



# Meta-analysis challenges a textbook example of status signalling and demonstrates publication bias

Alfredo Sánchez-Tójar<sup>1,2†</sup>\*, Shinichi Nakagawa<sup>3</sup>, Moisès Sánchez-Fortún<sup>1,4‡</sup>, Dominic A Martin<sup>2§</sup>, Sukanya Ramani<sup>1,5</sup>, Antje Girndt<sup>1,2</sup>, Veronika Bókony<sup>6</sup>, Bart Kempenaers<sup>7</sup>, András Liker<sup>8</sup>, David F Westneat<sup>9</sup>, Terry Burke<sup>4</sup>, Julia Schroeder<sup>1,2</sup>

<sup>1</sup>Evolutionary Biology Group, Max Planck Institute for Ornithology, Seewiesen, Germany; <sup>2</sup>Department of Life Sciences, Imperial College London, Ascot, United Kingdom; <sup>3</sup>School of Biological, Earth and Environmental Sciences, University of New South Wales, Sidney, Australia; <sup>4</sup>Department of Animal and Plant Sciences, University of Sheffield, Sheffield, United Kingdom; <sup>5</sup>Department of Animal Behaviour, Bielefeld University, Bielefeld, Germany; <sup>6</sup>Lendület Evolutionary Ecology Research Group, Plant Protection Institute, Centre for Agricultural Research, Hungarian Academy of Sciences, Budapest, Hungary; <sup>7</sup>Department of Behavioural Ecology and Evolutionary Genetics, Max Planck Institute for Ornithology, Seewiesen, Germany; <sup>8</sup>MTA-PE Evolutionary Ecology Research Group, University of Pannonia, Veszprém, Hungary; <sup>9</sup>Department of Biology, University of Kentucky, Lexington, United States

**Abstract** The status signalling hypothesis aims to explain within-species variation in ornamentation by suggesting that some ornaments signal dominance status. Here, we use multilevel meta-analytic models to challenge the textbook example of this hypothesis, the black bib of male house sparrows (*Passer domesticus*). We conducted a systematic review, and obtained primary data from published and unpublished studies to test whether dominance rank is positively associated with bib size across studies. Contrary to previous studies, the overall effect size (i.e. meta-analytic mean) was small and uncertain. Furthermore, we found several biases in the literature that further question the support available for the status signalling hypothesis. We discuss several explanations including pleiotropic, population- and context-dependent effects. Our findings call for reconsidering this established textbook example in evolutionary and behavioural ecology, and should stimulate renewed interest in understanding within-species variation in ornamental traits. DOI: https://doi.org/10.7554/eLife.37385.001

### Introduction

Plumage ornamentation is a striking example of colour and pattern diversity in the animal kingdom and has attracted considerable research (*Hill, 2002*). Most studies have focused on sexual selection as the key mechanism to explain this diversity in ornamentation (*Andersson, 1994*; *Dale et al., 2015*). The status signalling hypothesis explains within-species variation in ornaments by suggesting that these ornaments signal individual dominance status or fighting ability (*Rohwer, 1975*). Aggressive contests are costly in terms of energy use, and risk of injuries and predation (*Jakobsson et al., 1995*; *Kelly and Godin, 2001*; *Neat et al., 1998*; *Prenter et al., 2006*; *Sneddon et al., 1998*). These costs could be reduced if individuals can predict the outcome of such contests beforehand

#### \*For correspondence: alfredo.tojar@gmail.com

Present address: <sup>†</sup>Department of Evolutionary Biology, Bielefeld University, Bielefeld, Germany; <sup>‡</sup>Department de Biologia Evolutiva, Ecologia i Ciències Ambientals, University of Barcelona, Barcelona, Spain; <sup>§</sup>Biodiversity, Macroecology and Biogeography, University of Goettingen, Goettingen, Germany

**Competing interests:** The authors declare that no competing interests exist.

#### Funding: See page 14

Received: 09 April 2018 Accepted: 11 October 2018 Published: 13 November 2018

**Reviewing editor:** Diethard Tautz, Max-Planck Institute for Evolutionary Biology, Germany

© Copyright Sánchez-Tójar et al. This article is distributed under the terms of the Creative

Commons Attribution License, which permits unrestricted use and redistribution provided that the original author and source are credited. CC

**eLife digest** Many bird species have colourful, intricately patterned plumage. This ornamentation is generally believed to exist to attract partners. In the 1970s, however, scientists proposed an alternative idea, called the 'status signalling hypothesis'. This suggests that some birds have plumage ornaments that indicate the fighting abilities or dominance status of their bearers, much like the military badges worn by humans. These badges of status might evolve because fights, which commonly determine who gets valuable resources such as food, are a risky business. Individuals would greatly benefit from being able to predict the fighting abilities of any potential competitor and so avoid fights that they will probably lose.

Male house sparrows have a black patch on their throat, known as the bib, that has been considered to be a textbook demonstration of the status signalling hypothesis. However, most of the studies that support this idea studied small numbers of birds and used inconsistent methods. Furthermore, some recent studies have failed to replicate previous findings.

Sánchez-Tójar et al. collected data from several house sparrow populations across the world and systematically scrutinized the published literature to find all of the studies that tested the status signalling hypothesis in house sparrows. This revealed only weak evidence that the bib of male house sparrows signals the fighting abilities of its bearer. Instead, the published literature is a biased subsample; failures to replicate the hypothesis likely remain unpublished.

Currently, failures to replicate previous findings are generally deemed uninteresting, and so are not often published. By demonstrating the need to replicate findings robustly to avoid biasing conclusions, Sánchez-Tójar et al. thus join the call for a change in incentives and scientific culture. DOI: https://doi.org/10.7554/eLife.37385.002

using so-called 'badges of status' – that is, two potential competitors could decide whether to avoid or engage in aggressive interactions based on the message provided by their opponent's signals (*Rohwer, 1975*).

Patches of ornamentation have been suggested to function as badges of status in a wide range of taxa, including insects (*Tibbetts and Dale, 2004*), reptiles (*Whiting et al., 2003*) and birds (*Senar, 2006*). The status signalling hypothesis was originally proposed to explain variation in the size of mountain sheep horns (*Beninde, 1937; Geist, 1966*), but the hypothesis has become increasingly important in the study of variability in plumage ornamentation in birds (*Rohwer, 1975; Senar, 2006*). Among the many bird species studied (*Santos et al., 2011*), the house sparrow (*Passer domesticus*) has become the classic textbook example of status signalling (*Andersson, 1994; Searcy and Nowicki, 2005; Senar, 2006; Davies et al., 2012*). The house sparrow is a sexually dimorphic passerine, in which the main difference between the sexes is a prominent black patch on the male's throat and chest (hereafter '*bib*'). Many studies have suggested that bib size serves as a badge of status, but most studies are based on limited sample sizes, and have used inconsistent methodologies for measuring bib and dominance status (*Nakagawa and Cuthill, 2007; Santos et al., 2011*).

Meta-analysis is a powerful tool to quantitatively test the overall (across-study) effect size (i.e. the 'meta-analytic mean') for a specific hypothesis. Meta-analyses are therefore able to provide more robust conclusions than single studies and are increasingly used in evolutionary ecology (Gurevitch et al., 2018; Nakagawa and Poulin, 2012a; Nakagawa and Santos, 2012b; Senior et al., 2016). Traditional meta-analyses combine summary data across different studies, where design and methodology are study-specific (e.g. effect sizes among studies are typically adjusted for different fixed effects). These differences among studies are expected to increase heterogeneity, and therefore, the uncertainty of the meta-analytic mean (Mengersen et al., 2013). Meta-analysis of primary or raw data is a specific type of meta-analysis allows methodology to be standardized so that comparable effect sizes can be obtained across studies and is, therefore, considered the gold standard in disciplines such as medicine (Simmonds et al., 2005). Unfortunately, meta-analysis of primary data is still rarely used in evolutionary ecology (but see Barrowman et al., 2003; Richards and Bass, 2005; Krasnov et al., 2009), perhaps due to the difficulty of obtaining

the primary data of previously published studies until recently (*Culina et al., 2018; Schmid et al., 2003*).

An important feature of any meta-analysis is to identify the existence of bias in the literature (**Nakagawa and Santos, 2012b**; **Jennions et al., 2013**). For example, publication bias occurs whenever particular effect sizes (e.g. larger ones) are more likely found in the literature than others (e.g. smaller ones). This tends to be the case when statistical significance and/or direction of effect sizes determines whether results were submitted or accepted for publication (Jennions et al., 2013). Thus, publication bias can strongly affect the estimation of the meta-analytic mean, and distort the interpretation of the hypothesis (**Rothstein et al., 2005**). Several methods have been developed to identify this and other biases (**Nakagawa and Santos, 2012b**; **Jennions et al., 2013**); however, such methods are imperfect and dependent on the number of effect sizes available, and therefore should be considered as types of sensitivity analysis (**Nakagawa et al., 2017**; **Nakagawa and Santos, 2012b**).

Here, we meta-analytically assessed the textbook example of the status signalling hypothesis in the house sparrow. Specifically, we combined summary and primary data from published and unpublished studies to test the prediction that dominance rank is positively associated with bib size across studies. We found that the meta-analytic mean was small, uncertain and overlapped zero. Hence, our results challenge the status signalling function of the male house sparrow's bib. Also, we identified several biases in the published literature. Finally, we discuss potential biological explanations for our results, and provide advice for future studies testing the status signalling hypothesis.

#### Results

Overall, we obtained the primary data for seven of 13 (54%) published studies, and we provided data for six additional unpublished studies (*Table 1—Appendix 1*).

#### **Dominance hierarchies**

Mean sampling effort was 36 interactions/individual (SD = 24), which highlights that, overall, dominance hierarchies were inferred reliably across groups (*Sánchez-Tójar et al., 2018b*). The mean Elorating repeatability was 0.92 (SD = 0.07) and the mean triangle transitivity was 0.63 (SD = 0.28). Thus, the dominance hierarchies observed across groups of house sparrows were medium in both steepness and transitivity.

#### Meta-analytic mean

Our meta-analyses revealed a small overall effect size with large 95% credible intervals that overlapped zero (**Table 2**; **Figure 1**). Additionally, the overall heterogeneity ( $l^2_{overall}$ ) was moderate (53%; **Table 2**). Thus, our results suggested that generally, bib size is at best a weak and unreliable signal of dominance status in male house sparrows.

#### Moderators of the relationship between dominance rank and bib size

None of the three biological moderators studied (season, group composition and type of interactions) explained differences among studies (**Table 3**). Sampling effort (i.e. the ratio of interactions to individuals recorded) also was not an important moderator (**Table 3**).

### **Detection of publication bias**

There was no clear asymmetry in the funnel plots (*Figure 2*). Also, Egger's regression tests did not show evidence of funnel plot asymmetry in any of the meta-analyses (*Table 2*). However, published effect sizes were larger than unpublished ones, and the latter were not different from zero (*Table 4*; *Figure 3*). Additionally, we found that the overall effect size decreased over time and approached zero (*Table 4*; *Figure 4*).

#### Discussion

The male house sparrow's bib is not the strong across-study predictor of dominance status once believed. In contrast to the medium-to-large effect found in the previous meta-analysis (*Nakagawa et al., 2007*), our updated meta-analytic mean was small, uncertain and overlapped

## eLIFE Research article

**Table 1.** Studies used in the meta-analyses and meta-regressions testing the across-study relationship between dominance rank and bib size in male house sparrows.

More information is available in the data files provided (Sánchez-Tójar et al., 2018a).

Study ID	Reference	Population ID	Primary data?	Number of groups*	Total number of males <sup>†</sup>	Comments
1	Ritchison, 1985	Kentucky (captivity)	No	3	35	
2	Møller, 1987	Denmark (wild)	Yes	3	37	
3	Andersson and Åhlund, 1991	Sweden (captivity)	No	10	20	Estimate originally reported as statistically non-significant.
4	Solberg and Ringsby, 1997	Norway (captivity)	Yes	5	44	
5	Liker and Barta, 2001	Hungary (captivity)	Yes	1	10	
6	Gonzalez et al., 2002	Spain (captivity)	No	8	41	
7	Hein et al., 2003	Kentucky (wild)	Yes	4	39	
8	Riters et al., 2004	Wisconsin (captivity)	No	4	20	
9	Lindström et al., 2005	New Jersey (captivity)	No	4	28	Author shared processed data, but group ID was unavailable, so data were not re-analysed.
10	Bókony et al., 2006	Hungary (captivity)	Yes	2	19	
11	Buchanan et al., 2010	Scotland (captivity)	No	14 5	56 20	Groups were tested twice. Post-breeding estimates originally reported as statistically non-significant.
12	Dolnik and Hoi, 2010	Austria (captivity)	No	4 4	31 31	Groups were tested twice. Pre-infection estimates originally reported as statistically non-significant.
13	Rojas Mora et al., 2016	Switzerland (captivity)	Yes	14	56	
14	Lendvai et al.	Hungary (captivity)	Yes <sup>3</sup>	4	46	Unpublished data part of: <i>Lendvai et al., 2004; Bókony et al., 2012</i>
15	Tóth et al.	Hungary (captivity)	Yes <sup>3</sup>	3	35	Unpublished data part of: Tóth et al., 2009; Bókony et al., 2012
16	Bókony et al.	Hungary (captivity)	Yes <sup>3</sup>	4	26	Unpublished data part of: <b>Bókony et al., 2010; Bókony et al.,</b> 2012
17	Sánchez-Tójar et al.	Germany (captivity)	Yes <sup>3</sup>	4	95	Unpublished study conducted in 2014.
18	Sánchez-Tójar et al.	Lundy Island (wild)	Yes <sup>3</sup>	7	172	Unpublished study conducted from 2013 to 2016.
19	Westneat	Kentucky (captivity)	Yes <sup>3</sup>	10	40	Unpublished study conducted in 2005.

\*for primary data = yes, groups of birds containing less than four individuals were not included (see Materials and methods).

†Note: since most studies analysed more than one group of birds, the total number of males is different from group size in most cases (see below).

‡Information for the unpublished datasets is available in **Appendix 1—table 5**.

DOI: https://doi.org/10.7554/eLife.37385.003

zero. Thus, the male house sparrow's bib should not be unambiguously considered or called a badge of status. Furthermore, we found evidence for the existence of bias in the published literature that further undermines the validity of the available support for the status signalling hypothesis. First, the meta-analytic mean of unpublished studies was essentially zero, compared to the medium effect size detected in published studies. Second, we found that the effect size estimated in published studies has been decreasing over time, and recently published effects were on average no longer distinguishable from zero. Our findings call for reconsidering this textbook example in evolutionary

# eLIFE Research article

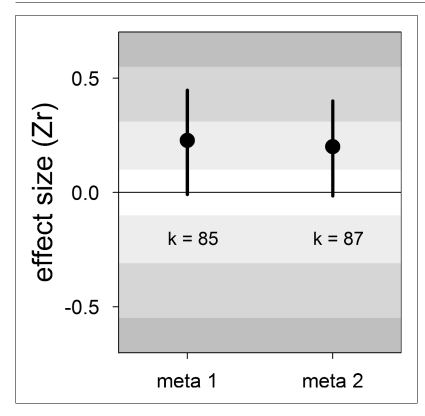
Meta-analysis	К	Meta-analytic mean [95% Crl]	l <sup>2</sup> population ID [ <b>95% Crl</b> ] (%)	l <sup>2</sup> study ID [95% Crl] (%)	l <sup>2</sup> <sub>overall</sub> [95% Crl] (%)	Egger's regression [95% Crl]
meta 1	85	0.23 [-0.01,0.45]	16 [0,48]	21 [0,51]	53 [33,73]	-0.13 [-0.59,0.27]
meta 2	87	0.20 [-0.01,0.40]	15 [0,46]	20 [0,49]	53 [34,74]	-0.12 [-0.55,0.28]

k = number of estimates; Crl = credible intervals;  $l^2 =$  heterogeneity.

DOI: https://doi.org/10.7554/eLife.37385.005

and behavioural ecology, and should stimulate renewed attention to hypotheses explaining withinspecies variation in ornamentation.

The status signalling hypothesis (*Rohwer, 1975*) has been extensively tested to try and explain within-species trait variation (e.g. reptiles: *Whiting et al., 2003*; insects: *Tibbetts and Dale, 2004*; humans: *Dixson and Vasey, 2012*), particularly plumage variation (*Santos et al., 2011*). Soon after the first empirical tests on birds, the black bib of male house sparrows became a textbook example of the status signalling hypothesis (*Andersson, 1994*; *Searcy and Nowicki, 2005*; *Senar, 2006*;



**Figure 1.** Forest plot showing the across-study effect size for the relationship between dominance rank and bib size in male house sparrows. Both meta 1 and meta 2 include published and unpublished estimates, with meta 2 including two non-reported estimates assumed to be zero (see section 'Meta-analyses'). We show posterior means and 95% credible intervals from multilevel meta-analyses. Estimates are presented as standardized effect sizes using Fisher's transformation (*Zr*). Light, medium and dark grey show small, medium and large effect sizes, respectively (*Cohen, 1988*). *k* is the number of estimates. DOI: https://doi.org/10.7554/eLife.37385.004

**Table 3.** Results of the multilevel meta-regressions testing the effect of several moderators on the relationship between dominance rank and bib size in male house sparrows.

Estimates are presented as standardized effect sizes using Fisher's transformation (Zr).

Meta-regression	Estimates	Mean [95% Crl]
meta 1	intercept	0.17 [-0.11,0.46]
(k = 85)	season	-0.11 [-0.41,0.21]
	group composition	0.14 [-0.34,0.59]
	type of interactions	0.33 [-0.17,0.91]
	$R^2_{marginal} =$	23 [2,48]
meta 2	intercept	0.15 [-0.10,0.45]
( <i>k</i> = 87)	season	-0.08 [-0.42,0.22]
	group composition	0.12 [-0.32,0.62]
	type of interactions	0.27 [-0.17,0.85]
	$R^2_{marginal} =$	20 [0,45]
sampling effort	intercept	0.24 [-0.15,0.55]
( <i>k</i> = 61)	sampling effort	0.11 [-0.49,0.74]
	sampling effort <sup>2</sup>	-0.14 [-0.77,0.43]
	$R^2_{marginal} =$	8 [0,24]

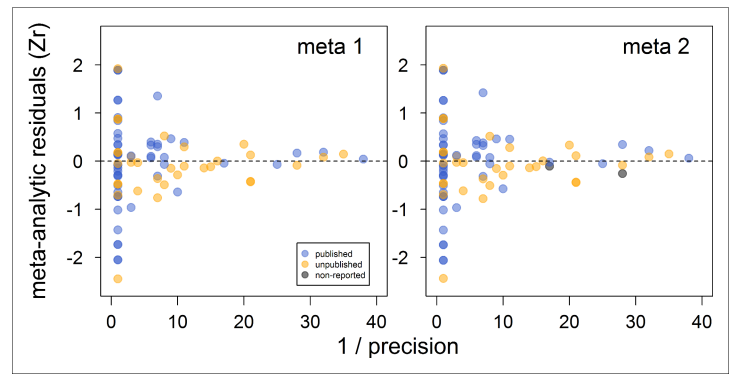
k = number of estimates; Crl = credible intervals;  $R^2_{marginal}$  = percentage of variance explained by the moderators. The factors season (non-breeding: 0, breeding: 1), group composition (mixed-sex: 0, male-only: 1), and type of interactions (all: 0, aggressive-only: 1) were mean-centred, and the covariates 'sampling effort' and its squared term were *z*-transformed.

DOI: https://doi.org/10.7554/eLife.37385.006

**Davies et al.**, 2012), an idea that was later confirmed meta-analytically (**Nakagawa et al.**, 2007). However, **Nakagawa et al.**, 2007 meta-analytic mean was over-estimated because only nine lowpowered studies were available (more in **Button et al.**, 2013). Here, we updated that meta-analysis with newly published and unpublished data. Our results showed that the overall effect size is much smaller and much more uncertain than previously thought. The status signalling hypothesis is thus no longer a compelling explanation for the evolution of bib size across populations of house sparrows.

Similar contradicting conclusions have been reported for other model species. An exhaustive review and meta-analysis on plumage coloration of blue tits (*Cyanistes caeruleus*) revealed that, after dozens of publications studying the function of plumage ornamentation in this species, the only robust conclusion is that females' plumage differs from that of males (*Parker, 2013*). Another example is the long-believed effect of leg bands of particular colours on the perceived attractiveness of male zebra finches (*Taeniopygia guttata*), which has been also experimentally and meta-analytically refuted (*Seguin and Forstmeier, 2012; Wang et al., 2018*). Finally, the existence of a badge of status in a non-bird model species, the paper wasp (*Polistes dominulus; Tibbetts and Dale, 2004*) has also been challenged multiple times (e.g. *Cervo et al., 2008; Green and Field, 2011; Green et al., 2013*), generating doubts about its generality. Our findings corroborate studies showing that abundant replication is needed before any strong or general conclusion can be drawn (*Aarts et al., 2015*), and highlight the existence of important impediments (e.g. *Dublication bias*) to scientific progress in evolutionary ecology (*Forstmeier et al., 2017; Fraser et al., 2018*).

Indeed, our results showed that the published literature on status signalling in house sparrows is likely a biased subsample. The main evidence for this is that the mean effect size of unpublished studies was essentially zero and clearly different from the mean effect size based on published studies, which was of medium size. Furthermore, this moderator (i.e. unpublished vs. published) explained a large percentage of the model's variance. In some of our own unpublished datasets, the relationship between dominance rank and bib size was never formally tested (D.F. Westneat and V. Bókony, *personal communication*, February, 2018), that is, our unpublished datasets are not all examples of the 'file drawer problem' (*sensu Rosenthal, 1979*). Egger's regression tests failed to



**Figure 2.** Funnel plots of the meta-analytic residuals against their precision for the meta-analyses used to test the across-study relationship between dominance rank and bib size in male house sparrows. Both meta 1 and meta 2 include published (blue) and unpublished (orange) estimates, with meta 2 including two additional non-reported estimates (grey; see section 'Meta-analyses'). Estimates are presented as standardized effect sizes using Fisher's transformation (*Zr*). Precision = square root of the inverse of the variance. DOI: https://doi.org/10.7554/eLife.37385.007

detect any funnel plot asymmetry, even in the meta-analyses based on published effect sizes only (*Appendix 2—table 1*). However, because unpublished data indeed existed (i.e. those obtained for this study), the detection failure was likely the consequence of the limited number of effect sizes available (i.e. low power) and the moderate level of heterogeneity found in this study (*Moreno et al., 2009; Sterne and Egger, 2005*).

An additional type of publication bias is time-lag bias, where early studies report larger effect sizes than later studies (*Trikalinos and Ioannidis, 2005*). We detected evidence for such bias

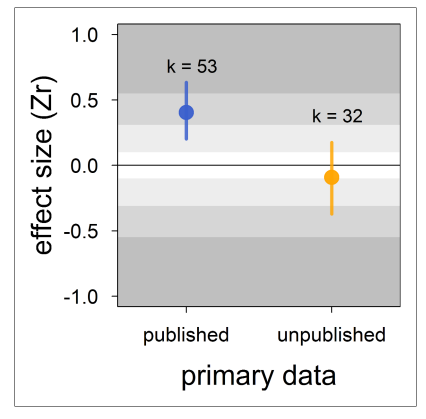
**Table 4.** Results of the multilevel meta-regressions testing for time-lag and publication bias in the literature on status signalling in male house sparrows.

Estimates are presented as standardized effect sizes using Fisher's transformation (Zr). Credible intervals not overlapping zero are highlighted in bold.

Meta-regression	Estimates	Mean [95% Crl]	
time-lag bias	intercept	0.26 [0.03,0.57]	
( <i>k</i> = 53)	year of publication	-0.21 [-0.41,-0.01]	
	$R^2_{marginal} =$	29 [0,66]	
published vs.	intercept	-0.09 [-0.37,0.18]	
unpublished ( $k = 85$ )	published <sup>a</sup>	0.50 [0.19,0.81]	
	$R^2_{marginal} =$	38 [0,68]	

k = number of estimates; CrI = credible intervals;  $R^2_{marginal}$  = percentage of variance explained by the moderators; <sup>a</sup> relative to unpublished. Year of publication was z-transformed.

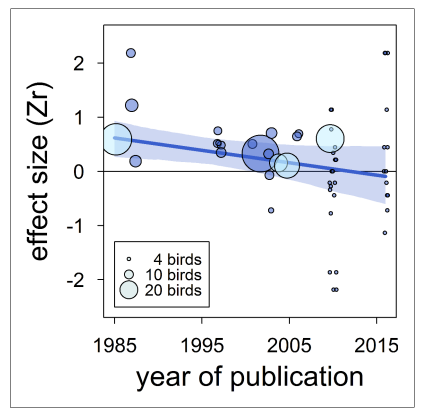
DOI: https://doi.org/10.7554/eLife.37385.010



**Figure 3.** Published effect sizes for the status signalling hypothesis in male house sparrows are larger than unpublished ones. We show posterior means and 95% credible intervals from a multilevel meta-regression. Estimates are presented as standardized effect sizes using Fisher's transformation (*Zr*). Light, medium and dark grey show small, medium and large effects sizes, respectively (*Cohen, 1988*). *k* is the number of estimates. DOI: https://doi.org/10.7554/eLife.37385.008

because the correlation between dominance rank and bib size in published studies has decreased over time and approached zero. Year of publication explained a large percentage of the model's variance, and accounting for year of publication resulted in a strong reduction of the mean effect size across published studies (Table 4 vs. Appendix 2-table 1). Time-lag bias has been detected in other ecological studies (Poulin, 2000); Jennions and Moller, 2002b), including a metaanalysis on status signalling across bird species (Santos et al., 2011). In the latter study, a positive overall (across-species) effect size persisted regardless of the time-lag bias, and no strong evidence for other types of biases was found (Santos et al., 2011). However, Santos et al., 2011 did not attempt to analyse unpublished data, so additional evidence is needed to determine the effect that unpublished data may have on the overall validity of the status signalling hypothesis across bird species. If effect sizes based on unpublished data for other species were of similar magnitude to those obtained for house sparrows, the validity of the status signalling hypothesis across species would need reconsideration. The existence of publication bias in ecology has long been recognized (Cassey et al., 2004; Jennions and Moller, 2002b; Palmer, 2000). Publication bias leads to false conclusions if not accounted for (Rothstein et al., 2005), and is, thus, a serious impediment to scientific progress.

In addition to estimating the overall effect size for a hypothesis, meta-analyses are also used to assess heterogeneity among estimates (*Higgins and Thompson, 2002; Higgins et al., 2003*). Understanding the sources of heterogeneity is an important step towards the correct interpretation of a meta-analytic mean, and can be done using meta-regressions (*Nakagawa and Santos, 2012b*). Here, we found that the percentage of variance that was not attributable to sampling error (i.e. heterogeneity) was moderate. This value is below the average calculated across ecological and evolutionary meta-analyses (*Senior et al., 2016*), and indicates that we accounted for large differences



**Figure 4.** The overall published effect size for the status signalling hypothesis in male house sparrows has decreased over time since first described (k = 53 estimates from 12 publications). The solid blue line represents the model estimate, and the shading shows the 95% credible intervals of a multilevel meta-regression based on published studies (see section 'Detection of publication bias'). Estimates are presented as standardized effect sizes using Fisher's transformation (*Zr*). Circle area represents the size of the group of birds tested to obtain each estimate, where light blue denotes estimates for which group size is inflated due to birds from different groups being pooled, as opposed to dark blue where group size is accurate. DOI: https://doi.org/10.7554/eLife.37385.009

among estimates. Our meta-regressions based on biological moderators explained 20–23% of the variance (**Table 3**). However, none of the biological moderators that we tested strongly influenced the overall effect size, possibly because of limited sample sizes.

The badge of status idea is more complex than typically portrayed (reviewed by Diep and Westneat, 2013). Badges of status are expected to be particularly important in large and unstable groups of individuals where individual recognition would otherwise be difficult (Rohwer, 1975). While the evolution of badges of status in New and Old World sparrows has been related to sociality (i.e. flocking) during the non-breeding season (Tibbetts and Safran, 2009), additional factors need to be involved if the signal is to function in reducing aggression but retaining honesty (Diep and Westneat, 2013). Our results, however, did not show any evidence for a season-dependent effect as the moderator 'season' (breeding vs. non-breeding) was not a strong predictor in our models. Badges of status are expected to function both within and between sexes (Rohwer, 1975; Senar, 2006). Indeed, we found little evidence that the status signalling function of bib size differed between male-only and mixed-sex flocks. Interestingly, when competing for resources, possessing a badge of status would be beneficial for both males and females. However, male but not female house sparrows have a bib. This sexual dimorphism suggests that the bib's function is likely more important when competing for resources other than essential, a priori non-sex-specific resources such as food, water, sand baths and roosting sites. Møller, 1988 and Pape Moller, 1989 reported that female house sparrows preferentially choose males with large bibs (but see Kimball, 1996), and bib size has been positively correlated with sexual behaviour (Veiga, 1996; Møller, 1990), which suggests that

the bib may play a role in mate choice. Furthermore, the original status signalling hypothesis posits that the main benefit of using badges of status would be to avoid fights, which should be particularly important when interacting with unfamiliar individuals (**Rohwer, 1975; Senar, 2006**). Although we did not have data to test whether unfamiliarity among contestants is an important pre-requisite for the status signalling hypothesis, we found no change in mean effect size when only obviously aggressive interactions were studied. In practice, testing whether the bib is important in mediating aggression among unfamiliar individuals is difficult because the certainty of the estimates of individual dominance increases over time as more contests are recorded, but so does familiarity among contestants.

There are some additional explanations for the small and uncertain effect detected by our metaanalyses. First, different populations might be under different selective pressures regarding status signalling. Indeed, the population-specific heterogeneity ( $I_{population ID}^2$ ) estimated in our meta-analyses was 15-16%, suggesting that population-dependent effects might exist. Second, although none of the moderators had a strong influence on the overall effect size, the study-specific heterogeneity estimated in our meta-analyses ( $l_{study ID}^2 = 20-21\%$ ) suggests that the uncertainty observed could still be explained by the status signal being context-dependent. However, context-dependence is often invoked post hoc to explain variation among studies, but strong evidence for it is lacking in most cases. Last, most studies testing the status signalling hypothesis in house sparrows are observational (Table 1), and the only two experimental studies conducted so far were inconclusive (Diep, 2012; Gonzalez et al., 2002). Thus, it cannot be ruled out that the weak correlation observed between dominance status and bib size is driven by a third, unknown variable. In this respect, it has been proposed that the association between melanin-based coloration (such as the bib; e.g. Galván et al., 2015; Galván and Alonso-Alvarez, 2017) and aggression is due to pleiotropic effects of the genes involved in regulating the synthesis of melanin (reviewed by Ducrest et al., 2008). Furthermore, bib size has been shown to correlate with testosterone, a hormone often involved in aggressive behaviour (Gonzalez et al., 2001) but this relationship has not been consistently observed (Laucht et al., 2010). Future studies should shift the focus towards understanding the function of bib size in wild populations and increase considerably the number of birds studied per group. The latter is essential because the statistical power of published tests of the status signalling hypothesis in house sparrows is alarmingly low (power = 8.5% for r = 0.20, Appendix 3) and lower than the average in behavioural ecology (Jennions, 2003).

Our analyses have several potential limitations. First, although the number of studies included in this meta-analysis is more than double that of the previous meta-analysis (*Nakagawa et al., 2007*), it is still limited. Also, it is likely (see above) that additional unpublished data are stored in 'file drawers' (*sensu Rosenthal, 1979*). Second, most tests included in this study were still low-powered in terms of group size (median = 6 individuals/estimate, range = 4–41), and the sample size is inflated because some of the published studies pooled individuals from different groups (*Figure 4*). Third, although our results showed little evidence of an effect of sampling effort on the overall effect size, the quality of the data on dominance and bib size may still be a potential factor explaining differences among studies. Fourth, experiments will normally yield larger effect sizes than observational studies because effects of confounding factors can be reduced (*Palmer, 2000*). Nonetheless, our systematic review only identified two studies where the status signalling hypothesis was tested experimentally in house sparrows (*Gonzalez et al., 2002*; *Diep, 2012*), preventing us from estimating the meta-analytic mean for experimental studies. Note, however, that the results of those experiments were inconclusive, and potentially affected by regression to the mean (*Forstmeier et al., 2017*).

In conclusion, our results challenge an established textbook example of the status signalling hypothesis, which aims to explain within-species variation in ornament size. In house sparrows, we find no evidence that bib size consistently acts as a badge of status across studies and populations, and thus, bib size can no longer be considered a textbook example of the status signalling hypothesis. Furthermore, our analyses highlight the existence of publication biases in the literature, further undermining the validity of past conclusions. Bias against the publication of small ('non-significant') effects hinders scientific progress. We thus join the call for a change in incentives and scientific culture in ecology and evolution (*Forstmeier et al., 2017; Ihle et al., 2017; Nakagawa and Parker, 2015; Parker et al., 2016*).

### **Materials and methods**

### Systematic review

We used several approaches to maximize the identification of relevant studies. First, we included all studies reported in a previous meta-analysis that tested the relationship between dominance rank and bib size in house sparrows (*Nakagawa et al., 2007*). Second, we conducted a keyword search on Web of Science, PubMed and Scopus from 2006 to June 2017 to find studies published after *Nakagawa et al., 2007*, using the combination of keywords ['bib/badge', 'sparrow', 'dominance/status/fighting']. Third, we screened all studies on house sparrows used in a meta-analysis that tested the relationship between dominance and plumage ornamentation across species (*Santos et al., 2011*) to identify additional studies that we may have missed in our keyword search. We screened titles and abstracts of all articles and removed the irrelevant articles before examining the full texts (*Supplementary file 1*). We followed the preferred reporting items for systematic reviews and meta-analyses (PRISMA: *Moher et al., 2009*); see 'Reporting Standards Documents'). We only included articles in which dominance was directly inferred from agonistic dyadic interactions over resources such as food, water, sand baths or roosting sites (*Appendix 1—table 1*).

#### Summary data extraction

Some studies had more than one effect size estimate per group of birds studied. When the presence of multiple estimates was due to the use of different statistical analyses on the same data, we chose a single estimate based on the following order of preference: (1) direct reports of effect size per group of birds studied (e.g. correlation coefficient), (2) inferential statistics (e.g. t, F and  $\chi^2$  statistics) from analyses where group ID was accounted for and no other fixed effects were included, (3) direct reports of effect size where individuals from different groups where pooled together, (4) inferential statistics from models including other fixed effects. When the presence of multiple estimates was due to the use of different methods to estimate bib size and dominance rank on the same data, we chose a single estimate per group of birds or study based on the order of preference shown in **Appendix 1—tables 1–3**. In each case, the order of preference was determined prior to conducting any statistical analysis, and thus, method selection was blind to the outcome of the analyses (more details in Appendix 1).

#### Primary data acquisition

We requested primary data (i.e. agonistic dyadic interactions and bib size measures) of all relevant studies identified by our systematic review. Additionally, we asked authors to share, if available, any unpublished data that could be used to test the relationship between dominance rank and bib size in house sparrows. We emailed the corresponding author, but if no reply was received, we tried contacting all the other authors listed. One study (*Møller, 1987*) provided all primary data in the original publication and, therefore, its author was not contacted. Last, we included our own unpublished data (*Appendix 1—table 5*).

Most studies recorded data from more than one group of birds (**Table 1**). For each primary dataset obtained, we inferred the dominance hierarchy of each group of birds from the observed agonistic dyadic interactions (wins and losses) among individuals using the randomized Elo-rating method, which estimates dominance hierarchies more precisely than other methods (**Sánchez-Tójar et al.**, **2018b**). We then used the provided measures of individual bib size (e.g. area outlined from pictures) or, if possible, calculated bib area from length and width measures following (**Møller**, **1987**). Subsequently, we estimated the Spearman's rho rank correlation ( $\rho$ ) between individual rank and bib size for each group of birds. For one study (**Buchanan et al.**, **2010**), we received the already inferred dominance hierarchies for each group of birds, which we then correlated with bib size to obtain  $\rho$ .

### Effect size coding

Regardless of their source (primary or summary data), we transformed all estimates (e.g.  $\rho$ , *F* statistics, etc) into Pearson's correlation coefficients (*r*), and then into standardized effect sizes using Fisher's transformation (*Zr*) for among-study comparison. We used the equations from **Nakagawa et al.**, **2007** and **Lajeunesse**, **2013**. Since log(0) is undefined, *r* values equal to 1.00 and -1.00 were transformed to 0.975 and -0.975, respectively, before calculating *Zr*. *Zr* values of 0.100, 0.310 and 0.549

were considered small, medium and large effect sizes, respectively (equivalent benchmarks from **Cohen**, **1988**). When not reported directly, the number of individuals (*n*) was estimated from the degrees of freedom. The variance in Zr was calculated as:  $V_{Zr} = 1/(n-3)$ . Estimates (k) based on less than four individuals were discarded (k = 33 estimates discarded).

#### **Meta-analyses**

We ran two multilevel meta-analyses to test whether dominance rank and bib size were positively correlated across studies. The first meta-analysis, in other words '*meta 1*', included published and unpublished (re-)analysed effect sizes (i.e. effect sizes estimated from the studies we obtained primary data from), plus the remaining published effect sizes obtained from summary data (i.e. effect sizes for which primary data were unavailable).

The second meta-analysis, in other words 'meta 2', tested the robustness of the results of meta 1 to the inclusion of non-reported estimates from studies that reported 'statistically non-significant' results without showing either the magnitude or the direction of the estimates (**Table 1**). Receipt of primary data allowed us to recover some but not all the originally non-reported estimates. Two 'non-significant' estimates were still missing. Thus, meta 2 was like meta 1 but included the two non-significant non-reported estimates, which were assumed to be zero (see **Booksmythe et al., 2017** for a similar approach). Note that non-significant estimates can be either negative or positive, and thus, assuming that they were zero may have either underestimated or overestimated them, something we cannot know from non-reported estimates. Meta-analyses based on published studies only are shown in Appendix 2.

We investigated inconsistency across studies by estimating the heterogeneity ( $l^2$ ) from our metaanalyses following **Nakagawa and Santos**, **2012b**.  $l^2$  values around 25, 50% and 75% are considered as low, moderate and high levels of heterogeneity, respectively (**Higgins et al., 2003**).

#### **Meta-regressions**

We tested if season, group composition and/or the type of interactions recorded had an effect on the meta-analytic mean. For that, we ran two multilevel meta-regressions that included the following moderators (hereafter 'biological moderators'): (1) 'season', referring to whether the study was conducted during the non-breeding (September-February) or the breeding season (March-August); (2) 'group composition', referring to whether birds were kept in male-only or in mixed-sex groups; and, (3) 'type of interactions', referring to whether the dyadic interactions recorded were only aggressive (e.g. threats and pecks), or also included interactions that were not obviously aggressive (e.g. displacements). Because only three of 19 studies were conducted in the wild (k = 12 estimates; **Table 1**), we did not include a moderator testing for captive versus wild environments. The three biological moderators were mean-centred following **Schielzeth**, **2010** to aid interpretation.

The ratio of agonistic dyadic interactions recorded to the total number of interacting individuals observed (hereafter 'sampling effort') is a measure of sampling effort that correlates positively and logarithmically with the ability to infer the latent dominance hierarchy (Sánchez-Tójar et al., 2018b). The higher this ratio, the more precisely the latent hierarchy can be inferred (Sánchez-Tójar et al., 2018b). For the subset of studies for which the primary data of the agonistic dyadic interactions were available (12 out of 19 studies; Table 1), we ran a multilevel meta-regression including sampling effort and its squared term as z-transformed moderators (Schielzeth, 2010). The squared term was included because of the observed logarithmic relationship between sampling effort and the method's performance (Sánchez-Tójar et al., 2018b). This meta-regression tested whether sampling effort had an effect on the meta-analytic mean: (i) a positive estimate would indicate that the meta-analytic mean. In contrast, (ii) a negative estimate would indicate that effect sizes were larger when based on unreliable estimates of dominance rank and hence provide evidence for the existence of publication bias.

For all meta-regressions, we estimated the percentage of variance explained by the moderators  $(R^2_{marginal})$  following (*Nakagawa and Schielzeth, 2013*).

#### **Random effects**

All meta-analyses and meta-regressions included the two random effects 'population ID' and 'study ID'. Population ID was related to the geographical location of the population of birds studied. We used Google maps to estimate the distance over land (i.e. avoiding large water bodies) among populations, and assumed the same population ID when the distance was below 50 km (13 populations; **Table 1**). Study ID encompassed those estimates obtained within each specific study (19 studies). Two studies tested the prediction twice for the same groups of birds (**Table 1**) and, within each population, some individuals may have been sampled more than once. However, we could not include 'group ID' and/or 'individual ID' as additional random effects due to either limited sample size or because the relevant data were not available.

#### **Detection of publication bias**

For the meta-analyses, we assessed publication bias using two methods that are based on the assumption that funnel plots should be symmetrical. First, we visually inspected asymmetry in funnel plots of meta-analytic residuals against the inverse of their precision (defined as the square root of the inverse of  $V_{Z_{t}}$  for each meta-analysis. Funnel plots based on meta-analytic residuals (the sum of effect-size-level effects and sampling-variance effects) are more appropriate than those based on effect sizes when multilevel models are used (Nakagawa and Santos, 2012b). Second, we ran Egger's regressions using the meta-analytic residuals as the response variable, and the precision (see above) as the moderator (Nakagawa and Santos, 2012b) for each meta-analysis. If the intercept of such a regression does not overlap zero, estimates from the opposite direction to the metaanalytic mean might be missing and hence we consider this evidence of publication bias (Nakagawa and Santos, 2012b). Further, we tested whether published estimates differed from unpublished estimates. For that, we ran a multilevel meta-regression that included population ID and study ID as random effects, and 'unpublished' (two levels: yes (0), no (1)) as a moderator. This meta-regression was based on meta 1 (i.e. it did not include the two non-reported estimates). We did not use the trim-and-fill method (Duval and Tweedie, 2000a; Duval and Tweedie, 2000b) because this method has been advised against when significant heterogeneity is present (Moreno et al., 2009; Jennions et al., 2013), as it was the case in our meta-analyses (see section 'Results').

Finally, we analysed temporal trends in effect sizes that could indicate 'time-lag bias'. Time-lag bias is common in the literature (*Jennions and Moller, 2002b*; *Poulin, 2000*), and occurs when the effect sizes of a specific hypothesis are negatively correlated with publication date (i.e. effect sizes decrease over time; *Trikalinos and Ioannidis, 2005*). A decrease in effect size over time can have multiple causes. For example, initial effect sizes might be inflated due to low statistical power ('winner's curse') but published more easily and/or earlier due to positive selection of statistically significant results (reviewed by *Koricheva et al., 2013*). We ran a multilevel meta-regression based on published effect sizes only, where 'year of publication' was included as a z-transformed moderator (*Nakagawa and Santos, 2012b*).

All analyses were run in R v. 3.4.0 (R Core Team, 2017). We inferred individual dominance ranks from agonistic dyadic interactions using the randomized Elo-rating method from the R package 'ani-Dom' v. 0.1.3 (Farine and Sánchez-Tójar, 2017; Sánchez-Tójar et al., 2018b). Additionally, we described the dominance hierarchies observed in the groups of house sparrows for which primary data was available. For that we estimated the uncertainty of the dominance hierarchies using the R package 'aniDom' v. 0.1.3 (Farine and Sánchez-Tójar, 2017; Sánchez-Tójar et al., 2018b) and the triangle transitivity (McDonald and Shizuka, 2013) using the R package 'compete' 3.1.0 (Curley, 2016). We used the R package 'MCMCglmm' v. 2.24 (Hadfield, 2010) to run the multilevel meta-analytic (meta-regression) models (Hadfield and Nakagawa, 2010). For each meta-analysis and meta-regression, we ran three independent MCMC chains for 2 million iterations (thinning = 1,800, burn-in = 200,000) using inverse-Gamma priors (V = 1, nu = 0.002). Model chains were checked for convergence and mixing using the Gelman-Rubin statistic. The auto-correlation within the chains was <0.1 in all cases. For each meta-analysis and meta-regression, we chose the model with the lowest DIC value to extract the posterior mean and its 95% highest posterior density intervals (hereafter 95% credible interval). We report all data exclusion criteria applied and the results of all analyses conducted in our study.

We provide all of the R code and data used for our analyses (Sánchez-Tójar et al., 2018a).

### Acknowledgements

AST and AG are grateful for the support of the International Max Planck Research School (IMPRS) for Organismal Biology. We thank Katherine L Buchanan, Sanh K Diep, Fabrice Helfenstein, Anna Kulcsár, Ádám Z Lendvai, Karin M Lindström, Thor Harald Ringsby, Alfonso Rojas Mora, Bernt-Erik Sæther, Emmi Schlicht, Erling J Solberg, Zoltán Tóth and Jarle Tufto for providing the primary data of published and unpublished studies. We thank Wolfgang Forstmeier, Lucy Winder, and Tim Parker and an anonymous reviewer for constructive feedback on the manuscript.

### **Additional information**

#### Funding

<b>U</b>		
Funder	Grant reference number	Author
Max-Planck-Gesellschaft	Open-access funding	Alfredo Sánchez-Tójar
Max-Planck-Gesellschaft	Funding captive house sparrow population	Bart Kempenaers
National Science Foundation		David F Westneat
Natural Environment Research Council	NE/N013832/1	Terry Burke
Volkswagen Foundation		Julia Schroeder
H2020 Marie Skłodowska-Curie Actions	CIG PCIG12-GA-2012- 333096	Julia Schroeder

The funders had no role in study design, data collection and interpretation, or the decision to submit the work for publication.

#### **Author contributions**

Alfredo Sánchez-Tójar, Conceptualization, Data curation, Software, Formal analysis, Validation, Investigation, Visualization, Methodology, Writing—original draft, Project administration, Writing review and editing; Shinichi Nakagawa, Conceptualization, Software, Supervision, Writing—review and editing; Moisès Sánchez-Fortún, Dominic A Martin, Sukanya Ramani, Antje Girndt, Veronika Bókony, Bart Kempenaers, András Liker, David F Westneat, Investigation, Writing—review and editing; Terry Burke, Julia Schroeder, Conceptualization, Supervision, Funding acquisition, Writing—review and editing

#### Author ORCIDs

Alfredo Sánchez-Tójar b http://orcid.org/0000-0002-2886-0649 Shinichi Nakagawa b http://orcid.org/0000-0002-7765-5182 Dominic A Martin b http://orcid.org/0000-0001-7197-2278 Antje Girndt b http://orcid.org/0000-0002-9558-1201 Veronika Bókony b http://orcid.org/0000-0002-2136-5346 Bart Kempenaers b http://orcid.org/0000-0002-7505-5458 David F Westneat b http://orcid.org/0000-0001-5163-8096 Terry Burke b https://orcid.org/0000-0003-3848-1244

#### **Decision letter and Author response**

Decision letter https://doi.org/10.7554/eLife.37385.028 Author response https://doi.org/10.7554/eLife.37385.029

### **Additional files**

#### Supplementary files

• Supplementary file 1. Decision spreadsheet of the systematic review. DOI: https://doi.org/10.7554/eLife.37385.011

- \_\_\_\_\_
- Transparent reporting form
- DOI: https://doi.org/10.7554/eLife.37385.012
- Reporting standard 1. PRISMA statement.

DOI: https://doi.org/10.7554/eLife.37385.013

#### Data availability

All data generated or analysed during this study are openly available at the Open Science Framework. We direct the reader to this project in the main text and the reference list. Link: https://osf.io/ cwkxb/ DOI: 10.17605/OSF.IO/CWKXB

The following dataset was generated:

Author(s)	Year	Dataset title	Dataset URL	Database and Identifier
Alfredo Sánchez- Tójar, Shinichi Na- kagawa, Moisès Sánchez-Fortún, Dominic A Martin, Sukanya Ramani, Antje Girndt, Vero- nika Bókony, Bart Kempenaers, An- drás Liker, David F Westneat, Terry Burke, Julia Schroe- der	2018	Supporting information for "Meta- analysis challenges a textbook example of status signalling and demonstrates publication bias"	http://doi.org/10.17605/ OSF.IO/CWKXB	Open Science Framework, 10.17605/ OSF.IO/CWKXB

#### References

- Aarts AA, Anderson JE, Attridge CJ, Open Science Collaboration. 2015. Estimating the reproducibility of psychological science. Science 349:aac4716. DOI: https://doi.org/10.1126/science.aac4716, PMID: 26315443
- Andersson S, Åhlund M. 1991. Hunger affects dominance among strangers in house sparrows. Animal Behaviour
  41:895–897. DOI: https://doi.org/10.1016/S0003-3472(05)80356-2

Andersson M. 1994. Sexual Selection. New Jersey: Princeton University Press.

Barrowman NJ, Myers RA, Hilborn R, Kehler DG, Field CA. 2003. The variability among populations of coho salmon in the maximum reproductive rate and depensation. *Ecological Applications* **13**:784–793. DOI: https://doi.org/10.1890/1051-0761(2003)013[0784:TVAPOC]2.0.CO;2

Beninde J. 1937. Naturgeschichte Des Rothirshes. Monographie Wildsiiugetiere IV. Leipzig: P. Schöps.

**Bókony V**, Lendvai ÁZ, Liker A. 2006. Multiple cues in status signalling: the role of wingbars in aggressive interactions of male house sparrows. *Ethology* **112**:947–954. DOI: https://doi.org/10.1111/j.1439-0310.2006. 01246.x

**Bókony V**, Kulcsár A, Liker A. 2010. Does urbanization select for weak competitors in house sparrows? *Oikos* **119**:437–444. DOI: https://doi.org/10.1111/j.1600-0706.2009.17848.x

- **Bókony V**, Seress G, Nagy S, Lendvai ÁZ, Liker A. 2012. Multiple indices of body condition reveal no negative effect of urbanization in adult house sparrows. *Landscape and Urban Planning* **104**:75–84. DOI: https://doi.org/ 10.1016/j.landurbplan.2011.10.006
- **Booksmythe I**, Mautz B, Davis J, Nakagawa S, Jennions MD. 2017. Facultative adjustment of the offspring sex ratio and male attractiveness: a systematic review and meta-analysis. *Biological Reviews* **92**:108–134. DOI: https://doi.org/10.1111/brv.12220, PMID: 26405787
- Buchanan KL, Evans MR, Roberts ML, Rowe L, Goldsmith AR. 2010. Does testosterone determine dominance in the house sparrow *Passer domesticus*? an experimental test. *Journal of Avian Biology* **41**:445–451. DOI: https://doi.org/10.1111/j.1600-048X.2010.04929.x
- Button KS, Ioannidis JP, Mokrysz C, Nosek BA, Flint J, Robinson ES, Munafò MR. 2013. Power failure: why small sample size undermines the reliability of neuroscience. *Nature Reviews Neuroscience* **14**:365–376. DOI: https://doi.org/10.1038/nrn3475, PMID: 23571845
- **Cassey P**, Ewen JG, Blackburn TM, Moller AP. 2004. A survey of publication bias within evolutionary ecology. *Proceedings of the Royal Society B: Biological Sciences* **271**:S451–S454. DOI: https://doi.org/10.1098/rsbl. 2004.0218

- Cervo R, Dapporto L, Beani L, Strassmann JE, Turillazzi S. 2008. On status badges and quality signals in the paper wasp *Polistes dominulus*: body size, facial colour patterns and hierarchical rank. *Proceedings of the Royal Society B: Biological Sciences* **275**:1189–1196. DOI: https://doi.org/10.1098/rspb.2007.1779
- **Cohen J.** 1988. *Statistical Power Analysis for the Behavioral Sciences*. Second edition. New Jersey: Taylor & Francis Inc.
- Culina A, Crowther TW, Ramakers JJC, Gienapp P, Visser ME. 2018. How to do meta-analysis of open datasets. Nature Ecology & Evolution 2:1053–1056. DOI: https://doi.org/10.1038/s41559-018-0579-2, PMID: 29915339
- Curley JP. 2016. compete: Analyzing Social Hierarchies. https://cran.r-project.org/web/packages/compete/index. html
- Dale J, Dey CJ, Delhey K, Kempenaers B, Valcu M. 2015. The effects of life history and sexual selection on male and female plumage colouration. *Nature* 527:367–370. DOI: https://doi.org/10.1038/nature15509, PMID: 26536112
- Davies NB, Krebs JR, West SA. 2012. An Introduction to Behavioural Ecology. Oxford: Wiley-Blackwell.
- **Diep SK**. 2012. The Role of Social Interactions on the Development and Honesty of a Signal of Status. University of Kentucky.
- Diep SK, Westneat DF. 2013. The integration of function and ontogeny in the evolution of status signals. Behaviour:1–30. DOI: https://doi.org/10.1163/1568539X-00003066
- Dixson BJ, Vasey PL. 2012. Beards augment perceptions of men's age, social status, and aggressiveness, but not attractiveness. *Behavioral Ecology* 23:481–490. DOI: https://doi.org/10.1093/beheco/arr214
- Dolnik OV, Hoi H. 2010. Honest signalling, dominance hierarchies and body condition in house sparrows Passer domesticus (Aves: Passeriformes) during acute coccidiosis. Biological Journal of the Linnean Society 99:718– 726. DOI: https://doi.org/10.1111/j.1095-8312.2010.01370.x
- Ducrest AL, Keller L, Roulin A. 2008. Pleiotropy in the melanocortin system, coloration and behavioural syndromes. Trends in Ecology & Evolution 23:502–510. DOI: https://doi.org/10.1016/j.tree.2008.06.001, PMID: 18644658
- **Duval S**, Tweedie R. 2000a. A nonparametric "Trim and Fill" Method of Accounting for Publication Bias in Meta-Analysis. *Journal of the American Statistical Association* **95**:89–98. DOI: https://doi.org/10.1080/01621459. 2000.10473905
- Duval S, Tweedie R. 2000b. Trim and fill: a simple funnel-plot-based method of testing and adjusting for publication Bias in meta-analysis. *Biometrics* 56:455–463. DOI: https://doi.org/10.1111/j.0006-341X.2000. 00455.x, PMID: 10877304
- Farine DR, Sánchez-Tójar A. 2017. aniDom: Inferring Dominance Hierarchies and Estimating Uncertainty.https:// cran.r-project.org/package=aniDom
- Forstmeier W, Wagenmakers EJ, Parker TH. 2017. Detecting and avoiding likely false-positive findings a practical guide. *Biological Reviews* **92**:1941–1968. DOI: https://doi.org/10.1111/brv.12315, PMID: 27879038
- Fraser H, Parker T, Nakagawa S, Barnett A, Fidler F. 2018. Questionable research practices in ecology and evolution. *PLOS One* **13**:e0200303. DOI: https://doi.org/10.1371/journal.pone.0200303, PMID: 30011289
- Galván I, Wakamatsu K, Camarero PR, Mateo R, Alonso-Alvarez C. 2015. Low-quality birds do not display highquality signals: the cysteine-pheomelanin mechanism of honesty. *Evolution* **69**:26–38. DOI: https://doi.org/10. 1111/evo.12549
- Galván I, Alonso-Alvarez C. 2017. Individual quality via sensitivity to cysteine availability in a melanin-based honest signaling system. The Journal of Experimental Biology **220**:2825–2833. DOI: https://doi.org/10.1242/jeb.160333, PMID: 28546507
- Geist V. 1966. The evolutionary significance of mountain sheep horns. Evolution 20:558–566. DOI: https://doi. org/10.2307/2406590
- Gonzalez G, Sorci G, Smith LC, de Lope F. 2001. Testosterone and sexual signalling in male house sparrows (Passer domesticus). Behavioral Ecology and Sociobiology **50**:557–562. DOI: https://doi.org/10.1007/s002650100399
- Gonzalez G, Sorci G, Smith LC, de Lope F. 2002. Social control and physiological cost of cheating in status signalling male house sparrows (*Passer domesticus*). Ethology **108**:289–302. DOI: https://doi.org/10.1046/j. 1439-0310.2002.00779.x
- Green JP, Field J. 2011. Interpopulation variation in status signalling in the paper wasp *Polistes dominulus*. Animal Behaviour **81**:205–209. DOI: https://doi.org/10.1016/j.anbehav.2010.10.002
- Green JP, Leadbeater E, Carruthers JM, Rosser NS, Lucas ER, Field J. 2013. Clypeal patterning in the paper wasp *Polistes dominulus*: no evidence of adaptive value in the wild. *Behavioral Ecology* **24**:623–633. DOI: https://doi.org/10.1093/beheco/ars226
- Gurevitch J, Koricheva J, Nakagawa S, Stewart G. 2018. Meta-analysis and the science of research synthesis. *Nature* **555**:175–182. DOI: https://doi.org/10.1038/nature25753, PMID: 29517004

Hadfield JD. 2010. MCMC methods for Multi-Response generalized linear mixed models: themcmcglmmrpackage. Journal of Statistical Software **33**. DOI: https://doi.org/10.18637/jss.v033.i02

- Hadfield JD, Nakagawa S. 2010. General quantitative genetic methods for comparative biology: phylogenies, taxonomies and multi-trait models for continuous and categorical characters. *Journal of Evolutionary Biology* 23:494–508. DOI: https://doi.org/10.1111/j.1420-9101.2009.01915.x, PMID: 20070460
- Hein WK, Westneat DF, Poston JP. 2003. Sex of opponent influences response to a potential status signal in house sparrows. *Animal Behaviour* **65**:1211–1221. DOI: https://doi.org/10.1006/anbe.2003.2132
- Higgins JP, Thompson SG. 2002. Quantifying heterogeneity in a meta-analysis. *Statistics in Medicine* **21**:1539–1558. DOI: https://doi.org/10.1002/sim.1186, PMID: 12111919

- Higgins JP, Thompson SG, Deeks JJ, Altman DG. 2003. Measuring inconsistency in meta-analyses. *Bmj* **327**:557–560. DOI: https://doi.org/10.1136/bmj.327.7414.557, PMID: 12958120
- Hill GE. 2002. A Red Bird in a Brown Bag: The Function and Evolution of Colorful Plumage in the House Finch. New York: Oxford University Press. DOI: https://doi.org/10.1093/acprof:oso/9780195148480.001.0001
- Ihle M, Winney IS, Krystalli A, Croucher M. 2017. Striving for transparent and credible research: practical guidelines for behavioral ecologists. *Behavioral Ecology* 28:348–354. DOI: https://doi.org/10.1093/beheco/ arx003, PMID: 29622916
- Jakobsson S, Brick O, Kullberg C. 1995. Escalated fighting behaviour incurs increased predation risk. Animal Behaviour 49:235–239. DOI: https://doi.org/10.1016/0003-3472(95)80172-3
- Jennions MD, Møller AP. 2002a. Publication Bias in ecology and evolution: an empirical assessment using the 'trim and fill' method. *Biological Reviews of the Cambridge Philosophical Society* **77**:211–222. DOI: https://doi.org/10.1017/S1464793101005875, PMID: 12056747
- Jennions MD, Moller AP. 2002b. Relationships fade with time: a meta-analysis of temporal trends in publication in ecology and evolution. *Proceedings of the Royal Society B: Biological Sciences* **269**:43–48. DOI: https://doi.org/10.1098/rspb.2001.1832
- Jennions MD. 2003. A survey of the statistical power of research in behavioral ecology and animal behavior. Behavioral Ecology **14**:438–445. DOI: https://doi.org/10.1093/beheco/14.3.438
- Jennions MD, Lortie C, Rosenberg M, Rothstein H. 2013. Publication and related biases. In: Koricheva J, Gurevitch J, Mengersen K (Eds). Handbook of Meta-Analysis in Ecology & Evolution. Princenton: Princeton University Press. p. 207–236. DOI: https://doi.org/10.1515/9781400846184-016
- Kelly C, Godin J-G. 2001. Predation risk reduces male-male sexual competition in the trinidadian guppy (*Poecilia* reticulata). Behavioral Ecology and Sociobiology **51**:95–100. DOI: https://doi.org/10.1007/s002650100410
- Kimball RT. 1996. Female choice for male morphological traits in house sparrows, *Passer domesticus*. *Ethology* **102**:639–648. DOI: https://doi.org/10.1111/j.1439-0310.1996.tb01155.x
- Koricheva J, Jennions MD, Lau J. 2013. Temporal trends in effect sizes: causes, detection, and implications. In: Koricheva J, Gurevitch J, Mengersen K (Eds). *Handbook of Meta-Analysis in Ecology & Evolution*. Princenton: Princeton University Press. p. 237–254. DOI: https://doi.org/10.1515/9781400846184-017
- Krasnov BR, Vinarski MV, Korallo-Vinarskaya NP, Mouillot D, Poulin R. 2009. Inferring associations among parasitic gamasid mites from census data. *Oecologia* 160:175–185. DOI: https://doi.org/10.1007/s00442-009-1278-0, PMID: 19189129
- Lajeunesse MJ. 2013. Recovering Missing or Partial Data from Studies: A Survey of Conversions and Imputations for Meta-analysis. In: Koricheva J, Gurevitch J, Mengersen K (Eds). *Handbook of Meta-Analysis in Ecology & Evolution*. Princenton: Princeton University Press. p. 195–206. DOI: https://doi.org/10.1515/9781400846184-015
- Laucht S, Kempenaers B, Dale J. 2010. Bill color, not badge size, indicates testosterone-related information in house sparrows. Behavioral Ecology and Sociobiology 64:1461–1471. DOI: https://doi.org/10.1007/s00265-010-0961-9, PMID: 20730125
- Lendvai AZ, Barta Z, Liker A, Bokony V. 2004. The effect of energy reserves on social foraging: hungry sparrows scrounge more. Proceedings of the Royal Society B: Biological Sciences 271:2467–2472. DOI: https://doi.org/ 10.1098/rspb.2004.2887
- Liker A, Barta Z. 2001. Male badge size predicts dominance against females in house sparrows1. The Condor 103:151–157. DOI: https://doi.org/10.1650/0010-5422(2001)103[0151:MBSPDA]2.0.CO;2
- Lindström KM, Hasselquist D, Wikelski M. 2005. House sparrows (*Passer domesticus*) adjust their social status position to their physiological costs. *Hormones and Behavior* **48**:311–320. DOI: https://doi.org/10.1016/j. yhbeh.2005.04.002, PMID: 15896793
- McDonald DB, Shizuka D. 2013. Comparative transitive and temporal orderliness in dominance networks. Behavioral Ecology 24:511–520. DOI: https://doi.org/10.1093/beheco/ars192
- Mengersen K, Gurevitch J, Schmid CH. 2013. Meta-analysis of primary data. In: Koricheva J, Mengersen K, Gurevit J (Eds). Handbook of Meta-Analysis in Ecology & Evolution. Princenton: Princeton University Press. p. 300–312. DOI: https://doi.org/10.1515/9781400846184-020
- Moher D, Liberati A, Tetzlaff J, Altman DG, PRISMA Group. 2009. Preferred reporting items for systematic reviews and meta-analyses: the PRISMA statement. *PLOS Medicine* **6**:e1000097. DOI: https://doi.org/10.1371/journal.pmed.1000097, PMID: 19621072
- Møller AP. 1987. Variation in badge size in male house sparrows Passer domesticus: evidence for status signalling. Animal Behaviour **35**:1637–1644. DOI: https://doi.org/10.1016/S0003-3472(87)80056-8
- Møller AP. 1988. Badge size in the house sparrow *Passer domesticus* Effects of intra- and intersexual selection. Behavioral Ecology and Sociobiology **22**:373–378.
- Møller A. 1990. Sexual behavior is related to badge size in the house sparrow Passer domesticus. Behavioral Ecology and Sociobiology 27:23–29. DOI: https://doi.org/10.1007/BF00183309
- Moreno SG, Sutton AJ, Ades AE, Stanley TD, Abrams KR, Peters JL, Cooper NJ. 2009. Assessment of regression-based methods to adjust for publication Bias through a comprehensive simulation study. BMC Medical Research Methodology 9:2. DOI: https://doi.org/10.1186/1471-2288-9-2, PMID: 19138428
- Morrison EB, Kinnard TB, STEWART IANRK, Poston JP, HATCH MI, Westneat DF. 2008. The links between plumage variation and nest site occupancy in male house sparrows. *The Condor* **110**:345–353. DOI: https://doi.org/10.1525/cond.2008.8470

- Nakagawa S, Cuthill IC. 2007. Effect size, confidence interval and statistical significance: a practical guide for biologists. *Biological Reviews* 82:591–605. DOI: https://doi.org/10.1111/j.1469-185X.2007.00027.x, PMID: 17 944619
- Nakagawa S, Ockendon N, Gillespie DOS, Hatchwell BJ, Burke T. 2007. Assessing the function of house sparrows' bib size using a flexible meta-analysis method. *Behavioral Ecology* 18:831–840. DOI: https://doi.org/ 10.1093/beheco/arm050
- Nakagawa S, Poulin R. 2012a. Meta-analytic insights into evolutionary ecology: an introduction and synthesis. Evolutionary Ecology 26:1085–1099. DOI: https://doi.org/10.1007/s10682-012-9593-z
- Nakagawa S, Santos ESA. 2012b. Methodological issues and advances in biological meta-analysis. *Evolutionary Ecology* 26:1253–1274. DOI: https://doi.org/10.1007/s10682-012-9555-5
- **Nakagawa 5**, Schielzeth H. 2013. A general and simple method for obtaining *R*<sup>2</sup> from generalized linear mixedeffects models. *Methods in Ecology and Evolution* **4**:133–142. DOI: https://doi.org/10.1111/j.2041-210x.2012. 00261.x
- Nakagawa S, Noble DW, Senior AM, Lagisz M. 2017. Meta-evaluation of meta-analysis: ten appraisal questions for biologists. *BMC Biology* **15**:18. DOI: https://doi.org/10.1186/s12915-017-0357-7, PMID: 28257642
- Nakagawa S, Parker TH. 2015. Replicating research in ecology and evolution: feasibility, incentives, and the costbenefit conundrum. BMC Biology 13:88. DOI: https://doi.org/10.1186/s12915-015-0196-3, PMID: 26510635
- Neat FC, Taylor AC, Huntingford FA. 1998. Proximate costs of fighting in male cichlid fish: the role of injuries and energy metabolism. *Animal Behaviour* 55:875–882. DOI: https://doi.org/10.1006/anbe.1997.0668, PMID: 9632474
- Palmer AR. 2000. Quasi-Replication and the contract of error: lessons from sex ratios, heritabilities and fluctuating asymmetry. Annual Review of Ecology and Systematics **31**:441–480. DOI: https://doi.org/10.1146/annurev.ecolsys.31.1.441
- Pape Moller A. 1989. Natural and sexual selection on a plumage signal of status and on morphology in house sparrows, *Passer domesticus. Journal of Evolutionary Biology* **2**:125–140. DOI: https://doi.org/10.1046/j.1420-9101.1989.2020125.x
- Parker TH. 2013. What do we really know about the signalling role of plumage colour in blue tits? A case study of impediments to progress in evolutionary biology. *Biological Reviews* 88:511–536. DOI: https://doi.org/10. 1111/brv.12013, PMID: 23279555
- Parker TH, Forstmeier W, Koricheva J, Fidler F, Hadfield JD, Chee YE, Kelly CD, Gurevitch J, Nakagawa S. 2016. Transparency in ecology and evolution: real problems, real solutions. *Trends in Ecology & Evolution* **31**:711– 719. DOI: https://doi.org/10.1016/j.tree.2016.07.002, PMID: 27461041
- Poulin R. 2000. Manipulation of host behaviour by parasites: a weakening paradigm? Proceedings of the Royal Society B: Biological Sciences **267**:787–792. DOI: https://doi.org/10.1098/rspb.2000.1072, PMID: 10819148
- Prenter J, Elwood RW, Taylor PW. 2006. Self-assessment by males during energetically costly contests over precopula females in amphipods. Animal Behaviour 72:861–868. DOI: https://doi.org/10.1016/j.anbehav.2006. 01.023
- **R Core Team. 2017.** R: A language and environment for statistical computing. *R Foundation for Statistical Computing*. Vienna,
- Richards TA, Bass D. 2005. Molecular screening of free-living microbial eukaryotes: diversity and distribution using a meta-analysis. Current Opinion in Microbiology 8:240–252. DOI: https://doi.org/10.1016/j.mib.2005.04. 010, PMID: 15939346
- **Ritchison G.** 1985. Plumage variability and social status in captive male house sparrows. *Kentucky Warbler* **61**: 39–42.
- Riters LV, Teague DP, Schroeder MB. 2004. Social status interacts with badge size and neuroendocrine physiology to influence sexual behavior in male house sparrows (Passer domesticus). Brain, Behavior and Evolution 63:141–150. DOI: https://doi.org/10.1159/000076240, PMID: 14726623
- Rohwer S. 1975. The social significance of avian winter plumage variability. Evolution 29:593. DOI: https://doi. org/10.2307/2407071
- **Rojas Mora A**, Meniri M, Glauser G, Vallat A, Helfenstein F. 2016. Badge size reflects sperm oxidative status within social groups in the house sparrow *Passer domesticus*. *Frontiers in Ecology and Evolution* **4**:67. DOI: https://doi.org/10.3389/fevo.2016.00067
- Rosenthal R. 1979. The file drawer problem and tolerance for null results. *Psychological Bulletin* **86**:638–641. DOI: https://doi.org/10.1037/0033-2909.86.3.638
- Rothstein H, Sutton A, Borenstein M. 2005. Publication Bias in Meta-Analysis: Prevention, Assessment and Adjustments. Chichester: Wiley. DOI: https://doi.org/10.1002/0470870168
- Sánchez-Tójar A, Nakagawa S, Sánchez-Fortún M, Martin DA, Ramani S, Girndt A, Bókony V, Kempenaers B, Liker A, Westneat DF, Burke T, Schroeder J. 2018a. Supporting information for "Meta-analysis challenges a textbook example of status signalling and demonstrates publication bias".
- Sánchez-Tójar A, Schroeder J, Farine DR. 2018b. A practical guide for inferring reliable dominance hierarchies and estimating their uncertainty. *Journal of Animal Ecology* 87:594–608. DOI: https://doi.org/10.1111/1365-2656.12776, PMID: 29083030
- Santos ESA, Scheck D, Nakagawa S. 2011. Dominance and plumage traits: meta-analysis and metaregression analysis. Animal Behaviour 82:3–19. DOI: https://doi.org/10.1016/j.anbehav.2011.03.022
- Schielzeth H. 2010. Simple means to improve the interpretability of regression coefficients. Methods in Ecology and Evolution 1:103–113. DOI: https://doi.org/10.1111/j.2041-210X.2010.00012.x

- Schmid CH, Landa M, Jafar TH, Giatras I, Karim T, Reddy M, Stark PC, Levey AS, Angiotensin-Converting Enzyme Inhibition in Progressive Renal Disease (AIPRD) Study Group. 2003. Constructing a database of individual clinical trials for longitudinal analysis. Controlled Clinical Trials 24:324–340. DOI: https://doi.org/10.1016/ S0197-2456(02)00319-7, PMID: 12757997
- Searcy WA, Nowicki S. 2005. The evolution of animal communication. Reliability and deception in signaling systems. New Jersey: Princeton University Press.
- Seguin A, Forstmeier W. 2012. No band color effects on male courtship rate or body mass in the zebra finch: four experiments and a meta-analysis. PLOS ONE 7:e37785. DOI: https://doi.org/10.1371/journal.pone. 0037785, PMID: 22685545
- **Senar JC**. 2006. Color displays as intrasexual signals of aggression and dominance. In: Hill G. E, McGraw K. J (Eds). *Bird Coloration: Function and Evolution*. London: Harvard University Press. p. 87–136.
- Senior AM, Grueber CE, Kamiya T, Lagisz M, O'Dwyer K, Santos ES, Nakagawa S. 2016. Heterogeneity in ecological and evolutionary meta-analyses: its magnitude and implications. *Ecology* 97:3293–3299. DOI: https://doi.org/10.1002/ecy.1591, PMID: 27912008
- Simmonds MC, Higginsa JPT, Stewartb LA, Tierneyb JF, Clarke MJ, Thompson SG. 2005. Meta-analysis of individual patient data from randomized trials: a review of methods used in practice. *Clinical Trials: Journal of* the Society for Clinical Trials 2:209–217. DOI: https://doi.org/10.1191/1740774505cn087oa
- Sneddon LU, Huntingford FA, Taylor AC. 1998. Impact of an ecological factor on the costs of resource acquisition: fighting and metabolic physiology of crabs. *Functional Ecology* 12:808–815. DOI: https://doi.org/ 10.1046/j.1365-2435.1998.00249.x
- Solberg EJ, Ringsby TH. 1997. Does male badge size signal status in small island populations of house sparrows, Passer domesticus? Ethology 103:177–186. DOI: https://doi.org/10.1111/j.1439-0310.1997.tb00114.x
- Sterne J, Egger M. 2005. Regression methods to detect publication and other bias in meta-analysis. In: Rothstein H, Sutton A, Borenstein M (Eds). *Publication Bias in Meta-Analysis2*. Chichester: John Wiley. p. 99–110. DOI: https://doi.org/10.1002/0470870168.ch6
- Tibbetts EA, Dale J. 2004. A socially enforced signal of quality in a paper wasp. Nature **432**:218–222. DOI: https://doi.org/10.1038/nature02949, PMID: 15538369
- Tibbetts EA, Safran RJ. 2009. Co-evolution of plumage characteristics and winter sociality in new and old world sparrows. *Journal of Evolutionary Biology* 22:2376–2386. DOI: https://doi.org/10.1111/j.1420-9101.2009. 01861.x, PMID: 19874438
- Tóth Z, Bókony V, Lendvai Ádám Z., Szabó K, Pénzes Z, Liker A. 2009. Kinship and aggression: do house sparrows spare their relatives? *Behavioral Ecology and Sociobiology* **63**:1189–1196. DOI: https://doi.org/10. 1007/s00265-009-0768-8
- Trikalinos T, Ioannidis JPA. 2005. Assessing the evolution of effect sizes over time. In: Rothstein H, Sutton A, Borenstein M (Eds). *Publication Bias in Meta-Analysis*. Chichester: John Wiley. p. 241–259. DOI: https://doi.org/ 10.1002/0470870168.ch13
- Veiga JP. 1993. Badge size, phenotypic quality, and reproductive success in the house sparrow: a study on honest advertisement. Evolution 47:1161–1170. DOI: https://doi.org/10.1111/j.1558-5646.1993.tb02143.x
- Veiga JP. 1996. Mate replacement is costly to males in the multibrooded house sparrow: an experimental study. *The Auk* **113**:664–671. DOI: https://doi.org/10.2307/4088987
- Wang D, Forstmeier W, Ihle M, Khadraoui M, Jerónimo S, Martin K, Kempenaers B. 2018. Irreproducible textbook "knowledge": The effects of color bands on zebra finch fitness. Evolution 72:961–976. DOI: https://doi. org/10.1111/evo.13459
- Whiting MJ, Nagy KA, Bateman PW. 2003. Evolution and maintenance of social status-signalling badges. In: Fox S. F, McCoy K, Baird T. A (Eds). *Lizard Social Behavior*. Baltimore: Johns Hopkins University Press. p. 47–82.

### **Appendix 1**

DOI: https://doi.org/10.7554/eLife.37385.014

### Information about data used in the study

Appendix 1—table 1. Summary of key differences in methodology among all studies (published and unpublished) testing the relationship between dominance rank and bib size in male house sparrows (N = 19 studies).

Variable	Levels	Number of studies	Order of preference*
Group composition	Males and females	11	-
-	Males only	8	-
Resource competed for	Food only	12	-
-	Food, water and roosting place	6	-
-	Females	1	-
Type of interactions	Aggressive only	12	-
-	Aggressive and non-aggres- sive	7	-
Interactions recording proto-	Live observations	11	-
- col	Video	6	-
-	Live and video observations	2	-
Type of bib size measured	Visible	14	1
-	Hidden	2	2
-	Both	3	-
Beak angle during measure-	90 <sub>deg</sub>	8	1
ment -	180°	3	2
-	Both	1	-
-	Unknown	7	-
Season	Non-breeding	13	-
-	Breeding	5	-
-	Both	1	-
Study location	Captive	16	-
-	Wild	2	-
-	Both	1	-

\*Order of preference used for the analyses (see main text). The order of preference was determined based on how frequently the method was used in previous studies.

DOI: https://doi.org/10.7554/eLife.37385.015

Appendix 1—table 2. List of the different methods used to estimate bib size in all studies (published and unpublished) testing the relationship between dominance rank and bib size in male house sparrows (N = 19 studies). Note that some studies used more than one method to estimate bib size.

Method to estimate bib size	Number of times used	Order of preference <sup>‡</sup>		
Area*	8	1		
Møller, 1987's equation	6	2		
Length and width†	3	2		
Appendix 1 table 2 continued on part page				

Appendix 1—table 2 continued on next page

#### Appendix 1—table 2 continued

Method to estimate bib size	Number of times used	Order of preference <sup>‡</sup>
Length only	2	3
Møller, 1987's drawings	1	4
Veiga, 1993's equation	1	5

\*Area was measured from pictures (N = 5 studies), by tracing and weighing (N = 2 studies), and by tracing and ranking (N = 1 study).

†If length and width were available, we estimated bib area using **Møller**, **1987**'s equation. ‡Order of preference used for the analyses (see main text). The order of preference was determined based on how frequently the method was used in previous studies.

DOI: https://doi.org/10.7554/eLife.37385.016

**Appendix 1—table 3.** List of the different methods used to infer dominance rank from dyadic interactions in published studies that tested the relationship between dominance rank and bib size in male house sparrows (N = 13 published studies, 11 different methods). Note that some studies used more than one method to estimate dominance rank and that unpublished studies are not included in this summary.

Method to infer dominance rank	Number of times used	Order of preference*
Proportion of contests won	4	4
Proportion of initiated contests	3	5
Kendall's linearity index	2	3
Proportion of contests won per dyad	2	6
Proportion of initiated contests won	2	6
David's score	1	1
I and SI	1	2
Landau's linearity index	1	3
Proportion of the received attacks won	1	7
Proportion of birds dominated	1	7
Proportion of contests won per dyad + linear assumption	1	7

\*Order of preference used for the analyses (see main text). The order of preference was determined based on both how frequently the method was used in previous studies and by taking into account the (expected) performance of each of the methods. First, higher order of preference was assigned to methods specifically designed for inferring linear dominance hierarchies (i.e. David's score, I and SI, Landau's and Kendall's linearity indices). We used the information available in **Sánchez-Tójar et al., 2018b** to rank David's score and I and SI as first and second methods in preference, respectively. Second, we ranked the remaining (proportion-based) methods based on how frequently they were used in previous studies. Importantly, the order of preference was chosen prior to conducting any statistical analysis, and thus, method selection was blind to the outcome of the analyses.

DOI: https://doi.org/10.7554/eLife.37385.017

**Appendix 1—table 4.** Additional comments on some of the published studies included in the meta-analysis.

Reference	Comments
Ritchison, 1985	According to the original publication, the total number of birds studied was 35, as opposed to the 25 individuals used in the meta-analyses of <b>Nakagawa et al., 2007</b> and <b>Santos et al., 2011</b> .
Appendix 1—tal	ble 4 continued on next page

Annen	div 1	—tab	lo 4 .	continue	4

Reference	Comments
Hein et al., 2003	The total number of birds included in our re-analysis of the primary data is smaller than that presented in the original publication. This is because our re-analysis only included fully identified individuals (e.g. birds missing rings could not be included).
Dolnik and Hoi, 2010	32 males were selected for the experiment, but one bird was excluded before the start of the experiment. Thus, $n$ was set to 31 individuals for this study.
Buchanan et al., 2010	96 birds were separated in 24 aviaries of four individuals each. The final <i>n</i> of several aviaries was less than four individuals, and therefore, these aviaries were not included in our meta-analyses (see main text, section 'Materials and Methods').
Rojas Mora et al., 2016	According to the primary data, one male did not interact, and thus, <i>n</i> was set to 59 individuals in Appendix 2.

DOI: https://doi.org/10.7554/eLife.37385.018

**Appendix 1—table 5.** Data descriptions for the unpublished data analysed in the metaanalysis.

#### Study

ID*	Data description
14	88 individuals were separated into four captive mixed-sex groups. Live observations after mild food deprivation were conducted to record agonistic dyadic interactions (i.e. fights) over (mostly) food for around one week in Feb 2003 (total = 1,563 fights). Bib length and width were measured for each male before the dominance observations using a ruler. More information can be found in <b>Lendvai et al., 2004</b> and <b>Bókony et al., 2012</b> .
15	61 individuals were separated into three captive mixed-sex groups. Live observations after mild food deprivation were conducted to record agonistic dyadic interactions (i.e. fights) over (mostly) food between Oct and Dec 2005 (two groups) and 2006 (one group; total = 2,003 fights). Bib area was measured for each male using standardized pictures taken after the dominance observations. More information can be found in <b>Tóth et al., 2009</b> and <b>Bókony et al., 2012</b> .
16	60 individuals were separated into four captive mixed-sex groups. Live and video observations after mild food deprivation were conducted to record agonistic dyadic interactions (i.e. fights) over (mostly) food for around two weeks per group between Oct 2007 and Feb 2008 (total = 6,641 fights). Bib length and width were measured for each male before the dominance observations using a ruler. More information can be found in <b>Bókony et al., 2010</b> and <b>Bókony et al., 2012</b> .
17	96 males were separated into four captive male-only groups. Videos after mild food deprivation were taken to record agonistic dyadic interactions (i.e. fights) over food for 10 days between Oct and Dec 2014 (total = 3,776 fights). Bib area was measured several times for each male (median = 3 times/male, range = 2 to 6) using standardized pictures taken from Oct to Dec 2014, and the mean bib area of each individual was used in the analyses.
18	453 individuals (215 females and 238 males) were observed in seven discrete sampling events in a wild population of house sparrows at Lundy Island, UK. Videos were taken to record agonistic dyadic interactions (i.e. fights) over food for 20 days between Nov 2013 and Dec 2016 (total = 11,063 fights). Bib length was measured several times for each male (median = 1 time/ male, range = 1 to 6) from Nov 2013 to Dec 2016 using a calliper, and the mean bib area of each individual in each sampling event was used in the analyses.
19	128 individuals were separated into 16 captive mixed-sex groups. Live observations after mild food deprivation were conducted to record agonistic dyadic interactions (i.e. supplants and hold-offs) over food between Mar and Apr 2005 (total = 5,496 fights). Bib length and width were measured for each male before the dominance observations using a calliper as in <b>Morrison et al., 2008</b> .
*Study	ID corresponding to <b>Table 1</b> in main text.
DOLL	

DOI: https://doi.org/10.7554/eLife.37385.019

### **Appendix 2**

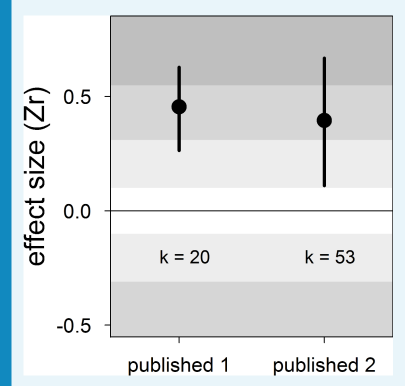
DOI: https://doi.org/10.7554/eLife.37385.020

### Meta-analyses based on published studies only

Appendix 2—table 1. Results of two multilevel meta-analyses to test the relationship between dominance rank and bib size in male house sparrows based on published studies only. Published 1 includes published effect sizes obtained from summary data, whereas published 2 includes published re-analysed effect sizes together with the remaining published effect sizes obtained from summary data. Additionally, the results of the Egger's regressions are shown. Estimates are presented as standardized effect sizes using Fisher's transformation (*Zr*). Credible intervals not overlapping zero are highlighted in bold.

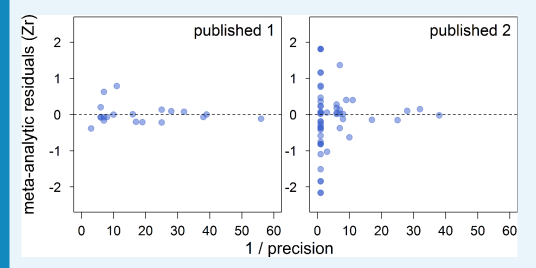
Meta- analysis	к	Meta-analytic mean [95% Crl]	l <sup>2</sup> population ID [ <b>95% Crl]</b> (%)	l <sup>2</sup> <sub>study ID</sub> [95% Crl] (%)	l <sup>2</sup> <sub>overall</sub> [ <b>95% Cri</b> ] (%)	Egger's regression [95% Crl]
Published 1	20	0.45 [0.26,0.63]	17 [0,51]	17 [0,53]	46 [15,78]	0.42 [-0.73,1.48]
Published 2	53	0.40 [0.11,0.67]	14 [0,46]	13 [0,42]	46 [17,72]	-0.25 [-0.73,0.26]

k = number of estimates; Crl = credible intervals;  $l^2$  = heterogeneity. DOI: https://doi.org/10.7554/eLife.37385.021



**Appendix 2—figure 1.** Forest plot showing the overall effect size of the relationship between dominance rank and bib size in male house sparrows based on published studies only. Published 1 includes published effect sizes obtained from summary data, whereas published 2 includes published re-analysed effect sizes together with the remaining published effect sizes obtained from summary data. We show posterior means and 95% credible intervals from multilevel meta-analyses. Estimates are presented as standardized effect sizes using

Fisher's transformation (*Zr*). Light, medium and dark grey show small, medium and large effect sizes, respectively (*Cohen, 1988*). *k* is the number of estimates. DOI: https://doi.org/10.7554/eLife.37385.022



**Appendix 2—figure 2.** Funnel plots of the meta-analytic residuals against their precision for the meta-analyses based on published studies only. Published 1 includes published effect sizes obtained from summary data, whereas published 2 includes published re-analysed effect sizes together with the remaining published effect sizes obtained from summary data. Estimates are presented as standardized effect sizes using Fisher's transformation (*Zr*). Precision = square root of the inverse of the variance. DOI: https://doi.org/10.7554/eLife.37385.023

### **Appendix 3**

DOI: https://doi.org/10.7554/eLife.37385.024

# Power analysis based on the estimated meta-analytic mean

#### R code used and explanations:

First, we need to clear up the memory and load the pwr library.

# clear memory

rm(list = ls())

# package needed

#### library(pwr)

Furthermore, we created a function to transform Zr values into r values. This is because our meta-analyses were based on Zr values, but the power analysis is based on r values.

# function to convert Zr to r

Zr.to.r<-function(Zr) {</pre>

r<-(exp(2\*Zr)-1)/(exp(2\*Zr)+1)

#### }

Power analysis

Next, we estimated the sample size necessary to find an effect size as small as the one estimated by our meta-analysis (Zr = 0.20). We used a significance level of 0.05, and the recommended 80% statistical power (**Cohen**, **1988**).

pwr.r.test(r = Zr.to.r(0.20), sig.level = 0.05, power = 0.8)

##

## approximate correlation power calculation (arctangh transformation)

##

## n = 198.3401

## r = 0.1973753

## sig.level=0.05

## power=0.8

#### ## alternative = two.sided

This shows that we would need the dominance rank and bib size of 198 individuals to find a significant r correlation of 0.20 with an 80% statistical power.

Additionally, we estimated the across-study statistical power of the tests on status signalling in house sparrows to compare it to the overall statistical power found in the behavioural ecology literature (*Jennions*, 2003).

pwr.r.test(n = 10, r = Zr.to.r(0.20), sig.level = 0.05)

##

## approximate correlation power calculation (arctangh transformation)

##

## n = 10

## r = 0.1973753

## sig.level=0.05

## power = 0.08474157

## alternative = two.sided

This shows that the statistical power of the sparrow literature on status signaling is as low as 8.5%, which is alarming.