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COMMENTARY

The opportunity for sexual selection: not mismeasured, just misunderstood

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Abstract

Evolutionary biologists have developed several indices, such as selection gradients (β) and the opportunity for sexual selection (I_s), to quantify the actual and/or potential strength of sexual selection acting in natural or experimental populations. In a recent paper, Klug et al. (J. Evol. Biol. 23, 2010, 447) contend that selection gradients are the only legitimate metric for quantifying sexual selection. They argue that I_s and similar mating-systembased metrics provide unpredictable results, which may be uncorrelated with selection acting on a trait, and should therefore be abandoned. We find this view short-sighted and argue that the choice of metric should be governed by the research question at hand. We describe insights that measures such as the opportunity for selection can provide and also argue that Klug et al. have overstated the problems with this approach while glossing over similar issues with the interpretation of selection gradients. While no metric perfectly characterizes sexual selection in all circumstances, thoughtful application of existing measures has been and continues to be informative in evolutionary studies.

Introduction

Biologists have long been interested in the prominent role that sexual selection plays in organismal evolution (Darwin, 1871; Fisher, 1929; Bateman, 1948). This key evolutionary process is thought to be responsible for many of the observed morphological and behavioural differences between the sexes (Andersson, 1994), including extreme ornaments that appear detrimental to survival (Promislow *et al.*, 1992) and those that contribute to macroevolutionary processes such as speciation (Qvarnstrom *et al.*, 2006; van Doorn *et al.*, 2009). Demonstrating the action of sexual selection can be difficult, and theoreticians have developed a number of metrics to measure the actual or potential strength of sexual selection under varying ecological and social conditions.

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In a recent review of several of these methods, Klug *et al.* (2010a) strongly advocated for a trait-based approach using sexual selection gradients or differentials and were critical of fitness variance-based approaches such as the opportunity for sexual selection ($I_{\rm s} = I_{\rm mates}$; Crow, 1958; Wade, 1979; Shuster & Wade, 2003) and the Bateman gradient (Arnold & Duvall, 1994), as well as population-based approaches such as the operational sex ratio (OSR; Emlen & Oring, 1977). Specifically, Klug *et al.* argued that (i) $I_{\rm s}$, the Bateman gradient and the OSR do not reflect actual selection on traits, (ii) these metrics may fail to capture meaningful variation in reproductive success and (iii) researchers inappropriately employ these metrics to quantify the strength of sexual selection.

We agree with several of the points raised by Klug *et al.* (2010a), but we disagree strongly with their conclusion that mating system measures such as I_s , the Bateman gradient and the OSR have no place in the study of sexual selection. In this paper, we argue that many of the flaws in the use of these metrics are overstated. We focus our discussion on the many uses of I_s , because it is

developed from a population-genetic selection framework, but also discuss other variance- and population-based metrics of sexual selection. We also suggest that Klug *et al.* take too restrictive a view of the goals of sexual selection studies, leading them to unnecessarily reject metrics based on population mating system parameters. Finally, we discuss some of the limitations in the use of selection gradients and suggest that over-reliance on this approach could create many of the same problems Klug *et al.* seek to avoid.

The utility of the opportunity for selection

We agree with Klug and colleagues that sexual selection gradients (β) and the opportunity for sexual selection (I_s) are neither equivalent nor interchangeable. Sexual selection gradients (derived from the more general concept of selection gradients) measure how a given trait covaries with relative fitness, measured in terms of mating success or offspring produced (Lande & Arnold, 1983; Arnold & Duvall, 1994) and therefore reflect the direct (and/or indirect) strength of sexual selection acting on that trait. In contrast, the opportunity for sexual selection (derived from the more general concept of opportunity for selection, c.f., Crow, 1958) measures the variance in mating success divided by the squared average in mating success (Wade, 1979; Wade & Arnold, 1980). Because this ratio measures the variance in relative fitness in the context of sexual selection, it is mathematically related to β and reflects the maximum potential strength of sexual selection in a particular population (Arnold & Wade, 1984a,b). Actual selection may not realize this upper boundary in natural populations because stochastic factors affecting mating patterns may decrease the actual selection on traits below this maximum; i.e. the correlation between phenotype and relative fitness rarely equals one (Shuster & Wade, 2003). As Klug et al. (2010a) correctly point out, the degree of discrepancy between the maximum possible strength of selection acting (described by I_s) and the measurable strength of selection acting on a trait (described by β) cannot be determined from I_s alone. However, it should also be noted that stochastic processes influence any measure of selection, regardless of whether such estimates are directly or indirectly measured (Wade & Shuster, 2010).

Thus, we agree with Klug *et al.* (2010a) that β and I_s measure different things, but these authors have misconstrued this difference to suggest that measures such as I_s are not worthwhile for understanding sexual selection. We maintain that the utility of each metric depends on the questions being asked. If one is exclusively interested in the evolution of specific traits or the potential evolutionary trajectories of one or more aspects of an organism's phenotype, then selection gradients are indeed the appropriate measure. If, however, one is interested in whether a sex difference in the variance in

relative fitness exists, or whether ecological, population and social factors that covary with fitness variance could shape patterns of selection and give rise to general patterns across classes of individuals, populations or species, then $I_{\rm s}$ and/or other measures often are more appropriate.

Although I_s measures the potential strength of selection, rather than the actual strength of selection acting on a trait, we believe this metric has two important uses for behavioural and evolutionary biologists. First, Is provides a useful summary statistic to describe a mating system (Arnold & Duvall, 1994; Shuster & Wade, 2003; Jones, 2009). Classical definitions of mating systems (e.g. Emlen & Oring, 1977) rely on terms such as 'monogamy' and 'polygyny' to describe the distribution of matings within a population. Modern molecular techniques can now identify actual patterns of mating and fertilization, and it is has become evident that verbal descriptors do a poor job of capturing the full diversity of reproductive affiliations present within and across populations (Hughes, 1998; Petrie & Kempenaers, 1998). Individuals can pursue different reproductive strategies with various outcomes; importantly, males and females may mate multiple times (or not at all), and this multiple mating may or may not serve to greatly increase variation in mating and/or reproductive success (Webster et al., 1995; Whittingham & Dunn, 2005; Freeman-Gallant et al., 2005). Measures of I_s effectively integrate the contributions of mate choice, social interactions, mate monopolization and other factors affecting mating patterns in a single value, thereby providing a concise description of the distribution of fertilizations (Vanpe et al., 2008; Krakauer, 2008; Mobley & Jones, 2007). We are uncertain why Klug et al. (2010a) find this usage of I_s 'disconcerting'.

Second, when properly employed, I_s is not only sufficient for measuring the potential for selection to occur, it is necessary for identifying whether sexual selection may operate at all (Hersch & Phillips, 2004). Estimates of I_s are therefore fundamental to examining ecological and social factors that have the potential to influence sexual selection. Because it is standardized by mean fitness, I_s is a unit-less fitness metric that can be compared across classes of individuals (e.g. males vs. females within a population) or across populations (e.g. those that show strong sexual dimorphism vs. those that do not). This measure describes the distribution of mating success within a population, and it is this variance in mating success that drives sexual selection (Arnold & Duvall, 1994; Arnold & Wade, 1984a; Shuster & Wade, 2003; Jones, 2009). Thus, I_s can be used to separate populations or conditions where sexual selection can have a strong evolutionary effect from those where it cannot. This application of I_s is useful for broad comparative studies seeking to understand general patterns of sexual dimorphism and ornamentation (Clutton-Brock et al., 1980; Bro-Jorgensen, 2007), and also for

theoretical and empirical studies seeking to understand how ecological and social factors shape the potential for sexual selection (Mobley & Jones, 2007; DuVal & Kempenaers, 2008; Webster *et al.*, 2007).

Klug et al. (2010a) specifically criticize the use of I_s when the traits relevant to mate choice or intrasexual competition are unspecified. We disagree and argue instead that the opportunity for selection's freedom from assumptions about specific targets of selection provides several advantages if the researcher is not focused exclusively on the evolution of a particular trait. Because $I_{\rm s}$ is calculated from measures of total mating success or, when available, measures of the mean and variance in offspring numbers from parentage analyses (Wade & Shuster, 2004; Shuster, 2009), it integrates the potential of all the organism's traits to contribute to selection. This feature is particularly useful when targets of selection are poorly known or are difficult to measure, or when patterns of actual selection may vary across time, space or social contexts (Moller & Jennions, 2002; Coleman et al., 2004; DuVal & Kempenaers, 2008).

Moreover, when I_s is calculated separately for males and females, these measures allow comparison of the potential strength of selection in each sex; the magnitude and sign of the sex difference in the opportunity for selection, $\Delta I = I_{\text{males}} - I_{\text{females}}$, identifies the direction and degree to which sexual dimorphism is expected to evolve (Shuster & Wade, 2003; Shuster, 2009). Bateman (1948) suggested that for many species the variance in mating success will be higher for males than for females, thus facilitating stronger sexual selection on male traits and behaviours used in inter-sexual attraction or intra-sexual competition. However, in many systems of particular interest to biologists, including brood parasites (Woolfenden et al., 2002), cooperative breeders (Hauber & Lacey, 2005) and sex-role-reversed species (Mobley & Jones, 2007), it may not be clear a priori whether males or females should exhibit higher reproductive potential. For example, behavioural evidence and the occurrence of male pregnancy suggested that the broad-nosed pipefish, Syngnathus typhle, was sex-role-reversed with respect to the intensity of sexual selection (Berglund et al., 1986). An examination of patterns of parentage showed that females displayed higher values of I_s and other mating system metrics compared to males, further supporting the occurrence of sex-role-reversal in S. typhle and validating Bateman's assertion that the sex experiencing stronger sexual selection should display higher variances in mating and reproductive success (Jones et al., 2000, 2005).

An extension of measures of the opportunity for selection is that I_s can be partitioned into component parts to further examine which aspects of mate acquisition or monopolization may be most important in determining the overall variance in reproductive success (Arnold & Wade, 1984a,b; Webster *et al.*, 1995). This partitioning can involve additive terms, such as the

relative contributions of within pair vs. extra pair paternity to male reproductive success (Webster et al., 1995, 2007; Whittingham & Dunn, 2005), or the relative opportunity for sexual selection from male-male competition and female mate choice (DuVal & Kempenaers, 2008). It can also include multiplicatively nested terms, such as the contributions of mate number, average mate fecundity and average paternity share (Webster et al., 1995; Freeman-Gallant et al., 2005; Webster et al., 2007; Krakauer, 2008). These terms, along with the associated covariance terms, indicate where most of the reproductive variance exists and whether trade offs may occur between fitness components, and therefore may indicate where selection has the greatest potential to act. Partitioning I_s in this way provides greater resolution of the source, the possible intensity and the mechanisms by which sexual selection operates (e.g. Webster et al., 2007), providing the foundation for trait-based studies aimed at measuring sexual selection on phenotypes that affect mating success through increased social status, mate attraction or mate quality.

The utility of other metrics: OSR and Bateman gradients

We have focused on I_s , but similar arguments can be made for other population- and mating-system-based metrics that Klug et al. (2010a) seek to discard. For example, the OSR measures the number of sexually active males relative to the number of sexually active females in a population, and therefore is a useful measure of the potential level of competition among individuals of one sex for mating access to individuals of the other sex (Emlen & Oring, 1977). We agree with Klug et al. that the OSR is an inaccurate measure of selection intensity because it does not measure selection at all. Moreover, estimates of OSR must be used with some caution because instantaneous estimates of the intensity of competition tend to over-estimate the overall intensity of sexual selection throughout the breeding season (Shuster & Wade, 2003; Shuster, 2009). However, OSR does provide useful information on instantaneous levels of intra-sexual competition, conditions likely to be perceived and responded to by individuals within field and laboratory populations alike (Berglund et al., 1986; Mobley & Jones, 2007). When the OSR is strongly biased towards one sex, individuals of that sex are likely to compete intensely for mates (Emlen & Oring, 1977). Selection favouring behavioural or morphological traits that lead to success in this competition is expected to occur under these circumstances, even if individual mating success does not remain constant over the entire breeding season (Shuster & Wade, 2003).

Similarly, the Bateman gradient (Bateman, 1948) measures the covariance between reproductive success (number of offspring) and mating success (number of mates), and so describes the extent to which total

reproductive success is tied to the ability to acquire mates (Arnold & Duvall, 1994; Jones, 2009). When this gradient is steep, then fitness is strongly tied to mating ability, and we expect strong selection on traits that help an individual acquire mates. In contrast, when the gradient is near zero, then total fitness is relatively independent of mating success and precopulatory sexual selection is unlikely to be acting strongly on any trait(s), measured or unmeasured. Moreover, comparison of the magnitude and sign of Bateman gradients estimated for each sex provide a quantitative measure of the degree to which sexual conflict may exist within and between species (Wade & Shuster, 2005). In each of these cases, metrics for the potential strength of sexual selection provide insights regardless of whether specific traits used in competition for mates have been identified.

Reliability of the opportunity for selection

Klug *et al.* (2010a) raise two key objections to I_s (as well as other measures of sexual selection). Specifically, they argue that (i) I_s is frequently misused as a measure of the strength of selection, and (ii) even when it is not misused, it is unreliable and provides a poor measure of selection. In this section, we address both of these claims.

The first claim rests on an assumption that most researchers calculating I_s are ultimately (and exclusively) interested in natural or sexual selection on specific phenotypic traits. Klug et al. correctly point out that there need not be a correlation between the opportunity for selection and the selection gradient for a given trait, and we agree with this important point. However, many if not most papers examining I_s are interested in the total potential for sexual selection rather than the evolution of specific traits. Klug et al. support their case with several quotes (their box 2) suggesting frequent misuse of I_s , but closer examination of the cited papers indicates that misuse of I_s is far less rampant than Klug et al. suggest. For example, statements from Shuster & Wade (2003) represent two quotations taken from a more than 500page book describing many ways that calculations of I (and I_s) can be useful for understanding organismal ecology and evolution. Shuster and Wade state: 'This is another reason why I_{mates} sets an *upper limit* on response to sexual selection: not all mating is differential with respect to male characters' (p. 31, emphasis in original). Similarly, the DuVal & Kempenaers (2008) quotation, cited as erroneous by Klug et al., is factually correct in describing I_s as a measure of the opportunity for selection and in pointing out that this measure is distinct from the trait-based evolutionary response to selection.

The second claim made by Klug *et al.* – that I_s and related measures do a poor job of quantifying or characterizing sexual selection – is argued using a series of thought experiments and simulations. In these examples, Klug *et al.* assert that the strength of selection acting on a specific trait depends only on how fitness covaries

with the trait (β) , and does not depend on the variance in relative fitness, estimated by mating success (I_s) , or on the numbers of competing males and females (OSR). For studies seeking to understand the evolution of specific traits, this is an excellent point. However, the simulations provided by Klug *et al.* do not negate the utility of I_s or OSR and are not sufficiently general to warrant removal of these measures from the evolutionary biologists' toolbox.

We disagree that the apparent discrepancies Klug et al. point out between the expected and observed performance of I_s (e.g. in their Fig. 2) invalidate it as a useful measure of sexual selection. Contrary to the authors' interpretation, we maintain that I_s does exactly what it should do in the hypothetical scenarios presented, in that $I_{\rm s}$ is high in scenarios where there is large potential for sexual selection to operate and low when there is little potential for sexual selection. Take, for example, the situation where the breeding sex ratio (OSR) becomes strongly biased towards males by exchanging one male for one female in a hypothetical small population (depicted in their Fig. 2d-f). Here, Klug et al. find fault with I_s because it is higher when the OSR is biased towards males, compared to when the OSR is unity (Fig. 2d), regardless of whether copulations are monopolized by a small number of males (Fig. 2e) or are 'as egalitarian as possible' (Fig. 2f) given the sex ratio constraint. To Klug et al., this result casts doubt on I_s, and they conclude that '[u]sing I_s as a proxy for actual selection is therefore problematic'. But there is nothing problematic here at all: Is is behaving exactly as it should, showing increasing potential for sexual selection (i.e. maximum strength of sexual selection) as one moves from a balanced OSR with weak bias in male mating success ($I_s = 0.67$), to a strongly male-biased (2:1) OSR with monogamous mating by those few males able to obtain mates ($I_s = 1.00$), to a strongly male-biased OSR combined with strong monopolization of mates by a small number of breeding males (i.e. polygyny, $I_s = 3.00$). Indeed, competition for mates under a biased breeding sex ratio was one of the two major mechanisms that Darwin (1871) proposed for sexual selection in socially monogamous species (e.g. Price, 1984), as the strength of selection depends not on how 'egalitarian' the system is, but rather on variance in reproductive/mating success. Is is a measure of the maximum covariance between trait values and mating success; it is a mathematical concept, not a sociological one, and there is no reason why I_s should coincide with qualitative assessments of whether a mating system is egalitarian or not.

Klug *et al.* also provide a second quantitative example using larger samples sizes (their Fig. 3), this time using simulations to calculate I_s and selection differentials (closely related to β) in populations of set size that differ in their OSR, and they present comparisons of these values using different rules to assign male mating success. Klug *et al.* observe that I_s and the selection differential are

not always strongly correlated with each other, and that I_s tends to vary based on the social system when underlying selection is held constant. In particular, in cases where selection was independent of mating system, either when there was no selection (their Fig. 3a) or when selection was constant and directional (their Fig. 3b), I_s increased considerably as the OSR became more male biased. Klug *et al.* take this as evidence that the relationship between I_s and the strength of sexual selection is too inconsistent for I_s to be used as a reliable proxy of sexual selection.

However, once again, and despite their claims to the contrary, here Klug et al. do an excellent job of showing how *I_s does* indeed correlate with the selection differential under biologically realistic conditions. First, in two of their five simulations (Fig. 3c,d), the selection differential and I_s correlate remarkably well; the intercept for I_s is consistently higher than it is for the selection differential, but this is as expected because the opportunity for selection will always be larger in magnitude than the selection differential (Crow 1958). Second, many of the simulations depicted are based on assumptions that are biologically unrealistic and/or uninformative. For example, in the first simulation (Fig. 3a), 'mating is random and no variation exists in the trait of interest' – if mating is indeed random, then selection is not occurring and the maximum strength of selection (I_s) will clearly be far greater than the selection differential (= 0) under any population scenario; this merely illustrates a well-known property of I_s and other measures for the opportunity for selection. Similarly, several simulations use unrealistically biased breeding sex ratios, which tend to generate unusually high values of Is, whereas most natural systems exhibit far lower values of I_s (Jones et al., 2001; Table 2, $I_s = 0.25-2.03$; Wade & Shuster, 2004; Table 1, $I_s = 1.36-8.99$; Tatarenkov et al., 2008, Table 4, $I_s = 0.16-$ 4.52).

Third, a simple explanation exists for all three of the uncorrelated examples: in each of these cases, Klug et al. have intentionally made the phenotypic difference between mating and nonmating males independent of the sex ratio, i.e. the relationship between OSR and the selection differential is flat (Fig. 3a,b,e,f). As is well known, for a covariance to be positive, both variables must have nonzero variances. However, in each of these simulations, the selection differential is a constant value, making the exercise of asking whether or not any variable covaries with the selection differential pointless. We suggest that this approach is similar to asking whether or not smoking causes lung cancer, but restricting attention to those people who do not have lung cancer. In that set of people, a researcher would find some people who do not smoke, some who smoke moderately, and some who smoke a great deal, but none of them would have lung cancer, so the researcher could conclude that smoking must not cause lung cancer. We find it telling that in the two cases in which Klug et al. did allow variation in the selection differential, I_s tracked the selection differential perfectly.

We believe Klug et al. mischaracterize the use of I_s in most studies of sexual selection, possibly because they rely too heavily on colloquial definitions of I_s (i.e. the degree of 'reproductive monopolization' in a population); in actuality, I_s is defined in terms of standardized variance (Arnold & Wade, 1984b,a; Shuster & Wade, 2003). Thus, although values for I_s may deviate from expectations based on colloquial definitions, those values are actually consistent with the more precise and mathematical definitions as the opportunity (or potential) for selection to operate. As emphasized above, Is is not a measure for actual selection on any trait(s), and most of the opportunity for sexual selection will remain unrealized if mating patterns are random with respect to male traits. We further suggest that if entirely random patterns of fertilization between males and females are a common feature of most biological systems, then overestimation of selection by opportunity for selection measures will be the least of our frustrations!

Selection gradients are not a panacea

Klug et al. advocate strongly for the use of direct, traitbased measures of sexual selection (e.g. sexual selection gradients), arguing that these measures more accurately capture the strength of sexual selection acting on a trait. We agree that the sexual selection gradient is a powerful statistical measure of the covariance between a trait and relative fitness, and that this is an excellent approach if one is interested in the extent to which sexual selection favours a particular trait. However, Klug et al. allude only briefly to some of the problems associated with selection gradients. First, this approach requires an assumption that a researcher has correctly identified and appropriately measured the key trait that is under selection. If, for example, the researcher has failed to measure the trait that is the target of sexual selection (e.g. those used by females to select a mate), then the conclusion would be that sexual selection is acting weakly on the traits that were measured, but selection on any unmeasured traits would go undetected despite being potentially strong. In contrast, I_s would indicate the potential strength of selection in the population; whether or not a high potential for sexual selection is typically matched by actual selection on phenotypic traits is an important empirical (but not theoretical) question that remains to be answered.

Second, the ability of researchers to detect selection gradients depends first on whether the variance in relative fitness within a population (i.e. the opportunity for selection) is sufficiently large for selection to operate on any trait, and secondarily on whether the traits that the experimenter identified are important. Hersch & Phillips (2004) have concluded 'that power to detect total selection (the selection differential) depends on sample

size and the strength of selection relative to the opportunity of selection'. These authors recommend that 'studies should be designed so that selection is analyzed on at least several hundred individuals, the total opportunity of selection be considered along with the pattern of selection on individual traits, and nonsignificant results be actively reported combined with an estimate of power'.

Third, and importantly, selection gradients are at their core a correlational analysis, and are therefore subject to the same difficulties that are inherent with any such approach, which include issues surrounding the study sample and degree of natural variation (Grafen, 1987, 1988). In particular, with a correlational selection gradient approach, it can be difficult to separate cause from effect: showing that a particular ornamental trait is correlated with mating success does not demonstrate that the trait causes high mating success. This difficulty can arise if the trait of interest is not a target of sexual selection (i.e. does not affect mating success), but is phenotypically or genotypically correlated with an unmeasured trait that is. Partial regression analysis can help circumvent this problem by separating direct from indirect selection on the trait (Lande & Arnold, 1983), but this does not help if the actual targets of selection have not been measured (as above). Moreover, it is possible that high mating success causes enhanced trait expression, as might be the case if expression of an ornament is mediated by hormones and increased attention from females leads to higher hormone levels (Safran et al., 2008), or the level of male courtship behaviour depends upon female proximity and interest (Patricelli & Krakauer, 2010). Thus, sexual selection gradients can suggest that a particular trait is a target of sexual selection; but if one truly wants to know whether the trait affects mating success, then experimental manipulation is needed (Lande & Arnold, 1983; Grafen, 1988; Wade & Kalisz, 1990).

Lessons learned

In summary, the review by Klug et al. attempts to clarify the distinctions among some metrics used to measure sexual selection in natural populations, and we agree with many of their points regarding limitations of population-based metrics that measure the opportunity for sexual selection. First, I_s describes the variance in relative mating success within a population, thus quantifying the maximum potential fitness differential available to drive sexual selection on one or more traits: researchers must be careful not to interpret I_s as a measure of the actual strength of selection acting on specific traits. This point warrants repeating; in fact, this is a warning that advocates of this measure have long made, beginning with Crow (1958). Second, although the dimensionless property of I_s allows comparisons between species or populations, such comparisons must be conducted carefully, as I_s is sensitive to differences in sample size as well as to researcher assumptions about the number of breeding adults in the population (Freeman-Gallant *et al.*, 2005). Direct comparisons across populations must be interpreted with caution to the extent that these factors differ substantially among studies.

Although we agree with some of the caveats raised by Klug et al. (2010a), we disagree strongly with their conclusion that measures such as Is, OSR, and the Bateman gradient are useless or misleading, or that they are regularly misused as substitutes for the strength of selection. Indeed, numerous studies have employed these measures to elucidate the operation of sexual selection, including studies by authors of the Klug et al. paper (e.g. Klug et al., 2010b; Anthes et al., 2010). Measures such as I_s can provide an integrative snapshot of the maximum fitness differentials available to drive selection on one or more traits, and can be used to better understand the effects of demographic, social and ecological factors on the maximum possible strength of sexual selection. I_s and related measures also provide a handy metric to compare across populations or mating systems. Further, I_s values are appropriate for some comparative studies, particularly those comparing the opportunity for selection between males and females or between distinct classes of males, because computational artefacts will be limited when all samples come from the same population. We also see great utility in the partitioning I_s into its relevant component parts, which can shed light on the effects of social and ecological factors on the potential for various selective mechanisms to operate (e.g. Webster et al., 2007). Finally, studies of sexual selection can be complicated by numerous factors, including but not limited to variation in female preferences, environments, densities and metapopulation dynamics (Cornwallis & Uller, 2010); the opportunity for selection is one valuable tool for making sense of these intricate real-world systems. Contrary to the claims made by Klug et al., we believe biologists are, in general, correctly interpreting studies involving the opportunity for selection. We hope this debate will further clarify the proper use and interpretation of measures of I_s .

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References

Andersson, M.B. 1994. *Sexual Selection*. Princeton University Press, Princeton, NJ.

- Anthes, N., David, P., Auld, J.R., Hoffer, J.N.A., Jarne, P., Koene, J.M. et al. 2010. Bateman gradients in hermaphrodites: an extended approach to quantify sexual selection. Am. Nat. 176:
- Arnold, S.J. & Duvall, D. 1994. Animal mating systems a synthesis based on selection theory. Am. Nat. 143: 317–348.
- Arnold, S.J. & Wade, M.J. 1984a. On the measurement of natural and sexual selection - applications. Evolution 38: 720-
- Arnold, S.J. & Wade, M.J. 1984b. On the measurement of natural and sexual selection – theory. Evolution 38: 709–719.
- Bateman, A.J. 1948. Intra-sexual selection in Drosophila. Heredity 2: 349-368.
- Berglund, A., Rosenqvist, G. & Svensson, I. 1986. Reversed sexroles and the parental energy investment in zygotes of 2 pipefish (Sygnathidae) species. Mar. Ecol. Prog. Ser. 29: 209-
- Bro-Jorgensen, J. 2007. The intensity of sexual selection predicts weapon size in male bovids. Evolution 61: 1316-1326.
- Clutton-Brock, T.H., Albon, S.D. & Harvey, P.H. 1980. Antlers, body size, and breeding group size in the Cervidae. Nature (London) 285: 565-567.
- Coleman, S.W., Patricelli, G.L. & Borgia, G. 2004. Variable female preferences drive complex male displays. Nature 428:
- Cornwallis, C.K. & Uller, T. 2010. Towards an evolutionary ecology of sexual traits. Trends Ecol. Evol. 25: 145-153.
- Crow, J.F. 1958. Some possibilities for measuring selection intensities in man. Human Biol. 30: 1-13.
- Darwin, C. 1871. The Descent of Man and Selection in Relation to Sex. John Murray, London.
- van Doorn, G.S., Edelaar, P. & Weissing, F.J. 2009. On the origin of species by natural and sexual selection. Science 326: 1704-
- DuVal, E.H. & Kempenaers, B. 2008. Sexual selection in a lekking bird: the relative opportunity for selection by female choice and male competition. Proc. R. Soc. B-Biol. Sci. 275:
- Emlen, S.T. & Oring, L.W. 1977. Ecology, sexual selection, and the evolution of mating systems. Science 197: 215-223.
- Fisher, R.A. 1929. The Genetical Theory of Natural Selection. Clarendon Press, Oxford.
- Freeman-Gallant, C.R., Wheelwright, N.T., Meiklejohn, K.E., States, S.L. & Sollecito, S.V. 2005. Little effect of extrapair paternity on the opportunity for sexual selection in savannah sparrows (Passerculus sandwichensis). Evolution 59: 422-430.
- Grafen, A. 1987. Measuring sexual selection: why bother? In: Sexual Selection: Testing the Alternatives (J.W. Bradbury & M.B. Andersson, eds), pp. 221-233. John Wiley & Sons, New York.
- Grafen, A. 1988. On the uses of data on lifetime reproductive success. In: Reproductive Success (T.H. Clutton-Brock, ed.), pp. 454-471. University of Chicago Press, Chicago, IL.
- Hauber, M.E. & Lacey, E.A. 2005. Bateman's principle in cooperatively breeding vertebrates: the effects of non-breeding alloparents on variability in female and male reproductive success. Integr. Comp. Biol. 45: 903-914.
- Hersch, E.I. & Phillips, P.C. 2004. Power and potential bias in field studies of natural selection. Evolution 58: 479-485.
- Hughes, C. 1998. Integrating molecular techniques with field methods in studies of social behavior: a revolution results. Ecology 79: 383-399.

- Jones, A.G. 2009. On the opportunity for sexual selection, the bateman gradient and the maximum intensity of sexual selection. Evolution 63: 1673-1684.
- Jones, A.G., Rosenqvist, G., Berglund, A., Arnold, S.J. & Avise, J.C. 2000. The Bateman gradient and the cause of sexual selection in a sex-role-reversed pipefish. Proc. R. Soc. Biol. Sci. B **267**: 677-680.
- Jones, A.G., Walker, D. & Avise, J.C. 2001. Genetic evidence for extreme polyandry and extraordinary sex-role reversal in a pipefish. Proc. R. Soc. Lond. B 268: 2531-2535.
- Jones, A.G., Rosenqvist, G., Berglund, A. & Avise, J.C. 2005. The measurement of sexual selection using Bateman's principles: an experimental test in the sex-role-reversed pipefish Syngnathus typhle. Integr. Comp. Biol. 45: 874-884.
- Klug, H., Heuschele, J., Jennions, M.D. & Kokko, H. 2010a. The mismeasurement of sexual selection. J. Evol. Biol. 23: 447-
- Klug, H., Lindstrom, K. & Kokko, H. 2010b. Who to include in studies of sexual selection is no trivial matter. Ecol. Lett. 13: 1094-1102.
- Krakauer, A.H. 2008. Sexual selection and the genetic mating system of Wild Turkeys. Condor 110: 1-12.
- Lande, R. & Arnold, S.J. 1983. The measurement of selection on correlated characters. Evolution 37: 1210-1226.
- Mobley, K.B. & Jones, A.G. 2007. Geographical variation in the mating system of the dusky pipefish (Syngnathus floridae). Mol. Ecol. 16: 2596-2606.
- Moller, A.P. & Jennions, M.D. 2002. How much variance can be explained by ecologists and evolutionary biologists? Oecologia 132: 492-500.
- Patricelli, G.L. & Krakauer, A.H. 2010. Tactical allocation of effort among multiple signals in sage grouse: an experiment with a robotic female. Behav. Ecol. 21: 97-106.
- Petrie, M. & Kempenaers, B. 1998. Extra-pair paternity in birds: explaining variation between species and populations. Trends Ecol. Evol. 13: 52-58.
- Price, T.D. 1984. Sexual selection on body size, plumage and territory variables in a population of Darwin's finches. Evolution 38: 327-341.
- Promislow, D.E.L., Montgomerie, R. & Martin, T.E. 1992. Mortality costs of sexual dimorphism in birds. Proc. R. Soc. Lond. Biol. Sci. 250: 143-150.
- Qvarnstrom, A., Haavie, J., Saether, S.A., Eriksson, D. & Part, T. 2006. Song similarity predicts hybridization in flycatchers. J. Evol. Biol. 19: 1202-1209.
- Safran, R.J., Adelman, J.S., McGraw, K.J. & Hau, M. 2008. Sexual signal exaggeration affects physiological state in male barn swallows. Curr. Biol. 18: R461-R462.
- Shuster, S.M. 2009. Sexual selection and mating systems. *Proc.* Natl Acad. Sci. USA 106: 10009-10016.
- Shuster, S.M. & Wade, M.J. 2003. Mating Systems and Strategies. Princeton University Press, Princeton, NJ.
- Tatarenkov, A., Healey, C.I.M., Grether, G.F. & Avise, J.C. 2008. Pronounced reproductive skew in a natural population of green swordtails, Xiphophorus helleri. Mol. Ecol. 17: 4522-4534.
- Vanpe, C., Kjellander, P., Galan, M., Cosson, J.F., Aulagnier, S., Liberg, O. et al. 2008. Mating system, sexual dimorphism, and the opportunity for sexual selection in a territorial ungulate. Behav. Ecol. 19: 309-316.
- Wade, M.J. 1979. Sexual selection and variance in reproductive success. Am. Nat. 114: 742-747.

- Wade, M.J. & Arnold, S.J. 1980. The intensity of sexual selection in relation to male sexual behaviour, female choice, and sperm precedence. Anim. Behav. 28: 446-461.
- Wade, M.J. & Kalisz, S. 1990. The causes of natural selection. Evolution 44: 1947-1955.
- Wade, M.J. & Shuster, S.M. 2004. Sexual selection: Harem size and the variance in male reproductive success. Am. Nat. 164:
- Wade, M.J. & Shuster, S.M. 2005. Don't throw Bateman out with the bathwater!. Integr. Comp. Biol. 45: 261–268.
- Wade, M.J. & Shuster, S.M. 2010. Bateman (1948): pioneer in the measurement of sexual selection. Heredity 105: 507–508.
- Webster, M.S., Pruett-Jones, S., Westneat, D.F. & Arnold, S.J. 1995. Measuring the effects of pairing success, extra-pair

- copulations and mate quality on the opportunity for sexual selection. Evolution 49: 1147-1157.
- Webster, M.S., Tarvin, K.A., Tuttle, E.M. & Pruett-Jones, S. 2007. Promiscuity drives sexual selection in a socially monogamous bird. Evolution 61: 2205-2211.
- Whittingham, L.A. & Dunn, P.O. 2005. Effects of extra-pair and within-pair reproductive success on the opportunity for selection in birds. Behav. Ecol. 16: 138-144.
- Woolfenden, B.E., Gibbs, H.L. & Sealy, S.G. 2002. High opportunity for sexual selection in both sexes of an obligate brood parasitic bird, the brown-headed cowbird (Molothrus ater). Behav. Ecol. Sociobiol. 52: 417-425.

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