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## JOURNAL OF THE ROYAL SOCIETY INTERFACE

### Repeated evolution of drag reduction at the air-water interface in diving kingfishers

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in diving kingfishers	

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#### **ABSTRACT**

Piscivorous birds have a unique suite of adaptations to forage under the water. One method aerial birds use to catch fish is the plunge dive, wherein birds dive from a height to overcome drag and buoyancy in the water. The kingfishers are a well-known clade that contains both terrestrially foraging and plunge-diving species, allowing us to test for morphological and performance differences between foraging guilds in an evolutionary context. Diving species have narrower bills in the dorso-ventral and sagittal plane and longer bills (size corrected data, n=71 species, p<0.01 for all), Although these differences are confounded by phylogeny (phylogenetically corrected ANOVA for dorso-ventral p=0.26 and length p=0.14), beak width in the sagittal plane remains statistically different (p<0.001). We examined the effects of beak morphology on plunge performance by physically simulating dives with 3D printed models of beaks coupled with an accelerometer, and through computational fluid dynamics (CFD). From physically simulated dives of bill models, diving species

have lower peak decelerations, and thus, enter the water more quickly, than terrestrial and mixed-foraging species (ANOVA p=0.002), and this result remains unaffected by phylogeny (phylogenetically corrected ANOVA p=0.05). CFD analyses confirm these trends in three representative species, and indicate that the morphology between the beak and head is a key site for reducing drag in aquatic species.

Keywords: plunge diving, avian hydrodynamics, beak, bow wave, Alcedinidae

TROLLING.

#### INTRODUCTION

Plunge diving has evolved in multiple flying species to facilitate transitioning between the air and water – two mediums of vastly different densities. Birds including gannets, terns, and boobies have mastered diving from air into water to access fish meters below the surface. Morphological adaptations likely compliment this foraging strategy in order to both improve dive efficiency and avoid damage on water entry. The shape of the kingfisher's bill has served as inspiration as a dragreducing structure for the Japanese Shinkansen Bullet train (1, 2). However, these functions have yet to be directly tested.

The conversion of gravitational potential energy to kinetic energy during the dive provides momentum for the bird to overcome body drag and buoyancy in order to dive deeper (3). Birds are particularly buoyant due to the layer of air trapped

between the body and the feathers, typically used for insulation (4), as well as body fat and the avian system of airsacs (5). In the diving species the Lesser Scaup (presumably already adapted to reduce drag), over 80% of work during a dive is to overcome the significant costs of body buoyancy (6).

Minimizing the energetic costs of drag have led to streamlined bauplans in swimming and flying animals (7-11). Bird beaks appear well-adapted to avoid both aerodynamic and hydrodynamic drag. Most beaks are relatively cone-shaped, with a small initial surface area relative to the direction of oncoming flow – thus reducing immediate profile drag. The gradual increase in cross-sectional area allows flow to remain laminar as it travels toward the wide middle-section of the animal.

While much work has focused on how shape influences drag across flying and swimming animals, less work exists examining morphological function at the air-water interface. Diving involves the animal rapidly transitioning between two fluids of different physical properties – from air, a relatively low density and viscosity fluid, to water, a higher density and viscosity fluid. Due to the high speed of entry, diving comes at the cost of an initial impact at the water's surface. Gannets reportedly dive from a height of 30 meters in the air– a fall resulting in a speed of 22 m/s when impacting the water (3). While these impact speeds could seriously damage a human entering feet-first (12), an avian injury due to water entry has not been reported. The neck musculature coupled with streamlined beak and skull help the gannet avoid injury by reducing impact forces (12). In fact, large decelerations due to water impact during diving may not occur in birds. Accelerometers mounted to free-living Cape Gannets sampling at 16 to 32 Hz detected no or minimal

deceleration due to impact during foraging dives. (3). Drag reduction due to morphology may help reduce immediate impact forces. The hydrodynamic shape of the avian bill may also reduce turbulence during the initial dive, which may help avoid visual or vibrational detection by the prey (13).

Recent work examining water piercing by geometric cones suggests that beak morphology may be selected on to reduce impact force, and thus, drag on entry (14). The lower the opening angle of the cone (or the tip angle), the lower impact forces and more smooth the transition between air and water (14). The opening angle of a cone (a) can be calculated as a= 2\*arcsin(r/s), where r is the radius of the base, and s is the length of the side from base to tip (also called 'slant height'). Thus, to decrease the angle of a cone, either the radius of the base (r) must decrease, or the length (s) must increase. If diving species of kingfisher are morphologically adapted to minimize drag, we would expect them to have longer bills with a narrower base relative to terrestrial species.

Kingfishers (Alcedinidae) are an ideal clade in which to explore morphological adaptations for diving. They comprise 114 species that encompass terrestrial, aquatic, and mixed (both terrestrial and aquatic) foraging strategies (15), allowing us to test function and morphology in an evolutionary context. Here, we examine beak morphology to elucidate patterns of streamlining in diving species. We test hydrodynamic properties of bird beak shape by simulating dives with scaled 3D printed plastic models of the birds. Printed models allow us for the first time to isolate shape from size. Lastly, we use Computational Fluid Dynamics to explore flow around the beak and head.

**Morphometrics** 

**METHODS** 

3D digital models of bird beaks were generously provided by the Mark My Bird project as 3D scans of specimens housed in the Natural History Museum at Tring and the Manchester Museum (See Appendix 1 for museum details and specimen IDs). Please see information in the appendix of (16) for details pertaining to scanning methodologies. The scans are available for download by request from markmybird.org. The scan of a Forest Kingfisher (*Todiramphus macleayii*) was obtained from a specimen in the Bangor University Brambell Natural History Museum. This scan was produced by Rowan Howe at the Pontio Innovation Centre with an Artec Spider (Artec Group, Luxembourg), with a standard resolution of 0.05 mm and mesh resolution of 0.1 mm. Mesh generation was accomplished with Artec Studio 9 (Artec Group, Luxembourg).

Morphometrics were measured directly from specimen scans, representing 71 species (Appendix 1; Figure 1). Beak width was measured as the linear distance between either end of the lower and upper mandible external hinge. Beak height was measured from the linear distance between the most dorsal and most ventral points where the beak meets the feathered portion of the head along the sagittal plane. Beak length was measured from the tip of the bill to the end of the mandible hinge (Figure 2).

The mass of the individual museum specimen prior to preservation is unknown. Body size from the literature was used as an estimation of representative body size for each specimen. Masses for each species were found in the CRC Handbook of Avian Masses (17). When available, average mass for a species was used. If male and female mass was reported separately, the two were averaged for subsequent analyses. Any species for which mass data was not available was excluded from this study.

# 3D model manufacturing

Thirty-one species were subsampled for functional testing, representing a variety of foraging strategies and body sizes across the kingfisher phylogeny (Figure 3). One beak model was printed for each of 31 species (Appendix 1).

Prior to 3D printing, scans were post-processed in Ultimaker Cura 3 to remove holes. To account for differences in drag due to body size, all scans were geometrically scaled to 9 cm from the tip to the posterior of the beak (Figure 2). Scans were finished by a transverse cut across the head of the animal at the end of the beak. This cut allowed us to incorporate the entire morphology of the beak alongside the joint where the beak meets the head.

3D prints were produced on an Ultimaker 3+ (Ultimaker, Cambridge, MA, USA) with a 0.4 mm nozzle size. Prints were produced with a layer height of 0.1 mm, infill density of 20%, and four gradual infill steps. Beaks were printed with

biodegradable plastic poly lactic acid (PLA) filament (RS Components Ltd, Northants, UK).

#### Physically Simulated dives

Beak models were attached to a closed 50 ml falcon conical centrifuge tube. The models were mounted to a 9 cm long wooden or plastic dowel to increase the distance between the beak and falcon tube 'dive body,' thus minimizing any effects of the tube shape and buoyancy during the initial entry phase of the dive. The tube contained an Axivity AX-3 triaxial accelerometer (Axivity Ltd, Newcastle, UK) sampling at 1600 Hz with a maximum value of  $\pm 16$  G. The accelerometer was oriented to the beak model with the negative x axis aligned with gravity, and the positive z axis oriented dorsally. The falcon tube was weighted to equalize the weight of every model and support beam to that of the largest model. The mass of each of the total structure including models totaled 71.1 grams.

A fishing line track mounted perpendicular to the water surface was used to maintain model orientation during the dive. The tube was fitted with plastic drinking straws on either side lengthwise and threaded on to fishing line. The dive tank was a 60 cm tall flower vase with an opening of 25 cm. (Figure 4A). A simulated dive was performed by dropping the model (beak pointed down) in to the tank along the fishing wire track from 75 cm above the surface of the water. To confirm acceleration was not impacted by the trackway, and the accelerometer gave

a reliable reading, the accelerometer gravity axis during the fall was double integrated, and resulted in the correct 75 cm.

The fishing line maintained orientation of the models vertically, although slight differences in entry angle along the dorso-ventral plane were apparent, leading to slight variation in deceleration values. To account for this, 10 drops were performed for each model. All acceleration analyses were done only on the vertical (orthogonal to the water surface) component. All accelerometer outputs were analyzed in a custom written Matlab script. For the purposes of this study, only the initial deceleration phase was analyzed – the time between when the beak has entered the water and has become fully submerged. At the time of submergence, the model experiences a maximum deceleration (Figure 4B).

Any outliers above 3 standard deviations were removed from subsequent analyses. Resulting analyses for inter-species comparisons used the average maximum deceleration for each model.

#### Statistical analyses

Each species was assigned to a foraging group based on behavior and diet descriptions in the Handbook of Birds of the World Alive (18). Three foraging groups were used: terrestrial, aquatic, or both. If a species could not be readily assigned to one of these groups, it was not included in the study.

For analyses of morphological characters, in order to meet assumptions of normality and homoscedasticity, all measurements were log-10 transformed prior to analyses. Morphometric characters were tested for size-dependence with a linear

regression between character and reported body size (all p<0.01). All three were adjusted for size by regressing log-10 adjusted values against log-10 adjusted body mass and calculating the residuals. The residuals were used for subsequent comparisons. An analysis of variance (ANOVA) tested for differences between foraging groups.

In order to account for phylogenetic effects, a phylogenetic tree was constructed based on Anderson et al. (19) (Figure 1). Binomial names according to the Jetz et al. (20) phylogeny were used. *Alcedo euryzona* was placed as sister taxa to it's conspecific A. peninsulae (18). To explore the relationship between foraging guild and performance, a sub-sampled phylogeny of the 31 tested species was constructed from the first phylogeny (Figure 3). These 31 species were selected to encompass a range of foraging guilds and body sizes across the phylogeny. In both phylogenies, branch lengths were set using arbitrary lengths using a Grafen transformation (21). We tested for differences in morphology and hydrodynamic function between foraging groups with a phylogenetically corrected ANOVA according to Garland et al.'s method (22). The phylogenetic ANOVA was implemented via the phytools package in R (23). Both morphometric and performance phylogenetic ANOVAs were calculated with 10,000 simulations. To elucidate differences between groups, a pairwise posthoc test was performed using a Holm correction.

Computational Fluid Dynamics

To simulate flow over the beak and head, a virtual flume was simulated using Autodesk CFD 2019. Digital models of *Ceyx*, *Dacelo*, *and Ceryle*, were used as representative taxa; two attributed to terrestrial and one to aquatic feeding strategies. To create suitable, watertight meshes for CFD, the scan data was manipulated via a combination of Autodesk Maya 2019 and Autodesk Meshmixer. First, models were aligned to world axes (anterior aligned to +x, dorsal to +y, and right-lateral aligned to +z), and scaled such that beak length equaled 9 cm in all specimens, so as to match the physical models used above and to remove size effects. Models were then cropped posterior to the beak, but anterior to the eye sockets, before holes were filled, and the models made solid. A smoothing pass was applied to remove erroneous spikes in the laser scan data, or to remove small sharp topography caused by errant feathers when the specimens were scanned. To avoid flow artefacts from a flat surface at the back of the head, the filled surface was extruded, and then deformed into a cone-shape consistent with the edges of the head (Figure 5a). This avoided any abrupt or complex transitions from laser scan to reconstructed posterior. The now watertight meshes were then downsampled using InstantMeshes (https://github.com/wjakob/instant-meshes (24)) to ~20,000 triangles (figure 5A & B).

The downsampled meshes were imported into Autodesk CFD 2019, where simulations were constructed in a similar manner to (25). A fluid volume was generated around the mesh, so as to create a virtual flume with walls sufficiently far from the mesh to avoid edge effects. Using standard materials in Autodesk CFD, properties of fresh water (density = 998.2 kgm<sup>-3</sup>, Viscosity = 0.001003 Pa-s) were

applied to the fluid volume. Kingfisher models were given properties of ABS Polycarbonate, though as the models were stationary and mass-less, the material properties of the kingfisher beaks had little to no impact on results. The anterior end of the flume was set as an input flow of  $5 \, \text{ms}^{-1}$ , approximately the same as for the physical simulations. A zero-pressure boundary condition was applied to the opposing, posterior end allowing flow through the flume at a uniform  $5 \, \text{ms}^{-1}$ . All other fluid boundaries were set to a slip/symmetry condition. Gravity was not included in the simulation. Meshing of the domain was carried out automatically prior to the simulation process (Figure 5C). A steady-state simulation was run until convergence, utilizing the SST k-Omega turbulence model. Results were calculated and visualized using Paraview 5.6, and are presented vertically for consistency with physical simulations above. We also calculated coefficient of drag:  $C_d = 2 \, F / \rho \, v^2 \, a^2$ , where  $\rho = 998.78$ ,  $v^2$  is velocity, and  $a^2$  is cross-sectional area at the widest point of the model.

#### **RESULTS**

#### Morphology

Diving, terrestrial, and both foraging groups differ significantly in beak morphology, but these results are confounded by phylogeny for beak length and depth.

After adjusting for body size, beak length differs between foraging groups (Fig. 6A; ANOVA  $F_{2.68}$ =13.67, p<0.001). Aquatic foraging kingfishers have longer

beaks than terrestrial kinglishers (Tukey HSD p<0.001), but aquatic foragers do not
differ from birds that forage in both (p=0.99). Terrestrial kingfishers have shorter
beaks than birds found in the 'both' category (p=0.003). These relationships are
confounded by phylogeny – foraging guilds are not statistically significantly
different in beak length (Phylogenetic ANOVA F=13.67, p=0.14).

Size-corrected beak depth differs significantly between foraging groups (Fig. 6B; ANOVA  $F_{2,68}$ =8.98, p<0.001). Aquatic foraging birds have shallower bills than terrestrial (p<0.001) and both (p=0.003) foraging groups, but terrestrial birds do not differ from birds that forage both ways (p=0.64). These significances are not resilient to phylogeny (Phylogenetic ANOVA F=8.79, p=0.255).

Lastly, size-corrected beak width differs between foraging groups (Fig. 6C; ANOVA  $F_{2,68}$ =48.97, p<0.001). Aquatic beaks are narrower than terrestrial (p<0.001) and both (p<0.001) groups. Terrestrial beaks do not differ significantly from birds that forage in both methods (p=0.944). After accounting for phylogenetic relatedness, beak width remains significantly different between groups (phylogenetic ANOVA F=48.97; p<0.001). Aquatic beaks remain significantly more narrow than terrestrial (pairwise phylogentically corrected p<0.001) and mixed (p=0.003) foraging groups. Terrestrial species do not differ significantly from birds that forage both aquatic and terrestrially (p=0.79).

*Performance – physical simulations* 

Beaks from aquatic foraging species exhibited lower average peak decelerations during water entry than both terrestrial and aquatic-terrestrial foraging species (Figure 7; ANOVA  $F_{28,2}$ =7.645, p=0.002). Aquatic and terrestrially foraging species dive deceleration were significantly different (Tukey HSD, p=0.002), while aquatic (p=0.92) and terrestrial (p=0.92) were not significantly different from foraging strategies that utilized both aquatic and terrestrial styles.

When phylogeny was accounted for, the difference in performance between foraging guilds remains significant (phylogenetic ANOVA, F=7.64, p=0.047). However, a pairwise posthoc test with a Holm correction (26) found marginal differences between aquatic and terrestrial foraging groups (p=0.084), terrestrial and both foraging groups (p=0.084), and no difference between aquatic and both foraging groups (p=0.78).

#### Performance - CFD

The CFD simulations indicate a higher anterior-posterior drag force in the terrestrially foraging taxa, *Ceyx erithaca* and *Dacelo novaeguineae* than the aquatic forager *Ceryle rudis*. However, while this drag force was particularly high in *Dacelo* (6.86N,  $C_d$  = 0.23), the terrestrial *Ceyx* (2.98 N,  $C_d$  = 0.17) experienced only slightly more drag force than the aquatic *Ceryle* (2.27 N,  $C_d$  = 0.23). The three simulated kingfishers also exhibited differences in dorso-ventral drag force, *Dacelo* and *Ceyx* both experience force in the negative horizontal direction (i.e. force pushing the head ventrally) of 1.54 N and 0.68 N respectively. The aquatic foraging *Ceryle* however, experienced 0.14 N of force in a positive horizontal direction (i.e. a force

acting to lift the head). Lateral forces were generally low, as would be expected, but were not zero due to asymmetries in the scan data.

Visualization of fluid velocity indicates that anterior to the head, at the posterior beak, is where most fluid is pushed forwards, generating pressure (or form) drag. The bow waves are smallest in *Ceryle*, and then *Ceyx*, extending only a limited distance in front of the beak. The *Dacelo* model produces a significant bowwave approximately twice the magnitude of the other models. This is most notable in the extensive areas of water being pushed forwards in front of the tip of the beak (Figure 8).

#### 

#### **DISCUSSION**

Our data shows that diving kingfishers have morphological adaptations associated with aquatic foraging. Further, aquatic foraging species beak shapes produce less hydrodynamic drag than terrestrial species, measured as lower peak deceleration during impact with the water, and as drag force in CFD simulations. Collectively, we find evidence that supports adaptations for improved diving performance in aquatically foraging kingfishers relative to terrestrial and mixed foraging species. While the exact values for deceleration and drag of our models have been normalized to size and are therefore not directly applicable to individual taxa, they do provide valuable relative information regarding potential selection for drag-reducing shape.

Beak width in aquatically foraging species is less than in terrestrially foraging species. Both length and depth also differ between foraging groups, but these patterns were not significant once phylogeny was taken in to consideration. Our study aligns with hydrodynamic expectations based on water piercing studies using geometrically perfect cones (12, 14). Diving species have beaks of lower base width, and tend toward longer beaks with lower base depth. (Figure 6). Additional morphological details not measured in this study likely contribute to dive performance, including the morphology of the head, body and wings of the bird. In Vincent et al.'s (14) recent work, the larger the radius of the cone base (r, corresponding to depth and width on our kingfishers), the higher the initial impact forces, due to increased frontal and surface area (12), which increase both pressure and friction drag respectively. This suggests that not only the shape of the beak, but the shape of the frontal area of the bird (which is generally wider than the beak) likely plays a role in plunge diving. Our CFD analyses demonstrate that it is the rapid increase of frontal area at the beak-head transition that generates the largest drag forces, and this transition is smoothest in the diving species *Cervle rudis* relative to terrestrial species – where a larger volume of water is accelerated in the direction of travel by the beak-head transition (Figure 8).

Our CFD models were similar, but not entirely in agreement with our physical experiments. *Dacelo novaeguineae*'s physical model dive force was 107% that of the CFD model (physical model = 7.4 vs. CFD = 6.9 N), *Ceyx erithaca* was 142% (4.2 vs. 2.3), and *Ceryle rudis* was 159% (3.6 vs. 2.3). Our CFD analysis was performed on models with hydrodynamically smoothed ends, unlike the physical

models mounted to a pole and accelerometer, and were also tested at slightly different velocities (4.5 physical models vs. 5 ms<sup>-1</sup> CFD). Most notably, the CFD was performed in a closed boundary, simulating movement within water, rather than transitioning between low density (air) and high density (water) fluid. The mechanics of such transitions are complex (14), including cavitation and splash, and are thus difficult to simulate. Thus, our CFD is likely not a precise measure of the initial water entry phase, but is useful for comparing general hydrodynamic form between taxa.

Notably, no apparent bow wave, where water is pushed forward in front of the animal (27, 28) appears at the tip of the *Ceyx* or *Ceryle* kingfisher bills in the CFD simulations (Figure 8A, B). However, a noticeable bow wave does appear at the beak-head joint (Figure 8C). The elongated beaks of diving birds, coupled with apparent beak-head streamlined morphologies, may delay the effects of this bow wave long enough to avoid detection by the prey. The larger, highly terrestrial forager, *Dacelo* displayed significantly greater bow waves, both in front of the beak-head joint, and even in front of the beak tip, which is broader and deeper than the other two taxa simulated.

Of interest are the resulting dorso-ventral drag forces in our CFD results produced by each beak, with the terrestrial forms *Ceyx* and *Dacelo* generating forces that push the head ventrally. *Ceryle*, meanwhile, generated only very small dorso-ventral forces, acting in the opposite direction. This may be the result of the more curved beak, in comparison with the straighter beaks of the other two models. The lower forces acting orthogonal to the direction of movement may be necessary for

the bird to travel straight when diving into the water. Reducing these dorso-ventral forces may be more important during diving, in a more viscous fluid, than in flight through air.

Further work examining the hydrodynamics of living birds may illuminate additional patterns. For example, our study examined only kingfisher dives with closed beaks, with particular interest at the air-water boundary. However, the kingfisher must open the bill to catch prey. At that point, the hydrodynamics of the bird are likely to be very different. Computational fluid dynamic modeling of aquatic striking snakes suggests that prey could become dislodged by a bow wave created by the open jaw of the snake (27). However, the shape of the kingfisher bill, particularly in aquatic foragers, is much longer, and would likely open to a lesser angle, than a striking snake, which may reduce any emergent bow-wave. CFD models in aquatic snakes suggest that larger prey sizes can offset the bow wave induced movements of the prey. Behavioral studies have shown that captive Pied kingfishers tend to select the larger available prey items (29), and the Common kingfisher selects prey within a discrete size range of 5 to 6 cm in length (30). This size selection may impede the hydrodynamic effects of displacement from the open bill. Size selection could also be due to prey availability, depth (31), or visual limitations, such as contrast or light refraction (32), during foraging.

Selection may act not only on the beak, but the entire frontal area of diving birds. Unlike the plunge diving gannets and terns, the kingfisher neck is noticeably shorter and the feathers appear to smoothly taper from the head to the body in the dive posture – potentially ensuring an entirely streamlined body. Further work

examining entire body morphology in live animals is necessary to better understand the potential for streamlining across species.

While we adjusted the overall shape of the models in order to test questions pertaining to shape, not size, we can use our deceleration values to estimate if the dive itself is enough to overcome buoyancy with a rough calculation. Buoyancy (N) is calculated as  $F_b = \rho * V_{bird} * g$ , where  $\rho$  is the change between air and water density (998.78 kg m<sup>-3</sup>), V<sub>bird</sub> is the volume of water displaced by the bird (i.e., the volume of the bird, m<sup>3</sup>), and g is gravity (9.81 m s<sup>-2</sup>). As a rough estimate, we can consider a spherical bird with a radius of 6 cm, which would have a buoyancy force of 2.1 N that must be overcome to submerge the bird. Our prints were scaled to the beak length of the largest species in the sample, the diving bird *Megaceryle maxima*, which weighs 325 grams and had a deceleration value of 7.36 m s<sup>-2</sup>. By F = m a, the impact force of the bird would be 2.392 N, - a force larger than the estimated buoyancy of our spherical bird, allowing total submergence. In contrast, the smallest diving species, *Alcedo pusilla*, has a mass of 13.3 grams, and had a deceleration value of 5.95 m s<sup>-2</sup>, resulting in an impact force of 0.43 N – not enough to overcome buovancy for a 6 cm radius bird. Our calculation of buovancy force is very rough, and does not account for the density of the animal or actual volumes. Diving species may be less buoyant than their terrestrial counterparts in part due to differences in body mass and ability to retain air under feathers (5, 33, 34), although this has not been tested in kingfishers. Birds can actively adjust their buoyancy by changing the amount of air stored in the respiratory system during a dive (35). Birds may use leg-

or wing-produced thrust to help counter buoyancy during a dive following initial submergence (36, 37).

Conflicting evolutionary demands are placed on beaks. For example, higher mechanical advantage in relation to more leaf-based diets appears to be a primary driver of beak shape in *Anseriformes* (38). Shape changes associated with increased bite force in the beaks of Darwin's finches also limit the use of the jaw during song production (39). Thus, it is important to keep in mind that the beak shapes tested here are likely also under selection for other behaviors, including bite force, burrow excavation, or territorial defense. Additionally, morphological variables not measured here likely contribute to aquatic diving performance, including beak surface structure (40) and position of the nares.

In conclusion, we showed that diving kingfishers have narrower beaks, and a tendency toward longer and more shallow beaks once phylogeny is accounted for when comparing to terrestrial species. Our physical simulations show that diving species beak shapes experience markedly less deceleration when entering the water, corroborated by CFD models. This repeated evolution of functionally and morphologically more hydrodynamic beaks across the kingfisher phylogeny suggests convergence on morphology to improve foraging success in diving birds. Our work may help further inspire engineering solutions, including robotics working at the air-water interface.

#### **AUTHOR'S CONTRIBUTIONS**

KEC conceived of the study. KEC, ROH, and PLF contributed to the design of the study and drafting the manuscript. KEC and ROH acquired the data for the physical tests and performed statistical analyses, PLF carried out computational fluid dynamics analyses. All authors gave final approval for publication and agree to be held accountable for all aspects of the work.

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#### **DATA ACCESSIBILITY**

Data is available as a supplemental file.

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#### FIGURE LEGENDS

**Figure 1:** The phylogeny of 71 kingfishers (Alcedinidae) used for morphometric analysis in this study, constructed as a sub-sample of Anderson et al. (2018). Coloured circles represent classified foraging group: blue are aquatic foraging (diving) species, grey are mixed (aquatic and terrestrial), and green are terrestrially foraging species. See text for details.

**Figure 2:** Morphometric variables collected for each species on the Common Kingfisher, *Alcedo atthis*. (A) Lateral view, (B) Dorsal view. Beak Length measurements were scaled to 9 cm in all 3D models to standardize for body size. See text for additional details.

**Figure 3:** Phylogeny of 31 species of kingfishers used for performance testing, subsampled from the phylogeny in figure (1) (Anderson et al. 2018). Coloured circles represent classified foraging group: blue are aquatic foraging (diving) species, grey are mixed (aquatic and terrestrial), and green are terrestrially foraging species.

**Figure 4: (A)** Diagram of diving tank set up. Dive tank was 60 cm tall with an opening of 25 cm. The dive body consists of a 50 ml falcon tube containing the accelerometer and additional weights as needed. The accelerometer was mounted with the negative x-axis aligned with gravity, and the positive z axis oriented perpendicular to the bird dorsally. The falcon tube was fitted with plastic drinking straws on either side, and the straws were threaded along fishing line to maintain the dive orientation perpendicular to the water surface. **(B)** Exemplar accelerometer data from three representative species: *Ceryle rudis* (pied kingfisher), *Dacelo novaeguineae* (Laughing Kookaburra), and *Ceyx erithaca* (Black backed kingfisher). Data is smoothed by taking a running average for 3 points, and is truncated before cavitation.

**Figure 5:** Kingfisher beak models and CFD domain. **(A)** Original scan data (above), and cleaned, smoothed, and scaled model (below) of *Ceryle* presented in lateral and posterior-lateral views. Models were cropped at the posterior most portion of the beak, then holes were filled, surfaces extruded, and final model then smoothed. **(B)** Original and cleaned meshes for *Ceyx, Ceryle*, and *Dacelo*, left to right. Grid represents 1 cm squares. **(C)** Meshed CFD domain.

**Figure 6: (A-C)** Residuals of morphological characters regressed against body mass, resulting in size-corrected beak morphometrics for kingfisher species classified as aquatic foragers (blue), mixed foragers (grey), and terrestrial foragers (green). **(D-F)** Uncorrected measurements for morphological characters (in mm). Size-corrected (A) beak length, (B), beak depth, and (C) beak width are all statistically significantly different between foraging guilds (ANOVA  $F_{2,68}$ =48.97, p < 0.001). Once phylogeny is accounted for, only beak width (C) remains significantly different between size-corrected aquatic and terrestrial species (phylogenetically corrected p < 0.001).

**Figure 7:** Average peak deceleration values measured for 3D printed scaled models of kingfisher beaks classified as aquatic foragers (blue), mixed foragers (grey), and terrestrial foragers (green). Aquatic and terrestrially foraging species dive deceleration are significantly different (ANOVA  $F_{28,2}$ =7.645, p=0.002, Tukey HSD, p=0.002). This result is not affected by phylogenetic relatedness (phylogenetic ANOVA, F=7.64, p=0.047).

**Figure 8:** Water velocity in the anterior-posterior direction in front of the head of (A) *Ceryle*, (B) *Ceyx*, and (C) *Dacelo*. Velocity scale is truncated to illustrate areas of high and low velocity. Note the much larger bow wave in front of the highly terrestrial *Dacelo*. Bottom: Static pressure around (D) *Ceryle*, (E) *Ceyx*, and (F) *Dacelo*.

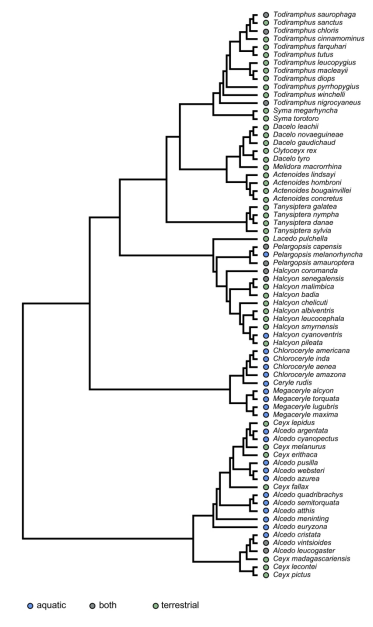
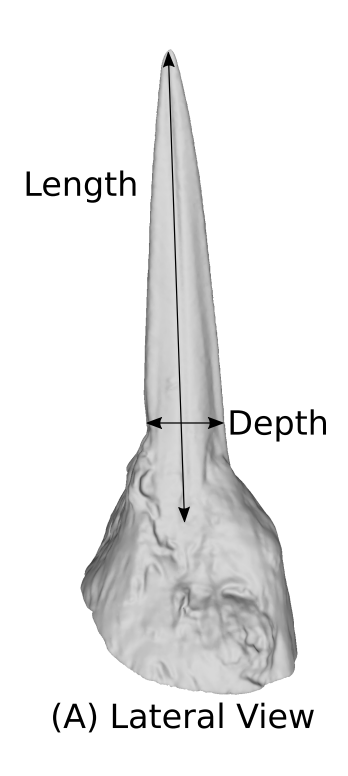
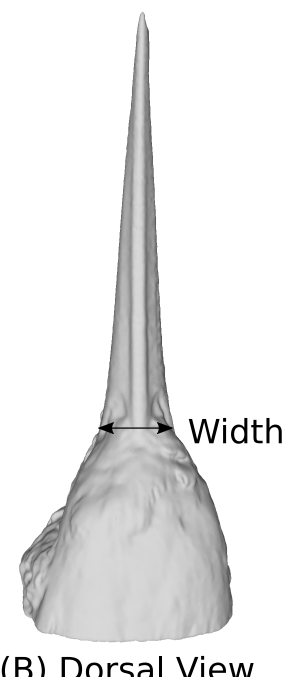


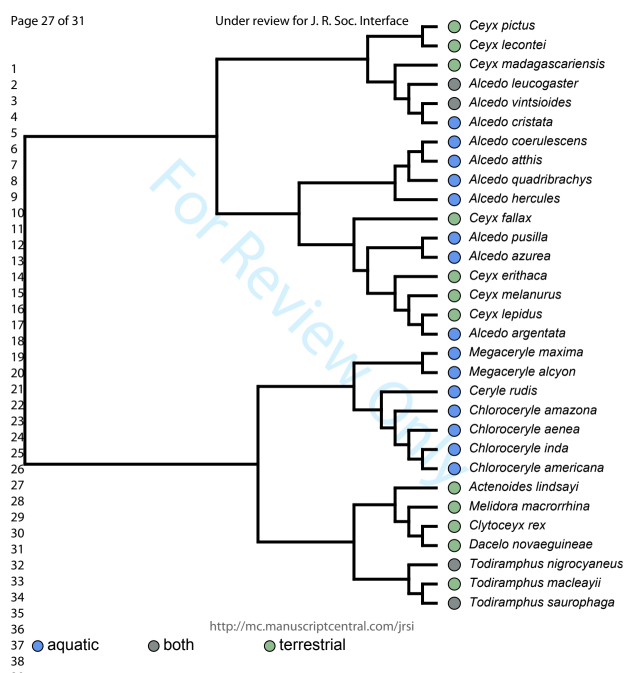
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159x279mm (300 x 300 DPI)





(B) Dorsal View



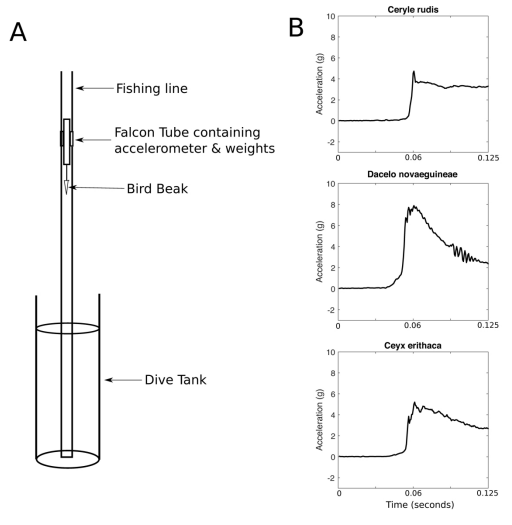


Figure 4: (A) Diagram of diving tank set up. Dive tank was 60 cm tall with an opening of 25 cm. The dive body consists of a 50 ml falcon tube containing the accelerometer and additional weights as needed. The accelerometer was mounted with the negative x-axis aligned with gravity, and the positive z axis oriented perpendicular to the bird dorsally. The falcon tube was fitted with plastic drinking straws on either side, and the straws were threaded along fishing line to maintain the dive orientation perpendicular to the water surface. (B) Exemplar accelerometer data from three representative species: Ceryle rudis (pied kingfisher), Dacelo novaeguineae (Laughing Kookaburra), and Ceyx erithaca (Black backed kingfisher). Data is smoothed by taking a running average for 3 points, and is truncated before cavitation.

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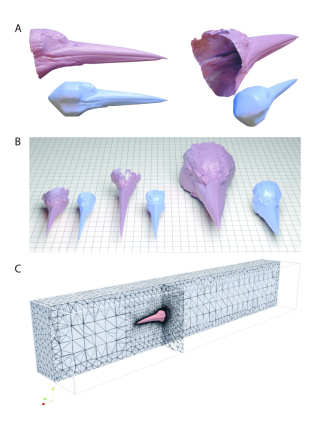


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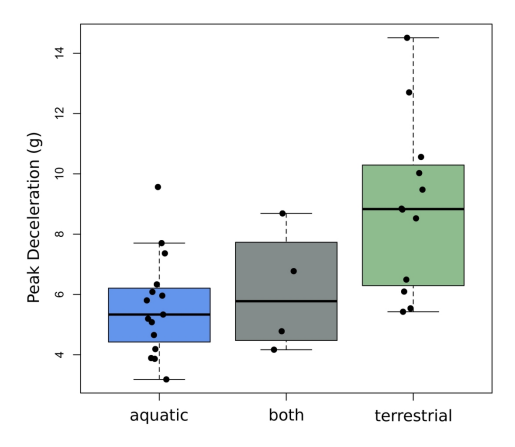


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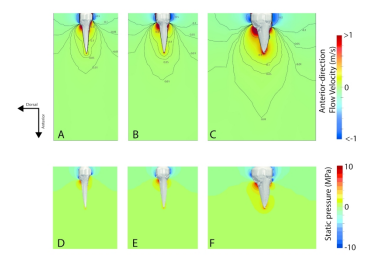


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