



Spiny and soft-rayed fin domains in acanthomorph fish are established through a BMP-gremlin-shh signaling network

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With over 18,000 species, the *Acanthomorpha*, or spiny-rayed fishes, form the largest and arguably most diverse radiation of vertebrates. One of the key novelties that contributed to their evolutionary success are the spiny rays in their fins that serve as a defense mechanism. We investigated the patterning mechanisms underlying the differentiation of median fin *Anlagen* into discrete spiny and soft-rayed domains during the ontogeny of the direct-developing cichlid fish *Astatotilapia burtoni*. Distinct transcription factor signatures characterize these two fin domains, whereby mutually exclusive expression of *hoxa13a/b* with *alx4a/b* and *tbx2b* marks the spine to soft-ray boundary. The soft-ray domain is established by BMP inhibition via *gremlin1b*, which synergizes in the posterior fin with *shh* secreted from a zone of polarizing activity. Modulation of BMP signaling by chemical inhibition or *gremlin1b* CRISPR/Cas9 knockout induces homeotic transformations of spines into soft rays and vice versa. The expression of spine and soft-ray genes in nonacanthomorph fins indicates that a combination of exaptation and posterior expansion of an ancestral developmental program for the anterior fin margin allowed the evolution of robustly individuated spiny and soft-rayed domains. We propose that a repeated exaptation of such pattern might underly the convergent evolution of anterior spiny-fin elements across fishes.

fin spine | acanthomorph | evolutionary key innovation | evo-devo | exaptation

Teleost fishes comprise ~50% of extant vertebrate species and display an astonishing diversity in body plans (1–4). Among the ~30,000 species of teleosts, the spiny-rayed fish—or *Acanthomorpha*—are evolutionarily the most successful lineage with over 18,000 species, representing approximately one third of all living vertebrates (1, 2, 5). Spiny-rayed fishes evolved relatively recently, during the Early Cretaceous (133 to 150 Mya) (6), and underwent their primary radiation after the Cretaceous–Paleocene mass extinction (ca. 66 Mya) when their lineage came to dominate many aquatic ecosystems (4, 6–10). One of the characteristics that has strongly contributed to the ecological and evolutionary success of the spiny-rayed fishes is fin spines in dorsal and anal median fins (2, 3, 11). Acanthomorph fin spines are mostly present on the anterior part of the dorsal, anal, and sometimes pectoral and pelvic fins and differ from soft rays by increased ossification, lack of segmentation, fusion of lateral half-segments (hemitrichia), and ending in a sharp point instead of bifurcating (11) (Fig. 1A). The main function of fin spines is to serve as a defense mechanism against gape-limited predators (2, 3, 11, 12), and as such, they strongly suggest a causal link to the success of the *Acanthomorpha*. Interestingly, anterior spines have evolved independently in other successful lineages of teleosts (2, 11–13), such as the Ostariophysians, in particular catfish and carps, underscoring their adaptive significance. However, in none of these lineages has this resulted in such persistent and pronounced individualization and modularization of separate median fin domains as present in acanthomorph fishes (*Discussion*).

In the acanthomorph dorsal and anal fins, the spiny and soft-rayed parts form distinct morphological and developmental units that behave as separate evolutionary modules (14). Examples of extreme morphological specialization of the spiny fin as compared to the soft rays are the Remora’s suction disk (15, 16), the Frogfishes’ ilicium/esca complex (17) and the dorsal part of the Triggerfishes’ “locking mechanism” (18). Species such as the Asian leaf fishes (e.g., *Nandus oxyrhynchus*) (19) further exemplify the divergence between spiny and soft-rayed fins. As many ambush-hunting fish, they have translucent soft-rayed fins and heavily pigmented spiny fins, whereby the transparency of the unpigmented soft rays enhances camouflage as slight undulations of those fin parts serve to keep the fish stationary. Altogether, this suggests a modularization that is distinct beyond the mere morphological difference between spines and soft rays and also determines pigmentation as well as function and further underscores the adaptive significance of individuated spine and soft-ray fin modules.

This individuation that affects a range of phenotypic traits is reminiscent of anatomical modules determined by master control genes that specify different ontogenetic outcomes for serially homologous elements. Examples of such systems are for instance the *hox* codes in the axial skeleton (20–22) or the hindbrain (23).

Significance

The “spiny fin,” comprising the anterior part of the dorsal and anal fins, is an evolutionary novelty that contributed to the success of the spiny-rayed fishes. This domain contains heavily ossified spines that serve as defense mechanism and differ from the posterior flexible soft rays. We show that the partitioning of the median fins into spines and soft rays is established through canonical developmental mechanisms responsible for the anterior–posterior patterning of appendages. Furthermore, the coloration of the anal fin in males appears to be genetically linked to soft-ray identity. Comparative analysis including nonacanthomorph fins indicates that the convergent evolution of fin spines across fishes likely involved the repeated exaptation of a deeply conserved developmental program from the anterior fin.

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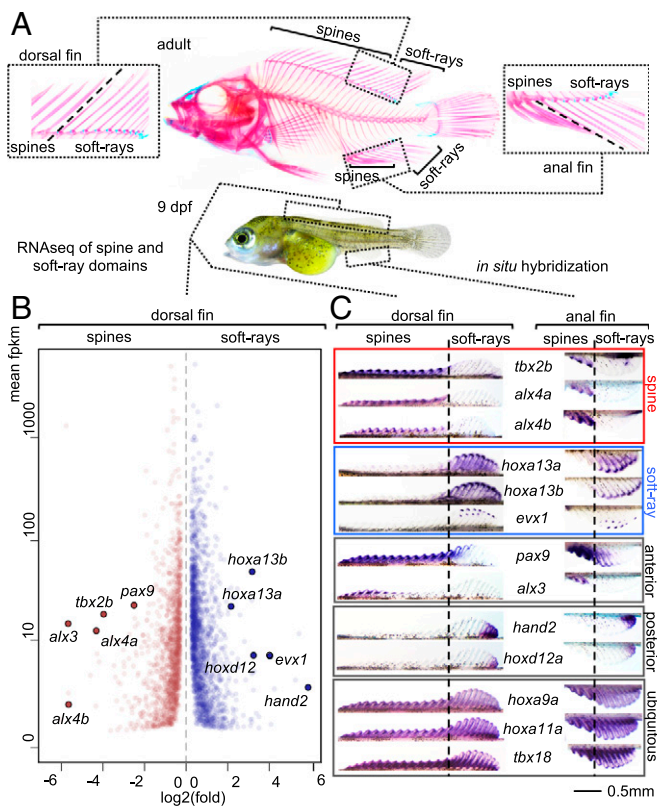


Fig. 1. (A) Skeleton of adult *A. burtoni* shows division of the dorsal and anal fins into spine and soft-ray domains. *Left Inset* shows transition of spiny and soft-ray domains for the dorsal fin, *Right Inset* for the anal fin. We investigated the developmental basis for the individuation of the spiny and soft-rayed fin domains using RNA-seq analysis (B) and in situ hybridization (C) using 8 to 9 dpf embryos. *Tbx2b* and *alx4a/b* define the fin spine territory while *hoxa13a/b* and *evx1* are expressed exclusively in soft rays. Additional transcription factor genes identified in the RNA-seq show an anterior (*alx3/pax9*) or posterior (*hand2/hoxd12*) bias but do not segregate with the spine to soft-ray transition. Expression of *hoxa9a*, *hoxa11a*, and *tbx18* is ubiquitous throughout the fins and underscores shared origins of spines and soft rays. Anterior is to the left. The skeleton and embryo in A are from ref. 25.

Thus, selector genes act upstream in the hierarchy of differentiation to initiate alternative downstream developmental trajectories for meristic structures (24).

In this work, we set out to unravel the developmental basis underlying the patterning of discrete spiny and soft-ray domains using the direct-developing cichlid fish *Astatotilapia burtoni* (25). Cichlids belong to the *Acanthomorpha* and possess a spiny fin. The established model-system zebrafish and medaka are not suited to address this question because zebrafish is not an acanthomorph, and medaka has secondarily lost the spiny fin. *A. burtoni* has the typical division of spine and soft-ray territories in dorsal and anal fins, as well as soft-ray-specific pigment pattern in males (egg spots). An understanding of the genetic basis for the specification of spine and soft-ray domains will help to elucidate the evolutionary origin of these modules at the base of the acanthomorph radiation as well as provide insight into how spines repeatedly emerged across fish clades as a diversity promoting trait.

Results

Mutually Exclusive *alx4/tbx2b* and *hoxa13a/hoxa13b* Expression Marks the Spine to Soft-Ray Boundary. We previously described the ontogeny of the spiny and soft-rayed domains in the dorsal and anal fins of *A. burtoni*. Spine and soft-ray territories differentiate simultaneously between 4 to 10 dpf (days postfertilization) from

continuous *Anlagen* located along the dorsal and ventral midline (25). The development of fin elements as either soft rays or spines reflects a binary developmental trajectory since intermediate forms do not occur. The partitioning of the fins into two morphologically discrete domains therefore suggests the existence of a code of “master control” genes that direct a developmental choice for the differentiation into soft rays or spines. We performed RNA-sequencing (RNA-seq) on prospective spiny and soft-rayed parts of the dorsal fin of 9 dpf embryos to identify differentially expressed transcription factor genes (Fig. 1 A and B). In the soft-rayed posterior part of the fin, *hoxa13a*, *hoxa13b*, *hoxd12*, *hand2*, and *evx1* are strongly up-regulated, while the spiny part of the fin shows strong expression of *alx4a*, *alx4b*, *alx3*, *tbx2b*, and *pax9*. To determine their specificity for spiny or soft-rayed fin domains, the expression of these genes was analyzed using whole mount in situ hybridization (Fig. 1C). In both dorsal and anal fins, we find a strong association of *hoxa13a/b* and *evx1*, and their anterior limit of expression marks the spine to soft-ray boundary. In line with its function in zebrafish (26), *evx1* is expressed in the forming segment boundaries of the soft rays. *Hoxd12* and *hand2*, however, associate with a more posterior part of the fin, away from the spine to soft-ray boundary. *Alx4a*, *alx4b*, and *tbx2b* associate with the spiny part of the fin and posteriorly demarcate the spine to soft-ray boundary. *Pax9* is expressed with an anterior bias but clearly overlaps the soft-ray territory while *alx3* is expressed in the anterior-most part of the spiny domain. Additional fin patterning genes *hoxa9a*, *hoxa11a*, and *tbx18* show ubiquitous fin expression and indicate a largely shared developmental program of the two fin domains, consistent with the spiny fin being a relatively young evolutionary modification. Analysis in a time series from 4 to 7 dpf shows that from 5 dpf onwards *alx4a* and *hoxa13a/hoxa13b* stably delineate spine and soft-ray domains whereas this is the case for *tbx2b* from 6 dpf onwards (SI Appendix, Fig. S1).

BMP Inhibition through *gremlin1b* Establishes the Soft-Ray Territory in Synergy with *shh*.

The division of fins into spiny and soft-ray domains reflects an anterior–posterior organization of the median fins. Therefore, we set out to investigate the role of canonical signaling mechanisms used to pattern the anterior–posterior axis of the appendages in the establishment of this division. In limbs and fins, *sonic hedgehog* (*shh*) secreted from a ZPA (zone of polarizing activity) is essential for correct anterior–posterior patterning (27–32). *Shh* expression in a posterior ZPA is an ancestral feature of gnathostome paired and median fins (27, 28, 33, 34). In *A. burtoni* dorsal and anal fins, we observe first *shh* expression in a ZPA starting at 5 dpf, becoming strongly expressed at 6 dpf, after which *shh* disappears from the ZPA and becomes expressed in the distal tips of the forming soft-ray and spine elements (Fig. 2B and SI Appendix, Fig. S2). Treatment during 4 to 6 dpf with the *shh* agonist SAG induces an anterior expansion of *hoxa13b* in the dorsal and anal fins while the expression of *alx4a* and *tbx2b* becomes more anteriorly restricted—indicating an anterior shift of the spine to soft-ray boundary (Fig. 2C and SI Appendix, Fig. S3). Analysis of the expression of *gli1*, which is a downstream target of *shh* and provides a read out for the range of *shh* signaling (27, 28), suggests that in untreated embryos at 6 dpf *shh* signaling extends anterior of the ZPA for the length of about two to three somites (Fig. 2B). That is, less than half the extent of the forming soft-ray domain, which develops over the width of 6 to 7 somites (SI Appendix, Fig. S1). Furthermore, inhibition of *shh* through treatment with the *shh* antagonist cyclopamine from 4 to 6 dpf fully abolishes *gli1* expression but does not lead to a strong displacement of the anterior–posterior position of the spine to soft-ray boundary as indicated by *alx4a/tbx2b* and *hoxa13b* expression (Fig. 2C and SI Appendix, Fig. S3) (although the expression levels of *hoxa13b* are decreased within the prospective soft-ray domain). Thus, this suggests that while *shh* appears capable of expanding the soft-ray

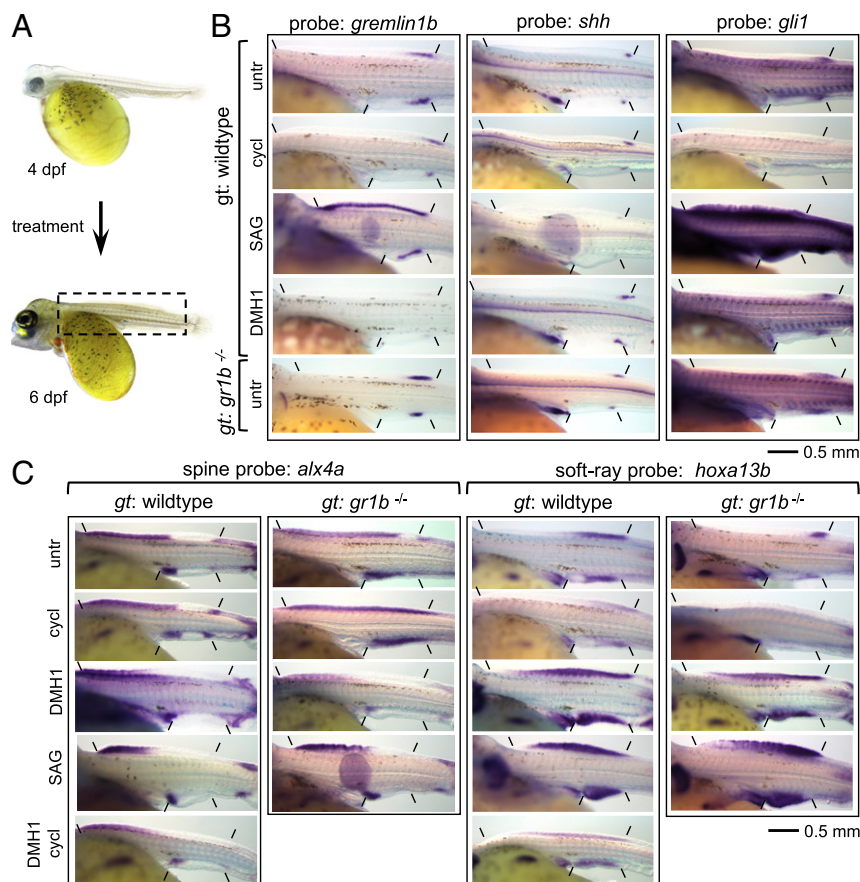


Fig. 2. The effect of interference with *shh* and BMP signaling on the establishment of spine and soft-ray territories. (A) All expression domains were analyzed at 6 dpf, all chemical treatments were initiated at 4 dpf. *Inset* in 6 dpf embryo shows the approximate extent of the part of the embryos shown in B and C. (B) Analysis of the expression of *gremlin1b*, *shh*, and *gli1* (indicated “probe”) in WT embryos treated with cyclopamine, SAG, DMH1, and in *gremlin1b*^{-/-} embryos. (C) Analysis of the spine to soft-ray transition using the spine marker *alx4a* and the soft-ray marker *hoxa13b* in treatment with cyclopamine, SAG, DMH1, and a combination of DMH1 and cyclopamine. Experiments were performed on WT and *gremlin1b*^{-/-} embryos (except the combination of DMH1 + cyclopamine) and observed on a minimum of 6/6 embryos per probe per treatment/genotype. Anterior and posterior limits of dorsal and anal fin *Anlagen* are indicated. gt: genotype, untr: untreated, cycl: cyclopamine. Anterior is to the left. The embryos in A are from ref. 25.

territory, the normal specification of the soft-ray domain occurs at least in part through another, *shh*-independent, mechanism.

In limbs, *shh* activates the secreted BMP antagonist *gremlin1* (35), which together with BMP4 provides a mechanism downstream of *shh* to regulate digit identity (36). In the dorsal and anal fins of *A. burtoni*, *gremlin1b* becomes expressed at 4 dpf. In the dorsal fin, its expression initially extends anterior of the vent but becomes subsequently restricted to approximately the soft-ray territory at 5 dpf and continues to regress more posteriorly during 6 and 7 dpf (Fig. 2B and *SI Appendix*, Fig. S2). No expression of the other gremlin homologs *gremlin1a*, *gremlin2*, or any other BMP antagonists investigated (*noggin1*, *noggin2*, *chordin*, *chordin-like-2*) was detected during early fin development (*SI Appendix*, Fig. S4). *BMP4* is expressed throughout the fin, although there may be a bias toward higher expression in the *gremlin1b* territory (*SI Appendix*, Fig. S2). We investigated the function of *gremlin1b* and BMP signaling during fin patterning by generating *A. burtoni* CRISPR/Cas9 knockout lines (*Materials and Methods*) and through gain of function by mimicking the inhibitory effect of *gremlin1b* with the small molecule BMP-receptor inhibitor DMH1. *Gremlin1b* knockout leads to a more posteriorly restricted *hoxa13b* domain of approximately the size of the *shh* signaling zone as inferred from *gli1* expression (Fig. 2C). At the same time, *alx4a* and *tbx2b* domains become expanded posteriorly, altogether indicating a posterior shift of the spine to soft-ray boundary. The gain of function approach induced the opposite effect with an anteriorly

expanded *hoxa13b* domain and anteriorly shifted *alx4a* and *tbx2b* domains (Fig. 2C and *SI Appendix*, Fig. S3). These anterior shifts are also induced in the *gremlin1b* knockout treated with DMH1, which therefore rescues the posterior shifts observed in untreated *gremlin1b*^{-/-} embryos (Fig. 2C). The posteriorized and anteriorized spine to soft-ray boundaries observed in *gremlin1b*^{-/-} and DMH1-treated embryos are maintained during development (*hoxa13b*-, *hoxa13a*-, *alx4a*-, and *tbx2b*-stained embryos shown at 9 dpf in *SI Appendix*, Fig. S3). Therefore, BMP inhibition in the posterior fin by *gremlin1b* is required for the delimitation of the *alx4a/tbx2b* and *hoxa13b* domains and influences the anterior–posterior position of the spine to soft-ray boundary.

In tetrapod limbs, *gremlin1b* is activated by, and acts downstream of, *shh* signaling (35). In *A. burtoni*, SAG treatment leads to widespread up-regulation of *gremlin1b* expression (Fig. 2B). Cyclopamine treatment however, reduces but does not eliminate posterior *gremlin1b* domain (Fig. 2B). This observation of *shh*-independent *gremlin1b* expression is consistent with its early activation at 4 dpf before *shh* in the ZPA becomes detectable (*SI Appendix*, Fig. S2). BMP inhibition by DMH1 strongly down-regulates *gremlin1b* expression, whereas it appears locally up-regulated in the *gremlin1b*^{-/-} embryos (Fig. 2B). This suggests that, as in limbs, BMP and *shh* are upstream of *gremlin1b* (36) but that in median fins these signaling pathways act in part redundantly. In the context of auto- and cross-regulatory interactions of these pathways, we observe that *shh* in the ZPA is

strongly down-regulated with SAG treatment and up-regulated with cyclopamine treatment, suggesting the presence of an autoregulatory negative feedback loop (Fig. 2B) as has also been observed during limb development (37). Furthermore, DMH1 treatment slightly enhances *shh* expression in the ZPA but does not increase signaling (as judged by *gli1* expression) to an extent that it explains the far anterior shift of the soft-ray to spine boundary (Fig. 2B). Altogether, these experiments suggest that *shh* and *gremlin1b* are acting independently upstream of the specification of the soft-ray domain.

We further tested this hypothesis by combining *shh* activation and inhibition conditions with *gremlin1b* knockout and BMP inhibition. Embryos treated with a combination of cyclopamine and DMH1 display a similar expansion of *hoxa13b* and reduction of *alx4a* and *tbx2b* domains as treatment with DMH1 alone (Fig. 2C and *SI Appendix, Fig. S3*), showing that BMP inhibition can posteriorize the fin independently of *shh*. In *gremlin1b*^{-/-} embryos treated with cyclopamine, the posterior residual patch of *hoxa13b* expression disappears completely and *alx4a* and *tbx2b* domains now extend throughout the length of the dorsal and anal fin, indicating a complete absence of a soft-ray domain (Fig. 2C and *SI Appendix, Fig. S3*). *Gremlin1b* knockout embryos treated with SAG resemble wild-type (WT) embryos treated with SAG (Fig. 2C and *SI Appendix, Fig. S3*), confirming that *hoxa13b* expansion and *alx4a/tbx2b* reduction can occur independent of BMP inhibition by *gremlin1b*. Therefore, the posterior soft-ray territory is synergistically patterned by *shh* and *gremlin1b*, whereby *gremlin1b* determines the position of the spine to soft-ray boundary in WT fish.

Interference with BMP Signaling Induces Homeotic Transformations of Soft Rays into Spines and Vice Versa. Next, we strived to assess the phenotypic consequences of interference with the *shh* and BMP pathways. Morphological differentiation between spine and soft-ray elements, as indicated by the presence of fin segments and the development of spine tips, first occurs in *A. burtoni* around 10 dpf (25). Cyclopamine and SAG treatments induced widespread pleiotropic effects outside of the fins and severely compromised embryonic viability beyond 8 dpf, that is, before the morphological differences between spines and soft rays are established and therefore preclude such morphological analyses. DMH1 treatment or loss of *gremlin1b* is, however, well tolerated with phenotypic consequences that appear primarily in the fins and thus allow further morphological analyses of the extent of spine and soft-ray territories. In the dorsal fins of *gremlin1b* mutants, we observe a posterior shift of the spine to soft-ray boundary caused by a homeotic transformation of the anterior soft rays into spines as indicated by the presence of a spiny tip, the absence of segmentation, and the anterior fusion of the hemitrichia (Fig. 3A and *SI Appendix, Fig. S5*) (WT/heterozygous ($n = 21$): 13 to 14 spines, 9 to 10 soft ray; *gremlin1b*^{-/-} ($n = 16$): 15 to 20 spines, 3 to 6 soft rays). In the anal fin, a similar posterior expansion of the spine domain is observed whereby only 3 to 4 soft rays are maintained (Fig. 3C and *SI Appendix, Fig. S5*) (WT/heterozygous ($n = 21$): 3 spines, 8 to 10 soft ray; *gremlin1b*^{-/-} ($n = 16$): 4 to 7 spines, 1 to 6 soft rays). The preservation of soft-ray identity in the posterior fin is consistent with the presence of a posterior patch of *hoxa13b* expression that arises in a *shh*-dependent manner in *gremlin1b*^{-/-} embryos (Fig. 2C). The inhibition of BMP signaling through DMH1 treatment for a 24-h window during 4 to 5 dpf results in the opposite phenotype in the dorsal fin with an anterior transformation of spines into soft rays (Fig. 3B and *SI Appendix, Fig. S5*) ($n = 7$, spines 3 to 10, soft rays 14 to 21). This treatment induces the same soft-ray expansion in a *gremlin1b*^{-/-} background (*SI Appendix, Fig. S6*) ($n = 5/5$). In the anal fin, no significant shift in number of soft rays and spines is observed ($P = 0.06$, two-sided t test, $n = 6$, spines 2 to 3, soft rays 9 to 11) (*SI Appendix, Fig. S5*), suggesting that additional genetic factors besides BMP

signaling determine the presence of the 3 anterior fin spines in the anal fin.

Altogether, the observed homeotic transformations of spines to soft rays and vice versa underpin that BMP inhibition by *gremlin1b* is a primary determinant of soft-ray identity as also suggested by the analysis of developmental marker genes.

***Gremlin1b* Mutants Display Homeotic Transformations in Anal Fin Coloration.** The individuation of the soft-rayed and spiny domains of the male anal fin in *A. burtoni* is also reflected in its coloration. The mouth-brooding African cichlids evolved egg spots, or “egg dummies,” apparently to increase the chances of fertilization during courtship (25, 38). The distribution of egg spots in the anal fin typically shows a bias toward the posterior side of the fin overlapping with the soft rays while being absent from the spiny part. To investigate whether egg spots are in fact part of the same genetic modules that determine soft-ray and spine development, we analyzed the presence of egg spots at 3 mo of age in WT/heterozygous and *gremlin1b*^{-/-} males derived from two *gremlin1*^{+/-} crosses. Comparison of mutant with WT or heterozygous fish (which are WT in appearance with respect to spine and soft-ray distribution) shows an altered distribution of egg spots on the fin. Concomitant with the posterior shift of the soft-ray domain, the egg spots in these fish are present more posteriorly, and egg spots were never observed to overlap with the spiny-fin domain. In the cross analyzed, WT and heterozygous fish have an average of 3.5 egg spots whereas *gremlin1b* homozygous mutant fish have an average of 2 egg spots (WT/heterozygous $n = 8$; *gremlin1b*^{-/-} $n = 9$, $P = 0.0002$, two-sided t test) (Fig. 3D and *SI Appendix, Fig. S5*). In the same cohort, WT and heterozygous male egg spots are present over 57% of the length of the fin, whereas this is reduced to 28% in homozygous *gremlin1b* mutant fish ($P = 9.6 \times 10^{-6}$, two-sided t test) (*SI Appendix, Fig. S5*). Therefore, the distribution of egg spots in the anal fin appears to be determined by the same upstream patterning mechanism as that inducing the soft-ray and spiny-fin domains, whereby the posterior reduction of the soft-ray domain results in a concomitant posterior shift in the presence of egg spots.

Analysis of the Dorsal Fin Pattern in Nonacanthomorph Spiny and Nonspiny Catfish. Anterior spines have convergently evolved in several clades of nonacanthomorph teleosts such as catfish and carps. We wanted to further understand the relationship between dorsal fin patterns and the repeated emergence of fin spines. Furthermore, the dorsal fin pattern of nonacanthomorphs could provide information concerning the evolutionary origin of the acanthomorph fin pattern. We thus compared the anterior–posterior patterning observed in *A. burtoni* with that in nonacanthomorph species with median fins consisting of soft rays only or in those with convergently evolved fin spines. The nonacanthomorph zebrafish possesses soft rays only, and *alx4a* is expressed in the anterior-most fin rays of the dorsal and anal fins (39), tentatively suggesting that the spine pattern derives from a domain originally confined to the anterior fin margin. Zebrafish, however, has a narrow dorsal fin that is restricted to the posterior part of the body and that is about the size of the *A. burtoni* soft-ray domain. This leaves open the possibility that wider and further anteriorly extending nonacanthomorph fins show a similar extended *alx4* domain as *A. burtoni*. We investigated the expression of *alx4a*, *hoxa13b*, and *gremlin1b* expression in embryos of the African catfish (*Clarias gariepinus*), which has an extended dorsal fin (Fig. 4A) comprised of soft rays only and lacks the typical anterior spine found in many catfish species. Consistent with its soft-ray identity, *hoxa13b* and *gremlin1b* expression extends anterior throughout most of the dorsal fin. As in zebrafish, *alx4a* expression is confined to the anterior fin margin. Analysis in South American *Ancistrus* catfish whose anterior-most dorsal fin element has convergently evolved

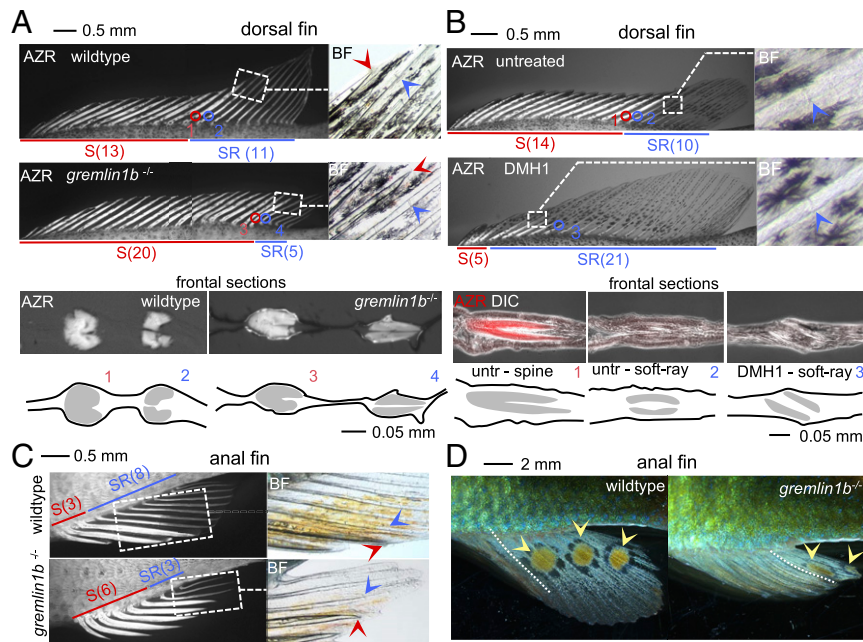


Fig. 3. Interference with BMP signaling induces homeotic transformations in dorsal and anal fins. (A–C) Fin morphology of dorsal (A and B) and anal fins (C) in WT (A–C), *gremlin1b*^{-/-} (A and C), and DMH1-treated (B) fish at approximately 1 mo postfertilization. Bony structures were visualized using Alizarin red (“AZR”), and fins were imaged using fluorescence microscopy. *Insets* shown (dashed boxes) were taken using brightfield microscopy (“BF”). In A and B, transversal sections at the level of the spine to soft-ray boundary are shown, which in A was imaged using fluorescence microscopy (Alizarin red fluorescence in white) and in B using fluorescence microscopy (Alizarin red fluorescence in false color red) and differential interference contrast microscopy (“DIC”) (in white). The dorsal fins of *gremlin1b*^{-/-} fish in A show an expanded spine domain (indicated by red line) and reduced soft-ray domain (indicated by blue line) indicating soft-ray to spine homeotic transformations. Alizarin red staining visualizes the heavier ossification of spines as compared to soft rays. *Insets* show spine (red arrowhead spine tip) and soft-ray (blue arrowhead segment boundary) characters, at the spine to soft-ray transition. Transversal sections through the spine to soft-ray boundary confirm the presence of fused and unfused hemisegments in spines and soft rays, respectively (section position is indicated with circles and numbers). The DMH1-treated fish shown in B show the opposite transformation displaying spine to soft-ray transformations. The *Inset* shows segments in the most anterior soft ray (blue arrowhead). Transversal sections confirm the presence of unfused soft-ray-like elements in the anterior fin. C shows a comparison of *gremlin1b*^{-/-} and WT anal fins showing soft-ray to spine transformations. *Insets* indicate spine and soft-ray characters at the spine to soft-ray boundary. A quantitative analysis of spine and soft-ray counts is provided in *SI Appendix, Fig. S5A*. (D) Egg spots are present on the soft-ray part of the anal fins of male *A. burtoni*. In *gremlin1b*^{-/-}, the egg spots have shifted posterior together with the soft-ray domain. A quantitative analysis of egg spots distribution is provided in *SI Appendix, Fig. S5B*. AZR: Alizarin red, BF: brightfield, S: spine, SR: soft ray. Anterior is to the left.

into a spine, for *hoxa13b*, *gremlin1b*, and *alx4a*, shows a similar pattern (Fig. 4B). In this species, the expression of all three genes overlaps in the first dorsal fin element, which will develop into a spine. Therefore, anteriorly limited expression of *alx4a* is also apparent in nonacanthomorph fish. This domain is, however, restricted much more anteriorly, and the *gremlin1b/hoxa13* domain extends along the anterior–posterior fin axis. Intriguingly, the anterior domain can coincide with the development of either a soft ray (as in zebrafish and *Clarias*) or a spine (as in *Ancistrus*) (*Discussion*).

Discussion

Spiny fins can be considered an evolutionary key innovation that arose as a novel module in the spiny-rayed fishes and added to the evolvability and thereby evolutionary success of the teleost body plan. Here, we show that the specification of spine and soft-ray domains during embryonic development is the result of a canonical signaling network involved in the patterning of the anterior–posterior fin axis, whereby posterior expression of *gremlin1b* and *shh* specify the soft-ray domains (Fig. 4C). In WT *A. burtoni*, the primary determinant of the spine to soft-ray boundary is BMP inhibition by *gremlin1b*, and alterations in BMP signaling induce homeotic transformations in fin identity. Interestingly, modulation of BMP levels is capable of inducing homeotic transformations in digits (40) and tooth identity (41). Therefore, spines and soft rays form another example of a deeply

homologous function of BMP signaling in “specifying discrete identities amongst meristic structures” (quotation from ref. 40).

During tetrapod limb development, *shh* and BMP inhibition via *gremlin1* are part of a regulatory loop including FGFs expressed in the distal ectoderm, which are required for ZPA survival (27–31, 35). We therefore investigated the potential role of FGFs in the establishment of soft-ray and spiny-fin domains. *Fgf16* is expressed along the anterior–posterior extent of the distal edge of the dorsal and anal fins and is slightly up-regulated by DMH1 and SAG treatment whereas it is somewhat down-regulated by cyclopamine treatment and in *gremlin1b*^{-/-} embryos (*SI Appendix, Fig. S7*). Altogether, this potentially indicates a conserved position of ectodermal FGF signaling downstream of *shh* and *gremlin1*. Treatment with the FGF antagonist BGJ398 from 4 to 7 dpf results in complete abortion of fin outgrowth, equally affecting spine and soft-ray domains and consistent with the relatively homogenous expression along the fin anterior–posterior axis. (*SI Appendix, Fig. S7*). Therefore, while important for fin outgrowth, ectodermal FGF signaling is not a major factor determining the anterior–posterior division of the dorsal and anal fins into spine and soft-ray territories.

In *A. burtoni*, the anterior–posterior pattern in dorsal and anal fins differs from that in their pectoral fins. In the latter, *hoxa13a/b* are expressed throughout the anterior–posterior extent of the fin (42, 43) and *alx4a/b* remain restricted to the anterior-most fin domain (39, 44). This appears to be a deeply conserved pattern that is for instance also present in shark pectoral fins (32, 45, 46).

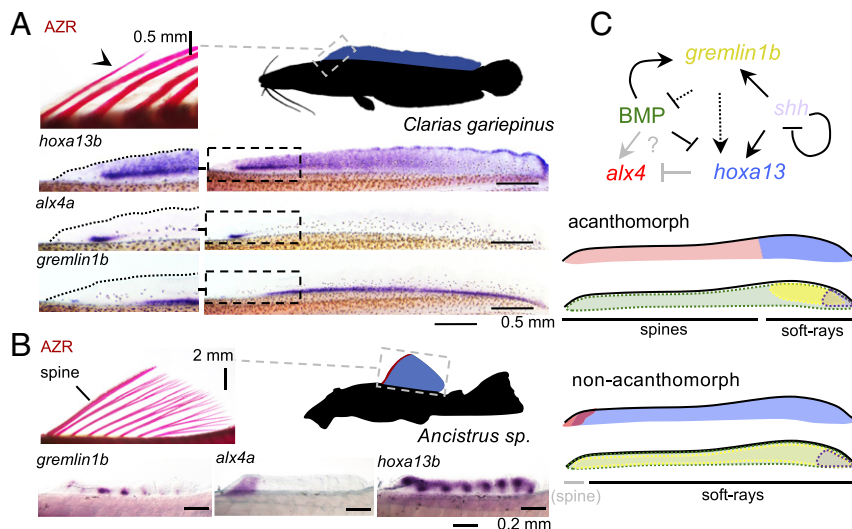


Fig. 4. Anterior–posterior dorsal fin patterning in acanthomorphs and nonacanthomorphs. (A) African catfish (*C. gariepinus*) have an extended dorsal fin (blue) comprised of soft rays only. The Alizarin red bone staining on the *Left* shows the anterior-most fin elements, and arrowheads indicate segment boundaries. In line with its soft-ray identity, *hoxa13b* and *gremlin1b* are expressed throughout most of the anterior–posterior fin axis. *Alx4a* expression is detected in a domain in the anterior fin similar to that in zebrafish (39). (B) *Ancistrus* catfish have a dorsal fin that is restricted to the anterior part of the trunk. This fin consists of posterior soft rays and a single anterior spine. *Hoxa13b* is expressed throughout the anterior–posterior extent of the fin, including the first elements, as is *gremlin1b*. *Alx4a* expression is confined to the spine and therefore may be involved in the individualization of this element compared to the posterior domain. (C) Model for the signaling network establishing the soft-ray domain in acanthomorph and nonacanthomorph teleosts. In acanthomorphs, the soft-ray domain is established via *gremlin1b*, which acts posteriorly in synergy with *shh* to activate *hoxa13* through the inhibition of BMP signaling. The absence of these posterior signals results in posterior expansion of *alx4* expression and the spine domain either through direct activation by BMP or loss of repression by *hoxa13* proteins. In nonacanthomorphs, the soft-ray signature extends throughout the anterior–posterior fin axis, and *alx4* is only expressed in the anterior fin margin, possibly related to the convergent evolution of spiny elements in nonacanthomorphs such as catfish. AZR: Alizarin red. Anterior is to the left.

Also, in *A. burtoni* pectoral fins *gremlin1b* is expressed throughout most of the anterior–posterior axis of the pectoral fin *Anlage* (*SI Appendix, Fig. S8*) and does not show the posterior bias observed in dorsal and anal fins. Overall, the patterning of the median fins in nonacanthomorph, *Clarias*, *Ancistrus*, and zebrafish (39) therefore resembles a pectoral fin pattern (although the median fin expression pattern of *gremlin1b* in zebrafish remains to be determined) and may therefore represent a shared ancestral pattern among median and paired fins that became modified in the median fins of spiny-rayed fish. This would have involved an expansion of the anterior pattern and a concomitant reduction of the soft-ray domain (Fig. 4C). Whether in the ancestral fin pattern *gremlin1b* acts to establish the posterior domain remains to be investigated by loss of function approaches in nonacanthomorphs. It is however suggestive that in *A. burtoni* *gremlin1b* loss does not lead to reduction of *hoxa13b* expression or expansion of *alx4a* expression in pectoral fins (*SI Appendix, Fig. S8*). This therefore might hint at a newly evolved posteriorizing role of *gremlin1b* in acanthomorphs median fins.

It is noteworthy that nonacanthomorphs frequently have a modified first fin element. For instance, in zebrafish and goldfish the first soft ray does not branch distally, and in many catfish species and carps a “spine” develops at this position. This suggests that an individuation of the anterior-most fin exists in nonacanthomorphs, which is consistent with the anterior domain of *alx4* expression in the fin margin of zebrafish (39), *Clarias*, and *Ancistrus* catfish. The tendency for more robustly ossified or spiny anterior fin ray elements is a trend present throughout fishes in both paired and median fins. Additional examples are the anterior fin spine in catfish (47) and sturgeon pectoral fins (48), robustly ossified anterior fin rays in tetrapodomorphs (49), the anterior fin spine that evolved convergently in chimaeras (50), and acanthodians (spiny sharks) and stem sharks (e.g., hybodonts) (51). It is plausible that convergently evolved spines all rely on the

same deeply homologous anterior fin individualization. Importantly however, this module appears restricted to the first few anterior-most fin elements only in all lineages except for the *Acanthomorpha*, which show a strong posterior expansion. Furthermore, spines in nonacanthomorph teleosts are different from those in acanthomorphs because the former initially develop as segmented elements that are indistinguishable from soft rays (47) (developing *Ancistrus* catfish dorsal fin shown in *SI Appendix, Fig. S9*). Therefore, in addition to the expansion of the anterior fin identity, a change in the downstream interpretation of this pattern (in the form of exaptation) was needed for the evolution of true fin spines and the consolidation of a robustly individualized anterior spiny-fin module in the acanthomorphs. Altogether, such changes in fin architecture allowed the emergence of the spiny-rayed fishes and initiated one of the most successful and diverse of vertebrate radiations.

Materials and Methods

In Situ Hybridization. In situ hybridization was carried out according to Woltering et al., 2009 (21), 2014 (52), 2015 (20), and 2020 (44). The reported shifts in expression domains in the inhibitor experiments and *gremlin1b*^{-/-} embryos were observed with complete penetrance.

Cloning of Probes. Probes were cloned in pGEMT (Promega A3600) vector using PCR from *A. burtoni*, *C. gariepinus*, or *Ancistrus* sp. embryonic copyDNA. A primer table is provided in *SI Appendix, Table S1*. The *A. burtoni* *hoxa11a*, *hoxa13a*, *hoxa13b*, *hoxd12*, and *alx4b* probes were described before (42, 44). Catfish sequences were identified by BLAST (basic local alignment search tool) against *C. gariepinus* and *Acistrus* sp. embryonal/larval RNA-seq libraries, and messengerRNA sequences for *alx4a*, *hoxa13a*, *hoxa13b*, *gremlin1a*, and *gremlin1b* were deposited in GenBank under accession nos. MW846856 to MW846866. Correct identification of “a” and “b” ohnologs was confirmed by generation of maximum likely hood gene trees and micro-synteny analysis (also reference *SI Appendix, Figs. S10 and S11* for *gremlin1* and *alx4*).

Small Molecule Treatment Experiments. Embryos were treated using the following concentrations: 1 μ M SAG (Selleckchem S7779) (dissolved at 10 mM in DMSO), 5 μ M cyclopamine (Selleckchem S1146) (dissolved at 50 mM in ethanol), 1 μ M DMH1 (Selleckchem S7146, dissolved at 20 mM in DMSO), and 1 μ M BGJ398 (Selleckchem S2183, dissolved at 10 mM in DMSO). Embryos were cultured in 30 mL equilibrated tap water (approximately pH 8, 9 $^{\circ}$ dH) with addition of 0.01 μ g/mL Methylene blue and penicillin/streptomycin (Sigma P4333) diluted 1:1,000 in \varnothing 8.5 cm plastic Petri dishes on an orbital shaker at 33 rpm at 28 $^{\circ}$ C in a heating incubator. Embryos were cultured at a maximum density of 20 embryos per dish (but usually less) and treated from 4 to 6 dpf for \sim 48 h (with the exception of BGJ398). Chemicals were added to the dish upon start of treatment, and embryos were kept in the same medium until the point of fixation (with 4% PFA buffered with 1 \times PBS (phosphate buffered saline) overnight at 4 $^{\circ}$ C, afterward storage in 100% ethanol at -20 $^{\circ}$ C). For the FGF inhibition experiment, embryos were cultured from 4 to 7 dpf for \sim 60 h in 1 μ M BGJ398, which was added at 4 dpf, and the embryos were kept in the same medium until the point of analysis. For the phenotypic analysis of DMH1 treatments, embryos were treated in 1 μ M DMH1 from mid 4 to mid 5 dpf for \sim 24 h and subsequently transferred to normal culturing medium and raised under standard conditions until the point of analysis. Mock treatments were performed using DMSO and ethanol, which do not result in phenotypic alterations.

RNA-Seq Analysis. RNA-seq was performed in triplicate using dissected soft-ray and spine territories of 9 dpf embryos using 10 individuals per sample. RNA was extracted using the ReliaPrep RNA Tissue Miniprep System (Promega Z6111) using the fibrous tissue protocol, and sequencing libraries were generated using TruSeq RNA Library Preparation Kit v2 (Illumina RS-122-2001). Samples were sequenced on an Illumina HiSeq2500 125 bp (base pairs) paired ends, and reads were demultiplexed and trimmed using Trimmomatic (version [v.] 0.36). The tuxedo pipeline for transcriptome assembly and quantification was used (53). Briefly, TopHat and Bowtie2 were used to map reads to the *A. burtoni* genome (v. 1.0). Cufflinks was used to assemble transcripts, to assemble a merged transcriptome, and to conduct differential gene expression analysis. Data (29,293 transcripts) were then imported into R (v. 3.6.3), and transcripts that showed no expression in at least one out of three replicates in at least one of the two groups (ray or spine) were excluded. Additionally, genes with extremely low expression (average FPKM [fragments per kilobase of transcript per million mapped reads] $>$ 0.5) were also excluded (20,592 transcripts, 17,733 of which were annotated and 17,597 were unique). Raw *P* values obtained from Cuffdiff were corrected for multiple testing using the false discovery method for transcripts. Raw sequence data have been deposited in National Center for Biotechnology Information (NCBI) Sequence Read Archive (SRA) under BioProject PRJNA718487 (54) with accession nos. SAMN18537261–SAMN18537266 (55–60).

Phenotype Analysis. Alizarin red staining was performed according to standard protocols and imaged under fluorescence microscopy. Spine and soft-ray counts given in *SI Appendix*, Fig. S5 were determined by manual inspection under a dissection binocular.

Animal Husbandry. *A. burtoni* were bred and collected at the University of Konstanz as previously described (25). *C. gariepinus* embryos were kindly provided by Fleuren and Nooyen, Aquaculture ID, Netherlands. *Ancistrus* sp. embryos were provided by private breeders. Animal experiments were carried out under 35-9185.81/G-18/32, *Tierforschungsanlage* (University of Konstanz) *Aktenzeige* T18/07.

Generation of CRISPR/Cas9 *gremlin1b* Mutant *A. burtoni* Lines. Guide RNAs (gRNA) were cloned in pT7gRNA and produced according to ref. 61 using oligos. gRNA-1 FW: TAGGACTCCAGCACTTCGTCGG, gRNA-1 RV: AAACCCGACGAAGTGTGGAGT, gRNA-2 FW: TAGGTTGTCTCCGATTCGTT, gRNA-2 RV: AACACAAGAATCGAGCAGCAA. A mixture of 1 to 2 nL of two gRNAs at 10 ng/ μ L each including Cas9 protein (NEB M0646T diluted 1/40) was injected at the one to two cell stage. Embryos were cultured individually in 6-well plates on an orbital skater at 28 $^{\circ}$ C in the presence of penicillin/streptomycin (Sigma P4333) diluted 1:1,000 and addition of methylene blue. Two independent lines were derived (*SI Appendix*, Fig. S12): *gremlin1b-stopCD38* has an in-frame premature stop codon introduced at codon 38; *gremlin1b Δ 740* has a 740-bp deletion including the 5' 339 bp and start codon. Both lines gave indistinguishable phenotypes. The *gremlin1b-stopCD38* was genotyped using fragment mapping on a capillary sequencer (3130xl Genetic Analyzer, Applied Biosystems) using a 40 cycle PCR with primers M13 tailed FW: CAGGAAACGCTATGACCACGCATATCTTCTACAGT-ATGG, RV: GTCTGCGTTGCTGCTCCGATTC followed by a second one-cycle labeling PCR with a HEX-labeled M13 FW primer: CAGGAAACAGTATGAC. The *gremlin1b Δ 740* allele was detected using standard PCR and gel electrophoresis using primers FW: CAGTGACAGTCGACACAGTAG, RV: GAC-GAGCACAATTTCTGGCTGTG.

Data Availability. RNA-seq dataset of *A. burtoni* 9 dpf fins and catfish gene sequences data have been deposited in NCBI SRA and NCBI GenBank (SRA: BioProject PRJNA718487 (54), SAMN18537261–SAMN18537266 (55–60); GenBank: MW846856–MW846866) (62–72).

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