Platyfish bypass the constraint of the caudal fin ventral identity in teleosts

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Abstract

Background: The caudal fin of teleosts is characterized by dorsoventral symmetry. Despite this external morphology, the principal rays of this appendage connect to bones below the notochord, indicating the ventral (hypochordal) identity of this organ. Results: Here, we report that this typical architecture of the caudal fin is not fully conserved in the platyfish (Xiphophorus maculatus) and the guppy (Poecilia reticulata), representatives of the Poeciliidae family. We show that in these species, 3–4 principal rays connect to bones above the notochord, suggesting an epichordal contribution. Consistently, as examined in platyfish, dorsal identity genes zic1/4 were highly expressed in these rays, providing molecular evidence of their epichordal origin. Developmental analysis revealed that the earliest rays above the notochord emerge at the 10-ray stage of fin morphogenesis. In contrast to zebrafish and medaka, platyfish and guppies display a mirrored shape of dorsal and ventral processes of the caudal endoskeleton. Our study suggests that an ancestral bauplan expanded in poeciliids by advancing its symmetrical pattern. Conclusion: The platyfish evolved a fin architecture with the epichordal origin of its upper principal rays and a high level of symmetry in the caudal endoskeleton. This innovative architecture highlights the adaptation of the teleost skeleton.

Key findings

1. In platyfish, principal rays of the caudal fin are associated with endoskeletal elements not only below but also above the notochord.
2. Dorsal identity markers zic1/4 are highly expressed in the three uppermost principal rays.
3. The epichordal principal rays begin to form at the developmental stage with 10 caudal rays.
4. The caudal fin of platyfish displays a striking internal symmetry of the endoskeleton.
INTRODUCTION

The fins are one of the most important innovations of fishes, since these appendages enable efficient locomotion in water. Among them, the caudal fin is particularly important to generate lift, propulsion power and to modulate maneuverability. The dorsoventral symmetry of this appendage has both functional and evolutionary implications. Ancestral fishes, possessed an asymmetrical fin, classified as “heterocercal,” with a larger dorsal lobe supported by the notochord or vertebral column, and a shorter ventral lobe. In the successive teleosts, the axial skeleton is withdrawn from the appendage, whereas the ancestral ventral lobe gives rise to a symmetrical “homocercal” fin. From the hydrodynamic perspective, dorsoventrally equilibrated morphology, in which the upper and lower halves are nearly equivalent in area and composition, could advance swimming performance. Approximately 96% of extant fish species are teleosts, reflecting the functional benefit of the homocercal tail.

The transition from the heterocercal to homocercal tail was associated with shortening and upward flexion of the notochord. Despite this modification and the external fin symmetry, the internal pattern of the associated endoskeleton largely retained its evolutionary origin. As formulated by Sallan, “the teleost caudal fin is actually the ventral lobe of the ancestral fin.” Thus, the symmetrical upper and lower fin halves derive from the ventral tissues of the axial skeleton, representing a “hypochordal” (below notochord) identity. Whether some teleost fish could bypass this limitation by increasing the contribution of “epichordal” (above notochord) tissues had not yet been investigated.

The fin comprises an array of exoskeletal elements, called lepidotrichia, which are classified as principal and procurrent rays. According to standard conventions, the principal caudal rays are described as the segmented and branched rays plus one upper and lower unbranched ray located at the lateral margin of each lobe of the fin (Figure 1A). Briefly, the number of principal rays is defined as the count of branched rays plus two. In practice, the principal rays reach to, or nearly to, the posterior fringe of the fin, whereby their relative length determines the fin shape, which can be forked, truncated or rounded. The procurrent rays are short and serve to widen the base of the fin. Consistent with the phylogenetic origin of the homocercal tail, anatomical studies of diverse teleosts have indicated that all the principal rays typically articulate with the hypochordal bones, which are positioned ventrally to the vertebral column, namely hypurals, parhypurals and haemal spines. Thus, with the exception of the dorsal procurrent rays, the principal part of the caudal fin is considered as an organ of ventral origin (Figure 1A). This interpretation is consistent with genetic studies in zebrafish, and lineage tracing analysis in medaka that demonstrated a continuous labeling of endoskeletal radials and their associated exoskeletal rays at specific anatomical positions. Based on these results it has been proposed that bones and their attached rays develop from the same population of mesoderm-derived mesenchyme. Thus, the positional identity of rays could to a certain degree be inferred from their attachment to the endoskeleton.

The homocercal tail is considered a beneficial trait for the 250 million years of teleost evolution, given the rich number of more than 30 thousand species. Following phylogenetic diversifications, certain taxonomic groups evolved unusual characteristics. Among them, the Poeciliidae family from the Cyprinodontiformes order, displays distinctive features that are rare among fishes. Indeed, these fishes do not reproduce by laying eggs, but instead, the fertilization is internal and females carry their offspring for approximately 3 weeks. During development, embryos are nourished either from a yolk or through a placenta analog, depending on species. To achieve the internal fertilization, the anal fin of males is transformed into the gonopodium. In addition, the caudal fin of males may carry secondary sexual traits, such as a ventral sword as seen in swordtails, or as a hypertrophied and elaborately patterned caudal fin, as seen in guppies. Members of the Poeciliidae family reveal a particular geographical adaptivity, which is highlighted by the fact that multiple independent populations have colonized habitats with extremely toxic conditions, such as hydrogen sulphide-rich springs in Central America. Thus, poeciliids display a high evolvability potential among teleosts. Whether the developmental symmetry of the caudal endoskeleton was subjected to evolutionary innovation remains yet insufficiently investigated in this group of fish.

A particularly valuable model organism from the Poeciliidae family is the platyfish (Xiphophorus maculatus). This species has attracted scientific interest for a century thanks to a great variation in pigmentation patterns, sex-
reversal ability, susceptibility to melanomas and complex behavior. The genome of this species has been sequenced and annotated, which renders it suitable for evo-devo research. Given the high evolutionary predisposition to novelties in the Poeciliidae family and the use of platyfish in genomic research, we have chosen to investigate whether the caudal skeleton has acquired any distinctive modifications in this species.

To better identify new traits, we compared them with the guppy (Poecilia reticulata) from the same Poeciliidae family, the Cyprinodontiformes order, and with the medaka (Oryzias woworae) from a sister order Beloniformes. (Figure 1B). The platyfish and the medaka belong to the same taxonomic monophyletic subseries Atherinomorpha, and their last common ancestor existed approximately 75 million years ago. For comparison, we also included the zebrafish (Danio rerio), the most common model fish in biomedical research. The platyfish and the zebrafish represent remote phylogenetic lineages that separated approximately 220–250 million years ago. Besides the anatomical characterization of the adult caudal skeleton, we also analyzed the developmental dynamics of the embryonic tail in platyfish. This morphological and embryological study highlights several unique features, suggesting that the platyfish evolved an exceptional bauplan of the homocercal fin, including epichordal components and an enhanced internal symmetry.

In this study, we attempt to distinguish between the superficial vs anatomical patterns. We use the terms “upper” and “lower” to describe the spatial position relative to the horizontal body axis, whereas “epichordal” and “hypochordal” to define the developmental dorso-ventral position relative to the notochord.

2 | RESULTS

2.1 | Comparison of caudal fin rays between platyfish and common model fish species

To characterize the skeleton of the caudal fin, we performed histological staining using Alcian blue & Alizarin red dyes. In two poeciliids, platyfish and guppies, the caudal fin has an oval form with a rounded margin, which...
FIGURE 2  Legend on next page.
contrasts with the forked fin shape of zebrafish and the straight-ended fin of medaka (Figure 2A–D). Examination of 6 male and 6 female platyfish revealed that the maximal length of the caudal fin reached approximately 11 mm, and the fin area was supported by 16–18 principal rays (average 17.5 ± .2; Table 1 and Figure 2D). Then, we counted procurrent rays, which are the short outermost unbranched rays at the dorsal and ventral edge of the appendage. The number of procurrent rays ranged between 4 and 7 (average 5.2 ± 0.2) at the dorsal side, and 5–9 (average 6.8 ± 0.3) at the ventral side (Table 1). We noticed a tendency that the fish with a lower number of principal rays had more procurrent rays, and inversely, so that the total number of all rays ranged from 28 to 31 (average 29.5 ± 0.3).

Interestingly, the zebrafish caudal fin is known to have comparable numbers of 16–19 principal rays and 5–7 procurrent rays at each of dorsal and ventral edges, as reported previously. However, despite nearly the same number of rays, the fin margin displayed opposite geometry, namely, convex (curved outwards) in platyfish vs concave (curved inwards) in zebrafish. This suggests that in these remotely related species, the growth of individual principal rays follows an opposite regulation along the dorso-ventral axis.

In medaka, the distal fin margin has a straight shape, and was supported by 11 principal rays (N = 6; 2 females and 4 males) (Figure 2B). The number of procurrent rays ranged between 4 and 5 (average 4.3 ± 0.2) at the dorsal side, and 5–6 (average 5.3 ± 0.2) at the ventral side. Together, the total number of rays was 20–22 (average 20.7 ± 0.4), consistent with a previous study.

Thus, although the medaka and the platyfish belong to sister orders (Figure 1B), they show a substantial variation in the number of caudal rays.

**TABLE 1** Number of rays in the caudal fin of adult platyfish

| Specimen | Fin length (mm) | Principal rays | Procurrent rays | Total number of rays |
|----------|----------------|----------------|-----------------|---------------------|
|          |                |                | Upper | Lower |                |
| Male 1   | 10.38          | 16             | 5      | 9      | 30               |
| Male 2   | 10.41          | 16             | 6      | 8      | 30               |
| Male 3   | 11.55          | 17             | 6      | 7      | 30               |
| Male 4   | 11.84          | 17             | 7      | 6      | 30               |
| Male 5   | 11.21          | 18             | 5      | 8      | 31               |
| Male 6   | 12.25          | 18             | 5      | 7      | 30               |
| Female 1 | 10.97          | 18             | 5      | 4      | 28               |
| Female 2 | 10.27          | 18             | 5      | 5      | 28               |
| Female 3 | 10.54          | 18             | 5      | 6      | 29               |
| Female 4 | 10.61          | 18             | 5      | 7      | 30               |
| Female 5 | 10.63          | 18             | 5      | 7      | 30               |
| Female 6 | 10.73          | 18             | 4      | 6      | 28               |
| Average  | 11.0 ± 0.2     | 17.5 ± 0.2     | 5.2 ± 0.2 | 6.8 ± 0.3 | 29.5 ± 0.3 |

Note: The fin length corresponds to the medial longest ray that was measured from the hypural plate to the distal tip on images of Alcian blue and Alizarin red stained specimens. Counting of rays was performed manually. The averages were calculated along with the standard error of the mean (SEM).

**FIGURE 2** Comparison of the caudal skeleton in adult zebrafish, medaka and platyfish (A–D). Tails of adult zebrafish (A), medaka (B), guppy (C) and platyfish (D) stained with Alcian blue and Alizarin red to detect cartilage and bone, respectively. The dorsal-most and ventral-most unbranched ray next to a branched ray represents the first and the last principal ray, respectively. The dashed-line frame indicates the area that is magnified in the right panel with the same letter followed by a prime symbol. In the magnified images of the caudal skeleton (A’–D’), the arrows point to the first and the last principal ray. (E–H) Schematic representations of the caudal skeleton based on histological staining. The epichordal (dorsal) bones are in purple, whereas hypochordal (ventral) bones in green. In zebrafish (E) and medaka (F), the principal rays (brown) articulate solely with hypochordal elements. In guppy (G) and platyfish (H), 3 upper principal rays articulate with epichordal bones. Zebrafish N = 6; Medaka N = 6; Guppy N = 5; Platyfish N = 12. CC, terminal compound centrum; Ep, epural; Hy, hypural; HyD, hypural diastema; HS, haemal spine; NS, neural spine; Ph, parahypural; PU, preural vertebra numerated from the posterior end; St, stegural (vestigial uroneural), Un, uroneural.
To compare the ray count of platyfish with that of another species from the same Poeciliidae family, we analyzed the skeleton of guppies (Figure 2C, C'). We found that in guppies, the number of principal rays ranged between 14 and 16 (N = 5), whereas the number of procurent rays was between 5 and 6 (average 5.4 ± 0.2) at the dorsal side, and 5–7 (average 6.2 ± 0.4) at the ventral side. Thus, X. maculatus has a few more principal rays than P. reticulata, suggesting variations of the caudal exoskeleton even among these closely related species.

2.2 | Atypical position of dorsal principal rays of the caudal fin in platyfish and guppies

In teleosts, the dorsal-most principal ray is known to align with the dorsal edge of the upper-most hypural, a bone of hypochordal identity (situated below the notochord). Consistently, in zebrafish and medaka, the first principal ray articulated with the upper-most hypural (Figure 2A, B, E, F). In platyfish, however, the first principal ray was carried by the neural spine of the preural-2 vertebra (Figure 2D, H). The assessment of 12 platyfish tails demonstrated that 3–4 of the upper principal rays articulated with epichordal bones, namely the epural and neural spines (4 fish with 3 such principal rays; 8 fish with 4 such principal rays). The same anatomical configuration was also observed in guppies (Figure 2C, G). These data demonstrate that the principal caudal fin ray field of poeciliids is not only of ventral origin, but includes a dorsal identity in its upper lobe, which is an atypical situation in teleosts.

Next, we determined the ventral extent of the caudal fin. In zebrafish and medaka, the last principal ray articulates with the haemal spine of the preural-2 vertebra (Figure 2A, B, E, F). A similar anatomical pattern was observed in platyfish and guppies (Figure 2C, D, G, H). Thus, in contrast to the dorsal margin of the principal rays, the ventral bounding remains conserved in all four fish species.

2.3 | Elevated expression of dorsal marker genes zic1/4 in the 3 uppermost principal rays of the platyfish caudal fin

Genetic studies in medaka suggested that zic1/4 genes, encoding zinc-finger type transcription factors, are required for the dorsal identity of the mesoderm in the trunk-tail region. Thus, the expression level of these two genes could serve to molecularly distinguish between the epichordal vs hypochordal origin of tissues. Accordingly, we identified zic1/4 orthologues in platyfish and conducted quantitative real-time PCR analysis using cDNA isolated from three parts of the caudal fin; (i) uppermost principal rays 1–3 (upR 1–3), which articulate with the epichordal bones; (ii) upper principal rays 6–8 (upR 6–8), which articulate with the hypural plate; (iii) lower principal rays 1–3 (loR 1–3), which articulate with the haemal spine and parhypural (Figure 3A, B). The latter sample unambiguously comprises tissues of ventral identity, which is predicted to show minimal zic1/4 expression, and thus, it can be used as calibrator for quantification of relative gene expression. To ensure the reproducibility of results, we performed normalization with two housekeeping genes, β-actin 2 (actb2) and polyadenylate-binding protein 1 (pabp) in the fin. These qRT-PCR analysis revealed several important results. First, we found that zic1 and zic4 transcripts showed approximately 7- to 9-fold increase in the uppermost triplet of principal rays, compared to the lowest triplet (Figure 3C, D). This relatively high zic1/4 expression in the upper principal rays 1–3 indicates the epichordal origin of this tissue, consistent with our anatomical observation. Second, the intermediate triplet, comprising principal rays 6–8, displayed low zic1/4 expression that was similar to values detected in the ventral-most principal rays (Figure 3C, D). This result demonstrates that these rays comprise the hypochordal tissue, as predicted from their articulation with the hypural. Third, analysis with both housekeeping genes yielded similar results, namely for zic1 in upR1-3, we recorded 11.9 and 12.3 normalized gene expression values relative to actb2 and pabp, respectively; for zic4, these numbers were 8.8 and 9.2 (Figure 3C, D). These independent quantifications reveal a robust reproducibility of our results. Taken together, we concluded that our qRT-PCR analysis provides a molecular indication for the epichordal origin of the three upper most principal rays in the platyfish caudal fin.

2.4 | Emergence of dorsal caudal rays during platyfish fin morphogenesis

The formation of the caudal skeleton has been well characterized in zebrafish and medaka. This process has not yet been described in platyfish. To understand the dorsal-ventral tissue contribution of the caudal fin in platyfish, we monitored developmental dynamics of ray formation. We dissected embryos from euthanized gravid females at different time-points, and exposed them to calcein, a fluorescent compound that is incorporated by cells at calcification sites, including
We classified the steps of caudal fin morphogenesis by counting rays. To determine the embryo size at each morphogenetic stage, we measured their standard length, defined as the distance from the head tip to the caudal peduncle, as previously established for zebrafish.

The correlation between the standard length and the number of caudal rays at different developmental stages was described by a nonlinear regression model ($N = 127$ embryos at different developmental stages; $R^2 = 0.8275$; $P$-value <0.0001) (Figure 4A). Embryos with the same number of rays varied in size: the highest variation was approximately 10% standard length difference for the group with 25/26 rays. This demonstrates that body growth and organ morphogenesis are not very tightly synchronized during in-ovario development, but individual variations are evident among platyfish embryos.

Calcein staining revealed that the earliest pair of rays formed in the middle of the emerging fin, which corresponds to the hypural diastema complex, as described below (Figure 4B). New rays are sequentially added at the upper and lower part of the fin, until the stage with 27/28 rays by the end of embryogenesis. Embryonic caudal fin rays were typically not yet branched, and thus, their prospective principal or procurent identity cannot be classified during development. This observation is consistent with postembryonic development in zebrafish, showing that ray branching morphogenesis occurs during juvenile growth.

Although calcein intensely stained early developing rays, the caudal endoskeleton became markedly labeled only from the 9/10-ray stage (Figure 4C). This result is consistent with the previous report in zebrafish that endochondral and dermal ray ossification occur independently from each other. At the 9/10-ray stage, two hypural plates were demarcated by calcein at the base of
the middle rays. Importantly, at least one ray was clearly located above the hypural edge at the 11/12 ray stage, suggesting a contribution from the epichordal tissue. At the 13/14-ray stage, 2 rays were positioned dorsally to the hypural plate (Figure 4C). We concluded that the tissue above the notochord starts to participate in the caudal fin formation after the stage with 10 rays.

2.5 | Developmental dynamics of hypural plate formation

To further investigate the developmental relation between the endoskeleton and exoskeleton, we performed histological analysis using Alcian blue staining for cartilage and Alizarin red for calcified bones. At early
FIGURE 5  The upper and lower hypurals develop during the formation of the 10 principal caudal rays (A-E) Whole platyfish embryos stained with Alcian blue and Alizarin red to label cartilage and bone, respectively, at subsequent developmental stages defined by the number of caudal rays from 2 to 10. At this development time-window no ossification of the skeleton is observed by Alizarin red staining. The labeling of the yolk (a dark spheric structure on the image) with Alizarin red might indicate the presence of calcium in this tissue. The framed box on each image encircles the embryonic tail, which is magnified in the images with a primed corresponding letter. (A’-E’) A higher magnification of the embryonic tail. The dashed-line frame depicts the tissue shown in a magnified view to the right. The earliest-forming pair of principal rays (red arrows) originate symmetrically around the hypural diastema (Hd) complex. New rays are added sequentially and symmetrically towards the upper and lower margin of the fin. (A’-E’) The magnified images focused on the development of the lower and upper hypural plates, which start to be ankylosed at their base. The dorsoventral branching of the caudal vasculature (red asterisk) is situated in the hypural diastema, in the notch between the upper and lower hypural plates. NT, notochord; Ep, epural; Hy, hypural plate; Hd, hypural diastema; Ph, parahypural; opc, opisthural cartilage. N = 7 (number of specimens).
stages up to 9/10 rays, the lepidotrichia were unstained and visible mostly by contrast imaging, whereas the notochord and the vertebral elements displayed cartilaginous staining (Figure 5).

As in other teleosts, the earliest-forming pair of rays was detected at the site of the hypural diastema complex, which participates in the establishment of dorsoventral caudal fin symmetry in adult fish. Indeed, the pioneering rays were formed by the connective tissue situated distally to the dorsal-ventral branching of the caudal vasculature, in the space between the lower (anterior) hypural plate (Hy\text{low}) and the upper (posterior) hypural plate (Hy\text{up}), which are compound elements in Cyprinodontiformes (Figure 5A). The latter hypural plate was smaller and only weakly visible at the 2-ray stage (Figure 5A), but it became strongly stained at the 4-ray stage (Figure 5B). At the 6- and 8-ray stage, all lepidotrichia were positioned distally to the hypural plates (Figure 5C, D). At the 10-ray stage, the upper-most ray was aligned with the upper edge of the hypural plate at the tip of the notochord (Figure 5E). Thus, until this time point, the caudal fin attached normally to the hypochordal tissue.

The analysis of the Alcian blue stained embryos revealed another interesting observation. At 2-ray stage, the terminus of the notochord was covered by intensively labeled cap, which may correspond to the opisthural cartilage (opc) (Figure 5A'). At the next developmental stage, this notochord-associated cartilage became linked to the upper hypural plate (Figure 5B'). Indeed, at subsequent developmental stages, the dorsal edge of the upper hypural plate encompassed the tip of the notochord, suggesting an unusual epichordal extension of this compound structure (Figure 5C', D', E'). We concluded that the opisthural cartilage fused with the hypural element during fin formation, which would be a new finding that has never been reported in any other species. This anatomical peculiarity might correlate with losing the positional boundary, which normally limits the lobe with principal rays to the hypochordal part.

The process of matrix calcification became visible by Alizarin red staining in more advanced embryonic stages (Figure 6). At the 18-ray stage, which corresponds to the typical number of principal rays in the adult fin, 4 rays articulated with the dorsal processes (Figure 6A). Subsequent fin development was associated with symmetrical addition of upper rays attached to the epichordal tissues and lower rays attached to the hypochordal tissues (Figure 6B, C). We concluded that new upper rays of the platyfish caudal fin are attached to the tissue above the notochord after the 10-ray stage.

2.6 Dorso-ventral symmetry of the adult caudal endoskeleton

To understand how the caudal endoskeleton was specialized in platyfish, we compared the morphology of the corresponding bones with three other fish species: zebrafish, medaka and guppies. We started from the posterior end, where the terminal compound vertebra provides the base for the dorsal uroneurals and ventral hypurals. While the zebrafish uroneural is a slender bone (Figure 2A', E), the medaka, as previously shown, lacks this element (Figure 2B', F). Similarly, guppies also did not have this bone in the skeleton (Figure 2C', G). In platyfish, we could detect only a vestigial outgrowth that corresponds to the rudimentary uroneural, also called stegural, consistent with our observations in embryos (Figure 2D', H). We concluded that the uroneural is not developed in the two analyzed species of Poeciliidae family.

Hypurals are flattened bones that form ventrally to the notochord, but become the most-posterior bones at the tip of the dorsally-flexed notochord. While zebrafish had five separate hypurals (Figure 2A', E) and medaka had two hypural plates (Figure 2B', F), the platyfish and guppy tails displayed only one hypural plate with a fan-like shape (Figure 2C', D', G, H). Our developmental analysis of platyfish indicated that this single plate originates from two initially independent hypural plates that started fusing anteriorly along the notochord at the 8-ray stage (Figure 5D). In adult platyfish, a foramen (an elongated oval gap) was present in the base of the hypural plate, corresponding to the remnant of the hypural diastema during embryonic development (Figures 5 and 6). Microcomputed tomography revealed that the hypural plate was fused with the terminal compound vertebra (Figure 7). Thus, the extensive fusion of the hypurals into one plate, which is ankylosed with the terminal vertebra, represents a remarkable feature of the platyfish.

The preural-1, which is the anterior part of the terminal compound centrum, is typically associated with the epural, a dorsal/epichordal bone, and the parhypural, a ventral/hypochordal bone. In zebrafish and medaka, the epural had a slender rod-like structure, whereas in platyfish and guppies, this bone had a flattened form (Figure 2). As opposed to zebrafish and medaka, in platyfish and guppies, the epural mirrored the shape of the ventrally located counterpart, the parhypural. In both Poeciliidae species, in the neural (dorsal) and haemal (ventral) spines of the preural-2 vertebra displayed a distally widened shape and carried blade-like plates with a mirrored appearance, which contrasts thin spines in zebrafish and medaka. Thus, in platyfish and guppies, the
The caudal skeleton displayed a higher level of dorsoventral symmetry.

Interestingly, we identified a difference between both Poeciliidae species in blade-like plates, which widen the base of the mentioned bones. These blade-like plates had a serrated edge in guppies (Figure 2C', G), whereas a straight margin in platyfish (Figure 2D', H). Furthermore, in guppies, the pattern of notches in these bony ridges was not identical on the dorsal and ventral counterparts. This variation in detailed serration led to a decreased dorsoventral symmetry of the skeleton in guppies, compared to platyfish. Nevertheless, both poeciliids displayed overall similar morphological modification of the epichordal and hypochordal spines, reinforcing the internal dorsoventral symmetry of their caudal endoskeleton.

Beside symmetry-boosting modifications of the dorsal spines in platyfish, we found that the number of vertebrae with supportive bones for the caudal fin was higher in platyfish and guppies than in the zebrafish and medaka. In the two latter species, the terminal compound centrum (CC) and two preurals (PU-2 and PU-3) participated in the support of the caudal rays, as reported (Figure 2A', E, F). In both poeciliids, beside these 3 vertebrae, another anterior preural 4 (PU-4) contributed to radials of the caudal fins. In 25% of examined platyfish specimens, neural spines of preural 5 (PU-5) articulated with the procurent upper rays (3 out of 12 specimens) (Figure 2D', H). Thus, the platyfish and guppy caudal fins were supported by the endoskeleton involving the terminal compound centrum and four to occasionally five separate preurals, offering a greater number of supports compared to zebrafish and medaka.

3 | DISCUSSION

The caudal fin of the teleost has evolved an external dorsoventral symmetry, characterized by a nearly mirrored distribution of principal rays in the upper and lower
lobes. This trait, however, is not reflected by the associated endoskeleton that retained its ancestrally derived ventral identity. Here, we provide evidence that the bauplan of the teleost fin is exceptionally innovative in two Poeciliidae species, the platyfish and the guppy, which bypassed the constraint of the ventral attachment of principal rays and achieved a high degree of internal symmetry.

Our main finding is that 3–4 dorsal principal rays articulate with the epural and neural spine, which are bones of epichordal origin. This contribution of the dorsally derived elements to the principal field of the caudal fin represents an atypical pattern in extinct and living teleosts, according to the broadly accepted conventions formulated by Schultze and Arratia. The epichordal identity of the 3–4 principal rays might be a distinctive trait for the Poeciliidae family. A recently described new species of Cyprinodontiformes, named Pseudorestias lirimensis, forms an articulation between the first principal ray and epural. Interestingly, the Cyprinodontiform fossil from the oligocene in France, †Prolebias delphinensis, reveals a configuration, in which the upper non-branched principal ray articulated with the neural spine. In this extinct species, the caudal fin consisted of 12 branched rays, whereby the upper-most branched ray connected to the epural. Thus, the involvement of epichordal tissue to the bauplan of the caudal skeleton might be common to the poeciliids and potentially to other families of the Cyprinodontiform order.

3.1 Hypothetical morphogenetic boundaries and organizers of the caudal fin

We do not yet know how poeciliids could bypass the constraint of the caudal fin ventral identity that is outlined by the notochord position. Developmental analysis can provide valuable cues about homologies emerging during ontogenesis. Studies of some teleosts have led to a hypothesis that a boundary landmark for the upper limit of principal rays corresponds to the opisthural cartilage, which is formed by the posterior tip of the notochord. In medaka, the opisthural cartilage has been reported to become ossified and fused to ural centrum 2 during fin development. In adult zebrafish, the opisthural cartilage remains non-ossified at the ventral
edge the procurent ray, above the first principal ray$^{4,53}$ (Figure 2A, E). We observed that in platyfish embryos, the opisthural cartilage fused with the hypural element during early fin formation (Figure 5A″−C″), which might be a unique characteristic among teleosts. This finding suggests that the anatomical landmark for the presumptive boundary between the upper principal and procurent rays becomes reduced already at the onset of fin development. It is tempting to hypothesize that the joining of notochord-associated boundary with a hypural element would permit or facilitate an evolutionary path for the formation of principal rays in the epichordal part of the body.

Developmental principles regulating external symmetry of the caudal fin, which is characterized by equal dorsal and ventral halves, have been recently addressed using transgenic reporters in zebrafish. In that study, monitoring of ray emergence indicates an interplay between three morphogenetic fields, namely a central organizer at the hypural diastema and two peripheral organizers at the upper and lower margins of the caudal fin. Platfish appear to adhere to the model of a central organizer, since our histological analysis show that rays are sequentially added from the hypural diastema outwards (Figures 5 and 6). In contrast, our data with calcein staining of platyfish embryos did not reveal any early separate emergence of bounding principal rays (Figure 4C). This suggests that in platyfish, the hypothetical peripheral organizers might be synchronized with the morphogenetic activity of the central organizer. Further research is warranted to elucidate the position of ray organizers relative to the caudal endoskeleton in platyfish.

### 3.2 | zic1/4 genes are markers of epichordal rays in the platyfish caudal fin

The exoskeletal fin rays differentiate together with the endoskeletal radials, the patterning of which occurs along the anterior-posterior axis during development. Cell lineage tracing analysis in medaka demonstrated that bones and their attached rays develop from the same population of mesoderm-derived mesenchyme.

In this study, we inferred the dorso-ventral identity of rays from their connection to the endoskeleton. Our developmental analysis of platyfish revealed that the first dorsally attached ray is formed starting at the 10-ray stage of the caudal fin. Given that 10 central rays are attached to the hypural plate, our interpretation of this finding is that dorsal and ventral rays become symmetrically distributed on each side during their formation.

The molecular basis of the mechanisms increasing the contribution of epichordal tissue in the caudal skeleton might involve the regulation of zic1 and zic4 expression. These zinc-finger transcription factors were highly expressed in the three uppermost principal rays as compared to the other rays, as shown by qRT-PCR. Studies in medaka demonstrated that zic1 and zic4 are epichordal marker genes. A mutation of the zic1/4 enhancer, called Double anal fin (Da), causes a ventralized phenotype of the body, in which caudal fin rays articulate with both hypurals and epurals. In another species, called Siamese fighting fish (Betta splendens; order Perciformes), a double-tail mutant has been associated with a deletion of a zic1/4 enhancer. Both in medaka and Siamese fighting fish, zic1 and zic4 were suppressed in double tail mutants, suggesting a conserved mechanism for dorsal duplication of the caudal fin lobe in teleosts. In platyfish, we found that zic1 and zic4 were expressed in the three upper principal rays, providing molecular evidence of the epichordal origin of these structures. How these master genes regulate the innovative bauplan of the platyfish tail requires further studies.

### 3.3 | Advancements of the caudal endoskeletal bauplan in Cyprinodontiformes

The next striking feature of the caudal skeleton in platyfish and guppies is a high degree of dorsoventral symmetry of the supporting bones. This observation is consistent with the main feature of Cyprinodontiformes, which evolved enhanced symmetry of its caudal endoskeleton, as reported for various species. In this order, the upper and lower hypural plates can stay separate or fuse to provide a more rigid support for the principal rays. Consistently, our developmental analysis of platyfish demonstrated that the hypural plate derives from two separate elements, namely a lower hypural plate (ie, hypurals 1 + 2 in zebrafish) and an upper hypural plate (ie, hypurals 3 + 4 + 5). They become ankylosed to form a single bone, starting at the 18-ray stage of the caudal fin, leading to the disappearance of hypural diastema. An anterior gap in this plate remains visible in platyfish, as reported for various species, and may permit the passage of caudal vasculature. Importantly, a symmetrical fan-like hypural plate provides a support for equidistant rays at the center of the fin. This regularly spaced sequence of rays contrasts with the typical situation of other teleosts, including zebrafish and medaka, in which a wider interspace between the lower and upper hypural groups is demarcated by a hiatus between the corresponding rays. Thus, as compared to the bauplan with separated
hypural elements, one hypural plate uniformly consolidates the structural support for the fin and it enhances the regularity of ray distribution.

The *Cyprinodontiformes* can be unambiguously recognized by the presence of a single, blade-like epural that mirrors the parhypural, and the widened neural and haemal spines. Consistently, the next level of the caudal endoskeletal symmetry in platyfish and guppies is represented by these structures. In both these species, the dorsal and ventral counterpart bones of preural centra display mirrored appearance and they symmetrically support upper and lower principal rays. Indeed, the shape of epural and parhypural bones of the terminal vertebra, as well as the neural and haemal spines of preural-2, were similarly flattened with a wider distal end, and with protruding blade-like ridges. In platyfish, these thin ridges had straight margins, whereas in the analyzed guppy species (*Poecilia reticulata*), they were serrated. Interestingly, a study of another guppy species, *Poecilia mexicana*, has shown that the counterpart bones bear straight edges, suggesting the existence of endoskeleton variations among the same genus. This observation supports a hypothesis about powerful evolvability within this taxonomic group.

From the functional perspective, the stunning internal symmetry of the platyfish and guppy caudal skeleton might further advance the hydrodynamic characteristics of the tail. This could be of advantage as compared to an internally asymmetrical bauplan of the typical homocercal fin. The biophysical properties could be compared between distinct types of caudal fins with a different degree of the endoskeletal symmetry. An interesting question is to determine how the evolutionarily innovative architecture of the poeciliid caudal fin translates to the swimming performance.

### 4 EXPERIMENTAL PROCEDURES

#### 4.1 Fish strains and animal procedures

The following fish were used for this study: zebrafish (*Danio rerio*) AB strain (Oregon) at approximately 3 cm standard length, platyfish (*Xiphophorus maculatus*) strain Bleeding Heart, Mickey Mouse or Gold platy at approx. 3.3 cm standard length, guppies (*Poecilia reticulata*) at approximately 3 cm standard length, medaka (*Oryzias wowitzae*) of approximately 2 cm standard length. The standard length was measured from the snout to the caudal peduncle, without including the caudal fin. Adult platyfish, guppies and medaka were purchased from a commercial aquarium fish vendor (Aqualand, Renens/Lausanne, Switzerland). Zebrafish were bred in our fish facility.

Animal euthanasia was performed by immersion in 300 mg/L tricaine solution (MS-222; Sigma-Aldrich) for approx. 10 minutes. Then, the hearts were dissected from the euthanized fish. This procedure was performed in accordance with Swiss regulations and was approved by the Cantonal Veterinary office of Fribourg, Switzerland.

Adult tails were cut with a sharp razor blade from euthanized fish after heart removal. Platyfish embryos were dissected from similarly euthanized gravid females using a pair of sterilized dissection scissors. The entire ovary was placed in system water and embryos were removed from follicles and manually dechorionated with tweezers.

#### 4.2 Histological staining

A two-color acid free staining technique with Alcian blue and Alizarin red solution was used as described previously. Two stock staining solutions were prepared as Part A and Part B and stored at room temperature. The Part A component contained 0.02% Alcian blue dissolved in 70% Ethanol with 60 mM MgCl2. Preparation of this solution required two steps. Firstly, 0.4 g Alcian blue 8GX (Sigma-Aldrich, A5268) was dissolved in 50 ml of 50% ethanol at 37°C on a shaker. After the powder has dissolved, 50 ml of 90% ethanol was added and mixed. In the last step, 5 ml of this solution was combined with 70 ml of 95% ethanol, 6 ml of 1 M MgCl2 and 19 ml water. The Part B consisted of 5% Alizarin red S (Sigma-Aldrich, A5533) dissolved in distilled water. The final double staining solution was prepared freshly before staining by mixing 10 ml of Part A with 200 µl of Part B.

The fins of adult fish were fixed in a petri dish in 4% paraformaldehyde in PBS for 2 days at room temperature. Embryos were fixed overnight at room temperature. Specimens were then rinsed and washed with PBS twice for 20–30 minutes each. For dehydration prior to staining, adult specimens underwent an ethanol series as follows: 70% - 80% - 90% - 100% for 1 hour each. For embryos, a 15-minute dehydration step in 50% ethanol sufficed. Fins and embryos were rocked overnight at room temperature in the Part A/Part B double staining solution, followed by rinsing in distilled water to remove excess dye. Specimens were subsequently bleached to remove pigmentation, using a 1:1 ratio of 2% potassium hydroxide and 3% hydrogen peroxide during 1–3 hours for adults and 20–30 minutes for embryos.

In order to clear the specimens to visualize the bones, adult fins were incubated in an enzyme solution of borax (3:7 saturated borax solution to distilled water) and
trypsin (0.1%; Sigma-Aldrich, T4799) during 3 days. Scales also stain lightly with the double stain solution, so these were removed in distilled water using forceps. For imaging and storage, fins and embryos were transferred to glycerol using a graded series of 0.5% potassium hydroxide and glycerol solutions (3:1, 1:1, 1:3; for at least 2 hours or overnight) and finally stored in 100% glycerol and imaged.

Color images were taken with a Leica AF M205 FA stereomicroscope equipped with a digital camera (Leica Microsystems, Wetzlar, Germany).

4.3 | Calcein labeling of embryos and developmental staging

0.2% calcein (Sigma-Aldrich, C0875) solution was prepared in system water. Due to calcein's strong acidifying affects, we adjusted the pH to 7.2 by adding an appropriate amount of 1 N sodium hydroxide. Live platyfish embryos were immersed in this solution for 15 minutes in petri dishes, rinsed several times in system water during approx. 30 minutes. The embryos were then euthanized in 300 mg/L tricaine solution. Fluorescent images of embryos were taken with a Leica AF M205 FA stereomicroscope using a GFP2 filter (Leica Microsystems, Wetzlar, Germany).

The standard length was measured from the snout to the caudal peduncle, not including the caudal fin and following the body axis, using ImageJ (NIH, Bethesda, Maryland, United-States). Developmental staging was based on caudal fin morphogenesis, which was assessed by counting caudal rays. These two parameters were plotted together using Prism (Graphpad, San Diego, California, United-States), where a nonlinear regression curve was calculated.

4.4 | Quantitative real-time polymerase chain reaction analysis

Ray triplets were excised from caudal fins with a scalpel: uppermost principal rays 1, 2 and 3 (upR 1–3), upper principal rays 6, 7 and 8, counted from the top (upR 6–8) and lower principal rays 1, 2 and 3, counted from the bottom (loR 1–3). Tissues from three fish were collected and pooled together per one sample. Each sample was reproduced in triplicates, deriving from other groups of three fish. Collected fin tissues were frozen immediately on dry ice and homogenized in Qiazol Lysis Reagent (Qiagen, Hilden, Germany) using a Polytron tissue homogenizer. RNA was extracted using chloroform and isolated from tissue debris using MaxXtract high density columns (Qiagen, Hilden, Germany). RNA was precipitated using isopropanol and resuspended in water. cDNA was synthetized using the Superscript IV reverse transcriptase (Thermo Fishers, Waltham, Massachusetts, United-States), following the manufacturer's protocol. qRT-PCR was performed using the Kapa SYBR Fast qPCR kit (Kapa Biosystems, Wilmington, Massachusetts, United-States) following the manufacturer’s guidelines. The following primers were used:

**actb2** (XM_005806049.2): Fw: 5'-CGTGCGGGGATATCATTTCGCCCTG-3'; Rev: 5'-ACAACCAGTGCGCGATTTCC-3'.

**pabp** (XM_005797394.2): Fw: 5'-CCAGAGTCTCTCCGCTCCAAG-3'; Rev: 5'-TGGGACAACCGCTGAGTTGG-3'.

**zic1** (XM_005797813.2): Fw: 5'-CACGTCGGACGCGGTATC-3'; Rev: 5'-GCCAGGGTTGGAGACTGTC-3'.

**zic4** (XM_005797812.2): Fw: 5'-GGGTTAAAGTGTCTGGAAG-3'; Rev: 5'-GGGTTAAAGTGTCTGGAAG-3'.

Normalized gene expression was calculated using the delta CT method. Gene expression levels were averaged over 3 cDNAs per ray type and two technical replicates. Results were plotted in Graphpad Prism (Graphpad, San Diego, California, United-States).

4.5 | Micro computed tomography

The tails collected from euthanized fish were fixed in 4% paraformaldehyde in a petri dish for 2 days at room temperature. The specimens were then dehydrated in 70% ethanol for one hour and transferred into a transparent plastic tube with 70% ethanol.

Micro CT images with a voxel size of 3 μm were acquired using a Bruker SkyScan 2211 X-ray microscopy platform, equipped with an 11-megapixel CCD panel. Image reconstruction was performed using NRecon software (Bruker Corporation, Germany) and Avizo (Thermo Fisher Scientific).

4.6 | Terminology

The terminology follows the conventions formulated by Schultze and Arratia.6,7,12

Centrum or vertebral centrum: A mineralized, ossified, or partly cartilaginous/ossified element that surrounds the notochord.

Compound centrum (CC): terminal vertebral centrum at the posterior region of the caudal endoskeleton comprising preural centrum 1 and ural centra.

Epural: Detached neural spine of a terminal compound vertebra that commonly supports one or more dorsal procurrent rays.
Hypural: Modified haemal spine (of an ural centrum) that has lost its haemal arch and canal.

Hypural diastema: Space positioned between the lower and upper hypural plates.

Opisthural cartilage (opc): Cartilage at the posterior extremity of the notochord. It can extend distally between the uppermost (unbranched) principal ray and the procurent fin ray above it. It is considered to define the upper limits of the field with principal caudal rajas.

Opisthural gap: A space between the upper bounding principal ray and the adjacent procurent ray. This gap corresponds to the upper limit of the caudal principal ray field.

Parhypural: The parhypural is the haemal spine of preural centrum 1, which is fused within terminal compound vertebra.

Preural centrum: Vertebral centrum of the caudal region preceding the ural centra, bearing both neural and haemal arches and usually both neural and haemal spines, each of which supports a caudal ray at its distal tip. Preural centra are numbered from the posterior-most to the anterior-most.

Procurent caudal ray: Procurent rays are short rays, shorter than the principal ones, which form the anterior series of lepidotrichia of median fins and which are associated with endoskeletal elements.

Principal caudal rays: Principal rays of the caudal fin are all the segmented and branched rays plus normally one unbranched but segmented ray located at the leading margin in each lobe of the fin.

Stegural: Modified anterior-most uroneural bearing membranous bony extension at its antero-dorsal border.

Uroneurals: Modified neural arches that extend from the ural centrum dorsally to the notochord.

**AUTHOR CONTRIBUTIONS**

Lana Rees: Investigation (equal); methodology (equal); project administration (equal); writing – review and editing (equal). Désirée König: Conceptualization (supporting); data curation (supporting); investigation (supporting); methodology (equal); writing – review and editing (equal). Anna Jaźwińska: Conceptualization (lead); funding acquisition (lead); investigation (equal); methodology (equal); resources (lead); supervision (lead); validation (equal); writing – original draft (lead); writing – review and editing (equal).

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**CONFLICT OF INTEREST**

The authors declare no conflict of interest.

**DATA AVAILABILITY STATEMENT**

All raw data and imaging files are available upon request from the authors.

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