Internal vertebral morphology of bony fishes matches the mechanical demands of different environments

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Abstract
Fishes have repeatedly evolved characteristic body shapes depending on how close they live to the substrate. Pelagic fishes live in open water and typically have narrow, streamlined body shapes; benthic and demersal fishes live close to the substrate; and demersal fishes often have deeper bodies. These shape differences are often associated with behavioral differences: pelagic fishes swim nearly constantly, demersal fishes tend to maneuver near the substrate, and benthic fishes often lie in wait on the substrate. We hypothesized that these morphological and behavioral differences would be reflected in the mechanical properties of the body, and specifically in vertebral column stiffness, because it is an attachment point for the locomotor musculature and a central axis for body bending. The vertebrae of bony fishes are composed of two cones connected by a foramen, which is filled by the notochord. Since the notochord is more flexible than bony vertebral centra, we predicted that pelagic fishes would have narrower foramina or shallower cones, leading to less notochordal material and a stiffer vertebral column which might support continuous swimming. In contrast, we predicted that benthic and demersal fishes would have more notochordal material, making the vertebral column more flexible for diverse behaviors in these species. We therefore examined vertebral morphology in 79 species using micro-computed tomography scans. Six vertebral features were measured including notochordal foramen diameter, centrum body length, and the cone angles and diameters for the anterior and posterior vertebral cones, along with body fineness. Using phylogenetic generalized least squares analyses, we found that benthic and pelagic species differed significantly, with larger foramina, shorter centra, and larger cones in benthic species. Thus, morphological differences in the internal shape of the vertebrae of fishes are consistent with a stiffer vertebral column in pelagic fishes and with a more flexible vertebral column in benthic species.

KEYWORDS
mechanics, statistical modeling, vertebral morphology

TAXONOMY CLASSIFICATION
Biodiversity ecology, Evolutionary ecology, Zoology
1 | INTRODUCTION

Many groups of fishes have evolved specialized forms for living close to the bottom of a body of water, called a benthic habitat, and for living in open water, a pelagic habitat (Burress et al., 2017; Friedman et al., 2020; Hollingsworth et al., 2013; Hulseey et al., 2013; Kusche et al., 2014; Ribeiro et al., 2018; Robinson & Wilson, 1994; Willacker et al., 2010). In benthic habitats, fishes spend much of their time in contact with the substrate and often use "lie-and-wait" predation strategies (e.g., flatfishes: Link et al., 2002). In pelagic habitats, fishes usually swim constantly and often evolve an elongate, streamlined form (Friedman et al., 2020; Tavera et al., 2018) that is thought to be advantageous for steady swimming (Lauder, 2015). Often, another category is added, termed demersal. In these habitats, fishes live in close proximity to the substrate, but do not typically sit directly on it. They often feed by sifting detritus through sand or scraping algae off of coral and rocks and tend to have deeper bodies (Friedman et al., 2020; Larouche et al., 2020).

This diversification in body shape across habitat categories was most thoroughly evaluated recently by Friedman et al. (2020). Across a large sample of fish species, they found subtle, but significant differences in body shape, particularly in benthic fishes. Compared to benthic fishes, pelagic and demersal fishes tended to have deeper bodies and a narrower range of body widths. Benthic fishes include both flatfishes, with extremely high body width, and elongate fishes, with more narrow bodies. Indeed, benthic fishes had the largest diversity of body shapes associated with the highest rate of body shape evolution (Friedman et al., 2020).

Within individual clades, the pattern of differences in species from benthic, demersal, and pelagic habitats is often present and can be even more pronounced than Friedman et al. (2020) found when considering many taxa. For example, cichlid species in the African Rift Lakes have repeatedly and convergently evolved streamlined shallow-bodied pelagic forms and deep-bodied benthic and demersal forms (Cooper et al., 2010; Muschick et al., 2012). Similar patterns have been seen in many different groups of fishes, including gnuts (Tavera et al., 2018), new world cichlids (Kusche et al., 2014), cyprinids (Hollingsworth et al., 2013), and in Carangaria, a large group of marine fishes that includes both flatfishes, an extreme benthic morphology, and billfishes, an extreme pelagic morphology (Ribeiro et al., 2018).

These morphological shifts can be observed even in individual species or genera, where their functional consequences are clearer. Within three-spine stickleback species Gasterosteus spp., shallower and deeper bodied ecomorphs have evolved multiple times in separate lakes. Ecomorphs that primarily feed on benthic prey and tend to stay close to the substrate (“demersal” in the classification we use) have deeper bodies, while those that feed in open water (“pelagic”) have more streamlined bodies (Schluter, 1993; Walker, 1997; Willacker et al., 2010). The pelagic ecomorphs had lower drag coefficients and could swim faster for longer than the demersal species (Blake et al., 2005). Similarly, bluegill sunfish (Lepomis macrochirus) have pelagic and demersal ecomorphs, often in the same lakes, where the pelagic morphs have more streamlined bodies and demersal morphs had deeper and wider bodies (Ehlinger & Wilson, 2006; Gerry et al., 2011). These morphological differences correspond to lower cost of transport during steady swimming in pelagic morphs (Ellerby & Gerry, 2011) and faster turning performance in demersal morphs (Gerry et al., 2012).

In this extensive body of literature, we see a consistent pattern in which pelagic species have more streamlined bodies and swim more constantly and at higher speeds than demersal and benthic species. In contrast, demersal and benthic species often have deeper bodies and greater turning performance than pelagic species. We hypothesized that the external morphological differences would be accompanied by internal differences. In particular, since these differences are related to locomotion, which is driven by the axial musculature that acts to bend the vertebral column, we hypothesized that they might be accompanied by differences in the morphology of the vertebral column. The vertebral column both serves to resist bending forces from the axial musculature and as an attachment point for those muscles. Overall body stiffness is thought to contribute to high speed, continuous swimming (Koob & Long, 2000; Summers & Long, 2006); we therefore predicted that pelagic species that swim continuously should have a morphology associated with a stiffer vertebral column than demersal or benthic species. Benthic and demersal species, in contrast, are often "lie-and-wait" predators (e.g., flatfishes: Link et al., 2002), attacking prey with rapid accelerations that require high body curvature (Akanyeti et al., 2017; Schwalbe et al., 2019).

Donatelli et al. (2021) recently examined how the internal shape of vertebræ in fishes affects the stiffness of the vertebral column. In particular, they showed that the shape of the intervertebral joints, which are filled by the notochord, has a substantial effect on the stiffness of the vertebral column. Teleost fishes have characteristic hourglass-shaped vertebral centra, called amphicoelous centra, consisting of anterior and posterior cones, joined at the tips (Laerm, 1976). The notochordal tissues (i.e., the notochordal cell mass, notochordal epithelium, and notochordal strand) fill the inside of the centra, including a hole through the middle called the notochordal foramen, and makes up most of the intervertebral joint (Symmons, 1979). The bony elements of the centra are joined via a complex of soft tissues (the encapsulating complex) including the external intervertebral ligament (EVL), elastica externa (EE), and a fibrous sheath (FS) (Symmons, 1979). Donatelli et al. (2021) found that vertebral segments with larger foramina and larger cone angles tended to be more flexible. Overall, the mechanical properties of the joints contribute to the stiffness of the vertebral column (Long, 1992; Long et al., 2004; Nowroozi & Brainerd, 2012, 2014; Porter & Long, 2010), although others have questioned whether fish bend enough during normal swimming for the intervertebral joints to have any effect (Nowroozi & Brainerd, 2013).

We therefore examined the variation in internal vertebral morphology in benthic, demersal, and pelagic fishes across actinopterygian fishes. To identify differences in vertebral morphology of these fishes due to habitat, we controlled for shared evolutionary history.
using the Rabosky et al. (2018) phylogenetic tree. Based on our hypothesis that internal morphological differences support behavioral differences across different habitat groups, we predicted that pelagic fishes should have a more closed internal vertebral morphology minimizing space for soft material and leading to a stiffer backbone, and that benthic and demersal species should have a morphology that allows for more flexibility in both mechanics and behavior.

2 | METHODS

We used micro-computed tomography (μCT) scans to measure six different vertebral features from 79 species across a trimmed version of the Rabosky et al. (2018) phylogeny (Figure 1). We classified these species into benthic (species in contact with the substrate most of the time), pelagic (open water swimmers), and demersal (species close to the substrate but not resting on it all of the time) habitats. For all of the species we examined that overlapped those from Friedman et al. (2020), we used their classification. For other species, we classified habitat based on descriptions of each species from guidebooks or relevant journal articles (Allen et al., 2002; Bailey, 1994; Basolo, 1990; Fine et al., 1987; Gilbert & Williams, 2002; Jaafar et al., 2004; Lowry et al., 2005; Magid, 1967; Matsui & Rosenblatt, 1987; McGinnis & Alcorn, 2006; Mérigoux et al., 1998; Page et al., 1991; Pearcy et al., 1982; Phomikong et al., 2015; Pietsch & Orr, 2015; Proctor & Lynch, 2011; Rodríguez-Olarte et al., 2011; van der Steen & Albert, 2018; Vasek et al., 2008).

We used histological data to compare the soft tissue anatomy across representative species from benthic, pelagic, and demersal habitats. No animals were sacrificed for histological analyses. Instead, all samples were donated from the Friday Harbor Labs Ichthyology collection. All specimens arrived fixed in 10% buffered formalin solution and stored in 70% ethanol (EtOH). The vertebral columns from Anoplarchus purpurascens (prickleback, demersal), Cymatogaster aggregata (perch, pelagic), and Myxocephalus polyacanthocephalus (sculpin, benthic) were carefully dissected and three full vertebrae near 50% total length were removed. These vertebrae were chosen as they are the most representative of each species’ vertebral morphology and not as influenced by other intrinsic factors such as head or tail shape. The vertebrae were decalcified in
a 10% EDTA solution for 10 days. After full decalcification, vertebrae were rinsed with dH\textsubscript{2}O for 48 h before undergoing an ethanol dehydration series to 95% EtOH. Each sample was then infiltrated and embedded with JB4 embedding media (14272-00) following Electron Microscopy Sciences JB4 embedding media protocol. Each species was sectioned at thicknesses between 3 and 3.5 μm with a glass knife, stained with Lee's Basic Fuchsin and Methylene Blue, and mounted to glass slides with permount. We imaged slides on a Nikon Eclipse E600 microscope with a Micropublisher 5.0 RTV camera. Whole slice histological images were tiled and stitched together and color balanced in Adobe Photoshop. This method is a composite that allows for high-resolution images to be taken of large histological slices while still viewing the whole section.

### 2.1 | Vertebral measurements

Most of the μCT scans were downloaded as image stacks off of Morphosource (https://www.morphosource.org/) or the “CT Scans - #ScanAllFish” (Adam Summers, https://osf.io/ecmz4/) database. Additional specimens were scanned at the Karel F. Liem Bioimaging facility using their Bruker Skyscan 1173 (Bruker microCT). Full identification of all scans is given in Table S1.

We measured vertebral features for vertebrae at 0.4–0.9 of the fish's standard length (BL), in increments of 0.1. We first measured the location of the snout and the final vertebra before the caudal fin to estimate standard length, then used that to calculate the locations at 0.4–0.9 BL. We then placed landmarks at seven points on sagittal sections through the center of vertebrae at each location if a vertebral centrum was present (Figure 2a). Based on those points, we computed the anterior and posterior cone diameter and angle, the notochordal foramen diameter, and the centrum body length (CBL) (formulas in Table 1). We also estimated fineness by dividing standard length by the maximum body width. All linear measurements were normalized to body length by dividing by the fish's standard length before we ran our statistical analyses.

### 2.2 | Statistical analysis

To examine differences in vertebral morphological measurements as they relate to pelagic, demersal, or benthic habitat categories, as well as to control for shared evolutionary history, we performed phylogenetic generalized least squares analyses (Adams & Collyer, 2018), with the phylogenetic effect modeled using a random Brownian correlation.

We first normalized the $x,y$ coordinates of each point ($P_1 - P_x$) for each vertebra (42 points in total) by subtracting the centroid of each individual vertebra, then dividing the coordinates by the standard length of each specimen. We aligned the coordinates using a generalized Procrustes analysis (Gower, 1975), followed by a multivariate phylogenetic generalized least squares analysis using a residual randomization procedure (Collyer & Adams, 2018) to identify differences relative to habitat and fineness. Following the initial multivariate test, we also tested the coordinates for each vertebra individually (seven points per vertebra), to examine differences along the body. Because fineness varies among habitats, we were not able to include an interaction effect between habitat and fineness in our statistical models.

We then computed the six shape parameters (Table 1) and took their mean values across all vertebrae for each species. We used a multivariate PGLS analysis to compare all the parameters simultaneously, then ran univariate tests on each parameter individually.

If we found that the habitat effect was significant, we then ran the same PGLS model on each pair of habitats separately to examine the pairwise differences. In this case, we controlled for multiple comparisons with a Bonferroni correction.

We fit the models using R (version 4.2.0; https://www.R-project.org/) with packages geomorph (version 4.0.4; Baken et al., 2021) and RRPP (version 1.3.0; Collyer & Adams, 2018). All code and data are available with DOI: 10.25833/772t-cw09 and on Github (https://tytell.github.io/BaxterVertEvol/). The main data table is available in Table S1.

### 3 | RESULTS

We examined 79 species of actinopterygian fishes based on publicly available CT scans, aiming to cover as many families as possible across the phylogeny. These species thus represented 68 different families. We classified the species’ habitats as benthic (22% of species in our data set), demersal (49%), or pelagic (29%). Marine species made up 66% of the data set, and freshwater species were 33%, and we had one anadromous species.

#### 3.1 | Mean morphology of vertebrae differ in fishes from different habitats

Species from different habitats had different fineness (Figure 3; $p = .048$). Pelagic species tended to be more elongate (higher fineness ratio) than benthic species, though, due to the Bonferroni correction, this difference was not significant.

Next, we compared the landmarks for all digitized points for all vertebrae (42 points for each species), using a generalized Procrustes analysis to align the points, and a multivariate residual randomization procedure to compare them in a phylogenetic context (Baken et al., 2021; Collyer & Adams, 2018). We measured vertebral postions from 0.4 to 0.9 BL in steps of 0.1 BL (six positions per species). Species from different habitats had significantly different vertebral morphology ($p = .045$; Table 2). Fineness was not significant ($p = .3$).

Overall, we found that these mean shape parameters are different in different habitats ($p = .001$; Table 3, Figure 4) and that they also depend on the fineness ratio ($p = .001$; Table 3). The notochordal foramen diameter differed among fishes from different habitats ($p = .001$; Figure 4a; Table 4), but did not depend on fineness ratio.
FiguRe 2  Vertebral centra measurements. (a) Measurements taken from vertebral centra, based on the numbered points. (a) d is foramen diameter, CBL is centrum body length, \(a_{anterior}\) is anterior cone angle, \(D_{anterior}\) is anterior cone diameter, \(a_{posterior}\) is posterior cone angle, \(D_{posterior}\) is posterior cone diameter. See Table 1 for formulas. (b–d) Rendering of vertebrae in lateral and frontal views from three representative species. Lateral views show are vertebrae bisected down the middle along the sagittal plane. (b) Myoxocephalus polyacanthocephalus (benthic), (c) Lumpenus sagitta (demersal), (d) Cymatogaster aggregata (pelagic).

TABLE 1  Summary of formulas for the vertebral morphology variables (Figure 2a)

| Variable                      | Symbol | Formula        |
|-------------------------------|--------|----------------|
| Notochordal foramen diameter  | \(d\)  | \(\|P_6 - P_7\|\) |
| Centrum body length           | CBL    | \(\frac{1}{2}(|P_{1x} - P_{3x}| + |P_{2x} - P_{4x}|)\) |
| Anterior cone angle           | \(a_{anterior}\) | \(\frac{|P_{1x} - P_{3x}|^2 + |P_{2x} - P_{4x}|^2}{|P_{1x} - P_{3x}| + |P_{2x} - P_{4x}|}\) |
| Anterior cone diameter        | \(D_{anterior}\) | \(\|P_3 - P_4\|\) |
| Posterior cone angle          | \(a_{posterior}\) | \(\frac{|P_{5x} - P_{7x}|^2 + |P_{6x} - P_{8x}|^2}{|P_{5x} - P_{7x}| + |P_{6x} - P_{8x}|}\) |
| Posterior cone diameter       | \(D_{posterior}\) | \(\|P_1 - P_2\|\) |

Note: \(P_i\) the \((x, y)\) coordinates of the \(i\)th point on each vertebra (Figure 2a), \(|x|\), length of the vector \(x\).

Pelagic fishes had smaller foramina than benthic fishes (\(p = .001\)). The posterior cone angle was also significantly different in fishes from different habitats (\(p = .01; Figure 4c; Table 4\)). Pelagic fishes had smaller posterior cone angles than benthic fishes (\(p = .005\)). Anterior cone angle was not significantly different in fishes in different habitats (\(p = .228; Figure 4b\)). Fineness ratio did not affect either cone angle significantly.

The CBL and anterior and posterior cone angles varied significantly among habitats (\(p < .011\) in all cases; Table 4) and also decreased significantly as fineness ratio increased (Figure 4d–f). Pelagic species had significantly longer vertebrae (larger CBL) than benthic (\(p = .011\)), but benthic species had larger anterior and posterior cones (\(p = .005\) and .006) than pelagic. These differences were small and may not be functionally relevant. Demersal species varied more than the other groups, and thus did not show any significant differences.

Figure 3 shows the mean shape of the internal parts of the vertebrae. Posterior cone angle (\(a_{posterior}\)) and foramen diameter (\(d\)) are both significantly smaller in pelagic species, while CBL is longer in pelagic species.
species. Both anterior and posterior cone diameters were slightly, but significantly, larger in benthic species.

3.2 Patterns of vertebral morphology along the body differ in fishes from different habitats

Our analysis showed that the way vertebral morphology varied along the body differed in our three habitat groups. To compare these patterns, we compared the landmarks for vertebrae at each location along the body among all of the species. We found that vertebrae at all locations were significantly different in different habitats ($p < .039$ in all cases; Table 2).

TABLE 3 Results of multivariate phylogenetic generalized least squares analyses for differences in overall means for each species relative to habitats and fineness.

| Effect      | df | Pillai's trace | Z    | p    |
|-------------|----|----------------|------|------|
| Habitat     | 2  | 0.621          | 3.48 | .001 |
| Fineness    | 1  | 0.515          | 4.57 | .001 |

Note: Significant effects are shown in bold. Variables included are centrum body length, foramen diameter, anterior and posterior cone angles and diameters. $N = 79$ species.

We then calculated the same shape parameters for each vertebra individually and plotted them as a function of location. Figure 6 shows mean ± standard error for posterior cone angle, foramen diameter, and CBL relative to position along the body. Posterior cone angle does not vary substantially along the body (Figure 6a). Foramen diameter tends to be largest near the mid-body and decreases toward the tail (Figure 6b). CBL also decreases toward the tail, particularly in benthic species (Figure 6c).

3.3 Diversity of soft tissue anatomy

To examine the underlying tissues more thoroughly, we also imaged tissue structures in vertebral centra using histological methods (Figure 7). We chose three representative fish species ($M. polyacanthocephalus$, benthic, Figure 7a–c; $A. purpurescens$, demersal, Figure 7d–f; $C. aggregata$, pelagic, Figure 7g–i) based on availability in the Friday Harbor Labs ichthyological collection. In all species, the amphicoelous vertebral centra are composed of cortical bone (CB), with two fluid vacuoles, and various notochordal tissues (Figure 6). Each vertebra has a pair of encapsulating complexes (EC) located on the anterior and posterior sides of the centra. These ECs are composed of three main tissues: (1) the EVL, a membranous ligament that attaches adjacent vertebrae, (2) the EE, a thin layer...
composed of elastin, and (3) a collagen-containing FS. The largest differences in soft tissue morphology were apparent in the intervertebral spaces for each species. The volume of notochordal tissues (NC) was lowest in *A. purpurescens*, the demersal representative, whereas both *C. aggregata* (pelagic) and *M. polyacanthocephalus* (benthic) had qualitatively similar notochordal volumes. Histological sectioning revealed clearly defined notochordal foramen in all three species. In *A. purpurescens*, the foramen is filled with a relatively thin notochordal cell mass composed of elongated notochordal cells ([Figure 7f](#) and an epithelium extending the length of the intervertebral junction. In both *C. aggregata* (pelagic) and *M. polyacanthocephalus* (benthic), we see rounded and elongated notochordal cells as part of the notochordal cell mass ([Figure 7c,i](#)). Similarly, the cortical bone of the vertebrae in pelagic and benthic representatives is much more robust than in our demersal representative.

The cellular morphology of the external intervertebral ligament was qualitatively similar across all three species but the EE and FS varies across species, and these were the two tissues responsible for changing tensile properties between vertebral elements. The FSs, sometimes referred to as the notochordal sheaths, of *C. aggregata* (pelagic, [Figure 7h](#) and *M. polyacanthocephalus* (benthic, [Figure 7b](#)) were similar and composed primarily of acellular tissue. In *A. purpurescens* (demersal, [Figure 7e](#)), the FS was smaller and thinner, though still acellular in morphology. The EE, the only elastin-containing tissue between vertebrae, was again most similar between the benthic and pelagic species and is notably asymmetrical in the demersal species ([Figure 7](#)).

| Effect                  | df  | $r^2$ | $F$  | $Z$  | $p$  | Pairwise comparisons<sup>a</sup> |
|-------------------------|-----|-------|------|------|------|---------------------------------|
|                         |     |       |      |      |      | $p - b^b$ | $d - b^c$ | $p - d^d$ |
| Fineness                |     |       |      |      |      |       |
| Habitat                 | 2   | 0.09  | 3.6  | 1.6  | .048 | 0.040 | 0.152 | 0.434 |
| Centrum body length     |     |       |      |      |      |       |
| Habitat                 | 2   | 0.11  | 6.8  | 2.6  | .007 | 0.011 | 0.287 | 0.416 |
| Fineness                | 1   | 0.23  | 29   | 3.6  | .001 | 0.001 | 0.135 | 0.144 |
| Foramen diameter        |     |       |      |      |      |       |
| Habitat                 | 2   | 0.29  | 15   | 3.5  | .001 | 0.001 | 0.135 | 0.144 |
| Fineness                | 1   | 0.03  | 2.9  | 1.4  | .081 | 0.943 |
| Anterior cone angle     |     |       |      |      |      |       |
| Habitat                 | 2   | 0.04  | 1.5  | 0.77 | .228 | 0.254 | 0.471 |
| Fineness                | 1   | 0.00  | 0.0059 | −1.6 | .943 |       |
| Posterior cone angle    |     |       |      |      |      |       |
| Habitat                 | 2   | 0.11  | 4.8  | 2.2  | .010 | 0.005 | 0.254 | 0.471 |
| Fineness                | 1   | 0.00  | 0.00028 | −2.1 | .989 |       |
| Anterior cone diameter  |     |       |      |      |      |       |
| Habitat                 | 2   | 0.22  | 20   | 3.7  | .005 | 0.005 | 0.583 | 0.563 |
| Fineness                | 1   | 0.24  | 44   | 4.0  | .001 | 0.006 | 0.515 | 0.725 |
| Posterior cone diameter |     |       |      |      |      |       |
| Habitat                 | 2   | 0.25  | 22   | 3.6  | .007 | 0.006 | 0.515 | 0.725 |
| Fineness                | 1   | 0.19  | 34   | 3.8  | .001 | 0.006 | 0.515 | 0.725 |

Note: Significant effects are shown in bold. df, degrees of freedom; $r^2$, a measure of the goodness of fit of the statistical model; $F$, statistical $F$ parameter; $Z$, a measure of the overall effect size; $p$, probability of seeing an effect that large due to random variation.

<sup>a</sup>The cutoff for a significant pairwise effect is $p < .016$ due to Bonferroni correction.

<sup>b</sup>Pelagic relative to benthic.

<sup>c</sup>Demersal relative to benthic.

<sup>d</sup>Pelagic relative to demersal.

DISCUSSION

We have shown that fish species that reside in benthic, pelagic, and demersal habitats have vertebrae with significantly different morphology. Mean notochordal foramen diameter and posterior cone angle were significantly different for fishes from the different habitats. CBL and the anterior and posterior cone diameters not only tended to be smaller in more elongate fishes but also showed some significant differences across habitats. In general, fishes from pelagic habitats tended to have shallower cones and narrower foramina than fishes from the other habitats, while benthic species have wider cones ([Figures 4a,b](#) and Figure 5). The morphology of the soft
tissue around and within the vertebral centra was different in our representative demersal species (Anoplarchus purpurescens) then the pelagic and benthic representatives (Figure 7).

On average, this means that pelagic fishes have less space for notochordal material and other flexible tissues, probably making their backbones passively stiffer. In both physical models and tests on excised vertebral columns, these shallower cones and narrower foramina are associated with stiffer intervertebral joints (Donatelli et al., 2021). The connection between body mechanics and swimming performance is still being examined, but in robotic models, adding stiff vertebral centra and reducing the amount of flexible notochordal material increased the stiffness of an artificial vertebral column and led to higher speed steady swimming (Long et al., 2011). More recently, increasing the stiffness of a tuna robot increased its swimming speed, up to an optimum, but then only decreased the speed slightly as stiffness increased further (Zhong et al., 2021). Thus, we suggest that these differences result in pelagic fishes having overall stiffer backbones, which may help them to swim continuously (Lauder, 2015). Conversely, demersal and benthic species have larger cones and relatively more space for notochordal material, making their backbones more flexible. Many benthic species, such as sculpins, are ambush predators, catching prey with high accelerations that are often accompanied by large body curvature, which could be facilitated by the larger cone angles in their vertebrae. Additionally, their more flexible vertebral columns may allow them more behavioral flexibility. For example, a greater range of speeds are possible if an animal can modulate its body stiffness (Wolf et al., 2020; Zhong et al., 2021). The more passively flexible bodies of demersal and benthic fishes may thus give them more active control of their body stiffness leading to a greater diversity of swimming modes. As these fish range from ambush predators like the sculpins to burrowers like flatfish, greater control of body stiffness would be advantageous.

Histology on three representative species revealed the diversity of soft tissues across depths, and while our examples represent specific species, they do point out key traits that are likely beneficial for their specific habitats. Our representative pelagic and benthic species have more robust skeletal structure and more notochordal material than the demersal species (Figure 6). Our pelagic representative, Cymatogaster aggregata, spends most of its time in the water column, swimming for prolonged periods of time, migrating, and avoiding predators. A robust vertebral column may contribute to an increase in vertebral stiffness that lends to more efficient continuous swimming—a beneficial trait for pelagic species. On the other hand, in benthic fishes, the more robust skeletal structure (Figure 6),
plus the overall greater posterior cone angle (Figure 4), may result in a vertebral column that is passively more flexible than the pelagic species, but can be selectively stiffened by contractions of the surrounding muscle. This would be ideal for bursts of acceleration to escape predators or catch prey – behavior commonly seen in *Myoxocephalus polyacanthocephalus* (our histological representative for benthic environments). Demersal fish, such as *Anoplarchus purpurascens*, appear to have characteristics that overlap both pelagic and benthic species, particularly an intermediate posterior cone angle (Figure 4). Many species of stichaeids, the family that includes *A. purpurascens*, swim through several types of complex environments, including burrowing into the sediment, and while they are constantly moving these elongate fishes are often after sedentary prey. The combination of a reduction in notochordal material and a less robust skeletal morphology (Figure 6) perhaps point to a generalized vertebral shape for a variety of locomotor modes such as navigating the seafloor, swimming up into the water column, burrowing, and other complex behaviors.

In our data set, internal vertebral morphology is related to body shape, but in a complex way. CBL (as normalized to overall body length) was shorter in more elongate fishes (higher fineness), reflecting the fact that these species often have more vertebral than less elongate species (Ward & Brainerd, 2007). Anterior and posterior cone diameters were also smaller in elongate fishes (Figure 4d–f, Table 4). These parameters were significantly different across habitats, with benthic and pelagic fishes relatively close to one another. Demersal species had shorter centra and smaller diameter cones, on average, but also had a wider variation in these parameters, likely reflecting the high variability in behavior in demersal species. Other parameters, including posterior cone angle and foramen diameter,
did not correlate significantly with fineness, but were significantly different in different habitats.

Pelagic species were significantly more elongate than benthic species (Figure 3), similar to what has been observed in previous studies (Burress et al., 2017; Cooper et al., 2010; Friedman et al., 2020; Gerry et al., 2011; Hollingsworth et al., 2013; Tavera et al., 2018; Willacker et al., 2010). For pelagic fishes with narrow and shallow bodies, there is less skin and muscle in the body’s cross-section than for benthic fishes. Thus, a smaller foramen diameter and larger cone angle of an individual vertebra may have a higher contribution to the mechanical properties of the whole body because they represent a larger fraction of the cross-section. This might mean that the whole-body properties of pelagic fishes are more influenced by the mechanics of the backbone than those of benthic fishes.

The pattern of vertebral shape along the body differs for fishes from different habitats (Figure 6). For all fishes, the foramen diameter has a maximum at some point along the body, and then tends to get smaller in more posterior and more anterior vertebrae. The foramen diameter tends to be more uniform along the body for pelagic fishes. CBL also varies along the body. Pelagic species again have vertebrae with relatively uniform length, whereas demersal and benthic species tend to have longer vertebrae more anteriorly and shorter vertebrae more posteriorly. The functional significance of these changes in vertebral shape along the body is hard to interpret. In benthic and demersal fishes, it may be that the anterior vertebral column is more flexible than the posterior portion because the space for flexible materials is smaller near the tail, whereas in pelagic fishes, the mechanical properties may be more uniform along the body. Though a stiffer body leads to faster speeds at low undulation frequencies, bodies that are more flexible toward the tail result in faster speeds at high frequencies and are also more efficient (Lucas et al., 2015). Mapping changes in vertebral morphology may therefore have implications in our understanding of locomotor efficiency, though other factors likely play a role in whole-body stiffness, like the mechanics of the skin and muscle as well as the shape of the body, which tends to vary in thickness particularly in benthic and demersal fishes (Friedman et al., 2020).

In contrast, demersal and benthic fishes tend to spend more time maneuvering around complex obstacles like coral reefs (Larouche et al., 2020). This often requires the body to bend with much higher curvature than during steady swimming. A more flexible backbone may permit these high curvatures. In addition, larger cone angles may allow the intervertebral joints to flex to a higher angle before the bone of the vertebra begins to limit bending (Nowroozi & Brainerd, 2013). Rapid turns or those with a small turning radius also typically require high curvature in the anterior body. For example, consider C-start escape responses, which require much higher anterior body curvature than steady swimming (Domenici & Blake, 1997; Gerry et al., 2012). This anterior curvature may be facilitated by the differences in vertebral morphology from anterior to posterior in benthic and demersal fishes (Figure 6).

Overall, the differences we observed are likely the result of selective processes that operate on the entire body as a functional unit, not selection for specific parameters that we measured. Moreover, body shape and vertebral morphology are likely to be coupled developmentally. The patterns we observe may therefore be a consequence of some other functional specialization (e.g., Gould & Lewontin, 1979), perhaps for body shape, but they are nevertheless relevant because they help distinguish species from different environments and may point to broader functional differences in those species.

5 | CONCLUSION

The internal morphology of the vertebrae of actinopterygian fishes differs across species with several overlapping patterns in fishes classified as benthic, demersal, and pelagic. These morphological differences were especially distinct between species from benthic and pelagic habitat groups. Overall, the differences seem to be consistent with commonly observed behavioral differences between benthic and pelagic fishes, where benthic fishes tend to maneuver more around complex substrates and do not tend to swim steadily, and pelagic fishes swim constantly but do not need to maneuver as much as benthic or demersal fishes. This work adds to the body of literature (Donatelli et al., 2021; Porter et al., 2007, 2014; Porter & Long, 2010) suggesting that creating a model of vertebral morphology in fishes has implications in modeling kinematic diversity. This will be especially powerful in developing an understanding of the kinematic patterns of extinct fishes or extant fishes that are difficult to keep.

AUTHOR CONTRIBUTIONS
Dana Baxter: Data curation (lead); formal analysis (equal); investigation (lead); software (supporting); visualization (supporting); writing – original draft (lead); writing – review and editing (equal).
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CONFLICT OF INTEREST
The authors declare that they have no competing interests.

OPEN RESEARCH BADGES

This article has earned an Open Data badge for making publicly available the digitally-shareable data necessary to reproduce the reported results. The data is available at https://tytell.github.io/BaxterVertEvol/.

DATA AVAILABILITY STATEMENT
All data and analysis scripts are available with DOI https://doi.org/10.25833/772t-cw09. The most recent version of the analysis scripts is also available at GitHub (https://github.com/tytell/BaxterVertEvol).

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