Title: Foretelling the flex - vertebral shape predicts behavior and ecology of fishes

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**Summary statement:** We modeled swimming kinematics and body mechanics of several fish species of varying habitat and body shape based on measurements of internal vertebral morphology.

**Data Availability**: The data underlying this article will be made available after proofs are finalized on GitHub at https://github.com/CDonatelli/FortellingTheFlex.

# Abstract

One key evolutionary innovation that separates vertebrates from invertebrates is the notochord, a central element that provides the stiffness *needed* for powerful movements. Later, the notochord was further stiffened by the vertebrae, cartilaginous and bony elements, surrounding the notochord. The ancestral notochord is retained in modern vertebrates as intervertebral material, but we know little about its mechanical interactions with surrounding vertebrae. In this study, the internal shape of the vertebrae – where this material is found – was guantified in sixteen species of fishes with various body shapes, swimming modes, and habitats. We used micro-computed tomography to measure the internal shape. We then created and mechanically tested physical models of intervertebral joints. We also mechanically tested actual vertebrae of five species. Material testing shows that internal morphology of the centrum significantly affects bending and torsional stiffness. Finally, we performed swimming trials to gather kinematic data. Combining these data, we created a model that uses internal vertebral morphology to make predictions about swimming kinematics and mechanics. We used linear discriminant analysis (LDA) to assess the relationship between vertebral shape and our categorical traits. The analysis revealed that internal vertebral morphology is sufficient to predict habitat, body shape, and swimming mode in our fishes. This model can also be used to make predictions about swimming in fishes not easily studied in the lab, such as deep sea and extinct species, allowing the development of hypotheses about their natural behavior.

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# 1 Introduction

One of the key evolutionary innovations that separates vertebrates from invertebrates is the notochord, a central element that provides the stiffness needed for fast and powerful locomotion (Annona et al., 2015; Koehl et al., 2000; Long, 1995; Long et al., 2002; Symmons, 1979). Later in evolutionary history, the notochord was mostly replaced by segmented vertebrae, though it is still present in all vertebrates (Annona et al., 2015). In bony fishes, the extent to which the notochord is present varies. In some fish, it simply makes up the intervertebral material. In others, it is the main structural component of the vertebral column and is present as a continuous tube running through the center of sometimes poorly mineralized vertebral centra. Because the vertebral column is important for swimming mechanics, it has been examined in several capacities to assess how anatomical variations impact the mechanics of fish swimming with one of the most commonly measured variables being bending stiffness. Studies of vertebral column bending stiffness have looked at single joints (Hebrank et al., 1989; Long, 1991; Long, 1995; Long et al., 1997; Nowroozi et al., 2012), as well as the entire vertebral column (Long et al., 2002; Porter et al., 2016). In this study, we build upon this literature by examining the functional effects of vertebral morphology, particularly the intervertebral elements (i.e., the notochord), as the relationship between these elements and swimming performance have not been systematically investigated.

Most fishes with discrete vertebral centra exhibit similar morphological characteristics within the centra themselves (Larem, 1975; Schaeffer, 1967). Though the external body of a centrum (the central core) is cylindrical, the interior has two opposite facing cones oriented such that the wide ends are anteriorly and posteriorly directed, and the narrow ends meet in the middle to form a canal through the center (Figure 1B-E). This results in an hourglass-like shape when a centrum is sectioned down the midsagittal plane (Figure 1E). Attached to the dorsal and ventral sides of the main body of the centra are the neural and hemal spines, respectively. These spines start to appear on centra near the head and continue all the way to the caudal-peduncle. Neural spines are present on most vertebra, while hemal spines appear only after the cloaca. Anterior to the caudal-peduncle, the ribs protrude ventro-laterally and serve as protection for vital organs. Smaller and closer to the centra body are small interlocking spines called zygapophyses. These small spines vary greatly between species, but generally interlock adjacent centra along the length of the body (Liem et al., 2001).

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Studies exploring the mechanical properties of the vertebral column and its anatomical components have resulted in many hypotheses about its role in swimming. Most of the work focusing on specific anatomy, like the intervertebral joints, has used larger swimmers such as the blue marlin (Makaira nigricans) (Hebrank et al., 1989; Long, 1991) or the saddleback dolphin (Delphinus delphis) (Long et al., 1997), partially because it is methodologically easier to make mechanical measurements on larger species. These studies have revealed that, in both the saddleback dolphin and the blue marlin, the stiffness of the intervertebral joint increases from the cranium to about three guarters down the length of the body, and then decreases from that point to the caudal region (Aleyev, 1977; Hebrank et al., 1989; Long, 1991). Hebrank et al. (1989) also found that the zygapophyses in the blue marlin increase stiffness substantially when bending dorso-ventrally, but less so when bending laterally (Hebrank et al., 1989). In short, vertebral morphology varies within an animal, and this variation has a direct impact on the mechanical properties of individual intervertebral joints. 

Regarding the vertebral column as a whole and its effect on swimming, Porter et al. (2016) found that the bending stiffness of the spiny dogfish (Squalus sucklei) vertebral column depended nonlinearly on frequency, suggesting that the vertebral column behaves as either a spring or a brake depending on swimming speed (Porter et al., 2016). Similarly, Long et al. (1995 and 2002) determined that the mechanical properties of the hagfish (Myxine glutinosa) notochord, in conjunction with the body's musculature, help the animal adjust its resonant bending frequency for more efficient swimming; in the sturgeon (Acipenser transmontanus), notochord angular stiffness, not morphology, is inversely correlated with swimming kinematics (Long, 1995; Long et al., 2002). Though these studies were mainly conducted on softer, more continuous structures, they reveal that the mechanics of the vertebral column is related to swimming performance. 

 In addition to the studies discussed above, there have been many more investigating the connection between swimming kinematics and the motion of the vertebral column, both as a whole and with a focus on individual intervertebral joints (Nowroozi and Brainerd, 2014; Porter et al., 2009; Porter et al., 2014; Porter et al., 2016). In a meta-analysis of 20 different studies, spanning 28 species, Nowroozi and Brainerd (2014) found that the vertebral column experiences increasing degrees of bending from cranium to caudal region during steady swimming in undulatory swimmers. Though these observations do not directly discuss the shape or structural components of the vertebral column, we can use this information about undulatory locomotion in conjunction with more detailed morphological and mechanical studies to create models of locomotion in different species.

In addition to tests on biological specimens, physical models have played an important role in understanding the vertebral column (Hirokawa et al., 2011; Lucas et al., 2015; Shelton et al., 2014). Hirokawa et al. (2011) found that changing the morphology of a biomimetic vertebral column causes changes in the swimming speed and frequency of a bioinspired robot, MARMT (Hirokawa et al., 2011). Specifically, they found that decreased intervertebral joint length increases stiffness in a vertebral column inspired tail. In swimming flexible plastic foils, Lucas et al. (2015) found that foils with uniform stiffness swam more slowly than those with a stiffness gradient from anterior to posterior. This, along with Nowroozi and Brainerd's 2014 meta-analysis, suggests that the way the vertebral column changes from head to tail could affect swimming performance (Lucas et al., 2015; Nowroozi and Brainerd, 2014). These studies also highlight the importance of using physical models to understand how individual morphological parameters affect swimming. By controlling for things such as material and individual variation, the results of studies using models can help us understand the links between specific aspects of internal vertebral morphology and swimming behavior.

Though many studies demonstrate that the mechanics of the vertebral column can have an important impact on swimming, very few of them have considered the internal shape of the vertebral centra and the role of the notochord. Because the notochordal material is much more flexible than the bony vertebrae, its properties may be more important in determining overall flexibility. Additionally, these studies have not examined multiple species to draw conclusions about how the common internal morphology of the vertebral column could relate to overall swimming behavior. In this study, we measured the vertebral morphology of sixteen species of fishes with diverse body shapes that live in habitats ranging from intertidal to subtidal. To evaluate the functional role of the internal vertebral morphology, we used micro computed tomography to quantify the internal shape of the vertebrae. We then created and mechanically tested physical models of intervertebral joints with morphologies spanning the measured range. Next, we mechanically tested actual vertebral joints dissected from a subset of the species in this study. Finally, using the data from swimming trials performed on thirteen species of fish, we created a statistical model that uses vertebral morphology to make predictions about the body shape, habitat, and swimming mode of these fishes.

- 93 Materials and Methods
- 94 Swimming Trials and Kinematics

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Study Specimens:

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We collected individuals from sixteen morphologically and ecologically diverse species of fish mostly
native to the Salish Sea surrounding the San Juan Islands (Washington, USA). Specimens were collected
using dip nets, beach seines, and otter trawls (Table 1) and then housed in sea tables connected to the
flow through system at Friday Harbor Labs according to University of Washington IACUC protocol 4238-
03.
Swimming Trials:
We filmed fish swimming around a track in a modified sea table (Figure 2). Once placed in the track,
individuals were given time to adjust to the new tank and then allowed to swim around the track at their
preferred steady swimming speed. We mounted a GoPro (GoPro Hero 5, GoPro, San Mateo, CA) above
the tank to capture a dorsal view of the fish swimming through the filming area. For each species, we
filmed five individuals swimming over several days until we had five steady swimming trials with at least
five tailbeats per trial if possible. Because some fish could swim through the filming area in less than five
tailbeats, we recorded more trials until we had a total of twenty-five tailbeats per fish.
Kinematic analysis:
Video data was processed using custom Matlab code that traces the midlines of the fish in each frame of
a swimming trial video and then uses the traces to extract kinematic parameters such as swimming
speed (distance traveled in body lengths (BL) per second), tail beat frequency (tail beats per second, Hz),
tail beat amplitude (BL), and stride length (the distance traveled per tail beat, BL). For the purposes of
this study, we only asked the code to output amplitude (Figure 3). We also examined patterns of long-
axis body twisting (wobble), in several elongate species used in this study. Wobble is a unitless ratio
describing the amount of twisting where 0 is no twisting and 1 is body twisting of 90 degrees (Donatelli
et al., 2017) (Figure 3).
CT Scanning and Morphometrics
Study Specimens:

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We collected morphological data from both museum and freshly fixed specimens representing sixteen

species of fishes. Individuals prepared for scanning were euthanized in a lethal dose (0.5g/L) of MS222

according to University of Washington IACUC protocol 4238-03 and stored in 70% ethanol for at least 24

1 2					
3 4 5 6	126	hours before scanning. For fish that were not available in the lab, we borrowed museum specimens			
	127	from the Burke museum (Table 1) (Burke Museum of Natural History and Culture, Seattle, WA.).			
7 8	128	Scanning:			
9	129	All fishes were scanned at the Karel F. Liem Bio-Imaging Center at Friday Harbor Laboratories (Friday			
10 11	130	Harbor, WA, USA). Before scanning, we labeled specimens with radiopaque markers for easy			
12 13	131	identification during post-scan processing. Then, we wrapped multiple individuals in cheesecloth soaked			
14	132	in 70% ethanol and packed them into a 3D printed plastic tube. Once packed, we scanned the tubes			
15 16	133	using a Bruker Skyscan 1173 (Bruker, Belgium, Germany) at 65 kV and 123 $\mu$ A with a voxel size ranging			
17 18	134	from 20.3 to 33.5 $\mu m$ depending on the size of the specimens (higher resolution used for smaller			
19	135	specimens).			
20 21 22	136				
22 23	137	We reconstructed the scans using NRecon software by Bruker (2016, Bruker, Belgium Germany). Once			
24	138	reconstructed, the stacks were converted into nrrd files using ImageJ (Schneider et al., 2012) and			
25 26	139	imported into Slicer 3D (BWH and Contributors, 2019). By examining nrrd files of each specimen in the			
27 28	140	three planes of traditional x-ray slices (Figure 1B) as well as a 3D reconstruction of the fish (Figure 1A, C)			
29	141	in Slicer 3D (BWH and Contributors, 2019), we were able to collect morphometric data for each			
24         25         26         27         28         29         31         32         33         34         35         36         37         38         41         42         43         44         45         46         47         48         50         53         54         55         56         57         58	142	specimen. In total, we scanned and collected morphometric data from 48 specimens, with three			
	143	specimens representing each of our sixteen species.			
	144				
	145	Morphometrics:			
	146	For each scanned specimen, we placed digital landmarks on multiple vertebrae along the length of the			
	147	body in Slicer 3D. Vertebrae were selected at 10% body length intervals (Figure 1A), resulting in			
	148	morphometrics of 8-9 vertebrae sampled per individual. We placed six markers on each selected			
	149	vertebra (Figure 1E) and converted the coordinates to measurements in body length (BL) units. Markers			
	150	1, 2, 3, 5, and 6 were placed on physical landmarks of the centrum (Figure 1E). Marker 4 was placed			
	151	between markers 5 and 6 in the center of the notochordal foramen and was used as the apex to			
	152	calculate centrum cone angle. We used measurements in terms of body length to correct for the			
	153	difference in size of our study specimens. The resulting measurements gave us length of the vertebral			
	154	centrum (CBL), diameter of the centrum's posterior facing cone (D), diameter of the centrum's central			
	155	canal (d), and angle of the centrum's posterior facing cone ( $ heta$ ) (Figure 1D). In addition to measuring the			
	156	vertebral column, we also assigned each species a body shape, habitat, and swimming mode based on			
	157	the literature and our own observations (Table 2) (Froese and Pauly, 2019; Lamb and Edgell, 2010).			
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2 3 4 5 6 7 8 9 10 11 12	158	
	159	Once we collected centrum measurements in Slicer, we then used Matlab to calculate the ratio of soft to
6 7	160	hard material in each vertebral centra. The area of the hard material (bone) was calculated using the
8	161	area of a triangle made using points that defined centrum length (i.e., points 2 and 3; Figure 1E), as well
9 10	162	as one of the points that defined foramen diameter (i.e., point 6; Figure 1E). We calculated the area of
11 12	163	bone on both the dorsal and ventral side of the centra and summed the results to get "bone area".
13 14	164	Similarly, the area of the soft material was calculated using the points that defined cone diameter (i.e.,
14	165	points 1 and 2; Figure 1E), as well as the point that defined the center of the foramen (i.e., point 4;
16 17	166	Figure 1E). We did these calculations for both anterior and posterior centrum cones and summed the
18 19	167	areas to get our "soft area" measurement. Once total areas were calculated, we divided the area
20	168	occupied by soft material by the area occupied by bone to get the ratio of soft to hard material (S:H).
21 22	169	
23 24	170	Material Testing
25	171	Physical models:
26 27	172	To investigate how the internal shape of vertebrae influences the mechanics of the vertebral joints, we
28 29	173	made simplified and scaled up physical models of the centra of a representative subset of the species.
30	174	We scaled the models up to roughly 20 times their natural size to work with them more easily and did
32	175	not include the spines or zygapophyses. To minimize parameters tested, all models had the same
33 34	176	external diameter and centrum length as well as a simplified cylindrical external structure (Figure 4B).
35	177	The parameters we chose to vary in the models were related to the shape of the internal hourglass-like
30 37	178	structure, including centrum angle, diameter of the centrum cone, and canal diameter (Figure 1A). We
38 39	179	chose three different centrum angles (70, 80, and 90 degrees), three different centrum cone diameters
40 41	180	(15, 20, and 25 mm), and two different canal diameters (2 and 4 mm) to match the variation exhibited
42	181	by specimens during preliminary data collection.
43 44	182	
45 46	183	We created models of single vertebral joints (centra – intervertebral space – centra; Figure 4) from two
47	184	different sets of hard and soft materials to ensure that any variation we saw was due to shape, and not
48 49	185	the material properties of the construction material. The first model centra were 3D printed on a
50 51	186	powder printer (ZCorp 310; 3D Systems, MA) to simulate the bony vertebrae, with Ecoflex 00-10
52	187	(Smooth-On Inc., Macungie, PA, USA) cast in the gaps between the vertebrae (yellow region; Figure 4B)
53 54	188	to simulate the intervertebral material (Figure 4C). For these models, we made three replicates of each
55 56 57 58	189	different combination (e.g., 70° centrum angle, 15mm centrum diameter, and 2mm canal diameter) to
59 60		http://mc.manuscriptcentral.com/icbiol

account for variation in casting. The second set of models were printed fully assembled with a multi-material printer (Figure 4D) (Connex Objet500; Stratasys Ltd., Eden Prairie, MN, USA). We used two materials: VeroClear for the centra and Tango+ for the intervertebral material, both of which are proprietary materials made by Stratasys specifically for their Connex printers. For the printed models, we used one replicate, as variation in printed models is 0.06% of model length with this system (Stratasys, 2018). 

We measured the bending and torsional stiffness of the single joint models using an Instron material testing system (Instron, Norwood, MA, USA). Custom rigs held the models in place with pins (Figure 4E) while each model was rotated or bent to a range of motion similar to or greater than what fish vertebrae would experience during steady swimming. We took the force displacement curves from the Instron and extracted the peak force for each trial using Matlab (Matlab R2018b, The MathWorks, Inc., MA, USA). 

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#### Real vertebral joints:

We measured the maximum bending force before failure of a vertebral joint for a subset of the species in this study, including Damalichthys vacca, Isopsetta isolepsis, Dasycottus setiger, Apodichthys flavidus, and Xiphister mucosus. We chose these species because they are large, thus vertebrae were easier to dissect out and mount in the material testing system. Three individuals of each species were euthanized with a lethal dose (0.5g/L) of MS222 according to University of Washington IACUC protocol 4238-03. Once euthanized, we dissected out sections of the vertebral column containing four centra with the first centra being the one closest to the start of the anal fin. We chose to use four rather than two centra so that we could consistently grip the small joints in our material testing system, though bending and failure occurred at only one joint (the joint between vertebrae 2 and 3 in this four-vertebrae prep). Once dissected out, we mounted vertebral sections in our material testing system with the anterior end being held stationary, and the other attached to the moving load cell. The load cell bent the samples laterally until the joint broke, and max load was recorded.

#### **Statistical Analysis**

For the physical models, we performed a multiple regression analysis in R using the Anova() functions to determine how the parameters we varied affected torsional or bending stiffness (Fox and Weisberg,

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1 2		
3 4 5	221	2019; R Core Team, 2020). We tested each parameter individually, as well as the interaction between
	222	that parameter and the angle to which we twisted or bent the models.
6 7	223	
, 8 9	224	We used a linear mixed effect model (Ime4 package, Imer()) ((Bates et al., 2015)) to ask how vertebral
$\begin{array}{c}1\\2\\3\\4\\5\\6\\7\\8\\9\\10\\11\\12\\13\\14\\15\\16\\17\\18\\9\\21\\22\\23\\24\\25\\26\\27\\28\\9\\30\\31\\32\\33\\45\\36\\7\\38\\9\\40\\142\\43\end{array}$	225	morphology predicted swimming kinematics (bending amplitude and wobble) and mechanics (torsional
	226	modulus and torsional stiffness). Our predictors were the measurements we took of the vertebrae down
	227	the length of the body (Figure 1E) with each row corresponding to a single vertebra. All measurements
15	228	were scaled (scale()) to take into account the different orders of magnitude between linear
16 17	229	measurements (centra length, centra diameter, and foramen diameter, in BLs) and angular
18 19	230	measurements (anterior and posterior centra angle, in degrees). We included individual and species as
20	231	random effects on the model intercept.
21	232	
22 23 24 25 26 27 28 29 30	233	To more easily discuss the relationship between individual morphometric variables and position we
	234	created linear models using the Im() command in R to describe. In comparing linear, quadratic, and cubic
	235	models and for all morphometric variables, the best fit was quadratic (Figure 5). We can then use
	236	coefficients from quadratic fits for each morphometric variable on each fish (intercept, slope, and
30 31	237	quadratic coefficients of the equation $y = p_2 x^2 + p_1 x + I$ ) as descriptors of change along the body.
31 32 33 34 35 36	238	These values will be represented with the tags "I", " $p_1$ ", and " $p_2$ " where I represents the intercept of the
	239	line, $p_1$ represents the slope, and $p_2$ represents the curvature (i.e., the higher the $p_2$ value the more
	240	extreme the curve of the quadratic fit).
37	241	
38 39 40 41 42 43 44 45 46 47 48 49	242	To ask how well morphology could predict categorical variables, we used linear discriminant analysis
	243	(LDA) (Kassambara, 2017; Martins, 2014). LDA is a method which uses the predictor variables
	244	(coefficients of the vertebral measurements down the length of the body in our case) to predict the
	245	class (body shape, swimming mode, or habitat) of an observation (species). We used the <i>lda()</i> function
	246	from the MASS package (Venables and Ripley, 2002) in R to run each of our models, and quantified
	247	accuracy as the percentage of correct predictions from applying the model to test data (a subset of 15%
	248	of the total dataset). We created three models to test three different categorical variables relating to
50 51	249	our data, including body shape, swimming mode, and habitat. For each of these models, our dependent
52 53	250	variable was the category and the independent variables were the coefficients of our quadratic fit lines
54	251	describing morphological and mechanical measurements along the length of the body. Each row of the
55 56 57 58	252	matrix corresponded to one individual, with columns representing measurements at each point along

1 2				
3 4 5 6 7 8 9	253	the length of the body. We also used LDA to predict feeding habitat on a larger group of fishes (Froese		
	254	and Pauly, 2019). This larger group of fishes was primarily used for the tree visualization in the		
	255	discussion, as that dataset only uses one individual from each species.		
10	256	Results		
1 2 3 4 5 6 7 8 9 10 1 12 13 14 15 16 7 18 9 20 1 22 3 24 25 26 27 28 9 30 1 32 33 4 3 5 6 37 8 9 40 1 42 3 44 5 46	257	Morphology: Overall, we found that vertebral morphology varies along the length of the body.		
13 14	258	Generally, vertebral measurements, with the exception of centrum length, increased from the cranium		
15 16 17	259	to around mid-body and then decreased from mid-body to caudal peduncle (Figure 5). For most species		
	260	and variables, the quadratic coefficient of the curve ( $p_2$ ) was negative, meaning the measurements were		
18 19	261	at their maximum roughly 50% of the way down the length of the body. For some species and variables,		
20 21	262	the slope ( $p_1$ ) was positive, or nearly zero.		
22	263			
23 24 25 26 27	264	Kinematics and mechanics: We found that some aspects of vertebral column morphology influenced		
	265	swimming mechanics and kinematics (Table 3). For wobble, tail beat amplitude, torsional stiffness (GJ),		
	266	and torsional modulus (G), at least some morphological variables were significant predictors (i.e., p <		
28 29	267	0.05). Species means for kinematics and kinematics had high standard deviations due to variance along		
30 31	268	the body (Table S1). Our analysis including position accounted for this variation.		
32	269			
<ul> <li>33</li> <li>34</li> <li>35</li> <li>36</li> <li>37</li> <li>38</li> <li>39</li> </ul>	270	Material Testing: For real fish joints, we found that as the ratio of soft material to bone (S:H) increased,		
	271	the force to break the joint decreased (p < 0.001) (Figure 6). For our models, we found that, as centrum		
	272	diameter increased, both bending stiffness and torsional stiffness significantly decreased (Figure 7; Table		
	273	S2). The effects of centrum angle were nonlinear. For bending, we observed peak stiffness in models		
40 41	274	with a centrum angle of 80 degrees. There was a significant effect of bending angle on the Tango+		
42 43 44 45 46 47 48 49 50 51	275	models (p = 0.0043), but not for the Ecoflex models (p = $0.3632$ ). For twisting, there was a trough in		
	276	stiffness with a centrum angle of 70 degrees (Figure 7). Similar to bending, twisting angle also had a		
	277	significant effect on Tango+ models (p = 0.0005), but not Ecoflex models (p=0.6623). Canal diameter did		
	278	not have a significant effect on bending or torsional stiffness. The results from bending or twisting to 15		
	279	degrees showed that the trends held for all angles of motion tested (Figure 7). We chose to show the		
	280	results of only the Tango+ models here, as the trends were more consistent (see Table S1). The variation		
52 53	281	in Ecoflex models is likely due to manufacturing imperfections, as those were hand cast rather than		
54	282	printed as one unit.		
55 56	283			

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LDA results: Our three LDA models showed significant separation between groups with different
swimming modes, habitats, and body shapes (Table 4; Figure 8). The greatest predictive power was
achieved when taking into account the centrum measurements taken along the length of the body
(Figure 1) and the calculated ratio of soft to hard material (Figure 6). Loadings for each of the predictors
can be found in Table S3.

# 290 Discussion

This study has revealed that we can use vertebral morphology to predict the biology of sixteen species of fishes. The degree of accuracy of our predictions varies, but it is often quite high (between 75-100%). Through our preliminary dataset of a larger variety of species, we have also shown that this method can be used to predict biology across the fish tree as well (Figure 9) with functional components increasing the predictive power from 70% accuracy to 79% accuracy. Using functional testing of real vertebral joints and physical models, we showed that internal morphology of vertebrae can affect the mechanical properties of the intervertebral joints (Figures 4 and 5), which may explain why these morphological parameters significantly predict swimming kinematics.

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# 300 Fish vertebral morphology varies among species

Many aspects of vertebral morphology differ among species with one of the most variable traits being number of vertebrae, which can be as many as 260 in elongate species (Mehta et al., 2010) and as few as 28 in more fusiform shaped fishes (Yokogawa, 2013) or even 16 in specialist species like ocean sunfish. In some species of fish, a greater number of vertebrae correlates with a greater curvature coefficient during C-starts (Brainerd and Patek, 2016). Though we did not measure vertebral count directly, centrum body length (CBL) (Figures 1E and 2A) can be used as a proxy since we measured it relative to the fish's body length. Our study species also differ in internal centrum morphology such as cone angle (alpha), cone diameter (D), and canal diameter (d). These species-specific differences in morphology correlate in a consistent way with body shape, habitat, and swimming mode (Figure 8; Table S3). For example, by looking at the coefficients of our LDs, we can see that the slope  $(p_1)$  of centrum body length and anterior diameter along the body both contribute heavily to the separation between behavioral and body shape groups, though for habitat and shape, diameter  $p_1$  is a greater contributor and, for swimming mode, diameter intercept (/) is a greater contributor. This could mean that there is a functional interaction between centrum length and anterior diameter that contributes to the overall shape and behavior of the fish. Generally, habitat and swimming mode have more highly 

316 contributing coefficients in common, such as the intercept of posterior cone angle and the intercept of317 the ratio of soft to hard material along the body.

These intervertebral morphological parameters have rarely been quantified, and our study represents the first time they have been measured and compared among multiple species. In particular, we have known about the notochordal foramen, which is the hole through the center of the vertebrae, from fossil species (Divay and Murray, 2013; Newbrey et al., 2013), but it has rarely been mentioned in extant species. Nowroozi et al. (2012) observed the foramen in the striped bass using histology. They noted that it may be possible for intervertebral fluid to flow through the canal, though it would take a great deal of pressure as the maximum diameter was 0.16 mm. In our study species, we observed the canal to be as wide as 1.1 mm, and we found canal diameter to be a significant predictor of both bending amplitude as well as wobble (Table 3). Perhaps the damping effect of the fluid flowing through differently sized foramen contributes to the steady swimming frequency fish choose as they bend and twist through the water. Future studies may employ fluid modeling to determine if and how fluid flow through the canal affects vertebral mechanics. 

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## 332 Vertebral morphology may be a proxy for other parameters

Using material testing, we showed that the internal vertebral morphology influences the mechanics of individual intervertebral joints in our printed models (Figure 7) and several connected joints in real fish vertebrae (Figure 6). Specifically, our physical models show that vertebrae with a smaller cone diameter are stiffer in both bending and torsion and real vertebrae show that more soft material in a joint means a lower force to break. When comparing morphology to swimming kinematics and mechanics of whole fishes, we found several parameters are significant predictors (Table 3). For example, centrum body length is a significant predictor of bending amplitude, wobble (body twisting), and torsional modulus and there are some interesting interactions to be explored involving anterior and posterior centra diameter and angle (Table 3).

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Our study shows that the change in vertebral morphology down the length of the body is a significant predictor of a fish's preferred swimming mode. The most systematic study of the mechanics of intervertebral joints to date found variation in vertebral morphology down the length of the body in the striped bass (Morone saxatilis), similar to the variation we describe here (Nowroozi and Brainerd, 2014; Nowroozi et al., 2012). This change in vertebral morphology, bending mechanics, and swimming 

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348 kinematics down the length of the body has been observed in other species as well (Hebrank et al., 349 1989; Long, 1991). Nowroozi observed that *M. saxatilis* do not laterally bend enough during swimming 350 to reach the maximum possible bending angle of individual intervertebral joints (Nowroozi and Brainerd, 351 2014; Nowroozi et al., 2012), though we show that a joint may not need to bend to max angle to have a 352 significant effect on behavior. This could indicate a more complicated relationship between the internal 353 morphology of the vertebral column, overall mechanics, and swimming.

355 Though internal measurements of the vertebral centra have generally been ignored when thinking 356 about the mechanics of the system, we have shown they are as significant as centrum length in 357 influencing mechanics (Table 3). Specifically, we have shown that including "functional" variables like 358 ratio of soft to hard material in our LDA models increases predictive power (e.g., the ability of the model 359 to predict categories like habitat). These functional variables may also link to other performance 360 metrics. For example, in robotic models, it has been shown that changing this ratio of hard to soft 361 material decreases bending stiffness which leads to a decrease in both speed and acceleration during 362 swimming (Hirokawa et al., 2011).

364 It is important to note that, while we have shown the relationship between intervertebral morphology 365 and swimming kinematics to be strong, we recognize that no anatomical structures work in isolation. 366 Our model allows us to accurately predict our biological classifications, but the morphology we 367 measured may be proxies for other anatomical parameters that have a more significant effect. We did 368 not include any measurement of neural and hemal spines, zygapophyses, or rib morphology, which can 369 all affect how the vertebral column interacts with connective tissue (Hebrank et al., 1989). We also 370 excluded parameters representing overall body shape, which can correlate with muscle mass and, thus, 371 the stiffness down the length of the body during swimming (Wardle et al., 1995). Our focus on internal 372 vertebral morphology provides insight on the impact of a centrum's stiff and flexible materials on 373 swimming kinematics. However, this internal structure likely works in conjunction with the vertebral 374 spines and connective tissues to influence the mechanics and kinematics of the vertebral column during 375 locomotion.

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Another interesting consideration is the material running through the notochordal foramen. As we 378 mentioned earlier, in some species, the foramen is large enough that the notochord is still present as a 379 continuous rod. In these species, the bony centra are quite thin and sometimes poorly mineralized. In

this case, one might predict that the material properties of the notochord, instead of the bony joints,
dominate the system. Our measurements of the internal centra diameter could also be a proxy for the
shape of the notochord. It may be of use for future work to create physical models representing a
continuous notochord to test whether bending and torsional stiffness change as the thickness of bony
centra is varied.

# 13 14 386 Vertebral morphology is a reliable metric for predicting kinematics in fishes

Though the vertebral column is a small part of fish anatomy in terms of volume, we have shown that it provides reliable parameters for predicting the way various fish species live, likely due to its effect on the mechanics and function of the body. The vertebral column is the main structural component of a fish's body, providing attachment points for muscle and skin. Though studies have shown that connective tissue and muscle play a role in mechanics, our results demonstrate that the internal morphology of the vertebral column also significantly contributes to fish swimming mechanics. A few other studies on vertebral morphology also suggest that the difference in shape and number of vertebrae may affect the behavior of an animal. For example, when reef native eels are behaviorally constrained by their environment, they show a decrease in morphological variation (Mehta et al., 2010). In our data, a decrease in slope  $(p_1)$  or the quadratic coefficient  $(p_2)$  indicates a decrease in variation along the body. Perhaps these data could be used to predict behavioral adaptability in different behavioral and body shape groups. 

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We have shown that internal vertebral morphology can be used to predict the swimming kinematics of fishes from different habitats which have different body shapes and swim using different modes. Models such as ours can also be used to make predictions about fish species for which we cannot collect kinematic data (Figure 8). For example, it is very difficult to measure swimming kinematics for deep sea fishes and impossible to do so for extinct species. By comparing the vertebral morphology of such species with related and more easily accessible organisms, one can predict how other animals may move or may have moved in their natural environments. Finally, with the recent spike in freely available CT scans of fishes and other vertebrates, morphological data is becoming accessible for a huge variety of species (Cross, 2017; Watkins-Colwell et al., 2018). We have shown that it is possible and very useful to describe how subtle variations in morphology play a significant role in locomotion. 

#### 54 <sup>410</sup> 55 417

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Vertebral morphology measured in this study. (A) 3D rendering of one of our scans (Apodichthys flavidus) with the ten evenly spaced points we took measurements from marked along the length of the body. (B) Single lateral slice through a selection of vertebrae generated with a micro-CT scanner. (C) 3D rendering of a pair of vertebrae. (D) Transverse diagram of a single vertebra showing the notochordal foramen in the center with anatomy marked (neural spines – neur., zygopophyses – zyg., hemal spines – hem.). (E). Lateral diagram showing the points we used to calculate centrum body length (CBL), centrum diameter (D), centrum cone angle (θ), and centrum canal diameter (d).

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Diagram of swimming track. The track was made in a modified sea table. Fish swam around the track until they were swimming at a consistent speed, then they were recorded with a GoPro in the filming area (white background).



Distance variables measured. (A) Dorsal View: Silhouettes of the body of Apodichthys flavidus swimming over several frames. Amplitude is the maximum distance the tail moves away from the axis of direction of motion (red line). (B) Transverse View: Wobble (W) is a ratio describing the magnitude of the torsional wave. In the transverse view, wobble is shown for the corresponding silhouettes in the dorsal view.



Diagram of physical models and torsional rig. We designed the models to mimic the morphological variation found in vertebrae, varying centrum angle ( $\theta$ ), centrum diameter (D), and canal diameter (d) (A, B). For the first set of models, we 3D printed centra on a powder printer (C). Then, we aligned matching centra and cast them in PVC with Ecoflex to simulate intervertebral material. For the second set of models, we 3D printed them fully assembled with a multi-material 3D printer (D). Both sets of models were tested using a custom rig (E) connected to an Instron material testing system.





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Load to failure increases significantly (p < 0.001) as the amount of soft material in the vertebral centra decreases. S:H ratio is the ratio of the soft material in the centra to the bony material by area.



Three of the six morphological parameters measured shown for four of sixteen species. In the majority of our study species, centrum body length increases anteriorly to posteriorly along the length of the fish until about 50% BL and then decreases posteriorly. A few species, including, Au. flavidus here, show an inverse trend (A). In contrast, most morphological variables, including foramen diameter and posterior cone angle (B,C), are relatively constant from head to tail. In most species, there is a peak at the midpoint of the body where the measurements reach their maximum or minimum values, though the prominence of that peak varies from species to species





Vertebral morphology can be used to predict attributes of fish biology. Percentages in axis labels are the percentage of between group variation described by that LD. Our LDA results show that the morphological variation in the vertebral column centra can be used to predict fish habitat (A), Body shape (B), and swimming mode (C). Ellipses are drawn at a 95% confidence level using a multivariate t-distribution. No ellipse is drawn for "Sandy Intertidal" in B, as there are only two individuals in that category.



Sample of predictive power of vertebral functional morphology based LDA models over a wider range of actinopterygian species. The top row of dots represents known habitat classifications and the bottom row represents predicted classifications. Correct predictions are highlighted in grey. Tree data was adapted from the fish tree of life database (Rabosky, Chang, Title, al. et, Alfaro, 2018)



Increases in centrum diameter result in decreases in both bending and torsional stiffness. Results from mechanical testing of Stratasys' printed models bent (A) and twisted (B) 15 degrees.

Title: Foretelling the flex - vertebral shape predicts behavior and ecology of fishes

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Keywords: centra morphology, vertebral mechanics, kinematics, swimming, statistical modeling

**Summary statement:** We modeled swimming kinematics and body mechanics of several fish species of varying habitat and body shape based on measurements of internal vertebral morphology.

**Data Availability**: *The data underlying this article are available on GitHub, at* [Link to be determined, a manuscript submission number is needed to create a Dryad Repository].

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# Abstract

One key evolutionary innovation that separates vertebrates from invertebrates is the notochord, a central element that provides the stiffness *needed* for powerful movements. Later, the notochord was further stiffened by the vertebrae, cartilaginous and bony elements, surrounding the notochord. The ancestral notochord is retained in modern vertebrates as intervertebral material, but we know little about its mechanical interactions with surrounding vertebrae. In this study, the internal shape of the vertebrae – where this material is found – was guantified in sixteen species of fishes with various body shapes, swimming modes, and habitats. We used micro-computed tomography to measure the internal shape. We then created and mechanically tested physical models of intervertebral joints. We also mechanically tested actual vertebrae of five species. Material testing shows that internal morphology of the centrum significantly affects bending and torsional stiffness. Finally, we performed swimming trials to gather kinematic data. Combining these data, we created a model that uses internal vertebral morphology to make predictions about swimming kinematics and mechanics. We used linear discriminant analysis (LDA) to assess the relationship between vertebral shape and our categorical traits. The analysis revealed that internal vertebral morphology is sufficient to predict habitat, body shape, and swimming mode in our fishes. This model can also be used to make predictions about swimming in fishes not easily studied in the lab, such as deep sea and extinct species, allowing the development of hypotheses about their natural behavior.



Figure 1: Vertebral morphology measured in this study. (A) Lateral 3D rendering of one of our scanned specimens, Apodichthys flavidus, with the ten evenly spaced points indicating where we took centrum measurements along the length of the body. (B) Single lateral slice through a selection of vertebrae generated with a micro-CT scanner.
(C) Lateral 3D rendering of a pair of vertebrae. (D) Transverse diagram of a single vertebra showing the notochordal foramen in the center and select anatomy marked (neural spines – neur., zygapophyses – zyg., hemal spines – hem.). (E). Lateral diagram showing the points we used to calculate centrum body length (CBL), centrum diameter (D), centrum cone angle (θ), and centrum canal diameter (d).

Table 1. Description of Fish: Short description of fishes used in this study. *L.macrochirus is a freshwater fish, so
the "Habitat" description does not necessarily apply. We chose to identify it as "Subtidal" since it does not deal
with the chaning tides in its natural pond habitat.

Genus	Species	Family	Habitat	Museum Number
Anoplarchus	insignis	Stichaeidae	Intertidal	UW 157117
Ammodites	personatus	Ammodytidae	Nearshore	UW 157013
Apodichthys	flavidus	Pholidae	Nearshore	N/A
Anoplarchus	purpurescens	Stichaeidae	Intertidal	N/A
Aulorhynchus	flavidus	Aulorhynchidae	Nearshore	UW 006155
Cymatogaster	aggregata	Embiotocidae	Nearshore	N/A
Lepomis	macrochirus	Centrarchidae	Subtidal*	UW 119953
Lumpenus	sagitta	Stichaeidae	Subtidal	UW 044716
Myoxocephalus	polyacanthocephalus	Cottidae	Subtidal	UW 027606
Ophidon	elongatus	Hexagrammidae	Subtidal	N/A
Phytichthys	chirus	Stichaidae	Nearshore	N/A
Pholis	laeta	Pholidae	Nearshore	N/A
Pholis	ornata	Pholidae	Nearshore	N/A
Ronqulis	jordani	Bathymasteridae	Subtidal	UW 045926
Xiphister	atropurpureus	Stichaeidae	Intertidal	N/A
Xiphister	mucosus	Stichaeidae	Intertidal	N/A



*Figure 2:* Diagram of swimming track. The track was made in a modified sea table. Fish swam around the track until they were swimming at a consistent speed, then they were recorded with a GoPro in the filming area (white background).



*Figure 3* Distance variables measured. (A) **Dorsal View:** Silhouettes of the body of *Apodichthys flavidus* swimming over several frames. Amplitude is the maximum distance the tail moves away from the axis of direction of motion (red line). (B) **Transverse View:** Wobble (W) is a ratio describing the magnitude of the torsional wave. In the transverse view, wobble is shown for the corresponding silhouettes in the dorsal view.

Table 2.	Description	of categories	assigned 1	to fish sp	ecies

воду зпаре		Habitat	
Deep	High aspect ratio fishes	Pelagic	Spends most of their time in the water column
Fusiform	Low aspect ratio fishes with a torpedo shape	Sandy Subtidal	Lives past the intertidal zone in sandy bottoms
Tadpole	Large head and quickly tapering body	Rocky Subtidal	Lives past the intertidal zone in rocky or shale bottoms
Elongate	Longer than average body and a circular cross section	Nearshore	Follows the tide in and out, often living in eel grass or other plants
Laterally Compressed	Long body and an extremely elliptical cross section	Sandy Intertidal	Lives in the sandy intertidal and often burrows under rocks
Eel-like	Extreme elongation (greater than 65 vertebrae) and an elliptical cross section	Rocky Intertidal	Lives in the rocky intertidal
Swimming Mode			
Pectoral	Swims mainly using their pectoral and	/or pelvic fins	
Caudal	Swims mainly using the posterior 1/3	of their bodies; Cara	angiform swimming
Body-Caudal	Swims using the majority of their bodi	ies; Anguilliform swi	mming





*Figure 4.* Diagram of physical models and torsional rig. We designed the models to mimic the morphological variation found in vertebrae, varying centrum angle ( $\theta$ ), centrum diameter (D), and canal diameter (d) (**A**, **B**). For the first set of models, we 3D printed centra on a powder printer (**C**). Then, we aligned matching centra and cast them in PVC with Ecoflex to simulate intervertebral material. For the second set of models, we 3D printed them fully assembled with a multi-material 3D printer (**D**). Both sets of models were tested using a custom rig (**E**) connected to an Instron material testing system.



*Figure 5.* Three of the six morphological parameters measured shown for four of sixteen species. In the majority of our study species, centrum body length increases anteriorly to posteriorly along the length of the fish until about 50% BL and then decreases posteriorly. A few species, including, *Au. flavidus* here, show an inverse trend (A). In contrast, most morphological variables, including foramen diameter and posterior cone angle (B,C), are relatively constant from head to tail. In most species, there is a peak at the midpoint of the body where the measurements reach their maximum or minimum values, though the prominence of that peak varies from species to species.

Table 3.	I ME results	Numbers listed	are n-values a	and highlighted	values are where	n < (1)(15)
	EIVIE I COUICOI	i tullio ci o lioteu	are p varaes		values are where	p • 0.00.

	Wobble	Amplitude	GJ	G					
Posterior diameter	0.113	<0.001	0.175	0.885					
Anterior diameter	<0.001	0.320	0.517	0.390					
Posterior Angle	0.083	<0.001	0.087	0.621					
Anterior Angle	0.524	<0.001	0.007	0.016					
Post diameter*angle	0.870	0.968	0.860	0.706					
Ant diameter*angle	<0.001	0.660	0.720	0.039					
Centra Length	0.144	<0.001	0.042	0.943					
Foramen Diameter	<0.001	0.004	0.383	0.125					



*Figure 6.* Load to failure increases significantly (p < 0.001) as the amount of soft material in the vertebral centra decreases. S:H ratio is the ratio of the soft material in the centra to the bony material by area.



*Figure 7.* Increases in centrum diameter result in decreases in both bending and torsional stiffness. Results from mechanical testing of Stratasys' printed models bent (A) and twisted (B) 15 degrees.

*Table 4.* LDA results. LD1 and LD2 describe the amount of variance between groups described by that LD. Accuracy is the percentage of correct predictions made using the model and test data (a subset of 15% of the total dataset).

Model	Measu body (0	rements C <sub>M</sub> )	along the	C <sub>M</sub> + ratio of soft to hard material along the body				
Results	LD1	LD2	Accuracy	LD1	LD2	Accuracy		
Mode	0.944	0.056	0.875	0.974	0.026	1.000		
Habitat	0.588	0.189	0.833	0.640	0.140	1.000		
Shape	0.419	0.247	1.000	0.432	0.292	1.000		





*Figure 8.* Vertebral morphology can be used to predict attributes of fish biology. Percentages in axis labels are the percentage of between group variation described by that LD. Our LDA results show that the morphological variation in the vertebral column centra can be used to predict fish habitat (A), Body shape (B), and swimming mode (C). Ellipses are drawn at a 95% confidence level using a multivariate t-distribution. No ellipse is drawn for "Sandy Intertidal" in B, as there are only two individuals in that category.



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*Figure 9:* Sample of predictive power of vertebral functional morphology based LDA models over a wider range of actinopterygian species. The top row of dots represents known habitat classifications and the bottom row represents predicted classifications. Correct predictions are highlighted in grey. Tree data was adapted from the fish tree of life database (Rabosky, Chang, Title, al. et, Alfaro, 2018)

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Title: Foretelling the flex - vertebral shape and swimming kinematics in elongate fishes

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Keywords: centra morphology, vertebral mechanics, kinematics, swimming, statistical modeling

**Summary statement:** We modeled swimming kinematics and body mechanics of several fish species of varying habitat and body shape based on measurements of internal vertebral morphology.

# Abstract

One key evolutionary innovation that separates vertebrates from invertebrates is the notochord, a central element that provides the stiffness needed for powerful movements. Later, the notochord was further stiffened by cartilaginous or bony elements surrounding it, the vertebrae. The ancestral notochord is retained in modern vertebrates as intervertebral material, but we know little about its mechanical interactions with surrounding vertebrae. In this study, we quantified the internal shape of the vertebrae where this material is found in sixteen species of fishes. We used micro-computed tomography to measure shape and then created and mechanically tested physical models of intervertebral joints. These models show that internal morphology of the centrum significantly affects bending and torsional stiffness. We also performed swimming trials to gather kinematic data. Combining these data, we created a model that uses internal vertebral morphology to make predictions about swimming kinematics and mechanics. We used partial least squares (PLS) regression to assess the relationship between vertebral shape and kinematics, body mechanics, and habitat. The analysis revealed that both mechanics and swimming kinematics are influenced by internal vertebral morphology. Our model can also be used to make predictions about swimming in fishes not easily studied in the lab, such as deep sea and extinct species, allowing the development of hypotheses about their natural behavior.

# **Supplemental Material**

### Table S1. Means and standard deviations for kinematics and mechanics data.

Enocioc	<b>Amplitude</b>		Wobble		<u> </u>	<u>3</u>	GJ		
species	<u>mean</u>	<u>sd</u>	<u>mean</u>	<u>sd</u>	<u>mean</u>	<u>sd</u>	<u>mean</u>	<u>sd</u>	
<u>A.insignis</u>	<u>0.0354</u>	<u>0.0163</u>	<u>0.2383</u>	0.0400	1.65E+06	1.82E+06	<u>0.0005</u>	2.11E-05	
<u>A.personatus</u>	<u>0.0394</u>	0.0202	<u>NA</u>	NA	NA	NA	<u>NA</u>	NA	
<u>A.purpureus</u>	<u>0.0431</u>	<u>0.0184</u>	<u>NA</u>	NA	NA	NA	<u>NA</u>	NA	
<u>Ap.flavidus</u>	<u>0.0551</u>	<u>0.0251</u>	<u>0.3349</u>	0.1665	6.13E+05	3.53E+05	0.0005	2.45E-05	
<u>Au.flavidus</u>	NA	<u>NA</u>	<u>NA</u>	NA	NA	NA	<u>NA</u>	NA	
<u>C.aggregata</u>	NA	<u>NA</u>	<u>NA</u>	NA	NA	NA	<u>NA</u>	NA	
<u>L.armatus</u>	NA	NA	<u>NA</u>	NA	NA	NA	<u>NA</u>	NA	
<u>L.macrochirus</u>	0.0451	<u>0.0124</u>	<u>NA</u>	NA	NA	NA	NA	NA	
<u>L.sagitta</u>	0.0379	0.0162	0.3634	0.1050	1.83E+06	2.31E+06	0.0006	2.61E-05	
<u>M.polyacanth</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>	NA	NA	NA	<u>NA</u>	NA	
<u>O.elong</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>	NA	NA	NA	<u>NA</u>	NA	
<u>P.chirus</u>	<u>NA</u>	<u>NA</u>	<u>NA</u>	NA	NA	NA	<u>NA</u>	NA	
<u>P.laeta</u>	<u>0.0550</u>	<u>0.0273</u>	<u>0.3380</u>	0.1186	8.56E+05	4.80E+05	<u>0.0005</u>	3.99E-05	
<u>P.ornata</u>	<u>0.0491</u>	<u>0.0256</u>	<u>NA</u>	NA	NA	NA	<u>NA</u>	NA	
<u>R.jordani</u>	<u>0.0206</u>	<u>0.0158</u>	<u>0.4379</u>	0.0655	1.66E+05	2.35E+05	<u>0.0006</u>	7.55E-05	
<u>X.atropurpureus</u>	<u>NA</u>	NA	<u>NA</u>	NA	NA	NA	<u>NA</u>	NA	
<u>X.mucosus</u>	<u>0.0568</u>	<u>0.0252</u>	<u>0.1944</u>	0.0053	4.90E+05	4.68E+05	<u>0.0007</u>	7.31E-05	

## Table S2. p-values of ANOVAs for material testing of physical models.

	Torsiona	l Stiffness	Bending Stiffness		
	Ecoflex	Tango+	Ecoflex	Tango+	
Angle of Motion	<0.0001	<0.0001	<0.0001	<0.0001	
D	0.0035	<0.0001	0.0013	<0.0001	
theta	0.3632	0.0043	0.6623	0.0005	
d	0.5112	0.3001	0.1605	0.8271	
Angle of Motion * D	0.4432	0.2319	0.0166	<0.0001	
Angle of Motion * theta	0.0701	0.9799	0.5472	0.0232	
Angle of Motion *d	0.6479	0.8787	0.3081	0.8778	

and the bound of the LD models manifest about the values mean a greater contribution to the LD axis.														
	Measurements along the body ( $C_M$ )							C <sub>M</sub> + S:H ratio along body						
	Hab	oitat	Mo	ode	de Shape			Habitat		Mode		Shape		
	LD1	LD2	LD1	LD2	LD1	LD2		LD1	LD2	LD1	LD2	LD1	LD2	
CBL.I	-0.033	0.781	1.303	0.511	-0.166	1.723		2.201	0.383	3.870	-0.099	-1.332	-0.963	
CBL.p1	1.064	-0.333	1.428	-0.547	1.090	-2.060		5.518	-1.412	12.413	-2.363	3.610	1.308	
CBL.p2	0.910	-0.242	-0.782	-2.040	-1.696	-4.642		3.677	1.392	5.808	-4.054	1.771	7.909	
Dant.l	-1.041	4.030	3.865	0.184	0.007	1.043		0.871	1.536	14.021	-1.338	-1.210	-0.683	
Dant.p1	-1.727	1.065	-0.523	-1.521	-1.248	1.022		-4.621	0.929	-5.119	0.337	-3.165	-0.294	
Dant.p2	-2.233	3.289	2.209	0.321	-0.613	1.553		0.302	0.177	7.806	0.843	-0.746	-3.357	
Dpost.l	2.332	-0.724	0.902	0.614	1.922	-0.870		2.138	-1.408	-2.100	2.659	2.714	-2.011	
Dpost.p1	3.104	-0.242	3.356	1.660	2.029	0.670		0.486	-0.870	-2.938	4.129	-0.642	-3.597	
Dpost.p2	2.123	-0.155	1.162	2.298	1.111	3.552		-0.567	-0.778	-7.112	4.924	-2.642	-7.758	
d.I	3,106	-2.383	-0.517	-1.365	2.827	-0.525		0.554	0.154	-2.863	-1.277	3.440	0.909	

Table S3. Scaling values for the LD models. Higher absolute values mean a greater contribution to the LD axis

60

d.p1

d.p2

alphaAnt.I

alphaAnt.p1

alphaAnt.p2

alphaPost.I

alphaPost.p1

alphaPost.p2

perc.l

perc.p1

perc.p2

0.887

1.500

0.477

-0.511

0.173

-0.957

-0.764

1.507

0.132

-1.153

-2.564

-0.800

-2.682

-0.374

1.411

-1.881

0.236

-0.020

-2.148

-0.217

-1.632

-1.919

0.208

0.004

0.253

0.202

-0.089

0.252

-1.184

-0.039

-0.881

-1.367

0.004

0.803

-0.175

-0.270

-0.590

-1.750

-1.081

0.637

-0.857

-1.532

-0.423

-0.061

-2.017

-2.216

-1.200

-2.401

-0.017

2.643

-0.647

4.705

2.692

-4.730

1.509

2.205

-4.057

0.915

-0.437

0.881

-0.827

-1.394

-1.016

-1.940

0.790

1.716

-1.526

0.450

0.877

0.475

-3.087

2.477

-6.476

7.442

2.980

6.062

2.814

-8.922

0.202

1.050

-11.833

0.998

-1.112

-0.047

-1.585

-3.303

2.055

-2.037

-2.039

2.388

0.294

-1.029

-0.354

1.532

-0.811

3.012

1.550

-1.754

1.258

1.666

-1.592

2.123

0.188

0.697

2.421

0.320

0.944

5.218

2.002

2.187

3.034

-0.930

0.378

1.600