Two developmental modules establish 3D beak-shape variation in Darwin's finches

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Changes in two different developmental modules establish beak shape variation in
Darwin’s finches

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Running title: Modularity in beaks shapes of Darwin’s finches

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Abstract

Bird beaks display tremendous variation in shape and size, which is closely associated with the exploitation of multiple ecological niches and likely played a key role in the diversification of thousands of avian species (1). While previous studies described the molecular mechanisms that regulate morphogenesis of the prenasal cartilage (2, 3), which forms the initial beak skeleton, much of the beak diversity in birds depends on variation in the premaxillary bone, which forms later in development and becomes the most prominent functional and structural component of the adult upper beak/jaw (4). Here, we studied the varied beak shapes of Darwin’s finches to understand the development and evolution of the premaxillary bone. We show that TGFβ receptor type II, β-catenin and Dickkopf-3, the top candidate genes from a cDNA microarray screen, are differentially expressed in the developing premaxillary bone in embryos of Darwin’s finches, in close correlation with their beak shapes. Furthermore, functional analyses in chick embryos demonstrated that these molecules form a regulatory network shaping the morphology of the premaxillary bone, independently of the network controlling the prenasal cartilage. Our results show that beak morphology is established by two different modules, the prenasal cartilage (during early development) and the premaxillary bone (during late development). We demonstrate that multiple molecules regulate these two modules and can independently alter their growth along different axes, thereby increasing the ability of the beak developmental program to generate variation. This modularity in developmental program may be a general mechanism by which morphological diversity can evolve.
Introduction

Modern evolutionary developmental biology postulates that adaptive morphological changes in adult organisms ultimately originate by altering particular developmental programs (5, 6). Thus, exploring cases in which the developmental pathways responsible for evolutionary changes can be identified and characterized is pivotal to our understanding of the origin of morphological diversity (7-9). In this study, we aimed to understand how changes in developmental controls of a morphological trait may constrain or facilitate diversification. To this end, we focused on unraveling the molecular and developmental mechanisms responsible for patterning the differences in the shapes of avian beaks—which are usually associated with differences in diet and ecological niche—by taking advantage of the natural diversity of beak shapes in the iconic Darwin’s finches.

Bird beaks are three-dimensional structures that show a tremendous amount of variation in shape along the depth, width, and length axes. Variation in beak shape has profound impacts in the ability of an organism to survive and reproduce in the wild and, thus, has played a major role in the radiation of thousands of species of birds, one of the most successful classes of vertebrates (1, 10-12). Therefore, its adaptive significance coupled to the extreme levels of diversity observed in nature make this trait ideal for tackling developmental and evolutionary questions about morphological diversification.

Adult beak morphology is determined by the development of two components, the prenasal cartilage (pnc; the ethmoid process of the nasal septum) followed by the premaxillary bone (pmx) from a separate condensation. Previous studies of pnc formation have identified two signaling molecules, Bmp4 and CaM, that regulate early differences in beak morphogenesis (2, 3). Comparable studies of the pmx are lacking, and are greatly needed for three reasons. First, it is
the most prominent functional and structural component of the adult bird upper beak/jaw (4).

Second, much of beak diversity in birds depends on variation in the pmx (4). Third, it is not known if molecules such as Bmp4 and CaM have the same roles at this crucially important stage of development, or the extent to which pnc and pmx formation are independent modules. Here, we took advantage of the beak shape differences in closely related species of Darwin’s finches to understand how variation in the pmx is generated and address fundamental questions about the evolution of beak shape diversity.

Darwin’s finches (Thraupinae, Passeriformes) of the Galapagos and Cocos Islands comprise a monophyletic group of fourteen closely related species that represent a classic example of adaptive radiation, niche partitioning, and rapid morphological evolution (13-16). In a relatively short period of time, this group has evolved a diversity of bill shapes adapted to exploit specific food items, particularly under conditions of food scarcity (16). Within the monopyletic genus Geospiza, the small, medium and large ground finches (G. fuliginosa, G. fortis, and G. magnirostris, respectively), which we refer to as “ground finches” in this article, have evolved a series of deep and broad beaks used to crush seeds. This series of ground finches contrasts with the more elongated and narrow beak shapes used by the large cactus and cactus finches (G. conirostris and G. scandens) to feed on nectar and pollen (Fig. 1A).

Here, we used a microarray screen results and comparative gene expression analyses in Darwin’s finch embryos followed by functional experiments in the chicken model system to determine the molecular and developmental mechanisms responsible for patterning shape differences in the avian beak. Our results show that beak morphology is controlled by two different developmental modules, the pnc (during early development) and the pmx (during late development). We find that multiple molecules are involved in regulating these two modules and
can independently alter their growth along different axes, thereby increasing the level of morphogenetic variability and potential for evolutionary change.

**Results and discussion**

During beak development, the *pnc* and the *pmx* condensations are established when the beak primordia form (4). The prenasal cartilage is the first skeletal structure to mineralize and establish species-specific beak shapes during early embryonic development (2, 4). As revealed by the expression pattern of the chondrogenic marker *Col2a1*, at embryonic stage 27 (st. 27), the *pnc* occupies a large portion of the developing upper beak primordia and explains differences in beak shape of the large and medium ground finches at this stage (Fig. 1B). However, its relative contribution to forming overall beak dimensions is significantly diminished by st. 30 (Fig. 1B) (2). At this later stage, the *pmx* condensation begins to expand and it is this structure that will ultimately determine the species-specific differences in adult bird beaks (4).

According to recent mechanical models, the *pmx* is the principal element of the adult bird upper beak responsible for dissipating and distributing forces generated during consumption of hard seeds (17, 18). Correspondingly, our analyses of micro-CT scan data showed that the adult large and medium ground finches have considerably larger *pmx* volumes, relative to their size, than the cactus finches and are, thus, ideal for analysis of variation in the *pmx* (Fig. 1C and Table S1). To determine when the species-specific differences in *pmx* are first established, we examined the expression of alkaline phosphatase, an osteogenic marker, in embryos of five species from the genus *Geospiza* at two critical stages of beak development, st. 27 (E5.5) and st. 30 (E6.5) (2, 3). In the species with the largest *pmx* volume, the large ground finch, alkaline phosphatase was expressed in the condensation of the *pmx* earlier than in any other species (st.
27) indicating that this species undergoes a heterochronic shift in the osteogenesis of this tissue. At the later st. 30, the pmx condensation in the large and the medium ground finches expands to occupy most of the upper beak primordium and expresses higher levels of osteogenic markers than size-matched cactus finches (Fig. 1D and Fig. S1). Thus, the results from this analysis showed that differences in adult pmx volume in Darwin’s finches correlate with the time, strength and place of expression of osteogenic markers during embryonic development.

Previously, we showed that two different molecules, Bmp4 and Calmodulin (CaM), regulate growth along different dimensions of the developing beak in Darwin’s finches (depth/width and length, respectively) by patterning the pnc element (2, 3). However, our functional tests showed that Bmp4 and CaM do not regulate morphogenesis of the pmx (2, 3). To identify genes, in an unbiased manner, that might explain the variation seen in the pmx of different species we took advantage of the previously conducted cDNA microarray-based screen in which we directly compared expression of several thousand transcripts from st. 26 upper beak primordia in Darwin’s finches (3). We searched for transcripts whose expression levels would correlate with the beak shapes of the large and the medium ground finches as they have considerably deeper and larger pmx than cactus finches. We identified three transcripts, TGFβ receptor type II (TGFβIIr), β-catenin and Dickkopf-3 (Dkk3), that were expressed at 12-15 fold higher levels in the large ground finches than in the reference species, the sharp-beaked finch (G. difficilis) (Table S2). These three new candidates represented significant developmental pathways and were not housekeeping or ribosomal genes.

TGFβIIr is a serine/threonine protein kinase that upon ligand binding initiates a series of phosphorylation events that can lead to the regulation of gene transcription (19). TGFβIIr-dependent pathway is important for craniofacial skeletal development in mammals and mutations
in this gene are associated with certain human craniofacial abnormalities (20, 21) but its function in morphogenesis of bird beaks has not been previously reported. \( \beta \)-catenin is a subunit of the cadherin protein complex and an integral component in the Wnt signaling pathway (22). While nuclear translocation of \( \beta \)-catenin in the osteogenic cells is both required and sufficient for terminal bone cell differentiation, the relationship between its expression level and osteogenic potential is unknown (23). Dkk3 encodes a secreted protein and is the most divergent member of the Dkk family in terms of sequence and function (24). Dkk3 does not have a reported function in craniofacial or skeletal development and, unlike the other members of the Dickkopf family, is not known to regulate Wnt signaling (reviewed in ref. 24).

We observed a striking correlation between adult beak morphology and expression of our three new candidate genes. The three genes were expressed at both higher levels and in broader domains in the large and the medium ground finches than in cactus finches, especially in the large ground finch, in which all three genes were expressed in most of the dorso-distal part of the upper beak primordium that accommodates the pmx condensation (Fig. 2). More specifically, at st. 27, the three molecules were strongly expressed throughout most of the beak mesenchyme (except in the prenasal cartilage) in the large ground finches, whereas they were confined to a much smaller region in the size-matched large cactus finches (Fig. 2A). By st. 30, both the large and medium ground finches expressed these molecules at much higher levels and in broader domains in the osteogenic beak mesenchyme than the corresponding large cactus and cactus finches, respectively (Fig. 2B). Notably, TGF\( \beta \)IIr and \( \beta \)-catenin accumulated in a restricted domain at the distal beak region in the large cactus and cactus finches in contrast to the broad domains for these genes found in the large and medium ground finches (Fig. 2B).
To determine the functional significance of the observed correlations, we used the RCAS replication-competent retroviral vector in the chicken embryo model to mimic the broader and stronger expression patterns of TGFβIIr, β-catenin, and Dkk3 seen in the large and medium ground finches (Fig. 3). Infection with a constitutively active version of the TGFβ Type I receptor (RCAS::Alk5*); with a construct driving expression of the stabilized version of β-catenin (RCAS::CA-β-catenin); and with a construct carrying the full-length chick homolog (RCAS::Dkk3), all led to a significant increase in both beak depth and length, relative to the uninfected controls, whereas beak width remained relatively unchanged (Fig. 3A, B, D, E, G). Most if not all increase in beak dimensions resulted from changes in the pmx element, as revealed by chondrogenic and osteogenic markers (Fig. 3A, B, D, and E). In addition, when we infected chicken embryos with a dominant-negative construct to decrease the levels of TGFβ signaling (RCAS::TGFβrΔ), we found a significant decrease in beak depth and length, whereas there was little effect on beak width (Fig. 3A, C, G). Likewise, this decrease in depth and length was a result of the diminished pmx dimensions (Fig. 3A, C).

These results differed from the significant increase in beak depth and width observed when Bmp4 signaling is upregulated in the chick embryonic beak with the RCAS::Bmp4 viral construct (Fig. 3A, F). Previous studies have found that Bmp4 does not elongate the beak (2, 25). Relative to wing length, the beaks of the large and medium ground finches are proportionally deeper and broader than in the basal sharp-beak finch G. difficilis (26-28). Interestingly, their beaks are also longer and thus, Bmp4 activity alone cannot fully explain morphological variation in this axis (29). Misexpression of the three new candidate molecules did not produce a marked effect on the development of pnc (Fig. 3A-E), whereas increased levels of Bmp4 led to a drastic expansion of the cartilage element and a decrease in pmx production and dimensions (Fig. 3A, F).
Therefore, the effect of \textit{Bmp4} upregulation on the final beak shape must be indirect, perhaps by providing extensive matrix support for the nascent \textit{pmx} later in development when expression of this gene is shut off. Together, these data suggest that \textit{TGF\beta IIr}, \textit{\(\beta\)-catenin}, and \textit{Dkk3}, in good correlation with their spatial and temporal expression, act by positively regulating the size of the \textit{pmx}, thereby specifying the final beak morphology in the large and medium ground finches (2).

Since \textit{TGF\beta IIr}, \textit{\(\beta\)-catenin}, \textit{Dkk3} displayed largely overlapping domains of expression in the beak primordia and were co-expressed in many of the same mesenchymal cells (Fig. S2), they could potentially be regulating each other’s expression during beak development. To investigate this possibility, we analyzed the effects of misexpressing each candidate molecule on other candidate genes as well as on \textit{Bmp4} (Fig. 4A-D). We found that upregulation of the \textit{TGF\beta IIr}-dependent pathway or \textit{\(\beta\)-catenin} caused a strong upregulation of \textit{Dkk3} expression (Fig. 4A). Conversely, downregulation of the \textit{TGF\beta IIr} pathway produced a decrease in the expression of \textit{Dkk3}, suggesting that \textit{Dkk3} is downstream of both \textit{TGF\beta IIr} and \textit{\(\beta\)-catenin} pathways (Fig. 4A). Furthermore, our analysis also demonstrated that \textit{\(\beta\)-catenin}, \textit{TGF\beta IIr}, \textit{CaM}, and \textit{Bmp4} do not regulate each other’s expression (Fig. 4B-D) suggesting that all these molecules can regulate beak development independently by altering different axes of growth.

In summary, \textit{Bmp4} and \textit{CaM} play important roles in the early expansion of the \textit{pnc} skeleton in ground and cactus finches, respectively (2, 3). This sets the stage, likely indirectly, for the later morphogenesis of the \textit{pmx}, which is patterned through the coordinated action of a small network of unrelated but interacting regulatory molecules, \textit{TGF\beta IIr}, \textit{\(\beta\)-catenin}, and \textit{Dkk3} (Fig. 4E) (2, 3). This pattern is consistent with previous heritability studies of single populations, which demonstrated the polygenic nature of beak shapes in Darwin’s finches (16, 29). These three new candidates not only contribute to beak morphogenesis but their expression and
functions help to comprehensively explain the observed morphological differences among the species we studied. For example, during the evolution of the extremely specialized “grosbeak” shape in the large ground finch, its depth increased by 92% relative to the basal condition in *G. difficilis* and *Bmp4* with all the new candidates contribute to this depth increase. However, its length also increased, albeit much more slowly, by about 30%, and this change cannot be explained by action of *Bmp4* alone, which does not regulate growth along this axis (2). Likewise, beak width increased by 87% and such increase cannot be explained by the activity of bone-specific genes alone (Fig. 5).

Taken together, our results show how changes in expression of multiple molecules regulating the formation of two distinct developmental modules, the *pnc* (in early development) and the *pmx* (in late development), can generate the morphological variation observed in the beaks of Darwin’s finches. Our experiments revealed that the three axes of beak growth—depth, length, and width—are regulated differently at these two stages of development, thereby enhancing the ability of the beak developmental program to generate variation on which natural selection can act. Since all modern birds share the same overall beak skeletal structure, while differing remarkably in size, proportions and curvature, our results provide a general developmental framework for understanding how the great diversity of beak shapes observed in nature is brought about developmentally. We suggest that the evolution of beak diversity has involved coordinated changes in multiple tissues and pathways. This is an example of how changes in the modular developmental program of an adaptive trait may facilitate diversification and may represent a general mechanism by which morphological diversity can evolve.
Materials and methods

Darwin’s Finches embryo collection and preparation.

Embryos of Darwin’s finches were collected according to regulations established by the Galapagos National Park using methods described in detail elsewhere (30). A total of 33 embryos were analyzed: *Geospiza magnirostris* (st. 27, n = 3; st. 30, n = 3), *G. fortis* (st. 27, n = 4; st. 30, n = 5), *G. fuliginosa* (st. 27, n = 3; st. 30, n = 3), *G. scandens* (st. 27, n = 3; st. 30, n = 5), *G. conirostris* (st. 27, n = 2; st. 30, n = 2).

In situ hybridizations and Immunohistochemistry

*In situ* hybridizations were performed as described before (30) using the *in situ* hybridization antisense probes for chicken. For immunohistochemistry, sections were blocked with 3% Bovine Serum Albumin (BSA) in PBS containing 0.1% Triton-X 100 for 1 hour, incubated overnight with primary antibody at 4 °C, washed in PBS, incubated for 1hr with secondary antibody, and washed with PBS. Immunostaining was performed using anti-*TGFβIIr* (sc-400; Santa Cruz), anti-*TGFβ1, β2, β3* (sc-146, sc-90, sc-82, respectively; Santa Cruz), anti-*β-catenin* (610153; BD Transduction Laboratories), and anti-*Dkk3* (kindly provided by Dr. Christof Niehrs). Antibodies were used at dilutions of 1:50 – 1:200. Reactions were visualized with Alexa Dye conjugated secondary antibodies (Molecular Probes) at 1:500 dilution in 3% BSA/PBS/Triton-X 100. For controls, sections were incubated with PBS instead of primary antibodies but no specific cellular staining was observed.

*Alkaline phosphatase*. Embryos were blocked with 3% Bovine Serum Albumin (BSA) in PBS containing 0.1% Triton-X 100 for 1 hour and incubated with an AP-conjugated secondary
antibody (Jackson ImmunoResearch). The signal was detected using a combination of NBT and BCIP to produce a purple precipitate.

**Functional experiments in chick embryos**

CDNA fragments containing a constitutively active form of the TGFβIr (Alk-5) (Addgene plasmid 14833) (31), a dominant negative form of TGFβIIr (Addgene plasmid 1176) (32), and the entire coding region of the chicken Dkk3 (gift of Dr. Chris Niehrs) were cloned into the SLAX-13 vector and then subcloned into RCAS(BP)A using methods described before (33). RCAS::CA-B-catenin and RCAS::Bmp4 constructs have been described previously (34, 35). Viruses were harvested, concentrated, and titered using methods described before (33). Fertilized eggs were obtained from SPAFAS (Norwich, CT), incubated at 37 °C, and staged according to Hamburger and Hamilton (36). Frontal nasal processes were infected at st. 24 and embryos were collected at stage 36. Embryos were fixed overnight in 4% paraformaldehyde, washed in PBS, stained with SYBR-safe (Invitrogen) and photographed under UV light under a Zeiss Discovery v8 Stereoscope (Carl Zeiss Inc). Frontal and lateral images were taken for each head, and measurements of different upper beak parameters were recorded using the Axiovision 4.6.3 software (Carl Zeiss Inc). Measurements were obtained in triplicate as follows: *Length*: from anterior part of nostril to posterior tip of egg tooth; *Depth*: perpendicular line passing through the anterior part of the nostril; *Width*: distance between the nostrils. The extent of viral infection was assayed by in situ hybridization with a viral specific probe (RSCH). Upregulation of β-catenin and of Dkk3 was monitored using the antibodies and in situ hybridization probe described above.

**(micro) Computed Tomography (CT) scans**
Darwin’s finches specimens were scanned at the Harvard CNS facility using an X-Tek XRA-002 micro-CT imaging system set at 75 kV. Specimens were mounted on a rotating table and a series of 3142 projections of 2000 by 2000 pixels covering 360 degrees was recorded. Volume and surface rendering was performed using AMIRA 5.0 (64-bit version, Computer Systems Mercury) for all specimens and the volume of the upper beak was extracted. As species differ in their body and head size rendering comparisons across species difficult, we calculated a multivariate indicator of overall size. To do so we used the $\log_{10}$ transformed wing chord length, tarsus length, head length, head width and head depth for each specimen as input into a factor analysis which is resulted in a single new factor hereafter referred to as ‘size’. $\log_{10}$ transformed beak volume was then regressed against ‘size’ and unstandardized residuals were extracted for comparison.

**Darwin’s finches microarray screen**

Details of the microarray production and data analysis are described elsewhere (3) and in the Supplementary Information.

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Fig 1. Contribution of the different skeletal structures to beak variation in Darwin’s finches. (A) Ground finches (G. fuliginosa, G. fortis and G. magirostris) have deep beaks used to crack seeds. In contrast, cactus finches (G. scandens and G. conirostris) use their elongated beaks to feed on pollen and nectar from flowers. (B) At stage (st.) 27 the prenasal cartilage (pnc) condensation, labeled with Col2a1, occupies a significant portion of the beak primordia in the large and medium ground finches, where is patterned by Bmp4 and Calmodulin (CaM). By st. 30, the pnc, labeled with Col2a1, occupies a smaller proportion of the beak relative to the developing premaxillary bone (pmx) and its role in generating species-specific beak morphologies decreases. (C) Micro-Computer Tomography (CT) scans indicate that interspecific variation in Darwin’s finches is caused mainly by differences in the amount of the pmx. Adults large and medium ground finches have larger pmx volumes than their size-matched large cactus and cactus finches, respectively. (D) These differences in adult pmx volume correlate with the
expression of the osteogenic marker alkaline phosphatase (*Alk. Phosp.*.) during embryonic development. In the large ground finch, alkaline phosphatase is expressed in the condensation of the *pmx* earlier than in any other species (st. 27). By st. 30, this marker is expressed at higher levels and in larger domains in the large and medium ground finches than in the size-matched large cactus and cactus finches, respectively. Arrow colors in B-D indicate species that have comparable body sizes but differ in beak morphology. Scale bar: 0.15 mm (stage 27) and 0.2 mm (stage 30) in (B); 0.2 mm in (C). Molecular tree from ref 27. Abbreviations: *pmx*, premaxillary bone; *pnc*, prenasal cartilage. *Pmx* volumes are corrected for body size.

**Fig. 2.** Variation in the premaxillary bone (*pmx*) in *Geospiza* correlates with the expression of *TGFβIIr*, *β-catenin* and *Dkk3*. (A) In the large ground finch, the skeletogenic condensation for the *pmx* appears earlier (st. 27) than in the other species showing a strong correlation with the earlier and broader expression of *TGFβIIr*, *β-catenin*, and *Dkk3*. At st. 30, the large and medium
ground finches have high expression levels of *TGFβIIr, β-catenin* and *Dkk3* in strong correlation with the volume of the developing *pmx*. Arrow colors in A and B indicate species that have comparable body sizes but differ in beak morphology. Scale bar: 0.1 mm in (A) and 0.2 mm in (B). Images of skulls are from ref. 15, with permission from the author. Abbreviations: *pmx*, premaxillary bone; *pnc*, prenasal cartilage
Fig. 3. Functional analysis of TGFβIIr, β-catenin, and Dkk3 in the chicken model system.

(A-F) Ultraviolet pictures of embryonic day 11 (HH st. 37) (A) wildtype chicken embryos and embryos infected with (B) RCAS::Alk5+, (C) RCAS::TGFβIIrΔ, (D) RCAS::CA-β-catenin, (E) RCAS::Dkk3, and (F) RCAS::Bmp4 constructs. We used RSCH, PTHrP-Rec, Col I and Col II, probes to reveal RCAS infection (RSCH), early osteoblasts (PTHrP-Rec), (overall bone (Col I)
and chondrocytes ($Col2a1$). Blue arrows indicate lower expression relative to wildtype specimens, red arrows indicate higher expression, and black arrows indicate no change.

(G) Histogram showing beak variation in wildtype and RCAS-infected chicken embryos. Embryos infected with RCAS::$Alk5^*$, RCAS::$TGF\beta r\Delta$, RCAS::$CA-\beta$-catenin, and RCAS::$Dkk3$ showed a significant change in their depth and their length relative to wild-type controls whereas the width remained unchanged (RCAS::$Alk5^*$: $n = 8$; $\mu_{\text{depth}} = 53.12 \pm 1.2$ (± s.d); $\mu_{\text{length}} = 51.39 \pm 2.06$; $\mu_{\text{width}} = 24.05 \pm 2.03$; $P_{\text{depth}} = 7.65 \times 10^{-6}$; $P_{\text{length}} = 0.0002$; $P_{\text{width}} = 0.53$; RCAS::$TGF\beta r\Delta$: $n = 9$; $\mu_{\text{depth}} = 36.49 \pm 2.85$; $\mu_{\text{length}} = 40.35 \pm 4.49$; $\mu_{\text{width}} = 22.43 \pm 1.07$; $P_{\text{depth}} = 3.32 \times 10^{-8}$; $P_{\text{length}} = 0.007$; $P_{\text{width}} = 0.1663$; RCAS::$CA-\beta$-catenin: $n = 9$; $\mu_{\text{depth}} = 52.92 \pm 1.51$; $\mu_{\text{length}} = 48.91 \pm 1.86$; $\mu_{\text{width}} = 24.11 \pm 0.82$; $P_{\text{depth}} = 1.21 \times 10^{-5}$; $P_{\text{length}} = 0.0079$; $P_{\text{width}} = 0.3364$; RCAS::$Dkk3$: $n = 15$; $\mu_{\text{depth}} = 54.39 \pm 1.76$; $\mu_{\text{length}} = 52.15 \pm 2.76$; $\mu_{\text{width}} = 24.43 \pm 1.19$; $P_{\text{depth}} = 2.02 \times 10^{-8}$, $P_{\text{length}} = 1.04 \times 10^{-5}$; $P_{\text{width}} = 0.1239$; WT: $n = 9$; $\mu_{\text{depth}} = 47.8 \pm 1.93$; $\mu_{\text{length}} = 45.6 \pm 2.59$; $\mu_{\text{width}} = 23.44 \pm 1.83$). Scale bar: 200 mm in whole-head images and 0.4 mm in sections (A-F).
**Fig. 4. Interaction of genes regulating beak development.** (A) Increasing levels of *Bmp4* didn’t have an effect on the expression of *Dkk3* relative to uninfected wildtype controls. However, upregulation of the *TGFβ* pathway and of *β-catenin* led to more cells expressing *Dkk3* relative to uninfected embryos. Conversely, downregulation of the *TGFβ* pathway caused a decrease in the number of cells expressing *Dkk3*. (B) Neither alterations in the *TGFβ* signaling pathway nor upregulation of *β-catenin* or *Dkk3* caused changes in the expression of *Bmp4* relative to wildtype embryos. (C) Similarly, alteration of *TGFβ* signaling and upregulation of *Bmp4* and *Dkk3* did not have an effect in the expression of *β-catenin* relative to wildtype embryos. (D) Likewise, upregulation of *β-catenin, Dkk3* or *Bmp4* did not have an effect in the expression of *TGFβIIr* compared to wildtype embryos. (E) *Bmp4* and *CaM* act independently to alter the growth of the prenasal cartilage whereas *TGFβIIr, β-catenin*, and *Dkk3* regulate the premaxillary bone. Scale bar: 0.4 mm in (A) and (B) and 0.2 mm in (C) and (D).
Fig. 5. The distinct beak morphologies in *Geospiza* are generated by differences in the strength, time, and place of expression of the molecules involved in beak development. (A) Species with deep beaks, such as the large ground finch, *G. magnirostris*, express higher levels of *Bmp4*, *TGFβIIr*, *β-catenin*, and *Dkk3*, whereas species with elongated beaks, such as the large cactus finch, *G. conirostris*, express higher levels of *CaM*. (B) Through their action on different skeletal tissues, the different molecules modify independent dimensions of growth and thereby, pattern the species-specific beak morphologies of Darwin’s finches. (C) The beak of the sharp-beaked finch, *G. difficilis*, represents a basal morphology for *Geospiza* (27, 28). Expression and function of *Bmp4*, *CaM*, *TGFβIIr*, *β-catenin*, and *Dkk3*, explain changes in the different beaks.
dimensions of the more derived species. Symbols used: + (positive effect); 0 (no effect); -
(negative effect). Abbreviations: pmx, premaxillary bone; pnc, prenasal cartilage. Measurements
in c were taken from ref. 32, corrected for wing length, and correspond to averages from males
that were collected in the islands where we obtained our samples. For the G. difficilis reference,
the analysis was performed with different populations (26) and since all the results showed the
same trend we used the population from Wolf because it represents one of the most basal
populations (28).