




Online repositories of photographs and videos provide insights into the evolution of skilled hindlimb movements in birds

Cristián Gutiérrez-Ibáñez ^{1,5✉}, Clara Amaral-Peçanha^{2,5}, Andrew N. Iwaniuk ³, Douglas R. Wylie ¹ & Jerome Baron^{2,4}

The ability to manipulate objects with limbs has evolved repeatedly among land tetrapods. Several selective forces have been proposed to explain the emergence of forelimb manipulation, however, work has been largely restricted to mammals, which prevents the testing of evolutionary hypotheses in a comprehensive evolutionary framework. In birds, forelimbs have gained the exclusive function of flight, with grasping transferred predominantly to the beak. In some birds, the feet are also used in manipulative tasks and appear to share some features with manual grasping and prehension in mammals, but this has not been systematically investigated. Here we use large online repositories of photographs and videos to quantify foot manipulative skills across a large sample of bird species (>1000 species). Our results show that a complex interaction between niche, diet and phylogeny drive the evolution of manipulative skills with the feet in birds. Furthermore, we provide strong support for the proposition that an arboreal niche is a key element in the evolution of manipulation in land vertebrates. Our systematic comparison of foot use in birds provides a solid base for understanding morphological and neural adaptations for foot use in birds, and for studying the convergent evolution of manipulative skills in birds and mammals.

¹Department of Biological Sciences, University of Alberta, Edmonton, Canada. ²Graduate Program in Physiology and Pharmacology Institute of Biological Sciences, Federal University of Minas Gerais, Belo Horizonte, Brazil. ³Department of Neuroscience, Canadian Centre for Behavioural Neuroscience, University of Lethbridge, Lethbridge, Alta., Canada. ⁴Department of Physiology and Biophysics, Institute of Biological Sciences, Federal University of Minas Gerais, Belo Horizonte, Brazil. ⁵These authors contributed equally: Cristián Gutiérrez-Ibáñez, Clara Amaral-Peçanha. ✉email: cgutierrez@ualberta.ca

The ability to grasp or manipulate objects with appendages has evolved repeatedly among land tetrapods¹. Because grasping and manipulating objects are characteristic of humans and nonhuman primates, the neural basis of these behaviors, and their association with primate evolution, including the brain, have received extensive attention^{1–4}. In the case of primates brain evolution, the development of skilled manipulation has been related to the evolution of specialized visual and motor circuits^{2,5}. Given the importance of skilled forelimb manipulation in a variety of behaviors⁴, a significant question is what drives the evolution of grasping and manipulation in some, but not all, species. In mammals, several selective forces have been proposed to explain the emergence of forelimb manipulation, including arboreal locomotion, digging, and prey handling^{1,4}. To better test this hypothesis, understanding the evolution of manipulation with the extremities in other vertebrates is needed, but this behavior has received relatively little attention outside the mammalian literature¹.

In birds, forelimbs have gained the almost exclusive function of flight, with grasping transferred predominantly to the beak^{6,7}. However, the absence of a second extremity limits the ability to manipulate objects with the beak. Consequently, many birds have evolved the ability to grasp and manipulate objects with their feet^{4,8}, including hawks, owls and falcons, which use their feet to capture and hold prey^{9,10}, as well as parrots, mousebirds and many songbirds^{8,11}. Given the diversity of clades in which pedal manipulation has evolved, birds represent a key comparison for understanding the evolutionary pathways by which pedal and manual dexterity have evolved in tetrapods. Unfortunately, there have been no systematic studies on the evolution of pedal dexterity in birds. A review of published reports by Sustaita et al.⁴ suggests that arboreality predates manipulative foot use in birds, although this was based on only a limited species sample (around 150 species). A broader approach is required to establish if the development of manipulative foot use aligns with the evolution of arboreality in birds. In addition, the extent to which manipulation skills vary among and within different avian clades is unknown. For example, some parrots use their feet to bring food to their beak and coordinate beak and foot movements for extractive foraging¹², but this does not appear to be true of all parrots¹³. Whether similar manipulation skills have evolved outside parrots is equally unclear.

In mammals, several studies have used direct observations of animals in captivity to evaluate differences in manipulative skills across species^{14–17}. While direct observations allow for a detailed study of manipulative skills, relatively few species can be examined this way, thus preventing the testing of hypotheses in a comprehensive evolutionary framework^{1,18,19}. An alternative approach is to use data deposited in digital databases by citizen scientists²⁰. This approach is particularly suitable for studies on birds, as a large (and growing) collection of pictures and videos are available: Macaulay Library at the Cornell Lab of Ornithology alone has >40 million pictures and videos of birds (<https://macaulaylibrary.org/>). Here, we use large online repositories of photographs and videos, as well as previous literature, to quantify foot manipulative skills of birds (Fig. 1a) and test several hypotheses about the selective pressures that give rise to skilled manipulation with the limbs in tetrapods.

Results

Our citizen-science approach (Fig. 1a) allowed us to obtain and score 3725 individual media files of birds using their feet to manipulate objects (Dataset 1) from a variety of sources (Fig. 1b). The observations encompassed 1054 species (i.e., close to 10% of all bird species) belonging to 13 orders and 64 families (Dataset 2). For

clades where we systematically searched for foot use in all species (see Methods), we found media of foot use behavior in 40 to 95% of species (Fig. 1c, d) in those clades. This large data set not only allowed us to assess if manipulation of objects with the feet was present, but also allowed us to quantify, in detail, (Fig. 1a, Table 1) the manipulative skills at the species or at the very least, genus level.

Arboreality drives the evolution of foot use in birds. First, we considered the absence or presence of foot use (for manipulation) at the family level to understand the origin of this behavioral trait in birds. The outer circle of dots in Fig. 2a shows the presence or absence of foot use, as well as families for which insufficient data were available (see Methods), in all 250 families of birds. We then performed an ancestral state reconstruction using a hidden Markov model. Our results show that the best supported model (Supplementary Fig. 1) is one where the transition from no foot use to foot use is indirect, through a “precursor” state, for instance, arboreality. Our results (Fig. 2a) show that the transition from an absence of foot use to this precursor occurred only once in the avian phylogeny at the base of the Telluraves, the clade that includes most of the small, arboreal neornithine birds²¹, and that after the transition to this precursor, foot use emerged independently at least 20 times (Fig. 2a). This includes independent origins of foot use for each of the raptor clades (falcons, owls, and hawks)⁹, mousebirds (Coliiformes), and parrots (Psittaciformes). We also found a single independent origin of foot use within the order Piciformes for a monophyletic clade comprising toucan-barbets (Semnornithidae), New-World barbets (Capitonidae), and toucans (Ramphastidae). Additionally, there were at least 14 independent gains of foot use within songbirds (Passeriformes). Most strikingly, our analysis recovers foot use as the most likely ancestral state of two large radiations of songbirds: the suborder Corvides²² and the superfamily Sylvioidea (Fig. 2a²³). Outside Telluraves, foot use is rare and has only evolved four times in <15 species. The most notable of these is the evolution of grasping and the ability to bring the foot and objects to the beak in eight species of swamphens that belong exclusively to the genus *Porphyrio*²⁴. Additionally, foot use has evolved in several individual species nested within larger clades: the greater coucal (*Centropus sinensis*, Cuculiformes), Australian brush-turkey (*Alectura lathami*, Galliformes), and snowy sheathbill (*Chionis albus*, Charadriiformes).

Differences in pedal manipulation skills and diet. Not only has foot use evolved independently multiple times in birds, but there are also significant differences in the manipulative skill among clades (Phylogenetic Generalized Least Squares (PGLS), $F_{5:1020} = 3.09$ $p < 0.01$; Supplementary Table 2, Fig. 2b, c). Most foot-using songbirds and piciforms have relatively simple manipulative skills (Fig. 2b, Supplementary Fig. 1b), consisting primarily of holding or clasping against a surface, with only a few species capable of grasping. The three raptorial orders have higher manipulative skills, associated with the widespread ability to grasp objects and, in many cases, bring objects to the beak (see below). Finally, parrots had the highest scores for our skill index. This is driven by the capacity of most parrots to grasp and bring object to the beak while also rotating their foot to manipulate objects, which includes not only food items but also tools and others non-food objects^{25,26}.

Next, we wanted to test if the evolution of foot use is related to a particular diet. First, we looked at the percentage of species that use their foot within each diet category (Fig. 2d, foot use, dark blue bars; yellow bars, no evidence of foot use). We also included all species that belong to families that use their feet, but for which we did not have any observations of foot use (Fig. 2d, light blue bars). This procedure is likely to overestimate the number of species that

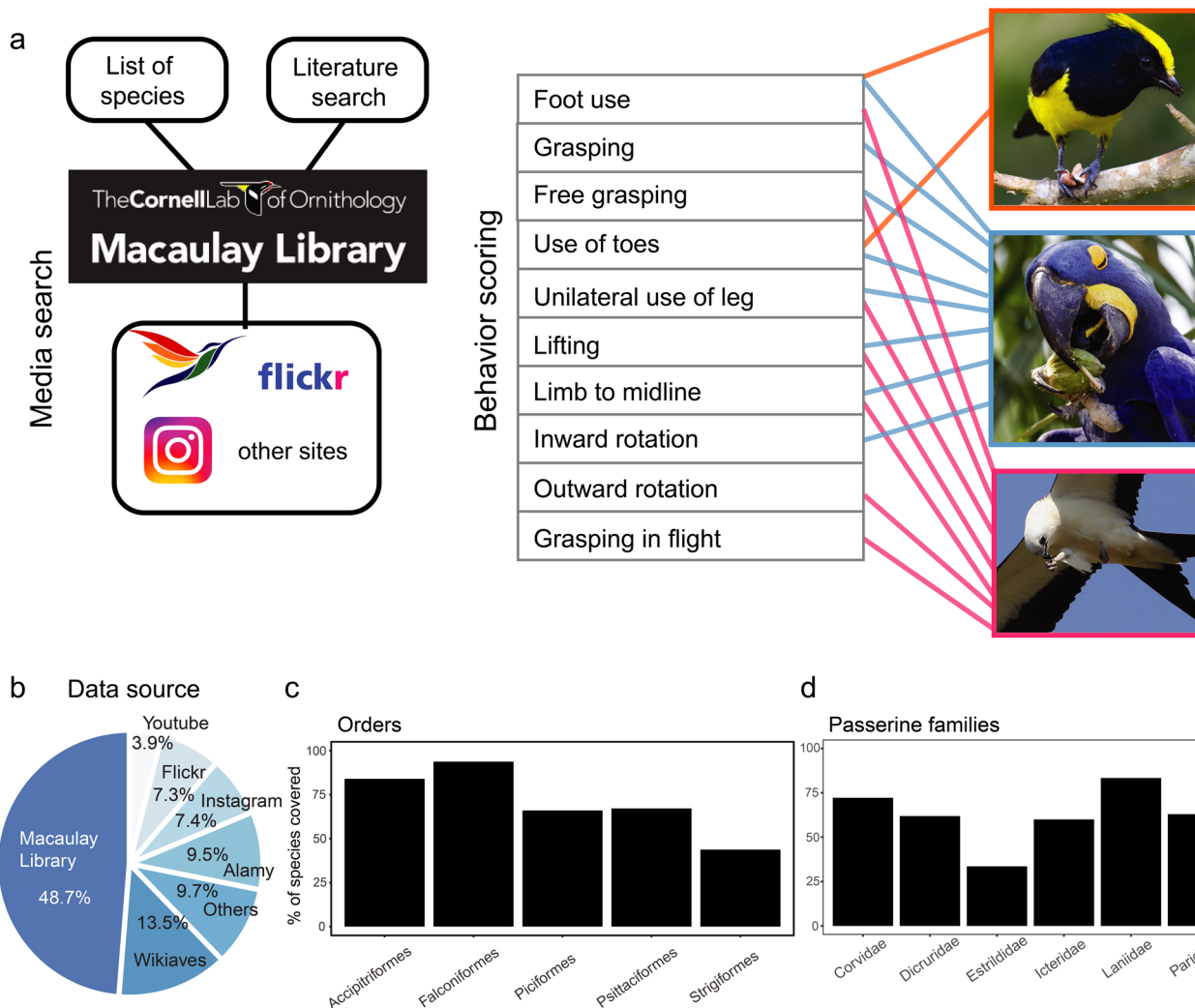


Fig. 1 Online bird media repositories are an effective source of behavioral data. **a** We combined exhaustive searches of media in clades known to use their feet with extensive literature searches to determine which clades use their feet to manipulate objects as well as compare manipulative skills among birds. Photographs, top to bottom: Sultan Tit (*Melanochlora sultanea*), Hyacinth Macaw (*Anodorhynchus hyacinthinus*), Swallow-tailed Kite (*Elanoides forficatus*). Photographer credits are listed in Supplementary Table 6. **b** Approximately 60% of media came from two citizen science bird media repositories (Macaulay Library and Wikiaves). **c, d** Species coverage for different orders and songbird families we searched systematically (all species). Coverage was at least 40% and as high as 95%.

Table 1 Behavioral scoring for each media of bird using their feet to manipulate objects.

Behavior	Description	Score
Foot use	The bird uses its claw to hold or grasp an object	0,1
Grasping	Claw/toes closed around an object	0,1
Free grasping	Claw/toes closed around an object while object is not in contact with any surface. Other parts of the claw/leg can be in contact with the surface.	0,1
Free grasping in flight	Object grasped during flight	0,1
Use of toes	Object is held or grasped with only some toes	0,1
Use of one leg	Only one leg is used to hold or grasp	0,1
Foot to beak	Claw is lifted to the beak	0,1
Foot to midline	Claw is brought to the midline	0,1
Inward rotation	The claw is rotated inward	0,1
Outward rotation	The foot is rotated outward	0,1

use their feet in some diet categories, but was included as our sampling was not uniform across birds. Although some diets seem to be associated with foot use (vertebrates, carrion), species with any diet can potentially manipulate objects with their feet. Because diet changes could have occurred after the origin of foot use, we

next performed an ancestral state reconstruction of diet in a large sample of birds to assess the ancestral diet at the main nodes where foot use has evolved independently (Supplementary Table 3). Here we found that the origin of foot use is associated with at least four different diets: vertebrates (reconstructed for the ancestors of the

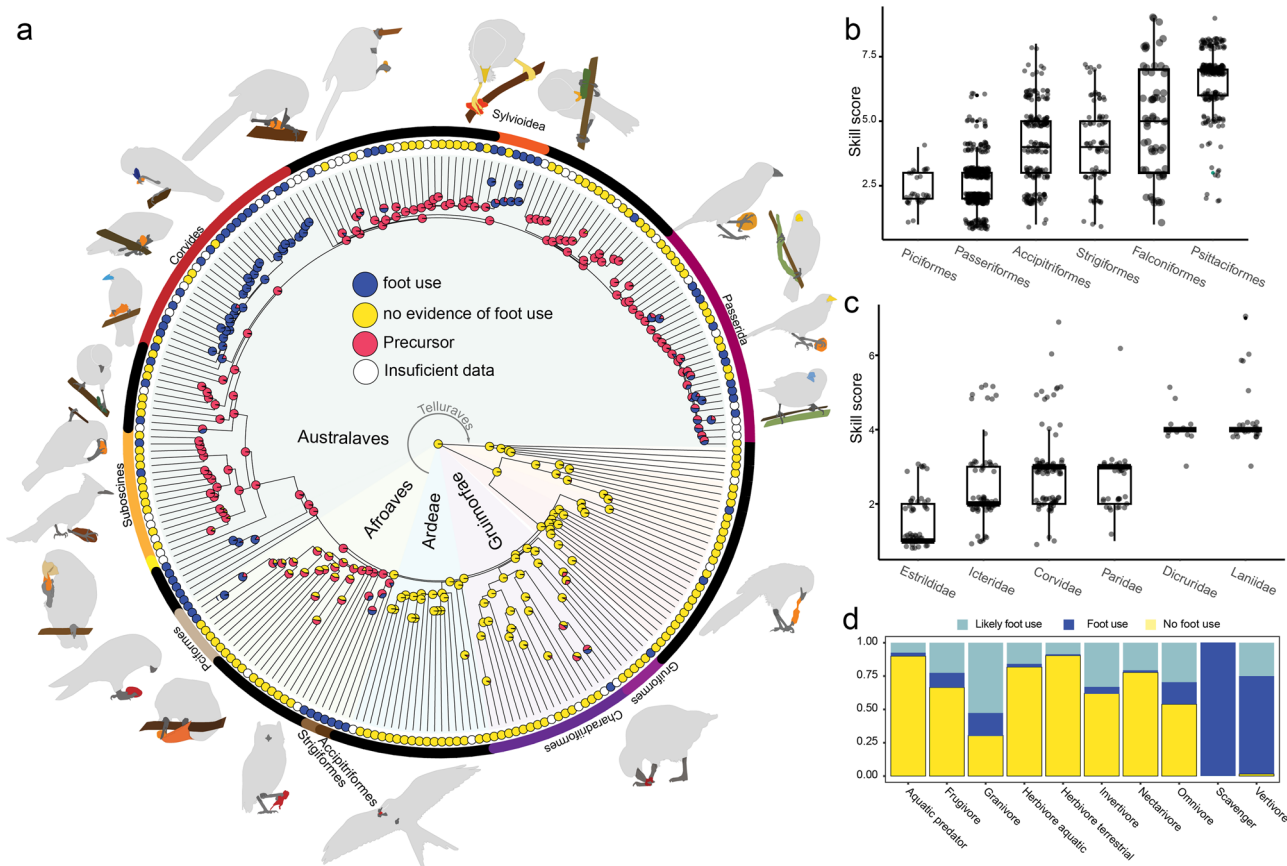


Fig. 2 After the transition to arboreality, foot use has repeatedly evolved in different clades, and is associated with multiple diets. **a** Ancestral state reconstruction with a precursor model shows that after a single evolution of a precursor state at the base of all core land birds (Telluraves), foot use has evolved repeatedly, at least 20 times. **b, c** Skill scores for different orders (**b**) and songbird families (**c**). The lower and upper hinges correspond to the first and third quartiles. Whisker extends no further than 1.5 the inter-quartile range, or distance between the first and third quartiles. **d** Percentage of species for each diet category where there is no evidence of foot use (yellow), evidence of foot use (dark blue) and likely foot use (light blue). Species are considered to belong to a particular diet category if >60% of the diet is of one type⁷². Likely foot use refers to species that belong to families where foot use was confirmed, but those species where not searched. In general, foot use is not associated with only one diet (see Supplementary Table 3).

three raptors orders), invertebrates (for the two large passerine radiations, Corvides and Sylvioidea), grains (parrots and at least two clades in the super order Passerida), and fruit (in South American barbets and toucans) (Supplementary Table 3, Supplementary Fig. 1a). These results indicate that the independent evolution of foot use in birds is not driven by a specific diet and that after the transition to arboreality, foot use can be co-opted to manipulate a variety of food items.

Foot use in raptors. Given that the three raptorial orders have evolved foot use independently but with similar diets and morphology⁹, we wanted to see if they shared similar skills in foot use. Figure 3 shows genus-level phylogenies and behavioral character matrices for the three raptor orders. Genera for which insufficient data were available were omitted (see Methods). Not surprisingly, foot use is widespread in all three orders and is the ancestral condition for each of them. Falcons and owls share similar skills; most genera are capable of free grasping and lifting objects to their beaks (Fig. 3a–c). In contrast, within Accipitriformes (hawks, eagles, and allies) the ability to lift objects to the beak while perching is rare and only partially present in a few clades (Fig. 3d). Curiously, many more accipitriform genera bring objects to their beaks during flight (e.g., Fig. 3a, bottom panel; Fig. 3d). Additionally, we found a convergence in how New and Old World vultures use their feet. In New-World vultures (Cathartidae, purple in Fig. 3d) and one of the Old World vulture

clades (Aegypiinae²⁷, green in Fig. 3d), most species are incapable of grasping objects and only hold prey against a surface. In other words, these two vulture clades share a rudimentary manipulation of food with their feet. In Old World vultures, this implies a loss of grasping ability, since Old-World vultures are nested within Accipitriformes, where grasping is widespread and ancestral. Thus, a change in diet, in this case from vertivore to scavenger, leads to a change in foot use skill. Finally, to examine other traits that may affect differences in manipulative skills among raptorial orders in more detail, we also compared their diets and body mass (Fig. 3e, f). Falcons and owls are similar in both body mass and diet, and on average are significantly smaller (PGLS, $F_{3,518} = 86.9$, $p < 0.0001$, Supplementary Table 2) than accipitriforms. The smaller mass of falcon and owls is reflected in a much higher percentage of species within these groups that feed on invertebrates (Fig. 3f), which likely explains the higher skill indices of these groups compared to accipitriforms.

Foot use varies among parrots. In contrast to falcons and owls, where foot use skills are similar in all species, foot use skills vary greatly among parrots. Our citizen science approach allowed us to score foot use in >70% of all parrot species, allowing us to explore foot use variations in detail (Fig. 4a). An ancestral state reconstruction (Fig. 4b) shows that while foot use is ancestral among parrots, it has been lost or reduced at least seven times independently. This includes the loss of foot use in several smaller

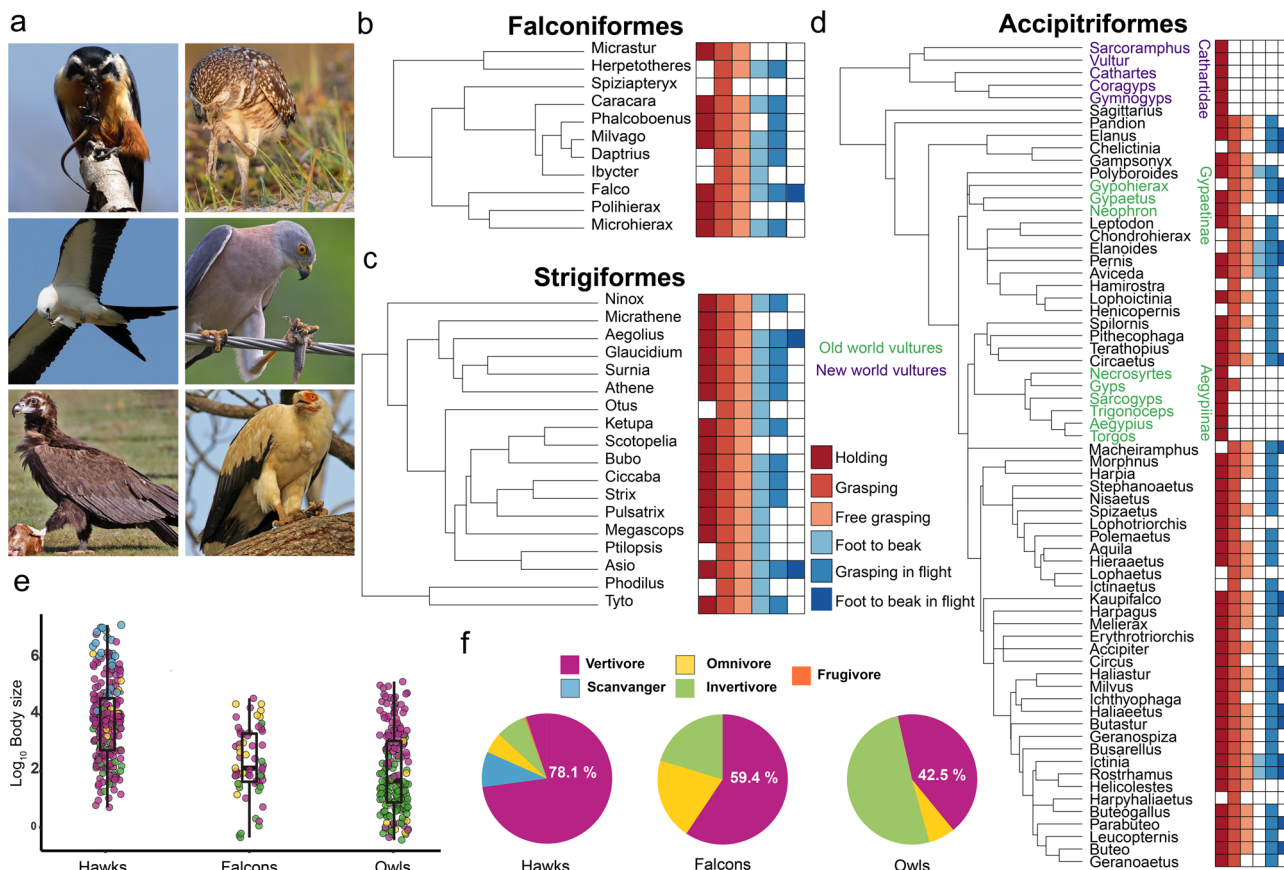


Fig. 3 Variation in foot use between and within raptors. **a** Examples of the three different raptor orders using their feet to manipulate objects. Top row, left to right: Collared Falconet (*Microhierax caerulescens*), Burrowing Owl (*Athene cucularia*). Middle row, left to right: Swallow-tailed Kite (*Elanoides forficatus*), Fiji Goshawk (*Accipiter rufitorques*). Bottom row, left to right: Palm-nut Vulture (*Gypohierax angolensis*), Cinereous Vulture (*Aegyptius monachus*). Photographer credits are listed in Supplementary Table 6. **b-d** Genus level phylogenies for falcons (**b**), owls (**c**) and hawks, eagles and New-World vultures (**d**) showing the different foot use behavior present for each genus. The key for the skilled foot use matrix is provided in (**c**), with colored squares reflecting the presence of each of the six behavioral elements. Falcons and owls have similar matrices in which the ability to grasp objects and bring them to the beak is widespread and likely ancestral. In contrast, the ability to bring the foot to the beak while perching (fourth column of the matrix) is only present in five genera of Accipitriformes, but the ability to bring the foot to the beak (sixth column) while flying is more widespread. Also shown is the convergent loss of grasping in New World vultures (in purple) and one of the Old World vulture clades (in green). **e** Body masses of the three orders of raptors. The color of each dot corresponds to the diet of each species as shown in **f**. The lower and upper hinges correspond to the first and third quartiles. Whisker extends no further than 1.5 the inter-quartile range, or distance between the first and third quartiles. **f** Percentage of each diet category in each raptor order. See methods for details on how dietary categories were assigned.

genera: *Forpus*, *Neophema*, *Neopsephotus*, *Touit*, *Melopsittacus*, *Cyclopsitta*, *Agapornis*, and *Micropsitta*. The lories and lorikeets (Loriini) had the greatest diversity in foot use; it is reduced or absent in several species (e.g., *Psitteteles*), but other species (e.g., *Trichoglossus*) have manipulative skills similar to other parrots, including the ability to grasp and bring objects to the beak. Loss of foot use is not clearly associated with one diet or niche (Supplementary Fig. 2a-d). Several of the clades where foot use has been lost or reduced are largely granivorous and feed primarily on the ground (e.g., *Neophema*, *Agapornis*, and *Melopsittacus*), but some perching and frugivorous genera also lost foot use (for example, *Touit*, Supplementary Fig. 2).

Most parrots grasp objects with the two external toes and/or turn their leg inward when bringing objects to their beaks (Fig. 4a). Our ancestral state reconstruction recovers this behavior as the ancestral state for all parrots (Fig. 4b, c). However, two clades have independently evolved the ability to grasp objects with the inner toes and turn their legs outwards when bringing objects to their beak (Fig. 4). One of these independent changes to outward rotation is at the base of the tribe Androglossini (*Amazona* and *Pionus* species, as well as related genera²⁸). The

second clade is Psittaculini (*Psittacula*, *Eclectus*, and related genera), although the racket-tails (genus *Prioniturus*, Fig. 4b, c) do not appear to rotate the foot outwards. As with loss of foot use, it is unclear whether the emergence of this new manipulative skill is associated with a particular diet, niche, or lifestyle (Supplementary Fig. 2).

Variation of foot use in songbirds. As mentioned above, we found repeated and independent evolution of foot use among passerines (at least 14 times, Fig. 2A), largely concentrated in oscines. Only two families of suboscines, Furnariidae and Oxyruncidae, exhibit foot use. In general, passerines have lower skill scores than orders in which foot use is common (Fig. 2b, c), which is reflected in their tendency to hold objects against the ground or perch but not grasping (Supplementary Fig. 3). The exceptions are families within the superorder Corvidae, where the ability to freely grasp objects occurs in many families and the ability to bring objects to the beak has evolved in at least three different families: shrikes (Laniidae), drongos (Dricuridae) and vangas (Vanguidae) (Fig. 5a, b). Outside of this clade, although there are a few species capable of grasping objects while hanging

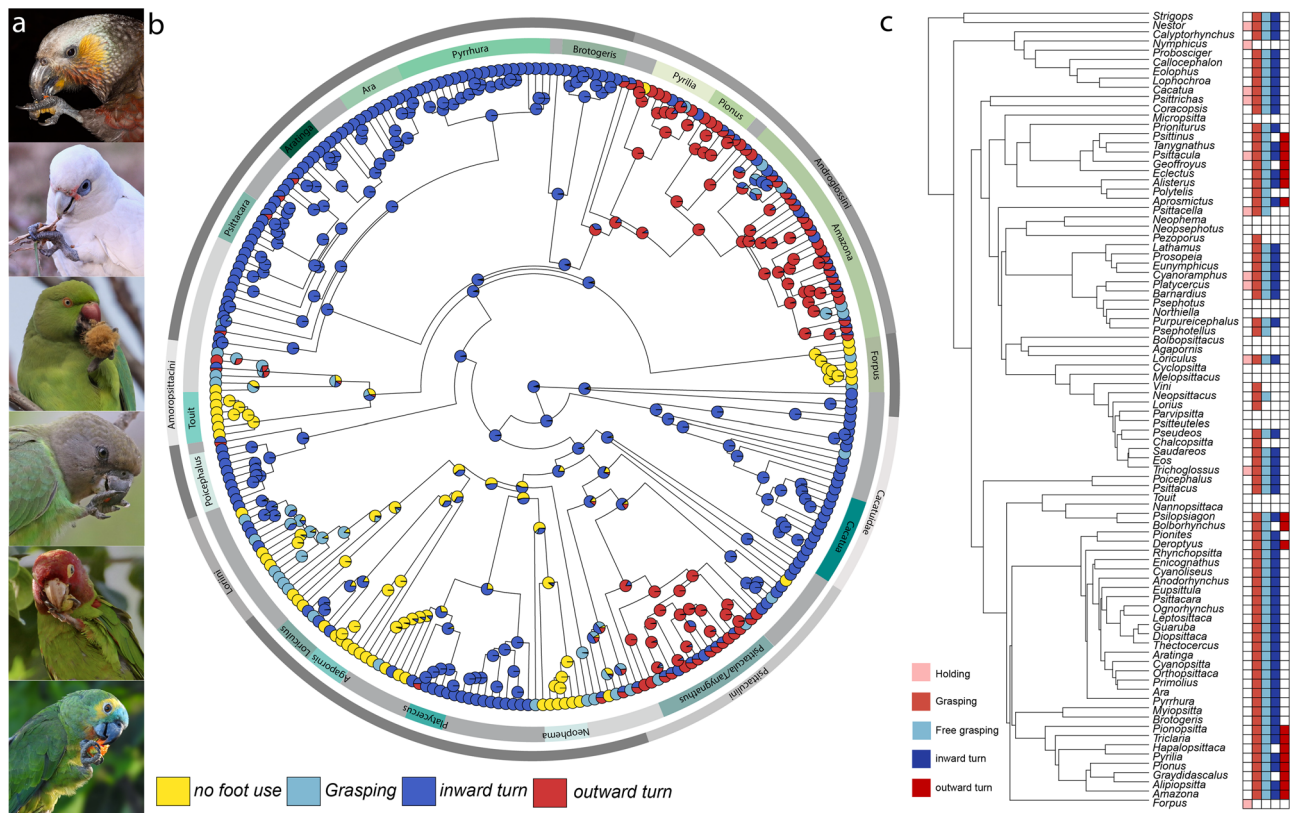


Fig. 4 Variation in foot use among parrots shows several instances of convergent evolution. **a** Examples of foot use among the main clades of parrots. From top to bottom: New Zealand Kaka (*Nestor meridionalis*), Red-masked Parakeet (*Psittacara erythrogenys*), Brown-headed Parrot (*Poicephalus cryptoxanthus*), Turquoise-fronted Parrot (*Amazona aestiva*), Little Corella (*Cacatua sanguinea*), Rose-ringed Parakeet (*Psittacula kramera*). Photographer credits are listed in Supplementary Table 6. **b** Ancestral state reconstruction of foot use in parrots shows that foot use is the ancestral state for all parrots, but this behavior has been lost at least five times. The ancestral state reconstruction also shows that an outward rotation of the foot when bringing the foot to the beak has evolved independently at least twice (dark red). **c** Genus level phylogeny of parrot shows in detail where foot use has been lost and where differences in foot use have emerged. Colored squares reflect the presence of each of five different behavioral elements.

from one foot (e.g., Aegithalidae, Remizidae, Paridae), no other species are capable of bringing an object to the beak while perching. Only one family, Callaeidae (New Zealand wattlebirds), is capable of free grasping.

To better understand what might drive differences in manipulation skill among passerines, we systematically searched for foot use in three closely related families within the superfamily Corvoidea: drongos (Dricuridae), shrikes (Laniidae) and crows, jays and magpies (Corvidae) (Fig. 5c–e). Most drongo and shrike species use their feet to grasp objects and are capable of free grasping and some drongos and shrikes can even bring objects to the beak with their foot (Fig. 5c, d). In contrast, grasping is rare in corvids: only one species appears to have the ability to free grasp (the yellow-billed chough, *Pyrrhocorax graculus*), and no corvids bring objects to their beaks with their feet (Fig. 5e). Corvidae differ from the other two families in that their diet is largely omnivorous, whereas drongos and shrikes are predators that feed on insects and small vertebrates (Fig. 5). Thus, the ability to manipulate objects with the feet is not associated with dietary breadth, but rather specific dietary types in songbirds.

Discussion

Here we have taken advantage of the vast number of pictures and videos of birds stored in online citizen science repositories to study the evolution of a largely ignored behavior in birds: skilled pedal manipulation. Our results show that this approach can result in extensive coverage (up to 95% of species in some clades, Fig. 1b), and that is sensitive enough to detect small and

previously undescribed differences in behavior, such as the differences in manipulation among raptor clades (Fig. 3), the emergence of new manipulation skills among parrots (Fig. 4), and the loss of foot use in some parrots (Fig. 4, Supplementary Fig. 3). While some studies have already used this resource to study different aspects of bird biology^{29,30}, the scale at which we employed this approach (covering 10% of all birds) is unprecedented. Further, this is the first study outside of mammals to quantify and compare manipulative skills in a broad phylogenetic context and test which factors might drive the evolution of limb manipulation across vertebrates¹⁴.

In mammals the evolution of manipulation with the forelimb is typically associated with arboreality, digging, and prey manipulation (Reviewed in 4). Here we show that the evolution of object manipulation with the feet in birds is a complex interaction of several putative selective forces. First, our results support the previous suggestion⁴ that the evolution of foot use in birds is greatly facilitated by the transition to arboreality. Our ancestral state reconstruction shows that the most likely scenario for the evolution of foot manipulation in birds is one where a transitional state is required, and that this state evolved once at the node that gave rise to the core landbirds, Telluraves. Given that this clade includes most of the small arboreal neornithine birds²¹ and arboreality is the ancestral state of all Telluraves³¹, we suggest that the transitional state required from the evolution of foot use is most likely an arboreal niche. Our ancestral state reconstruction of ancestral diet among the principal clades that have independently evolved foot use suggests that this behavior is related to

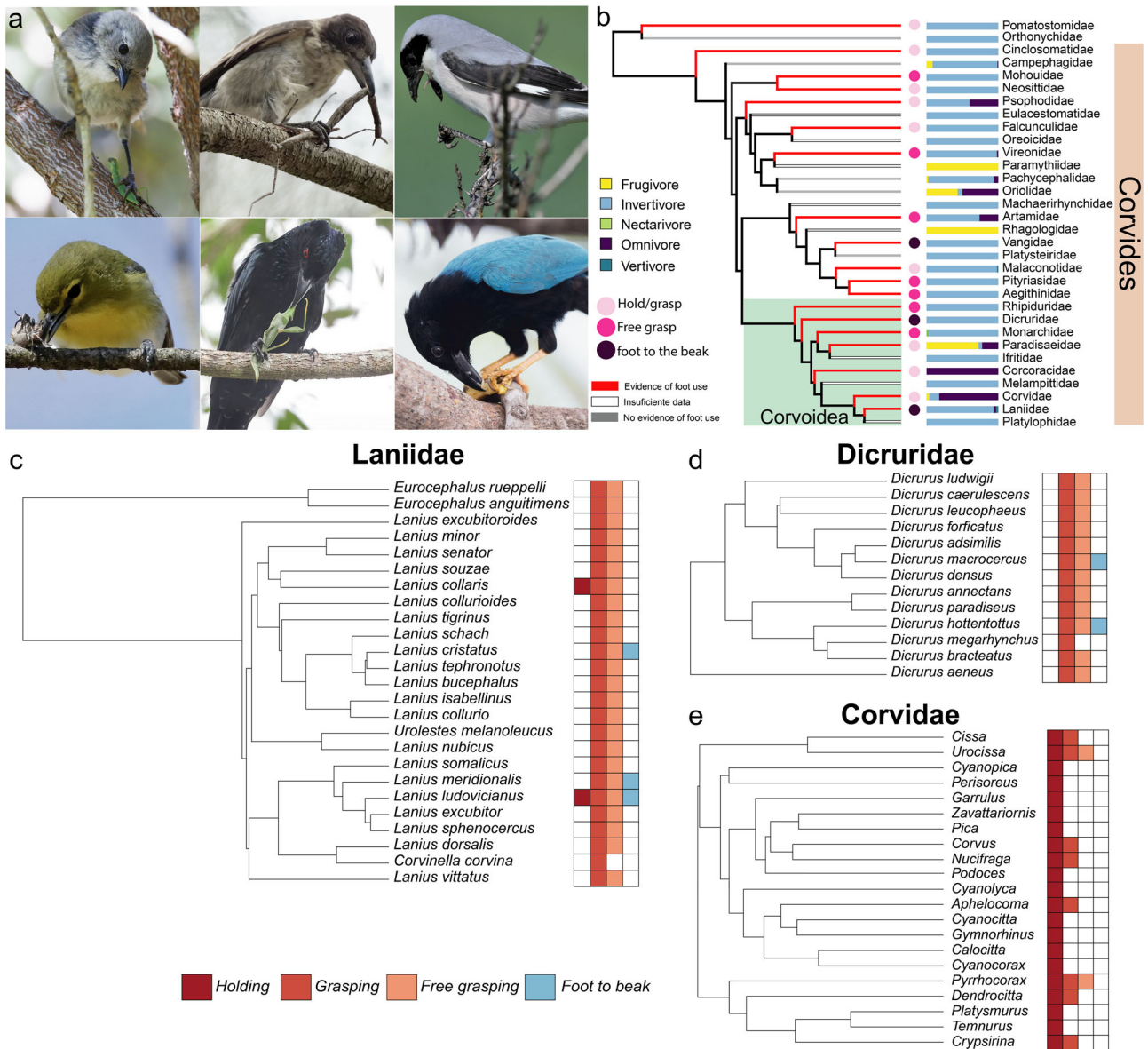


Fig. 5 Changes in foot use are driven by diet changes among the suborder Corvides. **a** Photographs showing examples of foot use among 6 of the 19 families within the superorder Corvides that use their feet to manipulate objects. Top row, from left to right: Whitehead (*Mohoua albigilla*), Pied Butcherbird (*Cracticus nigrogularis*), Lesser Gray Shrike (*Lanius minor*). Bottom row: Yellow-throated Vireo (*Vireo flavifrons*), Spangled Drongo (*Dicrurus bracteatus*), Yucatan jay (*Cyanocorax yucatanicus*). Photographer credits are listed in Supplementary Table 6. **b** Family level phylogeny of the suborder Corvides shows widespread foot use and that most of the Corvides families are insectivores, with only a few shifts to omnivory or frugivory. Bars represent the percentage of species for each within each diet category and the circles show the different manipulation skills of each family. **c**, **d** Show a character matrix for shrikes (Laniidae) and drongos (Dicuridae) adjacent to species level phylogenies. Both families are almost strictly invertivores and most species have the ability to free grasp and even bring object to the beak. In contrast, the character matrix at the genus level for crows and jays (**e**, Corvidae) shows that most species do not grasp.

different diets in different clades (Fig. 2, Supplementary Table 3). Thus, once birds evolved the morphological traits for perching, these traits were subsequently and repeatedly exapted for grasping and manipulating food items that included, but were not limited to, vertebrate and invertebrate prey. However, the relationship between foot use and diet is not straightforward. In the barbets and toucans, for example (Supplementary Fig. 1), frugivory is the ancestral state for a clade that includes three families that use their feet to manipulate objects, as well as two other families like the Asian (Megalaimidae) and African barbets (Lybiidae) that do not appear to use their feet to manipulate objects. Similarly, hornbills (Bucerotidae) have convergently evolved similar morphology and ecological niches to toucans³², but there is no

evidence that they use their feet to manipulate objects. Another example is that foot use evolved in relation to eating invertebrates in Corvides and Sylvioidea (Fig. 2a, Supplementary Table 3), but many other clades of passerines and Telluraves have diets based on invertebrates yet do not use their feet to hold or manipulate prey. In fact, several bird groups have evolved alternatives to using their feet to handle and feed on invertebrate and vertebrate prey. For example, trogons, coraciiform birds (e.g., kingfishers, motmots, bee-eaters), and roadrunners (*Geococcyx*, Cuculiformes) that feed on large insects and small vertebrates grasp their prey with the beak and beat and shake it repeatedly against a surface to kill and clean the prey before eating it whole³³. This same behavior can also be observed in many suboscines that feed

on large insects, such as antbirds (Thamnophilidae)³⁴ and flycatchers (Tyrannidae)³⁵. The evolution of skilled foot use in birds is therefore likely to be a product of diet (insects, small vertebrates), foraging or capture method (e.g., gleaning vs hawking), and morphology, but the relative contributions of each remain uncertain.

A key component of skilled manipulation in mammals is the ability to grasp and bring objects to the mouth^{1,3}. Although the transition to arboreality has resulted in the repeated evolution of foot use among birds, the ability to grasp, and particularly to bring objects to the beak, has evolved only a few times. This feature is restricted to owls, falcons (Fig. 3), mousebirds, parrots (Fig. 4) and three families within the suborder Corvidae (Fig. 5b). As discussed above, the evolution of foot use is related to several different diets and not necessarily in a consistent fashion. However, there is some evidence to suggest that the evolution of pedal grasping and bringing objects to the beak has its origin with a predatory diet. While our ancestral state reconstruction recovers granivory and frugivory as the likely ancestral states of extant parrots and mousebirds respectively (Supplementary Table 3), there is evidence that extinct, stem branches of both of these clades had a more predatory diet. Most stem group Coliiformes, as well as an extinct sister clade (Sandcoleidae) had morphological traits, such as shortened proximal phalanges in the second and fourth toe³⁶, that suggest they were adapted to capture and manipulate large objects, including prey. Also, stem Coliiformes had proportionally longer beaks than extant Coliiformes, as well as other beak adaptations, that suggest a less strictly frugivorous diet^{36,37}. Similarly, fossil evidence suggest that two of the stem pan-Psittaciformes clades had many raptor-like adaptations in the beak and foot^{38–40}. A raptorial diet for stem parrots is also supported by enhanced fat digestion and absorption in parrots, which is shared with the three raptorial orders⁴¹. Thus, while the ancestor of extant Psittaciformes and Coliiformes was likely a frugivore and/or granivore (Supplementary Table 3), it is possible that the earlier ancestors of both clades had a raptorial ecology that included the ability to grasp and bring objects to the beak, and that only later was this ability exapted for the manipulation of fruit and seeds. Nevertheless, it is possible that the ability to grasp and bring objects to the beak has evolved independently, and for different reasons in different groups of birds. In the case of parrots for example, it is possible that the combination of a zygodactyl foot, which allows for a firm grip³⁸, and a diet based on extractive foraging of seeds and fruits¹³ is what drove the evolution of this behavior.

The evolution of skilled manipulation in mammals is associated with changes to sensory and motor circuits, as well as adaptations of the skeleton and integument of the manus⁴². In the somatosensory system, mammals that use their forelimbs for haptic searching and complex object manipulation evolve glabrous skin and an increase in the number, acuity and sensitivity of touch receptors⁴³. Touch receptors can be found in avian feet⁴⁴ and share some similarities with receptors in the mammalian manus. For example, the plantar skin of owls has specialized tubercles, each of which contains a Herbst corpuscle (the avian equivalent of the mammalian Pacinian corpuscle), and the claw has a dual and detailed topographic representation in the anterior Wulst, which is equivalent to the somatosensory cortex in birds⁴⁵. Based on this, one would predict the same to be true of falcons, hawks, and parrots, but currently this is unknown. In the motor system, differences in forelimb manipulation skills among mammals are also correlated with anatomical changes. Whereas most mammals capable of manipulating objects with the hands have a corticospinal tract (direct projections from the cortex to the spinal cord), this is particularly developed in primates, where both ipsilateral and contralateral projections are present, and

cortical projections make direct contact with motoneurons in the spinal cord². An equivalent “corticospinal” projection does not appear to be well developed in birds⁴⁶, although many aspects of motor control in birds remains understudied. The closest to a corticospinal tract in birds has been found in the zebra finch (*Taeniopygia castanotis*), where the anterior Wulst projects to the spinal cord, but these fibers reach only to the level of C7, and are sparse⁴⁷. Corticospinal projections to the cervical spinal cord have also been suggested in owls⁴⁸, but in parrots, despite the high manipulative skills with the feet, there does not appear to be any direct projections from the Wulst or other parts of the pallium to spinal cord regions associated with hindlimb movements⁴⁶. Whether other birds that use their feet to manipulate objects have direct projections to the spinal cord is unknown. In fact, almost nothing is known about the control of voluntary movement in birds; it even remains unclear if birds possess a region that functions similarly to mammalian primary motor cortex⁴⁸.

Despite the gaps in our understanding of avian somatosensory and motor systems, it is clear that skilled manipulation has evolved in parallel in mammals and birds. In addition, arboreality and diet appear to have played a role in the evolution of skilled manipulation in birds, as has also been suggested for mammals. Owing to these parallels, further research into skilled hindlimb use in birds will provide new insights into the neural basis of skilled limb use more generally and may also aid in the interpretation of the behavior of extinct species. In birds, it is well established that both an arboreal niche and a raptorial diet are correlated with the morphology of the pedal phalanges^{4,49}, which has been used extensively to infer the raptorial lifestyle (and foot use) of fossil birds and non-avian dinosaurs³⁷. Nonetheless, it is unclear whether other diets or how foot use can be predicted from pedal morphology. The data presented here on the distribution of foot use along the avian phylogeny and the differences between clades in manipulative skills may set the stage for more accurate comparisons of pedal morphology with behavior and diet in birds thereby allowing for better prediction of the ecology and behavior of extinct species.

Materials and methods

Database construction. To study manipulation of objects with the foot in birds, we combined exhaustive searches of media in clades known to use their feet with extensive literature searches to determine which clades use their feet to manipulate objects. To determine which avian families exhibit skilled foot use, we first performed a literature search for reports of foot use. This included a systematic search of *Birds of the World*⁵⁰ for any report of foot use. Additionally, we performed full text searches of ornithological journals in the Biodiversity Heritage Library (BHL) for the phrases “a foot,” “under a foot,” “its foot,” “its feet,” “held under a foot,” and others. Along with previous reviews of the literature^{8,51}, we were able to collect references on foot use in 259 species of birds belonging to 85 families (Supplementary Table 7). However, some of these reports are anecdotal, one-time occurrences, or even mistaken. For example, Clark⁸ cited Skutch⁵² as evidence of foot use in trogons, but the cited reference is about a toucan species that uses its feet to manipulate objects. Other reports, such as that of the tooth-billed pigeon (*Didunculus strigirostris*), seem to be a repetition of a single report without any supporting evidence⁵³. To remedy this, for each of the species reported in the literature as using their feet to manipulate objects, we performed an in-depth search for media (described below). Because some of these species have very few pictures available (like the endangered tooth-billed pigeon, which has 0 in the Macaulay library and only a few outside of it), if no pictures of the species reported as using their feet were found, we then searched for foot use in additional species in the same family. To maximize the probability of finding foot use, we searched the top 5% of species with the most pictures in the Macaulay Library within that family. With this method we were able to confirm foot use in 59 of the 85 families where foot use has been reported. In total there were 26 families where foot use had been reported in the literature, but for which we could find no media showing foot use. These are families where foot use is either very rare or misreported. The former applies to families like herons (Ardeidae) and flycatchers (Tyrannidae), where a few species have been reported to use their feet, but no photos or videos of foot use were found. These families were not considered as having the ability to use their feet to manipulate object in our analysis. With this method, it is possible that

we missed some species that use their feet to grasp or manipulate objects, but it is likely that these are few and do not belong to any clade where foot use is widespread.

To quantify and compare pedal manipulation skills across species, we searched for videos or pictures of foot use associated with food or other types of object manipulation, an approach similar to previous studies⁵⁴. Based on previous literature^{8,51} we first systematically searched all species of those orders or families in which foot use is widely reported. These included all diurnal and nocturnal raptors (Strigiformes, Falconiformes, and Accipitriformes), seriemas (Cariamiformes), parrots (Psittaciformes), mousebirds (Coliiformes), and swamphens (genus *Porphyrio* and Gruiformes). We also systematically searched three closely related families in the order Piciformes: Ramphastidae (toucans), Capitonidae (New-World barbets) and Semnornithidae (toucan-barbets). Within the songbirds (Passeriformes), we systematically searched for six families in which foot use is widely reported: Corvidae (crows, jays, and allies), Paridae (tits, chickadees, and allies), Druridae (drongos), Icteridae (blackbirds, caticques, and grackles), Estrilidae (finches), and Laniidae (shrikes and allies). For the list of species we use the taxonomy of birdtree.org⁵⁵.

The sites used to search for photographs or videos were Macaulay Library (<https://www.macaulaylibrary.org>), Wikiaves (<https://www.wikiaves.com.br>), Google Images, Flickr, Alamy, Youtube, Twitter, and Instagram. We always searched pictures first in the Macaulay Library because it is a curated source for species identification and contains all bird species in the world. Up to 2000 media files (pictures and videos) were examined for each species. Only ~10% of the species in the Macaulay Library have >2000 media files (as of 2021, Supplementary Fig. 4e), which means that for 90% of the species examined, we looked at all media available in this data repository. The second main site was Wikiaves. It is also a curated source for species identification, but contains only Brazilian bird species. Then the species were searched on the following sites in this order: Google Images, Flickr, Alamy, YouTube, Twitter, and Instagram. The keywords used to search on these sites were: “Latin name”, “English name”, “Latin name + feeding”, “English name + feeding”, “Latin name + eating”, and “English name + eating”. Species with few photos (<500) in the first two sites were systematically searched using all keywords.

Detection thresholds. An important issue was to distinguish between species that do not use their feet for manipulation from those that do, but where only a limited number of photos and videos are available and therefore with a lower likelihood to detect the behavior. To address this problem, we estimated detection thresholds based on media availability. This requires knowing the number of files available for each species, which are published annually by the Macaulay Library (<https://www.macaulaylibrary.org/resources/media-target-species/>). For all our calculations, we used the March 2021 update because it is the closest to the dates when our searches occurred. We used the sum of all pictures and videos available for each species. First, we tested whether our skill index (see below) correlated with the number of media items. Although one could expect that more pictures would be associated with a higher number of different behaviors and therefore a higher skill index, we found no significant correlation (PGLS, $F_{1,1018} = 2.877$, p -value = 0.26, Supplementary Table 2, Supplementary Fig. 4c). We then calculated a “detection probability” for each species at the clade level (i.e., family or order), which represents the detection threshold of the number of media at which there was a 75% chance of detecting foot use (Supplementary Fig. 4a, b). For this purpose, we fitted a logistic regression for each clade with the number of media as the predictor using the `ggplot`⁵⁶ package in R⁵⁷.

We then used the lower 95% confidence interval of that curve to determine the number of media at which the 75% probability of detection threshold was crossed. In the case of parrots, because some species do not use their feet at all, we removed all species that had >500 media, but no foot use, to calculate the threshold. Supplementary Table 4 shows the calculated threshold for each systematically searched clade. The 75% threshold varies greatly among clades and was as low as 77 photos/videos in parrots and as high as ~2700 for some of the songbird families. We then used this threshold (rounded up conservatively, Supplementary Table 4) to determine in which species there were not enough media entries to determine if foot use was present (media below the threshold) and in which foot use was not present (media above the threshold). In the case of the Laniidae and Dicuridae, the number of species is too small to fit a logistic regression so we used a threshold calculated for the Corvoidea.

Prehensile behavior taxonomy and scoring. To record and quantify manipulative behavior with the feet, we followed previous research in mammals¹⁵ and calculated a pedal dexterity index. Table 1 and Fig. 1a show the 10 behaviors that were recorded. In each picture or video, we recorded the presence of any of these behaviors and assigned a score of one. Dataset1 shows the scores for each individual picture or video. A dexterity index for each species (Dataset 2) was then computed as the sum of all the behaviors present for each species, so the dexterity index for each species was a number between 0 and 10. Figure 1a shows three examples of this scoring. This scoring system allow us to compare the general skill level across species as well as the presence or absence of specific behaviors, such as grasping or the ability to lift objects to the beak.

Phylogenies and ancestral state reconstruction. For the family-level analysis, we used a family-level tree generated by Toda et al.⁵⁸. Briefly, in this tree the

backbone is based on a phylogenomic supertree⁵⁹. Relationships for passerines are based on Oliveros et al.²³. Family names correspond to Clements⁶⁰. Families missing in the Kimball et al.⁵⁹, backbone (Chionidae, Pluvianellidae, Pluvianidae, Ibidorhynchidae and Stercorariidae) were added using a midpoint rooting method implemented in `addTaxa`⁶¹, based on their phylogenetic position⁶². Other families with more uncertain positions, such as Semnornithidae and Teretistridae, were also included using the midpoint rooting method implemented in `addTaxa`^{63,64}. To build genus-level trees for owls (Strigiformes), hawks, eagles, vultures (Accipitriformes), and falcons (Falconiformes) we extracted 1,000 fully resolved trees from birdtree.org⁵⁵, and built a maximum clade credibility (MCC) tree using `phangorn`⁶⁵. The same procedure was used for the passerine families shown in Fig. 5 and Supplementary Fig. 3. We then pruned each tree until only one species per genus remained. In the case of parrots (Psittaciformes) the phylogeny used was a recently published MCC consensus phylogeny⁶⁶. Ancestral state reconstruction of foot use was performed on our family-level phylogeny, as well as on the maximum clade credibility tree at the species level for parrots (Supplementary Tables 1, 5), using the R package `corrHMM`⁶⁷, which implements a maximum-likelihood method that allows multi-state characters and polymorphic taxa. We then compared the fit of three different models: (1) an all-rates different matrix (ARD) model, in which all possible transitions between states receive distinct values; (2) a one-parameter equal rates (ER) model, in which a single rate is estimated for all possible transitions; (3) a symmetric (SYM) model in which forwards and reverse transitions between states are constrained to be equal. In the family-level reconstruction we also tested a fourth model, the precursor model of Marazzi et al.⁶⁸. In this precursor model (PREC), the observed state (foot use) could be exhibited only by a lineage that had transitioned from no foot use to the precursor state first. Thus, transitions from the no foot use directly to foot use were prohibited. In the family-level ancestral state reconstruction, families in which not enough data was available to assess the presence or absence of foot use were entered as unknown, which assigns an equal probability to both characters. Families entered as not enough data were those in which the total number of media entries for the whole family was <1500. This threshold was based on the upper range of detection thresholds calculated for individual orders and families (see above, Supplementary Fig. 4a, b, Supplementary Table 4). Because we found that at the family-level, the number of pictures per family in the Macaulay Library is strongly correlated with the research effort for each family (Supplementary Fig. 4d), this threshold not only reflects the probability of detect foot use behavior through pictures/video, but also in literature reports. Research effort for 8648 species of birds was obtained from Ducatez et al.⁶⁹, which corresponds to 206 of the 249 families of birds. Research effort and total number of media were added for each family.

Diet information for all species was obtained from Pigot et al.³², which used an updated version of the EltonTraits dataset^{70,71}. While updated, the ecology and diet of many birds species remains poorly known and therefore some of this data is inevitably inaccurate and may change in the future. In the Pigot et al.³², dataset, diet (trophic niche) is a categorical character in which a species is considered to belong to particular niche if >60% of the diet is of one type. Omnivores are species where no diet is >60%. These categories are shown in Fig. 2 and Supplementary Fig. 2 for parrots. We also used these categories to reconstruct the ancestral diets of the main groups in which foot use was expressed. For simplicity we used a MCC tree from⁷² but that also correspond to the Hackett backbone from birdstrees.org. Ancestral state reconstruction was also performed with `corHMM`, and, as described above, the best fitting model based on AIC values was used to extract ancestral diets (Supplementary Table 2). Because omnivory can be prevalent in some clades, we also used the original percentage base data from the EltonTraits⁷⁰ to reconstruct ancestral percentage of a given diet. For New-World barbets and toucans, we reconstructed the percentage of fruit and vertebrates (Supplementary Fig. 1c, d), while in parrots we reconstructed the percentage of seeds, fruit and nectar (Supplementary Fig. 2b-d). In this case, we treated diet as a continuous character and used the `contmap` function in the package `phytools`⁷³ to fit this character and ancestral states to the corresponding phylogenies. Phylogenetic Generalized Least Squares (PGLS) were performed using the R packages `ape`⁷⁴ and `nlme`⁷⁵.

Images of birds using their feet. All images of birds using their feet were from the Macaulay Library at Cornell University. Catalog numbers and photographer credits are listed in Supplementary Table 6.

Reporting summary. Further information on research design is available in the Nature Portfolio Reporting Summary linked to this article.

Data availability

All data are included in the electronic supplementary materials and Supplementary Data 1, 2.

Received: 6 March 2023; Accepted: 18 July 2023;

Published online: 15 August 2023

References

- Iwaniuk, A. N. & Whishaw, I. Q. On the origin of skilled forelimb movements. *Trends Neurosci.* **23**, 372–376 (2000).
- Whishaw, I. Q. Did a change in sensory control of skilled movements stimulate the evolution of the primate frontal cortex? *Behav. Brain Res.* **146**, 31–41 (2003).
- Karl, J. & Whishaw, I. Different evolutionary origins for the reach and the grasp: an explanation for dual visuomotor channels in primate parietofrontal cortex. *Front. Neurol.* **4**, 208 (2013).
- Sustaita, D. et al. Getting a grip on tetrapod grasping: form, function, and evolution. *Biol. Rev.* **88**, 380–405 (2013).
- Padberg, J. et al. Parallel evolution of cortical areas involved in skilled hand use. *J. Neurosci.* **27**, 10106–10115 (2007).
- Matsui, H. et al. Adaptive bill morphology for enhanced tool manipulation in New Caledonian crows. *Sci. Rep.* **6**, 22776 (2016).
- Navalón, G., Bright, J. A., Marugán-Lobón, J. & Rayfield, E. J. The evolutionary relationship among beak shape, mechanical advantage, and feeding ecology in modern birds. *Evolution* **73**, 422–435 (2019).
- Clark, G. A. Holding food with the feet in passerines. *Bird.-Band.* **44**, 91–99 (1973).
- McClure, C. J. W. et al. Commentary: defining raptors and birds of prey. *rapt* **53**, 419–430 (2019).
- Sustaita, D. Musculoskeletal underpinnings to differences in killing behavior between North American accipiters (Falconiformes: Accipitridae) and falcons (Falconidae). *J. Morphol.* **269**, 283–301 (2008).
- Demery, Z. P., Chappell, J. & Martin, G. R. Vision, touch and object manipulation in Senegal parrots *Poicephalus senegalus*. *Proc. R. Soc. B: Biol. Sci.* **278**, 3687–3693 (2011).
- O'Hara, M. et al. Extraction without tooling around—The first comprehensive description of the foraging- and socio-ecology of wild Goffin's cockatoos (*Cacatua goffiniana*). *Behaviour* **156**, 661–690 (2019).
- Forshaw, J. M. & Knight, F. *Parrots of the World*. (Princeton University Press, 2010).
- Iwaniuk, A. N., Pellis, S. M. & Whishaw, I. Q. The relationship between forelimb morphology and behaviour in North American carnivores (Carnivora). *Can. J. Zool.* **77**, 1064–1074 (1999).
- Iwaniuk, A. N., Pellis, S. M. & Whishaw, I. Q. The relative importance of body size, phylogeny, locomotion, and diet in the evolution of forelimb dexterity in fissiped carnivores (Carnivora). *Can. J. Zool.* **78**, 1110–1125 (2000).
- Iwaniuk, A. N., Nelson, J. E. & Whishaw, I. Q. The relationships between brain regions and forelimb dexterity in marsupials (Marsupialia): a comparative test of the principle of proper mass. *Aust. J. Zool.* **48**, 99–110 (2000).
- Whishaw, I. Q., Sarna, J. R. & Pellis, S. M. Evidence for rodent-common and species-typical limb and digit use in eating, derived from a comparative analysis of ten rodent species. *Behav. Brain Res.* **96**, 79–91 (1998).
- Peckre, L. R. et al. Food mobility and the evolution of grasping behaviour: a case study in strepsirrhine primates. *J. Exp. Biol.* **222**, jeb207688 (2019).
- Whishaw, I. Q. & Karl, J. M. in *Feeding in Vertebrates: Evolution, Morphology, Behavior, Biomechanics* (eds. Bels, V. & Whishaw, I. Q.) 159–186 (Springer International Publishing, 2019).
- Frigerio, D. et al. Citizen science and wildlife biology: synergies and challenges. *Ethology* **124**, 365–377 (2018).
- Sangster, G. et al. Phylogenetic definitions for 25 higher-level clade names of birds. *Avian Res.* **13**, 100027 (2022).
- Jönsson, K. A. et al. A supermatrix phylogeny of corvid passerine birds (Aves: Corvides). *Mol. Phylogenetics Evolution* **94**, 87–94 (2016).
- Oliveros, C. H. et al. Earth history and the passerine superradiation. *Proc. Natl Acad. Sci.* **116**, 7916–7925 (2019).
- Pellis, S. M. Head and foot coordination in head scratching and food manipulation by purple swamp hens (*Porphyrrio porphyrio*): rules for minimizing the computational costs of combining movements from multiple parts of the body. *Int. J. Comp. Psychol.* **24**, 255–271 (2011).
- Auersperg, A. M. I. et al. Social transmission of tool use and tool manufacture in Goffin cockatoos (*Cacatua goffini*). *Proc. R. Soc. B: Biol. Sci.* **281**, 20140972–20140972 (2014).
- Auersperg, A. M. I. et al. Combinatory actions during object play in psittaciformes (*Diopsittaca nobilis*, *Pionites melanocephala*, *Cacatua goffini*) and corvids (*Corvus corax*, *C. monedula*, *C. moneduloides*). *J. Comp. Psychol.* **129**, 62–71 (2015).
- Lerner, H. R. L. & Mindell, D. P. Phylogeny of eagles, Old World vultures, and other Accipitridae based on nuclear and mitochondrial DNA. *Mol. Phylogenet. Evol.* **37**, 327–346 (2005).
- Joseph, L., Toon, A., Schirtzinger, E. E., Wright, T. F. & Schodde, R. A revised nomenclature and classification for family-group taxa of parrots (Psittaciformes). *Zootaxa* **3205**, 26–40 (2012).
- Tubelis, D. P. & Sazima, I. Nuptial gifts among Brazilian cuckoos: an outline based on citizen science. *Ornithol. Res.* **29**, 188–192 (2021).
- Pyle, P. Examination of Macaulay Library images to determine avian molt strategies: a case study on hummingbirds. *Wilson J. Ornithol.* **134**, 52–65 (2022).
- Field, D. J. et al. Early evolution of modern birds structured by global forest collapse at the end-Cretaceous mass extinction. *Curr. Biol.* **28**, 1825–1831.e2 (2018).
- Pigot, A. L. et al. Macroevolutionary convergence connects morphological form to ecological function in birds. *Nat. Ecol. Evol.* **4**, 230–239 (2020).
- Remsen, J. V. & Robinson, S. K. A classification scheme for foraging behavior of birds in terrestrial habitats. *Stud. avian Biol.* **13**, 144–160 (1990).
- Rosenberg, K. V. Diet selection in Amazonian Antwrens: consequences of substrate specialization. *Auk* **110**, 361–375 (1993).
- Fitzpatrick, J. W. Form, Foraging Behavior, and Adaptive Radiation in the Tyrannidae. *Ornithological Monographs* 447–470 <https://doi.org/10.2307/40168298> (1985).
- Mayr, G. A new specimen of the Early Eocene *Masillacoli* *brevidactylus* and its implications for the evolution of feeding specializations in mousebirds (Coliiformes). *Comptes Rendus Palevol* **14**, 363–370 (2015).
- Mayr, G. Strigiformes (Owls), Coliiformes (Mousebirds), and Cavitaves (Trogon, Rollers, Woodpeckers, and Allies). in *Paleogene Fossil Birds* (ed. Mayr, G.) 197–225 (Springer International Publishing, 2022). https://doi.org/10.1007/978-3-030-87645-6_10.
- Ksepka, D. T. & Clarke, J. A. A new stem parrot from the Green River Formation and the complex evolution of the grasping foot in Pan-Psittaciformes. *J. Vertebrate Paleontol.* **32**, 395–406 (2012).
- Mayr, G. Psittacopasserines: Psittaciformes (Parrots) and Passeriformes (Passerines). in *Paleogene Fossil Birds* (ed. Mayr, G.) 177–195 (Springer International Publishing, 2022). https://doi.org/10.1007/978-3-030-87645-6_9.
- Mayr, G. A partial skeleton of a new species of *Tynskya* Mayr, 2000 (Aves, Messelaturidae) from the London Clay highlights the osteological distinctness of a poorly known early Eocene “owl/parrot mosaic”. *PalZ* **95**, 337–357 (2021).
- Wu, Y. et al. Genomic bases underlying the adaptive radiation of core landbirds. *BMC Ecol. Evol.* **21**, 162 (2021).
- The Evolution of the Primate Hand: Anatomical, Developmental, Functional, and Paleontological Evidence* (Springer, 2016).
- Verendeef, A. et al. Comparative analysis of Meissner's corpuscles in the fingertips of primates. *J. Anat.* **227**, 72–80 (2015).
- Martin, G. R. *The Sensory Ecology of Birds*. (Oxford University Press, 2017).
- Manger, P. R., Elston, G. N. & Pettigrew, J. D. Multiple maps and activity-dependent representational plasticity in the anterior Wulst of the adult barn owl (*Tyto alba*). *Eur. J. Neurosci.* **16**, 743–750 (2002).
- Webster, D. M. S., Rogers, L. J., Pettigrew, J. D. & Steeves, J. D. Origins of descending spinal pathways in prehensile birds: do parrots have a homologue to the corticospinal tract of mammals? *BBE* **36**, 216–226 (1990).
- Wild, J. M. & Williams, M. N. Rostral Wulst in passerine birds. I. Origin, course, and terminations of an avian pyramidal tract. *J. Comp. Neurol.* **416**, 429–450 (2000).
- Medina, L. & Reiner, A. Do birds possess homologues of mammalian primary visual, somatosensory and motor cortices? *Trends Neurosci.* **23**, 1–12 (2000).
- Hopson, J. A. Ecomorphology of avian and nonavian theropod phalangeal proportions: implications for the arboreal versus terrestrial origin of bird flight. in *New Perspectives on the Origin and Early Evolution of Birds: Proceedings of the International Symposium in Honour of John H. Ostrom*. (eds. Gauthier, J. A. & Gall, L. A.) 211–235 (Peabody Museum of Natural History, 2001).
- Cornell University. *Birds of the World* (Cornell Laboratory of Ornithology, 2022).
- Smith, G. The use of the foot in feeding, with especial reference to parrots. *Avic. Mag.* **77**, 93–100 (1971).
- Skutch, A. F. Life history of the keel-billed toucan. *Auk* **88**, 381–396 (1971).
- Collar, N. J. Natural history and conservation biology of the tooth-billed pigeon (*Didunculus strigirostris*): a review. *Pac. Conserv. Biol.* **21**, 186–199 (2015).
- Sustaita, D., Gloumakov, Y., Tsang, L. R. & Dollar, A. M. Behavioral correlates of semi-zygodactyly in Ospreys (*Pandion haliaetus*) based on analysis of internet images. *PeerJ* **7**, e6243 (2019).
- Jetz, W., Thomas, G. H., Joy, J. B., Hartmann, K. & Mooers, A. O. The global diversity of birds in space and time. *Nature* **491**, 444–448 (2012).
- Wickham, H. *ggplot2: Elegant Graphics for Data Analysis* (Springer-Verlag, 2016).
- R Core Team. *R: A Language and Environment for Statistical Computing*. <https://www.gbif.org/tool/81287/r-a-language-and-environment-for-statistical-computing> (2022).
- Toda, Y. et al. Early origin of sweet perception in the songbird radiation. *Science* **373**, 226–231 (2021).
- Kimball, R. T. et al. A phylogenomic supertree of birds. *Diversity* **11**, 109 (2019).

60. Clements, J. F. et al. *Downloadable Checklist | Clements Checklist*. <https://www.birds.cornell.edu/clementschecklist/download> (2022).
61. Mast, A. R. et al. Paraphyly changes understanding of timing and tempo of diversification in subtribe Hakeinae (Proteaceae), a giant Australian plant radiation. *Am. J. Bot.* **102**, 1634–1646 (2015).
62. Fain, M. G., Krajewski, C. & Houde, P. Phylogeny of “core Gruiformes” (Aves: Grues) and resolution of the Limpkin–Sungrebe problem. *Mol. Phylogenet. Evol.* **43**, 515–529 (2007).
63. Moyle, R. G. Phylogenetics of barbets (Aves: Piciformes) based on nuclear and mitochondrial DNA sequence data. *Mol. Phylogenet. Evol.* **30**, 187–200 (2004).
64. Barker, F. K., Burns, K. J., Klicka, J., Lanyon, S. M. & Lovette, I. J. Going to extremes: contrasting rates of Diversification in a recent radiation of New World passerine birds. *Syst. Biol.* **62**, 298–320 (2013).
65. Schliep, K. P. phangorn: phylogenetic analysis in R. *Bioinformatics* **27**, 592–593 (2011).
66. Smith, B. T. et al. Phylogenomic analysis of the parrots of the world distinguishes artifactual from biological sources of gene tree discordance. *Syst. Biol.* <https://doi.org/10.1093/sysbio/syac055> (2022).
67. Boyko, J. D., Beaulieu, J. M., Oliver, J. & Boyko, J. *corHMM 2.1: Generalized Hidden Markov Models. R Package Version 2.8*. <https://rdrr.io/cran/corHMM/> (2022).
68. Marazzi, B. et al. Locating evolutionary precursors on a phylogenetic tree. *Evolution* **66**, 3918–3930 (2012).
69. Ducatez, S., Sol, D., Sayol, F. & Lefebvre, L. Behavioural plasticity is associated with reduced extinction risk in birds. *Nat. Ecol. Evol.* **4**, 788–793 (2020).
70. Wilman, H. et al. EltonTraits 1.0: Species-level foraging attributes of the world’s birds and mammals. *Ecology* **95**, 2027–2027 (2014).
71. Tobias, J. A. & Pigot, A. L. Integrating behaviour and ecology into global biodiversity conservation strategies. *Philos. Trans. R. Soc. B: Biol. Sci.* **374**, 20190012 (2019).
72. Tobias, J. A. et al. AVONET: morphological, ecological and geographical data for all birds. *Ecol. Lett.* **25**, 581–597 (2022).
73. Revell, L. J. phytools: an R package for phylogenetic comparative biology (and other things). *Methods Ecol. Evol.* **3**, 217–223 (2012).
74. Paradis, E. et al. Package ‘ape’. *Anal. Phylogenet. Evol.* **2**, 47 (2019).
75. Pinheiro, J. et al. Package ‘nlme’. *Linear and Nonlinear Mixed Effects Models, Version 3*, <https://cran.r-project.org/web/packages/nlme/index.html> (2017).

Acknowledgements

We would like to thank Dr. Maude Baldwin for editorial comments. We would also like to thank Prof. Taichi Kato for finding many additional media of raptors using their feet. Funding for this work was obtained from The Natural Sciences and Engineering

Research Council of Canada (NSERC) by D.R.W. C.A.-P. received a scholarship from the National Council for Scientific and Technological Development (CNPq), Brazil.

Author contributions

C.G.-I., C.A.-P. and J.B. designed the research. C.G.-I. and C.A.-P. collected and analyzed behavioral data, and prepared figures. J.B. and D.R.W. acquired funding. C.G.I., C.A.-P. and A.N.I. wrote the manuscript with input from all authors.

Competing interests

The authors declare no competing interests.

Additional information

Supplementary information The online version contains supplementary material available at <https://doi.org/10.1038/s42003-023-05151-z>.

Correspondence and requests for materials should be addressed to Cristián Gutiérrez-Ibáñez.

Peer review information *Communications Biology* thanks the anonymous reviewers for their contribution to the peer review of this work. Primary Handling Editors: Luke Grinham and Christina Karlsson Rosenthal. A peer review file is available.

Reprints and permission information is available at <http://www.nature.com/reprints>

Publisher’s note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.



Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit <http://creativecommons.org/licenses/by/4.0/>.

© The Author(s) 2023