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Non-Model Systems in Mammalian Forelimb Evo-Devo

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Abstract

Mammal forelimbs are highly diverse, ranging from the elongated wing of a bat to the stout limb of the mole. The mammal forelimb has been a long-standing system for the study of early developmental patterning, proportional variation, shape change, and the reduction of elements. However, most of this work has been performed in mice, which neglects the wide variation present across mammal forelimbs. This review emphasizes the critical role of non-model systems in limb evo-devo and highlights new emerging models and their potential. We discuss the role of gene networks in limb evolution, and touch on functional analyses that lay the groundwork for further developmental studies. Mammal limb evo-devo is a rich field, and here we aim to synthesize the findings of key recent works and the questions to which they lead.

Keywords

Mammal; Forelimb; Evo-devo; evolution; development; Non-model

Mammals have adapted to a diverse array of habitats including but not limited to aerial, aquatic, arboreal, and subterranean environments. Key to mammals' success is the form and function of their forelimbs, which enables them to explore diverse environments. Like all tetrapods, the mammalian limb is divided into three segments: the stylopod (humerus), zeugopod (radius/ulna), and autopod (carpals, metacarpals, and digits). Though its basic patterning is a relative constant, the mammalian forelimb is highly adaptable within this standard framework. The stylopod, zeugopod, and autopod are present in all mammals, but their forms have diversified to allow species to fly, swim, climb, dig, and more. By reconciling development with evolutionary theory, evo-devo has the potential to reveal the mechanisms by which this diversification occurs, and thus has brought incredible

Author contributions

AOH wrote the manuscript, all other authors contributed edits. Thanks to the rest of the Sears lab for valuable discussions.

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Conflict of interest

The authors declare that they have no conflicts of interest.

advancements to our understanding of how mammals have evolved into disparate and successful forms[1–3].

To date, our knowledge of the mechanisms underpinning mammalian limb development have largely relied on mice due to their substantial benefits as a genetically tractable model organism. However, relying on a single generalist species in this way neglects much of the morphological and functional diversity that characterizes mammals' adaptation to a wide array of environments[4].

In recent years, this lack of diversity has begun to be remedied by further inclusion of non-model organisms [Box 1]. This increase in the number of viable organisms for evo-devo studies vastly expands the scope of our understanding of mammalian limb evolution. In this review, we highlight the contributions and potential of non-model organisms to improve our understanding of mammalian limb evo-devo. We note major recent findings across several mammal groups and suggest future directions for the field based on this increase in diversity and potential of study systems.

Non-Model Organisms in Evo-Devo: Precedent

The evo-devo perspective seeks to resolve the drivers of evolution by uniting the variation of developmental processes studied in the lab with the millions of years of evolution observed in nature and the fossil record[5–8]. This unification of biological scales is aided by the inclusion of a diverse array of model systems, which provide a fuller picture of how developmental processes differ and generate the variation necessary for natural selection. This is not a new goal; the importance of non-model organisms to evo-devo has long been recognized[8–12]. Excitingly, in the past decade, improved access to imaging and next-generation sequencing has finally allowed new models to begin to achieve their potential in evo-devo. In this section, we will explore how new models have been assimilated into evo-devo studies, offering an example of how the difficulties of developing new study systems can be overcome.

Bats:

Bats are an excellent example of how a non-model organism can be integrated into the field of evo-devo. As a diverse, specialized, and highly successful group, they have provided insights into the developmental drivers of limb ratios, membrane formation, sensory adaptations, and cranial evolution[13–20]. In the context of limb evolution, their elongated limbs and wing membranes provide a starting point to answer many questions about how limb ratios change and unique morphologies develop to allow colonization of new niches[20–22].

Recent work on bat limbs has improved our understanding of the origins and evolution of mammalian flight and highlights the important contributions of studies of modern and fossil anatomy to evo-devo. A study of the timing of ossification (bone formation) in developing bats found that the autopod and sternum ossify very late[23]. These regions are considered to be highly adaptable in bats, perhaps, in part, due to this delay in ossification. This may

represent one mechanism by which development can shape evolution; elements that ossify late during development may have a higher tendency to vary.

Fossil evidence has shown that the earliest bats likely had more limited flight capabilities than modern bats. The aerodynamic variables of a primitive fossil bat show that early chiropterans may have only relied on flight to commute, rather than hunt the way many modern bats do[22,24–26]. The evolution of flight style is thus an excellent target of further research, as the aerodynamic properties of bat wings widely vary, and this variation may be influenced by the autopod's delayed ossification[26]. How this may relate to regulatory changes that drive limb elongation has yet to be explored[17,20]. Further investigations of developmental timing and its evolutionary consequences may benefit from a reconstruction of the ancestral ossification sequence of bats or a direct comparison of ossification sequences between flight styles.

Ungulates:

The autopod of ungulates represents varying levels of digit loss, and thereby can be used as a tool for improving our understanding of evolution by the reduction of elements. Studies of ungulates have revealed that multiple mechanisms have evolved to reduce digit number, affecting both early and late developmental processes [Fig. 1][27]. For example, in horses and camels, digits are lost after initial patterning occurs due to expanded regions of cell death. In pigs and cows, however, changes in gene regulation of *Ptch1* alter early patterning, causing digit reduction by a wholly different mechanism[27,28].

Recent work has relied on embryos, fossils, and functional analyses to elucidate the extent and cause of digit reduction in horses. An analysis of embryonic material has found that horses go through a stage with five discrete digit condensations. Thus, even the most extreme case of digit reduction in ungulates shows evidence of a pentadactyl state[29]. This study also found that splint metapodials of horses are formed not only from the adjacent metapodials, but rather the fusion of both metapodials on each side[29]. Evidence for this has also been found in fossil horses, in which the splint metacarpals are clearly composed of two fused bones[30].

Analyses of physical stress and internal geometry performed on fossil horse limbs reveal a potential functional driver of digit loss. As body mass increases, the side digits of fossil horses fail to accommodate for increasing loads, and instead the middle digit must expand in size and alter its internal structure. Thus, increasing body size may have facilitated the loss of digits, as a single digit can functionally outperform multiple digits with the same total bone mass[31].

These new findings lead us to ask if all digit reduction mechanisms constitute a similar response to changes in body mass and show similar levels of incomplete digit loss, or if the functional pressures and developmental mechanisms of reduction are unrelated.

As exemplified by bats and ungulates, non-model organisms provide depth and clarity to evo-devo. Further diversification of study species thus can provide an increasingly holistic understanding of limb evolution.

Non-Model Organisms in Evo-Devo: New Systems

In the past few years, a wide array of species has been more thoroughly investigated, providing the foundation from which we can better understand the mechanisms underlying the diversification and specialization of mammal limbs.

Monotremes:

As an outgroup to placental and marsupial mammals, monotremes may provide insights into the ancestral mammalian limb condition. A pair of recent descriptive studies of echidna musculature and skeletal anatomy have revealed that the architectural diversity and range of motion in echidna limbs are highly reduced, potentially representing a conservative phenotype[32,33]. This conflicts with our understanding of monotreme limb development based on ossification sequences, which is highly derived in that the stylopod ossifies later than the zeugopod, a trait shared only by some moles[34]. Thus, monotremes offer a unique system to study mammalian limb evolution as they represent both an outgroup to most mammals and an instance of highly derived limb development. Further investigations of monotremes and comparisons with further outgroups may reveal how the elaboration of early developmental mechanisms has generated the diversity present today.

Moles:

Moles are of particular interest due to their extremely unique forelimb anatomy linked to their fossorial lifestyle[35,36]. Recent work has delved into the function of mole forelimbs, revealing how their unique stylopod and autopod function while walking and digging[37,38]. X-ray imaging has revealed that, while walking, moles rely heavily on their sesamoid false digit, allowing most true digits to be reserved for digging. [38]. This improved functional understanding informs studies of the autopod's development. The development of the sesamoid false digit has been attributed to a heterochronic shift in *Sox9* expression, but the extent to which this influences the rest of the autopod or other limb elements remains to be explored[39]. How, for example, might the development of the false digit and the functional consequences of reserving other digits for digging contribute to the robust nature of other autopod bones?

When burrowing, movement varies only in the distal portions of the limb when different behaviors are employed[37]. This may suggest that the stylopod is more conserved while the zeugopod is more variable. A developmental study of the mole stylopod noted that the abnormal morphology of the humerus is established when the cartilaginous scaffold is just forming. Later in development the humerus is twisted, contributing to the moles' unique cranially angled limb posture[38,40]. These findings suggest that, contrary to the expectations stated above, early limb development can be highly modified, but the mechanisms underlying this have not been investigated. Further investigations of the development and variation of these structures and the range of would help clarify our understanding of how the limb evolves.

Thus far, a limited amount of work has been done on mole development, but with newfound understanding of limb function in this group, there is greater incentive to investigate the development of their false digits, joint morphology, and limb shape[39–43].

Xenarthra:

Xenarthrans exhibit great extant and extinct ecomorphological diversity, making them particularly interesting to the study of evolution. An analysis of fossil and modern xenarthran humeri found that phylogenetically distant arboreal species converged on the same limb morphology. This was true for fossorial species as well. Furthermore, the evolutionary rates of fossorial humeri were much lower than those of arboreal species, suggesting that some underlying processes may bias the evolution of ecologically linked morphologies[44] This, coupled with a recent analysis of cortical compactness in slow arboreal mammals (including xenarthrans) and work studying bone microstructure, may lend insight into how skeletal architecture affects and is affected by development and evolution[44,45]. For example, an analysis of long bone structure via CT scanning of embryonic xenarthrans could reveal the timing of developmental processes that may underlie their unique limb morphologies.

By more thoroughly investigating non-model systems, we can begin to reveal the processes by which limb elements are elongated, reduced, deformed, and restructured [Fig. 1]. These mechanisms generate the variation that natural selection acts upon, allowing us to better understand the process of evolution.

Non-Model Organisms in Evo-Devo: Comparative Studies and Generating Variation

While recent work has improved the diversity of our study systems, there have been several advancements in our understanding of the generation of diversity through comparative development. This work has relied on traditional models but is steadily expanding to include other groups as well.

An analysis of transcriptomic data from opossum, mouse, pig, and bat limbs has shown that variation in gene expression levels increases between species as development proceeds[46]. This trend leads to further questions about how variation arises in development and what mechanisms generate it. The study of gene regulation offers a deeper look into these processes and has been applied across several systems[47–49].

Gene regulation and the networks of interactions that underlie it are important factors in developmental evolution[50–52]. Previous work in non-model systems has found that changes in regulatory networks and elements facilitate the elongation of limbs in bats and the retention of the interdigital membrane [Fig. 1][17,53,54]. Additionally, regulatory changes are thought to facilitate the reduction of digits in some ungulates, as the evolution of expression patterns of key genes that are conserved elsewhere seems to cause digit loss during autopod development[27,28,55]. In a comparison of mouse and pig limbs, significant rewiring of regulatory networks in the pig autopod were found, leading to differences in the expression of genes important to the development of the limb bud's

anterior-posterior axis[56]. Thus, network rewiring may be one mechanism driving digit reduction in artiodactyls. Further comparisons of regulatory networks within different species will advance our understanding of how regulatory changes facilitate the development and evolution of the limb.

Other work in mice has revealed how changes to development can overcome physical constraints on diversity. The growth plate, which facilitates bone elongation, has been the focus of many recent studies, as the proportions of long bones vary widely across mammals[28,57–60]. The secondary ossification center (SOC), located at the epiphyses of long bones of mammals, is closely associated with the growth plate, yet only recently has it been considered in an evo-devo context. A thorough study combining CT scanning, immuno-histochemistry, and structural measurements of tissues identified the potential evolutionary significance of this structure. The SOC shields the growth plate from physical stress during development, preventing unnecessary cell death and allowing mammal limbs to develop in otherwise intolerable conditions[61]. While this work was only done in mice, it implies that the SOC permits the development of the diverse array of limb morphologies present in mammals. Non-model systems may thus reveal the significance of this structure to the evolution of mammal limbs.

Non-Model Organisms in Evo-Devo: Future Questions

Recent work in non-model organisms and novel techniques tested in mice provide new ways to address various old and emerging questions of mammal limb evo-devo.

Regarding gene regulation, we now have the tools to ask how changes to the wiring of regulatory networks generate differences across a diverse array of species. This has begun to be addressed in bat and pig limbs but can be applied to the variety of models discussed here to understand the regulatory changes that may facilitate the formation of the mole humerus and other unique limb morphologies. Following the methods of Tissieres et al. (2020), it is now feasible to compare gene network evolution across different species and confirm the phenotypic effects of network rewiring in mice. Xenarthra and moles, for example, have much to offer to our understanding of regulatory evolution in the mammal autopod, as they offer additional and largely uninvestigated cases of variation by digit loss and sesamoid expansion, respectively [Fig. 1].

Recent advancements in our understanding of the function of the mole forelimb and the potential evolutionary consequences of the SOC can guide investigations of the evolutionary role of physical constraints. For example, we may now ask how the unique function and morphology of the mole forelimb influence its development and the development of adjacent structures. Hypotheses on the SOC's evolutionary influence from Xie et al. (2019) would greatly benefit from tests in other species. The presence, development, and morphology of the SOC should be studied in groups that represent diverse long bone phenotypes, like the moles, xenarthrans, ungulates, and monotremes. How, for example, are the unique functional constraints of fossorial or arboreal animals influenced by the SOC, which has only been studied in a general terrestrial species [Fig. 1]? After better understanding the SOC across

mammalian taxa, we can return to the mouse and attempt experimental manipulations to confirm its role as a mediator of physical constraints.

The potential of non-model organisms is vast. The difficulties of working with non-model groups are not negligible, but they can, in many cases, be overcome. More importantly, for a field focused on developmental generation of variation, any inclusion of diversity is a net benefit. In the past two years, the field of evo-devo has expanded the breadth of available models and produced new findings on how developmental processes affect evolution. The unification of these advancements has much to offer to evo-devo, and there is no doubt that these new tools, models, and theories will together advance our conceptualization of the evolution of life.

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Box 1:

Here we refer to non-model organisms as any organism outside of the typically studied species for a clade. Some groups have been more frequently studied in mammals (e.g., bats, opossums, pigs), but relative to mice, they lack the vast majority of tools needed to perform deep developmental studies, so we refer to them as non-model organisms as well.

Working with Non-Model Organisms:

A key challenge to the use of non-model organisms is the procurement of specimens. Bats, for example, are not easily maintained in lab conditions, and harvesting embryos from any lab-raised colonies can be highly inefficient. Ungulates are also logistically difficult to maintain as a lab animal. Other groups, like the monotremes and xenarthrans, are far too rare to generate large quantities of embryos. There are several ways this is overcome. Developmental studies of bats typically rely on field-caught specimens, as many species are substantially abundant that embryos can be sustainably collected. Field collection is also the source of pregnant moles used in developmental studies[39,40,42]. While ungulates are not easily maintained in a lab, their embryos can be sourced from several places, including farms, breeders, and museums [30,62]. The few developmental studies of monotremes rely on these types of sources as well.

Benefits of Museum Specimens:

Museum specimens can offer enormous amounts of data on species that are hard to access. One stellar example of this is a recent study that coordinated with museums around the world to CT scan a developmental series of the extinct thylacine, showing that even the rarest samples can be studied non-destructively[63]. Recent work on echidnas capitalized on historical thin sections supplemented by a breeding colony to reveal how the middle ear and jaw may have functioned during the transition from the ancestral state, demonstrating the power of studying unique mammals[64,65]. Similarly, studies of xenarthrans can and have relied on museum specimens for developmental analyses[66].

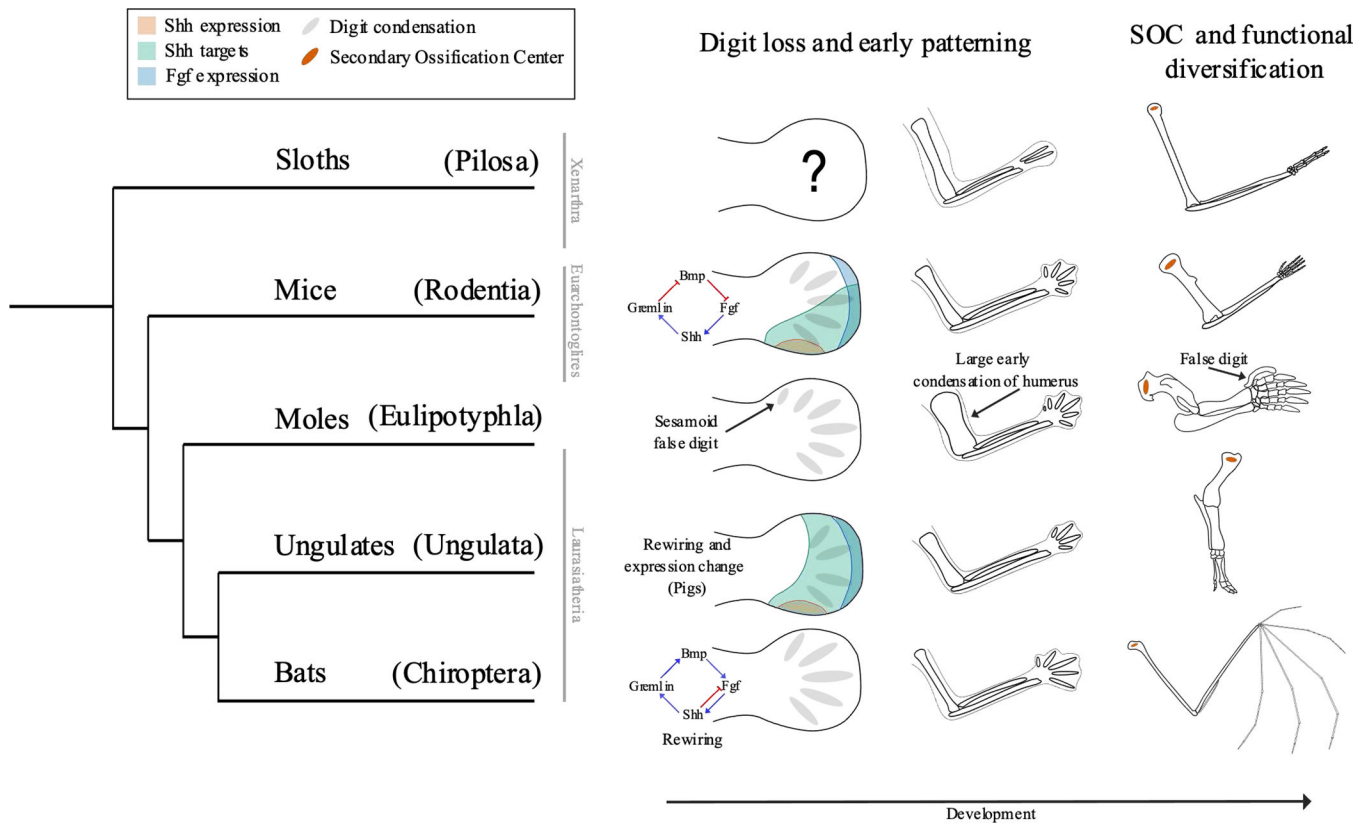


Figure 1: Diversity of Mammal Limb Development -

Schematic of limb development for sloths, mice, moles, ungulates, and bats through developmental time. Changes in developmental networks and expression patterns are indicated where known. Gray ovals in the hand plate and autopod represent early digit condensations. Orange ovals in the humerus represent secondary ossification centers. *Bmp* — Bone morphogenic protein; *Fgf* — Fibroblast growth factor; *Shh* — Sonic hedgehog. Expression patterns and networks from [56,17], respectively. Phylogeny based on [67–69].