

FoxP2 Expression in Avian Vocal Learners and Non-Learners

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Most vertebrates communicate acoustically, but few, among them humans, dolphins and whales, bats, and three orders of birds, learn this trait. *FOXP2* is the first gene linked to human speech and has been the target of positive selection during recent primate evolution. To test whether the expression pattern of *FOXP2* is consistent with a role in learned vocal communication, we cloned zebra finch *FoxP2* and its close relative *FoxP1* and compared mRNA and protein distribution in developing and adult brains of a variety of avian vocal learners and non-learners, and a crocodile. We found that the protein sequence of zebra finch *FoxP2* is 98% identical with mouse and human *FOXP2*. In the avian and crocodylian forebrain, *FoxP2* was expressed predominantly in the striatum, a basal ganglia brain region affected in patients with *FOXP2* mutations. Strikingly, in zebra finches, the striatal nucleus Area X, necessary for vocal learning, expressed more *FoxP2* than the surrounding tissue at post-hatch days 35 and 50, when vocal learning occurs. In adult canaries, *FoxP2* expression in Area X differed seasonally; more *FoxP2* expression was associated with times when song becomes unstable. In adult chickadees, strawberry finches, song sparrows, and Bengalese finches, Area X expressed *FoxP2* to different degrees. Non-telencephalic regions in both vocal learning and non-learning birds, and in crocodiles, were less variable in expression and comparable with regions that express *FOXP2* in human and rodent brains. We conclude that differential expression of *FoxP2* in avian vocal learners might be associated with vocal plasticity.

Key words: language; winged-helix/forkhead box; transcription factor; songbird; hummingbird; parrot; ring dove

Introduction

Human speech and birdsong share behavioral and neural similarities (Doupe and Kuhl, 1999; Brainard and Doupe, 2002; Wilbrecht and Nottebohm, 2003; Jarvis 2004). Both are learned during a critical period via the interaction of auditory and motor centers and require specialized cerebral structures, which are functionally lateralized (Bottjer and Arnold, 1985). Whereas innate dispositions to learn and produce species-appropriate sounds are present in both humans and birds (Pinker, 1994; Doupe and Kuhl, 1999), mutations that predominantly affect vocal communication seem rare (Fisher et al., 2003). The only gene identified is *FOXP2*, the mutated form of which has been autosomal-dominantly inherited through three generations by approximately half of the family members of the KE family (Lai et al., 2001). Affected family members have difficulty with the cor-

rect articulation of speech and score lower in receptive and grammatical language tests than their unaffected relatives. Execution of sequenced orofacial movements is also impaired (for review, see Fisher et al., 2003; Marcus and Fisher, 2003). In contrast, nonverbal cognitive abilities differ less dramatically between individuals with or without *FOXP2* mutations (Vargha-Khadem et al., 1995; Watkins et al., 2002).

FOXP2 belongs to the large winged-helix/forkhead box (Fox) transcription factor gene family (Kaestner et al., 2000), originally identified in *Drosophila* (for review, see Kaufmann and Knochel, 1996). Forkhead genes play pivotal roles in development and adulthood (Carlsson and Mahlapuu, 2002). In mammals, three of the four *FoxP* family members were originally isolated from mouse lung, where *Foxp1* and *Foxp2* act as transcriptional repressors (Shu et al., 2001; Lu et al., 2002). Analysis of *FOXP2* molecular evolution suggests that this gene has been the target of positive selection during recent primate evolution (Enard et al., 2002).

Although it remains to be shown how *FOXP2* acts in the brain, rapid progress has been made in demonstrating where *FOXP2* acts. Structural and functional brain anomalies of affected individuals carrying *FOXP2* mutations implicate the basal ganglia as a key affected brain region (Watkins et al., 1999; Belton et al., 2003). The striatum, a component of the basal ganglia, is also the site of high *FOXP2* expression in the developing human and rodent brain (Ferland et al., 2003; Lai et al., 2003; Takahashi et al., 2003). Because vocal learning in songbirds depends on a special-

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ized pathway through the basal ganglia, including the striatal vocal nucleus, Area X (Sohrabji et al., 1990; Scharff and Nottebohm, 1991), we asked: (1) Do birds that learn to vocalize differ in *FoxP2* brain expression patterns from birds that vocalize innately? (2) How do brain expression patterns of avian *FoxP2* and its close homolog *FoxP1* compare with expression patterns in mammals, and how might these relate to the evolution of vocal learning? and (3) How similar is the amino acid sequence of FoxP2 protein in vocal learners (human, zebra finch) compared with a non-learner (mouse)? To address these questions, we cloned the *FoxP2* and *FoxP1* genes of a commonly studied vocal learner, the zebra finch, and used it as a probe to evaluate expression patterns in the brains of eight species of avian “vocal learners,” two species of “vocal non-learners,” and a crocodylian, the closest living non-avian relative (Meyer and Zardoya, 2003), presumably also a vocal non-learner.

Materials and Methods

Nomenclature. For avian brain regions, we used the recently revised nomenclature proposed by the Avian Brain Nomenclature Forum (Reiner et al., 2004b) (<http://avianbrain.org/>). For FoxP2 nomenclature, we followed the convention proposed by the Nomenclature Committee for the forkhead family of genes (i.e., FOXP2 in *Homo*, Foxp2 in *Mus*, and FoxP2 in all other species, proteins in roman type, and genes and RNA in italics) (Kaestner et al., 2000).

Species. We studied 11 avian species, ordered genetically relative to zebra finch: songbirds: zebra finches (*Taeniopygia guttata*), Bengalese finches (*Lonchura striata domestica*), strawberry finches (*Amandava amandava*), song sparrows (*Melospiza melodia*), canaries (*Serinus canaria*), and black-capped chickadees (*Phoebastria atricapillus*); dove: ring doves (*Streptopelia risoria*); hummingbirds: rufous-breasted hermit (*Glaucois hirsuta*) and somber hummingbird (*Aphantochroa cirrochloris*); parrot/budgerigar (*Melospiticus undulatus*); and chicken: domestic chicken (*Gallus gallus*). Relationships were assessed from DNA–DNA hybridization melting temperature half-life indices (T50) and general taxonomy in the literature (Sibley and Ahlquist, 1990; Zann, 1996; Wada et al., 2004). The zebra, Bengalese, and strawberry finches all belong to three closely related subfamilies: Poepphilini, Lonchurini, and Estrilini, respectively, of the Estrinidae family of birds. The song sparrow is a close relative belonging to the Passeridae family (T50 = 8.5 removed), followed by the canary belonging to Fringillidae (T50 = 10.0 removed), then black-capped chickadee belonging to Paridae (T50 = 11.1 removed). Doves are the closest non-songbird relative examined (T50 = 21.6 removed), followed by hummingbirds, with the somber hummingbird and rufous-breasted hermit being two ancient species belonging to the only two family lineages Trochilidae and Phaethornithinae, respectively (Jarvis et al., 2000) (each T50 = ~22.5 removed), followed by parrots (T50 = 23.1 removed), and finally by chicken being the most distant from songbirds (T50 = 28 removed) (Sibley and Ahlquist, 1990). A crocodile (*Alligator mississippiensis*) was chosen as the closest living relative of birds (Meyer and Zardoya, 2003) for comparison.

Zebra finches were obtained from breeding colonies kept at Duke University, Tokyo Medical and Dental University, the Max-Planck Institute for Molecular Genetics Berlin, and City College of New York. Some birds were isolated ~12 hr overnight, followed by 1–2 hr silent conditions in the morning with the lights on. When necessary, singing attempts were interrupted by human approach to examine *FoxP2* expression in the absence of singing behavior. Brain sections from zebra finches that had sung a known quantity of undirected songs before perfusion and quiet controls and canaries that sang at different seasons of the year, song sparrows, hummingbirds, and parrots were obtained from the Jarvis laboratory collection (Jarvis et al., 1997; Jarvis and Nottebohm, 1997; Jarvis and Mello, 2000; Jarvis et al., 2000). For the seasonal canary experiment, we used 10 μ m cryostat-cut sections of brains of adult (20–32 months old) male canaries that were killed monthly between April 1995 and March 1996 (Jarvis and Nottebohm, 1997; Holzenberger et al., 1997). Song behavior [whether singing occurred, what type of singing

(stereotyped or plastic), and how much singing] of each bird was scored and recorded 1 hr before sacrifice. Quiet control black-capped chickadee brain sections were obtained from Dr. Tom Smulders (Jarvis laboratory) and Dr. David Sherry (University of Western Ontario, Ontario, Canada). Strawberry finches and additional canaries were purchased from a Tokyo breeder. Crocodile brain sections were obtained as part of another study by E.D.J. and K.W. in collaboration with Daphne Soares (University of Maryland). All animals were treated under the humane guidelines of the institutes where animals were bred and kept.

Cloning of FoxP2 and FoxP1 cDNAs from zebra finch brain. Primers specific for the mouse *Foxp2* sequence were used to amplify zebra finch *FoxP2* from adult male zebra finch brain total RNA. We obtained a fragment covering bp 114–959 of isoform III, relative to first start codon, with primers 5'-GACACCAGCTCTGAAGTAAGCACA-3' and 5'-GGTAGTC-GAGGAGGAATTGTTAGT-3' and the entire *FoxP2* ORF with primers 5'-ATGATGCAGGAATCTGCGACA-3' and 5'-TCATCCAGATCTTCAGATAAAG-3'. A 180 bp *FoxP1* fragment was obtained using degenerated primers 5'-GARTTYTAYAAARAAYGCNGANGT-3' and 5'-ATRTGNGCNACNG-CRTTYTCC-3'. All PCR products were examined on agarose gels, cleaned from nucleotides with the Qiaquick PCR purification kit (Qiagen, Chatsworth, CA) and cloned into the pGEMTeasy vector (Promega, Madison, WI). Inserts from 15 independent *FoxP2* clones and 6 *FoxP1* clones were then sequenced on both strands. We obtained additional cDNA and, thus, sequence for each gene using the SMART-RACE kit (Clontech, Palo Alto, CA). Sequence assembly and analysis was done with GCG 10.1 (Accelrys, Cambridge, UK) and the Staden package (Staden et al., 1998). All sequences were deposited into GenBank (accession numbers AY549148, AY549149, AY549150, AY549151, and AY549152) and the songbird cDNA database (<http://www.dbsr.duke.edu/songbird>). One full ORF *FoxP2* clone and one containing the fragment covering bp 114–959, relative to the first start codon, as well as the 180 bp *FoxP1* clone, were selected for the generation of riboprobes.

In situ hybridization and quantification. *In situ* hybridizations were performed according to two protocols using ³³P-labeled (Vortkamp et al., 1996) or ³⁵S-labeled (Mello et al., 1997) riboprobes. Both protocols yielded identical labeling patterns in the brain. Riboprobes were *in vitro* transcribed from T7 and SP6 promoter sides of the pGEMTeasy cloning vector containing the *FoxP2* and *FoxP1* cDNA clones. Slides were exposed to x-ray film (Bio-max; Eastman Kodak, Rochester, NY) for 2–3 d (³⁵S-labeled material) or 1–3 d (³³P). For species comparison and developmental studies, a set of 163 slides with sections from 11 different species and from zebra finches of 12 different developmental ages [embryonic stages 10, 23, 26, and 28 and featherbud stage embryos corresponding approximately to chick stages 34 and 37, and post-hatch days (PHDs) 15, 25, 35, 45, and 75 and adults >90 d] were hybridized at the same time with a *FoxP2* mastermix, with the same counts per minute-radioactive count per slide. For the seasonal comparisons, all sections were also hybridized with a master mix. *FoxP1* was hybridized on another day to avoid the possibility of cross-contamination. For *in situ* quantifications, the exposed film was placed under a high-power dissecting scope (Wild M420; Leica, Deerfield, IL) and scanned into a computer using a Spot III camera and Spot software version 3.2.4 (Diagnostic Instruments, Sterling Heights, MI). Images were transferred to Photoshop (Adobe, San Jose, CA) and converted to grayscale. Vocal nuclei and adjacent non-vocal areas, i.e., the surrounding brain subdivisions (caudal nidopallium ventrally adjacent to HVC; nonauditory arcopallium abutting the robust nucleus of the arcopallium (RA); nidopallium rostral to lateral magnocellular nucleus of the anterior nidopallium (IMAN) and the surrounding shell region; caudal striatum (CSt) immediately caudal to Area X) were outlined with a selection tool, and the average pixel density was calculated using the Photoshop histogram function.

To calculate ratios of differential expression in vocal nuclei relative to their surrounding brain subdivision, we divided the pixel density values of vocal regions by the pixel density values of the respective adjacent region, using comparably sized areas for quantification. When expression within a given vocal nucleus is the same as the expression of the region surrounding it, the ratio is 1; when the expression within the vocal nucleus is higher than expression in the region surrounding it, the ratio is >1; when lower, the ratio is <1.

Preparation of brain sections. Serial sagittal or frontal sections were cut either frozen on a rotary cryostat (8 μ m or 10 μ m), paraffin embedded and cut on a rotary microtome (8 μ m), or paraformaldehyde perfused and cut on a vibratome (40 μ m). Before embedding in paraffin, the brains were immersed in Serra's fixative (60% EtOH, 30% concentrated formaldehyde, and 10% acetic acid). The paraffin-embedded and the fresh-frozen brain sections postfixed in 4% paraformaldehyde were used for *in situ* hybridizations (ISH). The fresh-frozen sections (slide mounted) or free-floating vibratome sections were used for immunohistochemistry (IHC).

Northern blotting. Twenty micrograms of total RNA from adult male zebra finch brain and lung were separated on a 1% denaturing agarose gel and blotted onto a nylon membrane (Amersham Biosciences, Piscataway, NJ) overnight. The membrane was then incubated with a 32 P-labeled *FoxP2* fragment spanning bp 114–959 relative to the first start codon of isoform III at 65°C for 3 hr. The blot was washed and exposed to an MS-intensifying screen (Eastman Kodak), which was then scanned with a PhosphorImager (Molecular Dynamics, Sunnyvale, CA) and analyzed with ImageQuant software 5.2 (Molecular Dynamics).

Western Blotting. Brains from PHD 40 male zebra finches were crushed in liquid nitrogen and then lysed in hypotonic lysis buffer containing 0.1% NP-40 for 10 min. Extracts were then centrifuged for 10 min at 1500 \times g, and the resulting pellet was dissolved in Laemmli buffer and denatured for 5 min at 95°C. Nuclear extracts were separated by SDS-PAGE (8%), transferred to a polyvinylidene fluoride membrane (Roche, Indianapolis, IN), and blocked with 1% BSA in PBS/0.1% Tween 20 for 1 hr. The membranes were then incubated with a rabbit polyclonal Foxp2 antibody [1:150 dilution; raised against aa 613–715 of mouse Foxp2 and immunopurified against Foxp1 protein (Lu et al., 2002)] for 1 hr, followed by incubation with an HRP-conjugated anti-rabbit antibody (1:2000 dilution; Amersham Biosciences) for another 30 min. Binding was detected on x-ray films using an ECL detection system for HRP (Perkin-Elmer, Boston, MA).

Immunohistochemistry. In addition to the Foxp2 antibody described above, we used the following primary antibodies in double-labeling experiments, recognizing: human neuronal protein HuC/D (monoclonal; Molecular Probes, Eugene, OR), cAMP-regulated phosphoprotein of molecular weight 32,000 (DARPP-32; polyclonal; Santa Cruz Biotechnology, Santa Cruz, CA), polysialylated neural cell adhesion molecule (PSA-NCAM; monoclonal; AbCys, Paris, France), tyrosine hydroxylase (TH; polyclonal; Santa Cruz Biotechnology), parvalbumin (monoclonal; Swant, Bellinzona, Switzerland), anti-nitric oxide synthase (rabbit polyclonal; Zymed, San Francisco, CA), calbindin (rabbit polyclonal; Swant), and ChAT (polyclonal; Chemicon, Temecula, CA). Paraformaldehyde (4%)-perfused tissue was washed three or more times in 0.1 M phosphate buffer (PB) and preincubated for a minimum of 30 min with a permeabilization/blocking buffer (0.1 M PB, 2% skim milk, and 0.3% Triton X-100), followed by incubations with primary antibodies either for 1 hr at room temperature or for 24–72 hr at 4°C. Secondary antibodies were either fluorescently labeled (FITC, Cy3, Cy5) or biotinylated. Biotinylated antibodies were reacted with ABC (Vector Laboratories, Burlingame, CA) and visualized with DAB or DAB/cobalt (SigmaFast; Sigma, St. Louis, MO) reaction. Double labeling was performed using two different fluorescent secondary antibodies or using DAB to reveal one antigen and DAB with nickel enhancement for the second antigen. The double labeling was done sequentially unless the primary antibodies were raised in different species. To control for nonspecific binding of the secondary antibodies, control reactions were performed that included all steps, except incubation with the primary antibody.

Results

Cloning of zebra finch *FoxP2* and *FoxP1*

We initially amplified an 845 bp fragment of zebra finch *FoxP2* (*zfFoxP2*) from adult male zebra finch brain cDNA using primers designed from the mouse *Foxp2* (*mFoxp2*) sequence (see Materials and Methods). With subsequent 5' and 3' RACE (Rapid Amplification of cDNA Ends), we assembled 2830 bp of *zfFoxP2* mRNA that included 296 bp of the 5' untranslated region (UTR), the

entire ORF of 2207 bp, and 327 bp of the 3' UTR (GenBank accession numbers AY549148, AY549149, AY549150, and AY549151). To further confirm the *zfFoxP2* sequence, we sequenced 12 independent clones carrying the entire ORF amplified from adult male zebra finch brain cDNA. We found that two DNA segments, which we called *splice1* (71 bp) and *splice2* (60 bp), were either present or absent in these clones, suggesting the existence of four *zfFoxP2* mRNA isoforms, each different at the 5' end of the gene (Fig. 1A). *Splice1* introduces a stop codon at position 261 (relative to the first start codon), resulting in predicted protein isoforms III or IV that miss the first 92 amino acids. In human but not mouse, the *splice1* fragment also exists (Bruce and Margolis, 2002). *Splice2* introduces 20 additional amino acids in-frame into the predicted protein isoforms I and III. When the *splice2* fragment is absent, it results in isoforms II and IV. In human and mouse, *splice2* is apparently never spliced out. Reverse transcription (RT)-PCR with RNA from a variety of zebra finch tissues using primers at both ends of the alternatively spliced region generated products that matched the sizes expected for the isoforms (Fig. 1B,C). There were, however, differences between tissues, with isoform IV being predominant in muscle, II–IV in lung, and all four in brain and liver (Fig. 1C). Northern hybridization on adult zebra finch brain and lung total RNA revealed four transcripts of ~9.0, 6.5, 3.5, and 2.5 kb, respectively (Fig. 1D). The 9.0, 3.5, and 2.5 kb transcripts corresponded in size to the transcripts found in mouse (Shu et al., 2001), whereas the 6.5 kb transcript matched the size of the human transcript (Lai et al., 2001). The size of the two most abundant *zfFoxP2* transcripts of 9.0 and 6.5 kb suggests that they contain large amounts of regulatory sequence, perhaps to precisely regulate *zfFoxP2* translation, mRNA location, and mRNA stability.

To determine which protein isoforms are found in the zebra finch brain, we probed juvenile zebra finch brain extracts with an antibody raised against aa 613–715 of mFoxp2 (Lu et al., 2002) by Western blot. This antibody should recognize all four isoforms. We could exclude the existence of abundant levels of the short isoforms III and IV, because no protein corresponding to their predicted molecular weight (Fig. 1A) was detected (Fig. 1E). Thus, isoforms III and IV are present only in a small population of cells or at low levels across most cells. In zebra finch brain, one or both of the long isoforms (I and II) predominate, although we could not distinguish between their similar molecular weights of 77 and 79 kDa, respectively (Fig. 1E). For the mFoxp2 protein, a molecular weight in this range has been observed (Lu et al., 2002).

The *zfFoxP2* protein (isoform I) shares 98.2% identity with the human FOXP2 protein and 98.7% identity with mouse Foxp2 protein (supplemental Fig. 1A; available at www.jneurosci.org). This underscores the extreme degree of conservation of the *FoxP2* gene (Enard et al., 2002; Zhang et al., 2002), because ~320 million years ago is the latest time at which modern mammals and birds shared a common ancestor (Evans, 2000). At five amino acid positions that are identical in mice and men, *zfFoxP2* differs. At three additional positions, the mouse and zebra finch sequence are identical but the human sequence diverges. Of these three amino acids, one also exists in carnivores (Zhang et al., 2002) (supplemental Fig. 1A, amino acid framed by circle), one is common to primates (supplemental Fig. 1A, boxed amino acid), and one is unique to humans (supplemental Fig. 1A, amino acid framed by triangle). In an analysis of FOXP2 molecular evolution, the latter has been suggested to result from positive selection during recent primate evolution, indicating that human FOXP2 might have been pivotal for the development of

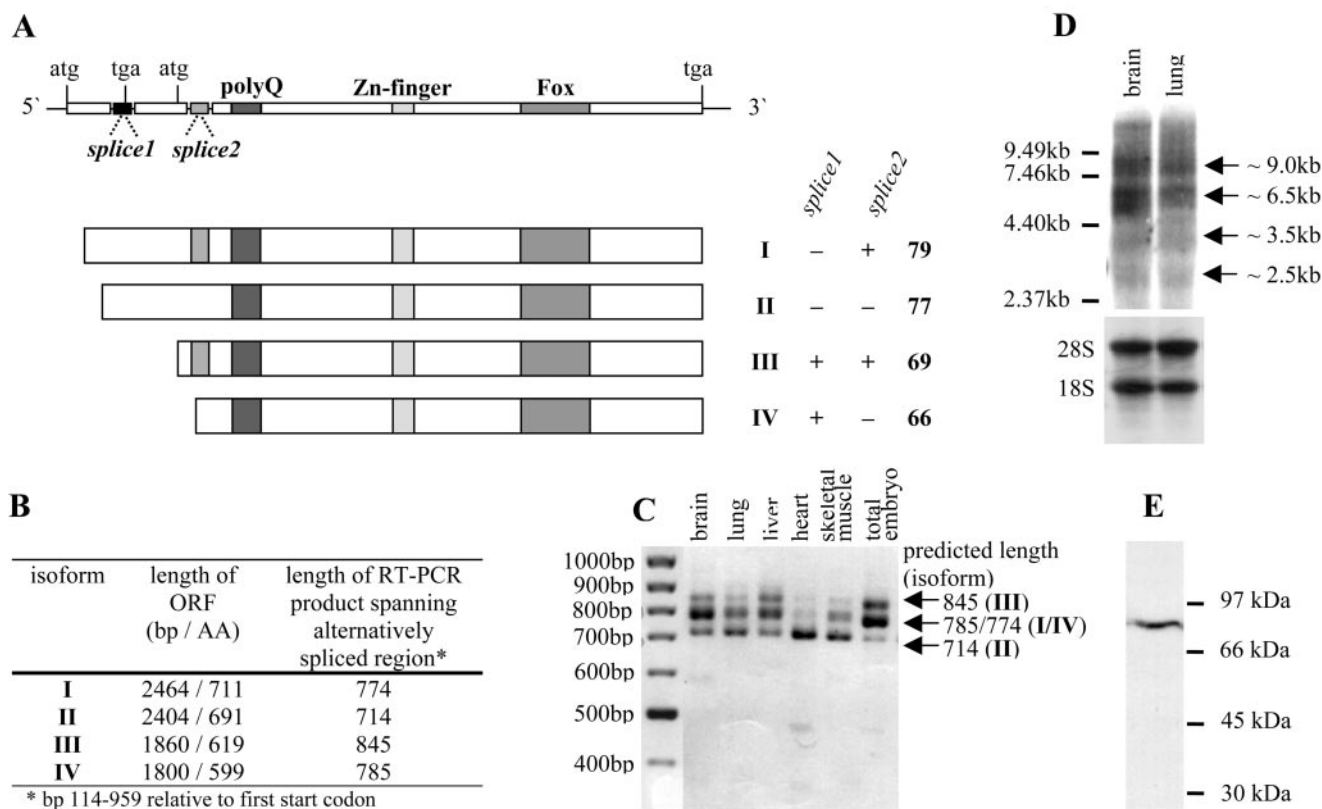


Figure 1. Identification of the zebra finch *FoxP2* (*zffoxp2*) mRNA. *A*, Schematic representation of the *zffoxp2* mRNA structure and its four predicted protein isoforms (I–IV). Positions of start (atg) and stop (tga) codons, the polyglutamine tract (polyQ), zinc finger (Zn-finger), and forkhead box (Fox) DNA-binding domains are shown. Two mRNA segments (*splice1* and *splice2*) are subject to alternative splicing. The presence (+) or absence (–) of *splice1* and *splice2* leads to variation in the length of ORFs. *Splice1* contains a stop codon that shifts the frame so that the ORF begins at the second atg, *splice2* inserts 60 bp in-frame into the coding region. The four predicted protein isoforms are shown. For the calculation of their theoretical molecular weight, we used Peptide Mass (<http://www.expasy.org/tools/peptide-mass.html>). *B*, Summary of length [in base pairs (bp) and amino acid (AA)] of the *zffoxp2* isoforms (I–IV) and the length of the RT-PCR products spanning the alternatively spliced region. *C*, RT-PCR on RNA of different zebra finch tissues spanning the alternatively spliced region, but not the entire ORF, yields DNA fragments of the expected sizes shown in *B*. *D*, Northern blot analysis of 20 μ g of total RNA from adult zebra finch brain and lung was performed with a 32 P-labeled DNA fragment spanning bp 114–959 (relative to the first start codon of isoform III). Ethidium bromide staining of 18S and 28S ribosomal bands demonstrates equal RNA loading. The different *zffoxp2* transcripts are indicated with arrows. *E*, Western blot analysis of 50 μ g of brain nuclear protein extract from a 40-d-old male zebra finch reveals a zffoxp2 protein corresponding in size to either isoform I or II, recognized by a polyclonal antibody raised against aa 613–715 of mouse Foxp2 (Lu et al., 2002).

human language (Enard et al., 2002). ZfFoxP2 lacks this human-specific amino acid change.

In addition, we cloned *zffoxp1*, the closest homolog to *zffoxp2*. With 5' and 3' RACE, we assembled 2412 bp of *zffoxp1* mRNA covering the ORF and 164 bp of the 3' UTR (GenBank accession number AY54952). ZfFoxP2 and zffoxp1 amino acid sequences are highly similar (supplementary Fig. 1*B*), with the biggest differences being that zffoxp1 misses the poly-glutamine stretch and 100 amino acids on the N terminus. For human FOXP1, an isoform that lacks the first 100 amino acids is reported (Banham et al., 1999), suggesting that we found a short zffoxp1 isoform. The high degree of similarity between zffoxp2 and zffoxp1 is consistent with their reported synergistic molecular function (Li et al., 2004).

Expression pattern of *zffoxp2*

In all brain regions that expressed *FoxP2* mRNA (ISH), the Foxp2-specific antibody (IHC) also strongly labeled nuclei, as is expected for a transcription factor.

Consistent with the reports from developing human and mouse brain (Shu et al., 2001; Lai et al., 2003; Takahashi et al., 2003), we detected *FoxP2* expression in the embryonic zebra finch brain as early as stage 26 (Hamburger and Hamilton, 1951;

Butler and Juurlink, 1987) (Fig. 2*A*). The highest expression was in the striatum and dorsal thalamus. This expression persisted throughout development (Fig. 2*B*) and was not restricted to vocal learners, because chickens also showed strong expression in the embryonic striatum (Fig. 2*C*). Closer examination at stage 34 revealed that the basal plate of the telencephalic vesicle, part of which gives rise to dorsal striatal areas in the adult, expressed *FoxP2* (Fig. 2*D*), as did the region that develops into the dorsal thalamus (data not shown). In the ventral midline of the mesencephalic vesicle, labeled cells appear to invade the laterally adjacent neuroepithelium (Fig. 2*E*). At limb levels of the spinal cord, cells that appear to be departing the roof plate and migrating to ventromedial regions expressed *FoxP2* (Fig. 2*F*). Expression was strong in the floor plate at this level, extending rostrally into the mesencephalon (Fig. 2*F*). The lateral margins of the hindbrain neuroepithelium and the region of the metencephalic/mesencephalic isthmus also strongly expressed *FoxP2*.

Throughout zebra finch post-hatch development and into adulthood the striatum and nuclei in posterior portions of the dorsal thalamus dominated expression (Fig. 3*A–F*). Expression levels in the striatum decreased slightly with age (Fig. 3*H*). Expression levels in pallial regions (i.e., those dorsal to the striatum) remained low throughout development and into adulthood (Fig.

3H). The prominent expression in the striatum and the dorsal thalamus was common to all birds, because this pattern was present in all species investigated and in both genders, regardless of whether they are vocal learners (Fig. 4A–G) or not (Fig. 4H), and even in a crocodile (Fig. 4I), the closest non-avian relative.

ZfFoxP2 expression in a striatal song nucleus during stages of vocal plasticity

In vocal learners, the dorsal striatum contains a nucleus that is part of the specialized song system, called Area X in songbirds, vocal nucleus of the anterior striatum [VAS; previously called VAP (Jarvis et al., 2000)] in hummingbirds, and magnocellular nucleus of the medial striatum [MMS; previously called LPOm (Striedter, 1994)] in parrots. This structure is part of a basal ganglia loop, the so-called anterior forebrain pathway (AFP) (Bottjer and Johnson, 1997; Durand et al., 1997; Farries and Perkel, 2002) and is essential for vocal learning (Sohrabji et al., 1990; Scharff and Nottebohm, 1991). During development, Area X in male zebra finches expressed more *FoxP2* mRNA than the surrounding striatum only at PHDs 35 and 50, the age at which zebra finches actively learn how to imitate song (Fig. 3C,D) (Tchernichovski et al., 2001). Before this period (at PHDs 15 and 25) and afterward, when birds crystallized their songs (PHD 75) and became adults (more than PHD 120), *FoxP2* expression in Area X did not differ from expression in the surrounding striatum. In adults, there was variability. Of 10 adult male zebra finches examined, 7 had expression levels in Area X similar to the region surrounding it, two slightly lower and one slightly higher. We could not find a variable (song complexity, amount of singing, or age at sacrifice) that could account for this adult variability. Female zebra finches, which do not learn song and do not have an Area X song nucleus, did not show any differential *FoxP2* expression in the region where Area X is located in males.

We also examined *FoxP2* expression in adult male canaries during different seasons of the year using a collection of canary brain sections described by Jarvis and Nottebohm (1997) (also see Materials and Methods). In July, August, and September, canaries expressed more *FoxP2* mRNA in Area X than in the region surrounding it (Fig. 5). These are the months when birds add new syllables into their song repertoire and song is more variable (Nottebohm et al., 1986; Leitner et al., 2001) than in the preceding breeding season, when song is stable. Breeding occurs in spring and can last through the end of June, and *FoxP2* expression during this time (sampled in April and May) did not differ from the surrounding region. This was also the case in October and January (Fig. 5).

To rule out that the developmental and seasonal changes in Area X *FoxP2* expression were the result of a generic feature of gene expression in this region, we compared the zebra finch glutamate receptor subunits *NR2B* and *mGluR2* (Wada et al., 2004) on adjacent sections to those that were probed with *FoxP2*. We found no differences in *mGluR2*. There were some developmental changes in *NR2B* expression in zebra finch Area X at PHD 25,

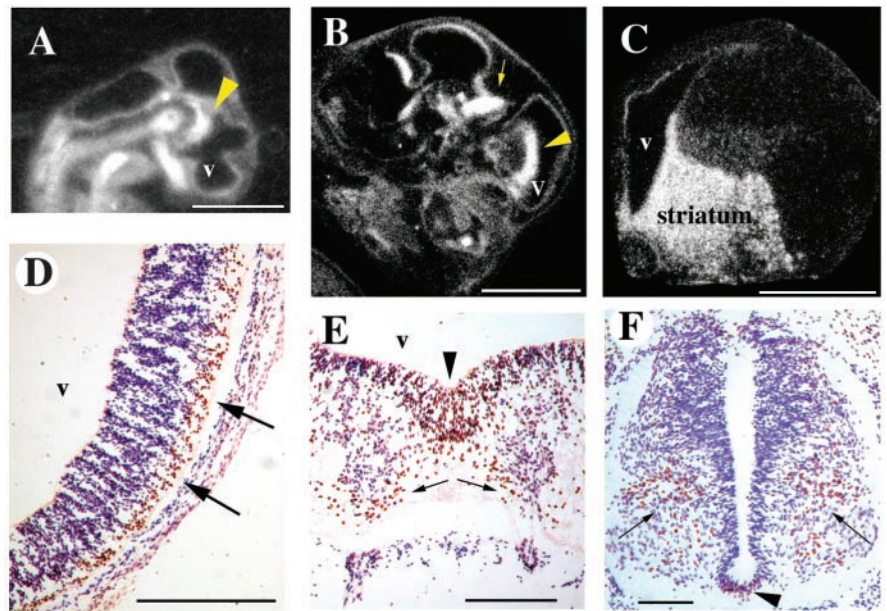


Figure 2. Embryonic *FoxP2* mRNA (A–C) and protein (D–F) expression. Sagittal sections through stage 26 (A) and 34 (B) zebra finch embryos show expression in presumptive striatum (arrowheads) and presumptive dorsal thalamus (arrow). The heads face toward the right. C, Embryonic chicken brain (embryonic day 13) had strong expression in the developing striatum and also in the pallial and subpallial germinal ventricular zone, shown in a frontal right hemisection. The *FoxP2* mRNA label appears white in dark-field illumination in A–C. D–F, *FoxP2* expression in a stage 26 zebra finch embryo frontal sections. FoxP2 immunoreactivity is brown, and cresyl violet-stained cells are purple/blue. D, A prominent band of FoxP2-positive cells is visible among cresyl violet-stained neurons in the ventrolateral telencephalic vesicle. E, The floor plate at the rostral end of the mesencephalic vesicle (arrowhead) has many FoxP2-expressing cells that seem to disperse laterally (arrows). F, At limb levels of the spinal cord, floor plate neurons expressed FoxP2 (arrowhead), as did a population of neurons in ventral cord (arrows). Scale bars: A–C, 2 mm; D–F, 100 μ m.

as expected from a previous report (Basham et al., 1999). However, the ratio of *NR2B* expression levels between Area X and the surrounding striatum remained similar at PHDs 35–75 (supplementary Fig. 2), the time when the *FoxP2* expression ratio was higher. In canaries, we observed no seasonal changes of *NR2B* expression in Area X, as was also shown previously (Singh et al., 2003).

We further tested whether some of the observed variability of *FoxP2* expression in Area X could be accounted for by singing activity. Singing strongly induces the expression of the immediate early gene *ZENK* (the avian homolog of mammalian *zif268/EGR-1/NGFI-A/krox24* gene) in Area X (Jarvis and Nottebohm, 1997). Moreover, the 5' flanking region of human *FOXP2* contains three predicted *EGR-1* (i.e., *ZENK*) binding sites (Bruce and Margolis, 2002). We found that for birds of similar age or season there were no significant differences in *FoxP2* mRNA expression between quiet control animals (quiet for at least 12 hr overnight) and animals that sang spontaneously [for 30 or 60 min for zebra finches ($n = 3$ each) and 1, 15, 30, or 60 min or 2, 4, or 6 hr for canaries ($n = 3$ each)], whereas *ZENK* was induced dramatically in zebra finches at PHD 65 or 150 by singing during the last 30 min before sacrifice (Jarvis and Nottebohm, 1997) (supplementary Fig. 3).

FoxP2 expression in Area X of adults of four other songbird species and in the corresponding region VAS in hummingbirds also differed relative to the surrounding striatum (Fig. 4). In chickadees and strawberry finches, both seasonal breeders (Langham, 1987; Smith, 1991), *FoxP2* expression was higher in Area X than in the surrounding striatum (Fig. 4A, B, J). In song sparrows and Bengalese finches, *FoxP2* expression was lower than the sur-

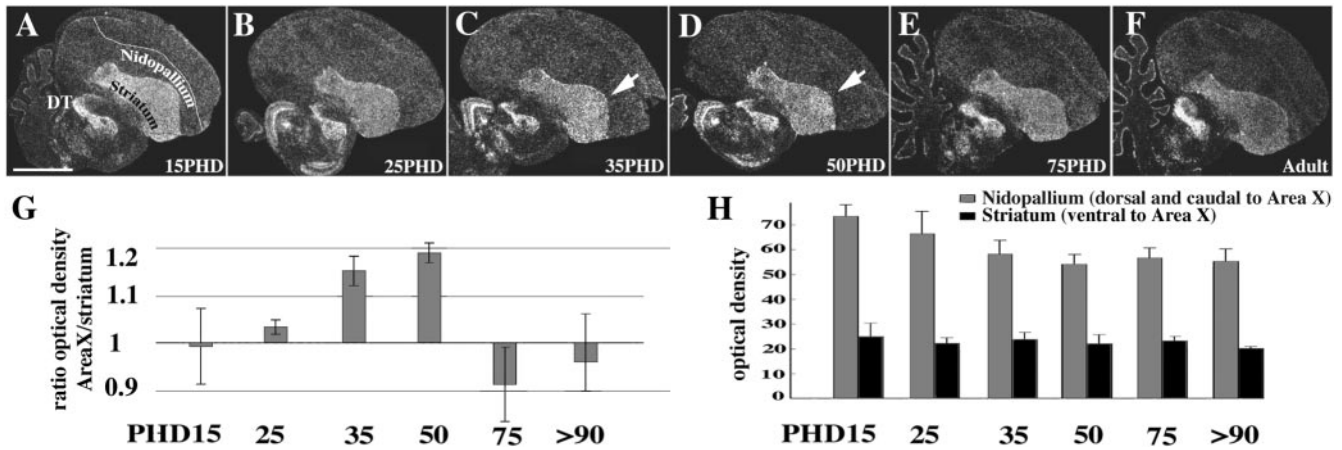


Figure 3. Differential *FoxP2* expression in Area X during post-hatch zebra finch development (A–F). Area X expressed more *zFoxP2* than the surrounding striatum only at PHDs 35 and 50 (C, D, arrowheads), which is the time when zebra finches learn to imitate song. G and H show the results of autoradiographic densitometric quantification of expression levels at the different ages ($n = 3$ for each age). The ratio of expression between Area X and the surrounding striatum increased during the phase when song imitation occurs on PHDs 35 and 50 (G). Absolute levels of *FoxP2* expression in the nidopallium did not change throughout development, whereas in the striatum (outside of Area X) they decreased slightly from PHDs 15 to 25 and reached adult levels by PHD 35 (H). Scale bar (in A): A–F, 2 mm.

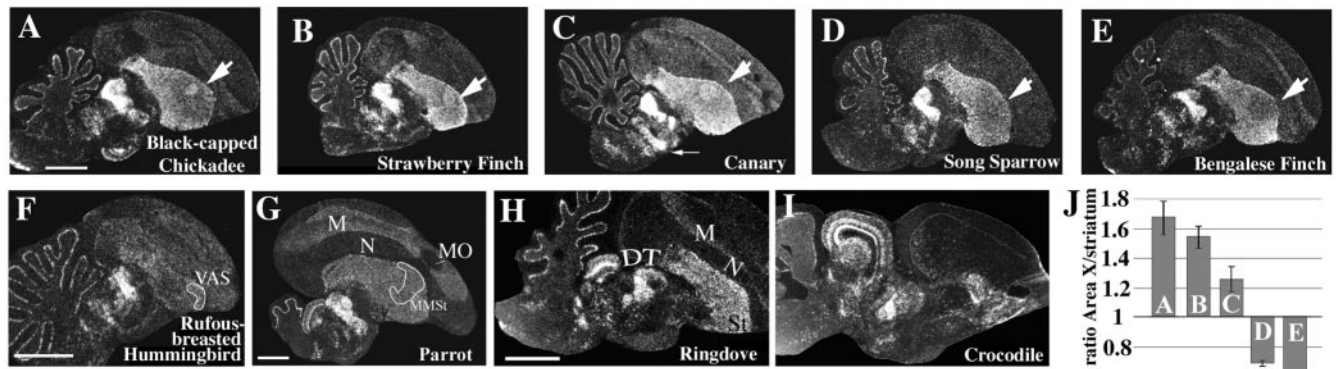


Figure 4. Different adult vocal learners (A–G), non-learners (H), and a crocodile (I) shared the *FoxP2* expression pattern in the striatum and dorsal thalamus (DT) but differed in expression levels in the striatal vocal nucleus (Area X/VAS/MMSt). Area X of chickadees (sampled in the fall), strawberry finches (sampled on long day photoperiod), and canaries (sampled in July) expressed more *FoxP2* in Area X than in the surrounding striatum (A–C), reflected in higher expression ratios (bars A–C in J). Area X of song sparrows (sampled in spring) expressed slightly less *FoxP2* than the surrounding striatum (D; bar D in J), as did Bengalese finch (E; bar E in J). The rufous-breasted hermit hummingbird (F) had slightly higher expression in the VAS, and the parrot (G) did not show a difference between vocal nucleus MMSt and the surrounding striatum. The adult ringdove (H), a bird that does not exhibit vocal learning and lacks telencephalic vocal nuclei, expressed high levels of *FoxP2* mRNA in the striatum and DT, as did a crocodile (I). The arrow in C points to the high levels of *FoxP2* expression in the substantia nigra pars compacta. M, Mesopallium; MO, oval nucleus of the mesopallium; N, nidopallium; St, striatum; VAS, vocal nucleus of the anterior striatum; MMSt, magnocellular nucleus of the medial striatum. Scale bars (in A for A–E; in H for H, I), 2 mm.

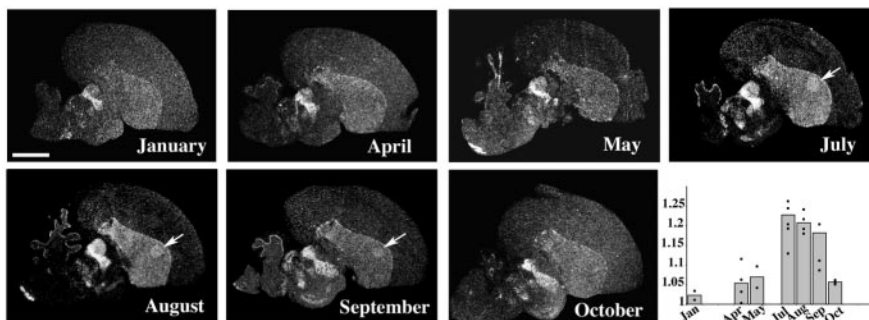


Figure 5. *FoxP2* expression in Area X of adult canaries varied seasonally. Area X expressed noticeably more *FoxP2* than the surrounding striatum only during the months of July, August, and September, resulting in higher ratios of Area X to striatum expression (the bar graph shows the mean ratios for each month, and superimposed points represent the values for individual birds).

rounding striatum (Fig. 4D,E,J). The chickadees were caught during the fall months (October and November), whereas the song sparrows were caught during late spring (April and May) (Jarvis et al., 1997), when song sparrows sing fewer variations of

song types and song is more stereotyped than in the fall (Smith et al., 1997). Bengalese finches are not strongly seasonal birds and breed opportunistically (Seiler et al., 1992), as do zebra finches, although the latter are also sensitive to photoperiod (Bentley et al., 2000). Rufous-breasted hummingbirds, captured near the end of their breeding season (Jarvis et al., 2000), showed slightly elevated levels of *FoxP2* in the hummingbird striatal vocal nucleus VAS (Fig. 4F). We did not find differential expression in MMSt of parrots (Fig. 4G). To address the source of the differences in *FoxP2* expression in Area X/VAS/MMSt among different species, we checked whether they might be related to differences in overall vocal syntax complexity, using the equations of Scharff and Nottebohm (1991). Scores of vocal syntax complexity are low when song elements are mostly rendered in an unvarying,

stereotyped manner. When songs consist of elements that are rendered in highly variable sequences, scores of syntax complexity are high. Vocal syntax complexity is low in strawberry finch, zebra finch, and somber hummingbird; intermediate in Bengalese finch, canary, and song sparrow; and high in rufous-breasted hermit hummingbird and budgerigar (K. Wada and E. D. Jarvis, unpublished observation). Thus, vocal syntax complexity cannot account for the observed *FoxP2* expression differences among the species (data not shown). Instead, the *FoxP2* expression pattern in chickadee, strawberry finch, and song sparrow are more consistent with the notion that during times of increased song stereotypy, as is usually observed during the breeding season, *FoxP2* is not upregulated in Area X, whereas outside of the breeding season, when song tends to be more plastic, *FoxP2* expression in Area X tends to be higher.

Hummingbirds and parrot differed with respect to pallial expression from the six songbird species investigated. In the hummingbird, the differential higher expression of *FoxP2* in the striatum relative to the pallium was less pronounced than in the other species. In the parrot, *FoxP2* expression in mesopallium was much higher relative to other pallial regions than it was in the other species tested. However, the AFP mesopallial song nucleus [MO; previously called HVo (Jarvis and Mello, 2000)] had low *FoxP2* expression (Fig. 4G). None of the other pallial vocal nuclei of the parrot, songbird, or hummingbird AFP (songbird IMAN like) or vocal nuclei of their motor pathways (songbird HVC like, used as a proper name, and RA like) expressed high levels of *FoxP2*.

Cellular identity of *zfFoxP2*-expressing cells

In adult zebra finch striatum, *Foxp2* immunoreactivity was characteristically seen in medium or small cells that were uniformly distributed throughout, except for one peculiarity. Small *Foxp2*-positive cells formed distinct, evenly spaced clusters in the part of the lateral striatum that abuts the pallial–subpallial lamina (PSL; previously called LMD), which separates the pallium from subpallium (Fig. 6A,B). More medially in the striatum these clusters formed a thin, continuous band (data not shown), matching the high levels of mRNA seen at the striatum side of the PSL (Fig. 3F). In pigeon striatum, similarly arranged patches contain dense ChAT-immunoreactive fibers (Medina and Reiner, 1994). In zebra finch, these *Foxp2*-immunoreactive cell clusters were, likewise, innervated by ChAT (Fig. 6C). The clusters were also visible in Nissl-stained material (Fig. 6D). A Hu antibody, which binds to an RNA-binding protein specifically present in young postmitotic and adult neurons (Barami et al., 1995), revealed that all *Foxp2*-immunoreactive brain cells were neurons, including the clusters at the PSL in the striatum (Fig. 6E,F). Some of the latter also expressed PSA-NCAM, a marker for cellular plasticity and migration (Durbec and Cremer, 2001) (Fig. 6G).

To determine whether the *FoxP2*-expressing neurons in the striatum belonged to a particular population of neurons, we used markers for the three classes of striatal interneurons (Reiner et al., 1998, 2004a) in conjunction with *Foxp2* IHC:ChAT to detect the large, aspiny cholinergic interneurons, nitric oxide synthase (nNOS) to detect the medium-sized aspiny interneurons that also contain somatostatin and NPY, and the calcium binding protein parvalbumin to detect another population of medium-sized aspiny interneurons that also contain GABA and the neurotensin-related hexapeptide LANT6 (Reiner et al., 1998, 2004a). Neither ChAT (Fig. 6J) nor nNOS (Fig. 6K) nor parvalbumin (Fig. 6L) were detected in the same neurons as *FoxP2*, suggesting that the striatal neurons that express *FoxP2* are pro-

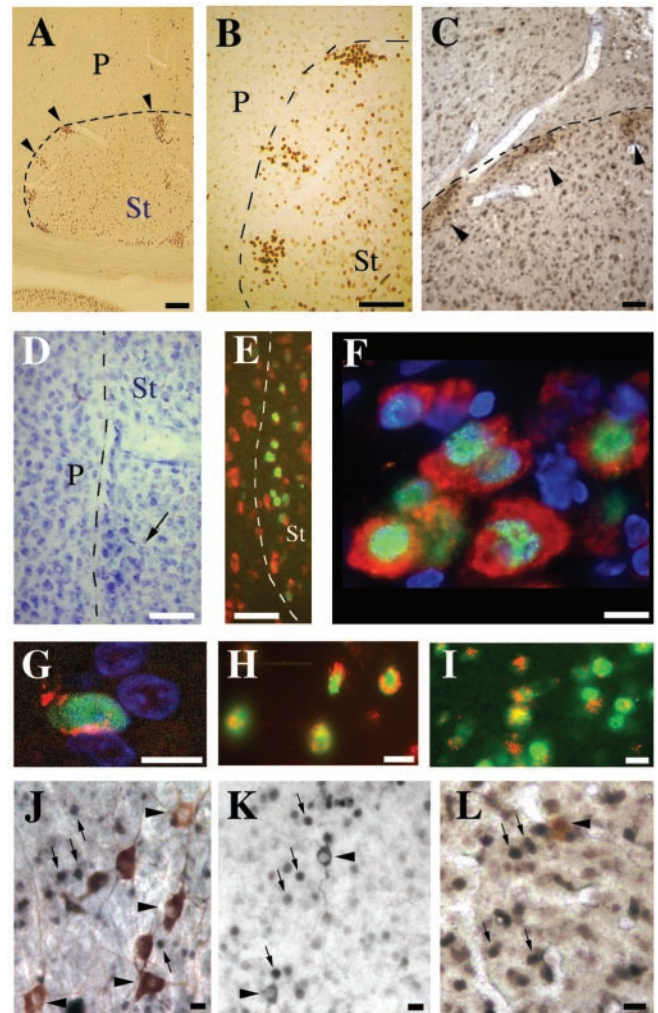


Figure 6. *FoxP2* expression in distinct populations of neurons in adult zebra finches. Low (*A*) and high (*B*) magnification of a sagittal section showing the dorsolateral extent of the subpallial–pallial (*P*) border with the striatum (*St*; black dashed line), where clusters of cells in the dorsal and lateral striatum express *FoxP2* (arrowheads; brown immunoreactivity). Dorsal is up, and rostral is to the right. *C*, These clusters (arrowheads; black-brown immunoreactivity) are characterized by dense ChAT fiber staining (lighter brown immunoreactivity). *D*, Clusters visualized with cresyl violet stain. *E*, *FoxP2*-immunoreactive cells within the clusters are neurons as shown by double labeling with fluorescent anti-Hu (red) and anti-*FoxP2* (green). *F*, Higher magnification in the dorsal thalamus shows that the cytoplasmic neuronal anti-Hu antibody (red) colocalizes with nuclear *FoxP2* antibody staining (green). *FoxP2*-negative nuclei can be seen in blue, stained with nuclear 4',6-diamidino-2-phenylindole DNA stain. *G*, Some *FoxP2*-positive cells are recognized by anti-PSA-NCAM antibody, a cell adhesion protein (PSA-NCAM, red; *FoxP2*, green; TOPRO3 nuclei, blue). *H*, Striatal neurons also coexpress DARPP-32 (red) and *FoxP2* (green) and appear to be innervated by TH-positive (red) terminals (*I*). Colabeling with neurochemical markers for three different striatal interneuron populations [ChAT (*J*), nNOS (*K*), or parvalbumin (*L*) (brown cytoplasmic labeling; arrowheads)] revealed that *FoxP2* (black nuclear labeling; arrows) was not expressed in these cell types. Scale bars: *A, B*, 100 μ m; *C–E*, 50 μ m; *F–L*, 10 μ m.

jection neurons rather than interneurons. It is known that the striatal neurons that project to the pallidum in birds, as in mammals, and striatal neurons that project to pallidal-like cells in Area X are the site of convergent nigral dopaminergic and cortical (i.e., pallial) glutamatergic input (Reiner et al., 1998, 2004a). DARPP-32 is thought to serve as a critical integrator of these two inputs onto the striatal projection neurons (Hemmings et al., 1995). Concordant with our expectation that *FoxP2* is expressed in striatal projection neurons, we found two indicators of dopa-

Table 1. Expression patterns of FoxP2 in subtelencephalic brain regions of adult zebra finch

Abbreviation	Subtelencephalic region	FoxP2
AN	Nucleus angularis	+
DM	Dorsomedial nucleus of the midbrain	–
DT	Dorsal thalamus (posterior nuclei)	+++
nIII	Cranial nucleus III (Edinger–Westphal)	–
Cn	Cuneate nucleus	+
Gn	Gracile nucleus	+
Gct	Substantia grisea centralis	+
Imc	Nucleus isthmi, pars magnocellularis	–
Ipc	Nucleus isthmi, pars parvocellularis	–
IO	Nucleus isthmo-opticus	+
La	Nucleus lateralis anterior thalami	+
LLi	Nucleus lemnisci lateralis	+
MC	Nucleus magnocellularis	+
MLd	Nucleus mesencephalicus lateralis, pars dorsalis	++
MnV	Motor part of trigeminal nucleus or V nucleus	–
MnX	Dorsal motor part of the vagus nucleus or X nucleus	–
nBOR	Nucleus of the basal optic root	–
nIX	Glossopharyngeal nucleus or IX nucleus	–
nVI	Abducens nucleus or VI nucleus	–
nXII	Hypoglossal nucleus or XII nucleus	–
OI	Nucleus olivaris inferior	+++
Omd	Nucleus nervi oculomotorii, pars dorsalis	–
OMdm,OMv	Nucleus nervi oculomotorii, pars dorsalis/ventralis	–
OMv	Nucleus nervi oculomotorii, pars ventralis	–
Ov	Nucleus ovoidalis	+
PAG	Periaqueductal gray	+
PMH	Nucleus medialis hypothalami posterior	+
PT	Pretectal nucleus	–
PTD	Nucleus pretectalis diffusus	–
PTM	Nucleus pretectalis medialis	++
PVN	Paraventricular nucleus	–
Rt	Nucleus rotundus	++
RPC	Nucleus reticularis pontis caudalis	+
R	Red nucleus	+
ST	Nucleus of the solitary tract	+
SNc	Substantia nigra, pars	+
SP	Nucleus subpretectalis	–
SPL	Nucleus spiriformis lateralis	–
T	Nucleus triangularis	++
VeD	Nucleus vestibularis descendens	+
VeL	Nucleus vestibularis lateralis	+
VTA	Ventral tegmental area	+

minergic innervation. Foxp2-immunoreactive striatal neurons coexpressed DARPP-32 (Fig. 6H), which is indicative of the presence of dopamine D1 receptors (Snyder et al., 1998), and immunoreactivity for TH, the synthetic enzyme for biogenic amines, was present in fibers around perikarya of neurons with Foxp2-immunoreactive nuclei (Fig. 6I).

ZfFoxP2 expression in subtelencephalic brain regions

Table 1 lists subtelencephalic structures that did or did not express FoxP2. For identification of subtelencephalic brain regions, we analyzed serial frontal and sagittal sections through the entire brain of male zebra finches and used the region-specific parvalbumin (Braun et al., 1985, 1991; Wild et al., 2001) and ChAT (Medina and Reiner, 1994) staining in adjacent series of sections as landmarks to ascertain the identity of brain regions that expressed FoxP2 (Fig. 7B, C, H, I). FoxP2 expression was prominent in dopaminergic midbrain regions that project to the basal ganglia, the substantia nigra/ventral tegmental area (Fig. 4C, small

white arrow), and in caudal regions of the dorsal thalamus (Figs. 3A–F, 4A–I, 5). In addition, FoxP2 was expressed in many regions that are involved in relaying and integrating ascending sensory information, including auditory regions [e.g., midbrain nucleus MLd (dorsal part of the lateral mesencephalic nucleus) (Fig. 7A, B) and thalamic nucleus ovoidalis (data not shown)], visual regions [e.g., afferent upper layers of midbrain optic tectum (Fig. 7A, F) and thalamic nucleus rotundus (Fig. 7D)], multimodal regions [e.g., layers 10 and 11 of the optic tectum (Fig. 7F)], and somatosensory regions [e.g., sensory trigeminal (data not shown)]. Prominent FoxP2 expression was observed in the Purkinje cells of the cerebellum (Figs. 3E, F, 4A–I, 7E) and the inferior olive (Fig. 7G), which gives rise to all the climbing fibers innervating the Purkinje cells. All species tested, including males and females, regardless of whether they learn their vocalization or not, expressed FoxP2 in these regions. In contrast, FoxP2 expression was not found in midbrain and brainstem motor control areas, such as the vocal nucleus DM (dorsomedial motor nucleus of the intercollicular region) (Fig. 7B, C), the hypoglossal vocal and tongue nucleus, nXII (Fig. 7H, I), and most other motor cranial motor nuclei (Table 1).

ZfFoxP1 expression

Similar to FoxP2, FoxP1 was expressed at high levels in the striatum and in the dorsal thalamus of zebra finches and other birds (Fig. 8A–F). Unlike FoxP2, FoxP1 expression in the striatal vocal nuclei (Area X or MMSt) was similar across development and season, across all songbirds tested, and in parrots (i.e., higher expression in the striatal vocal nucleus relative to the immediate surrounding striatum) (Fig. 8A–D, F). Also unlike FoxP2, within the pallium, FoxP1 was consistently and prominently expressed in the mesopallium in all avian species tested (Fig. 8A–F). Interestingly, for the three main songbird pallial vocal nuclei (IMAN, HVC, and RA), FoxP1 expression differed notably from the expression of the subdivisions in which these nuclei are embedded. HVC and RA strongly expressed FoxP1, whereas the surrounding territories did not. The reverse was true for IMAN, which did not express FoxP1, while the region around it did (Fig. 8A–D). This was consistent across songbird species. The parrot pallial analog of HVC, the central nucleus of the nidopallium, had noticeably higher levels than the surrounding nidopallium (Fig. 8F). In contrast to FoxP2, FoxP1 was never expressed in the Purkinje cells of the cerebellum. FoxP1 expression in the ring dove brain was similar to that of the songbirds and parrot, with the exception that there was no differential expression in the striatum and pallium, where vocal nuclei are found in vocal learners (Fig. 8E). A telencephalic expression pattern remarkably similar to that of the avian brain was found in crocodile (Fig. 8G), including high expression in striatal-like and mesopallium-like regions. This suggests that the general FoxP1 and FoxP2 expression patterns in vocally learning and non-learning birds were inherited from their common reptilian ancestor.

Discussion

We set out to answer three questions: (1) Is FoxP2 differentially expressed in the brains of avian vocal learners and non-learners? (2) How does FoxP2 expression in birds compare with FoxP2 expression in mammals? and (3) Does zebra finch FoxP2 bear molecular similarities to human FOXP2?

The answer to the first question is partly yes, partly no. Partly yes, because FoxP2 is differentially expressed in vocal nucleus Area X, a part of the special basal ganglia forebrain network required for vocal learning that vocal non-learners do not possess.

Moreover, *FoxP2* expression is elevated in Area X at the time when young zebra finches learn to imitate song and during the time when adult canaries remodel their songs. In addition, in adults of six different species, Area X (and in the equivalent structure VAS in the hummingbird) showed consistent differences in *FoxP2* expression, being either higher or lower than the surrounding striatum, in a pattern consistent with periods of change in vocal behavior. Lesions of Area X in zebra finches during vocal learning result in adult song production that is more plastic than when Area X is intact (Sohrabji et al., 1990; Scharff and Nottebohm, 1991), suggesting that Area X helps generate song stability. If *FoxP2* acts as a transcriptional repressor in the brain, as it does in the lung (Shu et al., 2001; Lu et al., 2002; Li et al., 2004), then the higher levels found during periods of vocal plasticity might suggest that *FoxP2* represses genes that are involved in neural stability in Area X. These findings and interpretation are compatible with a role for *FoxP2* in learned vocalization.

The answer to question (1) is partly no, because *FoxP2* was expressed in non-vocal striatal regions outside of Area X/VAS/MMSt of all 11 bird species examined and was similar regardless of whether or not they learn their vocalizations. Both vocal learners and vocal non-learners had similar developmental onset of *FoxP2* expression in comparable brain regions and equivalent expression patterns in adults. The strongest signal was consistently observed in the striatum of the basal ganglia, nuclei of the dorsal thalamus and mid-brain, the inferior olive, and the Purkinje cells of the cerebellum. Less intense, but consistent, expression was observed in various nuclei connected to these regions.

Our second question concerned the comparison of our data with those found in mammals. Recent publications of *FOXP2* expression in the human, rat, and mouse brain (Ferland et al., 2003; Lai et al., 2003; Takahashi et al., 2003) report essentially concordant striatal and subtelencephalic sites of *FoxP2* expression with those we and another group (Teramitsu et al., 2004) found in birds. We also noted a similar striatal expression pattern in a closely related reptilian species to birds, the crocodile. Birds expressed little *FoxP2* in pallial regions, with the exception of the mesopallium, and this varied between species. Mammals also expressed little *FoxP2* in pallial regions, with the exception of cortical layer 6 (Ferland et al., 2003). The expression of *FOXP1* in mammals (Ferland et al., 2003) is also concordant with the expression we and Teramitsu et al. (2004) found in birds for striatal and subtelencephalic sites. In pallial regions, relative to *FoxP2*, birds expressed more widespread *FoxP1*, with the highest expression occurring in the mesopallium and in vocal nuclei HVC

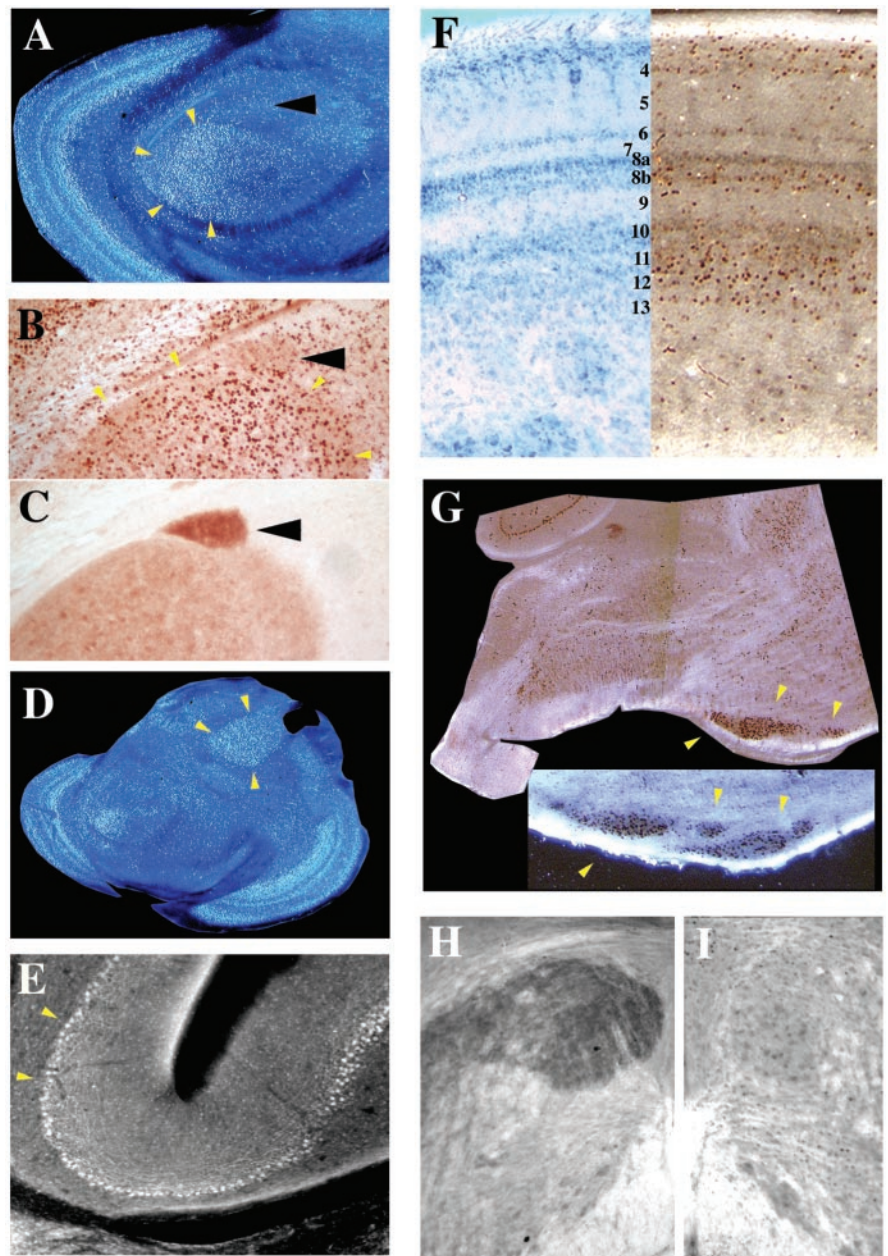


Figure 7. *FoxP2* expression in subtelencephalic regions was associated more with afferent sensory or multimodal areas rather than with pure motor areas. Auditory nucleus MLD (dorsal part of the lateral mesencephalic nucleus) expressed *FoxP2* (white dark-field label in *A* and brown label in *B*; both surrounded by yellow arrowheads). In contrast, the dorsomedial motor nucleus of the intercollicular region (DM), which controls vocalizations, showed little mRNA and immunoreactivity for *FoxP2* (*A*, *B*, black arrowheads) but strong parvalbumin immunoreactivity (*C*) (Braun et al., 1985). Also, *FoxP2*-immunoreactive cells were seen in the visual thalamic nucleus rotundus (*D*), cerebellar Purkinje cells (*E*), specific layers of the optic tectum in the midbrain (*F*), and brainstem nucleus inferior olive (*G*) but not in the tracheosyringeal portion of the nucleus of the hypoglossal nerve nXII (*I*). We took advantage of the strong parvalbumin immunoreactivity of nXII to unambiguously identify this nucleus (adjacent section to *I* stained with parvalbumin in *H*) (Wild et al., 2001). Immunoreactivity in dark-field images appears white, and in bright-field photomicrographs appears brown. *A*, *D*, and *E–G* are sagittal sections, rostral is to the right, and *B*, *C*, *H*, and *I* are frontal sections. Dorsal is up in both orientations.

and RA of songbirds, but notably low levels in the tissue surrounding HVC and RA. Mammals also expressed more widespread *FoxP1* levels in the pallium, cortical layers 3–5 during development, and also in layer 6 during adulthood (Ferland et al., 2003). These differences in cortical/pallial *FoxP2* expression between mammals and birds are difficult to interpret because direct homologies between most avian and mammalian pallial areas remain unresolved (Reiner

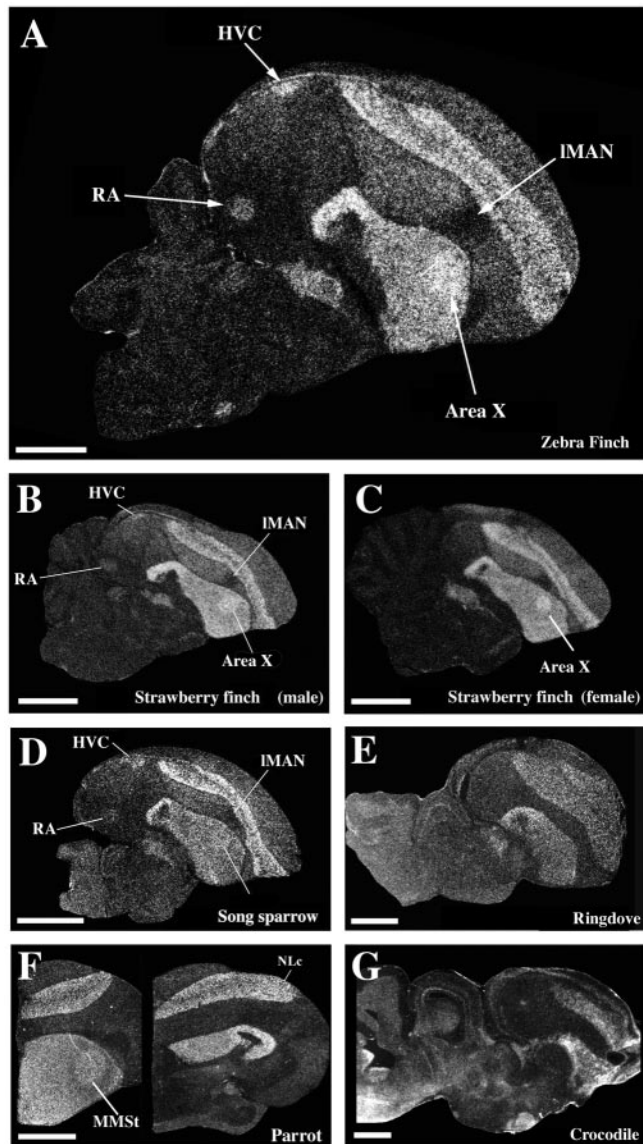


Figure 8. Expression pattern of *FoxP1* was distinct from but partially overlapping with that of *FoxP2*. *A*, *FoxP1*, like *FoxP2*, was expressed in the dorsal thalamus and striatum in adult zebra finches (*A*). In addition, it was expressed in vocal nuclei HVC, RA, and Area X (but not IMAN) at higher levels than their surrounding regions and in the mesopallium. Both male (*B*) and female (*C*) strawberry finches, male song sparrow (*D*), as well as the parrot (*F*) expressed more *FoxP1* mRNA in Area X (MMSt in parrot) than in the surrounding striatum. *E*, A vocal non-learner, the ring dove, also expressed *FoxP1* mRNA in the subpallial and pallial areas. *G*, The crocodile had a telencephalic pattern very similar to that of birds. All sections are sagittal, except the parrot sections in *F*, which are frontal. Scale bars: *A–D*, 1 mm; *E–G*, 2 mm.

et al., 2004b). The pallium of the avian telencephalon possesses a nuclear organization, whereas that in mammals shows a layered organization. There has been no predicted relationship between the avian mesopallium and cortical layer 6 that we are aware of (<http://avianbrain.org>). The main projection of the mesopallial vocal nuclei and other mesopallial areas (previously called hyperstriatum ventrale) are to arcopallial, nidopallial, and striatal areas (Durand et al., 1997; Csillag 1999; Brauth et al., 2001), whereas those of layer 6 in mammals are to the dorsal thalamus in addition to other cortical (pallial) layers (Ferland et al., 2003). Thus, our results suggest that high striatal and subtelencephalic *FoxP1* and *FoxP2* expression and restricted pallial expression in birds and mammals was inherited from

a common stem-amniote ancestor (Evans, 2000), but that their specific pallial patterns have either diverged or evolved independently.

We also answered question (3), whether *FoxP2* might be molecularly more similar to human *FOXP2* than to rodent *Foxp2*. We were motivated to ask this rather unlikely question because of recent reports (Enard et al., 2002; Zhang et al., 2002) that *FOXP2* has been the target of positive selection during recent primate evolution, resulting in a human-specific amino acid change that is thought to be related to language (i.e., human vocal learning). If this change was pivotal to the evolution of learned vocal communication in hominids, perhaps similar selection pressure acted on the supposedly independent evolution of vocal learning in birds. Although *zfFoxP2* clearly lacks the human-specific amino acid change, we cannot exclude this possibility. The fact that zebra finches, in contrast to mouse, have a 6.5 kb *zfFoxP2* transcript that corresponds in size to the human transcript raises the possibility that selection acted on the regulatory sequences.

Taken together, we conclude that the striking conservation of the *FoxP2* gene sequence and overall brain expression pattern in avian, reptilian, and mammalian brains, regardless of whether they learn to vocalize or not, confirms that *FoxP2* has a more general role than to enable vocal learning. *FoxP2* could be an ancient transcription factor involved in shaping cerebral architecture, perhaps via restriction of certain neuronal lineages, as reported recently for *Foxg1* (Hanashima et al., 2004). If *FoxP2* were involved in the development and maintenance and function of subtelencephalic and striatal sensory and sensory-motor circuits, this could create a permissive environment on which vocal learning can evolve if other factors come into play. Given the prominent role of many other forkhead transcription factors in early development, this is a likely scenario (Carlsson and Mahlapuu, 2002). Support of this notion also stems from the fact that regions of early *FoxP2* expression in the avian embryo are sources of inductive signals that organize adjacent neuroepithelium and neuronal migration during early development.

The common expression pattern of *FoxP2* in birds and mammals might provide valuable clues about what constitutes a “permissive environment” for vocal communication and evolution of vocal learning. Learning to imitate acoustic signals requires integration of sensory information with the desired motor output. The basal ganglia as well as the cerebellum in all vertebrates integrate afferent sensory information with descending motor commands and thus participate in the precise control of temporally sequenced muscle movements (Doyon et al., 2003). Both innate and learned avian and human vocalizations depend on such control (Doupe and Kuhl, 1999). Anatomical evidence suggests that the specialized regions for vocal learning in birds were elaborated from already modularly connected forebrain regions translating ascending auditory, somatosensory, and visual information into motor commands. Consistent with this, an AFP-like circuit apparently also exists in vocal non-learners (Farries, 2001). In humans, the basal ganglia and the cerebellum have attracted far less attention than the cortical speech and language areas, but there is increasing awareness that the basal ganglia and cerebellum are not only essential for the execution but might also be required for the acquisition of human vocal behavior (Lieberman, 2001; Marien et al., 2001).

Besides having the advantage of being able to study anatomical structures involved with learned vocal communication, our study also differs from the rodent and human studies in that we specifically investigated the distribution of *FoxP2* in cranial motor nuclei. It has been suggested that the speech and language pathology in humans with *FOXP2* mutations consists of an oro-

facial dyspraxia core deficit (for review, see Marcus and Fisher, 2003). This could be primarily attributable to a lack of muscle control over the speech apparatus. However, our data suggest that in birds *FoxP2* is, for the most part, expressed in afferent sensory pathways and in the striatal projection neurons, which are the site of convergence for both pallial and subpallial projections. Takahashi et al. (2003) also argue that in rats *FoxP2*-positive striatal neurons are projection neurons, based on selective *FoxP2* expression in the lateral ganglionic eminence but not the medial ganglionic eminence and the known precursor populations of these germinal regions (Parnavelas et al., 2002). In addition, many sites of *FoxP2* expression, such as the inferior olive–Purkinje cell pathway, the optic tectum, and the striatum, are known substrates for experience-dependent plasticity (Krupa and Thompson, 1997; Hyde and Knudsen, 2000; Doyon et al., 2003). This highlights the need for more studies investigating the role of ascending visual, auditory, and somatosensory information in complex learned motor skills such as birdsong and human speech.

In summary, our findings underscore the usefulness of the songbird system to study a gene implicated in human speech and language. *FoxP2* has a characteristic expression pattern in a brain structure uniquely associated with learned vocal communication, Area X in songbirds. In the rest of the brain of birds that learn to sing and in birds that do not, *FoxP2* expression predominates in sensory and sensory–motor circuits. These latter regions also express *FoxP2* in mammals and reptiles. We conclude that *FoxP2* may be important for establishing and maintaining brain pathways including, but not limited to, those essential for learned vocal communication.

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