Development/Plasticity/Repair

Dopaminergic Contributions to Vocal Learning

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Although the brain relies on auditory information to calibrate vocal behavior, the neural substrates of vocal learning remain unclear. Here we demonstrate that lesions of the dopaminergic inputs to a basal ganglia nucleus in a songbird species (Bengalese finches, Lonchura striata var. domestica) greatly reduced the magnitude of vocal learning driven by disruptive auditory feedback in a negative reinforcement task. These lesions produced no measureable effects on the quality of vocal performance or the amount of song produced. Our results suggest that dopaminergic inputs to the basal ganglia selectively mediate reinforcement-driven vocal plasticity. In contrast, dopaminergic lesions produced no measurable effects on the birds' ability to restore song acoustics to baseline following the cessation of reinforcement training, suggesting that different forms of vocal plasticity may use different neural mechanisms.

Key words: basal ganglia; Bengalese finch; dopamine; negative reinforcement; songbird; vocal learning

Significance Statement

During skill learning, the brain relies on sensory feedback to improve motor performance. However, the neural basis of sensorimotor learning is poorly understood. Here, we investigate the role of the neurotransmitter dopamine in regulating vocal learning in the Bengalese finch, a songbird with an extremely precise singing behavior that can nevertheless be reshaped dramatically by auditory feedback. Our findings show that reduction of dopamine inputs to a region of the songbird basal ganglia greatly impairs vocal learning but has no detectable effect on vocal performance. These results suggest a specific role for dopamine in regulating vocal plasticity.

Introduction

The brain relies on sensory information to guide the acquisition and maintenance of complex motor skills, including vocal behavior. Neurophysiological studies in mammals have shown that the activity of midbrain dopamine (DA) neurons reflects the difference between predicted and experienced reward, leading to the widespread hypothesis that such signals guide reinforcement learning (Schultz, 2007, 2013). However, establishing DA’s role in sensorimotor learning, including vocal learning, has proved challenging in both experimental and clinical settings. Neurotoxic lesions of DA inputs to the basal ganglia often result in severe motor performance deficits (Zhou and Palmiter, 1995; Beeler et al., 2010), complicating efforts to isolate DA’s specific contributions to motor learning. Moreover, because the mammalian striatum mediates a wide range of behaviors, dopaminergic inputs likely affect cognitive and behavioral processes other than those being assayed experimentally. Additionally, Parkinson’s disease, which includes dysfunction of the dopaminergic system, is associated with vocal performance and plasticity deficits (Ramig et al., 2008; Mollaei et al., 2013). However, because Parkinson’s disease involves pathologies that extend beyond a simple loss of dopaminergic neurons (Lang and Obeso, 2004), it is difficult to use clinical studies to pinpoint DA’s role in vocal behavior.

Songbirds provide a well-defined neural circuit in which to investigate DA’s role in vocal learning. As in a number of forms of mammalian behavioral plasticity (Ogura et al., 2005; Eckart et al., 2010), the basal ganglia are crucial for vocal plasticity in songbirds. Area X (Fig. 1a,b), the song system’s basal ganglia component, is a nucleus in the anterior forebrain pathway (AFP), a basal ganglia-thalamocortical loop long implicated in vocal learning (Sohrabji et al., 1990; Scharff and Nottebohm, 1991; Brainard and Doupe, 2000). Area X is necessary for song learning but not performance. Its destruction abolishes normal song learning in juveniles and degrades the adaptive modification (but not performance) of song in adults (Sohrabji et al., 1990; Scharff and Nottebohm, 1991; Fee and Goldberg, 2011; Ali et al., 2013). How-
we directly microinjected 6-OHDA into area X to reduce dopaminergic inputs to area X (ever, it remains unclear how auditory error signals are conveyed to area X, a basal ganglia nucleus critical for vocal learning. Figure 1. A song-specific basal ganglia nucleus receives strong dopaminergic input. a, The song system includes area X, a basal ganglia nucleus critical for vocal learning. b, A parasagittal section stained for TH shows heavy label within the basal ganglia (blue dotted line) with especially strong label in area X (borders of X indicated by white triangles). TH stain also shows dopaminergic cell bodies in the VTA/SNC (red triangles) and their ascending axons (yellow triangles). c, Experimental design (see Materials and Methods).

Figure 1. A song-specific basal ganglia nucleus receives strong dopaminergic input. a, The song system includes area X, a basal ganglia nucleus critical for vocal learning. b, A parasagittal section stained for TH shows heavy label within the basal ganglia (blue dotted line) with especially strong label in area X (borders of X indicated by white triangles). TH stain also shows dopaminergic cell bodies in the VTA/SNC (red triangles) and their ascending axons (yellow triangles). c, Experimental design (see Materials and Methods).

However, it remains unclear how auditory error signals are conveyed to area X (Peh et al., 2013; Vallentin and Long, 2015). In mammals, dopaminergic inputs to the basal ganglia convey error-related signals, suggesting that dopaminergic afferents to area X might guide vocal learning in songbirds (Doyle and Sejnowski, 1998; Graybiel, 2005; Turner and Desmurget, 2010; Fee and Goldberg, 2011; Leblois, 2013; Colombro, 2014). However, to our knowledge, no studies have directly addressed how DA contributes to vocal learning. We therefore reasoned that selectively lesioning dopaminergic inputs to area X would allow us to isolate DA’s contribution to learning without inducing performance deficits.

We used the neurotoxin 6-hydroxydopamine (6-OHDA) to reduce dopaminergic innervation within area X. In mammalian systems, 6-OHDA injections are commonly used to selectively eliminate dopaminergic fibers and cell bodies (Schober, 2004). The dopaminergic cells innervating area X originate in the VTA and SNCs and are spatially intermingled with dopaminergic neurons that project to other parts of the striatum (Person et al., 2008), precluding injection directly into the VTA/SNCs. Instead, we directly microinjected 6-OHDA into area X to reduce dopaminergic innervation of area X, but not surrounding striatum, and quantified the resulting effects on song performance and vocal learning (Fig. 1c).

Materials and Methods
All subjects were adult (>100-d-old) male Bengalese finches (Lonchura striata var. domestica). All procedures were approved by Emory University’s Institutional Animal Care and Use Committee.

Vocal learning paradigm and behavioral analysis
Adaptive changes in the pitch (fundamental frequency) of targeted syllables were driven using a disruptive auditory stimulus as described previously (Tumer and Brainard, 2007). Briefly, when the pitch of a particular “targeted” syllable was above (or below) a particular threshold, a blast of white noise was played through the speakers, a contingency previously shown to induce birds to lower (or raise) vocal pitch to avoid white noise playback. During reinforcement, a randomly selected subset of target syllables (10%) were selected as “catch trials” during which white noise was not played back, allowing quantification of holistic syllable features, such as sound amplitude. The frequency threshold for white noise was determined using the target syllable’s pitch distribution from songs produced the morning of the first white noise day (>25 song bouts). To drive pitch down (or up), the targeting software was set to trigger a 40–50 ms white noise blast whenever target syllable pitch was above the 10th percentile (or below the 90th percentile) of this distribution.

All behavioral experiments began with a 3 d baseline period in which no white noise playbacks occurred. Postlesion baseline occurred during post-surgery days 4–6 or 5–7. In prelesion experiments, birds were exposed to white noise training for 3 d. In postlesion experiments, training continued for at least 3 d plus up to 3 additional days to ensure that birds had sung at least 90% as many songs as during the prelesion white noise regime, allowing comparison of learning between prelesion white noise day 3 and the approximately trial-matched postlesion white noise day. This extra training means that the white noise day immediately preceding washout day 1 is not necessarily white noise day 3. Although across experiments lesions did not significantly affect the amount of song production on average (see Results), comparing trial-matched prelesion and postlesion days allows us to control for the effect of different amounts of vocal practice on the amount of learning in individual animals. Notably, as described in Results, all findings were qualitatively identical if prelesion and postlesion learning was compared on the same chronological day (white noise day 3) rather than the trial-matched day. Each bird’s postlesion trial-matched day was the day where (by the end of the day) birds had sung the lowest number of songs under the white noise regime as they had after prelesion day 3. Across birds postsham or postlesion, white noise training ended after days 3, 4, 4, 5 (for shams) and 2, 3, 3, 6 (for 6-OHDA), and for each trial-matched day the total number of songs was within 10% of the number produced after prelesion day 3. Daily targeting sensitivity (hit rate) had a median value of 92% across all experiments (range, 58%–99%). Daily targeting precision (1 – false-positive rate) had a median value of 93% across all experiments (range, 51%–100%). Across experiments neither sensitivity nor precision was significantly different between the prelesion and postlesion epochs (Kolmogorov–Smirnov tests, p > 0.25). All singing was undirected (i.e., in the absence of a female bird) throughout the white noise experiments.

After the last white noise day, we withdrew reinforcement and recorded song for 3 additional days to monitor spontaneous pitch restoration back to baseline, which typically occurs after several days (Tumer and Brainard, 2007; Warren et al., 2011). Throughout this paper, we refer to this time period as “washout” and the birds’ process of returning vocal acoustics to their baseline values as “restoration.” Washout occurred between 11 and 18 d after surgery, depending on the bird.

Although all singing was undirected (i.e., no female bird was present) during baseline, white noise training, and washout, we collected female-directed songs from 4 birds (3 pre- and post-6-OHDA lesion, one presham) to assess the effect of 6-OHDA lesion on social context-dependent changes in pitch variability (Sakata et al., 2008). We obtained directed songs 1–4 d after washout was concluded in both prelesion and postlesion conditions. Interleaved directed and undirected songs were...
collected as described previously (Sakata et al., 2008). We obtained 1207 directed (919 undirected) syllable iterations across 11 prelesion syllables and 707 directed (688 undirected) iterations across 8 postlesion syllables and analyzed an equal number of interleaved undirected songs per bird/condition (>30 syllable iterations per syllable and condition).

Custom-written MATLAB software (The MathWorks) was used for data analysis. Pitch changes were quantified in units of semitones as follows:

\[
s = 12 \cdot \log_2(h/b)
\]

where \(s\) is the pitch change (in semitones) of the syllable, \(h\) is the pitch (in Hertz) of the syllable, and \(b\) is the average baseline pitch (in Hertz) of the syllable. On each baseline, white noise day, and washout day, we quantified the pitch, amplitude, and spectral entropy of the targeted syllable in 100 song bouts spaced evenly throughout the day (or all songs when birds sang <100 songs on a given day), as described previously (Sober et al., 2008). To assess lesion-related changes in the quantity of song produced, for each bird we quantified the ratio of the mean number of song bouts per day after 6-OHDA or sham lesion to the mean number produced per day before lesion. We then tested whether the distribution of ratios from 6-OHDA-injected animals differed from both unity and the distribution of ratios from the sham-injected group using one- and two-sample Kolmogorov–Smirnov tests, respectively. These tests allow us to quantify whether neurotoxin injection had a consistent effect on the amount of song production relative to both preinjection behavior and any effects of sham injections.

To assess changes in vocal pitch during the washout period (i.e., after the cessation of white noise training), we fit an exponential decay model to the pitch data as follows:

\[
p(t) = p_{\text{initial}} \cdot e^{-\frac{t}{\tau}} + p_{\text{final}}(1 - e^{-\frac{t}{\tau}})
\]

where \(p_{\text{initial}}\) was set to the mean pitch on the final day of WN training, and fit parameters were \(\tau\) and \(p_{\text{initial}}\). Corresponding to the time constant of pitch restoration during washout and the asymptote of the exponential fit (final value of pitch if restoration were to reach equilibrium), respectively. We note that this exponential model is a generalization of a model we have used previously to quantify the time course of learning when songbirds experience real-time errors in the pitch of auditory feedback delivered via miniature headphones (Sober and Brainard, 2012); in that earlier model, \(p_{\text{initial}}\) was zero, because changes in vocal pitch were quantified relative to baseline error of zero.

**6-OHDA and sham lesions**

Subjects were randomly assigned to either the sham or 6-OHDA lesion group. Before injections, birds were anesthetized with ketamine, midazolam, and isoflurane, mounted in a stereotax at a 20° beak angle relative to the table surface, and small craniotomies were made above area X. Lesioned birds received bilateral injections of 11.8 mg 6-OHDA-HBr/ml (i.e., 8 mg freebase 6-OHDA/ml) and 2 mg ascorbic acid/ml (stabilizer) in a 0.9% NaCl solution into area X using a Drummond Scientific Nanoject II auto-nanoliter injector. During each injection, the pipet was lowered into the brain along a plane perpendicular to the table surface. We injected 13.8 nl at each location and waited 30 s before raising the pipet. For sham lesions, only vehicle (2 mg ascorbic acid/ml in 0.9% NaCl) was injected per the procedure described above. All birds recovered from surgery within a few hours and usually sang the next day.

We varied injection coordinates and volumes slightly between birds and hemispheres to optimize injection parameters. For detailed parameters for each bird, see Tables 1 and 2. Total injection volume in each hemisphere ranged from 124.2 to 179.4 nl for all hemispheres, except one (234.6 nl). Necrotic damage within area X was observed in only one hemisphere of one bird (the right hemisphere of Bird 3, which was the hemisphere receiving the largest total injection volume of 234.6 nl). This necrotic damage affected 8% of the total volume of area X in the affected hemisphere. As our results are unaffected by removing Bird 3 from our dataset, we have included it in our analysis. As described below in Histology, we also performed a number of other analyses to investigate whether 6-OHDA injections killed neurons in area X in cases where no necrosis was apparent.

To cover the greatest possible volume of area X while still injecting low volumes, we placed injections (13.8 nl each) on a 3 × 3, 3 × 4, or 4 × 4 grid. Each grid was located at a single dorsal-ventral (DV) coordinate between 3.1 and 3.4 mm and individual injections were evenly spaced anterior–posterior (AP) and medial-lateral (ML) coordinates between 5.1 and 6.3 mm and 0.9–2.2 mm, respectively. All AP and ML coordinates were relative to the posterior edge of a Y-shaped sinus visible beneath the inner skull layer, whereas DV coordinates were relative to the exposed brain surface. In three birds (and right hemisphere of a fourth), there was an additional injection outside of the main grid intended to hit

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**Table 1. Injection parameters for 6-OHDA lesioned birds**

<table>
<thead>
<tr>
<th>Bird ID</th>
<th>Hemisphere</th>
<th>Concentration (mg/ml)</th>
<th>Injected volume (nl)</th>
<th>AP coordinates (mm)</th>
<th>ML coordinates (mm)</th>
<th>DV coordinates (mm)</th>
<th>Days killed after surgery</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bird 1</td>
<td>Left</td>
<td>8</td>
<td>179.4</td>
<td>1.435</td>
<td>5.1, 5.5, 5.9, 6.3</td>
<td>0.9, 1.55, 2.0</td>
<td>3.18 (2.6 for Med_X) 18</td>
</tr>
<tr>
<td>Bird 1</td>
<td>Right</td>
<td>8</td>
<td>179.4</td>
<td>1.435</td>
<td>5.1, 5.5, 5.9, 6.3</td>
<td>0.9, 1.55, 2.0</td>
<td>3.18 (2.6 for Med_X) 18</td>
</tr>
<tr>
<td>Bird 2</td>
<td>Left</td>
<td>8</td>
<td>124.2</td>
<td>0.994</td>
<td>5.3, 5.8, 6.3</td>
<td>1.0, 1.6, 2.2</td>
<td>3.1</td>
</tr>
<tr>
<td>Bird 2</td>
<td>Right</td>
<td>8</td>
<td>124.2</td>
<td>0.994</td>
<td>5.3, 5.8, 6.3</td>
<td>1.0, 1.53, 2.05</td>
<td>3.4</td>
</tr>
<tr>
<td>Bird 3</td>
<td>Left</td>
<td>8</td>
<td>179.4</td>
<td>1.435</td>
<td>5.1, 5.5, 5.9, 6.3</td>
<td>0.9, 1.55, 2.2 (0.8 for Med_X) 3.18 (2.6 for Med_X) 21</td>
<td></td>
</tr>
<tr>
<td>Bird 3</td>
<td>Right</td>
<td>8</td>
<td>234.6</td>
<td>1.877</td>
<td>5.1, 5.5, 5.9, 6.3</td>
<td>0.9, 1.33, 1.77, 2.2</td>
<td>3.18 (2.6 for Med_X) 21</td>
</tr>
<tr>
<td>Bird 4a</td>
<td>Left</td>
<td>8</td>
<td>179.4</td>
<td>1.435</td>
<td>5.1, 5.5, 5.9, 6.3</td>
<td>0.9, 1.55, 2.2 (0.8 for Med_X) 3.18 (2.6 for Med_X) 21</td>
<td></td>
</tr>
<tr>
<td>Bird 4a</td>
<td>Right</td>
<td>8</td>
<td>179.4</td>
<td>1.435</td>
<td>5.1, 5.5, 5.9, 6.3</td>
<td>0.9, 1.55, 2.2 (0.8 for Med_X) 3.18 (2.6 for Med_X) 21</td>
<td></td>
</tr>
<tr>
<td>Bird 5</td>
<td>Left</td>
<td>8</td>
<td>124.2</td>
<td>0.994</td>
<td>5.1, 5.7, 6.3</td>
<td>0.9, 1.55, 2.2</td>
<td>3.18</td>
</tr>
<tr>
<td>Bird 5</td>
<td>Right</td>
<td>8</td>
<td>138</td>
<td>1.104</td>
<td>5.1, 5.7, 6.3 (4.8 for Med_X) 0.9, 1.55, 2.2 (0.8 for Med_X) 3.18 (2.6 for Med_X) 14</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Table 2. Injection parameters for sham-lesioned birds**

<table>
<thead>
<tr>
<th>Bird ID</th>
<th>Hemisphere</th>
<th>Injection parameters</th>
<th>Days killed after surgery</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bird 6</td>
<td>Left</td>
<td>Identical to Bird 5 but no 6-OHDA</td>
<td>17</td>
</tr>
<tr>
<td>Bird 6</td>
<td>Right</td>
<td>Identical to Bird 5 but no 6-OHDA</td>
<td>17</td>
</tr>
<tr>
<td>Bird 7</td>
<td>Left</td>
<td>Identical to Bird 2 but no 6-OHDA</td>
<td>18</td>
</tr>
<tr>
<td>Bird 7</td>
<td>Right</td>
<td>Identical to Bird 2 but no 6-OHDA</td>
<td>18</td>
</tr>
<tr>
<td>Bird 8</td>
<td>Left</td>
<td>Identical to Bird 1 but no 6-OHDA</td>
<td>27</td>
</tr>
<tr>
<td>Bird 8</td>
<td>Right</td>
<td>Identical to Bird 1 but no 6-OHDA</td>
<td>27</td>
</tr>
<tr>
<td>Bird 9</td>
<td>Left</td>
<td>Identical to Bird 3 but no 6-OHDA</td>
<td>23</td>
</tr>
<tr>
<td>Bird 9</td>
<td>Right</td>
<td>Identical to Bird 3 but no 6-OHDA</td>
<td>23</td>
</tr>
</tbody>
</table>

*AP and ML coordinates are on a grid such that each AP coordinate is paired with each ML coordinate in three or four rows. AP and ML coordinates are relative to Y0, and ML coordinates are offset to the left or right of Y0 depending on the hemisphere. DV coordinates are relative to the exposed brain surface. “Med_X” indicates a single extra injection that was intended (but not conclusively proven) to hit the medial-most portion of area X, where its pear shape comes to a dorsal and posterior point.

*In two additional birds, we performed unilateral 6-OHDA lesions and Th/MelATs to compare area X cell counts in sham- and 6-OHDA-lesioned hemispheres (see Materials and Methods). In these birds, we used identical injection coordinates and volumes as Bird 4.
Between each of the following steps, tissue was rinsed three times for 10 min after each step, except for blocking. Because all sections were cut parasagittally, medial and lateral subregions were designated by categorizing each section as belonging to either the medial or lateral half of area X. Using the binarization shown in b, we quantified the fraction of area X in which TH stain was reduced (see Materials and Methods). We also quantified the density of TH-positive fibers both within and outside the lesioned subregion of area X (“lesioned ROI” and “nonlesioned ROI,” respectively) in individual histological sections. Examples of lesioned and nonlesioned ROIs are shown as filled and empty squares in a, respectively. Within each section, we normalized the fiber density in the lesioned ROI by the density in the nonlesioned ROI from the same image. Histogram and error bars indicate the mean ± SEM of this measure across five 6-OHDA-injected birds and one sham. *p < 0.05 (two-sided Kolmogorov–Smirnov tests). We obtained the same result when raw (un-normalized) fiber density measures were used.

Histology

Each bird was perfused 14–23 d after 6-OHDA or sham lesion. Dissected brains were fixed overnight at 4°C in 4% formaldehyde, sunk in 30% sucrose for 1–4 d, and sliced in 40 μm sections on a microtome. We performed chromogenic TH stains on odd-numbered sections (to assess loss of area X catecholaminergic fiber innervation) and either Nissl stains (7 birds) or fluorescent NeuN and fluorescent TH stains (2 birds; one sham, one 6-OHDA) on even-numbered sections (to assess postlesion necrosis).

In two additional birds, we performed unilateral 6-OHDA lesions (one bird in left and one in right hemisphere) and perfused 11 d after surgery. We performed chromogenic tyrosine hydroxylase (TH) and fluorescent NeuN stains on alternating sections to compare area X cell counts in sham- and 6-OHDA-lesioned hemispheres together with the two bilaterally lesioned NeuN-stained birds mentioned above. All hemispheres in these birds used the same injection coordinates and volumes as Bird 4 in Table 1. These birds were housed singly after surgery (no other birds were present), and no behavioral data were collected from them.

For chromogenic TH immunohistochemistry, all steps used 0.2 M PBS (23 g sodium phosphate (dibasic) + 5.25 g sodium phosphate (monobasic) per 1 L deionized H₂O) as the solvent unless otherwise indicated. Between each of the following steps, tissue was rinsed three times for 10 min in 0.2 M PBS. Tissue was then incubated in 0.3% H₂O₂ for 30 min and 1% NaBH₄ for 20 min. It was then incubated at room temperature overnight in a solution containing primary antibody against tyrosine hydroxylase (Millipore MAB318; 1:2000) and NeuN conjugated to AlexaFluor488 (Millipore ABN78A4; 1:2000) in 1% normal horse serum and 0.5% Triton X-100. Tissue was then rinsed and incubated in biotinylated anti-mouse secondary (Vector Labs BA-2000; 1:200 + 0.5% Triton X-100) for 1 h at room temperature, rinsed, and incubated in streptavidin-AMCA (SA-5008; 5 μg/ml + 0.5% Triton X-100) for 1 h at room temperature. Sections were mounted and cover-slipped with Fluoro Gel with 1,4-diazabicyclooctane. Similar to chromogenic immunohistochemistry, 0.2 μm PBS was used as a solvent, and tissue was rinsed three times for 10 min after each step, except for blocking.

Image analysis

Lesion size and location. We quantified both the fraction of area X that exhibited reduced TH label and the extent to which the lesions affected different subregions of area X by measuring lesion-induced changes in the density of TH label. Images were acquired on a slide scanner (Meyer Instruments PathScan Enabler IV; 24 bit color, 7200 dpi, “sharpen more” filter, brightness, and contrast level 50). A custom-written ImageJ (version 1.47) macro was used to manually outline area X as an ROI on each TH-stained section.

As shown in Figure 2b, in each image an optical density (OD) threshold was established and then used to binarize the image so that each pixel within area X was categorized as belonging to either the “lesioned” (indicated by a lighter TH stain in that area) or “nonlesioned” subregion of area X. We use the terms “lesioned” and “nonlesioned” to differentiate subregions of area X in 6-OHDA-injected birds that do or do not exhibit a loss of TH label (i.e., these terms do not refer to 6-OHDA injected vs sham-lesioned animals). Because the level of background staining varied somewhat across sections, the OD threshold was set manually. We then used the binarized images to quantify the fraction of area X in that image that had been lesioned as follows:
dorsal, ventral, anterior, posterior, lateral, and medial subregions, as lesion in subregions of area X. To do so, we divided images of area X into location and behavioral data). Additionally, we quantified the fraction of ROI was treated as background signal, and OD ratio was computed as count for cross-section and cross-animal variations in stain density. We This ratio was calculated separately in each TH-stained section to ac-

where the lesioned and nonlesioned pixels are summed across the k sections of area X. This measure is summed across the two hemispheres, resulting in a single value of \( \alpha_{\text{total}} \) for each bird which was then compared with each bird’s learning behavior (see Relationship between lesion size/ location and behavioral data). Additionally, we quantified the fraction of lesion in subregions of area X. To do so, we divided images of area X into dorsal, ventral, anterior, posterior, lateral, and medial subregions, as shown in Fig. 2b. Thus, each of the six subregions comprised half of area X (e.g., the dorsal subregions included measurements from the dorsal half of area X in each section). We then calculated the fraction of each subregion that was lesioned (\( \alpha_{\text{dorsal}}, \alpha_{\text{ventral}}, \) etc.) using the procedure described above.

Alternate analysis of OD. In addition to the above analysis of lesion size and location, we performed an alternate analysis that did not rely on manually establishing an OD threshold. A custom-written ImageJ macro was used to manually outline area X as an ROI on each TH-stained section and place 0.5-mm-diameter ROI circles on representative areas of cortex (just dorsal to area X) and non-X-striatum (just posterior to area X near the dorsal border of the striatum). In some cases (e.g., Fig. 2a, right) loss of TH label extended slightly outside of the border of area X; the “non-X-striatum” ROI was positioned to exclude such areas. OD was quantified by converting the image to 8 bit grayscale and then measuring the average pixel value in each ROI.

To assess the effects of 6-OHDA injection into area X, we quantified OD\(_{\text{Area X}}\) the ratio of OD in the area X ROI to the non-X-striatum ROI.

This ratio was calculated separately in each TH-stained section to account for cross-section and cross-animal variations in stain density. We also performed an alternate analysis in which the OD of the cortex ROI was treated as background signal, and OD ratio was computed as \( \frac{\text{OD}_{\text{Area X}} - \text{OD}_{\text{cortex}}}{\text{OD}_{\text{Striatum}} - \text{OD}_{\text{cortex}}} \) within each stained section. This alternate technique yielded nearly identical results as the primary analysis. Quantifying the distribution of OD ratios in sham-lesioned animals (which is typically \( >1 \) because area X receives denser catecholaminergic input than the surrounding striatum) (Soha et al., 1996) allowed us to determine the 95% confidence interval of this metric in sham-lesioned brains. Any section from a 6-OHDA-injected animal with an OD ratio beyond the 95% interval therefore exhibits a significant reduction in TH staining density with \( p < 0.05 \).

Neuron counts. We quantified the number of surviving neurons following 6-OHDA (and sham) lesions in area X as well as in VTA/SNC, which sends a massive dopaminergic projection to the striatum, and in the locus ceruleus (LC), which may send a weak noradrenergic projection to area X (but see Mello et al., 1998; Castelino et al., 2007). We quantified the number of neurons in area X by imaging 4–10 sections from each of the four NeuN-stained brains (see Histology) at 40X magnification (0.75 zoom) using a Leica SP8 multiphoton microscope. We excluded 3 of the 29 images due to an imaging artifact. ImageJ was used to convert images to 8 bit grayscale, threshold based on pixel intensity to create a binary image, reduce noise (Remove Outliers in ImageJ), and separate touching cell bodies (Watershed in ImageJ). Cell bodies were counted using the Analyze Particles plug-in. Identical acquisition and processing parameters were used for all images.

To assess whether the number of NeuN-stained cells in area X differed between experimental conditions (6-OHDA vs sham), we performed a multinilinear regression analysis with birds and lesion condition as factors. Including each bird as a factor in the model increased our power by controlling for any between-bird differences. We combined data across NeuN-stained birds (one bilateral sham, one bilateral 6-OHDA, two unilateral 6-OHDA as described in Histology) and fit a standard multiple linear regression model (Kutner et al., 2005) as follows:

\[
y_i = \beta_0 + \beta_1 \cdot b_{2i} + \beta_2 \cdot b_{3i} + \beta_3 \cdot b_{4i} + \beta_4 \cdot C_i + e_i
\]

where \( y_i \) = cell counts for bird i and image j, \( C_i \) is an indicator variable to represent experimental condition (\( C_i = 1 \) if image i is from a 6-OHDA hemisphere, 0 if from a sham hemisphere), \( b_2-b_4 \) are indicator variables to represent bird-specific effects (\( b_2 = 1 \) if image i is from bird 2, 0 otherwise), \( \beta \) values are regression coefficients, and \( e \) is the residual error, assumed to be normally distributed. The term \( \beta_4 \cdot C \) represents the condition-specific effect after controlling for bird-specific effects (\( \beta_0 + \beta_2 + \beta_3 + \beta_4 \cdot b_{2i} + \beta_4 \cdot b_{3i} \)). Because indicator variables are 1 or 0 depending on the bird, the term \( \beta_4 \) represents the effect of Bird 1. To determine whether 6-OHDA-lesioned hemispheres had fewer cells in area X than sham-lesioned hemispheres, we performed a partial F test on whether \( \beta_4 \) is significantly different from zero after including the other factors.

To quantify neuron numbers in VTA/SNC and LC in each white noise trained bird, chromogenically TH-stained sections were imaged at 10X on an Olympus IX51 Widefield microscope with a Hamamatsu Orca ER CCD camera (for VTA/SNC) or an Axioplan widefield microscope with an Optronics camera (for LC). Ten sections containing VTA/SNC and two containing LC were imaged for each subject. Because sections were cut parasagittally, we did not attempt to identify the border between VTA and SNC. The most medial sections of VTA and the most lateral sections of SNC were not imaged because these regions contain few area X-projecting neurons (Person et al., 2008). Image acquisition parameters were held constant across subjects. In cases where the region being imaged was too large to fit into a single field of view, multiple images were taken and stitched together using the ImageJ Pairwise Stitching plug-in (Preibisch et al., 2009).

VTA/SNC and LC cell counts were performed using the ImageJ Cell Counter plug-in by four raters who were blinded to bird identity and treatment condition. Rater bias was quantified by having all raters count cells in the same histological sections and comparing mean counts across raters. The mean count from each individual was used to linearly scale all counts from that rater, with all correction terms having an absolute value of \( \pm 13\% \). Cell count results were qualitatively identical even if this correction term was not applied. As described in Results, cell counts from all birds revealed no significant difference in either VTA/SNC or LC. How-

However, a post hoc power analysis revealed that we would be unlikely to detect such change given the very small size of our neurotoxin injections, given that area X comprises \( \pm 10\% \) of the total volume of the basal ganglia (Karten et al., 2013) and that our lesions affected only part of area X. To perform the power analysis, we made two extremely conservative assumptions to put an upper bound on the number of catecholaminergic neurons that project to area X. First, we can assume that area X received 10% of the catecholaminergic input (the actual fraction is likely much lower given that both VTA and LC send inputs to the forebrain in addition to the striatum). Second, we can assume that 6-OHDA injections lesion will kill 100% of neurons that project to the affected region of the striatum (the actual fraction of neurons killed is likely significantly lower than this). Therefore, given that our lesions affected \( \pm 50\% \) of area X, we would expect that our lesions would kill at most 5% of catecholaminergic neurons projecting to area X (50% \( \times \) 10%), and likely much less.

We therefore performed a power analysis to quantify whether we would be likely to be able to detect a 5% change in neuron number. Across repeated measurements of the same TH-stained section, our cell counts had an SD of \( \sigma_{\text{action}} = 10 \) relative to the mean. We assessed the total number of TH\(^+ \) cells in the VTA by summing cell counts across \( n_{\text{sections}} = 10 \) histological sections. Assuming that cell counts of different sections represent independent measurements, we therefore expect that the total cell count for each bird has a SD of the following:
We then quantified the power of an analysis to detect a 5% difference in the number of TH \(^{-}\) neurons with a SD of 32% and a total of 5 measurements (5 birds per group). This analysis yielded a power of 0.07, indicating that we would only have a 7% chance of detecting such a difference. Our failure to detect a significant change in TH \(^{-}\) cell body number (see Results) is therefore unsurprising given the very small size of our neurotoxin injections within the basal ganglia.

Analysis of fiber density. For TH fiber density analysis in area X, sections were imaged at 63× using a Zeiss Axiosplan 2 Widefield microscope with an AxioCam HRC Color Camera. Lesioned and nonlesioned portions of area X within a single section were selected based on previously captured images (see Lesion size and location). Image acquisition parameters varied slightly between sections but were held constant for lesion-nonlesioned pairs within a single section.

Mammalian studies frequently induce unilateral lesions (e.g., injecting 6-OHDA into the striatum on one side and vehicle into the other side), allowing the experimenter to normalize the fiber density in the lesioned hemisphere to that in the opposite hemisphere to compensate for stochastic variations in TH stain intensity (i.e., animal-by-animal variation that is unrelated to the experimental condition). Because all birds used in our behavioral experiments received bilateral lesions of dopaminergic inputs to area X, we were unable to take this approach. Instead, we normalized the fiber density within the lesioned subregion of area X to the fiber density within a nonlesioned subregion in the same histological section, as described below. However, as described in the main text, we obtained qualitatively identical results when we did not perform this within-section density normalization.

Images were analyzed using ImageJ, with identical image-processing steps applied to every section. In each brain, we chose 10 tissue sections (five from each hemisphere) that contained both lesioned and nonlesioned subregions of area X. We then captured two images from each section, one from the lesioned and one from the nonlesioned subregion, and converted all images to 8 bit grayscale. To isolate TH-positive fibers, images were then bandpassed (FFT Bandpass Filter in ImageJ) to emphasize features with high spatial frequency (i.e., labeled axons) and then thresholded based on pixel intensity to create a binary image in which black pixels represented TH-stained fibers. After removing ouurring pixels (Remove Outliers in ImageJ), we then measured the density of TH-positive fibers by quantifying number of black pixels as a fraction of total pixels in the image. Fiber density from each lesioned subregion was then normalized to the density of the nonlesioned subregion in the same images. To assess the level of variation in this measure in a sham-lesioned bird, in one sham bird we randomly selected 5 of 10 ROIs to serve as the “lesioned” subregions.

High performance liquid chromatography (HPLC) In a separate set of adult (>100-d-old) male Bengalese finches (n = 6), we performed unilateral 6-OHDA lesions and used HPLC to compare area X DA and norepinephrine (NE) levels in lesioned and sham-lesioned hemispheres. Each bird received a 6-OHDA lesion in area X in one hemisphere and a sham lesion in the other hemisphere using the same procedure described in 6-OHDA and sham lesions. All hemispheres across birds used the same injection coordinates and volumes as Bird 4 in Table 1. We alternated which hemisphere was injected with 6-OHDA, so half the birds received lesions in the left and half in the right hemisphere. The birds were housed singly after surgery (no other birds were present), and we did not collect any behavioral data from these animals.

Fourteen days after surgery, we decapitated each bird, rapidly harvested the brains, flash-froze them in powdered dry ice 2–4 min after decapitation, and stored them at −80°C. Frozen brains were sliced into 300 μm parasagittal sections in a −12°C cryostat, placed on slides, briefly wet-thawed to room temperature (<20 s) to allow tissue to settle on the slide and placed on dry ice. In each hemisphere, we made 1-mm-diameter, 300-μm-thick circular tissue punches of area X in two sections using a previously described technique (Palkovits, 1973), placed both punches in a tube while still frozen and stored the sample at −80°C until tissue was analyzed for monoamine content. Area X was identified by observing the frozen and briefly wet-thawed sections with the naked eye and through a dissecting microscope using a bright light and dark surface to enhance contrast.

NE and DA concentrations were determined by HPLC with coulometric detection using established methods (Pozdeyev et al., 2008). Each sample was processed individually (one sample per hemisphere). Briefly, samples were first homogenized in 0.1 N HClO\(_4\) solution (containing 0.01% sodium metabisulfite and 25 ng/ml internal standard 3, 4-dihydroxybenzylamine HBr), and centrifuged at 13,000 × g for 15 min at 4°C. Supernatant fraction aliquots were injected into an UltraspHERE 5 μm ODS column, 250 × 4.6 mm (Hichrom) and separated with a mobile phase containing 0.1 μ sodium phosphate, 0.1 mM EDTA, 0.35 mM sodium octyl sulfate, and 7% (v/v) acetonitrile, pH 3.2. DA and NE amounts (ng/sample) were then quantified by comparison with internal standards, with a standard curve generated with 0.1–5 ng for each analyte. Protein (mg/sample) was determined using the Lowry protein assay with a standard curve generated with 0–95 μg BSA (Lowry et al., 1951).

Relationship between lesion size/location and behavioral data We used a stepwise regression procedure (Draper and Smith, 1998) to ask whether the magnitude and/or location of the loss of dopaminergic inputs to area X was predictive of the observed changes in learning behavior. To do this, we calculated the change in the absolute magnitude of learning due to 6-OHDA lesion as Δabsolute = μpost − μpre, where μpost and μpre are the magnitude of pitch change before and after lesion, respectively, on first baseline day and trial-matched postlesion day. We then asked which of seven measurements of lesion size and location (όventral, όventral, όanterior, όposterior, όanterior, όventral, όanterior) were significantly predictive of Δabsolute. Stepwise regression analysis provides a systematic method for testing, which predictor terms should be included in a multilinear model by beginning with an initial model and testing changes in the model’s predictive power that result from including or excluding individual predictor terms. We therefore applied this procedure to ask which, if any, of the seven candidate predictors should be included in a multilinear model that predicts Δabsolute. This analysis concludes when neither including nor excluding any additional terms significantly improves the model (p < 0.05 after Bonferroni correction). In an alternate analysis, we used the same seven candidate predictors to produce a model of Δrelative, which quantifies the fractional reduction (“percent decrease in learning”) in learning behavior after lesion:

\[
Δ_{\text{Relative}} = \left(1 - \frac{\mu_{\text{post}}}{\mu_{\text{pre}}}\right) \times 100\%
\]

In one 6-OHDA-lesioned bird, in the postlesion experiment, the bird made a small pitch change in the adaptive direction (i.e., υpost was negative). In this case, Δrelative was set to 100%.

Results

We injected 6-OHDA into area X of adult male Bengalese finches, measured the ensuing effects on song performance and vocal learning, and quantified the lesion-induced depletion of area X’s dopaminergic input. Following previous studies in mammals, we quantified dopaminergic innervation using an immunohistochemical stain for TH, the rate-limiting enzyme in the DA synthesis pathway (Fig. 1b). As shown in Figure 2a, 6-OHDA injections substantially reduced TH label within area X. To quantify the volume of area X affected by the lesion, we manually set an OD threshold for each image (see Materials and Methods; and Fig. 2b) and counted the fraction of pixels in which TH density fell below the threshold. As shown in Figure 2c, by this metric TH stain was reduced in 46%–68% of area X across birds, indicating that our lesions affected approximately half of the volume of the nucleus. Furthermore, to obtain a more direct measure of the lesions’ effects on dopaminergic innervations, we analyzed tissue at high magnification to quantify the prevalence of TH-positive axons within area X (Fig. 2d). We found that 6-OHDA injections
reduced TH-positive fiber density to 51%–84% of the normal value within the lesioned subregions of area X (Fig. 2e). In the analyses shown in Figure 2a–c, we manually set an OD threshold to delineate the lesioned and nonlesioned subregions of area X (see Materials and Methods). To verify that these results were not an artifact of this procedure, we also performed an alternate analysis of lesion volume that did not rely on manual thresholding. As shown in Figure 3, this alternate analysis similarly found that 6-OHDA injections led to significant reductions in TH stain in approximately half of area X.

In addition to optical imaging of TH-stained sections, we also quantified the extent to which 6-OHDA injections reduced DA concentrations within area X using HPLC. In a separate cohort of Bengalese finches (n = 6), we injected one hemisphere with 6-OHDA and performed a sham lesion on the other hemisphere. This design allowed us to control for interindividual differences in neurotransmitter levels. As shown in Figure 4a, left, 6-OHDA lesions reduced the concentration of DA by an average of 47.1% (range 9.3%–74.1%, mean concentration 206.1 and 390.5 ng DA/mg protein in lesioned and sham hemispheres respectively).

6-OHDA is toxic to both dopaminergic and noradrenergic neurons, and TH staining labels both types of neurons. It was therefore important to consider the possibility that any observed effects of 6-OHDA injections on both behavior and TH stain density might reflect changes in noradrenergic input as well as (or instead of) changes in DA. However, we think that this possibility is extremely unlikely. Catecholaminergic innervation of area X has previously been shown to be overwhelmingly dopaminergic (Mello et al., 1998; Gale and Perkel, 2005; Castelino et al., 2007). Studies of noradrenergic inputs to area X have reported that such inputs are either absent (Mello et al., 1998) or extremely weak (Castelino et al., 2007), and NE concentration within area X has been reported to be <3% of that of DA (Gale and Perkel, 2005). Consistent with these prior findings, NE concentrations assessed by HPLC were <2% as great as DA concentrations in sham hemispheres (Fig. 4a, right; mean 1.2%, range 0.5%–1.6%, mean concentration 4.9 and 4.3 ng NE/mg protein in lesioned and sham hemispheres, respectively), and furthermore were not significantly affected by 6-OHDA injections (Fig. 4b). Therefore, loss of noradrenergic inputs to area X is very unlikely to have affected our results.

Importantly, staining for the neuron-specific nuclear protein NeuN revealed that 6-OHDA injections did not reduce the number of neuronal somata within area X relative to sham injections (Fig. 5), suggesting that 6-OHDA injections reduced dopaminergic inputs without killing neuronal cell bodies in the basal ganglia. Additionally, we examined lesion-induced loss of dopaminergic neurons by counting TH-positive cell bodies in midbrain nuclei VTA/SNC and assessed lesion-related changes in noradrenergic neurons by counting TH-positive cells in the LC. Cell counts revealed no significant difference in either area (2655 ± 427 mean ± SD for VTA/SNC in sham birds; 2598 ± 369 for VTA/SNC in lesioned birds; 193 ± 40 for LC shams; 201 ± 48 for LC lesions; p > 0.8 in all cases, Kolmogorov–Smirnov test).

Because increased DA within area X during female-directed song is associated with reductions in acoustic variability (Sasaki et al., 2006; Leblois et al., 2010; Leblois and Perkel, 2012; Murugan et al., 2013) depleting DA with 6-OHDA injections might cause vocal variability to increase in directed song even if it does not affect the variability of undirected song (Fig. 6c). Quantitative acoustic analysis revealed that 6-OHDA injections caused no consistent changes in either the mean or variance of syllable pitch (Fig. 6b), sound amplitude, or spectral entropy (p > 0.25, Kolmogorov–Smirnov tests).
lesion and postlesion directed song from 4 birds (see Materials and Methods) and analyzed pitch variability. Consistent with prior results (Kao et al., 2005; Kao and Brainard, 2006; Sakata et al., 2008), we found a lower pitch SD in prelesion directed versus undirected song ($p < 0.05$, Wilcoxon signed-rank test; prelesion directed SD: mean 0.36 semitones, range 0.17–0.64 semitones; prelesion undirected: 0.47, 0.23–0.98). Interestingly, we did not find differences in either prelesion versus postlesion directed pitch SD or prelesion versus postlesion undirected pitch SD ($p > 0.05$, Wilcoxon signed-rank tests). Nor did we find a difference in postlesion directed versus undirected pitch SD ($p > 0.05$, Wilcoxon signed-rank test). The ~50% reduction in DA induced by 6-OHDA lesions (Figs. 2, 4) therefore appeared to have no significant effect on acoustic variability in either female-directed or undirected song.

However, we caution that these results are based on a relatively small dataset of directed song that is likely underpowered to detect subtle differences. Bengalese finches produce directed song much less readily than do zebra finches. Although we were able to collect some directed song (from 11 syllables prelesion and 8 syllables postlesion, with a mean of 110 and 88 iterations per syllable, respectively), attempting to collect more female-directed song would have significantly impeded our examination of 6-OHDA’s effect on learning because introducing female songbirds acutely reduces the total amount of song production. Importantly, we note that a recent study of the effects of 6-OHDA lesions of area X in zebra finches collected a much larger amount of female-directed song and reports significant variability changes after 6-OHDA lesions in area X (Miller et al., 2015).

In each bird, we compared vocal learning before and after either 6-OHDA or sham (saline) injections into area X (Fig. 1c). We evoked learning by providing disruptive auditory feedback (white noise blasts) conditional on the sung pitch of a particular syllable (Tumer and Brainard, 2007). In response to this reinforcement training, birds modify the pitch of the targeted syllable to avoid white noise, as shown in a representative prelesion experiment (Fig. 7a, black line, b). Following 6-OHDA injections, learning was greatly reduced in this bird (Fig. 7a, red line, c). Averaged across all subjects, the rate of learning decreased by >50% following 6-OHDA injections (Fig. 7d), whereas no reduction was seen following sham surgeries (Fig. 7e).

In addition to directly comparing the time course of learning across the first 3 d of white noise (Fig. 7a–f), we also quantified learning on trial-matched prelesion and postlesion days (see Materials and Methods) and similarly found reduced learning in 6-OHDA-lesioned (Fig. 7g) but not sham-lesioned birds (Fig. 7h). This analysis addresses a confound that could arise if we interpreted learning solely on a chronological basis (Fig. 7a–f). Because of normal variation in singing rate, birds sang a somewhat different number of songs by postlesion day 3 than by prelesion day 3. Hence, if a particular bird learned less by postlesion day 3 (compared with prelesion day 3), it could occur because that bird experienced substantially fewer learning trials, not because of DA lesion. Likewise, if a bird learned the same amount by postlesion day 3, it could occur because that bird experienced substantially more trials, masking a DA-dependent learning deficit. Therefore, it is crucial to compare prelesion/postlesion both chronologically (Fig. 7a–f) and using a postlesion day where birds had a similar number of trials (within 10%) as in the prelesion experiment (Fig. 7g–i). Additionally, because difference between $p$ values does not always correspond to a difference between effects (Nieuwenhuis et al., 2011), we directly compared the (post − pre) learning changes between conditions and found significantly reduced learning in 6-OHDA birds (Fig. 7i). To our knowledge, these results provide the first direct evidence that vocal learning in songbirds depends strongly on dopaminergic input to the basal ganglia.

As described in Materials and Methods, animals were randomly assigned to either the 6-OHDA lesion or sham group. Notably, two of the birds randomly assigned to the lesion group exhibited stronger prelesion learning than did their counterparts in the sham group (compare black symbols showing “prelesion” and “pre-sham” learning values in Fig. 7g and 7i, respectively). As a result, the prelesion learning data combined across subjects (Fig. 7d, black trace) exhibited noticeably greater learning than the sham data (Fig. 7e, black trace). To assess whether the apparent effects of 6-OHDA on learning could have arisen from a difference in prelesion/sham learning ability, we repeated the analysis in Figure 7d after excluding the two animals that exhibited the greatest prelesion learning. As shown in Figure 7f, this reanalysis yields comparable prelesion/sham learning (compare black traces in Fig. 7e and 7f) and, similar to the full dataset from lesioned animals, reveals a significant drop in learning ability following 6-OHDA injection, demonstrating that the 6-OHDA-dependent reduction in learning shown in Figure 7d, e was not an artifact of a difference in learning ability in the two subject groups before lesion or sham injections.
We used a stepwise regression procedure to quantify whether the size or location of the loss of dopaminergic input within area X predicted the magnitude of behavioral effects shown in Figure 7g. We found that none of the candidate predictor values, which included the total fraction of area X in which TH stain was reduced (α_total), as well as the fraction within six different subregions of area X (anterior, posterior, medial, lateral, dorsal and ventral), were significantly correlated with either the absolute or relative change in vocal plasticity (ΔAbsolutep, or ΔRelatedp, see Materials and Methods), either as individual predictors or in any combination. However, it should be noted that our dataset contains a somewhat limited range of lesion sizes (Fig. 2c), potentially limiting our ability to identify such effects. Notably, anatomical studies have shown a topographic mapping between different subregions of area X and downstream components of the song system (Luo et al., 2001), suggesting that different portions of area X might be dedicated to the modification of particular vocal parameters, such as pitch (Luo et al., 2001; Fee and Goldberg, 2011). Although our analysis did not produce positive evidence for such specificity, the spread of 6-OHDA within area X prevented us from fully assessing this idea by precisely confining lesions to particular subregions of the nucleus.

Following training, we turned off white noise playbacks and continued to monitor vocal acoustics for at least 3 d (see Materials and Methods). In contrast to the large deficits in learning observed during operant conditioning (Fig. 7), 6-OHDA lesions did not appear to impair spontaneous pitch restoration after learning. Both before and after lesion, the pitch of song changed monotonically toward the baseline (pretraining) value (Fig. 8a). Furthermore, quantifying the time constant of restoration demonstrated that pitch actually recovered significantly faster after 6-OHDA lesion than prelesion (τPrelesion = 2.15 d, τPostlesion = 0.87 d; Fig. 8b). However, the faster time constant of restoration postlesion does not necessarily reflect enhanced learning after 6-OHDA lesions. Rather, the observed difference in time constant may reflect the well-established phenomenon (Sober and Brainard, 2012; Kelly and Sober, 2014) that learning speed increases when the experienced sensory error is smaller relative to baseline. Indeed, in postlesion experiments, birds began restoration with a smaller error because learning was impaired (compare the last white noise day for prelesion and postlesion experiments in Fig. 8a). The fit parameter pFinal, which estimates the eventual equilibrium state of learning, was close to zero in both cases (pFinal = −0.04 semitones prelesion, pFinal = 0.05 semitones postlesion), suggesting that both prelesion and postlesion animals would have returned pitch to near the baseline value had washout been allowed to run for longer than the three post-white noise washout days shown in Figure 8a.

To control for the fact that learning speed depends on sensory error magnitude, we further examined the effects of DA lesion on pitch restoration by comparing postlesion restoration (Fig. 8, red traces) with data from specially selected subsets of experiments performed in the non-6-OHDA-lesioned condition. Specifically, we created these subsampled datasets by progressively eliminating the nonlesioned animals with the greatest learning on the last day of prelesion white noise playbacks, thereby producing a set of values that were more precisely confined to the range of learning observed in postlesion experiments (Fig. 6b). This approach enabled us to test whether the nonlesioned animals with the greatest learning on the last day of prelesion white noise playbacks showed a significant increase in learning rates compared to the entire sample. The results from this analysis revealed a significant difference between the two groups, with the nonlesioned animals showing a more rapid approach to the baseline value of learning (τPostlesion = 0.7, partial F test; see Materials and Methods). This suggests that, while lesions decreased dopaminergic inputs to area X (Figs. 2, 3, 4), 6-OHDA injections did not kill neurons with cell bodies within area X, n.s., Not significant.

**Figure 5.** 6-OHDA injections do not lead to neuron loss within area X. **a**, Representative NeuN-stained images from birds that received sham (top) and 6-OHDA (bottom) lesions. In each section, we counted the number of neuronal cell bodies (right column; see Materials and Methods). **b**, Area X images were taken from two bilaterally and two unilaterally lesioned birds (each 369 × 369 μm). Blue and red circles represent the number of cell bodies in individual sections. Open circles represent the values for the non-6-OHDA-lesioned condition. Specifically, we created these subsampled datasets by progressively eliminating the nonlesioned animals with the greatest learning on the last day of prelesion white noise playbacks, thereby producing a set of values that were more precisely confined to the range of learning observed in postlesion experiments (Fig. 6b). This approach enabled us to test whether the nonlesioned animals with the greatest learning on the last day of prelesion white noise playbacks showed a significant increase in learning rates compared to the entire sample. The results from this analysis revealed a significant difference between the two groups, with the nonlesioned animals showing a more rapid approach to the baseline value of learning (τPostlesion = 0.7, partial F test; see Materials and Methods). This suggests that, while lesions decreased dopaminergic inputs to area X (Figs. 2, 3, 4), 6-OHDA injections did not kill neurons with cell bodies within area X, n.s., Not significant.
white noise day until the remaining nonlesioned animals showed nearly identical vocal errors on the final white noise day as did the DA-lesioned population. Figure 8c shows a version of this analysis in which we compared all post-6-OHDA animals (red, n = 5 experiments) with data selected from prelesion, presham, and postsham animals so that the selected nonlesioned dataset ("Non-6-OHDA (selected)," n = 6 experiments; Fig. 8c, blue trace) had approximately the same initial error as the postlesion data (Fig. 8c, red trace). In an alternate version of this analysis (Fig. 8e), we selected the nonlesioned datasets only from postsham animals ("Postsham (selected)," n = 2 experiments). In both cases, these alternate analyses (Fig. 8d,f) yielded qualitatively the same results as those shown in the initial analysis (Fig. 8b), with significantly faster learning after 6-OHDA lesion (Fig. 8d,f, asterisks) and $p_{\text{max}}$ values very close to zero ($p_{\text{max}} = 0.004$ and 0.1 semitones for the nonlesioned data shown in Fig. 8d and Fig. 8f, respectively). Thus, although analysis of our relatively short washout period does not allow us to make strong conclusions regarding the effects of DA lesions on spontaneous error correction, our analyses clearly indicate that restoration back to pitch baseline is not impaired by reduction of dopaminergic inputs to area X, as is learning guided by white noise, and indeed may be facilitated by 6-OHDA lesions (see Discussion).

**Discussion**

Our experiments show that 6-OHDA injections into the songbird basal ganglia nucleus area X caused significant loss of DA inputs without causing detectable loss of neurons within area X. These dopaminergic lesions caused significant vocal learning deficits when pitch changes were driven by white noise reinforcement but did not result in measurable changes in song performance, song variability, or pitch restoration to baseline after white noise was discontinued. These results suggest dopaminergic inputs to the basal ganglia are critical for guiding vocal learning, at least when learning is driven by external reinforcement signals.

Although we took pains to precisely target 6-OHDA injections to area X, and indeed loss of TH stain was mostly confined to this nucleus, in some cases we observed loss of label in the striatum immediately surrounding area X (Fig. 5a). Importantly, the "shell" region immediately surrounding area X is hypothesized to be part of a circuit parallel to the classical song system shown in Figure 1a. Although cortical components of this parallel system appear to contribute to vocal learning (Iyengar et al., 1999; Person et al., 2008; Bottjer and Altenau, 2010), the involvement of area Xshell in learning remains to be directly tested. Because some spillover of 6-OHDA is inevitable, we cannot exclude the possibility that some of the observed effects on vocal learning reflect loss of dopaminergic input to the shell surrounding area X. However, we consider this possibility unlikely given that the loss of label outside of area X affected a very small fraction of the surrounding striatum.

Because female-directed song has lower acoustic variability and is associated with increased DA in area X (Sasaki et al., 2006; Leblois et al., 2010; Leblois and Perkel, 2012; Murugan et al., 2013), we expected to see increased vocal variability after 6-OHDA lesions. However, we did not observe a significant change in pitch variability in either direction following 6-OHDA injections into the songbird basal ganglia (Fig. 3a). Al- though cortical components of this parallel system appear to contribute to vocal learning (Iyengar et al., 1999; Person et al., 2008; Bottjer and Altenau, 2010), the involvement of area Xshell in learning remains to be directly tested. Because some spillover of 6-OHDA is inevitable, we cannot exclude the possibility that some of the observed effects on vocal learning reflect loss of dopaminergic input to the shell surrounding area X. However, we consider this possibility unlikely given that the loss of label outside of area X affected a very small fraction of the surrounding striatum.

The lack of an effect of lesions on social context-dependent variability may reflect the incomplete nature of our lesions, which spared a substantial number of DA terminals within area X. Alternatively, it is possible that neuromodulatory factors in addition to DA or neural circuits other than area X also contribute to context-dependent changes in variability and were able to compensate for lesion-induced changes.

Notably, a recent study analyzed vocal variability after 6-OHDA lesions of dopaminergic input to area X in zebra finches.
Figure 7. Removal of DA inputs to area X impairs reinforcement-driven vocal learning. **a**, In an example experiment, a bird received 3 d of training in which higher-pitched renditions of a syllable were punished by a disruptive auditory stimulus (see Materials and Methods). Black and red traces represent the pitch of the targeted syllable (mean ± SEM) before and after 6-OHDA injections, respectively, and illustrate a substantial reduction in learning magnitude following lesion. Pitch changes in the adaptive direction (downwards) are plotted as positive values. **b, c**, Prelesion (★) and postlesion (●) pitch distributions for the experiment shown in **a**. Gray bars represent the 3 d baseline pitch distribution. Dashed lines indicate the threshold for white noise playback (i.e., any pitches sung above that threshold received white noise). In every experiment, learning was driven in the same syllable and in the same direction prelesion and postlesion. **d**, Group data for lesioned (6-OHDA-injected) animals. Solid lines indicate the pitch of the targeted syllable as in **a**, except that here data are combined across \( n = 5 \) lesioned animals. Dotted lines indicate linear regression to pitch data. \( * p < 0.0001, \) significantly different slopes (\( F \) test). **e**, Group data for \( n = 4 \) sham-lesioned animals, plotting and testing conventions as in **d**. Slopes of pitch as a function of time are not significantly different \( (p = 0.48, F \) test). **f**, Alternate analysis of data from 6-OHDA-lesioned animals, excluding the two subjects who showed the greatest prelesion learning (see Results and **g**). Plotting and testing conventions as in **d**; pitch slopes are significantly different \( (* p < 0.0001, F \) test). **g**, Adaptive pitch change on the last white noise day in the prelesion experiment (relative to baseline) versus adaptive pitch change on a trial-matched white noise day in the postlesion experiment (not necessarily day 3; see Materials and Methods). \( * p < 0.05, \) significant difference in prelesion and postlesion learning magnitude (one-sided Wilcoxon signed-rank test). **h**, Adaptive pitch change on the last white noise day for sham-lesioned animals (conventions as in **g**). **i**, Direct comparison between sham and 6-OHDA learning changes. \( * p < 0.05 \) (two-sample \( t \) test). n.s., Not significant.

(Miller et al., 2015). Contrary to our hypothesis, this study found significant reductions in undirected (but not female-directed) song variability, similar to the decreases in vocal variability observed in Parkinson’s disease (Ramig et al., 2008). Our conflicting results may be attributed to two factors beyond the obvious difference in the species being studied. First, as noted in Results, we had a relatively small sample size of interleaved directed/undirected songs, lowering our statistical power to detect subtle differences in variability. Second, our average 6-OHDA dosage was slightly higher (1.3 \( \mu \)g; Table 1), and we quantified variability at later time periods (13–22 d after lesion for directed song analysis compared with 4–5 d after lesion in Miller et al., 2015), raising the possibility of a complex relationship between the extent of DA depletion, time course of compensation, and changes in vocal variability. The effects of DA depletion on vocal variability in Bengalese finches therefore remain to be clarified by future studies.

Although 6-OHDA lesions caused vocal learning deficits during white noise training (Fig. 7), restoration to baseline pitch was not impaired during the washout phase (Fig. 8), suggesting that DA might play different roles in distinct forms of vocal learning. Wholesale lesions or inactivation of area X or LMAN (the output nucleus of the AFP) severely degrade both white noise-driven pitch learning and spontaneous restoration back to baseline (Andalman and Fee, 2009; Warren et al., 2011; Charlesworth et al., 2012; Ali et al., 2013). Our data indicating that only the former process depends on dopaminergic inputs to area X suggest that the basal ganglia might mediate these two forms of vocal plasticity in distinct ways. Interestingly, lesions in caudal medial nidopallium, proposed as a candidate site for template song memory, disrupt restoration but spare noise-avoidance learning (Canopoli et al., 2014), the opposite pattern we observed following dopaminergic lesions to area X. These observations support the idea that “vocal plasticity” can be divided into different subtypes, each driven by distinct yet interacting neural processes.

Despite the above considerations, however, our data suggesting that restoration is less impeded by DA loss than is noise-driven learning should be treated with a great deal of caution.
Figure 8. Removal of DA inputs to area X does not impair pitch restoration. a, Combined data across five 6-OHDA-lesioned birds during the restoration period, after white noise was discontinued. Washout day 0 is the last day of white noise (not necessarily day 3; see Materials and Methods). Black and red represent prelesion and postlesion experiments, respectively. Prelesion and postlesion restoration data were fit with an exponential decay model (dashed lines; see Materials and Methods). Birds restored pitch toward baseline in both prelesion and postlesion experiments. After 6-OHDA lesions, birds began with a lower absolute pitch difference from baseline because of postlesion learning deficits (Fig. 7d). b, Fitted time constant \( \tau \) for prelesion and postlesion learning. Lower \( \tau \) indicates faster return to baseline. In postlesion experiments, birds restored pitch significantly faster than in prelesion experiments (*\( p < 0.05 \), permutation test). Error bars indicate 95% confidence intervals. c, d, Same analysis as in a, b, but selecting nonlesioned datasets so as to approximately equalize the initial error size (i.e., to approximately equalize error on the last white noise day). Here nonlesioned datasets are selected from prelesion, presham, and postsham subjects (see Materials and Methods). e, f, Same analysis as in c, d, but with nonlesioned datasets drawn only from postsham subjects. Note different vertical scale in d compared with that in b, f, a, c, e; SEM error bars are obscured by the plotted circles.

First, our 6-OHDA injections only partially ablated DA within area X, and the robust restoration observed after lesion might reflect residual DA function. Second, because washout experiments necessarily occurred after white noise training, it is possible that some form of compensatory plasticity occurred in the few days that elapsed between the onset of postlesion white noise training and the beginning of the washout period. Third, it is important to emphasize that the return to baseline is not necessarily auditory-guided and could in theory be mediated by a bird’s matching proprioceptive feedback to the baseline motor command. Finally, as shown in Figure 8a, c, e, we did not collect washout data for sufficient time for syllable acoustic restoration to fully return to baseline either prelesion or postlesion. Therefore, although the speed of restoration appears unimpeded by 6-OHDA lesions, the endpoint of restoration following DA depletion remains to be measured directly. Intriguingly, the analyses shown in Figure 8 suggest that pitch restoration may actually progress more quickly following DA lesions, even when selecting subsets of the data so as to equalize the initial error size (Fig. 8c–f). Although the caveats detailed above prevent us from drawing strong conclusions about DA’s role in vocal learning other than that driven by white noise, future studies could ask whether DA is necessary for error-corrective learning by providing a correctable auditory perturbation to baseline song without explicit external reinforcement (Sober and Brainard, 2009; Hoffmann et al., 2012).

Prior studies have identified potential mechanisms by which dopaminergic inputs to area X might mediate vocal learning. The nuclei of dopaminergic neurons that project to area X reside in the VTA/SNc complex, which in turn receives input from forebrain neurons that respond to perturbations of auditory feedback during singing (Mandelblat-Cerf et al., 2014), providing a candidate pathway by which sensory error signals might influence DA release within the basal ganglia. Furthermore, dopaminergic signaling plays a crucial role in mediating plasticity at corticostriatal synapses in both mammals and songbirds (Ding and Perkel, 2004; Surmeier et al., 2007; Leblois, 2013), suggesting that DA might mediate vocal learning by modulating changes in synaptic strength between cortical area HVC and spiny neurons in area X (Fee and Goldberg, 2011). Although our results strongly suggest that dopaminergic projections from VTA/SNc guide vocal plasticity, future studies (including recording from DA neurons that project to area X) are needed to investigate the nature of the signals conveyed by these projections.

Our lesions only partially eliminated dopaminergic inputs to area X, in contrast to studies in mammals in which injections of neurotoxins into the medial forebrain bundle produces near-complete ipsilateral loss of dopaminergic input throughout the striatum (Deumens et al., 2002). Specifically, our lesions reduced...
TH stain in approximately half of area X (Fig. 2c), and within the affected regions eliminated ~15%–50% of catecholaminergic axons (Fig. 2e), comparable with the fiber loss observed following intrastral 6-OHDA injections in mammals (Debeir et al., 2005). HPLC results similarly showed DA concentration dropping 47% on average (Fig. 4a). Because even these relatively modest reductions in dopaminergic innervation produced learning deficits, our results demonstrate that vocal learning is sensitive to disruptions of dopaminergic input to the basal ganglia and suggest that DA plays a crucial role in the processing of sensorimotor errors.

References


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