The same but different: setal arrays of anoles and geckos indicate alternative approaches to achieving similar adhesive effectiveness

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Abstract

The functional morphology of squamate fibrillar adhesive systems has been extensively investigated and has indirectly and directly influenced the design of synthetic counterparts. Not surprisingly, the structure and geometry of exemplar fibrils (setae) have been the subject of the bulk of the attention in such research, although variation in setal morphology along the length of subdigital adhesive pads has been implicated to be important in the effective functioning of these systems. Adhesive setal field configuration has been described for several geckos, but that of the convergent \textit{Anolis} lizards, comprised of morphologically simpler fibrils, remains largely unexplored. Here, we examine setal morphology along the proximodistal axis of the digits of \textit{Anolis equestris} and compare our findings to those for a model gecko, \textit{Gekko gecko}. Consistent with previous work, we found that the setae of \textit{A. equestris} are generally thinner, shorter, and present at higher densities than those of \textit{G. gecko} and terminate in a single spatulate tip. Contrastingly, the setae of \textit{G. gecko} are hierarchically branched in structure and carry hundreds of spatulate tips. Although the splitting of contacts into multiple smaller tips is predicted to increase the adhesive performance of a fiber compared to an unbranched one, we posited that the adhesive performance of \textit{G. gecko} and \textit{A. equestris} would be relatively similar when the configuration of the setal fields of each was accounted for. We found that, as in geckos, setal morphology of \textit{A. equestris} follows a predictable pattern along the proximodistal axis of the pad, although there are several critical differences in the configuration of the setal fields of these two groups. Most notably, the pattern of variation in setal length of \textit{A. equestris} is effectively opposite to that exhibited by \textit{G. gecko}. This difference in clinal variation mirrors the difference in the direction in which the setal fields of anoles and geckos are peeled from the substrate, consistent with the hypothesis that biomechanical factors are the chief determinants of these patterns of variation. Future empirical work, however, is needed to validate this. Our findings set the stage for future comparative studies investigating the functional morphology of these convergent adhesive apparatuses. Such investigations will lead to an enhanced...
INTRODUCTION

The remarkable ability of gecko digits to effectively adhere to a variety of surfaces using intermolecular forces has been of considerable multidisciplinary interest over the past two decades, prompting the generation of bio-inspired synthetic adhesives aimed at replicating their functional properties. Most species of gecko examined thus far, however, possess rather complex manifestations of the epidermal filaments that induce adhesion, perhaps adding substantial challenges to the biomimetic process. In light of this, several authors have noted that Anolis, a genus of lizards with convergently evolved fibrillar adhesive arrays, provides an alternative model system for the study of fibrillar adhesion because of its structurally simpler adhesive fibrils (Garner et al., 2019; Russell and Garner, submitted; Russell and Garner, in preparation). Gecko subdigital pads are composed of expanded scales (scansors; Figure 1a) that carry arrays of structurally hierarchical fibrils (setae) terminating in hundreds of small spatulate tips (spatulae). Anoline subdigital pads are also composed of expanded scales (lamellae; Figure 1b) that carry arrays of setae terminating in but a single large spatula. Exemplar setae of anoles have generally been described as being shorter and thinner than those of geckos (Delannoy, 2005; Maderson, 1970; Peterson, 1983; Ruibal & Ernst, 1965; Williams & Peterson, 1982). Although geckos and anoles differ in the dimensions and geometry of their adhesive setae, how these differences potentially influence adhesive performance at the level of the fiber or whole animal has received little attention.

The dimensions of a spatula, the presence of structural hierarchy, and the density of setae/spatulae are important contributors to the adhesive performance of fibrillar systems. For a single contact (i.e., spatula), larger contacts should induce greater adhesion than smaller ones (Arzt et al., 2003; Autumn et al., 2014; Johnson et al., 1971; Kendall, 1975). The concept of contact splitting predicts that the subdivision of an adhesive area (e.g., a subdigital pad) into multiple smaller contact points (e.g., setae) increases the overall adhesive force (Arzt et al., 2003; see Supporting Information). Some authors have also extended this formalism to predict that branched, hierarchical fibers should benefit further from contact splitting at multiple levels (i.e., the pad into setae and setae into multiple spatulae; Peattie & Full, 2007). Indeed, synthetic hierarchical fibers generate greater adhesion than unbranched fibers despite possessing similar contact areas (Murphy et al., 2009). Subdigital pads with greater setal/spatular density should also generate greater adhesion than those with lower setal/spatular density. Early work examining differences in the morphology of exemplar gekkotan and anoline setae found that geckos and anoles differ in many of these aspects (Peterson, 1983; Ruibal & Ernst, 1965; Stork, 1983; Williams & Peterson, 1982). Do these differences, however, translate into differences in whole animal adhesive performance between geckos and anoles? Several studies have documented anoline adhesive performance (Bloch & Irschick, 2005; Elstrott & Irschick, 2004; Garner et al., 2017; Irschick et al., 2005; Zani, 2000), but only two studies have directly compared static adhesive performance between geckos and anoles and surprisingly found their adhesive systems to be functionally similar on smooth substrates (Irschick et al., 1996; Ruibal & Ernst, 1965). Thus, it is not clear what is responsible for the functional similarity in these convergent (Russell and Garner, submitted), yet morphologically distinct, adhesive systems.

In addition to the structure and geometry of individual setae, the manner in which setae are configured along the proximodistal axis of the subdigital pad impacts the effective functioning of biological fibrillar adhesives. A series of critical studies demonstrated that the morphology of gecko adhesive setae varies predictably along the length of the subdigital pad (Delannoy, 2005; Johnson & Russell, 2009; Russell et al., 2007; Russell & Johnson, 2014). Most notably, setal length increases proximodistally both within scansors and between them, and base diameter decreases proximodistally within scansors. Variation in setal length and diameter can alter the bending stiffness of the fibers, which can directly impact the contact area the fibers are capable of generating (Ge et al., 2007; Greiner et al., 2007, 2009; Johnson & Russell, 2009; Spolenak et al., 2005). The predictable variation in setal length across the subdigital pad

**FIGURE 1** (a) Representative scanning electron micrograph of two scansors from the setal field of *Gekko gecko*, from Delannoy (2005). (b) Representative scanning electron micrograph of a lamella from the setal field of *Anolis aequestris*
spawned several functional hypotheses (Johnson & Russell, 2009): improvement of adhesion on rough substrates, inhibition of interference between adjacent setae during attachment, and permission of simultaneous detachment of the entire battery of setae on each scanner during active distoproximal hyperextension (the subdigital pad peeling mechanism exhibited by some geckos). The subdigital pads of *Anolis*, however, detach from surfaces employing the ancestral proximodistal pattern common to lizards in general (Russell & Bels, 2001). Thus, it is possible that they exhibit different patterns of setal length if such variation is driven by differences in the biomechanics of subdigital pad peeling.

Although setal form and density have been posited to be important contributors to adhesive performance, such parameters have rarely been considered collectively when contemplating the functional similarity of anoline and gekkotan adhesive systems. This is largely because no studies have comprehensively documented the morphology and configuration of anoline setal fields, even though some work has mentioned their general variability and patterning (Peterson, 1983; Ruibal & Ernst, 1965). Exploration of these topics holds promise for enabling morphological and functional comparisons with the gecko adhesive system focused on answering outstanding questions relating to fibrillar adhesion. With this in mind, we explored setal morphology and setal field configuration of *Anolis equestris* (Merrem, 1820) and compared them, in a functional context, with similar data previously recorded for a model gecko, *Gekko gecko* (Linnaeus, 1758) (Delannoy, 2005).

## METHODS

### Animals

Four *Anolis equestris* were obtained from Snakes at Sunset (Miami, FL USA). These were sacrificed via two-stage intraperitoneal injection of tricaine methanesulfonate (MS-222) (Conroy et al., 2009), fixed with 10% neutral-buffered formalin, and stored in 70% ethanol. Prior to euthanasia, anoles were cared for as described in Niewiarowski et al. (2008). All animals were weighed before euthanasia. Snout-vent length (SVL) was measured postmortem. All protocols involving animals were approved by the University of Akron IACUC Protocol #19-07-13 NGC.

### Sample preparation and morphometrics

Digits of *A. equestris* were prepared similarly to the methods described by Johnson and Russell (2009). Digit IV of both the right pes and manus was removed from each specimen and sectioned parasagittally parallel to the phalanges to yield a longitudinal transect of the setal fields for imaging via scanning electron microscopy (SEM). The longitudinal sections containing phalanges were critical point dried and attached to SEM stubs with carbon tape (Figure 2a). Samples were sputter coated (30 seconds) with gold palladium (Denton Vacuum Desk II; Moorestown, NJ USA) and viewed with a high-vacuum field-emission SEM (JEOL 7401 FESEM; JEOL USA Inc., Peabody, MA USA).

ImageJ (National Institutes of Health, Bethesda, MA USA) was used to measure a variety of characteristics of lamellae and setae as seen in longitudinal section: total setal length, length of the setal stalk, setal base diameter, setal apex diameter, setal resting angle, setal density, and lamella length (Figure 2b,c). Setal density was assessed by counting the number of setae along a length of a lamella and calculating an areal estimate of the number of setae; this was then extrapolated to the number of setae per mm². We observed setal spacing to be similar mediolaterally, rendering our estimate of setal density accurate. Lamella length was defined as the length of the portion of the lamella bearing fully spatulate setae, which includes the epidermal free margin.

Three lamellae from each of three regions along the subdigital pad (proximal, intermediate, and distal pad regions) were imaged at ~400× to enable measurement of lamella length (Figure 3a). In total, images of 21 distal, 15 intermediate, and 21 proximal lamellae were obtained. Some lamellae (mostly those positioned at intermediate stations along the digit) were not of sufficient quality for data acquisition. Images were taken at ~2000× along the entire proximodistal axis of these lamellae to enable measurement of setal features. Each lamella was visually subvided into three zones (proximal, intermediate, and distal lamella zones) by splitting the total images for a lamella roughly into thirds (Figure 3a). One image from each of these zones was selected for measurement of setal characters based on suitability for accurate measurement of their features. Subsampling of respective areas along pad regions and lamella zones was undertaken because of the high setal densities and limitations related to sample preparation. In many cases, an individual seta was not positioned correctly or of sufficient quality to accurately obtain all desired measurements. Thus, 5-10 setae were selected from each image on a per character basis (e.g., setal length, base diameter, etc.) for acquisition of morphometric data. Three repeat measurements were made for all dimensions for each seta measured and averaged to obtain accurate estimates. A total of 16,522 individual measurements were made.

### Estimation of effective bending stiffness of anoline setae

Collection of anoline setal morphometric data allowed the effective bending stiffness of anoline setae to be estimated. We found these setae to be noticeably tapered, requiring the modification of the traditional equation used to estimate bending stiffness of cylinders with fixed radius because the moment of inertia varies along the length of the fiber (Caliero et al., 2013). This modification is simply the multiplication of the bending stiffness of a fixed radius cylinder (k) by what we call the tapering ratio (t), which is the ratio of the apex radius (Rₐ) to the base radius (Rₖ). Thus, the effective bending stiffness (kₑ-eff) of an anoline seta can be calculated as

\[
kₑ-eff = k \cdot t = \frac{3\pi Rₐ^4 E}{4L^2} \cdot \frac{Rₖ}{Rₐ}
\]
where $E$ is Young’s modulus of $\beta$-keratin (~1 GPa) and $L$ is the length of the setal stalk. It is evident that more drastically tapered structures ($t \approx 0$) will have a significantly reduced effective bending stiffness. We use the above estimation to examine effective bending stiffness ($k_{\text{eff}}$) of setae along the proximodistal axis of anoline subdigital pads.

### 2.4 Predicted adhesive forces of Anolis equestris and Gekko gecko based on morphological data

Several authors (Arzt et al., 2003; Autumn et al., 2002, 2014; Huber et al., 2005; Spolenak et al., 2005) have modeled spatulae as
hemispheres to enable the employment of the Johnson-Kendall-Roberts (JKR) contact theory of elastic solids (Johnson et al., 1971) to estimate the pull-off force of a single spatula. The critical pull-off force \( F_c \) of a hemisphere in contact with a planar surface can be written as

\[
F_c = \frac{3}{2} \pi R \gamma
\]

(2)

where \( R \) is equivalent to the radius of the contact and \( \gamma \) is equivalent to the work of adhesion. Employing the approximate \( R \) of \( A. \) equestris...
spatulæ (0.375 μm) and a γ typical of van der Waals interactions (50 mJ m⁻²), we calculated the pull-off force of a single A. equestris spatula and multiplied it by the setal/spatular densities measured in this study to generate a predicted adhesive force per unit area for A. equestris subdigital pads. We then utilized similar data collected by Delannoy (2005) to predict adhesive force per unit area for the subdigital pads of Gekko gecko.

2.5 Statistical analyses

Setal morphometric data obtained from one high-magnification image were averaged to obtain mean setal morphometrics for that lamella zone. Mean effective setal bending stiffness \( k_{\text{eff}} \) was calculated based on mean morphometric values.

Prior to robust statistical analysis, the overall mean values of each of the setal morphological characters for each individual were regressed against the body mass and SVL of that individual using ordinary least squares (OLS) regression. There were no significant linear relationships between body size and any of the morphological parameters \((p > 0.05)\), permitting non-size-corrected data to be used in the analyses.

Analyses of variance (ANOVA) were used to compare setal morphometric and \( k_{\text{eff}} \) data as a function of pad region, lamella zone, autopodium (manus or pes), and all possible interactions. Interactions that were not significant after an initial analysis were removed. In the case of significant effects \((p < 0.05)\), post hoc Tukey’s Honest Significant Difference (HSD) tests were performed to determine where particular significant differences occurred. In the case of setal density and \( k_{\text{eff}} \) data were log-transformed to meet the homogeneity of variance and normality assumptions of ANOVA. Two variables, setal base diameter and setal resting angle, violated one of the assumptions of ANOVA and transformations were unsuccessful in alleviating this, requiring the use of nonparametric statistical methods. Variance between lamella zones was heterogeneous for setal base diameter, thus, Welch’s ANOVA was used to determine differences in base diameter as a function of pad region, lamella zone, or autopodium. In the case of significant factors, the Games-Howell nonparametric post hoc test was used to determine which groups were significantly different from one another. The residuals of setal resting angle were not normally distributed, thus, the Wilcoxon rank sum test was used to examine whether setal resting angle varied significantly as a function of pad region, lamella zone, or autopodium. In the case of significant factors, nonparametric pair-wise comparisons using the Wilcoxon method were performed post hoc and alpha values corrected using the sequential Bonferroni method. All statistical tables are included in the Supporting Information for reference.

3 RESULTS

3.1 Setal morphology of Anolis equestris

The setal morphology of A. equestris varies considerably across the subdigital pad. Setae range in length from 6.56 to 29.98 μm (Figure 3b), with base diameters ranging from 241 nm to 1.25 μm (Figure 3c). The setae of A. equestris are markedly tapered, with apex diameters generally being smaller than the corresponding base diameter. Setal apex diameters range from 98 to 603 nm (Figure 3d).

Setal resting angle varies from 28.27 to 130.15° (Figure 3e). Setal density ranges from 5.68 x 10⁵ to 2.20 x 10⁶ setae mm⁻² (Figure 3f). The length of lamellae varies from 93.37 to 270.80 μm (Figure 4).

3.2 Setal field configuration of Anolis equestris

Set length varies significantly as a function of pad region \((DF = 2, F = 15.60, p < 0.0001)\), lamella zone \((DF = 2, F = 25.77, p < 0.0001)\), autopodium \((DF = 1, F = 6.31, p = 0.015)\), and the interaction between lamella zone and autopodium \((DF = 2, F = 3.39, p = 0.042; Figure 3B, SI Tables S1, S5, and S10)\). For both autopodia, setae from the intermediate and distal pad region are significantly longer than those from the proximal pad region \((\text{distal vs. proximal: } p < 0.0001; \text{intermediate vs. proximal: } p = 0.0002)\), but setae of the distal and intermediate pad region are not significantly different in length \((p = 0.962)\). Setae are longest in the intermediate lamella zone \((\text{intermediate vs. distal and intermediate vs. proximal: } p < 0.0001)\), but setae of the distal and proximal zones are not significantly different in length \((p = 0.40)\). Setae of the intermediate lamella zones of the pes are significantly longer than those of the manus \((p = 0.014; Figure 3B)\). Trends in setal length among pad regions are identical for the manus and pes. Trends in setal length between lamella zones are similar for the manus and pes, although setae from the distal lamella zone are not significantly different in length than setae from the intermediate lamella zone in the manus \((p = 0.151)\).

Welch’s ANOVA revealed highly significant variation in setal base diameter depending on lamella zone \((DF = 2, F = 23.02, p < 0.0001)\) and marginally non-significant variation across pad regions \((DF = 2, F = 3.31, p = 0.05; Figure 3C, SI Tables S1, S6, and S11)\). There were no significant differences in setal base diameter as a function of autopodium \((p = 0.92)\). We observed significant decreases in setal base diameter proximodistally across lamella zones (all pair-wise comparisons \(p < 0.05\)), and a general trend of decreasing setal base diameter proximodistally along pad regions.

Setal apex diameter varies significantly as a function of autopodium \((DF = 1, F = 22.47, p < 0.0001)\), pad region \((DF = 2, F = 8.13, p = 0.0009)\), lamella zone \((DF = 2, F = 7.27, p = 0.0018)\), and the interaction between pad region and lamella zone \((DF = 4, F = 3.29, p = 0.018; Figure 3D, SI Tables S1, S5, and S12)\). Setal apex diameter is significantly greater for setae of the pes than those of the manus \((p < 0.0001)\). Setal apex diameter initially appears to decrease proximodistally between pad regions and lamella zones, but examination of the significant interaction term revealed that this was not the case. Setal apex diameter is significantly smaller in distal pad regions compared to proximal ones, but only for the proximal lamella zone \((p = 0.0003)\). There are no other significant differences in setal apex diameter between pad regions. In the proximal pad region, setal apex diameter is significantly smaller in the distal lamella zones compared to the proximal lamella zones \((p = 0.0003)\). Setae of the intermediate lamella zones of the proximal pad region do
not, however, vary significantly in setal apex diameter from the distal ($p = 0.504$) or proximal lamella zones ($p = 0.111$). Setae do not vary significantly in apex diameter along lamella zones in the intermediate and distal pad regions.

A Wilcoxon rank sum test revealed that setal resting angle varies significantly as a function of pad region (Chi-Square = 6.321, DF = 2, $p = 0.042$) and lamella zone (Chi-Square = 20.065, DF = 2, $p < 0.0001$; Figure 3E, SI Tables S2, S8, and S14), but not for equivalent regions of the autopodia ($p = 0.16$). Resting angle is significantly lower in the distal pad region than the proximal pad region ($p = 0.01$), yet setae of the intermediate pad region do not differ significantly in resting angle when compared with the proximal or distal pad regions (intermediate vs. proximal: $p = 0.248$, intermediate vs. distal: $p = 0.369$). Setal resting angle decreases significantly proximodistally among lamella zones (all pair-wise comparisons $P < $ Bonferonni adjusted alpha $\alpha$).

ANOVA revealed that setal density does not vary significantly as a function of pad region, lamella zone, autopodium, or any of their interactions (whole model: DF = 5, F = 5.016, $p = 0.11$; Figure 3F and SI Table S3).

Lamella length varies significantly along pad regions and between the manus and pes ($p = 0.015$ and $0.0009$, respectively; Figure 4, SI Tables S4, S9, and S15). Lamellae of the distal pad region are significantly shorter than those of the proximal and intermediate pad region (distal vs. proximal: $p = 0.041$; distal vs. intermediate: $p = 0.022$), whereas the length of lamellae of the intermediate pad region is not significantly different from that of lamellae of the proximal pad region ($p = 0.985$). Lamellae of the pes are significantly longer than those of the equivalent regions of the manus ($p = 0.003$).

Effective setal bending stiffness ($k_{eff}$) varies significantly as a function of pad region (DF = 2, F = 23.07, $p < 0.0001$) and lamella zone (DF = 2, F = 50.65, $p < 0.0001$; Figure 5, SI Tables S2, S7, and S13). Setal $k_{eff}$ does not differ significantly between autopodia (DF = 1, F = 0.6, $p = 0.44$). Across pad regions, setal $k_{eff}$ becomes significantly smaller proximodistally (all pair-wise comparisons $p < 0.05$). Setal $k_{eff}$ of distal and intermediate lamella zones is significantly less than that of the proximal lamella zones (both comparisons $p < 0.0001$), whereas setal $k_{eff}$ of distal and intermediate lamella zones are not significantly different ($p = 0.0632$).

3.3 Predicted adhesive forces of Anolis equestris and Gekko gecko based on morphological data

Using the JKR contact theory of elastic solids, we estimated that a single spatula of Anolis equestris with a radius of 0.375 μm is capable of generating 0.09 μN of pull-off force, assuming a work of adhesion ($\gamma$) typical of van der Waals interactions (50 mJ m$^{-2}$). When multiplied by the range of setal densities observed in this study, this results in 0.09-0.14 N mm$^{-2}$ of normalized adhesive force per unit area. Using the range of spatular diameters and densities measured by Delannoy (2005), Gekko gecko is estimated to generate between 0.05 and 0.25 N mm$^{-2}$ of normalized adhesive force per unit area.
4 | DISCUSSION

4.1 | Setal field configuration of Anolis equestris – an overview

Many setal morphological features of A. equestris vary predictably along the proximodistal axis of the subdigital pad (Figure 6). The general trends are as follows. Overall, setal length increases proximodistally along pad regions, but is maximal in intermediate lamella zones. Setal base diameter and setal resting angle decrease proximodistally along pad regions and lamella zones. Lamella length decreases markedly proximodistally along the subdigital pad. As a consequence of variation in setal length and setal base diameter, effective setal bending stiffness ($k_{eff}$) decreases significantly along the proximodistal axis of both the subdigital pad and lamella zones, indicating that setae of the distal regions of the subdigital pad and lamellae are considerably more flexible than those situated more proximally. Setal length of intermediate lamella zones is significantly greater for the pes than the manus but there are no significant differences between the effective bending stiffness ($k_{eff}$) of the setae of the manus and pes.

4.2 | Comparative morphology of Anolis equestris and Gekko gecko setae and its functional implications

With regard to fibrillar adhesion, the Tokay gecko (Gekko gecko) is arguably the most frequently studied gekkotan. This makes it an excellent candidate for morphological comparison with a similarly sized species of Anolis. Comparing data for a representative individual of Gekko gecko (Delannoy, 2005) and Anolis from our present study, the setae of A. equestris are 3-5 times shorter and 2-7 times narrower at
their base than those of G. gecko (Figure 7a,b). Setal resting angle in A. equestris ranges from being similar to that of G. gecko to nearly twice as great (Figure 7c) (Delannoy, 2005), and setal density of A. equestris is between 40 and 88 times greater than that for G. gecko (Figure 7d) (Delannoy, 2005). Although we had difficulty reliably obtaining images of the spatulae of A. equestris along the proximodistal axis of the subdigital pad, they appear to be approximately 0.75 μm wide, and are thus likely between 3 and 9 times greater in width than those of G. gecko (Delannoy, 2005).

Although both geckos and anoles have greater tip densities than other organisms employing fibrillar adhesion (Arzt et al., 2003; Garner et al., 2019; Ruibal & Ernst, 1965; Williams & Peterson, 1982), the setae of G. gecko and A. equestris differ in terms of their overall morphology. A single G. gecko seta terminates in hundreds of spatulae (Delannoy, 2005), whereas that of A. equestris carries only one. Thus, the setae of G. gecko exhibit contact splitting, consistent with the idea that the subdivision of a single fiber should increase its adhesive force potential. If the only difference between the setae of geckos and anoles was the presence of structural hierarchy, the magnitude of gecko adhesion would certainly be greater than that of anoles, as has been demonstrated for synthetic fibrillar systems (Murphy et al., 2009). The data collected here, however, demonstrate that gecko and anole setae not only vary in their hierarchical structure but also in their density, setal size, and spatula width. The predicted adhesive force of A. equestris (0.09-0.14 N mm⁻²) falls well within the range of that estimated for G. gecko (0.05-0.25 N mm⁻²).

Thus, our theoretical calculations indicate that the benefit of structural hierarchy exhibited by individual G. gecko setae does not translate into overall greater adhesive performance compared to A. equestris because of the greater density of setae present on the

FIGURE 7  Setal length (a), setal base diameter (b), setal resting angle (c), and setal density (d) of A. equestris and G. gecko setae as a function of pad region and lamella/scansor zone. All presented data were obtained from the pes. Gekko gecko data are from Delannoy (2005). Error bars are ±1 standard error.
subdigital pads of *A. equestris* compared to *G. gecko* (Irshick et al., 1996). These findings provide a potential morphological explanation for the similarity in whole-organism adhesive performance between anoles and geckos on smooth substrates (Irshick et al., 1996; Ruibal & Ernst, 1965).

Of course, structural hierarchy has been implied to influence a number of other properties of fibrillar systems, including self-matting, fiber fracture, flaw tolerance, and adhesion to rough substrates (Persson, 2003; Spolenak et al., 2005; Yao & Gao, 2006). Thus, any functional consequences of the absence of structural hierarchy of anoline setae may become more apparent in other contexts. Anecdotal observations also suggest functional disparities between geckos possessing complex manifestations of the adhesive apparatus, such as *Gekko gecko* (Russell & Gamble, 2019), and anoles during adhesive locomotion on vertical or inverted, low friction substrates (A.M. Garner and A.P. Russell, personal observations). Such disparities are likely not a function of the individual fibrillar structure and function, but rather how entire batteries of setae are deployed and controlled using higher-order anatomy (e.g., musculotendinous systems) and physiology (e.g., neurological control). Future interdisciplinary comparative investigations regarding gekkotan and anoline attachment in both static and dynamic contexts are clearly necessary to further explore these topics.

### 4.3 Setal field configuration in anoline and gekkonid adhesive apparatuses – functional implications seen through a comparative lens

Historically, most authors have focused on the geometry and characteristics of a single exemplar seta and how this, by extrapolation, relates to the function of the entirety of the adhesive apparatuses of squamates. However, setal variation along the subdigital pad and individual lamellae also has the potential to influence the performance and functional properties of fibrillar adhesive systems (Johnson & Russell, 2009; Russell & Johnson, 2007, 2014). In light of this, we turn from consideration of comparisons of setal morphology of *A. equestris* and *G. gecko* to comparisons of the configuration of their setal fields.

Although having some similarities, the setal field configuration of *A. equestris* and *G. gecko* is quite different. Most strikingly, the setae of *A. equestris* are longest in the intermediate lamella zone, whereas in *G. gecko*, setae increase progressively in length proximodistally along scansor zones. Examination of histological sections from other species of *Anolis* and SEM micrographs of the lamellae of *A. equestris* (Figure 8) demonstrate that the shorter proximal setae may be covered by the dorsal aspect of the epidermal free margin of the immediately preceding (i.e., more proximal) lamella when the subdigital pad is placed in contact with the substrate. Thus, the setal field exposed to and in contact with the substrate exhibits decrease in setal length from intermediate portions of lamellae to those situated more distally, effectively opposite to the trend observed in *G. gecko*. Trends in setal base diameter are also markedly different. In *A. equestris*, setal base diameter decreases proximodistally along the zones of the lamellae and in succeeding pad regions, whereas in *G. gecko*, setal base diameter increases proximodistally along pad regions. From one scansor zone of *G. gecko* to the next, there is little proximodistal variation in setal base diameter, although it appears to be maximal in the intermediate scansor zone of the proximal pad region.

Variability in the trends of setal length and diameter between geckos and anoles has the potential to influence variation in their material properties along the subdigital pad. For example, the proximodistal increase in *gecko* setal length is thought to reduce bending stiffness along scansors. As such, setae situated more distally on a scansor may be able to achieve greater substrate contact and induce greater adhesive force than those that are more proximally situated (Johnson & Russell, 2009). Although there are
marked differences between trends in setal length and diameter along scansors/lamellae in *G. gecko* and *A. equestris*, trends in the effective bending stiffness ($k_{\text{eff}}$) of the setae of these two species are similar (i.e., there is a general decrease in $k_{\text{eff}}$ proximodistally along each lamella and along the digit as a whole; Figure 9). Furthermore, the high aspect ratio of biological fibrillar adhesives is thought to considerably diminish the effective elastic modulus ($E_{\text{eff}}$) of fibril arrays to that within the range of tacky materials (Autumn et al., 2006). Employing the formalism introduced by Autumn et al. (2006) with slight modifications to account for tapering (see SI for details), we found that the average $E_{\text{eff}}$ of anoline setal arrays is estimated to range between 50.4 kPa and 2.9 MPa along the proximodistal axis of the subdigital pad. Although the dimensions and configuration of anoline setae (length, diameter, resting angle, and density) result in a wide range of $E_{\text{eff}}$, eight of nine regions examined have $E_{\text{eff}}$ less than 2 MPa, which is within the range of tacky materials used for pressure-sensitive adhesives (e.g., Sylgard 184, polydimethylsiloxane Bartlett et al., 2012; Khanafer et al., 2008). As such, the major differences between gecko and anole setal field configuration likely have little, if any, consequences for differential contact and adhesion along the proximodistal axis of the subdigital pad. More complex manifestations of the gekkotan adhesive apparatus like that of *G. gecko*, however, possess higher-order anatomical structures, such as subphalangeal hydrostatic cushions and musculature that permit distoproximal hyperextension of the digits, that all anoles examined thus far lack (Russell, 2017; Russell & Gamble, 2019). Such anatomical differences may influence the overall compliance and geometry of the subdigital adhesive system (King & Crosby, 2015), but it is not clear how these anatomical differences may interact with setal morphology and setal field configuration to impact adhesion.

Setal field configuration also has the potential to impact other properties of fibrillar adhesive systems beyond their material properties. Setal length variation along gecko scansors, for example, has been implicated in permitting effective release of subdigital pads during distoproximal hyperextension (distal-to-proximal subdigital pad peeling exhibited by some geckos) (Johnson & Russell, 2009). Many geckos employ distoproximal peeling of their subdigital pads from the substrate and it has been calculated that the graded variation in setal length along the proximodistal axis of each scansor is consistent with all setae of that single scansor attaining their critical angle of detachment simultaneously, as opposed to successively for each setal row (Johnson & Russell, 2009). *Anolis* lizards, however, peel their digits proximodistally (Russell & Bels, 2001). The patterning of setal length in *Anolis equestris* is virtually opposite that observed in geckos and is thus consistent with the simultaneous release hypothesis, although more empirical work is needed to corroborate this. Among geckos, some genera may employ proximodistal hyperextension, but comprehensive investigations of their locomotor kinematics and setal field configuration are lacking (Russell & Gamble, 2019).

![Figure 9](https://example.com/figure9.png)

**Figure 9**  Effective bending stiffness ($k_{\text{eff}}$) of *A. equestris* and *G. gecko* setae as a function of pad region and lamella/scansor zone. *Gekko gecko* data are from Delannoy (2005).
Furthermore, variation in bending stiffness could also influence setal release; setae that are less stiff should reach their critical detachment angles more easily than stiffer ones. Thus, studies of setal field configuration and subdigital pad biomechanics in broader phylogenetic contexts are critical for further exploration of these topics.

5 | CONCLUSIONS

Anolis lizards have long been considered a model system for the study of adaptive radiation and evolutionary key innovations (Losos, 1994, 2011), yet the microstructure of their adhesive subdigital pads, one of two proposed key innovations (Losos, 2011), have remained relatively understudied. Herein, we report on our comprehensive examination of the setal morphology and setal field configuration of a model crown giant anole, Anolis equestris, and make comparisons with similar data collected for a model gecko, Gekko gecko (Delannoy, 2005). The setae of A. equestris are generally thinner, shorter, and present in higher densities than those of G. gecko. Although contact splitting is predicted to increase the adhesive performance of a branched fiber compared to an unbranched one, we discovered that, by taking into account the configuration of their entire setal fields, the predicted adhesive performance of G. gecko and A. equestris setal fields is relatively similar. Anoles compensate for the lack of branched setae by carrying more setae per unit area. These findings highlight the importance of incorporating morphological variability into functional hypotheses, extrapolations, and analyses. Furthermore, we found that the pattern of variation in setal length is effectively opposite in A. equestris to that of G. gecko. Considering this difference in clinal variation mirrors the difference in the direction in which the setal fields of geckos and anoles are peeled from the substrate, our findings are consistent with the hypothesis that biomechanical factors are the chief determinants of these patterns of variation. Future empirical work, however, is needed to validate this hypothesis. Our work here sets the stage for further comparative studies that explore the functional differences between two convergent squamate adhesive systems and the connections among form, function, and environment as they relate to biological fibrillar adhesive systems.

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CONFLICT OF INTEREST

The authors declare that they have no conflict of interest.

DATA AVAILABILITY STATEMENT

Data used in this study are available in the figshare repository at: https://doi.org/10.6084/m9.figshare.12637019.

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SUPPORTING INFORMATION
Additional supporting information may be found online in the Supporting Information section.

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