RESEARCH ARTICLE



Morphological characterization of HVC projection neurons in the zebra finch (*Taeniopygia guttata*)

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1 | INTRODUCTION

The HVC nucleus of the zebra finch telencephalon is a crucial link in the song production pathway, sending descending projections from the forebrain to the brainstem and ultimately driving muscles important for vocal production (Bottjer, Glaessner, & Arnold, 1985; Goller & Cooper, 2004; Nottebohm, Kelley, & Paton, 1982; Nottebohm, Stokes, & Leonard, 1976; Wild, 1993). Within that region, the premotor cell population that projects to the downstream motor cortical region RA (robust nucleus of the arcopallium) are referred to as HVC_(RA) neurons. Additionally, another group of cells (HVC_(X) neurons) project to the avian basal ganglia (Dutar, Vu, & Perkel, 1998; Kubota & Taniguchi, 1998; Mooney, 2000). HVC_(X) neurons are thought to be critical for song acquisition during development (Scharff, Kirn, Grossman, Macklis, & Nottebohm, 2000; Scharff & Nottebohm, 1991; Sohrabji, Nordeen, &

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Abstract

Singing behavior in the adult male zebra finch is dependent upon the activity of a cortical region known as HVC (proper name). The vast majority of HVC projection neurons send primary axons to either the downstream premotor nucleus RA (robust nucleus of the arcopallium, or primary motor cortex) or Area X (basal ganglia), which play important roles in song production or song learning, respectively. In addition to these long-range outputs, HVC neurons also send local axon collaterals throughout that nucleus. Despite their implications for a range of circuit models, these local processes have never been completely reconstructed. Here, we use *in vivo* single-neuron Neurobiotin fills to examine 40 projection neurons across 31 birds with somatic positions distributed across HVC. We show that $HVC_{(RA)}$ and $HVC_{(X)}$ neurons have categorically distinct dendritic fields. Additionally, these cell classes send axon collaterals that are either restricted to a small portion of HVC ("local neurons") or broadly distributed throughout the entire nucleus ("broadcast neurons"). Overall, these processes within HVC offer a structural basis for significant local processing underlying behaviorally relevant population activity.

KEYWORDS

axon collaterals, premotor, sequence generation, songbird, vocal production

Nordeen, 1990), but the role of these neurons in the production and maintenance of adult song is less well understood (Scharff et al., 2000). During singing, both HVC projection neuron types exhibit sparse bursts that form sequences of activity at the network level, with different neurons precisely active at different moments throughout the song (Hahnloser, Kozhevnikov, & Fee, 2002; Hamaguchi, Tschida, Yoon, Donald, & Mooney, 2014; Kozhevnikov & Fee, 2007; Long, Jin, & Fee, 2010; Lynch, Okubo, Hanuschkin, Hahnloser, & Fee, 2016; Picardo et al., 2016; Vallentin & Long, 2015). The importance of local process-ing within HVC for the generation of these neural sequences remains controversial, in part because the structure of circuits within HVC are only partially understood (Kornfeld et al., 2017; Kosche, Vallentin, & Long, 2015; Mooney & Prather, 2005; Stauffer et al., 2012).

A major gap in our knowledge of HVC function stems from an incomplete picture of the local wiring within that nucleus, especially with respect to the axon collaterals of excitatory projection neurons. These local processes have been shown to form interconnections



within HVC (Kornfeld et al., 2017; Kosche et al., 2015; Mooney & Prather, 2005) and have been strongly implicated in the generation of sequences of neural activity (Galvis, Wu, Hyson, Johnson, & Bertram, 2017; Gibb, Gentner, & Abarbanel, 2009; Hanuschkin, Diesmann, & Morrison, 2011; Jin, 2009; Long et al., 2010). Although the basic morphological characteristics of HVC neurons have been investigated through bulk labeling (Benton, Cardin, & DeVoogd, 1998; Fortune & Margoliash, 1995; Nixdorf, Davis, & DeVoogd, 1989; Vates, Broome, Mello, & Nottebohm, 1996; Yip, Miller-Sims, & Bottjer, 2012) or partially reconstructed single-cell fills (Dutar et al., 1998; Katz & Gurney, 1981; Mooney, 2000), surprisingly little is known about their anatomy because these reconstructions were incomplete. To better understand the impact of single neurons within the circuit, we generated the first complete reconstructions of HVC projection neuron morphology through in vivo single-cell labeling and cell tracing within the entire volume of HVC. From these reconstructions, we characterize dendritic morphology of individual neurons as well as a wide range of projection patterns of axon collaterals that extend throughout the volume of that structure. Both HVC(RA) and HVC(X) neurons have significant local axonal projections, whose projection patterns form a basis for local processing within this sequence-generating cortical circuit.

2 | MATERIALS AND METHODS

2.1 Animals

We used adult (>90 days post hatch) male zebra finches that were obtained from an outside breeder and maintained in a temperature- and humidity-controlled environment with a 12/12 hr light/dark schedule. All animal maintenance and experimental procedures were performed according to the guidelines established by the Institutional Animal Care and Use Committee at the New York University Langone Medical Center.

2.2 Surgery

To conduct two-photon guided cell labeling, we employed a previously described in vivo microscopy preparation (Kornfeld et al., 2017). Briefly, animals were anesthetized with 1–3% isoflurane in oxygen, and a retrograde dextran tracer conjugated to tetramethylrhodamine (fluoro-Ruby, MW: 10,000; Invitrogen, Carlsbad, CA) was injected into either RA or Area X. To gain optical access to HVC, a craniotomy (1 mm \times 1 mm) was prepared using stereotaxic coordinates, the dura was removed, and a cover glass (#0 thickness, Warner Instruments, Hamden, CT) was implanted over the exposed brain with dental acrylic (Cooralite Dental MFG, Diamond Springs, CA). A small metal head plate with two tapped holes was implanted at the anterior part of the skull using dental acrylic to enable head fixation during two-photon imaging.

2.3 | Two-photon guided cell labeling

All cell fills were conducted while the animal was anesthetized (1-3% isoflurane in oxygen). On the day of the experiment, a region of dense retrograde labeling in HVC was defined using two-photon microscopy

(MOM, Sutter Instrument Company, Novato, CA), and a small pipette access hole was drilled in the cover glass adjacent to this region. In each animal, one to two cells were targeted under two-photon guidance and loaded with 3% Neurobiotin (Vector Labs, Burlingame, CA) in 150 mM K-Gluconate using a glass electrode (4–6 M Ω) in a juxtacellular configuration by applying 250 ms current pulses (2-15 nA; 2 Hz) for \sim 10 min (Kornfeld et al., 2017; Narayanan et al., 2014; Pinault, 1996). Pipettes were visualized by adding 40 µM of Alexa 488 (Invitrogen) to the internal solution. The activity of $HVC_{(RA)}$ and $HVC_{(X)}$ neurons was monitored during the fill process using a Multiclamp 700B amplifier (Molecular Devices). Successful Neurobiotin loading was often accompanied by transfer of the Alexa fluorophore into the cell body, providing visual confirmation. Following single-cell dye injection, the pipette was slowly withdrawn from the cell and removed from the brain. Kwik-Cast (World Precision Instruments, Sarasota, FL) was then applied over the exposed tissue and the animal was removed from anesthesia.

2.4 Histological procedures

After 1-3 hr following the completion of juxtacellular filling, birds were anesthetized with pentobarbital sodium and perfused transcardially with 4% w/v paraformaldehyde (EMS, Hatfield, PA). Brains were removed from the skull using a surgical scoop, immersed in 4% paraformaldehyde for 3-5 days to achieve thorough fixation, and incubated in phosphate buffer for an additional 1-3 days to decrease endogenous peroxidase activity. To prepare sections, the brain was cut along the midline, mounted on the sagittal surface with cyanoacrylate, and stabilized with 3% agarose. Parasagittal sections (100 µm thickness) of HVC were cut using a vibratome (Leica VT1000S). Slices were washed five times with phosphate buffer and treated with 3% H₂O₂ to further reduce endogenous peroxidase activity. Slices were then immersed overnight at 4°C in a solution containing avidin/biotin complexes and 0.5% Triton X-100 in phosphate buffer (Vector Labs and Sigma-Aldrich, St. Louis, MO respectively) to tag the Neurobiotin with peroxidase complex. On the following day, slices were washed five times with phosphate buffer and then immersed in a solution containing 2.3 mM diaminobenzidine (DAB, Sigma-Aldrich) and 0.01% H₂O₂ in phosphate buffer to label processes containing Neurobiotin. Slices were then washed and mounted on slides with Vectashield (Vector Labs) or Mowiol (Sigma-Aldrich) mounting medium.

2.5 | Post-histological imaging and reconstruction

A full description of our imaging and reconstruction procedures was published previously (Kornfeld et al., 2017). Only well-filled $HVC_{(RA)}$ neurons were selected for reconstruction, specifically those in which the soma, dendrite, and axon were all labeled without interruptions and with clearly labeled dendritic spines and presynaptic boutons. The cells that passed these criteria were imaged using light microscopy with a custom-designed high-resolution mosaic/optical-sectioning brightfield microscope system (Oberlaender, Bruno, Sakmann, & Broser, 2007; Oberlaender et al., 2009). Neuronal branches in each histological section were reconstructed in 3D using NeuroMorph (Oberlaender et al.,

2007). Reconstructions were performed without prior knowledge about the projection target (apart from the descending axon), and branches were designated as belonging to axons or dendrites based on the presence of boutons or spines, respectively. Automated tracing results from each histological section were manually proof-edited using FilamentEditor (Dercksen, Hege, & Oberlaender, 2014), a custom-designed program based on Amira visualization software (FEI-Visualization Sciences Group). The NeuroMorph and FilamentEditor tools enable tracings that are independent of the experience of the human operator, with an interobserver variability of \sim 20 μ m per 1 mm axonal length (Dercksen et al., 2014).

2.6 Visualization and analysis of reconstructions

Light micrographs were captured using the microscope described in the previous section or a Zeiss AxioObserver Inverted SK-2. All visual renderings of reconstructed dendrites and axons were generated in Amira. Polar plots were generated by computing the total axonal length in 20° angular bins centered on the soma of each neuron reconstruction.

To generate an average HVC reference frame, we first manually traced HVC contours of all animals used in this study using microscope images (AX70, 10 \times objective UPIanFI) of the fluorescence of retrogradely labeled HVC projection neurons in all sagittal sections containing HVC. Contours were aligned across sections by spacing them 100 µm apart along the mediolateral axis and aligning by translations and rotations in the sagittal plane using traced neuron morphology, blood vessels, and the brain surface as references. The aligned HVC contours were then used to compute a closed surface by 3D Delaunay triangulation and defined as the HVC boundary surface. Next, we computed the center of mass and the three principal axes of the HVC boundary surface of each animal and aligned the boundary surfaces of all animals using these coordinates and orientations. The extent of the HVC boundary surface along the three principal axes was defined as follows: for the first and second principal axes (i.e., the long axis, corresponding approximately to the mediolateral axis, and the medium axis, corresponding approximately to the anterior-posterior axis), the boundary surface was intersected with a plane centered on the center of mass and the plane normal defined by the direction of the third principal axis (i.e., the shortest axis, corresponding approximately to the dorsal-ventral axis). The maximum extent of the resulting intersection contour was then computed along the first and second principal axis, respectively. For the third principal axis, the boundary surface was intersected with a plane defined by the center of mass and the first principal axis. Finally, each neuron morphology was translated and rotated according to the center of mass and principal axes of the corresponding HVC boundary surface in the same animal, and variability in the extent of HVC along the three axes was minimized by scaling the neuron morphology linearly along each dimension according to the difference of the individual HVC boundary surface and the average extent along each axis. Finally, we transformed all aligned neuron morphologies to a coordinate system aligned with the anterior-posterior and mediolateral axes of the bird brain (Guest, Seetharama, Wendel, Strick, & Oberlaender, 2017). We measured the average angle between the first

principal axis and the sagittal plane, as defined by the brain sections, and rotated all aligned neuron morphologies accordingly (i.e., by 28.5°). For visualization purposes, we computed an average HVC boundary surface. We sampled all aligned HVC boundary surfaces every 50 μ m along the average extent of the first principal axis, and in 10° steps, computed the average location of these samples, and generated a

2.7 | Quantification of spatial parameters of dendrites and axons

closed boundary surface by 3D Delaunay triangulation.

For the following analyses, all axonal processes outside the boundaries of HVC were excluded. Total dendritic and axonal lengths were measured using the FilamentEditor in Amira. "Primary branches" (Table 1) refer to the number of dendritic branches that emanate directly from the cell body. Branch nodes are bifurcation points along the dendrite or axon and were quantified using the FilamentEditor in Amira. Onedimensional spatial extent of dendrites and axons was quantified in Amira, first by aligning the soma to the origin, then computing length in 10 or 50 µm voxels for dendrites and axons, respectively, and plotting the distribution along the three principal axes. The radial dendritic extent was defined as the length of all dendrite segments in $1-\mu m$ thick concentric shells centered on the soma. HVC innervation proportion was computed for each cell by first replacing the line segments describing its axonal arbor by 100 µm radius tubes (i.e., reflecting the minimal average radius of potential postsynaptic projection neurons) (Table 1), yielding a closed surface (tube surface). The volume of the tube surface within the boundaries of HVC was measured and was divided by the total volume of HVC in that animal. This yielded a number between 0 and 1 (HVC innervation proportion), which roughly reflects the fraction of HVC projection neurons whose dendritic fields are intersected by the axon. Differences in morphological properties between HVC(RA) and HVC(X) neurons were assessed using the Wilcoxon rank-sum test.

2.8 Clustering of neurons based on dendritic and axonal projection patterns

To determine whether HVC projection neurons form morphologically defined clusters, we extracted parameters describing the 3D projection patterns of their dendrites and axons within HVC. For clustering of dendrites, we used four parameters: the 90% dendritic extent along the mediolateral and anterior-posterior axes within HVC, the 90% radial dendritic extent, and the total dendritic length. For clustering of axons, we used four parameters: the 90% axonal extent along the mediolateral and anterior-posterior axes within HVC, the HVC innervation proportion, and the total axonal length beyond 200 μ m distance from the soma. Computing the 90% extent as opposed to the full extent made this analysis less sensitive to sparse outlier processes.

Clustering of all neurons using dendrite and axon parameters occurred separately. We used the OPTICS algorithm (Ankerst, Breunig, Kriegel, & Sander, 1999; Oberlaender et al., 2012) to sort all neurons based on their mutual proximity in 4D feature space. This allowed identifying groups of neurons forming regions of high density in feature



TABLE 1 Features of HVC projection neuron dendrites. Statistical differences were assessed using the Wilcoxon rank-sum test

ID	Proj. target	90% M-L extent (μm)	90% A-P extent (μm)	90% D-V extent (μm)	90% radial extent (μm)	Path length (mm)	Primary branches	Branch nodes	Dendrite cluster
35	RA	60	90	90	68	2.78	5	35	Small
38	RA	120	100	110	88	4.27	7	53	Small
43	RA	110	70	110	82	3.15	5	55	Small
45	RA	140	130	110	102	4.00	6	44	Small
48	RA	130	100	90	85	4.53	7	52	Small
54	RA	120	120	120	98	4.84	5	58	Small
57	RA	80	100	90	80	2.88	4	39	Small
63	RA	110	90	130	98	2.72	3	37	Small
66	RA	80	110	120	84	2.72	6	30	Small
67a	RA	110	110	70	84	4.64	5	70	Small
67p	RA	110	80	70	78	2.68	6	32	Small
78	RA	80	70	100	70	1.56	8	20	Small
80	RA	60	70	100	68	2.31	5	37	Small
132p	RA	120	180	110	122	4.86	4	40	Large
135a	RA	70	100	100	77	2.32	4	29	Small
135p	RA	80	120	110	87	2.45	4	38	Small
136a	RA	90	120	100	89	3.15	6	42	Small
136p	RA	110	100	50	72	2.18	5	38	Small
140a	RA	100	100	110	88	3.07	6	40	Small
140p	RA	100	110	120	92	4.30	3	53	Small
142a	RA	90	80	60	66	1.35	4	20	Small
142p	RA	110	100	70	81	4.27	8	69	Small
$AVG\pm \mathit{SD}$		99 ± 22	102 ± 24	97 ± 21	85 ± 13	$\textbf{3.23} \pm \textbf{1.06}$	5.3 ± 1.4	42 ± 13	
22	х	160	190	160	134	4.99	4	47	Large
81	х	110	190	180	137	5.61	12	23	Large
91	х	120	220	120	130	7.09	5	64	Large
97	х	150	240	160	152	4.95	6	47	Large
103	х	130	190	180	129	7.12	10	67	Large
106	Х	140	210	110	131	6.04	7	56	Large
110	х	140	150	160	123	5.59	6	49	Large
127a	Х	160	210	170	144	6.34	4	52	Large
127p	х	160	160	150	135	4.36	7	30	Large
128a	х	140	220	150	140	5.80	5	60	Large
128p	х	160	200	130	133	4.52	4	60	Large
129a	х	160	260	140	153	5.54	6	49	Large
129p	Х	180	210	100	131	3.86	4	37	Large
130a	Х	170	220	160	150	7.23	11	59	Large
130p	Х	170	180	160	130	7.88	4	83	Large
132a	Х	190	220	150	147	8.15	10	87	Large

(Continues)

TABLE 1 (Continued)

ID	Proj. target	90% M-L extent (μm)	90% A-P extent (µm)	90% D-V extent (μm)	90% radial extent (μm)	Path length (mm)	Primary branches	Branch nodes	Dendrite cluster
$AVG\pmSD$		153 ± 21	204 ± 28	149 ± 23	137 ± 9	$\textbf{5.94} \pm \textbf{1.27}$	$\textbf{6.6} \pm \textbf{2.7}$	54 ± 17	
RA vs. X p-value		$1.7 imes 10^{-6}$	$2.7 imes 10^{-7}$	4.9×10^{-6}	$2.1 imes 10^{-7}$	$2.4 imes 10^{-6}$	0.25	0.03	
28	RA/X	60	100	90	71	2.01	4	25	Small
75	RA/X	170	150	180	142	2.83	7	28	Large
Small AVG \pm SD		96 ± 23	99 ± 17	96 ± 21	82 ± 10	$\textbf{3.10} \pm \textbf{1.03}$	5.3 ± 1.4	42 ± 14	
Large AVG \pm SD		152 ± 22	200 ± 30	148 ± 25	137 ± 9	5.71 ± 1.42	6.4 ± 2.6	52 ± 17	
Small vs. Large p-value		$7.0 imes 10^{-7}$	$6.7 imes10^{-8}$	$3.1 imes 10^{-6}$	$7.8 imes10^{-8}$	$2.4 imes 10^{-6}$	0.27	0.06	

space, which we identified as putative clusters. Neurons with lowmutual proximity to neurons in these putative clusters were labeled as "ambiguous." We decided to assign these "ambiguous" neurons to one of the two putative clusters based on their distance to the center of the two clusters, while accounting for potentially different variability within each cluster (Narayanan et al., 2015; Rojas-Piloni et al., 2017). Specifically, we defined the distance of neuron *i* to cluster *k* as:

$$d_{i,k} = \sqrt{\left(\vec{f}_{i} - \vec{\mu_{k}}\right)^{\mathsf{T}} \mathbf{C}_{k}^{-1} \left(\vec{f}_{i} - \vec{\mu_{k}}\right)}$$

Here, $\vec{f_i}$ is the 4D feature vector of neuron *i*, $\vec{\mu_k}$ is the mean feature vector of cluster *k*, and C_k^{-1} is the inverse of the covariance matrix of cluster *k*. Hence, because we identified two putative clusters for each parameter set using the OPTICS algorithm, two distance values (i.e., one for each cluster) were computed for each "ambiguous" neuron. We then assigned each "ambiguous" neuron to the cluster with the smallest distance without requiring a predetermined cutoff distance.

3 | RESULTS

To characterize the properties of local processes of HVC projection neurons, we conducted single-cell Neurobiotin fills (Narayanan et al., 2014; Pinault, 1996) in vivo under two-photon guidance (Figure 1a-c; see Materials and Methods). In a total of 81 birds, we attempted to fill 59 HVC(RA) neurons and 34 HVC(X) neurons. Following histological processing and imaging (Figure 1d), we selected only the neurons whose dendrites and axons were completely and darkly labeled, a data set acquired from 31 birds and including 22 HVC(RA) neurons, 16 HVC_(X) neurons, and 2 neurons projecting to both regions (Kittelberger, 2002). In this data set, we include 13 $\text{HVC}_{(\text{RA})}$ neurons and 2 $\text{HVC}_{(\text{RA}/\text{X})}$ neurons that were previously analyzed as part of a separate anatomical study (Kornfeld et al., 2017). As a population, the somata of the targeted cells spanned most of the mediolateral and rostrocaudal extent of HVC (Figure 1e). In contrast, their dorsal-ventral distribution was biased toward superficial neurons, due to limitations on imaging depth in our tissue (Figure 1f). For each neuron, we fully reconstructed all local processes including dendrites and axon collaterals (Figure 2). Long-range projections leading to RA and/or Area X were clearly visible

in these neurons, enabling their identification. A single axon branch leading from HVC to RA often exited the posterior boundary of HVC (Figure 2a,e), while an axon branch from an $HVC_{(X)}$ neuron (Figure 2b,f) took a more variable path, exiting from the anterior, medial, or ventral



FIGURE 1 Two-photon guided single-cell filling. (a) *In vivo* imaging preparation for targeted cell fills. $HVC_{(RA)}$ or $HVC_{(X)}$ neurons are targeted under visual guidance through a cover glass implanted over HVC. (b) In a juxtacellular configuration, positive current pulses are used to fill neurons with Neurobiotin, often eliciting spikes. (c) Example image of a cell being filled with Alexa 488 and Neurobiotin *in vivo*. (d) Brightfield micrograph of the filled neuron in (c) after histological processing. Scale bars: 10 μ m. (e) Relative locations of 40 reconstructed somata mapped onto an average HVC reference frame such that the long axis of HVC is defined along the horizontal. (f) An orthogonal view perpendicular to the long axis of HVC



FIGURE 2 Anatomical reconstruction of HVC projection neurons. (a, b) Micrographs show a projection of 100 μ m sagittal sections of a DAB-stained HVC_(RA) neuron (a) and HVC_(X) neuron (b). Scale bars: 100 μ m. (c, d) Complete reconstructions of the above neurons shown in the sagittal plane. Scale of reconstructions is matched with micrographs in (a, b). (e, f) Same cells as in (c) and (d), shown in the horizontal plane. Shaded region marks the area shown in the micrographs (a, b). Dendrites are represented with black lines, and axons are either red (HVC_(RA)) or blue (HVC_(X)). Dotted lines mark HVC boundaries in (a, b, e, f). Arrows indicate descending axons

boundary (Fortune & Margoliash, 1995). Additionally, three cells in our data set (one $HVC_{(RA)}$ and two $HVC_{(X)}$ neurons) sent two descending axon branches originating from the same hillock to the downstream nucleus.

3.1 | HVC_(RA) and HVC_(X) neurons have distinct dendritic morphologies

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Our approach enables us to visualize and reconstruct all neuronal processes of individual cells throughout the entire volume of HVC, thus avoiding any possible bias associated with analyzing partial reconstructions (Dutar et al., 1998; Fortune & Margoliash, 1995; Mooney, 2000). Our first step was to examine the dendrites of projection neurons in order to better understand the spatial extent of synaptic inputs that could arrive onto each cell type. Dendrites were similar in shape within a cell type, with dendritic arbors consistently more compact in HVC_(RA) neurons than in HVC_(X) neurons (Figure 3). We went on to quantify and compare the morphological properties of HVC_(RA) and HVC_(X) dendrites in several ways. First, to measure the span of dendrites, we quantified and plotted the 1D spatial extent of each arbor along the mediolateral, rostrocaudal, and dorsoventral axes (Table 1 and Figure 4a,b). We found that averaging across the population, HVC_(X) dendrites have a 1.5-fold greater span than HVC_(RA) dendrites in the mediolateral axes, and a 2-fold greater span in the rostrocaudal axis. Across the population, HVC_(RA) dendrites have a similar span in all three axes, lending to a spherical shape, while HVC_(X) dendrites are



FIGURE 3 Dendrites of HVC projection neurons. Each panel shows the dendrite of an $HVC_{(RA)}$ (red), $HVC_{(RA/X)}$ (green) or $HVC_{(X)}$ (blue) neuron projected on the horizontal plane. All $HVC_{(RA)}$ neurons are designated as "small" with the exception of 132p, and all $HVC_{(X)}$ neurons were designated as "large" (see Figure 4 and Materials and Methods). One $HVC_{(RA/X)}$ neuron each fell into the "small" and "large" categories

slightly more elongated in the rostrocaudal axis (Table 1, Figure 4). We also measured total dendritic length of each cell and found that HVC_(X) dendrites on average have almost a 2-fold greater length than HVC_(RA) dendrites (Table 1, Figure 4c). Averaging across the population, HVC_(X) dendrites gave rise to an average of one more primary branches than HVC_(RA) dendrites. The complexity of the dendritic arbor, as measured by the correlation between total path length and number of branch nodes, was similar across all neurons (r = .78, $p < 10^{-6}$).

Our next step was to determine whether we could morphologically classify dendrites based on their spatial properties. We defined four parameters (total dendritic length, 90% extent along the mediolateral and anterior-posterior axis, and 90% radial extent, Figure 4a-c) and used a density-based cluster algorithm (Ankerst et al., 1999) to determine whether cells could be assigned to discrete groups (Oberlaender et al., 2012). Our analysis yielded two morphological categories, which appeared as distinct clusters in a principal component analysis (Figure 4d). One cluster consisted primarily of neurons with large dendritic fields, as measured by their radial extent, the extent across the three principal axes of HVC, and total path length (Table 1), and we hence referred to neurons in this cluster as "large." The "large" cluster contained all HVC_(X) neurons (see also Figure 3). HVC_(RA) neurons—with one exception—were part of the remaining cluster, which we termed "small," due to their compact dendritic span (Figure 3 and Table 1). In addition, one HVC_(RA/X) neuron was assigned to each cluster.



FIGURE 4 Spatial properties of $HVC_{(RA)}$ and $HVC_{(X)}$ dendrites. (a, b) One dimensional profile of dendritic length along three axes for $HVC_{(RA)}$ (a) and $HVC_{(X)}$ (b) neurons. Red and blue traces are from example neurons, grey traces are from all remaining individual neurons from each group in our data set. The center of each plot (0 µm) denotes soma location. (c) Total dendritic path length organized by projection target. Filled circles represent examples from (a) and (b). (d) The first two principal components of four spatial parameters of dendrites. The two clusters as designated using the OPTICS algorithm (see Materials and Methods) are represented by closed circles and open circles; red $-HVC_{(RA)}$; blue $-HVC_{(X)}$; green $-HVC_{(RA/X)}$

3.2 Bouton morphology of local projections in HVC

After examining the dendritic structure of HVC projection neurons, we shifted our focus to potential presynaptic partners by examining local axonal boutons. Our first step was to assess the distribution of boutons along these wires, which correlate with the presence of presynaptic terminals (Kornfeld et al., 2017). Local collaterals of both HVC(RA) and HVC_(X) neurons were lined with prominent swellings, which were readily visible along DAB-stained collaterals (Figure 5a). We manually counted boutons along collateral branches of 7 HVC(RA) and 9 HVC(X) neurons and computed the bouton density of each branch (number of boutons/length of axon; Figure 5b). Across all branches, $\mathsf{HVC}_{(\!\mathsf{RA}\!)}$ and $\mathsf{HVC}_{(\!\mathsf{X}\!)}$ neurons had similar bouton densities (HVC_{(RA)}: 32 \pm 2 boutons/mm; HVC_{(X)}: 36 \pm 2 boutons/mm, mean \pm SEM; Figure 5c), suggesting that they form connections with other neurons in HVC at similar frequencies. We previously reported that the density of boutons did not vary with soma distance in HVC(RA) neurons (Kornfeld et al., 2017) and found the same to be true for HVC_(X) neurons (Pearson's correlation r = -.20; p = .11). However, boutons from both projection types appeared to sometimes be clustered (Figure 5a), which was further evident from nonuniform spatial distributions of boutons within an axon (Figure 5d).

3.3 | Two categories of local axonal innervation

Once we confirmed that boutons are prevalent along collaterals, we moved on to examine and compare the structure of collaterals of

 $\mathsf{HVC}_{(\mathsf{RA})}$ and $\mathsf{HVC}_{(X)}$ neurons. In nearly all cells, collaterals were completely confined within the borders of HVC, even in cases where the soma was close to the edge of the nucleus (e.g., Figure 6, Cell 38). In two HVC_(X) neurons, however, collateral branches extended outside of HVC and terminated ${\sim}500~\mu m$ rostral and caudal to the boundary of the nucleus, into HVC shelf (Kelley & Nottebohm, 1979). Cell 22 sent 7.2 mm of collaterals into rostral and caudal shelf, and Cell 103 sent 4.5 mm of collaterals into caudal shelf (Figures 7 and 9). A general feature of both projection types is that their local collaterals appeared to fill the dorsoventral extent of the nucleus. In the horizontal plane, however, an axon forming several collaterals emanated from the soma at distinctive orientations along the mediolateral and rostrocaudal axes (Figures 6-8). Across the population, we did not observe a clear bias toward a particular orientation relative to the soma. As with the dendrites, we observed almost a 2-fold greater total axon length of HVC_(X) neurons compared with HVC(RA) neurons, and the span of HVC(X) collaterals was larger along all measured dimensions (M-L, D-V, A-P) (Table 2).

Our next step was to determine if, like the dendrites, axon collaterals of HVC projection neurons could be organized into distinct morphological categories. We again focused specifically on parameters that describe the spatial properties of these projections and their innervation of HVC. For each cell, we quantified and plotted the 1D spatial extent of the axon along the three principal axes (Table 2 and Figure 10a,b). We chose to exclude dorsoventral extent from our sorting analysis, as this parameter was relatively consistent from cell to cell (Table 2).





FIGURE 5 Bouton density of HVC_(RA) and HVC_(X) collaterals. (a) Example micrographs of HVC_(RA) (top) and HVC_(X) (bottom) axon collaterals. Scale bar: 5 µm. Arrowheads mark the location of boutons. (b) Bouton densities from 170 branches in 7 HVC_(RA) neurons and 95 branches in 9 HVC_(X) neurons. Mean density for each cell is shown as red and blue horizontal line, respectively. (c, d) Histograms showing bouton densities for all branches (c) as well as inter-bouton distance (d) for each cell type from (b)

We quantified distal axonal length (>200 µm Euclidean distance from soma), previously shown to preferentially target other projection neurons (Kornfeld et al., 2017) as a measure of broad axonal innervation, and used this as our third parameter (Figure 10c, left). Finally, for our fourth parameter, we computed an HVC innervation proportion for each cell (see Materials and Methods), which reflects the upper bound proportion of neurons in HVC whose dendritic fields overlap with the axonal field (Figure 10c, right). The cluster algorithm yielded two clusters distinguished by the span of their axonal fields, leading to morphological categories that we refer to as "local" and "broadcast" neurons (Figure 10d and Table 2). "Broadcast" neurons had a higher extent along all principal axes than "local" neurons (M-L: 1.6-fold higher; A-P: 1.5-fold higher; D-V: 1.3-fold higher; p < .005 in all cases), 1.9-fold more path length within HVC ($p = 1.0 \times 10^{-4}$), and innervated a larger fraction of HVC (1.4-fold higher

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HVC innervation ratio, p = .01). In contrast to the dendrites, we found that "local" and "broadcast" neurons were not as strictly associated with projection type: the "local" group consisted of 21 HVC_(RA), 6 HVC_(X), and 1 HVC_(RA/X), whereas the "broadcast" group consisted of 1 HVC_(RA), 10 HVC_(X), and 1 HVC_(RA/X) (Figures 6–8). Therefore, 6 out of 16 HVC_(X) neurons have similar axon morphologies to HVC_(RA) neurons in terms of their spatial distribution within HVC, whereas 10 HVC_(X) neurons exhibit distinctly broader projections that innervate a larger volume of HVC. To visualize and compare how "local" and "broadcast" neurons in each category onto an average geometrical reference frame (see Materials and Methods; Figure 11a,b). "Broadcast" axons occupy a relatively large volume and span most of the mediolateral and rostrocaudal extent of HVC, whereas the "local" axons are more spatially confined and occupy a smaller volume in the nucleus (Figure 11c,d, Table 2).

4 | DISCUSSION

We used *in vivo* two-photon guided single cell labeling to acquire a data set of 40 HVC projection neurons to examine the projection patterns of their dendrites and local axon collaterals in three dimensions. This data set contains the first complete reconstructions of local processes across the entire volume of HVC. By characterizing these morphological properties in detail, we have gained insight into the impact of individual neurons on the circuits involved in the acquisition and production of a well-characterized complex sensorimotor behavior.

4.1 | Two-photon targeted cell labeling

Our approach of using in vivo microscopy to assist cell targeting/filling enabled several experimental advantages. First, we were able to choose the cells we targeted based on their projection type by injecting a fluorescent retrograde tracer in either RA or Area X. Furthermore, by observing the field of retrogradely labeled neurons, we were able to target cells in specific regions of HVC, which made it easier to generate a data set that spanned the mediolateral and rostrocaudal extent of the nucleus. Second, the visualized approach helped to direct the tip of the electrode during the process of cell filling, which was especially useful when targeting $HVC_{(RA)}$ neurons, which have relatively small somata (7-10 µm diameter). Third, we were able to monitor the flow of internal solution from the pipette tip throughout the experiment, ensuring that it was not clogged before attempting to inject Neurobiotin. One limitation to this approach is that our fills are biased toward neurons in the dorsal half of HVC, as dextran-labeled neurons were only visible up to ${\sim}400~\mu m$ ventral from the surface, and the ventral-most boundary of HVC extends ${\sim}550~\mu\text{m}$ ventral. It is possible that neurons closer to the ventral boundary send collaterals dorsally, mirroring what we observe from cells in our data set, however, we cannot rule out the possibility that the collaterals of ventral neurons exhibit different projection patterns altogether.



FIGURE 6 Local processes of $HVC_{(RA)}$ neurons. Each panel shows the dendrite (black) and local axons (red) of all $HVC_{(RA)}$ neurons projected on the horizontal plane and its position within HVC with accompanying polar plots quantifying axonal distribution. Arrows mark descending axons to RA. "Local" (L) and "broadcast" (B) designations given in parentheses (see Figure 11 and Materials and Methods)

4.2 | HVC_(RA) neurons: Functional implications for sequence generation

The primary function of $HVC_{(RA)}$ neurons is to provide premotor commands to the song production pathway (Nottebohm et al., 1976). During singing, the majority of $HVC_{(RA)}$ neurons are sparsely active at moments that are highly stereotyped across song renditions (Hahnloser et al., 2002). Different $HVC_{(RA)}$ neurons are often active at different time points, forming a sparse sequence that spans the entirety of the song (Hahnloser et al., 2002; Kozhevnikov & Fee, 2007; Lynch et al., 2016; Picardo et al., 2016). Several models have been proposed to explain how HVC_(RA) premotor sequences are generated during singing. In one class of models, precisely timed bursts are the direct result of connectivity between HVC_(RA) neurons, which have been proposed to form a feed-forward network (Long et al., 2010). An alternative model suggests that sequential activity may be generated by a distributed network involving structures outside HVC, where precisely timed bursts during song are driven by thalamic afferents onto HVC_(RA) neurons rather than local connections within the nucleus (Hamaguchi, Tanaka, & Mooney, 2016).



FIGURE 7 Local processes of $HVC_{(X)}$ neurons. Each panel shows the dendrite (black) and local axons (blue) of all $HVC_{(X)}$ neurons projected on the horizontal plane and its position within HVC with accompanying polar plots quantifying axonal distribution. Arrows mark descending axon to Area X. "Local" (L) and "broadcast" (B) designations given in parentheses (see Figure 11 and Materials and Methods)

Our current findings provide an anatomical basis for local excitatory connectivity that may play an important functional role within HVC. We show that nearly all 22 HVC_(RA) cells have elaborate axon collaterals that exclusively innervate HVC and likely make many connections, as supported by the prevalence of putative presynaptic terminals. We were able to estimate the total boutons per HVC_(RA) cell by multiplying the axonal length and bouton density. We find that, on average, an HVC_(RA) neuron forms 401 ± 52 boutons. Given that there are ~40,000 HVC_(RA) neurons (Walton, Pariser, & Nottebohm, 2012; Wang, Hurley, Pytte, & Kirn, 2002), we estimate that the entire population of HVC_(RA) neurons forms ~16 million local boutons, suggesting a significant impact on network activity in HVC during song production.

To better understand the role of intrinsic HVC circuitry, we must first establish the postsynaptic partners of the $HVC_{(RA)}$ neurons, which cannot be determined through the light microscopic approach used in this study. Paired recordings *in vitro* have found sparse evidence for $HVC_{(RA)}$ - $HVC_{(RA)}$ connections (Kosche et al., 2015; Mooney & Prather, 2005), motivating a number of alternative models in which disynaptic inhibition within HVC is sufficient to generate song-related sequences (Armstrong & Abarbanel, 2016; Gibb et al., 2009). In a previous study, using an electron microscopic approach capable of revealing the identity of postsynaptic partners of $HVC_{(RA)}$ neurons, we found that 17% of



FIGURE 8 Local processes of HVC_(RA/X) neurons. Each panel shows the dendrite (black) and local axons (green) of both HVC_(RA/X) neurons projected on the horizontal plane and its position within HVC with accompanying polar plots quantifying axonal distribution. Arrows mark descending axons to RA and Area X. "Local" (L) and "broadcast" (B) designations given in parentheses (see Figure 11 and Materials and Methods)



FIGURE 9 HVC_(X) dendrites and collaterals can extend into HVC shelf. (a) Fluorescent micrograph of a sagittal section of Cell 103. Scale bar: 100 μ m. (b) Brightfield micrograph of the area indicated with a white rectangle in (a). Inset shows a high magnification image from the boxed region. Scale bars: 25 μ m and 5 μ m (inset). (c) Reconstructed dendrite (black) and axon collaterals (blue) from the boxed region in (a). Scale bar: 25 μ m

synapses made by HVC_(RA) axons form connections onto other HVC_(RA) neurons (Kornfeld et al., 2017) and that the total density of these connections is ~75 synapses/mm (Kornfeld et al., 2017). With this data set, we update our previous estimate (Kornfeld et al., 2017) for the average axonal length of HVC_(RA) collaterals $(13.31 \pm 1.44 \text{ mm mean} \pm SEM)$ by including data from an additional nine neurons. We now estimate that each individual HVC_(RA) neuron forms 170 ± 18 synapses onto other HVC_(RA) neurons, which suggests that the synaptic connections exist to allow for the possibility that this feed-forward circuit architecture could enable the generation of song-related sequences within HVC. The total number of synaptic partners remains elusive, because little is known about the relative number of connections made between a given pair of HVC_(RA) neurons (Kornfeld et al., 2017).

Additionally, in our previous study, we observed that $HVC_{(RA)}$ $HVC_{(RA)}$ connections often occurred in the portions of the axons that were distant from the soma. This regional specialization, coupled with the relatively compact nature of $HVC_{(RA)}$ axonal field, places specific spatial bounds on the location of $HVC_{(RA)}$ postsynaptic partners—proximal axons are not likely to form robust $HVC_{(RA)}$ - $HVC_{(RA)}$ synapses and axons often do not reach across the entire span of the nucleus. Furthermore, we find that collaterals of both $HVC_{(RA)}$ and $HVC_{(X)}$ neurons are heterogeneously biased in orientation with respect to their soma, suggesting that excitatory signals during singing propagate without a bias toward a particular axis in HVC.

Our results are also relevant in light of previous data suggesting that HVC is composed of multiple processing centers with distinct behavioral roles. In these studies, microlesions to specific subregions of HVC led to selective deficits in song structure (Basista et al., 2014; Galvis et al., 2017). Despite these compelling findings, we were unable to see differences between the anatomical properties of individual premotor neurons across the extent of HVC. As a result, the differences observed following perturbation may result from other factors, such as the specific downstream targets of these neurons within RA, for instance. Additionally, functional observations (Stauffer et al., 2012) as well as perturbation studies (Poole, Markowitz, & Gardner, 2012; Stauffer et al., 2012) have suggested an anterior–posterior bias in the wiring of HVC. In our data set, axonal collaterals sometimes send significant projections in the mediolateral direction (e.g., Figure 6), indicating the existence of long-range connectivity across the long axis of HVC. Future studies using more precise methods to monitor and perturb these long-range axonal collaterals may hold the key to understanding their role.

4.3 | HVC_(X) neurons: Local processing of basal-ganglia projecting neurons within HVC

HVC_(X) neurons project to the avian basal ganglia and are primarily involved in delivering an efference copy of motor activity to the anterior forebrain pathway. Although HVC_(X) neurons are essential for song learning (Scharff & Nottebohm, 1991; Sohrabji et al., 1990), their role in adult song production is less clear (Scharff et al., 2000). Our study suggests four potentially new roles for the HVC(X) network. First, the possibility exists that HVC_(X) neurons may have an impact on vocal production in adult zebra finches, given the fact that HVC_(X) neurons have such an extensive network of axon collaterals in HVC. Considering the average total axon collateral length and bouton density of $HVC_{(X)}$ neurons, we estimate that on average, an HVC_(X) neuron forms 755 \pm 91 boutons. There are \sim 10,000 HVC_(X) neurons in the nucleus (Walton et al., 2012), so the population of $HVC_{(X)}$ neurons may form ~7.5 million boutons in HVC. Given that HVC_(X) neurons display activity patterns similar to HVC_(RA) neurons during song production (Hahnloser et al., 2002; Long et al., 2010; Vallentin & Long, 2015) and may form many connections with HVC(RA) neurons, despite a paucity of such interconnections having been observed in earlier studies (Mooney & Prather, 2005), future work is needed to determine whether $HVC_{(X)}$ neurons play a role in the premotor patterns leading to the generation of singing behavior. Second, we find that $HVC_{(X)}$ neurons can exist either as "local" or "broadcast" neurons. Future functional measurements of these neurons during singing combined with anatomical reconstructions could potentially shed light on the roles of different morphological classes in song production. Third, we found a subpopulation of $HVC_{(X)}$ neurons that innervate a region adjacent to HVC, known as HVC shelf (Kelley & Nottebohm, 1979). HVC shelf is an auditory region that receives input from field L (Fortune & Margoliash, 1995; C. V. Mello, Vates, Okuhata, & Nottebohm, 1998; Vates et al., 1996), which is the primary forebrain relay for pathways originating in the auditory thalamus (Bonke, Bonke, & Scheich, 1979; Fortune & Margoliash, 1992; Karten, 1968). Therefore, the HVC(X)-to-shelf synapse could be transmitting motor-related signals to the auditory system, similar to a recently characterized population of HVC neurons projecting to Avalanche (Roberts et al., 2017). Fourth, two HVC neurons send longrange projections to both X and RA. Such dual projectors have been



TABLE 2 Features of HVC projection neuron axons

ID	Proj. target	90% M-L extent (µm)	90% A-P extent (µm)	90% D-V extent (µm)	Path length (mm)	Path length beyond 200 μm (mm)	Branch nodes beyond 25 μm	HVC innervation	Axon cluster
35	RA	800	250	350	6.77	2.74	24	0.23	L
38	RA	650	450	350	20.00	11.94	62	0.51	L
43	RA	600	450	150	7.35	2.99	22	0.22	L
45	RA	650	550	350	15.56	9.21	42	0.51	L
48	RA	600	500	200	16.66	7.81	73	0.57	L
54	RA	350	200	250	5.89	1.06	18	0.22	L
57	RA	850	500	300	10.83	5.72	27	0.49	L
63	RA	750	450	300	9.88	4.93	30	0.46	L
66	RA	700	350	300	12.42	5.73	38	0.46	L
67a	RA	750	500	400	24.36	19.17	44	0.47	L
67p	RA	900	700	350	22.24	16.28	37	0.50	В
78	RA	1000	350	300	5.97	3.21	13	0.40	L
80	RA	700	400	400	9.75	3.38	30	0.58	L
132p	RA	650	700	400	29.17	21.99	57	0.54	L
135a	RA	650	350	550	10.47	6.37	25	0.35	L
135p	RA	850	500	350	13.40	8.57	31	0.54	L
136a	RA	550	500	250	13.79	6.75	41	0.48	L
136p	RA	800	450	400	6.58	2.77	24	0.31	L
140a	RA	150	200	100	3.31	0.17	32	0.09	L
140p	RA	500	450	350	13.39	7.42	47	0.34	L
142a	RA	650	350	250	7.19	3.75	25	0.44	L
142p	RA	600	350	200	10.68	5.81	29	0.53	L
$AVG\pmSD$		668 ± 184	432 ± 131	$\textbf{311} \pm \textbf{99}$	12.53 ± 6.61	$\textbf{7.17} \pm \textbf{5.66}$	35 ± 15	$\textbf{0.42}\pm\textbf{0.13}$	
22	Х	550	550	450	21.06	17.14	53	0.54	L
81	х	650	650	300	16.09	9.02	34	0.40	L
91	Х	650	600	350	16.30	11.80	21	0.49	L
97	Х	1150	600	400	22.35	18.89	39	0.66	В
103	Х	1050	500	400	21.24	17.64	31	0.79	В
106	х	850	750	500	16.10	12.53	29	0.37	В
110	Х	400	450	300	6.64	2.55	18	0.24	L
127a	Х	650	550	250	15.39	11.09	38	0.41	L
127p	Х	400	250	150	4.59	2.26	7	0.15	L
128a	Х	950	650	400	27.06	21.23	51	0.49	В
128p	Х	1200	600	450	30.37	23.20	48	0.58	В
129a	Х	1100	800	450	26.69	20.39	47	0.47	В
129p	Х	1150	550	450	38.88	32.81	58	0.68	В
130a	х	1050	700	350	22.80	19.03	32	0.61	В
130p	Х	1100	700	350	32.07	25.60	52	0.65	В
132a	Х	1000	650	350	17.90	14.64	27	0.44	В

(Continues)





TABLE 2 (Continued)

ID	Proj. target	90% M-L extent (µm)	90% A-P extent (µm)	90% D-V extent (μm)	Path length (mm)	Path length beyond 200 μm (mm)	Branch nodes beyond 25 μm	HVC innervation	Axon cluster
$AVG\pmSD$		869 ± 277	597 ± 130	369 ± 89	$\textbf{20.97} \pm \textbf{8.91}$	16.24 ± 8.03	37 ± 14	$\textbf{0.50} \pm \textbf{0.17}$	
RA vs. X p-value		0.03	$4.2 imes 10^{-4}$	0.046	0.003	0.002	0.48	0.15	
28	RA/X	950	600	300	16.07	11.15	39	0.46	В
75	RA/X	550	500	300	27.75	15.52	91	0.59	L
L AVG \pm SD		630 ± 171	441 ± 126	307 ± 97	12.90 ± 6.82	$\textbf{7.53} \pm \textbf{5.58}$	36 ± 19	0.41 ± 0.14	
$B\;AVG\pmSD$		1038 ± 109	650 ± 85	396 ± 58	$\textbf{24.48} \pm \textbf{6.86}$	19.45 ± 5.94	41 ± 10	$\textbf{0.56} \pm \textbf{0.12}$	
B vs. L p-value		$1.5 imes 10^{-6}$	$2.4 imes 10^{-5}$	0.003	$1.0 imes 10^{-4}$	$2.6 imes 10^{-5}$	0.10	0.01	

B: broadcast type, L: local type. Statistical differences were assessed using the Wilcoxon rank-sum test.

previously described in one existing data set (Kittelberger, 2002), but the unique function of such neurons—if any—remains elusive. In our study, we present two such neurons and further find that one neuron appears to have morphological similarities to HVC_(X) neurons and the other appears more similar to an HVC_(RA) neuron (Rojas-Piloni et al., 2017). Whether HVC_(RA/X) neurons are an example of errant wiring or a distinct neuron cell type with a specified functional role within the network remains unknown. In conclusion, we show that there is significant innervation within HVC from both major projection cell types, providing the substrate for local processing that could potentially play a crucial role in vocal learning and production. Furthermore, these connections can help to inform our understanding of the neural mechanisms of sequence generation more broadly (Harvey, Coen, & Tank, 2012; G. B. Mello, Soares, & Paton, 2015; Pastalkova, Itskov, Amarasingham, & Buzsaki, 2008; Schmidt et al., 2017; Schmitt et al., 2017).



FIGURE 10 Spatial properties of $HVC_{(RA)}$ and $HVC_{(X)}$ axons. (a, b) One dimensional profile of axonal length along three axes for $HVC_{(RA)}$ (a) and $HVC_{(X)}$ (b) neurons. Red and blue traces are from example neurons, grey traces are from all remaining individual neurons from each group in our data set. The center of each plot (0 μ m) denotes soma location. (c) Distal axon path length and HVC innervation proportion (see Materials and Methods) organized by projection target. Filled circles represent examples from (a) and (b). (d) The first two principal components of four spatial parameters of axons. The two clusters as designated using the OPTICS algorithm (see Materials and Methods) are represented by closed circles and open circles; red $-HVC_{(RA)}$; blue $-HVC_{(RA)}$; green $-HVC_{(RA)X}$

a 'Local' axons b 'Broadcast' axons C 'Local' axon fields 500 µm

FIGURE 11 Broadcast axons innervate a larger volume than local axons. (a, b) Axons of three local (a) and broadcast (b) neurons registered within the average HVC reference frame (dotted line). (c, d) Schematic of local (c) and broadcast (d) axonal innervation of HVC

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CONFLICT OF INTEREST

The authors have no conflicts of interest.

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