**Drosophila FoxP Mutants Are Deficient in Operant Self-Learning**

Ezequiel Mendoza¹, Julien Colomb², Jürgen Rybak³, Hans-Joachim Pflüger², Troy Zars⁴, Constance Scharff¹, Björn Brembs².⁵,⁶

¹ Inst. Biol. – Behavioral Biology, Freie Universität Berlin, Berlin, Germany; ² Inst. Biol. – Neurobiology, Freie Universität Berlin, Berlin, Germany; ³ Neuroethology, Max Planck Institute for Chemical Ecology, Jena, Germany; ⁴ Biological Sciences, University of Missouri, Columbia, Missouri, United States of America; ⁵ Institut für Zoologie - Neurogenetik, Universität Regensburg, Regensburg, Germany

**Abstract**

Intact function of the Forkhead Box P2 (FOXP2) gene is necessary for normal development of speech and language. This important role has recently been extended, first to other forms of vocal learning in animals and then also to other forms of motor learning. The homology in structure and in function among the FoxP gene members raises the possibility that the ancestral FoxP gene may have evolved as a crucial component of the neural circuitry mediating motor learning. Here we report that genetic manipulations of the single *Drosophila* orthologue, dFoxP, disrupt operant self-learning, a form of motor learning sharing several conceptually analogous features with language acquisition. Structural alterations of the dFoxP locus uncovered the role of dFoxP in operant self-learning and habit formation, as well as the dispensability of dFoxP for operant world-learning, in which no motor learning occurs. These manipulations also led to subtle alterations in the brain anatomy, including a reduced volume of the optic glomeruli. RNAi-mediated interference with dFoxP expression levels copied the behavioral phenotype of the mutant flies, even in the absence of mRNA degradation. Our results provide evidence that motor learning and language acquisition share a common ancestral trait still present in extant invertebrates, manifest in operant self-learning. This ‘deep’ homology probably traces back to before the split between vertebrate and invertebrate animals.

**Introduction**

The Forkhead Box P2 (FOXP2) transcription factor is the first gene discovered to be involved in the development of speech and language [1,2]. The gene reveals signs of recent selection in the hominin lineage [2,3] and natural variation in the FOXP2 gene has been found to alter grey matter concentrations in patients with schizophrenia [4]. In fact, different polymorphisms have been found to dissociate between autism spectrum disorders on one side and schizophrenia on the other [5–9] but see also [10]. The FoxP2 gene sequence is highly conserved in vertebrates, its expression is largely concordant and numerous experiments indicate that FoxP2 is important for modulating the neural circuits involved in vocal learning [11–14]. For instance, RNAi-mediated knock-down of FoxP2 gene expression in zebra finch area X, a basal ganglia structure necessary for song learning, results in an incomplete and inaccurate imitation of tutor song [13]. Recent reports broaden the functional role of FoxP genes among vertebrates to also cover other forms of motor learning and other paralogues, particularly FoxP1 [11,12,15–22]. These findings together suggest a role of FoxP genes primarily in the acquisition of skilled coordination of the movements that underlie effective vocal communication and other movements (‘motor learning hypothesis’)[11–13,16–19,23–26]. Both paralogues’ importance for motor learning indicates that the single ancestral gene may once have provided this function alone.

Learning to produce vocalizations by imitation is a form of motor learning that proceeds slowly from ‘babbling’ (in humans) and ‘subsong’ (in birds) towards full-fledged language and ‘crystallized’ song. This type of motor learning has been classified as a form of operant learning [27–29]. That is, first exploratory, highly variable actions are initiated (i.e., babbling or subsong) and then sensory feedback shapes the initiation of future behavior, reducing its variability (i.e., crystallized song, speech) [14,30]. A particular variant of operant learning in tethered flying *Drosophila* [31,32] parallels some features of vocal learning: the animal first initiates highly variable, exploratory actions (i.e., turning maneuvers to the left and to the right), then sensory feedback shapes the initiation of future behavior, reducing its variability (i.e., turning maneuvers only to one side, see Fig. 1). We tested the hypothesis that operant learning may share homologous aspects with vocal learning using the dFoxP gene. Our evidence suggests that the *Drosophila* FoxP2 orthologue, dFoxP, plays a critical role in operant self-learning, a form of motor learning that shares conceptually analogous features with vocal learning.

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*Email: bjoem@brembs.net

† These authors contributed equally to this work.
Cloning of dFoxP isoforms and PCR

Primers where designed using the software Primer3 0.4.0 (http://frodo.wi.mit.edu/). The first set of primers was designed as follows: FoxPlisoMor (5'-AGTATTCGAGGATGCGACAAG-3') and FoxPlisoArev (5'-CAAAACGGAAGGATTTTTGGA-3'), set on the gene CG16899, which amplified a 1347 bp product of Isomform A (NCBI acc. # JN160729) and a 1718 bp product of an intron- retention isoform (NCBI acc. # JN160730) of the FoxP gene. For sequencing exon 8 in the different strains, which is the one that changes between Isomform A and Isomform B of the FoxP gene of Drosophila (designated as CG32937 in the Flybase), we used: FoxPlisoFor (5'-AGAATTCGAGGATCTGTCGAC-3') and FoxPlisoBrev (5'-TATAATTTCCGAAATCAGGACC-3'), which amplified a 532 bp fragment of the last exon (NCBI acc. # JN160731 and NCBI acc. # KF192848-KF192876). For all PCRs we used 1 µl of un-diluted cDNA of each strain. We ran a gradient PCR (50–65°C) with wild type Drosophila cDNA to determine the optimal annealing temperature. The PCR conditions were 94°C for 5 min, denaturation at 94°C for 5 s, annealing for 30 s at 60°C for Isomform A and 56.7°C for Isomform B, elongation for 2 min for Isomform A and 40 s for Isomform B at 72°C, 35 cycles, a last elongation at 72°C for 10 minutes. With this information we designed primers to amplify the full length FoxP isoforms of Drosophila melanogaster adding a restriction site for BamHI in the forward primer followed by the start sequence and FLAG-tag at the C'-terminal part of the protein followed by a Stop codon and EcoRI restriction site. We used the same forward primer for all isoforms, since the first six exons are shared. The forward primer we used was: sFOXPdm_for (5'-GGATCCTCGGCAATTCTGATCGATCGGATTGCAATCCGAGGATGCGACAAG-3') which amplified a 1347 bp fragment (NCBI acc. # KF206330). For the intron retention isoform we used the eFOXPr99_rev (5'-GGCGAATTCCTAATTTACGTGCTGCTATCTCTGTTTGACGCACATACCACCG-3') which amplified a 1374 bp fragment (NCBI acc. # KF206331). For all PCRs we used 1 µl of un-diluted cDNA of heads of Canton S wild type strain. For amplification of the full Open reading frame of the FoxP isoforms we used the High Fidelity PCR Enzyme Mix (#K0192; Lot 00116896). We used 5 µl of un-diluted cDNA in 50 µl final volume. We employed a HLA-PCR with the conditions 96°C for 1 min; followed by 5 cycles of 96°C for 25 seconds, 65°C for 45 seconds and 72°C for Immitude; followed by 25 cycles of 96°C for 25 seconds, 60°C for 45 seconds and 72°C for 1 minute; followed by 6 cycles of 96°C for 25 seconds, 55°C for 1 minute and 72°C for 2 minutes and a last cycle at 72°C for 10

mRNA extraction

Around 50 heads of the different fly strains were placed separately in 1.5 ml Eppendorf tubes, frozen in liquid nitrogen and stored at −80°C until mRNA extraction. Tissue was homogenized in 200 µl of TRIZOL reagent (Invitrogen) using a pellet pestle tip attached to a pellet pestle motor (Sigma Z359947 – 100EA; Z359971 – 1EA) and mRNA extraction was performed according to the manufacturer’s instructions. Residual DNA was then degraded using Turbo DNA – free kit (Ambion) following the manufacturer’s instructions. mRNA yield was determined by UV spectroscopy at 260/280 nm with Nanodrop (PEQLab Nanodrop, Spectrophotometer ND-1000). For cDNA synthesis, the same quantities of mRNA (1200 ng) were taken and Super Script III Reverse Transcriptase (Invitrogen, 18090-099) using Random Hexamers primers were used for first strand synthesis following the manufacturer’s instructions. As a negative control a minus Reverse transcriptase reaction was also run with the same quantity of mRNA. For QPCR the cDNA was diluted 1:10 in Nuclease free water (Ambion, AM9932).

Materials and Methods

Fly strains

All flies were kept on standard cornmeal/molasses medium [33] at 25°C and 60% humidity with a 12 hr light/12 hr dark regime [34]. FlyBase (http://flybase.org) insertion lines [P(RS5)FoxP]-3.95kb, FlyBaseID: FBti0030257; PBac[WH]FoxP[307346], FlyBaseID: Fbit0031415; PBac[PB]c03619, FlyBaseID: Fbit0044630) were crossed into a wild type strain. For amplification of the full Open reading frame of the FoxP gene of Drosophila (designated as CG32937 in the Flybase), we used: FoxPlisoFor (5'-AGAATTCGAGGATCTGTCGAC-3') and FoxPlisoBrev (5'-TATAATTTCCGAAATCAGGACC-3') which amplified a 1347 bp fragment (NCBI acc. # JN160729) and a 1718 bp product of an intron- retention isoform (NCBI acc. # JN160730) of the FoxP gene. For sequencing exon 8 in the different strains, which is the one that changes between Isomform A and Isomform B of the FoxP gene of Drosophila melanogaster adding a restriction site for BamHI in the forward primer followed by the start sequence and FLAG-tag at the C'-terminal part of the protein followed by a Stop codon and EcoRI restriction site. We used the same forward primer for all isoforms, since the first six exons are shared. The forward primer we used was: sFOXPdm_for (5'-GGATCCTCGGCAATTCTGATCGATCGGATTGCAATCCGAGGATGCGACAAG-3') which amplified a 1347 bp fragment (NCBI acc. # KF206330). For the intron retention isoform we used the eFOXPr99_rev (5'-GGCGAATTCCTAATTTACGTGCTGCTATCTCTGTTTGACGCACATACCACCG-3') which amplified a 1374 bp fragment (NCBI acc. # KF206331). For all PCRs we used 1 µl of un-diluted cDNA of each strain. We ran a gradient PCR (50–65°C) with wild type Drosophila cDNA to determine the optimal annealing temperature. The PCR conditions were 94°C for 5 min, denaturation at 94°C for 5 s, annealing for 30 s at 60°C for Isomform A and 56.7°C for Isomform B, elongation for 2 min for Isomform A and 40 s for Isomform B at 72°C, 35 cycles, a last elongation at 72°C for 10 minutes. With this information we designed primers to amplify the full length FoxP isoforms of Drosophila melanogaster adding a restriction site for BamHI in the forward primer followed by the start sequence and FLAG-tag at the C'-terminal part of the protein followed by a Stop codon and EcoRI restriction site. We used the same forward primer for all isoforms, since the first six exons are shared. The forward primer we used was: sFOXPdm_for (5'-GGATCCTCGGCAATTCTGATCGATCGGATTGCAATCCGAGGATGCGACAAG-3') which amplified a 1347 bp fragment (NCBI acc. # KF206330). For the intron retention isoform we used the eFOXPr99_rev (5'-GGCGAATTCCTAATTTACGTGCTGCTATCTCTGTTTGACGCACATACCACCG-3') which amplified a 1374 bp fragment (NCBI acc. # KF206331). For all PCRs we used 1 µl of un-diluted cDNA of heads of Canton S wild type strain. For amplification of the full Open reading frame of the FoxP isoforms we used the High Fidelity PCR Enzyme Mix (#K0192; Lot 00116896). We used 5 µl of un-diluted cDNA in 50 µl final volume. We employed a HLA- PCR with the conditions 96°C for 1 min; followed by 5 cycles of 96°C for 25 seconds, 65°C for 45 seconds and 72°C for Immitude; followed by 25 cycles of 96°C for 25 seconds, 60°C for 45 seconds and 72°C for 1 minute; followed by 6 cycles of 96°C for 25 seconds, 55°C for 1 minute and 72°C for 2 minutes and a last cycle at 72°C for 10
minutes, PCR products were examined in 0.5% EtBr agarose gels, the specific bands were cut from the gel, and purified using QIAquick Gel Extraction Kit (Qiagen, Cat. 28706). PCR products were then cut using Fast digest BamHI and EcoRI enzymes (Fermentas) and cloned into µDINAS.1 (→ vector (Invitrogen) cut with the same enzymes. These plasmids were transformed into One Shot Top 10 Escherichia coli chemically competent cells (Invitrogen, C404010) and colonies with ampicillin (100 µg/ml) resistance were selected on agarose plates. Clones with the specific insert were picked and grown overnight, at 37°C in 3 ml LB/ampicillin medium in a shaker. Plasmids were purified using Invisorb Spin Plasmid Mini Two columns (Invitek, Ref 1010140300) as described by the manufacturer. Finally, the inserts were fully sequenced and analyzed.

Sequencing of premature stop in 3955 strain

We used the FoxPIsoBf6or primer (described above) and a reverse primer in the RS P element (SPI 5′-CA-CAACCTTTGCTCTTCAACAAG-3′) to amplify a fragment of 440 bp. We sampled 2 µl of undiluted cDNA of the FoxP9535 strain. The PCR conditions were 94°C for 5 min, denaturation at 94°C for 5 s, annealing for 30 s at 55°C; elongation for 3 min at 72°C, 35 cycles, and a last elongation at 72°C for 10 minutes. The PCR product was examined in 1% EtBr agarose gel, the specific band was cut from the gel, and purified using QIAquick Gel Extraction Kit (Qiagen, Cat. 28706) using the manufacturer’s instructions and sequenced the fragment using the FoxPIsoBf6or primer (NCBI acc. # KF198510).

Cloning of QPCR fragments and QPCR

We designed QPCR primers specifically to distinguish between the different FoxP isoforms of Drosophila melanogaster (see Fig. 2b for the position of the primers in the gene) and the hypertrophic discs gene (hyd, CG9484). Primers where designed using the software Primer3 0.4.0 (http://frodo.wi.mit.edu/). Primers for hyd covered an intron-exon boundary, the intron retention FoxP isoform spans from exon 6 to the intron between exon 6 and 7, isoform A and B forward primer was set between exon 6 and 7 (isoform A) and exon 6 and 8 (isoform B). The settings in Primer3 were: melting temperatures between 58°C and 62°C (ATc<1°C), GC content between 40 and 60% and amplicon length between 90 and 120 base pairs. The size, specificity and annealing temperature was tested in a gradient PCR and checked with gel electrophoresis. The PCR conditions were 94°C for 5 min, denaturation at 94°C for 5 s, annealing for 30 s at 55–65°C; elongation for 30 s at 72°C, 35 cycles, and a last elongation at 72°C for 10 minutes. PCR products were examined in 2% EtBr agarose gels, the specific bands were cut from the gel, and purified using QIAquick Gel Extraction Kit (Qiagen, Cat. 28706). PCR products were then cloned into pGEMTeasy plasmid (Promega, Cat. A1360). These plasmids were transformed into One Shot Top 10 Escherichia coli chemically competent cells (Invitrogen, C404010) and colonies with ampicillin (100 µg/ml) resistance were selected on agarose plates. Clones with the specific insert were picked and grown overnight, at 37°C in 3 ml LB/ampicillin medium in a shaker. Plasmids were purified using Invisorb Spin Plasmid Mini Two columns (Invitek, Ref 1010140300) as described by the manufacturer. Finally, the inserts were fully sequenced and analyzed.

Additionally, two different house-keeping genes, EF1 and RPL32 [37] were used to normalize the data. The different sets of primers used are: EMIsoA3for (5′-ACGCAGCTACGTGGAAGAAC-3′) and EMIsoA3rev (5′-CA-TATATGTCGACCATCCCAA-3′) which amplify a 98bp fragment, from 1100–1107 bp of the ORF of isoform A (NCBI acc. # KF198509); IsolRPCRfor (5′-TCAGTGGAGAATGGAATTTC-3′) and IsolRPCRrev (5′-CATATTATGTCGACCATCCCAA-3′) which amplify a 98bp fragment, from 1100–1107 bp of the ORF of isoform B of FoxP, between exon 6 and exon 7 (NCBI acc. # KF198508); HydOPlfor (5′-AGCATCGGAAATGGATGCAG-3′), HydOPlrrev (5′-AGCATCGGAAATGGATGCAG-3′) and HydOPlrev (5′-AGCATCGGAAATGGATGCAG-3′) which amplify a 97 bp fragment, from 951–1047 bp of the ORF of hyd (NCBI acc. # KF198506); and EF1 and RPL32 primers published in [37].

For the QPCR analysis we used an Mx3005P system and the MxPRO QPCR program (Stratagene; Agilent Technologies, U.S.A.). Every sample was run in triplicate in a 96-well plate in a total volume of 25 µl. The mixture contained 12.5 µl of Kapa SYBR Fast Universal mix (Cat. No 07-KK4600-01 Code KM1100), 0.5 µl of Kapa SYBR Fast Rox Low (50x) Code KD 4601), 5 µl of 1:10 diluted cDNA, 450 nM of each primer and 4.5 µl Nuclease Free water. The QPCR conditions were: 95°C for 10 minutes, followed by 40 cycles of 95°C for 30 seconds, annealing/elongation at 60°C (for intron retention, isoform B, hyd and RPL32), 61°C (EF1) and 65°C (isoform A) for 30 seconds. The efficiency of each gene was determined for each treatment with the slope of a linear regression model using the MxPRO QPCR program. A standard curve was generated for each gene using the cloned QPCR fragment in pGem-T easy vector and as internal normalization between plates we used the 1×105 dilution of the standard curve. We determined relative expression levels through normalization of the EF1 house keeping gene [37] which gave the best coefficient of correlation for RNAi and mutant strains (comparing EF1, RPL32 and hyd) using the BestKeeper software tool. Relative expression levels were determined with the comparative cycle time (Ct) method. All primers used in this study amplified the cDNA with similar efficiency (E = 100e−8−8%) in a validation experiment.

ANOVA and a Tukey’s Multiple Comparison post-hoc test was used to identify significant differences between strains in the FoxP expression with each isoform.

Data mining of sequences of FoxP subfamily members

mRNA sequences of 80 FoxP subfamily members of invertebrates and vertebrates were downloaded from NCBI (http://www.ncbi.nlm.nih.gov/gene). We used the ORF finder from NCBI (http://www.ncbi.nlm.nih.gov/projects/gorf) to locate and cut all open reading frames.

Phylogenetic analysis

We created alignments across all species (see Table 1 for a list) using the European Bioinformatics Institute version of ClustalW [38] (http://www.ebi.ac.uk/Tools/clustalw2). The resulting.aln file was downloaded and opened with the Bioedit program (V.7.0.9; Tom Hall, Ibis Biosciences, Carlsbad, CA9208). and saved as a.phy file for further analysis. The Phylip program (V.3.69; Joe Felsenstein, Department of Genome Sciences at the University of Washington) was used to do bootstrapping (Seqboot.exe program of Phylip) using the default parameters; Distance Matrix (dndist.exe program of Phylip) using the default parameters; we used the neighbor joining method (neighbor.exe program Phylip) using Hydrop magnesium Ribululose as the outgroup and rooting the tree. To determine the consensus tree of all FoxP
Figure 2. Insertion mutants of the Drosophila FoxP gene affect isoform expression and flight performance. a, Rooted phylogenetic tree using the genomic sequence of the Drosophila FoxP gene (see Materials and Methods). The single invertebrate FoxP gene probably corresponds to the ancestral form from which the four vertebrate genes have arisen by serial duplication. b, Location of the three insertions (black triangles) and qRT-PCR primer pairs (grey arrows) on the genomic structure of the dFoxP gene (left). Structure of the three cloned transcripts (right). IR: intron retention; FH: Forkhead-Box Domain. c, Expression levels of the three dFoxP isoform mRNAs in heads of Canton S wildtype flies and of the three insertion lines using qRT-PCR. Horizontal lines denote averages, individual circles, triangles and squares constitute different biological replicates. d, Total flight duration for Canton S, FoxP and FoxP insertion lines (25%-75% and 15%-85% intervals).
sequences we used the consense.exe program of Phylip changing the parameters of the Hydra m. outgroup and treated the tree as rooted. We used the nexus-formatted text output of the Consense program of Phylip to generate the phylogenetic tree with the online interactive tree of life (iTOL) software (http://itol.embl.de)[39].

Stationary flight preparation

After briefly immobilizing 24–48 h old female flies by cold-anesthesia, head and thorax were glued (Sinfony Indirect Lab Composite, 3 M ESPE, St. Paul, MN, USA) to a triangle-shaped copper hook (diameter 0.05 mm) either the day before the experiment (learning tests) or 1–2 hours before the experiments (flight performance). The animals were then kept individually in small moist chambers containing a few grains of sucrose until the experiment [34].

Flight performance measurement

Flight performance was measured as described previously [40]. The fly, glued to the hook as described above, was attached to the experimental setup via a clamp to allow stationary flight. For observation, the fly was illuminated from behind and above (Schott, 150 Watt, 15 V) and fixed in front of a polystyrene panel. Additionally, it was shielded by another polystyrene panel from the experimenter. Tarsal contact with a bead of polystyrene prevented flight initiation before the experiment started. To initiate flight, the polystyrene bead was removed and the fly gently stimulated by a puff of air. The time until the fly ceased flying was recorded (initial flight). The fly was stimulated with a puff of air each time it stopped flying. When flight was not resumed even after three consecutive stimulations, the experiment was completed and the total flight time recorded. Every stimulus after the first one, to which the fly showed a flight response, was recorded. The person scoring the flight time was unaware of the genetic identity of the animal. All animals were included in the study, including those which did not show any flight behavior (recording a flight time of zero seconds and zero flight initiations). Flight performance measurements were conducted after an initial observation during learning pilot experiments.

Learning experiments

The core device of the set-up is the torque compensator (torque meter)[41]. It measures a fly’s angular momentum around its vertical body axis, caused by intended flight maneuvers. The fly, glued to the hook as described above, was attached to the torque meter via a clamp to accomplish stationary flight in the center of a cylindrical panorama (arena, diameter 58 mm), which was homogeneously illuminated from behind. The light source was a 100 W, 12 V tungsten-iodine bulb. For green and blue illumination of the arena, the light was passed through monochromatic broad-band Kodak Wratten gelatin filters (#47 and #99, respectively) [42]. Filters were exchanged by a fast solenoid within 0.1 s. Alternatively, the arena was illuminated with daylight by passing it through a blue-green filter (Rosco “surfblue” No. 5433). The transmission spectrum of the Rosco blue-green filter used in this study is equivalent to that of a BG18 filter (Schott, Mainz) and constitutes an intermediate between the Kodak blue and green filters [42]. An analogue to digital converter card (PCL812; Advantech Co.) fed the yaw torque signal into a computer which stored the trace (sampling frequency 20 Hz) for later analysis. Punishment was achieved by applying heat from an adjustable infrared laser (StockerYale Lasiris SNP series, LAS-SNF-XXXX-8308; 825 nm, 150 mW), directed from behind and above onto the fly’s head and thorax. The laser beam was pulsed (approx. 200 ms pulse width at ~4 Hz) and its intensity reduced to assure the survival of the flies. The entire experimental procedure can be seen on video [34].

A second set-up was used by a different experimenter (JC) to first independently reproduce some of the FoxP3 results and then perform the habit formation experiments (see below). Instead of the torque compensator described above, we used a torque meter as described elsewhere [43], to measure yaw torque. The remaining components of the set-up were analogous to the ones described above, with a color switch (implemented using the voice coil actuator of a computer hard disk drive [44], instead of a solenoid), analogous to the ones described above, with a color switch (implemented using the voice coil actuator of a computer hard disk drive [44], instead of a solenoid), an analog to digital converter card (ADC-USB-120FS, measurement computing Inc., 10 Commerce Way, Norton, MA 02766, USA), which transformed the analog signal into a 12 bit digital signal that feeds into the computer, and software to control the experiment and record the data (LabView, National Instruments Germany GmbH, Ganghoferstraße 70 b, 80339 München).

Operant self-learning was performed as previously described [31,32]. The direction for straight flight in all experiments at the torque meter was determined as the central value exactly between the maximum left and right turning yaw torque elicited by an optomotor stimulus. The fly’s spontaneous yaw torque range was then divided into ‘left’ and ‘right’ domains at this value. There were no patterns on the arena wall, but the illumination was spectrally restricted by a blue-green daylight filter. During training, heat was applied whenever the fly’s yaw torque was in one domain and switched off when the torque passed into the other. Punishment of yaw-torque domains was always counter-balanced. In the test phases, heat was permanently switched off and the fly’s choice of yaw torque domains recorded.

The type of operant world-learning used in this study is a modification of self-learning, as described previously [45,46]. During training, the fly was heated whenever its yaw torque passed into the domain associated with punishment, as in self-learning. In addition, whenever the fly switched yaw torque domains, not only temperature but also arena coloration were changed (from green to blue or vice versa). Thus, yaw torque domain and color served as equivalent predictors of heat. In the test phases, heat was permanently switched off and only the fly’s choice of yaw torque domains/colors recorded. Punishment of the color/yaw-torque combination was always counterbalanced. It has been shown previously that self- and world-learning, despite both being operant conditioning procedures, engage different biological learning mechanisms [32,47,48].

Habit formation was tested also as described previously [46]: for the first 26 minutes of the experiment the color filters were present, providing sixteen minutes of training. Thereafter, the two color filters were removed from the light source and replaced by a single blue-green filter, as in the self-learning experiments. After a short, 60s familiarization training, yaw torque preference was measured with the heat permanently switched off.

The color/yaw torque domain preference of individual flies was quantified as the performance index: \( PI = (t_{a}-t_{b})/(t_{a}+t_{b}) \). During
Table 1. List of sequences used for the phylogenetic analysis of FoxP subfamily members.

| Sequence | Accession numbers | bp   |
|----------|-------------------|------|
| Sequence 1: FoxP_Nasonia_vitripennis | XM_001599977.1 | 2067 bp |
| Sequence 2: FoxP_Anopheles_gambiae | XM_565765.3 | 1098 bp |
| Sequence 3: FoxP_Hydra_magnipapillata | XM_002164074.1 | 1746 bp |
| Sequence 4: FoxP_Strongylocentrotus_purpuratus | DQ286749.1 | 1747 bp |
| Sequence 5: IsoAApis | NM_001104949.1 | 2177 bp |
| Sequence 6: IsoATribolium | LOC657917 | 2706 bp |
| Sequence 7: IsoABombyxmolito | BGIBMGA004582-TA^ | 1239 bp |
| Sequence 8: FoxPIADrosophila | NM_001104256.1 | 1329 bp |
| Sequence 9: FoxP2_Gorilla_gorilla | AF512948 | 2142 bp |
| Sequence 10: FoxP2_Homo_sapiens | NM_148898.3 | 2148 bp |
| Sequence 11: FoxP2_Mus_musculus | NM_053242.4 | 2145 bp |
| Sequence 12: FoxP2_Danio_rerio | NM_001030082.1 | 2043 bp |
| Sequence 13: FoxP2_Rattus_norvegicus | XM_002729286.1 | 2097 bp |
| Sequence 14: FoxP2_Gallus_gallus | XM_001232321.1 | 1237 bp |
| Sequence 15: FoxP2_Pan_troglodytes | NM_001090920.2 | 2151 bp |
| Sequence 16: FoxP2_Canis_familiaris | XM_860734.1 | 2133 bp |
| Sequence 17: FoxP2_Xenopus_laeviss | NM_001095669.1 | 2121 bp |
| Sequence 18: FoxP2_Bos_taurus | NM_001205569.1 | 2130 bp |
| Sequence 19: FoxP2_Macaca_mulatta | NM_00133021.1 | 2145 bp |
| Sequence 20: FoxP2_Sus_scrofa | NM_001113049.1 | 2130 bp |
| Sequence 21: FoxP2_Felis_catus | NM_001131377.1 | 2124 bp |
| Sequence 22: FoxP2_Papio_anubis | NM_001168922.1 | 2196 bp |
| Sequence 23: FoxP1_Gallus_gallus | NM_001024827.1 | 2061 bp |
| Sequence 24: FoxP1_Bos_taurus | NM_001083689.1 | 2025 bp |
| Sequence 25: FoxP1_Oryctolagus_cuniculus | XM_002713313.1 | 1734 bp |
| Sequence 26: FoxP1_Callithrix_jacchus | XM_002758547.1 | 2034 bp |
| Sequence 27: FoxP1_MONODELPHIS_DOMESTICA | XM_001364178.1 | 2127 bp |
| Sequence 28: FoxP1_Macaca_mulatta | XM_001084998.2 | 2034 bp |
| Sequence 29: FoxP1_Canis_familiaris | XM_858603.1 | 2034 bp |
| Sequence 30: FoxP1_Pan_troglodytes | XM_001140904.1 | 2034 bp |
| Sequence 31: FoxP1_Omithorhynchus_anatinus | XM_001509776.1 | 2061 bp |
| Sequence 32: FoxP1_Equus_caballus | XM_001498190.1 | 1806 bp |
| Sequence 33: FoxP1_Homo_sapiens | NM_032682.4 | 2034 bp |
| Sequence 34: FoxP1_Xenopus_laeviss | NM_001095533.1 | 1624 bp |
| Sequence 35: FoxP1_Rattus_norvegicus | NM_001034131.1 | 2136 bp |
| Sequence 36: FoxP1_Mus_musculus | NM_001197321.1 | 2118 bp |
| Sequence 37: FoxP4_Rattus_norvegicus | NM_001109588.1 | 867 bp |
| Sequence 38: FoxP4_Canis_familiaris | XM_538914.2 | 1875 bp |
| Sequence 39: FoxP4_Macaca_mulatta | XM_001082913.2 | 2004 bp |
| Sequence 40: FoxP4_Sus_scrofa | XM_001926847.1 | 2064 bp |
| Sequence 41: FoxP4_Pan_troglodytes | XM_518463.2 | 2058 bp |
| Sequence 42: FoxP4_Papio_anubis | NM_001168744.1 | 2043 bp |
| Sequence 43: FoxP4_Bos_taurus | XM_002684495.1 | 2058 bp |
| Sequence 44: FoxP4_Pongo_abelii | XM_002816867.1 | 2043 bp |
| Sequence 45: FoxP4_Equus_caballus | XM_001501047.2 | 2049 bp |
| Sequence 46: FoxP4_X_haemae | NM_001095615.1 | 1926 bp |
| Sequence 47: FoxP4_Danio_rerio | NM_001199491.1 | 2091 bp |
| Sequence 48: FoxP4_X_tropicalis | NM_001077187.2 | 1938 bp |
| Sequence 49: FoxP4_Homo_sapiens | NM_001012426.1 | 2043 bp |
| Sequence 50: FoxP4_Mus_musculus | NM_001110824.1 | 2058 bp |
training periods, t_h indicates the time the fly is exposed to the heat and t_a the time without heat. During tests, t_a and t_h refer to the times when the fly chose the formerly (or subsequently) unpunished or punished situation, respectively. Thus, a PI of 1 means the fly spent the entire period in the situation not associated with heat, whereas a PI of -1 indicates that the fly spent the entire period in the situation associated with heat. Accordingly, a PI of zero indicates that the fly distributed the time evenly between heated and non-heated situations and a PI of 0.5 indicates that 90 of the 120 s in that period were spent in the unpunished situation.

The experiments are fully automated and computer-controlled. Each fly was used only once. The time-course of each experiment was divided into consecutive periods of 2 minutes duration. Depending on whether heat was applied during such a period, it was termed a training period (heating possible) or a test period (heat off). Standard experiments consisted of two pre-test periods (labeled PI1 and PI2), four training periods (PI3, PI4, PI6 and PI7) and three memory test periods (PI5, PI8 and PI9). For experiments with extended training, the experimental time course was essentially repeated such that in total four additional training periods (PI9, PI10, PI12, PI13) followed test-PI8, as well as one additional test period (PI11). In these extended-training experiments, the color filters were removed after period 13 and period 14 was a 60 s familiarization training without colors. The final PI15 and PI16 were scored as memory tests. Depicted are always the PI’s of the first two minutes after the last training period, i.e., PI8 for all standard experiments (Fig. 3, 4 and 5a, b) and PI15 for the extended-training experiments testing habit formation (Fig. 5d).

All data are expressed as means ± SEM. Statistical differences between groups were tested with a Mann-Whitney U-Test or a non-parametric Kruskal-Wallis ANOVA with subsequent two-sided, Bonferroni corrected, posthoc-tests. The PIs for the genetic control strains (elav-driver and FoxP-RNAi-effector lines, respectively, crossed to Canton S) for the FoxP RNAi experiments (Fig. 5) were pooled because they did not differ in their performance and were all different from zero when tested with a t-test for single means (Fig. S2, raw data deposited at figshare, DOI 10.6084/m9.figshare.740444, for close scrutiny of all the training and test PIs of all groups and individual flies).

* Bombyx mori data base.

Table 1. Cont.

| Sequence | Accession numbers | bp |
|----------|------------------|----|
| Sequence 51: FoxP3_Mus_musculus | NM_001199347.1 | 1290 bp |
| Sequence 52: FoxP3_Homo_sapiens | NM_014009.3 | 1296 bp |
| Sequence 53: FoxP3_Rattus_norvegicus | NM_001108250.1 | 1290 bp |
| Sequence 54: FoxP1_Taeniopygia_guttata | NM_001076698.1 | 2052 bp |
| Sequence 55: FoxP4_Taeniopygia_guttata | JN160732 | 2007 bp |
| Sequence 56: FoxP2_Taeniopygia_guttata | NM_001048263.1 | 2136 bp |
| Sequence 57: FoxP3_Macaca_mulatta | NM_001032918.1 | 1296 bp |
| Sequence 58: FoxP3_Sus_scrofa | NM_001128438.1 | 1296 bp |
| Sequence 59: FoxP3_Bos_taurus | NM_001045933.1 | 1296 bp |
| Sequence 60: FoxP3_Felis_catus | NM_001083952.1 | 1293 bp |
| Sequence 61: FoxP2_Melopsittacus_undulatus | AY466101 | 2130 bp |
| Sequence 62: FoxP2_Arctonyx_collaris | EU076391.1 | 2121 bp |
| Sequence 63: FoxP2_Asseliscus_stoliczkanus | EU076392 | 2151 bp |
| Sequence 64: FoxP2_Coelops_fritthii | EU076393.1 | 2232 bp |
| Sequence 65: FoxP2_Capra_hircus | EU076394.1 | 2199 bp |
| Sequence 66: FoxP2_Chimarrogale_himalayica | EU076395.1 | 2202 bp |
| Sequence 67: FoxP2_Chaerophon_plicatus | EU076396 | 2163 bp |
| Sequence 68: FoxP2_Cynopterus_sphinx | EU076397.1 | 2283 bp |
| Sequence 69: FoxP2_Equis_asinus | EU076398.1 | 2118 bp |
| Sequence 70: FoxP2_Hippodermis_armiger | EU076400 | 2202 bp |
| Sequence 71: FoxP2_Megaderma_spasma | EU076401 | 2123 bp |
| Sequence 72: FoxP2_Miniopterus_schreibersii | EU076402 | 2151 bp |
| Sequence 73: FoxP2_Myotis_ricketti | EU076403 | 2151 bp |
| Sequence 74: FoxP2_Oryctolagus_cuniculus | EU076404 | 2151 bp |
| Sequence 75: FoxP2_R.ferrumequinum | EU076405 | 2133 bp |
| Sequence 76: FoxP2_Rousettus_leschenaultii | EU076407 | 2262 bp |
| Sequence 77: FoxP2_Taphozous_melanopogon | EU076409 | 2154 bp |
| Sequence 78: FoxP2_Tylonycteris_pachypus | EU076410 | 2151 bp |
| Sequence 79: FoxP2_Trachemys_scripta | EU076411 | 2106 bp |
| Sequence 80: FoxP2_R._luctus | EU076406 | 2130 bp |

* Bombyx mori data base.
doi:10.1371/journal.pone.0100648.t001
Brain anatomy

Brains of 5–15 day old female flies were dissected and wholemounts were treated as described previously [49]. In brief, to visualize neuropil structures, immunostainings using the synaptic marker antibody nc82 (the Developmental Studies Hybridoma Bank; DSHB; this mouse monoclonal antibody binds to an epitope of the bruchpilot gene product [50]) were performed. Brains were fixed in 4% paraformaldehyde in phosphate-buffered solution (PBS; 0.1 M, pH 7.4) for 30 min on ice. Then the brains were washed with PBS containing 0.2% Triton X-100 (PBST) for 60 min (3 times for 20 min) at room temperature (RT). After being blocked in 5% normal goat serum (NGS) in PBST (PBST-NGS) for 60 min at RT, the brains were incubated in 1:30 nc82 antibody in PBST-NGS for 2 days at 4°C. Then the brains were washed for 60 min at RT and incubated in 1:200 goat anti-mouse Alexa Fluor 488 (Invitrogen) for 2 days at 4°C. Afterwards, brains were washed several times, ten minutes each, and mounted in Vectashield (Vector, Burlingame, CA) using custom-made slides and coverslips with standard 0.170 μm thickness.

Confocal images were taken with a Zeiss LSM510 confocal microscope (Carl Zeiss, Jena, Germany) using the Ar-Kr 488-nm laser line and a 40x water immersion (C-Apochromat) objective (NA: 1.2, Carl Zeiss) resulting in a voxel resolution of approximately 0.3×0.3×1 μm for a single section. In all confocal scans we used a pixel resolution of 1024×1024 in xy-axis and an 8 bit intensity resolution. The wholemount brains were thus scanned in tiled stacks, the central brain and each optic lobe separately.

Confocal image stacks were imported into the 3D software Amira (version 4.1; Visage Imaging, Berlin; San Diego, CA, USA), using the three-dimensional visualization and segmentation modules. In the case of tiled images, stacks were combined with the ‘Merge’ module. Neuropil borders were either manually segmented or semi-automatically using the 3D reconstruction module, compiled using the ‘LabelField Editor’ in the ‘Amira Image Segmentation’. The segmentation results were image stacks of type ‘LabelField’ that assign a label, which represents a distinct (brain) structure, to each voxel. The labels were rendered as surfaces for each neuropil and the ‘Statistics’ module was used for volumetric measurements.

The major neuropils of the Drosophila brain were defined according to the atlas proposed by Chiang et al. [51]. Only

Figure 3. The mutant line FoxP3955 was impaired in operant self-but not world-learning. a1, In operant self-learning the fly, tethered to a torque meter, could operate a punishing heat-beam with its yaw torque. Torque of one domain (e.g., ‘right’) may turn the heat on and the other (e.g., ‘left’) off, or vice versa. a2, Self-learning performance indices in a two-minute test with the heat permanently switched off immediately after eight minutes of training showed a significant impairment of FoxP3955 mutant flies compared to wild type Canton S (CS) control animals (Mann-Whitney U-Test, U = 101.5, p < 0.003).

b1, In operant world-learning, the fly still operated the heat with its yaw torque, but the coloration of the environment changed with the heat as well, allowing for the colors to indicate both heat and torque domain. b2, World-learning performance indices in a two minute test with the heat permanently switched off immediately after eight minutes of training. There was no significant difference in performance between the two strains (Mann-Whitney U-Test, U = 201.5, p > 0.5). Numbers in bars denote number of animals throughout.

doi:10.1371/journal.pone.0100648.g003

Figure 4. Deficiency ED5438 uncovers the FoxP3955 self-learning phenotype. a, Genomic region of dFoxP gene. The deficiency deletes all exons of the dFoxP locus up until the 5-SZ-3955 insertion, which was used to generate the deficiency, as well as 52 upstream genes. ED5438 leaves the downstream gene hyperplastic disks (hyd) intact. b, Operant self-learning performance indices in a two-minute test with the heat permanently switched off immediately after eight minutes of training showed a significant impairment of FoxP3955/ED5438 flies compared to control animals in which either the deficiency or a Canton S chromosome was crossed over the 5-SZ-3955 insertion (Kruskal Wallis ANOVA, H(2, N = 52) = 10.13; p < 0.007; two-sided, Bonferroni-corrected post-hoc p-values indicated in the graph).

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neuropils where the synaptic marker nc82 clearly defined the subneuropil borders were selected for volumetric measurements. These are 7 bilateral neuropils, among them the output areas of the optic and olfactory peripheral nervous system, i.e. optic tubercle/glomeruli and the mushroom bodies, respectively (medulla, lobula, lobula plate, mushroom bodies, AL – antennal lobes). Additionally, four non-bilateral neuropils belonging to the central complex (ellipsoid body, noduli, fan-shaped body, protocerebral bridge) were reconstructed. The remaining neuropils of the central brain (i.e. the protocerebral lobes) were summarized as one neuropil. All values are expressed as a percentage of the sum of all brain neuropils measured.

Amira surface files were exported as wavefront (.obj) files. Wavefront files of neurons and neuropil surfaces were imported with the Adobe 3D Reviewer to Adobe Acrobat Pro Extended (Adobe Systems, Inc.) [32]. The images in the PDF version (Figure 6a) of this publication can be viewed by using the 3D viewer mode of the Acrobat Reader (version 9 and higher, which is freely available at http://get.adobe.com/de/reader/).

The Principal Component Analysis (PCA) was performed on the symmetric correlation matrix (Pearson’s method) of the volume percentage data using the R statistics package (http://r-project.org). PCA is a method that allows identifying the major patterns of correlations among multivariate data and thus condensing as much of the variation in the data as possible in few dimensions –

Figure 5. Drosophila FoxP full length isoform B is required for operant self-learning. a, Flies expressing an RNAi construct targeting the FoxP isoform B were impaired in operant self-learning, compared to the genetic controls (pooled, see Materials and Methods; Mann-Whitney U-Test, U = 378.5, p < 0.03) in a 2-minute test immediately after eight minutes of training. See raw data or Fig. S2 for separated control groups. b, Both flies expressing an RNAi construct targeting the FoxP isoform B, as well as genetic control flies performed well in operant world-learning (Mann-Whitney U-Test, U = 420.5, P < 0.2). c, No reduction in the expression of isoform B mRNA using qRT-PCR in flies expressing an RNAi construct targeting the FoxP isoform B. Canton S data are identical to those in Fig. 2c, as experiments were performed simultaneously. d, Mutant FoxP^{P955} flies were impaired in habit formation. In a 2-minute self-learning test (i.e. without colors) after 16 minutes of training in world-learning (i.e. with colors), FoxP^{P955} flies showed a significantly reduced preference for the previously unpunished turning-maneuvers, compared to wild type control animals (Mann-Whitney U-Test, U = 186.0, p < 0.04). However, the mutant flies nevertheless showed a significant performance index (t-test against zero: df: 23; t = 2.54; p < 0.02).

Figure 6. Subtle morphological alterations in the brains of FoxP^{P955} mutants. a, Three-dimensional surface renderings of typical fly brains from wild type Canton S (a1) and FoxP^{P955} mutants (a2). PLOS ONE can only handle 3D PDF figures as part of the supplementary files. Hence, the 3D functionality for Figure 6 is available as Figure S3. Alternatively, a fully functional PDF will be hosted on BBs website. b, Quantitative volumetric analysis of eleven major neuropils (M – medulla, L – lobula, LP – lobula plate, MB – mushroom bodies, AL – antennal lobes, FB – fan-shaped body, OT – optic tubercle, EB – ellipsoid body, OG – optic glomeruli (purple in a), PB – protocerebral bridge, N – noduli) revealed a significant reduction in the volume of the optic glomeruli in FoxP^{P955} flies (Mann-Whitney U-Test, U = 2.0, p < 0.002). The volume of the remaining neuropils (denoted PL – protocerebral lobes) did not differ significantly. Asterisk – significant difference with a Bonferroni-corrected level of p < 0.004. Black stripes – median, boxes – 25–75% percentiles, whiskers – total range. Grey boxes indicate FoxP^{P955}, white boxes Canton S. c, Principal Components Analysis of the volumetric data. Plotted are the factor loadings of the individual flies on the two first components. Colored bars indicate means and standard errors (PC). Factor loadings are significantly different between Canton S and FoxP^{P955} for PC1 (Mann-Whitney U-Test, U = 52.0, p < 0.04), but fail to reach significance for PC2. Number of brains analyzed: 7 (Canton S) and 9 (FoxP^{P955}).
the principal components (PC). Factor loadings for each individual fly were plotted for the first two (2D plot) or three (3D plot movie) components. Mean and standard error of the loadings were calculated for each of the three components and plotted as well. The ellipsoids in the 3D representation represent the 70% confidence interval using the covariance between the three first components.

Results

Flight performance and dFoxP expression in three insertion lines

The four different FoxP genes in vertebrates probably arose from serial duplications of a single ancestral FoxP gene after the separation of the invertebrate clades (Fig. 2a). The invertebrate FoxP orthologue corresponds most closely to the ancestral form of the gene at the base of the bilateria [36], thus lending itself to investigating the depth of the functional conservation among the members of the FoxP gene family. Two FoxP isoforms generated by differential splicing have been described for the fruit fly Drosophila melanogaster [36] and we discovered a third transcript, generated by intron retention (Fig. 2b, c). All three isoforms differ in the sequence of the Forkhead Box domain, presumably resulting in different DNA binding properties [53].

We studied dFoxP gene expression in three fly lines each with a distinct insertion in or near the last dFoxP exon, but otherwise genetically highly similar to the Canton S control strain (lines FoxP3955, FoxP03746 and FoxP063619, Fig. 2b). We performed quantitative polymerase chain reaction following reverse transcription (qRT-PCR), to uncover the effects of the insertions on dFoxP gene expression. None of the insertion lines showed any deviation in the level of the neighboring gene, hypenastic disc (hyd). In contrast, the three lines differed with respect to expression of the three dFoxP isoforms (Fig. 2c). We could not detect any expression of the intron retention isoform in the lines FoxP03746 and FoxP063619. Line FoxP03746 showed elevated expression of isoform A and line FoxP0635, where the P-element had inserted into the last exon of dFoxP, does not show any significant difference in any of the isoforms. Genome and RNA sequencing revealed that the inserted P-element of line FoxP0635 is transcribed within exon 8, resulting in a premature stop codon, such that the putative isoform B protein is truncated at position 486 (about 93% of the original size) and its final amino acid is changed to an alanine instead of B protein is truncated at position 486 (about 93% of the original size) and its final amino acid is changed to an alanine instead of

Flies with manipulated dFoxP expression are specifically impaired in operant self-learning

One prediction of the hypothesis that the ancestral FoxP gene played a role in the forms of motor learning most similar to vocal learning is that manipulations of the Drosophila FoxP gene should affect learning with reference to its own body, ‘self-learning’ [47,48], more than other forms of operant learning [32]. In the operant self-learning paradigm that we used [31], the tethered fly’s attempts to turn are measured by a torque meter and divided into two domains, roughly corresponding to ‘left’ and ‘right’ turns, respectively (Fig. 3a1). Fixed in space in a featureless environment, the fly alternates between turning directions in a highly variable, random-like fashion [54]. Making punishment by an infrared heat beam contingent on one set of maneuvers (e.g., right turning attempts), leads to a reduction in the variability of the behavior as the fly restricts its turning maneuvers to the unpunished side (i.e., left turning attempts). This reduction outlines the application of the heat beam after a given training period and the flies restrict their yaw torque to the previously unpunished side, even when the heat is permanently switched off [31,32].

After eight minutes of training, wild type flies of the strain Canton S spent a significantly larger fraction of the 2-minute test without heat initiating previously unpunished turning attempts (82.5 ± 5.2%) than FoxP3955 mutant flies (57.1 ± 6.9%; Fig. 3a2), with a structural mutation in the last exon of the dFoxP gene, but with wild type expression levels. To test if the flies possessed the general capabilities to solve these kinds of learning tasks, i.e., generate variable yaw torque, sense and learn to avoid the heat and restrict their yaw torque in the absence of heat, we tested FoxP3955 flies in a very similar experiment, with only a single, crucial difference. Adding a solenoid with green and blue color filters to the setup allowed us to alternate the coloration of the fly’s environment between the two colors together with the switch in the turning attempts. For instance, left turning attempts might have led to green coloration and right turning attempts to blue coloration or vice versa. This alteration allowed the FoxP3955 mutant flies to use the colors as external cues (‘world-learning’ [47,48]) instead of their own behavior (‘self-learning’) to predict the heat punishment, rescuing their performance (Fig. 3b). This result demonstrates that FoxP3955 mutant flies are able to generate variable yaw torque, sense and learn to avoid the heat and restrict their yaw torque in the absence of heat.

The 5-SZ-3955 insertion had been introgressed into the Canton S genetic background in order to minimize the chance of genetic aberrations elsewhere potentially causing a phenotype. We used an additional strategy classically used to map a phenotype to a specific mutation. We crossed both FoxP3955 mutant flies and Canton S control flies over a deficiency which deleted all exons of the FoxP3955 gene up until the 5-SZ-3955 insertion site, as well as 52 other, upstream genes (Fig. 4a). In parallel, we tested the progeny of a Canton S - FoxP3955 cross for a potential dominant effect of the 5-SZ-3955 insertion. The heterozygote F1 offspring of each of the three crosses were tested in operant self-learning as described above. The recessive self-learning phenotype of the FoxP3955 mutation was uncovered by the deficiency (Fig. 4b), indicating that indeed the 5-SZ-3955 insertion is the cause for the mutant phenotype of these flies. Together with the truncated isoform B expression in FoxP3955, the above results encouraged us to attempt a regulatory manipulation of dFoxP expression. We hypothesized that isoform B may be specifically required for operant self-learning.

To test this hypothesis, we targeted isoform B by expressing a specific RNAi construct under the control of the pan-neuronal clav-GAL4 driver line. These flies did not appear impaired in their flight ability and hence their flight performance was not quantified. Consistent with our hypothesis, subjecting flies expressing an RNAi-construct targeting isoform B to self- and world-learning yielded a phenocopy of the mutant data (Fig. 5a, b). We assessed the effectiveness of the RNAi method using qRT-PCR, but the level of isoform B mRNA was unaltered in the experimental flies (Fig. 5c). Genome sequencing revealed two polymorphisms in the RNAi target region (NCBI Acc. # KF192040-KF192876), potentially explaining the lack of a knock-down (see discussion).
Insertion line dFoxP3955 is impaired in habit formation

Crystalization of birdsong shares a number of aspects with habit formation, a process induced by extended operant conditioning [55]. In humans, the mutations in the FOXP2 gene that lead to the developmental dyspraxia speech phenotype cause morphological alterations in cortico-striatal circuits [12,17]. These circuits are also involved in habit formation [35,36] and express FoxP2 in songbirds [57]. In flies, extended operant world-learning also produces habit formation: removing the color filters after sixteen minutes of training (but not after eight minutes) reveals a preference for the previously unpunished yaw torque domain [46].

Using flies with the FoxP3955 insertion, we tested the hypothesis that the FoxP-dependent process of self-learning also underlies habit formation, but is inhibited during non-extended world-learning. Consistent with our hypothesis, FoxP3955 mutants showed a significant impairment in habit formation after extended world-learning (Fig. 5d). Interestingly, the FoxP3955 mutant flies do show a significant performance index after extended operant world-learning.

Subtle morphological alterations in the brains of FoxP3955 mutant flies

FoxP mutations cause neuroanatomical alterations in multiple species [12,16,17,57–59]. In flies, the general brain anatomy, structure, shape and relative position of the major neuropils did not differ markedly between wildtype and FoxP3955 mutant flies (Fig. 6a). However, quantitative volumetric analysis of eleven major neuropils revealed that the most conspicuous of the recently identified ‘optic glomeruli’ [60] were significantly smaller in the mutant flies compared to Canton S controls (Fig. 6b). Moreover, a Principal Components Analysis (PCA) performed on the relative volume data of all neuropils also distinguished the FoxP3955 mutant and control groups, in spite of spatial overlap of some individuals (Fig. 6c; for PCA results with data from optic glomeruli excluded, see Fig. S4). These results indicate that in flies, as in vertebrates, experimental manipulations of FoxP expression can lead to alterations in brain morphology.

Discussion

Genetic manipulation of the dFoxP locus

We selected three insertions within the dFoxP locus. Prior to testing, all three isoforms had been introgressed into the Canton S wild type strain, homogenizing the genetic background of all four lines. Two of the lines (FoxP39476 and FoxP39519) did not express any detectable level of the novel intron-retention isoform mRNA. These two lines also showed a significant impairment in flight initiation and maintenance, suggesting that this isoform may be involved in flight performance (Fig. 2). A recent study supports the notion of a broader function of the different dFoxP isoforms in motor control [61] above and beyond operant self-learning. Coincidentally, this study and ours also serve to suggest that the dFoxP-dependent phenotype described in another, yet more recent report [62] may potentially also be due to motor issues, in contrast to the assertions of those authors. The operant self-learning phenotype of FoxP3955 mutant flies (Fig. 3) was uncovered in the heterozygous stage over a deficiency spanning the FoxP3955 allele, solidly tying the phenotype to this dFoxP manipulation (Fig. 4).

Line FoxP3955 expressed a mutated isoform B mRNA, such that the putative isoform B protein was truncated and the C-terminal amino acid altered, potentially affecting protein function. There did not appear to be any change in the regulation of isoform B expression in this line as the expression levels were similar to the CS control strain on the mRNA level. This structural modification with its specific effect on operant self- but not world-learning is a reminder that many specific behavioral mutants are often due to structural mutations, commonly affecting only a subset of a gene’s isoforms [63]. Targeting isoform B with a specific RNAi construct in order to probe for regulatory effects of dFoxP, yielded a phenocopy of the FoxP3955 mutant behavioral phenotype, albeit without any detectable knock-down of the mRNA (Fig. 5). We cannot rule out that the knock-down did take place although at undetectable levels due to the already low levels of isoform B expression. The alternative also remains that hypothetical extra-neuronal expression could have masked any knock-down in the neuronal tissue. Finally, mismatches between the RNAi construct sequence and the target sequence can bias the RNAi process towards posttranscriptional silencing rather than mRNA degradation [64–67]. Sequencing of the strains used for the RNAi experiments indeed revealed two such polymorphisms leading to mismatches of the target region with the RNAi construct (see Results), potentially biasing the RNAi process towards sequestration.

It is thus plausible that the RNAi method employed here affected the protein, but not the overall mRNA levels of isoform B, explaining the behavioral phenotype. In the review process of this manuscript, an antibody against dFoxP isoform B was described [61], such that it will now be possible to test this hypothesis. Interestingly, none of the six currently available RNAi lines targeting the dFoxP locus show any detectable knock-down of any of the three isoforms (replicated in two different laboratories, each using a different pan-neuronal driver line, manuscript in prep.), despite one of them also showing a behavioral phenotype in two studies [61,62], as well as mRNA knock-down using semi-quantitative PCR in the Lawton et al. work [61]. As we have observed similar dFoxP mRNA knock-downs using semi-quantitative PCR which were not confirmed by qPCR, we would cautiously speculate that the knock-down observed by Lawton et al. may be a false positive as well, with the behavioral phenotypes in both reports potentially explained by polymorphic mismatches in the target region. For these reasons, we can only tentatively conclude that besides the structural manipulations, also regulatory manipulations of only isoform B expression may affect operant self-learning. The most parsimonious explanation of the common phenotypes after various dFoxP manipulations is that dFoxP is indeed necessary for operant self-learning.

Intact dFoxP gene required for normal brain development

Our volumetric analyses suggest a role of Foxp3955 on brain development in Drosophila, analogous to the role FOXP2 plays in vertebrates [3,4,57,68,69]. Specifically, flies with the mutant Foxp3955 allele have smaller optic glomeruli (Fig. 6). So far, very little is known about the functional role of optic glomeruli and, thus, of their relevance for operant self-learning. It has been hypothesized that optic glomeruli may form information hubs by virtue of containing the terminals of many projection neurons within the ipsilateral brain hemisphere [51]. Optic glomeruli have been discovered in dipterans and it is not known if other insect orders possess optic glomeruli. Nevertheless, it is interesting to note that a conspicuous cluster of AmFoxP-expressing neurons is located near the optic lobes of the honeybee [70]. Notably, no significant change was observed in the protocerebral bridge, a neuropil in which a recently developed dFoxP-GAL4 driver line was reported to express [61].

It deserves to be emphasized, however, that the overall structure of the brains of FoxP3955 mutant flies appears completely normal to the human eye. Only numerical analyses are capable of detecting...
the subtle changes this mutation causes to brain anatomy. More severe alleles of the dFoxP locus have been observed to lead to more severe anatomical defects (Troy Zars, unpublished observation).

FoxP is specifically involved in self-learning

We targeted dFoxP with our genetic manipulations because of the conserved role of the FoxP gene family in vocal learning. We then tested these manipulations each in two very similar operant learning paradigms that differ in their conceptual similarity to language acquisition.

Operant self-learning in Drosophila parallels the operant feedback structure of vocal learning in humans or songbirds in that no other external cues are contingent upon the feedback (Fig. 1). This conceptual similarity is now supported by the parallel biological similarity of both vocal and operant self-learning requiring FoxP function (e.g., [11–13,23,71] and Figs. 3, 4, 5). Interestingly, Protein Kinase C (PKC), the only other known molecular component of the self-learning mechanism [32,72,73], has also been implicated in vocal learning in songbirds [74,75].

The technically all but identical operant world-learning task, in contrast, not only differs conceptually from vocal learning – the behavior is also controlling an external cue (the colors) – but the biological requirements are also different: neither dFoxP (Figs. 3, 5) nor PKC [32] function is necessary. Recent experiments reporting unaffected Pavlovian conditioning in FoxP3955 mutant flies [62] confirm the notion that FoxP is specifically required for self-learning. PKC is not required for other forms of world-learning in flies either [76,77], but in other model systems, the data are less straightforward, with varying requirements of varying PKC isoforms for various phases of memory induction and/or maintenance having been reported, depending on preparation, type of training, time of testing, brain region and method of PKC manipulation. To our knowledge, there is only one experiment where PKC requirement has been compared between self- and world-learning. This experiment in mice supports the view that PKC-activity in the cerebellum is not required for world-learning but for self-learning [72], analogous to the requirements in flies [32]. Conversely, the components that are required for operant world-learning in flies are the same that are required for many other forms of learning in many other model systems, such as classical (Pavlovian) conditioning, sensitization or some forms of habituation, e.g., the type 1 adenylyl cyclase [78–90], encoded by the fly gene rutabaga, which is dispensable for self-learning [32,73].

Thus, we postulate that the converging evidence from multiple model systems concerning PKC and FoxP provides first insights into a core set of mechanisms that are specifically required for operant self-learning, and not for other forms of (associative) learning, such as world learning.

Self-learning and habit formation share the same biochemical substrate

We found that not only self-learning but also habit formation is impaired in FoxP3955 mutant flies (Fig. 5), indicating that habit formation may be mediated by the same biochemical processes as operant self-learning, as had been hypothesized before [46,47]. If wild-type flies are trained in world-learning for eight minutes (i.e., with colors) and then tested for their turning preference in the absence of the colors, there is no preference, demonstrating a hierarchical learning system where external cues are preferentially memorized/retrieved (world-learning) over behavioral cues (self-learning) [46–48]. However, if training in the world-learning situation (i.e., with colors) is extended to 16 minutes and then the same preference test for turning is performed (i.e., without the colors), then a preference for turning towards the previously unpunished direction can be observed [46]. Because self-learning is manifest already after eight minutes of training (i.e., without colors), the presence and learning of the colors during training must have inhibited self-learning, apparently via a neuropil in the fly brain termed mushroom-bodies [46]. Extended training can overcome this inhibition and lead to habit formation. These habits lead to reduced generalization in ‘habit interference’ experiments when an orthogonal behavior is used to control the colors after extended world-learning [46]. Here, we show that flies with a mutated dFoxP gene are impaired in habit formation (Fig. 5), i.e., extended operant world-learning leads to a reduced preference for the previously unpunished yaw torque domain in a test without colors. Interestingly, the preference for the unpunished turning direction in the mutant flies is statistically significant, suggesting that perhaps flies with the mutated gene product are still able to encode some memory, albeit at greatly reduced efficiency. This finding also indicates that perhaps extended self-learning (i.e., without colors) might also be able to yield significant learning scores in FoxP3955 mutant flies. We will therefore test these flies not only in extended self-learning, but also in habit interference experiments in future work.

Vocal learning has been characterized as an automatization of behavior, akin to habit formation in non-vocal mammals [53]. Analogous to the conceptual and biological similarity of self-learning across taxa, both vocal learning and habit formation in vertebrates also share biological substrates in cortico-striatal circuits, where FoxP2 is expressed [12,17,55–57]. Our finding that the FoxP3955 allele is also involved in habit formation in invertebrates not only further supports the homology between PKC/FoxP-mediated operant self-learning in flies and vocal learning in birds and humans, but also prompts the hypothesis that habit formation may require a FoxP gene in vertebrates as well.

Self-learning as an exaptation for language acquisition

Thus, we can draw from several bodies of evidence spanning multiple vertebrate and invertebrate model systems and humans, when we conclude that our results strongly support the hypothesis that the FoxP-dependent component of language evolved from an ancestral operant self-learning mechanism. Interestingly, preliminary data from non-vocal vertebrate motor learning further corroborate this hypothesis [91,92]. Moreover, following this extension of the ‘motor learning hypothesis’ of FoxP function [23,24], one may predict language and/or motor skill deficits in patients with mutations in other FOXP paralogues, anticipating that some of the ancestral function is conserved. Indeed, the symptoms of patients with mutations in the closest paralogue of FoxP2, FOXP1, include language and motor skill impairments [18,19,21,22]. In fact, the Drosophila isoform probably involved in self-learning is not present in FoxP2 genes, but in FoxP1 [36]. Because FoxP gene products form homo- as well as heterodimers [93,94], it is tempting to speculate that all three brain-expressed FoxP paralogues (1, 2 and 4 [95]) may perform similar tasks in a degenerate fashion, with the neuronal circuits controlling language being more susceptible to disruption than other FoxP-expressing circuits.

Autism spectrum disorders and schizophrenia are being discussed as diametric malfunctions on a continuous scale of ‘sense of self’ [5,96]. Given the implication of members of the FoxP gene family in both disorders [5–10,18,22,97,98], it is interesting to note that the operant self-learning mechanism appears to be selectively engaged when the content of the learning
task concerns the organism itself and not when it concerns non-self entities (Figs. 3, 3).

In contrast to Chomsky’s assertion that it is “completely meaningless to speak of extrapolating this concept of operant to ordinary verbal behavior” [99], we were able to show that a transcription factor known for its specific involvement in language is also involved in operant conditioning, specifically operant self-learning. This result is consistent with Skinner’s description of language as an operant behavior [27, 100], while nevertheless espousing Chomsky’s view of inborn components in language acquisition [99]. In particular, our results emphasize the speech component in the evolution of language: “In many respects the story of the evolution of language must begin with the evolution of serial motor activity and its nervous control” [101].

Supporting Information

Figure S1 Four aspects of flight behaviour in wild type and mutant flies. For panels A-C the black squares indicate the median, the boxes signify the 25 and the 75 percentiles, and the error bars range from the 15 to the 85 percentiles. A – Flight duration until the first stop for wild type (Canton S, yellow box), FoxP955 (orange box) and the other two insertion lines (grey boxes). B – Average duration of flight bouts for wild type and mutant lines. C – Total flight duration as in Fig. 1. D – Mean number of stimuli to which the flies responded with flight bouts before they did not respond to 3 consecutive stimuli (errors are s.e.m.s). Numbers in bars = number of animals. Asterisks = significant differences with a Bonferroni-corrected level of p < 0.002. Black stripes = median, boxes = 25–75% percentiles, whiskers = total range. Grey boxes indicate FoxP955, white boxes Canton S. c, Principal Components Analysis of the volumetric data. Plotted are the factor loadings of the individual flies on the two first components. Colored bars indicate means and standard errors (PC). Factor loadings are significantly different between Canton S and Foxg955 for PC1 (Mann-Whitney U-Test, U = 52.0, p < 0.04), but fail to reach significance for PC2. Number of brains analyzed: 7 (Canton S) and 9 (Foxg955).

(EPS)

Figure S2 Separating the genetic control groups still shows the RNAi effect in operant self-learning. Both genetic control lines show significant learning scores of 0.37 and above (t-tests against zero, elav/+; t = 2.26, p = 0.038; RNAi/+; t = 3.53, p = 0.004). In contrast, the flies from the strain in which the elav driver expressed the isoform B-specific RNAi construct in all neurons, fails to reach even half of the lowest control score with a PI of less than 0.17, which cannot be distinguished from a PI of zero (t-test against zero; t = 0.141, P = 0.166).

(EPS)

Figure S3 Subtle morphological alterations in the brains of FoxP955 mutants. a, Three-dimensional surface renderings of typical fly brains from wild type Canton S (a1) and FoxP955 mutants (a2). In the online version, clicking on the reconstructions will activate the 3D features of the figure and allow for interactions with the object in space. The different neuropil areas can be selected in the pop-up menu. b, Quantitative volumetric analysis of eleven major neuropils (M – medulla, L – lobula, LP – lobula plate, MB – mushroom bodies, AL – antennal lobes, FB – fan-shaped body, OT – optic tubercle, EB – ellipsoid body, OG – optic glomeruli (purple in a), PB – protocerebral bridge, N – noduli) revealed a significant reduction in the volume of the optic glomeruli in FoxP955 flies (Mann-Whitney U-Test, U = 2.0, p < 0.002). The volume of the remaining neuropils (denoted PL – protocerebral lobes) did not differ significantly. Asterisk indicates significant difference with a Bonferroni-corrected level of p < 0.004. Black stripes = median, boxes = 25–75% percentiles, whiskers = total range. Grey boxes indicate FoxP955, white boxes Canton S. c, Principal Components Analysis of the volumetric data. Plotted are the factor loadings of the individual flies on the two first components. Colored bars indicate means and standard errors (PC). Factor loadings are significantly different between Canton S and Foxg955 for PC1 (Mann-Whitney U-Test, U = 52.0, p < 0.04), but fail to reach significance for PC2. Number of brains analyzed: 7 (Canton S) and 9 (Foxg955).

(PDF)

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Author Contributions

Conceived and designed the experiments: BB. Performed the experiments: EM JC JR HP TZ CS BB. Analyzed the data: CB BB JR JC EM. Contributed reagents/materials/analysis tools: TZ HP. Wrote the paper: BB CS. Have seen and commented on the manuscript: EM JC JR HP TZ CS BB.

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