}y/z \times 100.$

palGFP-labeled neurons were examined to determine whether they expressed immunoreactivity for TH (Fig. 1A-A'').

Double immunoperoxidase staining for GFP and μ -opioid receptor. All the sections containing single palGFP-labeled and TH-immunopositive neurons were incubated overnight with a mixture of 0.4 μ g/ml affinitypurified rabbit antibody to GFP (Tamamaki et al., 2000) and 2 μ g/ml affinity-purified guinea pig antibody to μ -opioid receptor (MOR) (Kaneko et al., 1995) in PBS-XCD. The incubation medium for the sections immunolabeled for TH further contained 10% (v/v) normal mouse serum to block the reaction with bound mouse antibody. After a rinse with PBS-X, the sections were incubated for 2 h with 10 μ g/ml biotinylated anti-rabbit IgG donkey antibody (Millipore) in PBS-XCD, and then for 1 h with avidin-biotinylated peroxidase complex (ABC Elite; Vector) in PBS-X. After a rinse with PBS, the bound peroxidase was developed black by reaction for 10-30 min with 0.02% (w/v) diaminobenzidine-4HCl, 10 mM nickel-ammonium sulfate and 0.0003% (v/v) H₂O₂ in 50 mM Tris-HCl, pH 7.6 (Fig. 1B). The reaction was terminated by a rinse with PBS containing 2% (w/v) sodium azide, followed by two rinses in PBS. Subsequently, the sections were further incubated for 2 h with 10 μ g/ml of biotinylated anti-guinea pig IgG donkey antibody (Millipore) in PBS-XCD, and then for 1 h with the avidin-biotinylated peroxidase complex in PBS-X. Finally, the second bound peroxidase was visualized violet-red by reaction for 10-30 min with 0.01% (w/v) Trisaminophenol methane, 0.07% (v/v) p-cresol and 0.003% (v/v) H₂O₂ in 50 mM Tris-HCl, pH 7.6 (Fig. 1D, E) (Kaneko et al., 1994). All the above incubations and reactions were performed at room temperature. The stained sections were serially mounted onto the gelatinized glass slides, dried up, washed in running water, dried up again, cleared in xylene, and finally coverslipped with mounting medium MX (Matsunami).

Reconstruction of single dopamine neurons. The cell body, dendrites, and projecting main axon of stained dopamine neurons were reconstructed under a microscope attached with camera lucida apparatus. In particular, the striatal axon fibers were photographed with a digital camera attached to the microscope with a $20 \times$ objective, and then traced on the photograph with software CANVAS 9 (ACD Systems International) in a computer. The axon fibers in a section were reconstructed one by one onto a parasagittal plane, and superimposed to create the medial view in the computer. For the frontal and dorsal views, the extent of axon fibers in each 40- μ m-thick section was superimposed on frontal and horizontal planes, respectively, with software ImageJ (NIH; http://rsb.info.nih.gov/ ij/). The length of axons was obtained separately in the striosome and matrix compartments by calculating the length of traced lines with "Object Information" function in software Canvas 9.

Localization of cell bodies. The location of infected neurons was determined by cytoarchitecture. After tracing of infected neurons and axon fibers, the coverslips were removed from glass slides and the mounted sections were counterstained for Nissl with Neutral Red (Acros Organics). In addition, control parasagittal sections of rat mesencephalon were serially stained for Nissl and immunostained for calbindin D-28k and TH (supplemental Fig. 1, available at www.jneurosci.org as supplemental material). These control sections were used as a reference for defining



Figure 2. The distribution of the eight neurons labeled with palGFP Sindbis virus vector and immunopositive for tyrosine hydroxylase in the substantia nigra. The locations were projected onto a parasagittal plane 2.25 mm lateral to the midline. Each encircled numeral indicates the position of the neuron with the same number in Table 1. cp, Cerebral peduncle; ic, internal capsule; Zl, zona incerta.

ventral and dorsal tiers of the SNc by comparing cytoarchitecture with chemoarchitecture of calbindin and TH as described previously (Prensa and Parent, 2001).

Statistical analysis. The difference between dorsal and ventral tier of SNc dopamine neurons was determined using the two-tailed Student's *t* test and two-tailed Mann–Whitney *U* test.

Results

Selection of neurons for single-cell tracing

In the present study, we stereotaxically injected Sindbis virus vector for palGFP expression bilaterally into the SN of 42 rats, and thus of 84 hemibrains. The neurons for single-cell tracing were selected as follows: first, the hemiencephalic sections including the injection sites were examined under epifluorescence microscope. The hemibrains which contained only 1-10 palGFPlabeled SN neurons were selected. Seventy nigral neurons in 25 hemibrains were thereby processed for further analysis. Second, 70 palGFP-labeled neurons were examined for the expression of TH immunofluorescence, a marker for dopaminergic neurons in the SN (Fig. 1A-A'', C-C''). Twenty-one of 70 neurons were positive for TH immunoreactivity. Finally, the hemiencephalic sections containing these dopaminergic neurons were further stained by the immunoperoxidase method with anti-GFP antibody (Fig. 1B, D-G). Of the 21 stained neurons, the axons of 13 neurons were entangled or overlapped in the striatum with those of another neurons, and thus inseparable for reconstruction.



Figure 3. Camera lucida reconstruction of SNcd neuron #1.*A*, *B*, The axon fibers in the striatum (*A*) and dendrites (*B*) in the SNc were projected onto a parasagittal plane and superimposed from the medial side. *C*, The dorsal and frontal views of the intrastriatal axonal arborization were reconstructed and compared with the medial view. Red and blue lines in the striatum indicate the axon fibers located in the striosome and matrix compartments, respectively. Red fibers at the most rostral portion in *A* were mostly located in the MOR-positive subcallosal streak. *D*, The axon gave rise to only minor collaterals in the external segment of the GPe. ac, Anterior commissure; cc, corpus callosum; cp, cerebral peduncle; CPu, caudate—putamen (neostriatum); Hpc, hippocampus; ic, internal capsule; LV, lateral ventricle; ml, medial lemniscus; ot, optic tract; STh, subthalamic nucleus; str, superior thalamic radiation; Th, thalamus; Zl, zona incerta.

Thus, only 8 neurons were selected for the present morphological analysis of single neurons. In 5 cases of them (Table 1, neurons #2, 5-8), a hemibrain contained only one TH-immunopositive neuron. The other 3 neurons were found in a hemisphere, but fortunately separable from one another (Table 1, neurons #1, 3, 4). Since TH-negative SN neurons (mainly located in the SNr) had little, if any, axonal projection to the striatum, it was easy to

separate axons of TH-positive neurons from those of TH-negative neurons.

These reconstructed neurons were subdivided into two groups based on the location of their cell bodies in the SNc (Fig. 2). By identifying the location of the cell bodies with Nissl counterstaining and by comparing Nissl cytoarchitecture with calbindin-immunoreactive structure in control parasagittal sections (supplemental Fig. 1, available at www.jneurosci.org as supplemental material), five neurons (Table 1, TH-positive neurons #1-5) were classified as neurons of the dorsal tier of the SNc (SNcd), and three (#6-8) were as neurons of the ventral tier (SNcv). The region around each palGFP-positive neuron was darkly immunostained, probably due to the presence of extracellularly leaked palGFP (Fig. 1*B*), suggesting an extremely strong expression of protein by the subgenomic promoter of Sindbis virus. However, no spread to adjacent cell bodies was observed.

Intrastriatal axons and striosome–matrix organization of the striatum

Simultaneously with GFP immunoperoxidase staining, immunoreactivity for MOR was visualized in violet-red (Fig. 1D, E, arrows), and used as a marker for the striosome compartment of the striatum (Kaneko et al., 1995). As shown in Figure 1, D and E, some axon fibers of palGFPlabeled TH-positive SN neurons seemed to prefer the matrix or striosome compartment to the other compartment. These intrastriatal axons were relatively thick and seemed continuous to varicosities, making it difficult to count varicosities (Fig. 1F). Furthermore, axon fibers located in the striosome compartment showed little morphological difference compared with those in the matrix compartment.

To examine the effect of the palmitoylation signal on the morphology of intrastriatal axon fibers, we compared palGFPlabeled axon fibers and those labeled with unmodified green fluorescent protein (GFP). As shown in Figure 1, F and G, there was no obvious difference between the two fibers. Since intrastriatal axon fibers were not completely labeled with GFP (supplemental Fig. 2, available at www. jneurosci.org as supplemental material),

we could not compare global properties such as the total axonal length, but local characteristics between palGFP-labeled axon fibers and well labeled fibers with GFP. For example, the fiber width and interbranch interval of 80 randomly selected GFP-labeled axon fibers were $0.82 \pm 0.11 \ \mu\text{m}$ and $31.2 \pm 19.4 \ \mu\text{m}$, respectively. These data were not statistically different ($p \ge 0.72$ by the two-sided *t* test) from those of 80 palGFP-labeled intrastri-

atal axon fibers, the fiber width and interbranch interval of which were 0.83 ± 0.13 μ m (mean \pm SD) and $30.9 \pm 21.6 \mu$ m, respectively (for further detail, see supplemental data, available at www.jneurosci. org as supplemental material). Thus, as far as we could compare, no significant difference was detected between the GFPlabeled and palGFP-labeled axon fibers.

Axons of single nigrostriatal dopaminergic neurons

The axons of all eight nigrostriatal dopaminergic neurons were isolated and reconstructed by camera lucida drawings as shown in Figures 3–7. The representative axonal distribution of neuron #2 is shown in the photographs of supplemental Figure 3, available at www.jneurosci.org as supplemental material. In all the reconstructed neurons, the main axon emerged from the cell body or proximal dendrite in the SNc without local axon collaterals around the cell body, then passed through the internal capsule, and then headed directly to the striatum. All but one of the axons emitted very short curly collateral projections in the GPe before entering the striatum (Figs. 3D, 5D-7D, neuron #2 was the exception). No such collaterals were observed in the GPi. In contrast to the minimal collateral projections within extrastriatal structures, the individual nigrostriatal axons formed widely spread and highly dense axonal bushes within the striatum (Figs. 3A-7A, 3C-7C). The axonal arborizations reconstructed in the striatum were more extended and much denser than those of the nigrostriatal axons described previously (Gauthier et al., 1999; Prensa and Parent, 2001), suggesting that the present method using the palGFPexpressing Sindbis virus vector was a



Figure 4. Camera lucida reconstruction of SNcd neuron #2. This neuron did not emit axon collateral in the GPe, but took a loop-making meandering course through the subthalamic nucleus (STh; **D**). Main axons meandering in the STh were found only in neurons #2 and #4. See the legend of Figure 3 for further detail.

highly sensitive and efficient technique for visualizing the arborization of single axons. The eight SN neurons had a total axonal length, on average, of ~467,000 μ m (Table 1, #1–8), whereas the seven representative SN neurons of the previous study (Prensa and Parent, 2001) had that of 36,000 μ m (see supplemental Table 1, estimated from the published data, available at www.jneurosci.org as supplemental material), <1/10 of the previous studies was largely underestimated. The intrastriatal axonal arborizations of neurons #1, 2, 5, 7 and 8 innervated mainly the dorsal part of the striatum, and those of neurons #3, 4 and 6 mainly the middle and ventral parts of the striatum and the accumbens nucleus.

The five SNcd neurons had total axonal lengths of $140,000-780,000 \ \mu m$ (Table 1, #1–5). The typical wide spread arborizations of intrastriatal axons was observed in dopamine neurons #1, #2, #4 and #5, and those of neurons #1 and #2 were reconstructed in Figures 3 and 4, respectively. The arbors occupied oval volumes that extended 1.5–2.0 mm rostrocaudally, 0.7–1.0 mm dor-

soventrally and 1.0–1.5 mm mediolaterally. The total axonal lengths of these four neurons ranged from 280,000–780,000 μ m (Table 1). The striatal volume covered with the dense axonal plexus was estimated to be 0.09–1.12 mm³ (Table 2) and reached a maximum of 5.7% of the total neostriatal volume on one side (19.81 ± 0.63 mm³, n = 3). SNcd dopamine neuron #3 was an exceptional case with a smaller and mediolaterally flat axonal bush, shown in Figure 5. This arbor formed a small terminal field in the fundus of the caudoputamen as a main bush with a large diameter of ~500 μ m and a smaller bush arbor extending rostroventral to the main bush. Although this neuron had the smallest axonal bushes in the present study, the total axonal length in the striatum was 140,000 μ m (Table 1).

The three SNcv neurons had total axonal lengths of $500,000 - 610,000 \ \mu m$ (Table 1, #6–8). Two of them (#7 and #8) are illustrated in Figures 6 and 7, respectively. The arbors of SNcv neurons #6 and #8 formed single oval volumes with a major diameter of >1 mm as had shown in SNcd dopamine neurons. Neuron #7 formed two axonal bushes, large and small ones, similar to SNcd

Table 2. Volume occupied by TH-positive axons and axon fiber density in the striatum

TH-positive neuron	Volume occupied by axons (in mm ³) ^a			Axon density (in μ m/mm ³) b		Striosome-
	In matrix (a)	In striosome (b)	Total (c)	In matrix (x/a)	In striosome (y/b)	sity ratio ([y/b]/[x/a])
1	0.94	0.18	1.12	678	802	1.18
2	0.59	0.07	0.66	681	854	1.25
3	0.08	0.01	0.09	1691	1130	0.67
4	0.25	0.03	0.28	997	1155	1.16
5	0.59	0.06	0.65	703	294	0.42
6	0.53	0.06	0.59	843	1370	1.64
7	0.42	0.07	0.49	693	3139	4.53
8	0.36	0.08	0.44	1310	1715	1.31
Average \pm SD	0.47 ± 0.26	0.07 ± 0.05	0.54 ± 0.30	950 ± 371	1307 ± 851	1.52 ± 1.27

^aIn a section, the matrix or striosome area occupied by the axon fibers was measured in a polygon circumscribed to the axonal arborization. The volume in a section was calculated by multiplying those areas with the thickness of the section, and these volumes in single sections were summed up to the whole volume.

^bData x and y were taken from Table 1.



Figure 5. Camera lucida reconstruction of SNcd neuron #3. Of the eight neurons examined in the present study, this neuron had the smallest axonal arborization in the striatum. See the legend of Figure 3 for further detail.

neuron #3, although the arborization was much bigger than that of neuron #3.

Comparison between single-axon arbors innervating the striosome and matrix compartments of the neostriatum

Most intrastriatal axon fibers of the palGFP-labeled dopaminergic neurons were distributed nonselectively in the striosome and matrix compartments (Fig. 1D), but some fibers appeared to innervate either the striosome or the matrix compartment preferentially (Fig. 1E). Because the labeled fibers possessed many ambiguous varicosities in the striatum (Fig. 1F, G) and synaptic specializations are not often found at large varicosities but frequently at the thinner parts of dopaminergic axons (Pickel et al., 1981; Arluison et al., 1984; Freund et al., 1984; Zahm, 1992), the intrastriatal length of axon fibers, instead of the density (or the number) of varicosities, was compared between striosome and matrix compartments to evaluate quantitatively the contribution of single dopaminergic neurons to each compartment.

The striosome compartment including the subcallosal streak has been reported to occupy 10-13% of the total neostriatal volume in the rat (Johnston et al., 1990; Desban et al., 1993). Two SNcd neurons #3 and #5 sent only 4-5% of their axon fibers to the striosome compartment (Table 1), and thus appeared to prefer the matrix compartment over the striosome compartment. In contrast, all three SNcv neurons preferred the striosome compartment, although proportionally the striosome innervation constituted only a minority of their total axonal length (16-43%). On average, five SNcd neurons sent $10.8 \pm 5.2\%$ (mean \pm SD) of axon fibers to striosomes, and three SNcv neurons

projected 27.0 \pm 11.5% of their axonal arbors to striosomes. The p-value of difference between the two groups was 0.0554 by a two-tailed Student's t test, just failing statistical significance. We further examined whether or not SNcd and SNcv neurons showed the difference in local preference for either compartment by evaluating their axonal preference in the regions where the axon fibers were distributed (Table 2). We separately measured the striosome and matrix volumes which were covered by the projecting axon fibers, and then calculated the axon fiber density in the striosome and matrix compartments. In 3 of the 5 SNcd neurons and all of the 3 SNcv neurons, the axon fiber density in the striosome compartment was higher than that in the matrix compartment (Table 2, striosome-matrix density ratio of >1). Furthermore, the striosome-matrix density ratio of SNcv neurons were significantly higher than that of SNcd neurons (p = 0.0357 by the two-tailed U test).Thus the latter comparison partly supported the previously reported finding that SNcv neurons innervate the striosome/patch compartment and that SNcd neurons project preferentially to the matrix compartment (Gerfen et al., 1987a).

Notably, even for a given nigrostriatal neuron, the axonal preference for compartments was different between its axonal bushes. SNcd neuron #3 and SNcv neuron #7 both had an isolated small axonal bush that were preferentially distributed in striosomes, despite an overall wide distribution of the entire axon (Figs. 5, 6). This pattern indicates that the target preference of a single axon could be different depending on the striatal terminal fields it innervates, and that dopamine-containing innervation might not be so simple as

hypothesized in the simple dopamine-dependent learning model of the basal ganglia circuit.

In summary, although each nigrostriatal dopamine neuron showed some preference for the striosome or matrix compartment, all the dopamine neurons innervated both the striosome and matrix compartments. Furthermore, even a single dopamine axon altered its preference for the compartments in a field-byfield manner within the striatum.

Discussion

Analysis of the entire axonal arborizations of single dopaminergic nigrostriatal neurons with a new virus tracer demonstrated that the intrastriatal arbors of nigrostriatal neurons were much wider and denser than previously reported (Gauthier et al., 1999; Prensa and Parent, 2001), where single nigrostriatal axons were examined by a labeling method with a small injection of biotinylated dextran-amine into the rat SN. Moreover, although some axon arbors exhibited local preferences for one or the other compartment, all the single nigrostriatal axons innervated both striosome and matrix compartments of the striatum. A further critical



Figure 6. Camera lucida reconstruction of SNcv neuron #7. The axons of this SNcv neuron showed the highest preference for the striosome compartment in the present results and made two bushes in the neostriatum. Red fibers at the top of *A* were located in MOR-positive subcallosal streaks. See the legend of Figure 3 for further detail.

finding in the present study is that, except for negligible arbors in the GPe, dopaminergic neurons here had very little arborization outside of the striatum.

Compared with previous reports on the nigrostriatal projection (Gauthier et al., 1999; Prensa and Parent, 2001), the largest difference found was the extremely high density of the intrastriatal arborization of the nigrostriatal axons we reconstructed. This difference may be a consequence of much higher sensitivity of the present single-cell tracing method using the palGFP Sindbis virus vector. It is possible that palGFP expression might have affected the axonal arborization of dopamine neurons, given that overexpression of GFP with a palmitoylation signal of GAP-43 N terminus [1–14] was reported to induce filopodia formation on the surface membrane of COS-7 cells and on the dendrites of cultured hippocampal neurons (Gauthier-Campbell et al., 2004). However, this possibility is unlikely for the following reasons (see also supplemental data, available at www.jneurosci.org as supplemental material). (1) The effect reported was on dendritic membranes, but not on axonal membranes. In the present study, neither filopodia-like nor growth cone-like structures were observed



Figure 7. Camera lucida reconstruction of SNcv neuron #8. Although the cell body of this neuron was located at the cell cluster region of the SNcv, the axonal arborization was very similar to that of SNcd neuron #1 (Fig. 3). See the legend of Figure 3 for detail.

on axon fibers of TH-positive neurons. (2) Even on the dendritic membrane of palGFP-expressing neurons, we detected no morphological differences compared with the previous morphological studies of SNc neurons (Juraska et al., 1977; Preston et al., 1981; Tepper et al., 1987). (3) In a comparative study using GFP Sindbis virus vector, GFP-positive nigrostriatal axon fibers showed characteristics similar to those of palGFP-positive fibers, although the axon fibers were not completely visualized throughout their arborization (supplemental Fig. 2, available at www. jneurosci.org as supplemental material). No difference in the width or interbranch interval of axon fibers was detected between palGFP- and GFP-labeled axon fibers. Therefore, we conclude that palGFP expression seemed to have little, if any, morphological effect on the dopaminergic axons, and thus that naive nigrostriatal dopaminergic neurons may have as widely spread and highly dense axonal arborization in the neostriatum as the neurons infected with palGFP Sindbis virus.

A second major difference from previous findings concerns the nigrostriatal innervation of striosomes and matrix. The topography of the substantia nigra itself has been reported to provide the organization of the nigrostriatal projection, whereby neurons in the SNcd project chiefly to the matrix, whereas those in the SNcv send axons mainly to the striosome/ patch compartment in the rat brain (Gerfen et al., 1987a). In addition, differential nigrostriatal projections in association with striosome and matrix compartments have been reported in cat and primate brains (Jimenez-Castellanos and Graybiel, 1987; Langer and Graybiel, 1989). A segregation of these projections was partially supported by the present findings. There was some preference of SNcd neurons for the matrix compartment and some preference of SNcv neurons for striosomes, but we found that single SNcd or SNcv neurons basically innervated both the striosome and matrix compartments. This means that identical dopamine signals can be sent simultaneously to both striosome and matrix neurons.

Some reports favor a projection from striosomes to SNc dopaminergic neurons, although a recent report on monkey brain is in disagreement (Levesque and Parent, 2005): Favoring striosome-to-SNc projections are the findings that fluorescent retrograde tracers injected into rat mesencephalon including the SNc but not the SNr, mostly retrogradely labeled neurons in the striosome/patch compartment (Gerfen, 1985); and immunoreactivity for substance P, which was produced by both patch and matrix neurons, is distributed mainly in the neuropil of the SNr, whereas immunoreactivity for preprodynorphin, expressed more intensely in striosome

neurons, is distributed more densely in SNc neuropil than SNr neuropil (Lee et al., 1997). Thus, it seems most likely that striosomes contribute to the production of a dopamine signal by SNc neurons (Schultz, 1998; Doya, 2000). This dopamine signal is sent back to the neostriatum for the long-term modification of corticostriatal and/or thalamostriatal synapses (Calabresi et al., 2000, 2007), resulting in the updates of responsiveness of striatal neurons. The present results indicate that a large number of matrix and striosome neurons are affected by the single dopamine signal emitted by a single SNc neuron.

The present results indicate that, on average, 2.7% of neostriatal neurons (Table 2) $(0.54 \text{ mm}^3/19.81 \text{ mm}^3 = 0.027)$ and, at the maximum, 5.7% of neurons (1.12/19.81 = 0.057) are under a strong influence of a single dopamine neuron. Since the rat neostriatum contained 2,790,000 neurons in one side (Oorschot, 1996), and the volume innervated by a single dopamine neuron was, on average, 2.7% of the total neostriatal volume in the present study, ~75,000 striatal neurons are estimated to be influenced by a single dopaminergic neuron. The whole striatum might not need to receive a single dopamine signal, because the basal ganglia circuit is functionally segregated into several parallel circuits such as motor, oculomotor, prefrontal, and limbic circuits (for review, see Alexander et al., 1990). However, it is known that electrical synapses are formed between SNc dopaminergic neurons (Grace and Bunney 1983; Vandecasteele et al., 2005), suggesting a mechanism for synchronization or coactivation of dopaminergic neurons. If a single dopamine signal covers the whole striatum, the present results indicate that at least 37 dopaminergic neurons (19.81/0.54 \approx 37) have to synchronize and convey the same dopamine signal.

Because the rat SNc contained 3500-7200 neurons in one side (Andén et al., 1966; Halliday and Tork, 1986; Oorschot, 1996), it is estimated that a neuron in the rat neostriatum might be under the influence of 95-194 dopaminergic neurons on average $(3500-7200 \div (19.81/0.54) \approx 95-194)$. This highly overlapping innervation of dopaminergic neurons raises the question of its necessity. The question might partially be answered by the fact that animals have many different kinds of reward-based signals, such as those related to appetitive and sexual behaviors. However, since it is unlikely that animals have a hundred kinds of reward-based signals, another justification seems necessary for this redundancy. The large margin of safety in dopaminergic neurons has been reported in human brains diagnosed as idiopathic Parkinson's disease, as extensive loss (>70%) of neurons in the SNc occurs in most patients before parkinsonian symptoms appear (Hirsch et al., 1988). Since neurons are labile elements and are relatively easily lost from many causes, it is a good strategy for the system composed of such unreliable elements to maintain a high level of redundancy. Thus our findings have important implications for an understanding of the clinicopathology of Parkinson's disease and related neurodegenerative disorders.

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