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Kyoto University
Transgenic songbirds with suppressed or enhanced activity of CREB transcription factor

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Abstract

Songbirds postnatally develop their skill to utter and to perceive a vocal signal for communication. How genetic and environmental influences act in concert to regulate the development of such skill is not fully understood. Here we report the phenotype of transgenic songbirds with altered intrinsic activity of CREB transcription factor. By viral vector mediated modification of genomic DNA, we established germ-line transmitted lines of zebra finches, which exhibited enhanced or suppressed activity of CREB. Although intrinsically acquired vocalizations or their hearing ability were not affected, the transgenic birds showed reduced vocal learning quality of their own songs and impaired audio-memory formation against conspecific songs. These results thus demonstrate that appropriate activity of CREB is necessary for the postnatal acquisition of learned behavior in songbirds, and the CREB transgenic birds offer a unique opportunity to separately manipulate both genetic and environmental factors that impinge on the postnatal song learning.
Significance

In highly socialized animals such as humans or songbirds, individuals postnatally develop their skills to communicate with conspecifics under the social influence. Both genetic and environmental influences play a crucial role in the development of such abilities, but dissection of the influences has been difficult, because genetic manipulation of avian species is still a challenging issue. In this study, we applied transgenic technology to songbirds along with an experimental song training paradigm to separately manipulate both genes and social environment, and found that appropriate activity of CREB is necessary for the postnatal song learning in songbirds.
The development of behavioral traits in animals is influenced both by intrinsic and extrinsic factors. The contributions of the genetic and environmental factors on the development of such behaviors have often attracted public interest, i.e. the “nature versus nurture” debate; however, tangible dissection of the magnitude of the contributions of such factors has been difficult. The songbird’s skill to vocalize and to perceive a birdsong, a vocal signal for intra-species communication, is one of the prominent skills that require both genetic and environmental factors for the development(1, 2). During postnatal developmental periods of zebra finch (Taeniopygia guttata), juvenile birds hear songs of their conspecifics and store this information inside their brain in order to acquire the knowledge to utter and to perceive vocal signals(3–5). Inadequate auditory experience during the postnatal development results in abnormalities of their songs(3, 6–8), which often results in a reduced communication or mating performance(9–11). On the other hand, even the songs of birds reared in acoustic isolation contain species-specific syllable elements(7, 12) and such birds prefer the songs of conspecifics over those of different songbird species(13, 14). Moreover, it has been reported that some characteristics of vocal traits are heritable(15, 16). Hence, both genetic and environmental influences are necessary for developing the functional neural network required for the proper song vocalization and perception in songbirds.

Neural activity-dependent gene transcription is one of the key mechanisms by which postnatal experience can affect the expression of genes in the neural systems(17, 18). Among the transcription factors that regulate the activity-dependent gene transcription in neurons, the cAMP response element-binding protein (CREB) is one of the most well studied(19–22). Various kind of external stimuli induces phosphorylation of CREB, the modification of which is required for the transcription of target genes(19,
CREB functions as a molecular hub to regulate neuronal gene transcription depending on the neuronal activity (23), and is known to play a pivotal role in neuronal plasticity and memory formation in various species (24–27). In songbirds, however, a previous histological study has found that activation of CREB occurs in the brain regions responsible for vocalization and discrimination of songs in adult zebra finches after they hear songs (28), but its role in vocal learning or auditory discrimination of songs had not been analyzed.

Since the postnatal environment can be controlled experimentally, songbirds have been an ideal experimental animal to study how one’s ability develops according to postnatal experiences (1, 29–31). Furthermore, they are one of the rare species that exhibit imitation of vocal signals, a behavior that is thought to be important for the acquisition of languages in humans (32). Moreover, the skill to communicate with learned vocal patterns are culturally transmitted through social interaction (33, 34). These features make songbirds as a rare and promising experimental system to study not only the requirement of genetic and environmental factor, but also how social or cultural influence stimulates the individual development of a behavior. Nevertheless, the lack of efficient methods to manipulate the genome of songbirds has hampered the research needed to reveal the contribution of the genetic factor to the development of communicative ability. Recently, however, transgenic expression of exogenous genes became possible by the development of virus-mediated transgenic technology in songbirds (35). This technology has allowed us to study in detail the genetic and environmental influences on the song acquisition of zebra finches. Taking this approach, we have established germ-line transmitted lines of zebra finch that express mutant forms of CREB molecules. Here, we show that formation of auditory memories
against conspecific songs and the acquisition of own song is impaired in those
transgenic birds, although the basal hearing ability or the acoustic quality of intrinsic
vocalizations are not altered. These birds demonstrate that appropriate activity of CREB
is necessary for song learning and offer an opportunity to separately manipulate both
genetic and environmental factor influence in the acquisition of postnatally learned
behaviors.

Results

Generation of the transgenic zebra finches with modified CREB activity. By
regulating the expression of multiple genes, transcription factors that show neural
activity-dependent gene transcription, including CREB, can alter the proteomic
landscape depending on environmental influences(24, 36). To substantiate the link
between CREB-mediated gene transcription and the quality of the postnatally acquired
communicative ability of songbirds, we manipulated the activity of CREB in zebra
finches by transgenically expressing mutant CREB molecules. Introduction of
transgenes was performed by injecting lentiviral vectors bearing the transgene into the
early embryo in fertilized zebra finch eggs(35) (SI Material and Methods). Exogenously
expressed mutant CREB1 genes harboring the amino acid substitution affect the activity
of CREB-mediated gene transcription by forming heterodimer with the endogenous
CREB molecule(37). In the present study, we tried to express these mutant zebra finch
CREB1 under the control of a human SYN1 (synapsin I) promoter which restricts the
transgene expression to neurons(38). Transgenic lines expressing the phosphorylation-
deficient form of CREB (DN; S119A, equivalent to mouse CREB-S133A) and the
countitively active form of CREB (Actv; Y120F, equivalent to mouse CREB-Y134F)
were established (Figs. 1 and S1). Germ-line transmission and the expression of the
transgenes (EGFP-CREB) in their offspring were observed in 20 out of 1473 virus-
injected eggs. By crossing these 20 founder birds (11 for DN and 9 for Actv) with wild-
type (WT) birds, a total of 116 DN and 103 Actv transgenic G1-offspring were
obtained. Integration of the transgene into genome (Figs. 1B and S1B), and brain
expression of transgenes were observed in the G1-offspring (Figs. 1C and S1A). As
observed in the transgenic quails (38), the expression of transgene was observed
throughout the NeuN expressing cells (Fig. 1D). These birds express exogenous CREB
mutants in addition to the endogenous CREB (Fig. 1E).

G1-offspring of neither genotype showed any significant difference in their body
weight compared to that of the WT control birds (Fig. S2A). Also, their brain
morphology looked unchanged, and the size of their song nuclei (HVC, Area X, robust
nucleus of the arcopallium) did not show any significant difference (Figs. S2B, C). We
next asked whether these birds show any difference in their hearing ability or behavioral
reactions to stimuli, by observing the reactions against the increasing volume of sounds
(Fig. S3). When the change in behavioral reaction was analyzed by calculating the
frequency of vocalization (“calls”, a short vocalization different from “songs”(39)),
birds of all genotypes equally discriminated the increased volume of white noise
between 45 dB and 47 dB (Fig. S3B). Notably, we found that whereas WT and DN
birds suppressed calling behaviors in response to the volume-change of 45 dB to 47 dB,
Actv birds tended to increase them (Fig. S3C). These observations indicate that
although the basal hearing ability needed to discriminate the sounds was unchanged
between the transgenic birds, Actv birds tend to show a differential reaction in response to stimuli compared with WT and DN birds.

Through maintaining the transgenic lines, we noticed that although the transgenic birds look normal, they tend to die abruptly. We next analyzed the difference of survival ratios among the genotypes. Being kept in our rearing condition, in which all birds were separated from the parents after the fledge and were kept in a sound-proof chamber through ~ 30 to 140 days post hatch (dph), G1-offspring of both genotypes lived significantly shorter compared to WT birds (Fig. S4A). After they become sexually matured, we tried to obtain G2 birds by crossing G1-TgN to WT birds. However, we found that the fertilization rates of G1-TgN were remarkably lower in both sexes (Fig. S4B). Even for the pairs that succeeded to leave G2-offspring, more days were needed to obtain chicks compared to WT (Fig. S4C). Because we were unable to obtain sufficient numbers of G2-offspring, further behavioral analysis was performed using the male G1-offspring from multiple transgenic lines.

**Transgenic zebra finches showed altered intrinsic activity of CREB.** The activity of CREB mediated gene transcription was assayed by a lentivirus based transcription-reporter construct (Fig. 2A). In this construct, a constitutive *PGK1* (human phosphoglycerate kinase 1) promoter expresses an infection-reference gene (flag-tagged Histone-2B, H2B-flag); whereas in the other direction, a minimal promoter expresses a reporter gene (turboGFP, tbGFP). The expression of the reporter gene is influenced by the CREB binding sequence (CRE), inserted just upstream of the minimal promoter in LV-CREB-reporter constructs. By normalizing the quantified number of the transcribed reporter mRNAs using those of internal infection-reference, these constructs can
reliably reflect the activity of CREB irrespective of the transfected number or the genome-inserted locus of the reporter constructs (Fig. 2B). Injecting these lentivirus based transcription-reporter into the brains of transgenic birds, we compared the intrinsic activity of CREB between WT, DN and Actv birds. CREB-mediated gene transcription was significantly suppressed in DN birds and augmented in Actv birds, compared with WT birds (Fig. 2C). Quantitative RT-PCR analysis of the relative expression of endogenous genes in G1-offsprings further revealed the effect of CREB mediated gene transcription in these transgenic finches. As observed with the transgenic mice expressing mutant CREBs(23), the expression of many genes were increased or decreased in DN and Actv birds compared to WT birds (Fig. S5). Specifically, in both DN and Actv birds, the expression levels of genes of which human homologues have CREs in the promoter region (CRE+ genes) were significantly changed, compared to those genes without CRE (CRE– genes; Figs. 2D and S5). Collectively, these results demonstrate that CREB-mediated gene transcription was misregulated in both DN-and Actv-CREB expressing transgenic finches.

Deficit of memory formation in zebra finch with mutated CREB. CREB plays a pivotal role in neuronal activity-dependent gene regulation and neural plasticity in various species(24–26, 40). For example, disrupting CREB function is known to suppress formation of long term memory in fear conditioned mice(41). To assess whether the transgenic manipulation of CREB activity has any effect on the memory formation in songbird, we analyzed the process of associative auditory memory formation in WT and the transgenic birds. To this end, a standard classical auditory conditioning paradigm was used to assess formation of memory after training. For
employing the classical fear memory conditioning in zebra finches, we developed an auditory song-conditioning test for songbirds (Fig. 3A; SI Material and Methods). In this test, a subject finch was isolated in a sound-proof chamber, and 5 songs of zebra finches, recorded from 5 different individuals unfamiliar to any of the subjects, were played through a speaker in a random order; after one particular song (CS; conditioned song stimulus), calls of a crow were presented (US; unconditioned stimulus), whereas the other songs (Cont.; control song stimulus) were followed only by an interval of silence. The presentation of a crow’s call caused freezing behavior (conditioned response), which was reflected in a significant decrease in their behavior (such as calling) throughout the training blocks (TB; Fig. 3B). These freezing behaviors to the crow’s call seemed to be intrinsic, because the zebra finches used in this study had never heard such calls before. At the 4th training block (TB4), WT birds began to decrease call behavior in response to CS, indicating that they began to associate the appearance of US with CS (Fig. 3C). By contrast, DN birds did not show any significant difference in their change in behavioral responses to CS compared with those to the control songs (DN; TB1, $P = 0.21$; TB4, $P = 0.67$; Student’s paired $t$ test; Fig. 3C). Actv birds did not show significant change in behavior, either (Actv; TB1, $P = 0.45$; TB4, $P = 0.60$; Fig. 3C). At the training blocks on the next day, the WT birds showed the conditioned response even at the beginning session (TB5), indicating that the memory was retained to the next day. In contrast, the DN birds failed to show conditioned responses, even at the final training block (DN; TB5, $P = 0.22$; TB8, $P = 0.92$; Fig. 3C); and the Actv birds showed conditioned responses only at the final block (Actv; TB5, $P = 0.12$; TB8, $P < 0.001$; Fig. 3C). Similar results were obtained when the presented song stimulus was changed to another set of songs, in order to exclude the possibility
that such behaviors were specific to a particular song (Figs. S6A, B). The decrease in the 
learning in DN and Actv-TgN birds was not due to motor defects or impaired auditory 
perception, because significant behavioral responses to US were observed for each 
genotype in every training block (Fig. 3B), indicating that the memory formation 
associating CS with US was specifically impaired in DN and Actv-TgN birds.

Although both DN and Actv birds displayed difficulties of song memory 
formation, we noticed a trend that DN and Actv birds react differently to US. Although 
presentation of US caused a significant reduction in call behavior (Fig. 3B) to DN and 
Actv birds, Actv birds showed significantly more number of calling, suggesting that 
they are behaviorally more active after the presentation of US compared to WT and DN-
TgN birds (Fig. S6C; WT, \( P < 0.0016 \); DN, \( P < 0.014 \); Tukey’s post-hoc analysis; two-
way ANOVA, \( F(2, 288) = 5.27 \), genotype factor, \( P < 0.0058 \)). Together, these data 
show that transgenic manipulation of the CREB transcriptional activity altered the 
memory formation in adult birds, consistent with a reported role of CREB in memory 
formation in other animal models(24–26).

At the beginning of the 4th training block (TB4), when the behavioral 
association between CS and US was observed, phosphorylation of CREB at serine 119 
(equivalent to serine 133 in mouse CREB), which activate its transcriptional 
activity(42), was detected in the brains of auditory conditioned WT birds. We observed 
pCREB signal in various brain regions. One of the brain regions that showed differential 
phosphorylation of CREB was the basal ganglia including Area X, the nucleus essential 
for song learning(43, 44) (Fig. S7A). During the conditioning, we did not observe the 
singing of the subjects; this phosphorylation was similarly observed when the subjects 
were auditory conditioned in a dark chamber, indicating that such phosphorylation was
not caused by singing\(^{(45)}\) (Fig. S7B). Rather, these signals may be caused by perception of auditory stimuli\(^{(28)}\), or by motor behavior of subjects such as vocalization of calls\(^{(46)}\), or by neuromodulators such as dopamine\(^{(47)}\). This phosphorylation was suppressed by injection of STO609 (20 \(\mu\)M), a selective inhibitor of Calmodulin-dependent protein kinase kinase (CaMKK), which is known to suppress the activity dependent phosphorylation of CREB through suppressing the activity of calcium-calmodulin-dependent protein kinase IV\(^{(48)}\), into the basal ganglia prior to the trainings (Fig. S7C).

The basal ganglia is known to play an important role in the learning of sequential motor behavior or in selecting the action in classical conditioning\(^{(49)}\), not only in rodents, but also in avian species\(^{(50, 51)}\). We therefore investigate the role of CREB activation in basal ganglia to the auditory conditioning formation. We injected vehicle or STO609 bilaterally into the basal ganglia of WT birds prior to the auditory conditioning task, and analyzed the effect of these manipulations on the memory formation (Figs. S7D–F). The injected drug seems to spread to parts of Area X and the surrounding striatum (Fig. S7C, SI Material and Methods). The injection of vehicle or STO609 did not affect the freezing response against the presentation of the unconditioned stimulus in the next day sessions (Fig. S7D). However, injection of STO609 prior to the conditioning training abolished the conditioned response in the next day (Fig. S7E; STO609; TB5, \(P = 0.12\); TB8, \(P = 0.41\), Student’s paired \(t\) test); whereas the vehicle-treated birds showed normal conditioned responses (Fig. S7E; Vehicle). Thus, the pharmacological method to suppress CREB-activation in a local brain structure reproduced the results of CREB transgenic animals, indicating that CREB was involved in the formation of the conditioned response in this experiment.
**Impaired vocal learning in transgenic zebra finches.** Next, we analyzed how the genetic manipulation of CREB activity affects song development, which requires social learning during the postnatal period. Postnatal song acquisition was assayed by use of our song training paradigm, which allows us to compare the accuracy with which song of a tutor (used repeatedly to different juveniles) can be copied by individuals of different genetic backgrounds (Fig. 4A; *SI Material and Methods*). We found that the relative qualities of acquired songs between the transgenic birds and WT birds, tutored by a common male tutor, were strongly affected by the genotype (Figs. 4B–F and S8A). Birds with DN-CREB expression developed songs with severely reduced similarity scores compared to WT controls (Fig. 4C; one-way ANOVA: $F(2, 85) = 4.10, P < 0.0004$; DN, $P < 0.0008$, Dunnett’s *post-hoc* test). On the other hand, birds with Actv-CREB expression developed songs that showed no significant difference compared with those of WT birds (Fig. 4C; Actv, $P = 0.87$). Both WT and Actv birds, but not the DN birds ($P = 0.11$), showed an increase in song quality during development (between 60 and 140 dph) (Fig. 4D). Song-similarity analysis based on the similarity of an entire motif (52), a stereotyped temporal sequence of syllables, also yielded a similar result (Figs. 4E, F and S8B). In contrast to songs, call is known to be acquired mainly intrinsically (15, 39), although some feature of calls are modified postnatally through learning (53). Notably, we did not observe any significant genotype effect on the similarity of tutee’s calls against the tutor’s calls (Figs. 5A, B and S8C; Kruskal-Wallis test, $P > 0.50$), nor the acoustic quality of calls (Fig. 5C). These results indicate that postnatally acquired behavior was specifically affected in the mutant CREB transgenic birds. Collectively, the findings demonstrate that genetic manipulation of intrinsic
factor, the activity of CREB, differentially affected the postnatal song development
even within shared environmental conditions.

Discussion

The transgenic technology has been applied to a wide variety of animal species to study
the effect of genetic involvement on animal behaviors and development. However,
because the early embryogenesis of avian species have specific features different from
other animals, generating transgenic avian species is still challenging(38, 54). Recently,
transgenic lines of zebra finch that express GFP was generated by injecting lentiviral
vectors into the early embryos(35). Using this approach, we obtained several lines of
zebra finches expressing mutated CREBs. In addition to the existing methods, the new
transgenic technology in songbirds will be a strong tool to study how specific genes
influence the acquisition of behaviors. For example, by generating transgenic zebra
finches with suppressed CREB activity, we showed that birds expressing DN-CREB
develop songs with poor copying quality of tutor’s songs (Fig. 4). Transgenic
manipulation of the genome allows a uniform expression of transgenes in the entire
population of cells involved in executing or learning certain behaviors. In a certain
situation, this method is better than other methods such as local injection of
pharmacological reagents or viral vectors into the brain, which are unable to control the
extent of diffusion or the efficiency of transfection among the cell population. For
example, the transgenic strategy seems to be particularly advantageous in the present
study because manipulations of the activity of CREB in a subset of the neuronal
population has been shown to lead compensation of the disturbed function by the
surrounding population that is not affected by the treatment(55). One caveat that should
be mentioned about our present study is the transgenic strategy employed in this study
results in the expression of transgenes in a wide population of neurons throughout
development, owing to the activity of the synapsin promoter used to express the
transgene. Another concern is the possibility of disrupting the expression of endogenous
genes especially near the integration locus of transgenes, which may cause some
difference of phenotypes among the transgenic lines. Further technical refinements
using transgenic strategies already applied in other animals, such as the application of
cell type specific promoters or inducible promoters, or knock-in of transgene into a
specific locus, may specify the neuronal circuitries or the time window of plasticity
involved in the development of the postnatally learned behaviors in songbirds.

Although the effect on the intrinsic phenotypes still needs to be analyzed, the
results presented here indicate that activation of the CREB signaling pathway is
essential for the proper song learning in postnatal periods. In other animal models,
CREB has been known to function as a positive regulator of memory formation by
regulating the activity dependent gene transcription in neurons(24, 26, 27, 40, 56).
Transgenic rodent models have shown that expression of dominant negative mutant of
CREB suppresses whereas constitutive active mutants enhance some forms of memory
acquisition(40, 56). Our results are basically in line with the idea that CREB is a key
molecule to regulate postnatal learning in animals, and provide evidence that songbirds
also utilize CREB to auditory and vocal learning of birdsongs. We observed that the
transgenic zebra finches expressing DN-CREB (DN; S119A) developed songs with
severely reduced copying accuracy. On the other hand, the birds expressing the active
form of the CREB mutant (Actv; Y120F) showed songs with comparable copying
accuracy to those of WT birds (Fig. 4). Since the song learning may be well-optimized
through the evolutionary processes of songbirds, it is possible that our experimental
methods did not allow the enhancement of song learning in Actv birds. Whether the
chronic enhancement of CREB activity actually shows enhanced or accelerated learning
of songs or auditory memory in a different context remains to be determined in further
studies. In contrast to song development in juvenile birds, we observed impaired
auditory memory formation in adult Actv birds, similarly to DN birds (Fig. 3). Genetic
manipulation of CREB function in rodents sometimes causes discrepancies in the
evaluation of behavioral studies, because enhancement of CREB phosphorylation in
mice does not necessarily result in the enhancement of memory formation, depending
on the expression level of proteins or the testing context(40, 57–59). In our hearing
threshold analysis (Fig. S3), although Actv birds showed normal auditory ability to
discriminate the change of sound volume, we noted that the reaction to the auditory
signal presentation was different; i.e., whereas WT and DN birds showed a reduction in
their number of calling behavior, Actv birds showed a significant increase in it (Fig.
S3C). Similarly, Actv birds were more active in their response to the unconditioned
stimulus (US) in the auditory conditioning test (Fig. S6C). The difference in the reaction
to sound stimuli may be partly attributable to the disturbed formation of conditioned
responses in the Actv-CREB birds.

Although the efficiency is still low, the transgenic technology has opened the
door to perform molecular genetic studies on songbirds(35). Additional technologies,
such as gene knock-out or conditional expression of transgenes, will clarify the
contribution of genes and environments, as well as how these are intermingled, to the
development of the sophisticated ability for song acquisition in songbirds. Because we can manipulate both genetic and environmental factors, as shown in this study, songbirds may provide valuable knowledge as to how environments affect the development or disorders of an animal’s behaviors.
**Materials and Methods**

Also see *SI Materials and Methods* for detailed descriptions.

**Animal care and treatment.** To generate transgenic zebra finches, freshly laid eggs were collected from nests, and lentiviral vectors were microinjected around the central portion of the embryos as described earlier (35). Germ-line transmission of transgene was analyzed by performing PCR-mediated genotyping of the offspring which were produced by crossing the virus-injected bird with WT birds (G1-generation). Transgene expression was further checked by RT-PCR and by immunostaining of brain sections.

**Song similarity analysis and Behavioral analysis.** At each developmental time point, birds were isolated in a sound-proof chamber; and their vocalizations were recorded with a microphone. The songs and calls were analyzed with Sound Analysis Pro 2011 (SAP2011), using the similarity batch mode. Behavioral analysis was done as described previously (34). Only male birds were used for the behavioral analysis. For the auditory conditioning, birds were isolated in a sound proof chamber and their responses against song presentation were video recorded. For normalization of the call behavior to the CS, the number of call responses during the 1-min period before the CS presentation was subtracted from the number of call responses of the 1-min period after (and during) the CS, and divided by the sum of the values before and after. For the US, the number of call behaviors during the 1-min period before the CS presentation was subtracted from the number of call responses of the 1-min period after (and during) the US, and divided by the sum of the values. Statistical analysis was performed using the paired t-test on raw values before and after the stimulus presentation (without normalization).
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Author contributions

K.A. conceived the project, performed all the experiments, analyzed data, and wrote the paper. S.M. assisted histological experiments and analyzed data. D.W. discussed the data and provided feedback on the manuscript.

The authors declare no conflict of interest.

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**Figure Legends**

**Fig. 1. Generation of the transgenic zebra finches with modified CREB activity.**

(A) Schematics of the transgenes used to generate transgenic zebra finches. SYN1, human synapsin 1 promoter; WPRE, woodchuck hepatitis virus post-transcriptional regulatory element. (B) PCR analysis of genome integration and expression of transgenes. Genome DNA (upper) and total RNA (lower) of WT and transgenic line (TgN) were collected from an adult bird from 3 transgenic lines for each. (C) Images of sagittal sections from WT and G1-TgN birds immunostained against EGFP showing the expression of transgene in G1-TgN birds. Scale bar, 1 mm. See Fig S1A for the anatomical profiles. (D) High-magnification image of nidopallium, showing signals of EGFP (green), and Neu-N (red, neuronal nucleus marker), and DAPI. Scale bar, 20 μm. (E) Quantitative RT-PCR analysis of RNA collected from WT and the transgenic birds. Data from primer sets which amplify endogenous CREB1 (right) and endogenous and exogenous CREB (left) are shown. Bar graph shows the mean ± SEM of relative expression values normalized to the WT. * P < 0.05, n.s. P > 0.5 Dunnett’s post-hoc test.

**Fig. 2. Transgenic zebra finches show misregulated CREB mediated gene transcription.** (A) Schematics of the lentivirus (LV) based transcription-reporter constructs. A constitutive PGK1 (human phospho-glycerate kinase) promoter expresses an infection-reference gene (H2B-flag, flag-tagged Histone-2B). In the other direction, a minimal promoter expresses a reporter gene (tbGFP-PEST, turboGFP fused to PEST sequence), whose expression is influenced by the presence of CREB binding sequence
(CRE), in the LV-CREB-reporter. (B) HEK293T cells transfected either with LV-CREB-reporter or LV-Control-reporter at the multiplicity of infection (MOI) 1, 0.1, 0.01, and were treated with vehicle (0.1% DMSO) or 100 μM forskolin in order to stimulate the cAMP dependent activation of CREB. Each reporter activity was quantified by dividing the amount of tbGFP-PEST by the amount of H2B-flag, both of which were quantified by quantitative RT-PCR. Stimulus dependent changes in CREB activity were calculated by dividing the reporter activity of LV-CREB-reporter by those of LV-Control-reporter. *\( P < 0.0001 \) against each vehicle treated cells. Bar graph shows mean ± SEM, \( n = 4 \) independent experiments. (C) Activity of CREB mediated gene transcription in transgenic birds. LV-CREB-reporter and LV-Control-reporter were injected into Area X and the reporter activities were quantified for each subject. WT, \( n = 10 \); DN, \( n = 6 \); Actv, \( n = 5 \) birds. Bar graphs indicate mean ± SEM, *\( P < 0.05 \), Dunnett’s post-hoc test. (D) Quantitative RT-PCR analysis of endogenous RNA collected from WT and transgenic birds (\( n = 11 \) birds for each genotype). Un-paired \( t \) test; Bar graph shows mean ± SEM of the absolute log\(_2\) value of relative amount of expression against WT, comparing gene with (\( n = 47 \)) and without (\( n = 32 \)) CREs. See Fig. S5 for details.

Fig. 3. **Deficits in auditory memory formation in transgenic zebra finches.** (A) Experimental timeline (upper panel) and the schematics of the experiment of 1 training block (lower panel). (B, C) Results of the auditory conditioning experiments. Behavioral reaction against control song stimulus (Cont., dotted lines) and unconditioned stimulus (US, solid lines; B) or conditioned song stimulus (CS, solid lines; C) are shown. Change in call behavior number after the presentation of stimuli (Cont., US, and CS) are
normalized and shown for each genotype (WT, left; DN, middle; Actv, right). Mean ± SEM are shown. Asterisks indicate a significant difference in the call response before and after the presentation of each stimulus, $P < 0.05$, Student’s paired $t$ test; WT, $n =$ 25; DN, $n = 25$; Actv, $n = 25$. See also Fig. S6 for the raw number of the call behavior, and the experiment performed with another set of song stimuli.

**Fig. 4. Acquisition of tutor’s song in transgenic zebra finches.** (A) Top panel: experimental timeline. Bottom panel: schematics of the experiment. Juvenile male birds were moved from their home cages and kept in a sound-proof chamber with a live male finch (tutor). The same tutor bird was used multiple times for comparisons. (B) Examples of the sonograph of the tutor bird’s song (Tutor’s song) and that of the 140-dph birds of different genotype, reared with the same tutor (Tutee’s song). (C) Similarity score of tutee’s songs at 140 dph, calculated from the similarity of each syllable. (D) Developmental changes in the similarity score. (E, F) Similarity score of tutee’s songs calculated from the similarity of the entire motif. * $P < 0.001$, one-way ANOVA, Dunnett’s post-hoc test, n.s., $P > 0.67$ against WT. Summarized values from 85 tutees (WT, $n = 39$; DN, $n = 23$; Actv, $n = 23$) tutored by 5 tutors are shown. Boxes and whiskers show the respective median and 25–75 and 10–90 percentiles. Line graphs shows mean ± SD.

**Fig. 5. Call vocalization in transgenic zebra finches.** (A) Examples of the sonograph of the calls of the 140-dph birds of different genotype. (B) Similarity scores between tutor’s and tutee’s call. (C) The differences of acoustic features of calls between the genotype. Summarized values from 85 tutees (WT, $n = 39$; DN, $n = 23$; Actv, $n = 23$)
tutored by 5 tutors are shown. Boxes and whiskers show the respective median and 25–75 and 10–90 percentiles. Line graphs shows mean ± SD, n.s., $P > 0.15$ against WT, Kruskal-Wallis test.