



On the evolution of extreme structures: static scaling and the function of sexually selected signals

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ARTICLE INFO

Article history:

Received 28 January 2018

Initial acceptance 26 June 2018

Final acceptance 5 July 2018

MS. number: A18-00080R

Keywords:

animal signal
fossil
scaling
sexual selection

The 'positive allometry hypothesis' predicts that ornaments and weapons of sexual selection will scale steeply when among-individual variation in trait size is compared with variation in overall body size. Intuitive and striking, this idea has been explored in hundreds of contemporary animal species and sparked controversy in palaeobiology over the function of exaggerated structures in dinosaurs and other extinct lineages. Recently, however, challenges to this idea have raised questions regarding the validity of the hypothesis. We address this controversy in two ways. First, we suggest the positive allometry hypothesis be applied only to morphological traits that function as visual signals of individual body size. Second, because steep scaling slopes make traits better signals than other body parts, we propose that tests of the positive allometry hypothesis compare the steepness of the scaling relationships of focal, putative signal traits to those of other body parts in the same organism (rather than to an arbitrary slope of 1). We provide data for a suite of 29 extreme structures and show that steep scaling relationships are common when structures function as signals of relative body size, but not for comparably extreme structures that function in other contexts. We discuss these results in the context of animal signalling and sexual selection, and conclude that patterns of static scaling offer powerful insight into the evolution and function of disproportionately large, or extreme, animal structures. Finally, using data from a ceratopsid dinosaur and a pterosaur, we show that our revised test can be applied to fossil assemblages, making this an exciting and powerful method for gleaning insight into the function of structures in extinct taxa.

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Understanding how morphology scales with body size is one of the most pervasive topics in organismal biology (Gould, 1966, 1974a, 1974b; Dial, Greene, & Irschick, 2008; Huxley, 1932; Schmidt-Nielsen, 1984; Templeton, Greene, & Davis, 2005; Thompson, 1917; Voje, 2016; West & Brown, 2005; West, Brown, & Enquist, 1997). The reason for this is simple – virtually every measurable aspect of an organism scales with body size. Some relationships hold across hundreds of species, spanning multiple orders of magnitude in overall size (e.g. Kleiber's law: Kleiber, 1932; Rubner's surface rule: Rubner, 1883; Von Bertalanffy, 1957; Cope's rule: Stanley, 1973; Rensch's rule: Abouheif & Fairbairn, 1997; Blanckenhorn, Meier, & Teder, 2007; Fairbairn, 1997). Others

account for transformations in shape arising during ontogeny (e.g. brain/body mass: Cock, 1966; Gould, 1974a, 1977; Dyar's law: Dyar, 1890). Here we focus on 'static' allometry, scaling that occurs among individuals of the same age sampled from within populations (Cheverud, 1982; sensu Cock, 1966; Pélabon et al., 2013).

Perhaps the most striking pattern in the study of static scaling is the observation that many extreme products of sexual selection – ornaments of choice and weapons of intrasexual competition – scale steeply with body size (Bonduriansky & Day, 2003; Eberhard, 1998; Egset et al., 2012; Emlen & Allen, 2003; Emlen, 1996; Fromhage & Kokko, 2014; Gould, 1974b; Hongo, 2007; Kelly, 2005; Kodric-Brown, Sibly, & Brown, 2006; Miller & Emlen, 2010; Painting & Holwell, 2013; Shingleton, Frankino, Flatt, Nijhout, & Emlen, 2007; Shingleton, Mirth, & Bates, 2008; Simmons & Tomkins, 1996; Stern & Emlen, 1999; Voje, 2016; Wilkinson, 1993). Specifically, when examined on a log scale, the relationship

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between the size of these structures and body size is greater than one ('positive allometry') (Gould, 1966; Huxley & Teissier, 1936; Kerkhoff & Enquist, 2009; Shingleton & Frankino, 2013; Voje, 2016). These steep scaling relationships cause ornaments and weapons to attain extraordinary proportions in the largest individuals, inspiring descriptions such as 'extreme', 'exaggerated' (Darwin, 1871) and 'bizarre' (Gould, 1974b) (Fig. 1).

Early studies of static scaling often focused on the products of sexual selection, including cervid antlers (Gould, 1973; Huxley, 1932; Thompson, 1917), fiddler crab (*Uca*) chelae (Huxley, 1932) and beetle (Scarabaeidae) horns (Bateson & Brindley, 1892; Paulian, 1935). Since then, hundreds of sexually selected structures have been examined, and the overwhelming majority scale steeply with body size (Emlen, 2008; Emlen & Nijhout, 2000; Knell, Naish, Tomkins, & Hone, 2013b; Kodric-Brown et al., 2006; e.g. Otte & Stayman, 1979; Petrie, 1988, 1992; Voje, 2016). In fact, the link between steep scaling and exaggerated ornaments and weapons is so widespread that many consider the steepness of static allometry indicative of the intensity of sexual selection acting on a structure (e.g. stalk-eyed fly (Diopsidae) eyestalks: Baker & Wilkinson, 2001;

frog (Anura) forelimbs: Schulte-Hostedde, Kuula, Martin, Schank, & Lesbarrères, 2011; earwig forceps: Simmons & Tomkins, 1996), and testing of this 'positive allometry' hypothesis is frequently used to infer a sexual selection function when natural observation is unattainable (e.g. trilobite spines: Knell & Fortey, 2005).

The positive allometry hypothesis has, however, been met with resistance. Bonduriansky (2007) noted that the near universality of this pattern may be an artefact of the structures researchers elect to study. That is, when studies focus on morphological scaling, scientists seek the extremes, so the literature is biased in favour of steep scaling relationships (Emlen & Nijhout, 2000; Emlen, 2008; Kodric-Brown et al., 2006). Some extreme structures known to function as sexually selected ornaments, such as elaborate plumage in birds, do not scale positively with body size (Cuervo & Møller, 2001), nor do many genitalic traits, despite the fact that some experience strong selection for increased size (Bertin & Fairbairn, 2007; Blanckenhorn, Kraushaar, Teuschl, & Reim, 2004; Voje, 2016). Indeed, considering the full range of sexually selected structures, including those that are not extreme in size, reveals that slopes are frequently shallow or negative (Bonduriansky, 2007).

Furthermore, at least a few naturally selected structures, such as long bones in large mammals (Bertram & Biewener, 1990; Christiansen, 1999) and cranial horns in lizards (Bergmann & Berk, 2012), also scale positively with body size (Voje, 2016). Clearly, sexual selection need not lead to the evolution of steep scaling, and other agents of selection, such as locomotion and predator defence, occasionally lead to positive static scaling. Where, then, does this leave the positive allometry hypothesis?

We argue that steep static-scaling relationship slopes can be powerful clues to trait function, particularly when combined with other morphological measures of among-individual variation (e.g. trait-specific coefficients of variation; see below). In this context, we suggest much of the controversy and inconsistency in the literature stems from two sources. First, the positive allometry hypothesis has been applied to all sexually selected structures, when, in fact, the logic holds only for a particular subset: sexually selected signal structures where the size of the structure functions as an honest signal of the body size or resource-holding potential of their bearers. Second, tests of the positive allometry hypothesis often rely on demonstrating a slope significantly greater than one. While rich in historical precedent, this approach fails to incorporate the signalling function of these structures. We propose future studies ask not whether the slope is greater than one, but rather whether the slope is relatively steeper for the focal signal structure than it is for other, more typically proportioned, nonsignal-related body parts. It is the relative increase in slope that allows these structures to function effectively as signals, and appropriate tests should incorporate this into their methods.

We summarize literature on animal signalling to show why positive allometry is likely when structures evolve as signals of body size, and why these structures are predicted to scale more steeply with body size than other, nonsignal structures measured in the same individuals. By the same logic, we explain why other types of extreme structures, such as those used in prey capture or locomotion, should not scale more steeply than other body parts.

We test these predictions by comparing the slopes of a suite of extreme morphological structures (14 signal, 15 nonsignal; Supplementary Table S1) to slopes of more typically proportioned 'reference' structures within the same organism (rather than the traditional comparison to isometry, see below), and show that relatively steep slopes are common for structures that function as sexually selected signals but not for comparably extreme structures that function in other, nonsignalling contexts.

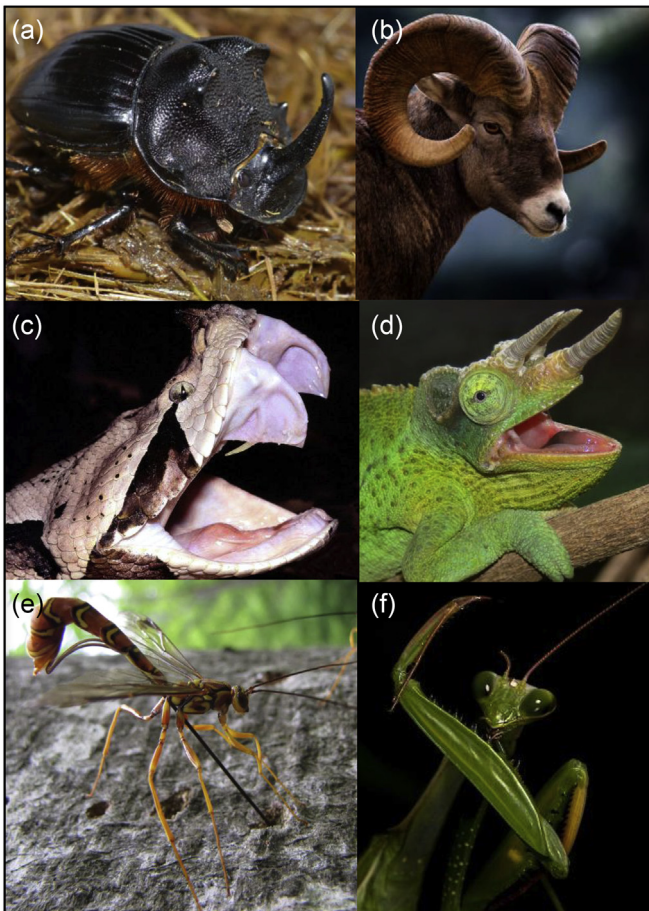


Figure 1. Extreme nonsignal (ns) and sexually selected (ss) signal structures: (a) dung beetle horns (Scarabaeidae, ss; photo credit: Bernard Dupont); (b) bighorn sheep horns (*Ovis canadensis*; ss; photo credit: Jeremy Weber); (c) gaboona viper fangs (*Bitis gabonica*; ns; photo credit: Brimac the 2nd); (d) Jackson's chameleon horns (*Triceros jacksonii*; ss; photo credit: Bengimint444); (e) ichneumon wasp ovipositor (Ichneumonoidea; ns; photo credit: Seney Natural History Association); (f) praying mantis forelimbs (Mantodea; ns; photo credit: Oliver Koemmerling (creativecommons.org/licenses/by-sa/3.0/deed.en)). Photographs licenced under creativecommons.org/licenses/by/2.0/deed.en unless otherwise specified.

METHODS

Specimen/Structure Selection and Morphological Measures

All species with putatively 'extreme' structures – hereafter referred to as 'focal structures' (see [Appendix](#), Identifying Extreme Morphology, for our classification of 'extreme') – and adequate sample size ($N \approx 10$) were surveyed from the Phillip L. Wright Zoological Museum at the University of Montana (MT, U.S.A.), the Museum of Comparative Zoology at Harvard (MA, U.S.A.) and the Emlen Lab Entomological Collection (MT, U.S.A.). Surveying all species that met these criteria allowed for a relatively unbiased sample of both taxa and structure type. However, since most sexually selected structures in insects are beetle horns (reviewed in [Emlen, 2008](#)), the invertebrates surveyed here appear somewhat Coleoptera biased. Six additional data sets were sourced specifically for this analysis: Jackson's chameleons, *Triceros jacksonii*, for the presence of both an extreme signal (horns) and nonsignal (tongue) structure; large bee flies (*Bombylius major*), sabre wasps, *Rhyssa persuasoria*, and peacock moths, *Saturnia pyri*, for the presence of sexually selected nonsignal structures; and ceratopsids (*Protoceratops andrewsi*) and pterosaurs (*Rhamphorhynchus muensteri*) to test the described methods on fossil data sets. Finally, note that while the species/structures surveyed here were unbiased relative to the sampled collections, the collections may have been biased either in taxa or in favour of particularly exaggerated structures. If true, then the results presented here, and their interpretation, may be limited to a particular subset of extreme morphology.

Focal structures of extant species were categorized as 'sexually selected signals' (i.e. structures used by potential mates or competitive rivals as visual signals of the bearer's overall condition/quality; [Andersson, 1994](#); [Bradbury & Vehrencamp, 2011](#)) or as 'nonsignal' structures using relevant behavioural studies from the literature (see [Supplementary Table S1](#)). When literature on the focal species was unavailable, studies in closely related species were used to infer trait function. Bill function in the American pelican, *Pelecanus erythrorhynchos*, was inferred from its sister species, *Pelecanus occidentalis* ([Bels et al., 2012](#); [Kennedy, Taylor, Nádorník, & Spencer, 2013](#); [Orians, 1969](#); [Schreiber, Woolfenden, & Curtsinger, 1975](#)). Lantern function in the Malagasy lantern bug, *Zanna madagascariensis*, was inferred from several other Fulgoridae species with similar head morphology ([Hogue, 1984](#); [Urban & Cryan, 2009](#)). Snout function in the elephant shrew, *Elephantulus fuscus*, was inferred from two species of the same genus with similar rostral morphology, *Elephantulus brachyrhynchus* and *Elephantulus myurus* ([Kingdon, 1974](#); [Kratzing & Woodall, 1988](#)). Horn function in dung beetles (*Sulcophanaeus menelas*, *Phanaeus saphirinus*, *Onthophagus lanista*) was inferred from both a comprehensive review of horn function in beetles ([Eberhard, 1980](#)) and empirical studies of dung beetle mating systems (e.g. [Emlen, Marangelo, Ball, & Cunningham, 2005](#); [Moczek & Emlen, 2000](#)). Hindleg function in frog-legged beetles, *Sagra buqueti*, was inferred from a closely related species with similar leg morphology and mating behaviour ([Katsuki, Yokoi, Funakoshi, & Oota, 2014](#); [O'Brien, Katsuki, & Emlen, 2017](#)). Finally, the function of focal traits in extinct species was inferred from key publications focused on 'bizarre' morphology in the fossil record ([Knell & Sampson, 2011](#); [Knell, Naish, Tomkins, & Hone, 2013a](#); [Hone, Wood, & Knell, 2016](#); but see [Padian & Horner, 2011, 2013, 2014](#)).

Reference structures were then chosen for each species as structures that could be consistently measured across all samples and that lacked an obvious functional connection with the focal structure. These criteria appear adequate in choosing reference structures. However, we recognize the limitation of using a single reference structure and encourage the use of multiple reference

structures per organism in future application of the described methods. Doing so will better capture the scaling relationship of 'typical' (i.e. nonsignal) traits and help mitigate impact of accidentally choosing inappropriate reference structures.

Measures of overall body size were based on established, taxon-specific methods for estimating body size. For species where established estimates of body size were unavailable, we adopted methods from closely related taxa. A summary of study species names, sample sizes, relevant morphological information (e.g. focal structure, reference structure, body size measures) and literature used to establish sexually selected signal/naturally selected nonsignal function is provided in [Supplementary Table S1](#).

Dung beetles (*S. menelas*), earwigs, mantidflies (*Climaciella brunnea*), large bee flies, sabre wasps and wildebeest (*Connochaetes taurinus*) were measured using photographs (including scale bars) and ImageJ 1.50i software (National Institutes of Health, Bethesda, MD, U.S.A.). Dung beetles (*S. menelas*), earwigs, mantidflies, large bee flies and sabre wasps were photographed using a 16.2 megapixel Nikon D5100 DSLR camera mounted on a binocular stereo microscope (Leica S6D) set at a fixed distance. Wildebeest were photographed using a 14.2 megapixel Nikon D3100 DSLR camera set at a fixed distance designated to minimize perspective effects (i.e. approximating orthographic projection). All other extant species were measured using digital callipers.

Measures of ceratopsians (*P. andrewsi*) and pterosaurs (*R. muensteri*) were collected directly using digital callipers, from photographs of specimens including scale bars, or from the literature when appropriate, to maximize the number of available specimens (see [Tables A1, A2](#)).

Statistical Analyses

Statistical analyses were performed in R 3.3.2 ([R Core Team, 2016](#)). Measurements were log transformed and mean-standardized prior to analysis. Ordinary least squares (OLS) regression was used to assess scaling relationship slope ([Kilmer & Rodríguez, 2017](#); [Smith, 2009](#); [Warton, Duursma, Falster, & Taskinen, 2012](#); [Warton, Wright, Falster, & Westoby, 2006](#)). For every species, focal structure size and reference structure size were regressed on body size in separate models. Analyses of covariance (ANCOVA) were then used to compare regression slopes of focal structure size on body size (β_{focal}) to regression slopes of reference structure size on body size ($\beta_{\text{reference}}$) within the same species (i.e. to determine whether or not there was a significant interaction between body size and trait group (focal/reference) in explaining trait size). (Differences in intercept were not analysed, since all data were mean-standardized prior to analysis.) In addition, we collected slope estimates (β_{focal} and $\beta_{\text{reference}}$) from each model and constructed 95% confidence intervals. These 95% confidence intervals were then compared between focal and reference structures within the same species.

We calculated mean β_{focal} for sexually selected signal structures and compared them to mean β_{focal} calculated for nonsignal structures using Welch's *t* test. We also calculated mean $\beta_{\text{reference}}$ for species with sexually selected signal structures and compared them to mean $\beta_{\text{reference}}$ for species with exaggerated nonsignal structures using Welch's *t* test. We constructed 95% confidence intervals around mean $\beta_{\text{reference}}$ for species with sexually selected signal structures and 95% confidence intervals around mean $\beta_{\text{reference}}$ for species with nonsignal structures and compared them. We calculated the difference between β_{focal} and $\beta_{\text{reference}}$ ($\Delta\beta_{\text{focal-reference}}$) for each species. We compared mean $\Delta\beta_{\text{focal-reference}}$ for species with sexually selected signal structures to mean $\Delta\beta_{\text{focal-reference}}$ for species with nonsignal structures using Welch's *t* test. We constructed 95% confidence intervals around mean $\Delta\beta_{\text{focal-reference}}$ for

sexually selected signal structures and mean $\Delta\beta_{\text{focal-reference}}$ for nonsignal structures and compared them.

Coefficients of variation were calculated for every structure. We calculated mean coefficient of variation across all signal structures and compared them to the mean coefficient of variation across all nonsignal structures using 95% confidence intervals and Welch's *t* test.

RESULTS

Results of species-level analyses are summarized in Table S1, including slope estimates (β_{focal} and $\beta_{\text{reference}}$) and adjusted R^2 values for all models, differences between β_{focal} and $\beta_{\text{reference}}$ ($\Delta\beta_{\text{focal-reference}}$), ANCOVA results, 95% confidence intervals surrounding β_{focal} , $\beta_{\text{reference}}$ and $\Delta\beta_{\text{focal-reference}}$, and coefficients of variation. For the majority of species with sexually selected signal structures, β_{focal} was significantly greater than $\beta_{\text{reference}}$ (Table S1, Fig. A1). For two of these species, white-tailed deer, *Odocoileus virginianus*, and wildebeest, β_{focal} was greater than $\beta_{\text{reference}}$, but 95% confidence intervals surrounding these estimates were overlapping and the ANCOVA showed no significant difference between β_{focal} and $\beta_{\text{reference}}$. In pronghorn antelope, *Antilocapra americana*, 95% confidence intervals surrounding β_{focal} and $\beta_{\text{reference}}$ were overlapping, but ANCOVA showed a (slightly) significant difference between β_{focal} and $\beta_{\text{reference}}$. Earwigs, on the other hand, displayed nonoverlapping 95% confidence intervals surrounding β_{focal} and $\beta_{\text{reference}}$, but the ANCOVA showed no significant difference between β_{focal} and $\beta_{\text{reference}}$. For all species with exaggerated nonsignal structures, β_{focal} and $\beta_{\text{reference}}$ were either not significantly different, or $\beta_{\text{reference}}$ was significantly higher than β_{focal} (Table S1, Fig. A2). Unlike extreme sexually selected signal structures, extreme nonsignal structures appear to scale similarly to reference structures within the same organism. Mean slope (β_{focal}) of all exaggerated sexually selected signal structures was greater than the mean slope (β_{focal}) of all nonsignal structures ($t_{13,543} = -3.835$, $P < 0.01$) and 95% confidence intervals were nonoverlapping (95% CI mean β_{focal} for sexually selected signal structures [1.709, 4.56]; 95% CI mean β_{focal} for nonsignal structures [0.374, 0.783]). Mean $\Delta\beta_{\text{focal-reference}}$ for sexually selected signal structures was significantly greater than mean $\Delta\beta_{\text{focal-reference}}$ for nonsignal structures ($t_{14,164} = -4.787$, $P < 0.01$; Fig. A3) and 95% confidence intervals did not overlap (95% CI mean $\Delta\beta_{\text{focal-reference}}$ for sexually selected signal structures [1.072, 3.831]; 95% CI mean $\Delta\beta_{\text{focal-reference}}$ for nonsignal structures [-0.501, 0.078]).

Coefficients of variation were significantly higher for extreme sexually selected signal structures (mean = 15.444, 95% CI [9.325, 21.562]) than for nonsignal structures (mean = 5.351, 95% CI [3.263, 7.438]) ($t_{16,043} = 3.37$, $P < 0.01$; Fig. A4).

DISCUSSION

Within species, sexually selected signal structures scaled steeply with body size (Table S1, Fig. A1). In the majority of sexually selected species surveyed here, the scaling relationship of the signal (β_{focal}) was significantly steeper than that of the reference structure ($\beta_{\text{reference}}$). Surprisingly, this pattern did not hold for white-tailed deer or wildebeest. In these species, β_{focal} was greater than $\beta_{\text{reference}}$, but there was no significant difference between β_{focal} and $\beta_{\text{reference}}$. Similarly, for earwigs, the ANCOVA showed no significant difference between β_{focal} and $\beta_{\text{reference}}$, but β_{focal} was greater than $\beta_{\text{reference}}$ and 95% confidence intervals surrounding these estimates were nonoverlapping (Table S1, Fig. A1). These results may be an artefact of relatively small sample size (e.g. $N < 18$ for white-tailed deer) and/or biased sampling (e.g. hunters favouring largest antlered males in sampled populations), since previous work has shown positive allometry and/or strong

selection for these, and similar, weapons (e.g. Kruuk et al., 2002; Lundrigan, 1996; Melnycky, Weladji, Holand, & Nieminen, 2013; Simmons & Tomkins, 1996). Alternatively, these structures may function strictly as weapons (i.e. tools) of intrasexual competition, not as visual signals of quality. If true, then steep scaling between weapon and body size is not expected (McCullough, Miller, & Emlen, 2016, see below). Overall, our results for sexually selected signal structures are consistent with previous work showing that these types of extreme structures tend to be positively allometric (Bonduriansky & Day, 2003; Emlen, 2008; Green, 1992; Kodric-Brown & Brown, 1984; Kodric-Brown et al., 2006; Petrie, 1988, 1992; Simmons & Tomkins, 1996; Voje, 2016).

Every exaggerated nonsignal structure measured scaled with a slope that was either less than, or not significantly different from, that of the reference structure (Table S1, Fig. A2). In addition, across species, the scaling relationship (β_{focal}) of sexually selected signal structures was significantly steeper than that of nonsignal structures ($t_{11,902} = -3.23$, $P < 0.01$). Even within the same organism, nonsignal structures scaled at a shallower rate than sexually selected signals. In Jackson's chameleon, for example, where both an extreme sexually selected signal, horn length, and an extreme nonsignal prey capture structure, tongue length, were surveyed, horn size scaled at a much steeper rate compared to the reference structure than did tongue size (Table S1, Fig. 2).

Why Signals Should Scale More Steeply Than Other Body Parts

Many studies have considered what makes a good signal (reviewed in Bradbury & Vehrencamp, 2011; Maynard Smith & Harper, 2003; Searcy & Nowicki, 2006). In the context of sexual selection, receivers are often females who use variation in signal expression as a basis for mate choice, or males who use these signals to determine the resource-holding potential (i.e. fighting ability) of rival males (Andersson, 1994; Bradbury & Vehrencamp, 2011; Hardy & Briffa, 2013). In both cases, information encoded in the signal pertains to the overall genetic quality and/or condition of the bearer (reviewed in Neff & Pitcher, 2005).

Although any phenotype could, in principle, be used as a signal (provided it is detectable and variable across individuals), some make more effective signals than others. The best signals are conspicuous – bigger or brighter than other body parts (Bradbury & Vehrencamp, 2011). However, it is not just the structure that must be conspicuous. Variation in the expression of that structure is key to mate and rival assessment, and the more pronounced the differences, the better. For this reason, signal structures are often selected to be more variable in their expression than other, surrounding, nonsignal structures (Alatalo, Höglund, & Lundberg, 1988; Cuervo & Møller, 2001; Emlen, Warren, Johns, Dworkin, & Lavine, 2012; Fitzpatrick, 1997; Petrie, 1992; Pomiankowski & Møller, 1995; Rowe & Houle, 1996; Simmons & Tomkins, 1996; Tazzyman, Iwasa, & Pomiankowski, 2014; Wallace, 1987). Hyper-variability in trait size amplifies associated variation in male quality, making these otherwise subtle differences easier to see (Hasson, 1991; Tazzyman et al., 2014; Wallace, 1987).

Effective signals must also be honest. If poor-quality males can cheat by producing effective signals, then reliability of the signal plummets and receivers should focus on other traits. One form of honesty arises when the growth of signal traits is condition sensitive (Biernaskie, Grafen, & Perry, 2014; Bonduriansky & Day, 2003; Bonduriansky, 2006; Grafen, 1990; Iwasa, Pomiankowski, & Nee, 1991; Johnstone, 1997; Kodric-Brown et al., 2006; Nur & Hasson, 1984; Pomiankowski, 1987; Zeh & Zeh, 1988). Condition-sensitive growth of signal structures may 'capture' genetic or environmental variation underlying overall quality, making these signals virtually impossible to fake (Miller & Moore, 2007; Rowe & Houle, 1996;

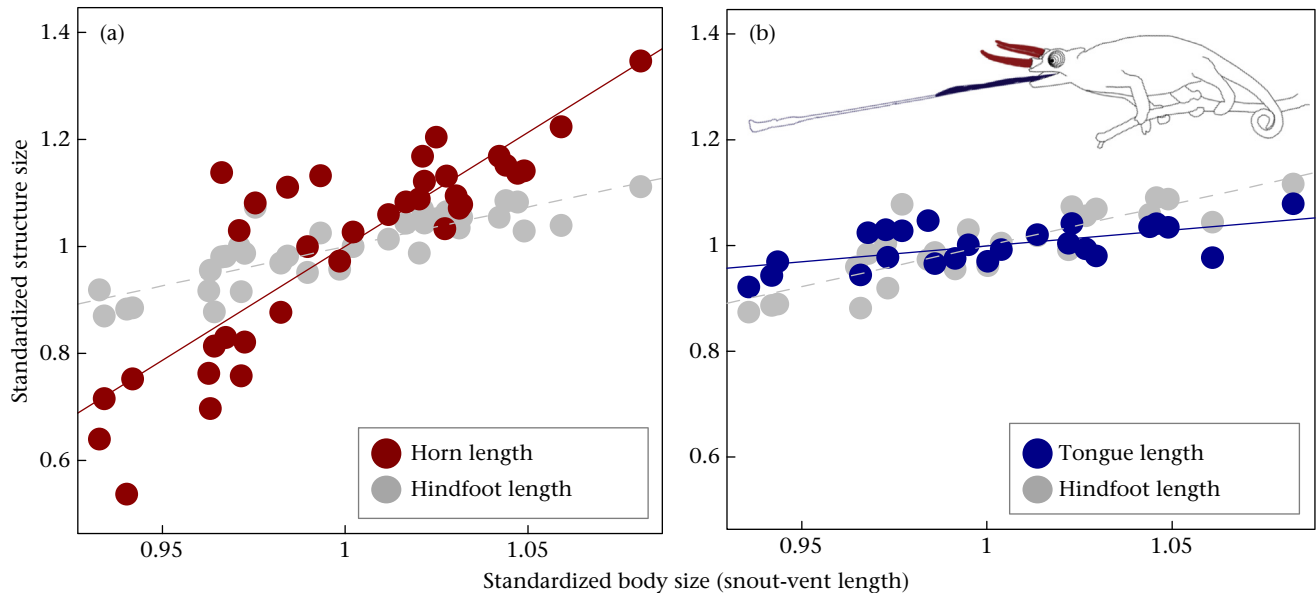


Figure 2. Static scaling relationships for (a) an extreme sexually selected signal structure (horn length, \bullet ; $N = 40$) and (b) an extreme naturally selected nonsignal structure (tongue length, \bullet ; $N = 25$) in Jackson's chameleons, *Triceros jacksonii*. \bullet : focal structures. \bullet : reference structures. Lines represent ordinary least squares regression of standardized log structure size on standardized log body size. In Jackson's chameleon, the extreme sexually selected signal (horn length) scales at a significantly steeper rate than the reference structure (hindfoot length); the extreme nonsignal structure (tongue length) does not; 95% CI for horn length [3.358, 5.159], tongue length [0.251, 0.949] and hindlimb length [1.13, 1.979]. Drawing: D. J. Emlen.

Wilkinson & Taper, 1999). Indeed, sexually selected signal structures are notoriously sensitive to stress, parasite load and nutrition (Cotton, Fowler, & Pomiankowski, 2004; Ezenwa & Jolles, 2008; Gosden & Chenoweth, 2011; Hamilton & Zuk, 1982; Izzo & Tibbetts, 2015; Knell & Simmons, 2010; Kruuk et al., 2002; Skarstein & Folstad, 1996).

Hypervariability through heightened condition sensitivity causes structures to be reliable and informative as signals of quality (Andersson & Iwasa, 1996; Andersson & Simmons, 2006; Andersson, 1994; Bradbury & Vehrencamp, 2011), and these basic characteristics are shared by a wealth of sexually selected signals (reviewed in Bradbury & Vehrencamp, 2011). When information contained in a sexually selected signal involves individual differences in the size of a structure, and when among-individual variation in condition or genetic quality manifests as differences in overall body size, then selection for increasingly effective signals should lead to the evolution of not just higher trait-specific coefficients of variation, but also to a relatively steeper scaling relationship slope (Biernaskie et al., 2014; Green, 1992; Kodric-Brown & Brown, 1984; Kodric-Brown et al., 2006; Petrie, 1988). The steeper the slope, the more variable the focal structure will be relative to surrounding body parts. Mechanistically, when variation in condition is driven by differential access to nutrition, then the evolution of heightened condition-sensitive growth in a particular structure, relative to others, will also manifest as an increase in the steepness of the slope for that structure (Emlen et al., 2012; Lavine, Gotoh, Brent, Dworkin, & Emlen, 2015; Mirth, Frankino, & Shingleton, 2016; Shingleton & Frankino, 2013). Thus, for this particular subset of signal structures, the positive allometry hypothesis should hold. Indeed, the steeper the scaling relationship slope, the better the signal will be, leading to the evolution of larger and larger structures with steeper and steeper patterns of static scaling.

A few exceptions should be noted, however. First, body size is not always correlated with overall genetic quality or condition, as is the case for many fishes (Bolger & Connolly, 1989) and birds (Cuervo & Møller, 2009). In these species, signals are still expected to be condition sensitive and hypervariable. However, because condition is not correlated with body size, differences in the

relative sizes of signal structures may not covary with body size (e.g. Bonduriansky & Day, 2003; Cuervo & Møller, 2009; Fitzpatrick, 1997; Pomfret & Knell, 2006). (This was true for several focal nonsignal traits, and several reference traits surveyed here (indicated by low adjusted R^2 values; Table S1). Indeed, future analyses may benefit from choosing reference structures that more tightly covary with body size.) Similarly, signals that vary in other ways besides size (e.g. colour, behaviour, chemical signals) are also not expected to scale with body size. Finally, sexually selected traits that do not function as signals (e.g. peacock moth antennae, measured here; Table S1, Fig. A2), are not predicted to scale steeper than reference structures, since hypervariation and/or condition sensitivity may actually decrease performance. This includes sexually selected weapons that function only as tools of battle and not as signals of quality, condition or resource-holding potential (McCullough et al., 2016). For these structures, trait expression should be proportional across the entire population, even when selection favours large relative trait sizes. Large structures may display especially high scaling relationship intercepts compared to other traits in the body, but since there is no hypervariation and/or heightened condition sensitivity, the slope should not differ from that of a reference structure. Consequently, we suggest much of the confusion regarding the link between positive allometry and sexual selection can be resolved by recognizing that the positive allometry hypothesis applies only to those structures that act as visual signals of among-individual variation in condition or genetic quality and, in fact, it applies only to a subset of these, signals whose information involves differences in signal size in species where quality is approximated by variation in overall size. For these structures, sexual selection is predicted to drive the evolution of extreme trait size and unusually steep scaling.

Testing the Positive Allometry Hypothesis Against Reference Structures, Rather Than Isometry

We suggest three reasons for testing the positive allometry hypothesis in comparison with reference structures, rather than

with isometry. First, inferring signal function for a structure that scales steeply only makes sense if that structure scales more steeply than other body parts. Steep scaling relationship slopes are relevant because they cause structures to be better signals than other surrounding body parts. The properties that make them effective signals are relative: they are more variable and more condition sensitive in their growth than other body parts. Sexual selection favours receivers who pay attention to these structures because, by doing so, individuals make more informed decisions than they would if they focused on other body parts. Consequently, the pattern that matters for inferring a sexually selected signal function is the difference in slope between the putative signal and other, nonsignal, structures.

Second, detecting hyperallometry in a focal structure without comparing the slope to a control can be misleading. It is possible for nonsignal structures to scale steeply. Indeed, in our sample of nonsignal exaggerated structures, gaboon viper, *Bitis gabonica*, fangs, elephant shrew snouts and mantidfly forelegs all scaled with relatively steep slopes (i.e. $\beta > 1$), but the reference structures were hyperallometric too (Fig. A2, Table S1). Had we focused only on the absolute value of the scaling relationship slope we would have erroneously inferred a signal function for these structures when, in fact, their scaling relationship slopes were no different from those of surrounding body parts. These structures lack the critical properties of an informative signal despite being hyperallometric.

Finally, comparing measured slopes with isometry places undue emphasis on the estimated slope per se. Isometry may be intuitive in principle, but actually detecting it, or rejecting it, depends a lot on the particular landmarks selected, the units of measurement involved and the chosen measure of body size (Bookstein, 1989; Jungers, Falsetti, & Wall, 1995; e.g. Mosimann & James, 1979). For this reason, focusing tests of the positive allometry hypothesis exclusively on rejection of a slope of one may be misleading, especially in the context of interspecific comparisons where landmarks and measures of body size/condition often differ (e.g. Bolger & Connolly, 1989; Jakob, Marshall, & Uetz, 1996; Peig & Green, 2010). Focusing instead on the slopes of focal structures compared to those of reference structures delivers an internally controlled assay for the properties of a structure's expression that matter. Significant increases in the slope of a focal structure relative to other body parts means that the focal structure has the predicted properties of a signal, and we suggest this constitutes evidence in favour of a function for that structure as a sexually selected signal.

Diversity of Exaggerated Morphology

Not all sexually selected structures are signals, but many experience strong selection for increased size. In arthropods with low population density, for example, males search for receptive females and selection can lead to the evolution of elaborate antennae and/or enlarged eyes (e.g. peacock moth antennae, measured here; Table S1). This results in pronounced sexual dimorphism in relative trait size and, in some species, exaggerated male sensory structures (Andersson, 1994; Bertin & Cézilly, 2003; Lefebvre, 2000; Thornhill, 1981). Similarly, antagonistic coevolutionary arms races arising from conflict between males and females can drive rapid evolution of genitalia (Arnqvist & Rowe, 2002, 2005; Brennan, Clark, & Prum, 2009; Parker, 1979; Simmons, 2014). In both contexts, sexual selection drives the evolution of extreme size, but these structures do not function as signals. There is little covariance between trait variation and fitness and, thus, no benefit in traits being hyper-variable or extra condition sensitive. For these traits, steep scaling slopes are not expected (e.g. Eberhard, 1998, 2010; Hosken & Stockley, 2004).

Exaggerated size can also arise through natural selection as, for example, in some locomotor, prey capture and feeding structures (reviewed in Lavine et al., 2015). Appendages such as praying mantis forelimbs and antlion mandibles function like levers, snapping closed to grasp prey. For these species, longer forelimbs or mandibles perform better than shorter ones both because they move faster at their tips and because they sweep through a larger 'kill zone' (Loxton & Nicholls, 1979; Maldonado, Levin, & Pita, 1967). However, like sensory and genitalic structures of sexual selection, large size in these naturally selected structures is not related to a signal function. There is no benefit to hypervariability or heightened condition sensitivity, and steep scaling relationship slopes are not expected.

Here, we provide measures of static allometry for 15 extreme nonsignalling structures (Table S1, Fig. A2). None are sexually dimorphic, and none scaled more steeply than other, typically proportioned, body parts. Jackson's chameleons provide perhaps the best example of all, since males in this species have both types of extreme structure: three horns on the head that function as a signal of competitive ability (Bustard, 1958) and an elongated tongue used to capture prey. Even though the tongue is relatively larger than the horns, tongues scaled with a slope that was shallower than the reference structure. Horns, in contrast, scaled disproportionately steeply (Fig. 2). Clearly, the evolution of extreme structures need not entail relative increases in static allometry slope, and steep slopes, when they occur, can provide valuable clues to a sexually selected signal function.

Inferring Function for Extreme Structures in Extinct Taxa

Unlike most organisms described above, the behaviour of extinct taxa cannot be observed. Even so, lines of evidence can be drawn from static, morphological data to provide testable hypotheses of behaviour (Hone & Faulkes, 2014). For example, hypotheses surrounding mechanical function, such as those involving anchors for musculature or levers that increase moment arms, can be assessed (and potentially rejected) using data from fossils (e.g. Hone, Naish, & Cuthill, 2012; Knell & Fortey, 2005). Similarly, we maintain that the use of static scaling relationship slopes and coefficients of variation may provide a means for inferring a sexually selected signal function for extreme morphology in the fossil record.

Static scaling relationships have been used already to infer function in the fossil record (Gould, 1973; Hone et al., 2016; Knell & Fortey, 2005). However, such inferences remain controversial (e.g. Padian & Horner, 2011, 2013, 2014; Hone & Mallon, 2017; Knell & Sampson, 2011; Knell et al., 2013a; Mallon, 2017). One issue is that collecting multiple individuals from the same fossil locality and horizon (i.e. a single population) is difficult. Sample sizes are often small or gathered from animals separated in space and/or time, and animals are rarely sexed (e.g. Hone & Mallon, 2017). As a result, detection of even fundamental patterns in morphology, such as sexual dimorphism, remains elusive (Mallon, 2017; but see Sengupta, Ezcurra, & Bandyopadhyay, 2017). Another issue is that distinguishing between different signal functions is often difficult. Social dominance and sexually selected signals, for example, are often confluent, and distinguishing between them is complex. In addition, the co-option of extreme structures to multiple functions, thereby exposing them to multiple patterns of selection, may further confound these data (e.g. dugong, *Dugong dugon*, tusks; Anderson, 1979; Domning & Beatty, 2007).

Despite these limitations, we suggest behaviour can be inferred from the fossil record using the methods and logic described above. We predict that when focal structures act as signals of overall body size, both the slope of the static scaling relationship and the coefficient of variation will be steeper/greater in the putative signal structure than in reference structures used as controls. As 'proof of

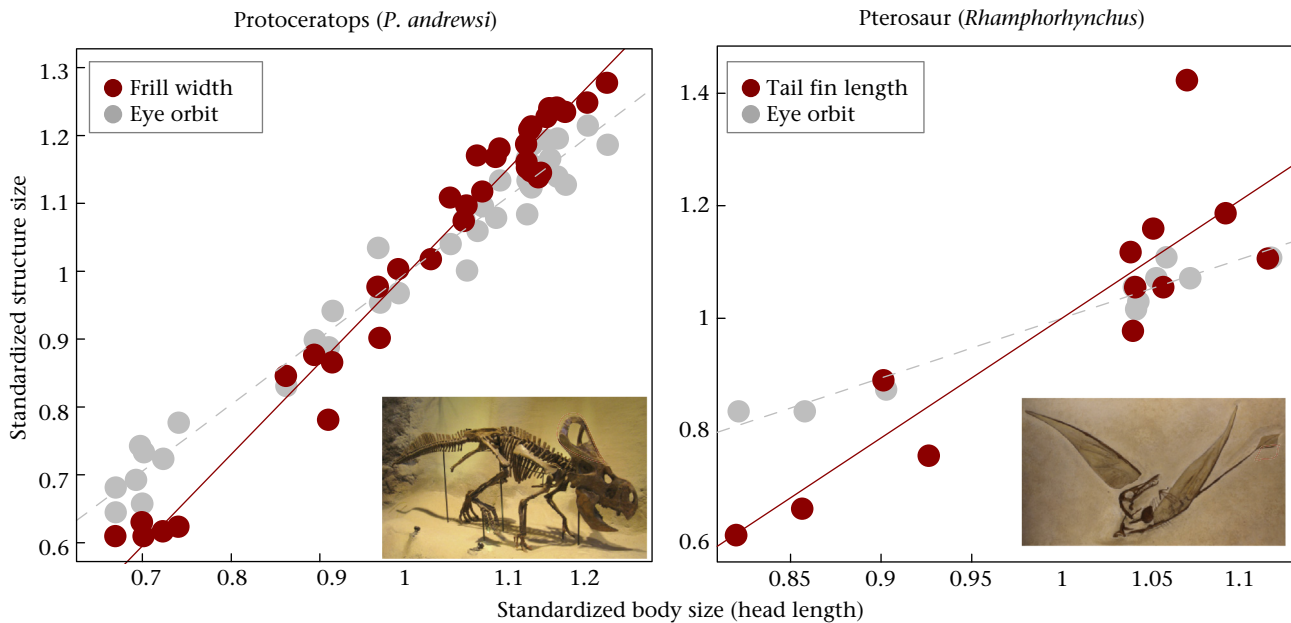


Figure 3. Static scaling relationships for extreme putative sexually selected signal structures in ceratopsians (*Protoceratops andrewsi*, $N = 38$; photo credit: FunkMonk) and pterosaurs (*Rhamphorhynchus muensteri*, $N = 10$; photo credit: M0tty (gnu.org/licenses/fdl-1.3.en.html; creativecommons.org/licenses/by-sa/3.0/deed.en)). ●: putative signal structures. ●: reference structures. Lines represent the ordinary least squares regression of standardized log structure size on standardized log body size. In both species, the scaling relationship of the putative signal trait is steeper than that of the reference trait (*P. andrewsi*: 95% CI for slope of focal structure [1.173, 1.353]; 95% CI for slope of reference structure [0.925, 1.039]; *R. muensteri*: 95% CI for slope of focal structure [1.332, 2.930]; 95% CI for slope of reference structure [0.871, 1.262]), consistent with a history of selection for a hypervariable sexually selected signal. Inlaid photographs display study species with focal structures highlighted in red.

concept' for this approach, we included two putative sexually selected signal structures from the fossil record in our analyses, the enlarged cephalic frill of the ceratopsian dinosaur *P. andrewsi* (adapted and expanded from Hone et al., 2016), and the tail vane of the pterosaur *Rhamphorhynchus*. In both cases, the focal structure scaled more steeply with body size and had a higher coefficient of variation than reference structures measured in the same individual (Fig. 3, Table S1), implying a signalling function.

Overall, we believe this method useful for inferring extreme structure function in the fossil record (perhaps even more useful when analysed in conjunction with other patterns in morphology; e.g. changes in complexity during ontogeny, high variation in trait shape and size between species lineages). Both morphological scaling relationships and coefficients of variation can be reliably measured in fossil specimens, even when sample size is small. We recommend the use of these methods in subsequent analyses of extreme or 'bizarre' morphology in the fossil record, and are hopeful that they might provide insight into the ongoing debate regarding sexual selection in nonavian dinosaurs.

Overall, we suggest that when applied specifically and exclusively to disproportionately large animal structures that function as signals of overall body size, and when assessed through comparison with surrounding nonsignal structures rather than through detection of an estimated slope greater than 1, the positive allometry hypothesis holds. Sexually selected signal structures are predicted to – and, in fact, appear to – scale more steeply with body size than nonsignal structures. For this reason, we suggest that relative patterns of trait scaling offer powerful clues to trait function, particularly when combined with other measures of trait expression such as trait-specific coefficients of variation.

Author Contributions

Conceptualization: D. M. O. and D. J. E.; Methodology: D. M. O.; Formal Analysis: D. M. O.; Investigation: D. M. O., C. E. A., M. J. V., D.

H., R. K., A. K. and S. C.; Resources: D. M. O., M. J. V. and D. J. E.; Data Curation: D. M. O.; Writing – Original Draft: D. M. O., D. H. and D. J. E.; Writing – Reviewing and Editing: D. M. O., C. E. A., M. J. V., D. H., R. K., A. K., S. C. and D. J. E.; Visualization: D. M. O., C. E. A. and D. J. E.; Supervision: D. J. E.; Project Administration: D. M. O.; Funding Acquisition: D. J. E.

Competing Interests

We have no competing interests to report.

Data Archive

Data sets supporting this article are available upon request.

Acknowledgments

We thank the National Science Foundation for funding (OEI-0919781), Libby Beckman, Sandra Chapman, Judith Chupasko, Natalie Cooper, Tannis Davidson, Amy Henrici, Rick Hunter, Julian Kiely, Matthew Lemanna, Carl Mehling, Hana Merchant, Robert Portela Miguez, Robert Niese, José Rosado, Daniela Schwartz-Wings, Justyna Slowiak, Zhao Qi and Stig Walsh for access to specimens and data. Thank you to Brenden Holland, Emy Roberge and Kellie Sorenson for help with collection of Jackson's chameleon data. We are also grateful to Erica McAlister, Alessandro Giusti and Gavin Broad for allowing access to the Diptera, Lepidoptera and Hymenoptera collections at the Natural History Museum, London, U.K. (NHMUK).

Supplementary Material

Supplementary Material related to this article can be found at <https://doi.org/10.1016/j.anbehav.2018.08.005>.

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Appendix

Identifying Extreme Morphology

Many extreme structures appear self-evident. Some, like beetle horns, are massive in absolute and relative size and few would contest their designation as extreme. Others are more ambiguous. Butterfly wings, for example, rarely earn the title of extreme yet, their ontogenetic growth and relative size are more akin to that of beetle horns than other insect wings (Nijhout & Emlen, 1998). Examples like this highlight the ambiguity surrounding extreme morphology and the subjective nature of categorizing structures as extreme. This uncertainty, in part, stems from the lack of established criteria for designating a structure as extreme. For over a century, researchers have explored the evolution of extreme morphology (Andersson, 1994; reviewed in Darwin, 1871; Emlen, 2008). Yet, to our knowledge, not once has the term 'extreme' been defined.

Recognizing and limiting bias is a vital component of biological research and, given the large body of work dedicated toward putatively extreme structures, we believe a consistent method for identifying these structures is needed. Here we suggest three (potentially overlapping) categories of extreme – ontogenetically, statically and evolutionarily extreme – and provide guidelines for assigning structures to each category.

Ontogenetically extreme

Ontogenetically extreme structures are those displaying rates of growth, often occurring in bursts close to reproductive maturity, that outpace other surrounding structures. Examples include the horns of beetles and the wings of lepidopterans, both of which grow to drastic proportions during the same time frame as other, more typically proportioned structures (Nijhout & Emlen, 1998). Ontogenetically extreme should be distinguished by rates of growth that are faster than those of reference structures within the same organism.

Statically extreme

Statically extreme structures are disproportionately larger than other structures when sampled across same stage (generally adult) individuals within a population. Relative size of a focal trait can be assessed by comparing the size of the focal trait to other analogous traits in the same sex (e.g. harlequin beetle, *Acrocynus longimanus*, forelegs are relatively larger than midlegs or hindlegs; Zeh, Zeh, & Tavakilian, 1992) or by comparing the size of the same trait across sexes (e.g. harlequin beetle forelegs are disproportionately larger in males than they are in females; Zeh et al., 1992). Statically extreme structures should be distinguished by comparing slopes and/or intercepts of the static scaling relationships (trait size versus body size) of the focal and reference traits.

Evolutionarily extreme

Evolutionarily extreme structures are extreme when compared with homologous structures in closely related organisms. Examples include the hindlegs of jerboas, which are relatively longer than the hindlegs of their quadrupedal ancestors (Dipodidae: Miljutin, 2008; Wu et al., 2014) and the raptorial forelimbs of mantidflies (Mantispidae: Ohl, Barkalov, & Xin-Yue, 2004). Evolutionarily

extreme structures can be distinguished by (1) comparing static scaling relationships (slopes and/or intercepts) of individuals sampled from populations of ancestral and derived species; (2) comparing mean relative trait size of ancestral and derived species (e.g. Wu et al., 2014); and/or (3) by mapping changes in trait size onto a phylogeny and testing for lineage-specific changes in relative trait size (Wu et al., 2014).

Table A1Sources for *Protoceratops andrewsi* data

Source	Specimen	Number
Dodson, 1976	AMNH	6419
Dodson, 1976	AMNH	6434
Dodson, 1976	AMNH	6430
Dodson, 1976	AMNH	6251
Dodson, 1976	AMNH	6431
Dodson, 1976	AMNH	6486
Dodson, 1976	AMNH	6432
Dodson, 1976	AMNH	6428
Dodson, 1976	AMNH	6409
Dodson, 1976	AMNH	6480
Dodson, 1976	AMNH	6444
Dodson, 1976	AMNH	6485
Dodson, 1976	AMNH	6408
Dodson, 1976	AMNH	6433
Dodson, 1976	AMNH	6429
Dodson, 1976	AMNH	6439
Dodson, 1976	AMNH	6441
Dodson, 1976	AMNH	6477
Dodson, 1976	AMNH	6417
Dodson, 1976	AMNH	6425
Dodson, 1976	AMNH	6413
Dodson, 1976	AMNH	6414
Dodson, 1976	AMNH	6438
Dodson, 1976	AMNH	6466
Dodson, 1976	AMNH	6467
Handa, Watabe, & Tsogtbaatar, 2012	MPC	100/539
Hone, Farke, Watabe, Shigeru, & Tsogtbaatar, 2014	MPC	100/534
Hone et al., 2014	MPC	100/526 B
Hone et al., 2014	MPC	100/526 C
Fastovsky et al., 2011	MPC	100/530 A
Fastovsky et al., 2011	MPC	100/530 B
Fastovsky et al., 2011	MPC	100/530 C
Fastovsky et al., 2011	MPC	100/530 D
Fastovsky et al., 2011	MPC	100/530 E
Fastovsky et al., 2011	MPC	100/530 F
Fastovsky et al., 2011	MPC	100/530 G
Fastovsky et al., 2011	MPC	100/530 H
Unpublished photos	IVPP	23899
Unpublished photos	IVPP	Unnumbered medium
Unpublished photos	IVPP	Unnumbered small
Unpublished photos	ZPAL	MgD-II/2b
Unpublished photos	ZPAL	MgD-II/5
Unpublished photos	CMNH	9185
Unpublished photos	NHM	5134
Unpublished photos	NHM	6442
Unpublished photos	NHM	6440
Unpublished photos	AMNH	6418
Unpublished photos	AMNH	6637
Unpublished photos	AMNH	6422
Unpublished photos	AMNH	6485

AMNH = American Museum of Natural History (New York, U.S.A.); MPC = Mongolian Palaeontological Centre (Ulaanbaatar, Mongolia); IVPP = Institute of Vertebrate Palaeontology and Palaeoanthropology (Beijing, China); ZPAL = Zoological Institute of Paleobiology, Polish Academy of Sciences (Warsaw, Poland); CMNH/CM = Carnegie Museum of Natural History (Pittsburgh, PA, U.S.A.); NHM = Natural History Museum (London, U.K.).

Table A2Sources for *Rhamphorhynchus muensteri* data

Source	Specimen	Number	Wellnhofer number
Wellnhofer, 1975	BSP	1960.I.470	9
Wellnhofer, 1975	BSP	1938.I.503	11
Wellnhofer, 1975	Eichstaett		28
Wellnhofer, 1975	YPM	1778	33
Wellnhofer, 1975	SMF	R 4128	43
Wellnhofer, 1975	CM	11429	53
Wellnhofer, 1975	BSP	1907 I 37	60
Wellnhofer, 1975	SOS	3558	77
Wellnhofer, 1975			102
Hone, 2012	NHM	W1198z0077/0001	
Direct measurement	TMP	2008.041.0001	
Direct measurement	MBR	3650.3	
Direct measurement	BMNS	21	
Measured from photo	NMS	G.1994.13.1.	
Measured from photo	TPI	1012	

BSP = Palaeontological Museum, Munich (Munich, Germany); YPM = Yale Peabody Museum (New Haven, CT, U.S.A.); SMF = Forschungsinstitut und Naturmuseum Senckenberg (Frankfurt, Germany); CMNH/CM = Carnegie Museum of Natural History (Pittsburgh, PA, U.S.A.); SOS = Jura Museum (Eichstätt, Germany); NHM = Natural History Museum (London, U.K.); TMP = Royal Tyrell Museum of Palaeontology (Drumheller, AB, Canada); MBR = Museo Argentino de Ciencias Naturales (Buenos Aires, Argentina); BMNS = Brazoport Museum of Natural Science (Clute, TX, U.S.A.); NMS = National Museums of Scotland (Edinburgh, U.K.); TPI = Thanksgiving Point Institute (Museum of Ancient Life, Lehi, UT, U.S.A.).

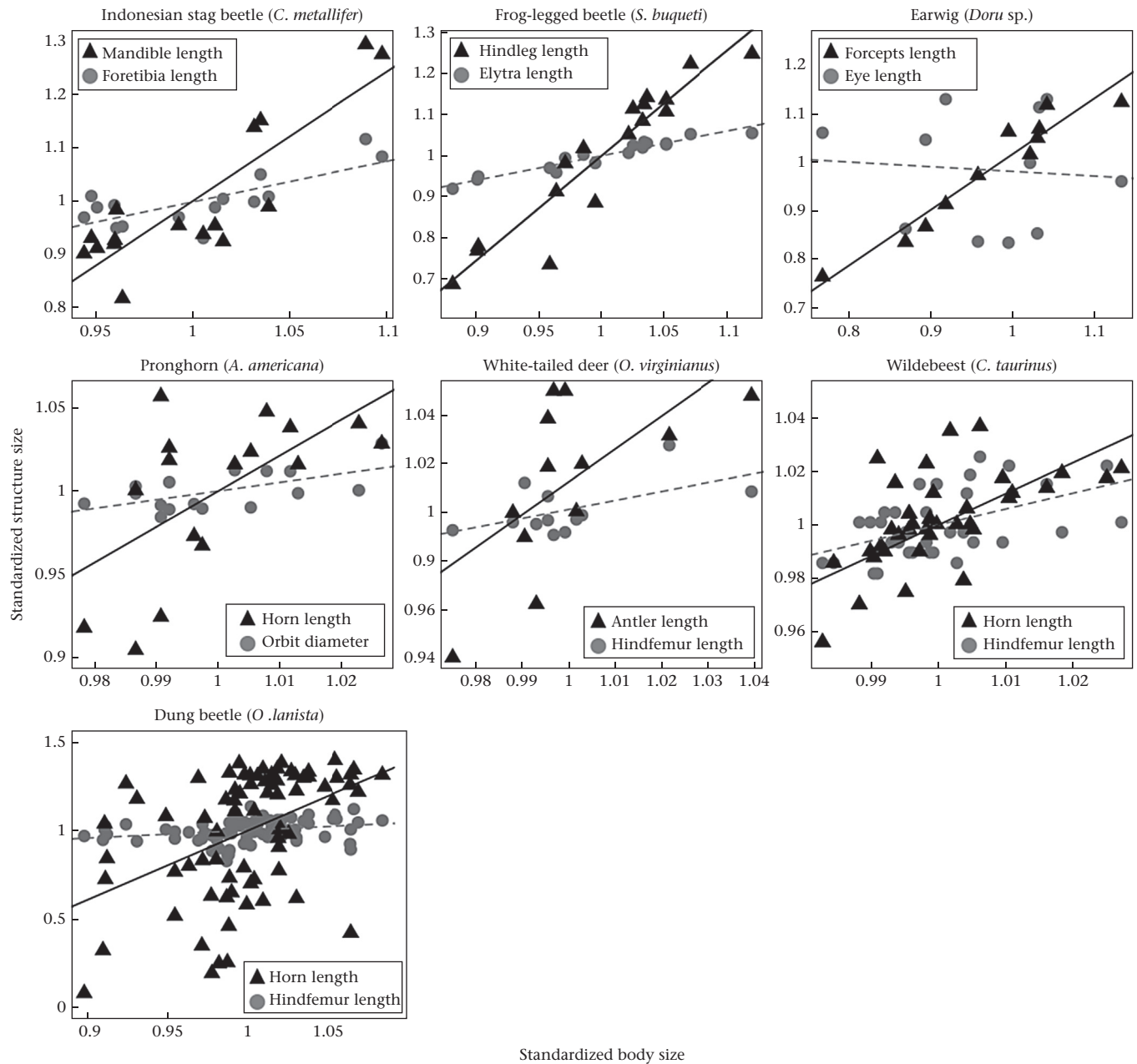


Figure A1. Scaling relationships for extreme sexually selected signal structures in Indonesian stag beetles, *Cyclommatus metallifer*, frog-legged leaf beetles, *Sagra buqueti*, earwigs (*Doru* sp.), pronghorns, *Antilocapra americana*, white-tailed deer, *Odocoileus virginianus*, wildebeest, *Connochaetes taurinus*, and dung beetles, *Onthophagus lanista*. Lines represent ordinary least squares regression of log standardized structure size on log standardized body size (slope estimates and sample sizes reported in Table S1). —▲—: focal traits; —●—: reference traits.

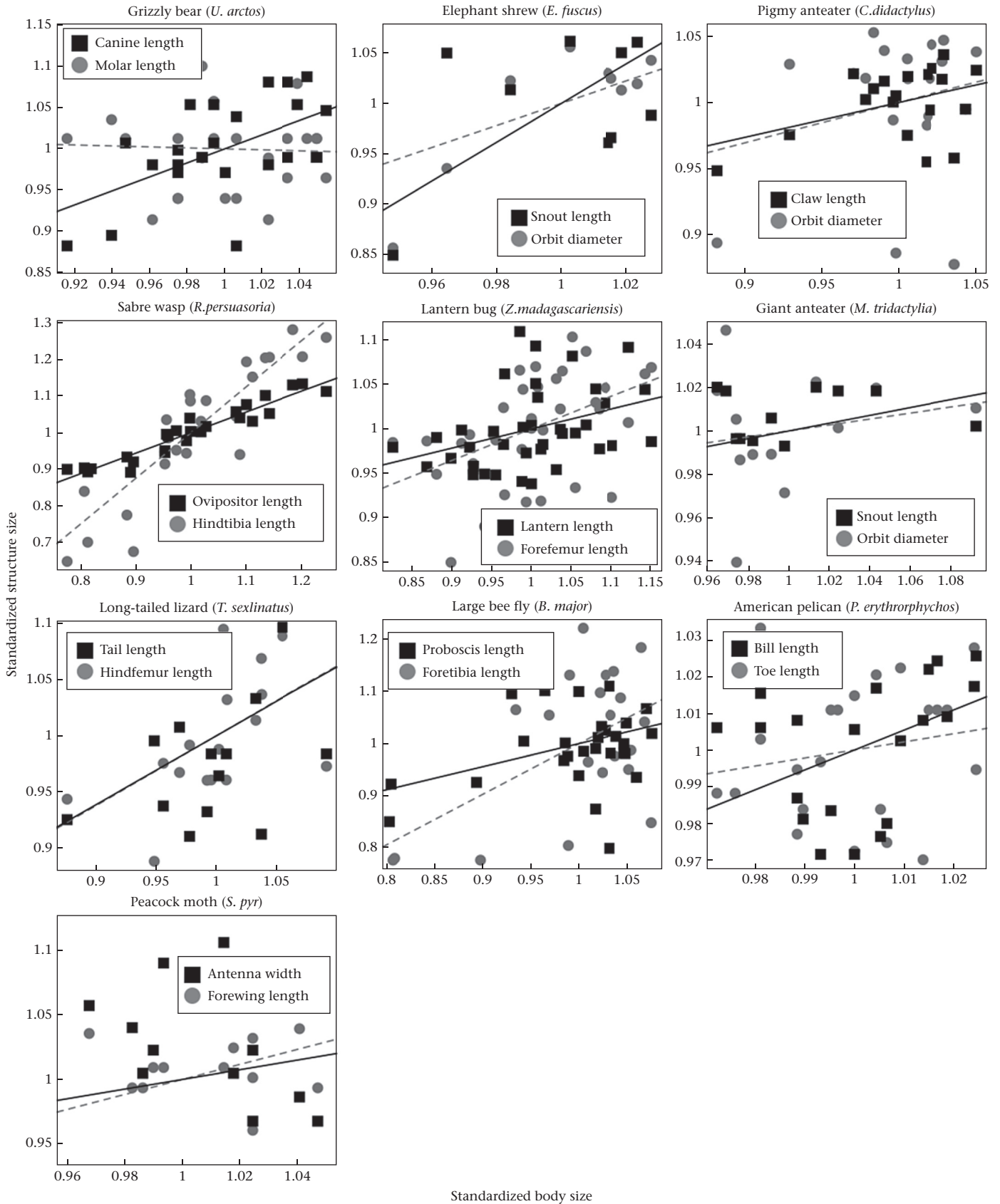


Figure A2. Scaling relationships for extreme naturally selected/nonsignal structures in grizzly bears, *Ursus arctos*, elephant shrews, *Elephantulus fuscus*, pigmy anteaters, *Cyclops didactylus*, sabre wasps, *Rhyssa persuasoria*, lantern bugs, *Zanna madagascariensis*, giant anteaters, *Myrmecophaga tridactylia*, long-tailed lizards, *Takydromus sexlinatus*, large bee flies, *Bombylius major*, American pelicans, *Pelecanus erythrorhynchus*, and peacock moths, *Saturnia pyri*. Lines represent ordinary least squares regression of log standardized structure size on log standardized body size (slope estimates and sample sizes reported in Table S1). —■—: focal traits; —●—: reference traits.

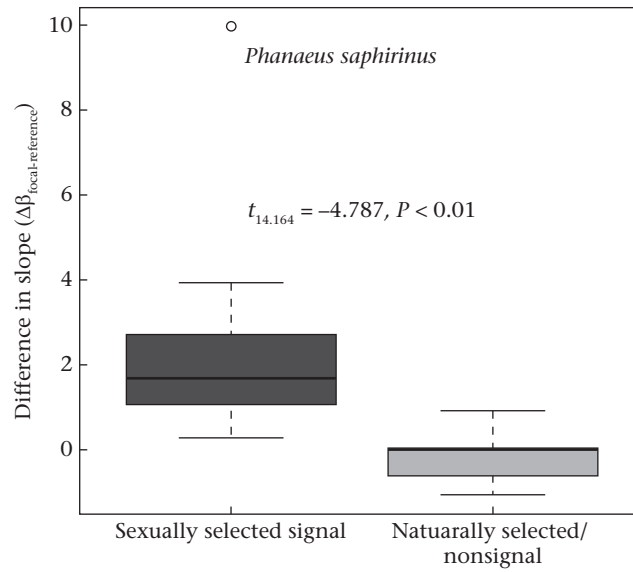


Figure A3. Comparison of $\Delta\beta_{\text{focal-reference}}$ (difference between the scaling relationship slope of focal traits and reference traits) between extreme sexually selected signal traits ($N = 14$) and extreme nonsignal selected traits ($N = 15$). Dung beetles, *Phanaeus saphirinus*, noted as outlier.

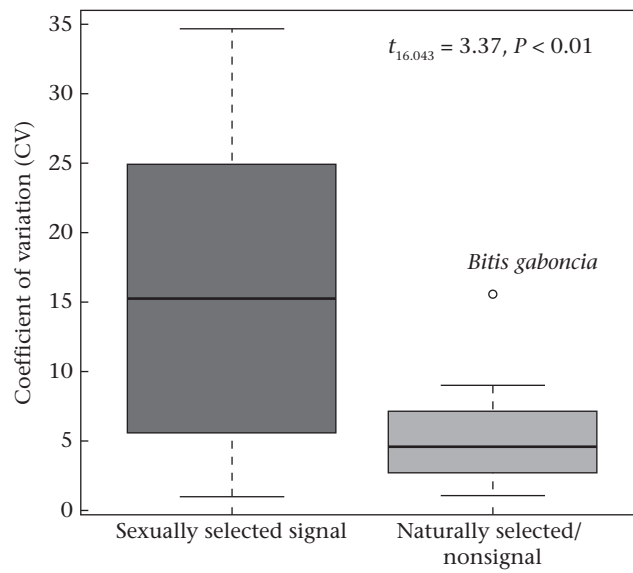


Figure A4. Comparison of coefficients of variation (CV) between extreme sexually selected signal traits ($N = 14$) and extreme nonsignal selected traits ($N = 15$). Gaboon viper, *Bitis gabonica*, noted as outlier.