SYMPOSIUM

Functional coupling in the evolution of suction feeding and gill ventilation of sculpins (Perciformes: Cottoidei)

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Synopsis

Suction feeding and gill ventilation in teleosts are functionally coupled, meaning that there is an overlap in the structures involved with both functions. Functional coupling is one type of morphological integration, a term that broadly refers to any covariation, correlation, or coordination among structures. Suction feeding and gill ventilation exhibit other types of morphological integration, including functional coordination (a tendency of structures to work together to perform a function) and evolutionary integration (a tendency of structures to covary in size or shape across evolutionary history). Functional coupling, functional coordination, and evolutionary integration have each been proposed to limit morphological diversification to some extent. Yet teleosts show extraordinary cranial diversity, suggesting that there are mechanisms within some teleost clades that promote morphological diversification, even within the highly integrated suction feeding and gill ventilatory systems. To investigate this, we quantified evolutionary integration among four mechanical units associated with suction feeding and gill ventilation in a diverse clade of benthic, primarily suction-feeding fishes (Cottoidei; sculpins and relatives). We reconstructed cottoid phylogeny using molecular data from 108 species, and obtained 24 linear measurements of four mechanical units (jaws, hyoid, opercular bones, and branchiostegal rays) from micro-CT reconstructions of 44 cottoids and 1 outgroup taxon. We tested for evolutionary correlation and covariation among the four mechanical units using phylogenetically corrected principal component analysis to reduce the dimensionality of measurements for each unit, followed by correlating phylogenetically independent contrasts and computing phylogenetic generalized least squares models from the first principle component axis of each of the four mechanical units. The jaws, opercular bones, and branchiostegal rays show evolutionary integration, but the hyoid is not positively integrated with these units. To examine these results in an ecomorphological context, we used published ecological data in phylogenetic ANOVA models to demonstrate that the jaw is larger in fishes that eat elusive or grasping prey (e.g., prey that can easily escape or cling to the substrate) and that the hyoid is smaller in intertidal and hypoxia-tolerant sculpins. Within Cottoidei, the relatively independent evolution of the hyoid likely has reduced limitations on morphological evolution within the highly morphologically integrated suction feeding and gill ventilatory systems.

Introduction

To understand the evolution of complex functional anatomical systems, we must consider how evolutionary forces act on a system at all levels of functional complexity. Studies of functional evolutionary morphology typically examine the evolution of structures in the context of performance of a specific function. However, although some structures appear to have evolved for a singular purpose, such as the baleen of whales or the suction disc of remoras, most structures perform many functions. This reflects a fundamental reality of whole organisms—most...
ubiquitous behaviors (e.g., eating, breathing, and locomotion) require coordination of many structures of the head and body (Wainwright 2007; Walker 2010). The overlap of structures among multiple functions is known as functional coupling. The few existing studies that address macroevolution of structures that perform multiple functions have centered around the idea that functional coupling limits morphological evolution. For example, rates of cranial morphological evolution were shown to be much lower in cichlids that use mouth brooding as opposed to other types of parental care (Tsuboi et al. 2015). Presumably, this is because the additional function of brooding constrains the morphology of the buccal cavity, which also must be used for eating and breathing (Hoey et al. 2012; Tsuboi et al. 2015). Additionally, a classic example of evolutionary innovation in fishes is the evolution of specialized pharyngeal jaws in cichlids and wrasses (Liem 1973; Kaufman and Liem 1982; Wainwright 2006). Many teleost fishes use their oral jaws for both prey capture and prey processing (e.g., manipulation and mechanical breakdown), but many also have specialized pharyngeal jaws used for prey processing (Wainwright 2006). In cichlids (Cichlidae) and wrasses (Labridae), such functional decoupling of prey capture and processing is considered a key innovation that has contributed to morphological diversity within these clades (Liem 1973; Wainwright 2006). Although these examples are compelling, there are counter-examples in which functional coupling is associated with increased diversity. The most striking example is the elytra of beetles (Coleoptera). Beetles exhibit extraordinary morphological diversity and species richness (Lawrence et al. 2011), and yet their key innovation, modified forewings (elytra), are a prime example of functional coupling, because they are used in both protection and flight (Dudley 2002; Johansson et al. 2012). Such functional coupling does not seem to have limited morphological diversification of beetles. Similarly, teleost fishes are the most diverse clade of vertebrates with more than 33,000 extant species (Eschmeyer and Fong 2019), and they have two major innovations: powerful suction feeding and an efficient double-pump system of gill ventilation. These two systems are highly functionally coupled, with both using many of the same structures, and yet this does not appear to have limited morphological or species diversity within teleosts.

Morphological integration can occur across many levels of biological organization and is central to the discussion of functional anatomical evolution. The term morphological integration refers to any functional coordination or evolutionary covariation among structures (Olson and Miller 1958; Cheverud 1996; Klingenberg 2008). This definition is intentionally broad (Cheverud 1996; Klingenberg 2008), because it encompasses the many areas of study necessary to develop a holistic understanding of integration, including fields as disparate as quantitative genetics, functional morphology, biomechanics, developmental biology, and paleontology. However, such a generalized definition makes it difficult to standardize terminology for the study of morphological evolution across a broad range of disciplines. Here, we use a functional anatomical framework to discuss these terms and concepts, and we demonstrate how they can be holistically applied to a specific example—cranial evolution in teleost fishes. For the study of the evolution of complex anatomical systems, we recognize four levels of morphological integration (Table 1). We use phenotypic integration to refer to any correlation or covariation of size or shape of structures across an individual or population. The term “phenotypic” is meant to reflect the convention that a phenotype is defined at the level of the individual. Functional coordination concerns multiple structures that work together to perform a specific function. Functional coupling, as defined previously, comes into play when a structure is involved with the performance of more than one function. At the interspecies level is evolutionary integration, which refers to correlation and co-variation among structures across taxa and evolutionary history.

### Table 1 Types of morphological integration

<table>
<thead>
<tr>
<th>Term</th>
<th>Definition</th>
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<tbody>
<tr>
<td>Morphological integration</td>
<td>Any coordination, covariation, or correlation among structures (Cheverud 1996; Klingenberg 2008)</td>
</tr>
<tr>
<td>Phenotypic integration</td>
<td>Correlation or covariation in size or shape of structures within an individual or population (Parsons et al. 2011)</td>
</tr>
<tr>
<td>Functional coordination</td>
<td>Multiple structures working together to perform a single function with a high degree of biomechanical coordination (Schwenk and Wagner 2001; Collar et al. 2014)</td>
</tr>
<tr>
<td>Functional coupling</td>
<td>Overlap in structures used to perform two or more separate functions (Wainwright 2007; Tsuboi et al. 2015)</td>
</tr>
<tr>
<td>Evolutionary integration</td>
<td>Correlation or covariation in size or shape of structures across evolutionary history (Olson and Miller 1958; Cheverud 1996; Monteiro and Nogueira 2010; Claverie and Patek 2013)</td>
</tr>
</tbody>
</table>

We define types of morphological integration in an evolutionary and functional context.
All levels of morphological integration have the potential to influence the evolution of musculoskeletal structures through complex interactions. These interactions are primarily discussed through the lens of morphological diversity (disparity), with most types of integration historically considered to constrain the extent of morphological evolution by either limiting the morphospace that a structural system can occupy or by reducing the rate of morphological change (Felice et al. 2018). For example, functional coordination has been thought to constrain morphological evolution, because structures that must move in a coordinated manner may be shaped by similar evolutionary influences for performance of a function (Schwenk and Wagner 2001; Collar et al. 2014; Roberts et al. 2018). As mentioned, functional coupling has been shown to constrain morphological evolution (Hoey et al. 2012; Tsuboi et al. 2015), because sets of structures performing multiple functions are limited by mechanical and physiological constraints of all of the functions performed (Lauder 1981; Schaefer and Lauder 1996; Walker 2010). Phenotypic and evolutionary integration are products of complex interactions among genes, developmental processes, performance (including functional coordination and multifunctionality), and evolutionary forces acting on them (Cheverud 1982; Hulsey et al. 2005; Young and Hallgrímsson 2005; Klingenberg 2008; Goswami and Polly 2010; Parsons et al. 2011, 2018). Therefore, there is not always a clear relationship between evolutionary integration and functions of structures (Goswami 2006; Claverie and Patek 2013; Klingenberg and Marugán-Lobón 2013). However, a high degree of evolutionary integration can indicate that a system is constrained by covariation that limits evolutionary variability of morphology within clades (Collar et al. 2014). A lack of evolutionary covariation among structures or parts of structures is termed evolutionary modularity, and the tendency for structures or modules to evolve independently within a functional system can be considered a source of relaxed constraint on morphological disparity in some systems (Goswami and Polly 2010; Parnell et al. 2012; Claverie and Patek 2013; Goswami et al. 2014).

To unite the concepts of evolutionary integration with functional coordination and coupling, we group structures into mechanical units (=functional modules; Breuker et al. 2006; Klingenberg 2008; Córdoba and Cocucci 2011) based on their mechanical connections and their tendency to move together in a mechanically coordinated manner. Mechanical units may show internal phenotypic or evolutionary integration, but this is not necessary to recognize or define them (Breuker et al. 2006; Klingenberg 2008), which is why we avoid the term module. Here, we define four mechanical units: (1) jaws, (2) hyoid, (3) opercular bones, and (4) branchiostegals. The jaws, hyoid, and opercular bones are essential for two functionally coupled systems of teleosts: suction feeding and gill ventilation (Lauder 1980, 1983). Suction feeding is the ancestral feeding strategy for jawed vertebrates (Wainwright et al. 2015), and its ubiquity among actinopterygians (Lauder 1982) and early-branching sarcopterygians (e.g., coelacanths, Lauder 1980; lungfishes, Bemis and Lauder 1986; Bemis 1987a) makes it critical to studies of vertebrate evolution. Suction feeding involves rapid expansion of the buccal chamber, creating negative pressures that draw water and prey into the mouth. Powerful suction feeding with jaw protrusion is considered to be a key innovation that lead, in part, to morphological diversity and species richness of teleosts (Wainwright et al. 2015; Wainwright and Long 2017). Rapid buccal expansion requires movements of the jaws, hyoid, and opercular bones. The expansive phase of suction feeding begins with an increase in gape (Lauder 1985), accomplished by depression of the lower jaw (dentary and anguloarticular, dn and an in Fig. 1) caused by posterodorsal rotation of the opercular bones (opercle, subopercle, and interopercle, op, so, and io in Fig. 1; Liem 1970; Anker 1974; Lauder 1982). Subsequently, protrusion of the upper jaw (pm and mx in Fig. 1), lateral expansion of the suspensorium, and ventral depression of the hyoid (cha and chp in Fig. 1) expand the volume of the oral cavity and further depress the lower jaw. Dorsal rotation of the neurocranium and retraction of the pectoral girdle by epaxial and hypaxial muscles, respectively, also contribute to buccal expansion (Lauder 1985; Camp and Brainerd 2014, 2015). Pump gill ventilation is the most common form of gill ventilation among teleosts, with other strategies including ram ventilation (swimming with an opened mouth to force water over the gills) and aerial ventilation (Brainerd and Ferry-Graham 2006). Pump ventilation relies on changes in water pressure driven by structures of the buccal and gill chambers. These chambers act as pumps, expanding and compressing cyclically to produce unidirectional water flow over the gills (Hughes and Shelton 1958; Hughes 1960). Movements of the buccal and gill chambers coordinate with one another, but the chambers perform different tasks to move water over the gills. The buccal pump generates negative pressure to draw water into the mouth during inhalation, followed by
positive pressure to force water from the mouth over the gills. The gill chamber pump generates negative pressure to draw water over the gills and into the gill chamber, followed by positive pressure to force water out of the gill opening during exhalation. The buccal pump primarily includes the jaws and hyoid, whereas the gill chamber pump consists of the opercular bones and the fourth mechanical unit, the branchiostegals (br in Fig. 1).

In addition to their functional coupling, suction feeding and gill ventilation in teleosts show evidence of high functional coordination and evolutionary integration among the structures involved (Sanford and Wainwright 2002; Gibb and Ferry-Graham 2005; Collar et al. 2014; Kane and Higham 2015). Given the aforementioned associations of functional coupling, functional coordination, and evolutionary integration with constraint on morphological evolution, it is perhaps surprising that there is so much diversity in cranial morphology among teleosts. However, we predict that many teleost clades have sources of relaxed morphological constraint within these complex systems, which allows morphological diversification to occur. We examined this possibility in Cottoidei (sensu Smith and Busby 2014), which includes sculpins, poachers, snailfishes, lumpers, and sandfishes and about 859 valid species to date (Eschmeyer and Fong 2019) in 13 families (Smith and Busby 2014; van der Laan et al. 2014). Cottoidei are ecologically diverse, ranging from intertidal to deep sea habitats, from sit-and-wait predators to active hunters, and from small to large sizes at maturity (Knope and Scales 2013; Buser et al. 2017). However, almost all cottoids are demersal suction feeders and pump gill ventilators, with variation in prey choice and feeding performance (Norton 1995; Napazakov and Chuchukalo 2005; Thedinga et al. 2006; Glubokov and Orlov 2008). Cottoidei has been the subject of extensive phylogenetic study and revision (Smith and Wheeler 2004; Ramon and Knope 2008; Knope 2013; Smith and Busby 2014; Buser and López 2015). Here, we use available molecular data to reconstruct cottoid phylogeny and test four hypotheses of evolutionary integration among the four mechanical units defined above (Fig. 2). Our first hypothesis (H1; Fig. 2A) is that the three mechanical units associated with rapid expansion of the buccal cavity during suction feeding will show significant evolutionary integration, based on their high functional coordination. Our second hypothesis (H2; Fig. 2B) is that mechanical units of the buccal and gill chambers will show significant within-chamber evolutionary integration but lack between-chamber integration, based on the tight within-chamber functional coordination during ventilatory pumping (Fig. 2B). Our third hypothesis (H3; Fig. 2C) is that the pattern of evolutionary integration among these units will reflect development, with the dermal skeleton of the jaws, opercular bones, and branchiostegals showing significant evolutionary integration, based on their similar developmental origins as dermal bones (except for the articular part of the anguloarticular) relative to the endochondral origin of the hyoid (de Beer 1937; Hall 2015). Our last hypothesis (H4; Fig. 2D) is that all mechanical units will show significant evolutionary integration based on the high degree of functional coupling between suction feeding and gill ventilation (Liem 1970; Lauder 1980, 1983). To relate our morphological measurements to ecology, we use data from studies of sculpin diet (Norton 1995), habitat (Eschmeyer et al. 1983), and hypoxia tolerance (Mandic et al. 2009, 2013) to test associations among ecological variables and mechanical units within a subset of cottoids examined.
Fig. 2 Four hypotheses of evolutionary integration among mechanical units of the skull. (A) H1: Mechanical units associated with suction feeding (jaws, hyoid, and opercular bones) will show significant evolutionary integration, based on functional coordination necessary for rapid buccal expansion, during which the upper jaw protrudes (blue) and the lower jaw (blue) is depressed by dorsal opercular rotation (green) and hyoid retraction (orange). (B) H2: Mechanical units of buccal and gill chambers will show significant within-chamber evolutionary integration but lack between-chamber integration, based on within-chamber functional integration during ventilatory pumping. (C) H3: The jaws, opercular bones, and branchiostegal rays will show significant evolutionary integration, based on their similar developmental origins relative to the hyoid. (D) H4: All mechanical units will show significant evolutionary integration, based on the high degree of functional coupling between suction feeding and gill ventilation.
Materials and methods

Phylogenetic reconstruction

We reconstructed a phylogeny of Cottoidei using mitochondrial and nuclear gene sequence data for 106 species of cottoids and two outgroup species, *Hexagrammos decagrammus* and *H. stellari* (Online Appendix Table SA1). We used a molecular dataset from Knope (2013), which included sequences from mitochondrial gene cytochrome b (cytb) and the first nuclear intron of the S7 ribosomal protein (S7) for 99 cottoid species. We supplemented these data with sequences from GenBank (Benson et al. 2013), including cytb from seven additional species of cottoids and the two outgroup species as well as S7 for the two outgroup species. We also included GenBank sequences from mitochondrial gene cytochrome c oxidase I (COI) for a subset of 72 taxa. Sequences from each gene were aligned individually using MUSCLE (Edgar 2004) on the EMBL–EBI bioinformatics web tool (Li et al. 2015). The complete alignment had 86.7% amplicon and 69.3% base pair coverage. Best-fit nucleotide substitution models were chosen using jModelTest v2.1.7 (Guindon and Gascuel 2003; Darriba et al. 2012) based on AIC values calculated for 24 candidate models.

We constructed trees using MrBayes v3.2.3 (Ronquist et al. 2012) on the CIPRES Science Gateway v3.3. We computed 36 MrBayes runs with seven gene partitions and an MCMC chain length of 20,000,000. Nucleotide substitution models were chosen for our seven partitions based on jModelTest results. We used a general time-reversible model (Tavaré 1986) with invariable sites and rate variation among sites (GTR + I + G) for S7 and cytb (codon positions 1, 2, and 3). For the COI gene, we used a general time-reversible model with equal state frequencies (SYM) for codon position 1, the F81 model (Felsenstein 1981) for codon position 2, and a general time-reversible model with rate variation among sites (GTR + G) for codon position 3. Each of the 36 MrBayes runs were evaluated for quality in Tracer v1.6.0 (Rambaut et al. 2014), and the 20 runs with the highest effective sample size for the mean likelihood score were used to construct 20 maximum clade credibility (MCC) trees in Tree Annotator v1.8.0. We determined which tree topologies best explained our molecular data with a Shimodaira–Hasegawa test using the SH.test function in the phangorn R package (Shimodaira and Hasegawa 1999; Schliep 2011; R Core Team 2016), and the results are listed in Online Appendix Table SA2.

Morphological measurements

We measured cranial skeletal elements from 44 cottoids and one outgroup taxon (*Hexagrammos stellari*) ranging in head length from 6 to 35 mm. We collected and measured 17 species from marine habitats near Friday Harbor Laboratories on San Juan Island, Washington (Cornell University IACUC 2013-0017) and accessioned them into the Cornell University Museum of Vertebrates (Online Appendix Table SB1). Individuals were identified to species using Eschmeyer et al. (1983). We supplemented these with measurements from specimens already in the collection (*Rhamphocottus richardsonii* [CU54050], *Xeneretmus latifrons* [CU71854], and *Cottus cognatus* [CU78131]). We imaged the specimens using micro–computed tomography (micro-CT) at the Cornell University Biotechnology Resource Center’s Imaging Facility (GE eXplore CT-120 or an Xradia Versa XRM-520). Each fish was scanned from the pectoral girdle to the anterior-most point of the head. Voxel size ranged from 14 to 50 μm³, with smaller specimens scanned at lower voxel sizes. DICOM files are available for public download on Morphosource.org. We further supplemented our dataset with 25 cottoid scans downloaded from Morphosource.org (Online Appendix Table SB1). We imported DICOM or JPG files from each scan into OsiriX (version 6.0.2 64-bit; Rosset et al. 2004) or Horos software (Nimble Co LLC, Annapolis, MD, USA) and used the “3D multiplanar reconstruction” tool to view the 3D reconstruction simultaneously in three orthogonal planes. To obtain linear measurements from each structure, we began with the specimen oriented such that we could view the sagittal, transverse, and frontal planes. We then rotated the planes until the entire length of the structure was in view in one plane by overlaying the axes of the other two planes (using the “Axis” tool) in the measurement plane, in which we measured the dimensions of the structure.

We collected linear measurements in this manner from the four previously noted mechanical units of the head: jaws, hyoid, opercular bones, and branchiostegals. Jaw measurements were: length of the maxilla, maximum height of the maxilla, length of the premaxilla, length of the ascending process of the premaxilla, lengths of the dentary from the anterior tip to the posterior tip of the dorsal and ventral processes, length of the anguloarticular from the anterior tip to the jaw joint, height of the anguloarticular at the jaw joint, and distance between the left and right jaw joints. Hyoid measurements were: length and height of the anterior ceratohyal, length...
and height of the posterior ceratohyal (=epihyal of some authors; see Grande and Bemis 1998: 23), and length of the urohyal. Opercular bone measurements were: length of the opercle from the joint with the hyomandibula to the posterior tip of the opercle and length of the opercle from the joint to the posterior margin of the opercle (horizontal from the joint with the hyomandibula), height of the opercle vertically from the joint with the hyomandibula, lengths of the subopercle from the joint with the interopercle to the fork and to the posterior tip of the subopercle, and length of the interopercle from the anterior to posterior tips. We did not measure the preopercle, because it is functionally associated with the suspensorium rather than the operculum. Branchiostegal measurements were: lengths of the first six branchiostegal rays from their articulations with the ceratohyal to their posterior tip. Each species examined has six branchiostegal rays, except for Dasycottus setiger and Artedius harringtoni, which have seven.

For size correction, we measured the length of the neurocranium from the anterior tip of the nasals to the posterior tip of the basioccipital, width of the neurocranium between the left and right joints of the neurocranium and hyomandibula, and maximum vertical height of the neurocranium. Although both the suspensorium and pectoral girdle are involved with suction feeding, we did not include them in our analyses due to substantial confounding factors, including the different developmental origins of the many suspensorial bones (Hulsey et al. 2005) and the role of the pectoral girdle in locomotion.

**Testing hypotheses of evolutionary integration**

To test for evolutionary correlation and covariation among the four mechanical units, we conducted the following analyses: (1) phylogenetic size correction of all measurements, (2) a phylogenetically corrected principal component analysis (pPCA) for each mechanical unit to reduce the dimensionality of the data, (3) correlations among phylogenetically independent contrasts computed from the first principle component axes (PC1) of each mechanical unit pPCA, and (4) tests of covariation among PC1 axes of each unit using phylogenetic generalized least squares (PGLS) models. We considered mechanical units to show evolutionary integration if both correlation of PIC contrasts and covariation of PC1 axes based on PGLS models were significant at $P = 0.05$. Analyses were conducted in R (R Core Team 2016) using the MCC tree from the best MrBayes run, as well as 500 trees randomly sampled from the posterior distribution of trees from this same MrBayes run (with 40% burn-in removed), with each tree successively used for all four analyses (Full code available on DataDryad, DOI:10.5061/dryad.gv3r544). We ultrametricized each tree using the *chronopl* function in the *ape* package (Paradis et al. 2004) and pruned each tree to include only the 45 taxa from our study using *drop.tip* in *ape* (Paradis et al. 2004). For size correction, we used the *phyl.resid* function in the *phytools* package (Revell 2009, 2012) to obtain residuals from PGLS regressions of each jaw, hyoid, opercular, and branchiostegal measurement against the geometric mean of head length, head width, and head height as a metric for head size. These residuals were used as phylogenetically size corrected data in our subsequent character analyses, as recommended by Revell (2009). For each of the four mechanical units, we reduced the dimensionality of measurements by performing a pPCA using the *phyl.pca* function in the *phytools* package (Revell 2009, 2012). We then extracted the PC1 axis. We used each PC1 to compute phylogenetically independent contrasts for each mechanical unit using the *pic* function in *ape* (Felsenstein 1985; Paradis et al. 2004). Correlations through the origin (Garland et al. 1992) were performed among contrasts of the four mechanical units, the jaws, hyoid, opercular bones, and branchiostegals, using the *cor.origin* function in the *PHYLOGR* package (Diaz-Uriarte and Garland 2014). Figure 3 summarizes these results, including biplots of PC1 PIC contrasts (lower left), and the distribution of correlation coefficients from each of the contrast pairs across 500 trees (upper right). To confirm that significant PIC correlations were associated with significant covariation between units, we tested for covariation using the *GLS* function in the *nlme* package (Pinheiro et al. 2018), with a Brownian Motion correlation structure computed from the phylogeny using the *corBrownian* function in *ape* (Paradis et al. 2004). The results of the PIC correlations and PGLS models are provided in Table 2 for both the MCC tree and for the posterior distribution of 500 trees from the best MrBayes run. We provide continuous character maps of the PC1 axis for each mechanical unit, made using the *contMap* function in *phytools* (Revell 2012), in Supplementary Fig. SA2.

To test whether we had sampled enough trees from the posterior distribution of trees from MrBayes, we randomly sampled 1–500 PGLS covariation coefficients from the above analysis of 500 trees and plotted mean coefficient against number of trees included. To test whether enough taxa were sampled, we resampled our pool of taxa 35 times to include a range of 10–45 species. We computed PGLS models
Table 2 Summary of statistical analyses

<table>
<thead>
<tr>
<th>Mechanical Units</th>
<th>r</th>
<th>P</th>
<th>Covariance</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Jaws vs. hyoid</td>
<td>−0.1846</td>
<td>0.2141</td>
<td>−0.2931</td>
<td>0.2246</td>
</tr>
<tr>
<td>Jaws vs. oper</td>
<td>0.3330*</td>
<td>0.0228</td>
<td>0.4080*</td>
<td>0.0254</td>
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<tr>
<td>Jaws vs. branch</td>
<td>0.5641*</td>
<td>3.633E−5</td>
<td>0.4568*</td>
<td>5.443E−5</td>
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<td>Hyoid vs. oper</td>
<td>−0.2020</td>
<td>0.1733</td>
<td>−0.1559</td>
<td>0.1833</td>
</tr>
<tr>
<td>Hyoid vs. branch</td>
<td>−0.3983*</td>
<td>0.0056</td>
<td>−0.2032*</td>
<td>0.0067</td>
</tr>
<tr>
<td>Oper vs. branch</td>
<td>0.5142*</td>
<td>2.179E−4</td>
<td>0.3399*</td>
<td>3.023E−4</td>
</tr>
<tr>
<td>Jaws vs. hyoid</td>
<td>−0.2985 (±0.0089)</td>
<td>0.0975 (±0.0125)</td>
<td>−0.5377 (±0.0187)</td>
<td>0.1026 (±0.0127)</td>
</tr>
<tr>
<td>Jaws vs. oper</td>
<td>0.2964* (±0.0014)</td>
<td>0.0444 (±0.0013)</td>
<td>0.3819* (±0.0022)</td>
<td>0.0494 (±0.0013)</td>
</tr>
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<td>Jaws vs. branch</td>
<td>0.5637* (±0.0015)</td>
<td>4.841E−5 (±7.054E−6)</td>
<td>0.4854* (±0.0020)</td>
<td>7.037E−5 (±9.222E−6)</td>
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<tr>
<td>Hyoid vs. oper</td>
<td>−0.2212 (±0.0031)</td>
<td>0.1459 (±0.0062)</td>
<td>−0.1640 (±0.0031)</td>
<td>0.1548 (±0.0063)</td>
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<tr>
<td>Hyoid vs. branch</td>
<td>−0.4186* (±0.0022)</td>
<td>4.013E−3 (±2.559E−4)</td>
<td>−0.2058* (±0.0012)</td>
<td>4.891E−3 (±2.934E−4)</td>
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<tr>
<td>Oper vs. branch</td>
<td>0.5114* (±0.0021)</td>
<td>3.904E−4 (±5.072E−4)</td>
<td>0.3420* (±0.0018)</td>
<td>5.091E−4 (±1.347E−4)</td>
</tr>
</tbody>
</table>

Correlation coefficient for correlations of independent contrasts (r) and coefficients from PGLS models (covariance), with respective P-values. The top set of values is based on the MCC tree from the best MrBayes run. The bottom set of values are means based on a posterior distribution of 500 trees (after burnin) from the same MrBayes run, with 95% confidence intervals included in parentheses. Significant correlations are indicated with an asterisks.

Ecological analysis

To relate size of mechanical units to ecology, we used previously collected ecological data to conduct phylogenetically corrected ANOVAs on subsets of our taxa. We grouped 13 species of sculpins according to diet data from Norton (1995; Online Appendix Table SB2). Norton reported species diets by quantifying percent wet mass of each diet item within stomach contents of at least 16 individuals per species. We assigned each species a diet type, based on the prevalence of elusive, grasping, or easy prey in their diet. According to Norton, elusive prey escape predation through locomotion, grasping prey cling to substrate, and easy prey can be captured with minimal resistance. We also assigned 21 species a habitat type (intertidal or subtidal), according to Eschmeyer et al. (1983). We also determined which species showed high levels of hypoxia tolerance using data for 11 species from Mandic et al. (2009, 2013). We tested for associations between diet, habitat, and hypoxia tolerance and each of the four mechanical units with phylogenetically corrected ANOVA models using the GLS function in the nlme package (Pinheiro et al. 2018), with a Brownian motion correlation structure computed from the phylogeny using corBrownian in ape (Paradis et al. 2004), along with the anova function (R Core Team 2016).

Results

Phylogenetic reconstruction

Based on the Shimodaira–Hasegawa test of 20 MCC trees, we identified one tree as the “best tree,” with the topology that best explained our molecular data (Online Appendix Fig. SA1), and we used this tree and its posterior distribution of trees for all phylogenetic comparative methods. Four trees showed a significant difference from the best tree in the ability of their topologies to explain the molecular data. All trees recovered the Cottoidei clade (sensu Smith and Busby 2014) with high support, and 13 trees recovered Cottoidea (sensu Yabe 1985; Smith and Busby 2014; Online Appendix Table SA2). Our best tree (Online Appendix Fig. SA1) generally agrees with previous studies, particularly because we recovered a large monophyletic group of marine sculpins (Knope 2013 [unnamed clade]; Smith and Busby 2014 [Psychrolutidae]; Buser and López 2015 [unnamed clade]). However, our placement of the monotypic genera Rhamphocottus, Jordania, and Leptocottus within Cottoidea conflicts with previous studies (Knope 2013; Smith and Busby 2014). These problematic taxa are typically recovered at various positions among the non-psychrolutid cottoids with low node support (Knope 2013; Smith and Busby 2014), which is also the case in our analysis.

Character analyses

In the pPCA of mechanical units across the distribution of 500 trees, the PC1 axis explained 75.29–79.05% of the variance (77.78% median; 79.33%...
with MCC tree) for jaws and was mainly associated with variance in the dentary, anguloarticular length, maxilla, and premaxilla length. In pPCAs of hyoid measurements, PC1 explained 46.16–64.99% of the variance (50.70% median; 58.78% with MCC tree) and was mainly associated with variance in length of the anterior ceratohyal and the urohyal. In pPCAs of opercular bone measurements, PC1 explained 50.75–64.78% of the variance (55.92% median; 59.69% with MCC tree) and was mainly associated with variance in length of the subopercle from the joint with the interopercle to the posterior tip. In

Fig. 3 Results of phylogenetically independent contrasts. The results of PIC correlations for each pair of mechanical units are shown in the lower left and upper right, with images of the structures from a micro-CT reconstruction in the center. We computed phylogenetically independent contrasts of PC1 scores from each mechanical unit pPCA to conduct correlations among our four mechanical units. Using the best MCC tree, we created bivariate plots of PC1 contrasts (lower left). Trendlines are shown for significant relationships. The distributions of correlation coefficients (r) across a posterior distribution of 500 trees are summarized by histograms (upper right). Among Cottoidei, we found significant positive evolutionary correlation among the jaws, opercular bones ("oper"), and branchiostegal rays ("branch"), with the hyoid showing no positive integration with any of the other mechanical units, and the hyoid was negatively correlated with the branchiostegal rays. Asterisks denote statistically significant correlations.
mainly elusive and grasping prey had larger jaws than species that ate easy prey \( (P = 0.0131) \), subtidal sculpins had larger hyoids than intertidal sculpins \( (P = 0.0275) \), and sculpins that were hypoxia tolerant had smaller hyoids than those with lower hypoxia tolerance \( (P = 0.0074) \). The opercular and branchiostegal mechanical units were not significantly associated with any of the ecological traits examined.

**Discussion**

Within Cottoidei, we found that the jaws, opercular bones, and branchiostegal rays show evolutionary integration (Table 1), as evidenced by significant evolutionary correlations among the PC1 axes of the pPCAs (Table 2 and Fig. 3). This is congruent with our third hypothesis based on the similar developmental origins of cranial dermal bone (dermal components of the jaws, opercular bones, and branchiostegal rays) relative to endochondral bone (hyoid). The coordinated evolution of mechanical units with similar development origins indicates that developmental pleiotropy within teleost skulls may produce developmental modularity on an evolutionary scale. Within Cottoidei, the independent evolution of the hyoid relative to other mechanical units of the skull may reduce limitations on morphological evolution of the suction feeding and gill ventilatory systems. Evolution of morphology within Cottoidei might otherwise be constrained by the high degree of functional coupling and functional coordination seen in teleost skulls (Hughes 1960; Lauder 1983, 1985; Collar et al. 2014). The relatively independent evolution of the hyoid in Cottoidei likely contributes to ecomorphological diversity of this clade.

**Morphological integration and suction feeding**

Suction feeding is widespread across teleost fishes, and the associated structures show substantial morphological diversity, despite the constraints of functional coordination, evolutionary integration, and coupling with other functions such as gill ventilation and occasionally parental care (Hoey et al. 2012; Tsuboi et al. 2015) or locomotion (Fish 1987; Pietsch and Grobecker 1987). Therefore, there must have existed sources of relaxed constraint on morphological evolution within the suction feeding system throughout the history of teleosts. The independent evolution of the hyoid relative to other mechanical units used for suction feeding is likely one of these mechanisms within Cottoidei. We speculate that independent evolution of the hyoid probably occurs within other clades of teleosts because the hyoid plays several key roles in suction feeding,

Fig. 4 Resampling analyses. To determine whether enough trees and taxa were sampled for consistent PGLS covariation coefficients, we randomly resampled PGLS covariation coefficients computed from 500 trees and plotted mean coefficient against number of taxa included (top left), and we randomly resampled our pool of taxa to include 10–45 species and plotted mean coefficient against number of taxa included (bottom right). PGLS covariation coefficients became increasingly consistent with increasing number of taxa and trees, demonstrating that our sampling was sufficient. Each plot represents a regression between the mechanical unit labeled vertically and the unit labeled horizontally.

pPCAs of the branchiostegal measurements, PC1 explained 76.87–84.71% of the variance (82.44% median; 82.15% with MCC tree) and was loaded highly for all six branchiostegal ray lengths. Our plots of number of trees and number of taxa sampled against mean PGLS covariation coefficients showed a strong convergence toward the values reported in Table 2 (Fig. 4), and therefore, we concluded that we had sampled enough taxa and trees for this study.

PC1s of the pPCAs of the jaws, opercular bones, and branchiostegals were significantly positively evolutionarily correlated with one another \( (P < 0.05; \text{Table 2}) \). However, PC1 of the hyoid was not significantly positively correlated with any of the other mechanical units in any of the trees (Table 2 and Fig. 3). PC1 of the hyoid was significantly negatively correlated with PC1 of the branchiostegals \( (P < 0.005; \text{Table 2}) \). For phylogenetic ANOVAs with ecological traits (Fig. 5), sculpins that ate mainly elusive and grasping prey had larger jaws...
and its relative size in different species likely is linked to its relative importance in fishes with different diets. For example, suction performance is thought to be maximized when peak suction corresponds with peak gape, which is made possible by hyoid depression that lags behind jaw depression (Gibb and Ferry-Graham 2005; Bishop et al. 2008; Wainwright et al. 2015), and therefore species with a larger hyoid can generate larger negative pressures during the peak-gape phase of suction. The hyoid is also critical to intraoral prey transport after the expansive phase of suction feeding, and often undergoes large excursions during prey processing and swallowing (Gillis and Lauder 1995).

In our analysis of cottoid diet data from Norton (1995), phylogenetic ANOVAs of 3 different prey types ("elusive prey," "grasping prey," and "easy prey") among 13 sculpin species (Fig. 5) showed that fishes that mostly eat grasping or elusive prey had larger jaws than those of fishes that eat easy prey (easy prey have "no major morphological defenses against capture once discovered;" Norton 1995: 64). This finding is consistent with Norton's conclusion that jaw size is largest in suction feeders that primarily eat elusive prey (Norton 1995). The increase in jaw size is likely due to a greater reliance on jaw protrusion to create a larger buccal volume change (Norton 1991, 1995; Carroll et al. 2004; Higham et al. 2007), increased hydrodynamic force (Holzman et al. 2008b; Wainwright and Longo 2017), and increased “jaw ram,” in which jaw protrusion serves the additional purpose of bringing the opening of the mouth closer to the prey (Longo et al. 2016).

There is interest in evolutionary integration and functional coordination of body movements with suction feeding (Holzman et al. 2008a; Kane and Higham 2015; Longo et al. 2016), but studies that empirically test for integration among cranial units used in suction feeding are rare (Collar et al. 2014). Although we rejected our hypothesis of tight evolutionary integration among three major mechanical units of suction feeding (jaws, hyoid, and opercular bones: Fig. 2A), we demonstrated evolutionary integration of two mechanical units used in suction feeding (jaws and opercular bones). Opercular bones depress the lower jaw during the expansive phase of suction feeding. The opercle is extraordinarily diverse among teleosts, showing internal evolutionary and developmental modularity (Kimmel et al. 2017), and its correlated evolution with the jaws of sculpins may influence the diversity of suction feeding kinematics within this group. Together with the branchiostegal rays, the opercular bones also form the lateral wall of the gill chamber and contribute its expansion during suction feeding (Liem 1978). However, gill chamber expansion, facilitated by abduction of the opercular bones and lateral and ventral expansion of the branchiostegal apparatus, has almost no known influence on suction feeding pressures (Lauder 1980, 1983) and does not impact suction feeding performance other than potentially reducing backflow of water into the buccal chamber (Liem 1978). Therefore, although evolutionary correlation between jaw and opercular size may result from selection for suction feeding performance due to the linkage of opercular elements to lower jaw depression, it is unlikely that the correlation between
jaw and branchioostegal size plays a role in the evolution of suction feeding.

Morphological integration and gill ventilation

The branchioostegal rays and the membrane they support are a major source of diversity in gill ventilatory morphology (Baglioni 1907; Hughes 1960; McAllister 1968; Farina et al. 2015; Farina and Bemis 2016), and therefore it is surprising that they show such strong evolutionary correlation with other bones of the skull, especially the jaws. Based on gill ventilatory function, we hypothesized that the mechanical units of the buccal and gill chambers would show a high degree of evolutionary integration within chambers and little integration between chambers (H2; Fig. 2B). However, we did not find the expected integration between the jaws and hyoid as the primary mechanical units of the buccal chamber. The evolutionary integration between the mechanical units of the gill chamber (opercular bones and branchioostegals) may be due to the shared role of these structures in expanding and contracting the gill chamber during ventilation (Hughes 1960).

Major innovations within Cottoidei are related to their convergent invasion of the intertidal zone in multiple lineages (Ramon and Knop 2008; Buser et al. 2017). Oxygen availability is not as stable in the intertidal as in the subtidal, and therefore many intertidal sculpins have evolved hypoxia tolerance, air breathing, and the ability to walk between tide pools (Mandic et al. 2009, 2013; Bressman et al. 2018). In the subset of species analyzed, the hyoid was larger in subtidal sculpins and in species that had a low tolerance for hypoxia (Fig. 5). In benthic fishes such as cottoids, hyoid depression and elevation are the primary components of buccal expansion and compression (Hughes 1960). A large hyoid may provide a larger ventilatory stroke volume by producing a higher amplitude of buccal pumping. This would be efficient for subtidal fishes that can breathe slowly due to relatively stable oxygen availability, but intertidal fishes tend to rely on rapid rates of ventilation (Martin 1991) instead of high stroke volume. The negative evolutionary correlation between the hyoid and branchioostegal rays implies that sculpins with smaller hyoids tend to have larger branchioostegal rays, although branchioostegal size was not significantly predicted by hypoxia tolerance or tidal zone in our ecological analyses, suggesting that this negative evolutionary correlation is not indicative of an ecological trade-off.

Morphological integration and functional coupling

The overlap in mechanical units associated with suction feeding and gill ventilation in teleosts offers opportunities to examine co-evolving structures in coupled functions. Some suction-feeding vertebrates, including lungfishes and some elasmobranchs, decreased coupling between feeding and ventilatory structures through anatomical specializations of the suction feeding system (Bemis and Lauder 1986; Bemis 1987a, 1987b; Motta et al. 2002), yet teleost fishes have maintained an extremely close anatomical coupling between the two functions. Suction feeding is not simply an amplified gill ventilatory cycle, as evidenced by differences in the kinematics and pressure profiles observed during ventilation and suction feeding, including substantial differences in the relative magnitudes of pressures between the buccal and gill chambers (Hughes 1960; Liem 1970; Lauder 1980, 1983). Also, suction feeding often involves protrusion of the upper jaws and movement of some post-cranial elements, but these are less common in gill ventilation. Therefore, there are likely separate selective pressures shaping morphology and kinematics of these coupled functional systems. The relatively independent evolution of the hyoid may provide some of the evolutionary flexibility needed to respond to these selective pressures for both functions within Cottoidei and possibly within other teleost clades. This is supported by the association of the jaws and hyoid with different ecological characteristics (diet and hypoxia tolerance, respectively) related to both feeding and respiration. Differing developmental mechanisms acting upon endochondral and dermal bone also likely play a large role in the relatively independent evolution of the hyoid. Although it is difficult to directly link development to the evolution of function, it is likely that developmental modularity contributes to the evolutionary flexibility of this system (Albertson et al. 2003; Hulsey et al. 2005), allowing different units to respond independently to selection for different types of functional performance. We argue that investigations of the evolution of functional anatomical systems, particularly in vertebrates with complex musculoskeletal configurations, should simultaneously consider the constraining influence of functional coordination, evolutionary integration, and functional coupling whenever possible.

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**Supplementary data**

Supplementary data are available at ICB online.

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